

National Greenhouse Gas Inventory Report of JAPAN

April, 2012

**Ministry of the Environment, Japan
Greenhouse Gas Inventory Office of Japan (GIO), CGER, NIES**

Center for Global Environmental Research



National Institute for Environmental Studies, Japan



National Greenhouse Gas Inventory Report of JAPAN (2012)

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Foreword

On the basis of Article 4 and 12 of the United Nations Framework Convention on Climate Change (UNFCCC) and Article 7 of the Kyoto Protocol, all Parties to the Convention are required to submit national inventories of greenhouse gas emissions and removals to the Secretariat of the Convention. Therefore, the inventories on emissions and removals of greenhouse gases and precursors are reported in the Common Reporting Format (CRF) and in this National Inventory Report, in accordance with UNFCCC Inventory Reporting Guidelines (FCCC/SBTA/2006/9) and Decision 15/CMP.1.

This Report presents Japan's institutional arrangement for the inventory preparation, the estimation methods of greenhouse gas emissions and removals from sources and sinks, the trends in emissions and removals for greenhouse gases (carbon dioxide (CO₂); methane (CH₄); nitrous oxide (N₂O); hydrofluorocarbons (HFCs); perfluorocarbons (PFCs); and sulfur hexafluoride (SF₆)) and precursors (nitrogen oxides (NO_x), carbon monoxide (CO), non-methane volatile organic compounds (NMVOC), and sulfur dioxide (SO₂)). Supplementary information under Article 7.1 of the Kyoto Protocol is presented as well.

The structure of this report is prepared in line with the recommended structure indicated in the Annex I of UNFCCC Inventory Reporting Guidelines (FCCC/SBSTA/2006/9) and the Annotated outline of the National Inventory Report including reporting elements under the Kyoto Protocol, prepared by the UNFCCC secretariat.

The Executive Summary focuses on the latest trends in emissions and removals of greenhouse gases in Japan. Chapter 1 deals with background information on greenhouse gas inventories, the institutional arrangement for the inventory preparation, inventory preparation process, methodologies and data sources used, key source category analysis, QA/QC plan, and results of uncertainty assessment. Chapter 2 describes the latest information on trends in emissions and removals of greenhouse gases in Japan. Chapters 3 to 8 provide the detailed estimation methods for emissions and removals respectively, described in the *Revised 1996 IPCC Guidelines*. Chapter 9 comprises current status of reporting of the emissions from sources not covered by IPCC guidelines. Chapter 10 provides the explanations on improvement and recalculation (data revision, addition of new sources, etc.) made since the previous submission, and Chapters 11 through 15 provide supplementary information under Article 7.1 of the Kyoto Protocol.

Annex offers additional information to assist further understanding of Japan's inventory. The background data submitted to the secretariat provides the complete process of estimating Japan's inventory.

For the latest updates or changes in data, refer to the web-site (URL: www-gio.nies.go.jp) of the Greenhouse Gas Inventory Office of Japan (GIO).

April, 2012
Low-carbon Society Promotion Office
Global Environment Bureau
Ministry of the Environment

Preface

The Kyoto Protocol accepted by Japan in June 2002 targets the reduction of six greenhouse gases (GHGs): carbon dioxide (CO₂); methane (CH₄); nitrous oxide (N₂O); hydrofluorocarbons (HFCs); perfluorocarbons (PFCs); and sulfur hexafluoride (SF₆). Quantified targets for reductions in emissions of greenhouse gases have been set for each of the Annex I parties including Japan. The target given to Japan for the first commitment period (five years from 2008 to 2012) is to reduce average emissions of greenhouse gases by six percent from the base year (1990 for carbon dioxide, methane and nitrous oxide, and 1995 for HFCs, PFCs, and sulfur hexafluoride). At the same time, the Annex I parties were required to improve the accuracy of their emission estimates, and to prepare a national system for the estimation of anthropogenic emissions by sources and removals by sinks of the aforementioned greenhouse gases by one year prior to the start of the commitment period (2007). The GHGs inventories have been therefore authoritative data for Japan in reporting its achievement of the Kyoto Protocol's commitment.

The GHGs inventory of Japan including this report represents the combined knowledge of over 70 experts in a range of fields from universities, industrial bodies, regional governments, relevant government departments and agencies, and relevant research institutes, who are members of the Committee for the Greenhouse Gas Emissions Estimation Methods established by the Ministry of Environment in November 1999 and has been often held since then.

In compiling GHGs inventories, the Greenhouse Gas Inventory Office of Japan (GIO) would like to acknowledge not just the work of the Committee members in seeking to develop the methodology, but other experts who provided the latest scientific knowledge, the industrial bodies and government departments and agencies that provided the data necessary for compiling the inventories. We would like to express our gratitude to the Low-carbon Society Promotion Office of the Global Environment Bureau of the Ministry of the Environment, for their support to GIO.

This is the third time to submit the inventory in the first commitment period to the secretariat of the United Nations Convention on Climate Change (UNFCCC). Getting many feedbacks from internal and external reviewers, we have made further efforts to improve this report. We hope this report will be used widely and accurately as an index of what Japan should accomplish with regard to emission reductions, and as an index that shows the extent of measures implemented against global warming of Japan.

My appreciation also extends to Mr. Hiroshi ITOH who worked with us until July 2011, Mr. Kiyoto TANABE, a GIO researcher, Ms. Makiko YAMADA and Ms. Kyoumi ISHIGAMI, our assistants, who supported us with the smooth operation of GIO.

April, 2012



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Abbreviations

Executive Summary of National GHGs Inventory Report of Japan 2012

E.S.1. Background Information on GHGs Inventories, Climate Change and Supplementary Information Required under Article 7, Paragraph 1, of the Kyoto Protocol

This National Inventory Report comprises the inventory of the emissions and removals of greenhouse gases (GHGs), indirect GHGs and SO₂ in Japan for FY1990 through to FY2010¹, on the basis of Articles 4 and 12 of the United Nations Framework Convention on Climate Change (UNFCCC) and supplementary information for FY2008 through to FY2010 required under Article 7, Paragraph 1 of the Kyoto Protocol.

Japan's estimation methodologies of GHGs inventories are in line with the *Revised 1996 IPCC Guidelines for National Greenhouse Gas Inventories* (hereafter, *Revised 1996 IPCC Guidelines*) which was developed by the Intergovernmental Panel on Climate Change (IPCC). In addition, the *Good Practice Guidance and Uncertainty Management in National Greenhouse Gas Inventories (2000)* (hereafter, *GPG (2000)*) and the *Good Practice Guidance for Land Use, Land-Use Change and Forestry* (hereafter, *GPG-LULUCF*) are applied to improve transparency, consistency, comparability, completeness and accuracy of the inventory.

Annual inventory is reported in accordance with the *UNFCCC Reporting Guidelines on Annual Inventories* (FCCC/SBSTA/2006/9) adopted by the Conference of the Parties. Supplementary information under Article 7, Paragraph 1 of the Kyoto Protocol is reported in accordance with "Annotated outline of the National Inventory Report including reporting elements under the Kyoto Protocol" prepared by the UNFCCC secretariat.

¹ "FY" (Fiscal Year), from April of the reporting year through March of the next year, is used because CO₂ is the primary GHGs emissions and estimated on a fiscal year basis. "CY" stands for "Calendar Year".

E.S.2. Summary of National Emission and Removal Related Trends, and Emission and Removals from KP-LULUCF Activities

2.1. GHG Inventory

Total GHGs emissions in FY2010² (excluding LULUCF) were 1,258 million tonnes (in CO₂ eq.). They increased by 4.4% compared to the emissions in FY1990³ (excluding LULUCF). Compared to the emissions in the base year under the Kyoto Protocol⁴, they decreased by 0.3%.

It should be noted that actual emissions of HFCs, PFCs, and SF₆ in the period from CY1990 to 1994 are not estimated (NE)⁵.

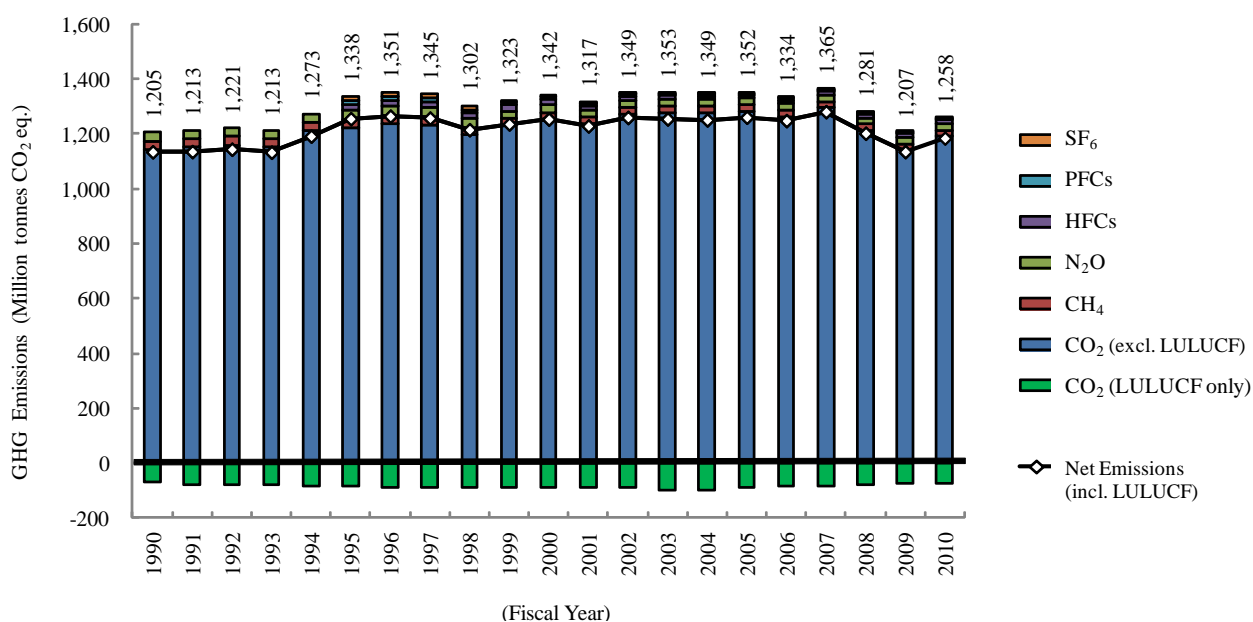


Figure 1 Trends in GHGs emission and removals in Japan

² The sum of CO₂, CH₄, N₂O, HFCs, PFCs and SF₆ emissions converted to CO₂ equivalents multiplied by their respective global warming potential (GWP). The GWP is a coefficient by means of which greenhouse gas effects of a given gas are made relative to those of an equivalent amount of CO₂. The coefficients are subjected to the *Second Assessment Report* (1995) issued by the Intergovernmental Panel on Climate Change (IPCC).

³ The sum of CO₂, CH₄ and N₂O emissions converted to CO₂ equivalents multiplied by their respective GWP.

⁴ Japan's base year under the Kyoto Protocol for CO₂, CH₄, N₂O emissions is FY 1990, while FY 1995 is the base year for HFCs, PFCs, and SF₆ emissions.

⁵ Potential emissions are reported in Common Reporting Format (CRF) for CY 1990 to 1994.

Table 1 Trends in GHGs emission and removals in Japan

[Million tonnes CO ₂ eq.]	GWP	Base year of KP	1990	1991	1992	1993	1994	1995	1996	1997	1998	1999	2000	2001
CO ₂ (excl. LULUCF)	1	1,144.1	1,141.2	1,150.1	1,158.6	1,150.9	1,210.7	1,223.7	1,236.6	1,231.5	1,195.9	1,230.9	1,251.6	1,236.4
CO ₂ (incl. LULUCF)	1	NA	1,071.0	1,072.7	1,081.4	1,070.9	1,129.0	1,141.6	1,149.9	1,144.5	1,109.1	1,143.8	1,163.8	1,148.5
CO ₂ (LULUCF only)	1	NA	-70.2	-77.4	-77.1	-80.0	-81.7	-82.1	-86.8	-87.1	-86.9	-87.1	-87.8	-87.9
CH ₄ (excl. LULUCF)	21	33.4	32.0	31.8	31.5	31.2	30.6	29.7	29.0	27.9	27.1	26.5	25.9	25.1
CH ₄ (incl. LULUCF)	21	NA	32.0	31.8	31.5	31.3	30.6	29.7	29.0	27.9	27.1	26.5	25.9	25.1
N ₂ O (excl. LULUCF)	310	32.6	31.6	31.1	31.3	31.0	32.2	32.7	33.6	34.3	32.8	26.4	29.0	25.5
N ₂ O (incl. LULUCF)	310	NA	31.7	31.2	31.4	31.1	32.3	32.7	33.7	34.4	32.8	26.4	29.0	25.6
HFCs	HFC-134a: 1,300 etc.	20.2	NE	NE	NE	NE	NE	NE	20.3	19.9	19.9	19.4	19.9	18.8
PFCs	PFC-14: 6,500 etc.	14.0	NE	NE	NE	NE	NE	NE	14.2	14.8	16.2	13.4	10.4	7.9
SF ₆	23,900	16.9	NE	NE	NE	NE	NE	NE	17.0	17.5	15.0	13.6	9.3	6.0
Gross Total (excl. LULUCF)		1,261.3	1,204.9	1,213.0	1,221.4	1,213.2	1,273.5	1,337.5	1,351.4	1,344.8	1,302.3	1,323.4	1,341.9	1,317.1
Net Total (incl. LULUCF)		NA	1,134.8	1,135.7	1,144.3	1,133.3	1,191.9	1,255.6	1,264.8	1,257.8	1,215.5	1,236.3	1,254.2	1,229.2

[Million tonnes CO ₂ eq.]	GWP	2002	2003	2004	2005	2006	2007	2008	2009	2010	Emission increase from the base year of KP	Emission increase from 1990 (2010)	Emission increase from 1995 (2010)	Emission increase from previous year (2010)
CO ₂ (excl. LULUCF)	1	1,273.5	1,278.6	1,278.0	1,282.3	1,263.1	1,296.3	1,213.2	1,142.3	1,191.9	4.2%	4.4%	-	4.4%
CO ₂ (incl. LULUCF)	1	1,184.4	1,180.4	1,180.3	1,191.5	1,178.1	1,212.2	1,134.5	1,070.4	1,118.8	-	4.5%	-	4.5%
CO ₂ (LULUCF only)	1	-89.1	-98.2	-97.7	-90.7	-85.0	-84.2	-78.7	-71.9	-73.2	-	4.3%	-	1.8%
CH ₄ (excl. LULUCF)	21	24.2	23.7	23.2	22.9	22.5	22.1	21.5	20.9	20.4	-38.8%	-36.2%	-	-2.1%
CH ₄ (incl. LULUCF)	21	24.2	23.7	23.2	22.9	22.5	22.1	21.5	20.9	20.4	-	-36.2%	-	-2.1%
N ₂ O (excl. LULUCF)	310	24.8	24.5	24.5	24.1	24.1	22.8	22.8	22.6	22.1	-32.4%	-30.3%	-	-2.2%
N ₂ O (incl. LULUCF)	310	24.8	24.5	24.6	24.1	24.1	22.8	22.8	22.6	22.1	-	-30.5%	-	-2.2%
HFCs	HFC-134a: 1,300 etc.	13.7	13.8	10.6	10.5	11.7	13.3	15.3	16.6	18.3	-9.7%	-	-9.9%	10.3%
PFCs	PFC-14: 6,500 etc.	7.4	7.2	7.5	7.0	7.3	6.4	4.6	3.3	3.4	-75.8%	-	-76.1%	4.2%
SF ₆	23,900	5.6	5.3	5.1	4.8	4.9	4.4	3.8	1.9	1.9	-89.0%	-	-89.0%	0.6%
Gross Total (excl. LULUCF)		1,349.1	1,353.0	1,348.9	1,351.5	1,333.6	1,365.3	1,281.3	1,207.4	1,258.0	-0.3%	4.4%	-5.9%	4.2%
Net Total (incl. LULUCF)		1,260.1	1,254.8	1,251.2	1,260.8	1,248.6	1,281.1	1,202.6	1,135.5	1,184.8	-	4.4%	-	4.3%

* NA: Not Applicable

* NE: Not Estimated

* LULUCF: Land Use, Land-Use Change and Forestry

2.2. KP-LULUCF Activities

Japan reports supplementary information on Afforestation/Reforestation (AR), Deforestation (D), Forest management (FM) and Revegetation (RV) as LULUCF activities under Article 3, Paragraphs 3 and 4 of the Kyoto Protocol. The breakdown of emissions and removals to each activity in the first commitment period of the Kyoto Protocol is shown in Table 2. For detailed information, see Chapter 11.

Table 3 Accounting summary for activities under Articles 3.3 and 3.4 of the Kyoto Protocol (CRF Information Table)

GREENHOUSE GAS SOURCE AND SINK ACTIVITIES	BY	Net emissions/removals				Accounting Parameters	Accounting Quantity
		2008	2009	2010	Total		
		(Gg CO ₂ equivalent)					
A. Article 3.3 activities							
A.1. Afforestation and Reforestation						-1230.68	
A.1.1. Units of land not harvested since the beginning of the commitment period		-389.54	-415.03	-426.11	-1,230.68	-1230.68	
A.1.2. Units of land harvested since the beginning of the commitment period							
A.2. Deforestation		2,456.72	3,115.09	4,822.89	10,394.70	10394.70	
B. Article 3.4 activities							
B.1. Forest Management (if elected)		-45,388.77	-49,005.55	-53,251.78	-147,646.10	-147646.10	
3.3 offset						9,164.02	
FM cap						238,333.33	
B.2. Cropland Management (if elected)	NA	NA	NA	NA	NA	NA	
B.3. Grazing Land Management (if elected)	NA	NA	NA	NA	NA	NA	
B.4. Revegetation (if elected)	-77.78	-1081.76	-1112.34	-1130.14	-3324.24	-233.34	

- ※ The net removals by FM after application of 3.3 offset are lower than the upper limit (13 Mt-C times 5 (238,333 Gg-CO₂)) given in the Appendix to decision 16/CMP.1.
- ※ Since the total anthropogenic GHG emissions by sources and removals by sinks in managed forests since 1990 are larger than the net source of emissions incurred under Article 3.3, the offset rule according to paragraph 10 of the Annex to decision 16/CMP.1 is applied to Japan.
- ※ Methodologies for estimation and accounting of Article 3.3 and 3.4 activities are continuously reviewed. The values in Table 11-2 are estimated by using the current methodologies, and are only reported but not accounted for in the 2012 submission since Japan elected accounting for the entire commitment period. The issuance of removal units from LULUCF activities under the Kyoto Protocol is to be performed at the end of the first commitment period.
- ※ The total values and results of summing up each element are not always the same because of the difference in display digit.

E.S.3. Overview of Source and Sink Category Emission Estimates and Trends

3.1. GHG Inventory

The breakdown of GHGs emissions and removals in FY2010 by sector⁶ shows that the Energy accounts for 91.1% of total GHGs emissions. It is followed by the Industrial Processes (5.2%), the Agriculture (2.0%), the Waste (1.7%) and the Solvents and Other Product Use (0.01%).

Removals by the LULUCF in FY2010 were equivalent to 5.8% of total GHGs emissions.

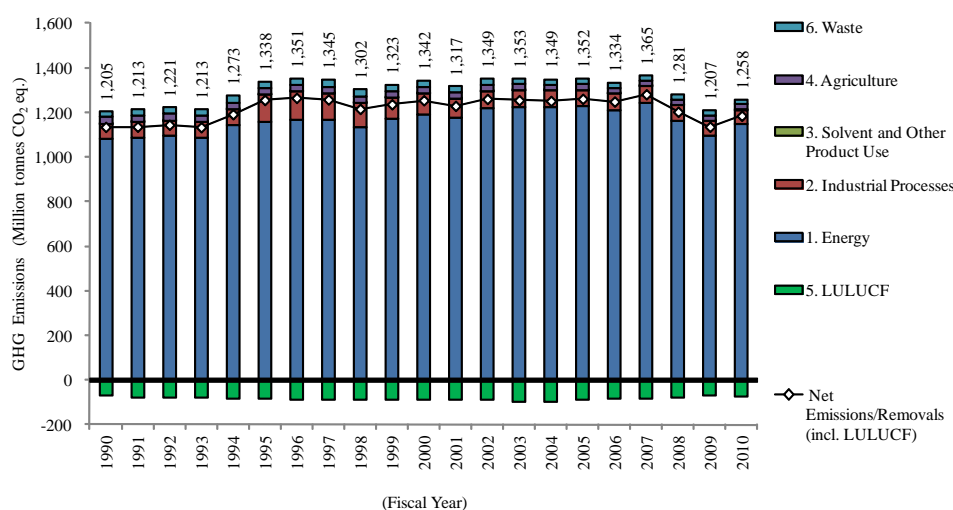


Figure 2 Trends in GHGs emissions and removals in each category

Table 4 Trends in GHGs emissions and removals in each category

[Million tonnes CO ₂ eq.]	1990	1991	1992	1993	1994	1995	1996	1997	1998	1999	2000
1. Energy	1,079.0	1,086.8	1,094.2	1,087.7	1,143.7	1,156.8	1,168.9	1,165.8	1,135.6	1,171.0	1,190.9
2. Industrial Processes	68.6	68.9	68.8	67.6	69.8	121.3	123.5	120.1	108.6	95.3	94.4
3. Solvent and Other Product Use	0.3	0.4	0.4	0.4	0.4	0.4	0.4	0.4	0.4	0.4	0.3
4. Agriculture	31.3	31.2	31.2	31.1	30.7	30.0	29.4	28.7	28.3	27.9	27.6
5. LULUCF	-70.1	-77.3	-77.1	-79.9	-81.6	-82.0	-86.7	-87.0	-86.8	-87.1	-87.7
6. Waste	25.8	25.7	26.8	26.4	28.9	29.0	29.3	29.7	29.3	28.9	28.7
Net Emissions/Removals (incl. LULUCF)	1,134.8	1,135.7	1,144.3	1,133.3	1,191.9	1,255.6	1,264.8	1,257.8	1,215.5	1,236.3	1,254.2
Emissions (excl. LULUCF)	1,204.9	1,213.0	1,221.4	1,213.2	1,273.5	1,337.5	1,351.4	1,344.8	1,302.3	1,323.4	1,341.9

[Million tonnes CO ₂ eq.]	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010
1. Energy	1,178.0	1,217.8	1,223.5	1,223.3	1,226.9	1,208.3	1,241.9	1,161.1	1,097.4	1,145.6
2. Industrial Processes	84.4	78.0	76.7	73.9	73.8	75.8	74.4	70.8	63.7	65.9
3. Solvent and Other Product Use	0.3	0.3	0.3	0.3	0.3	0.2	0.2	0.1	0.1	0.1
4. Agriculture	27.4	27.1	26.9	26.7	26.5	26.4	26.1	25.9	25.6	25.5
5. LULUCF	-87.9	-89.0	-98.2	-97.7	-90.7	-85.0	-84.1	-78.7	-71.9	-73.2
6. Waste	27.1	25.9	25.6	24.7	24.0	22.7	22.7	23.3	20.6	20.9
Net Emissions/Removals (incl. LULUCF)	1,229.2	1,260.1	1,254.8	1,251.2	1,260.8	1,248.6	1,281.1	1,202.6	1,135.5	1,184.8
Emissions (excl. LULUCF)	1,317.1	1,349.1	1,353.0	1,348.9	1,351.5	1,333.6	1,365.3	1,281.3	1,207.4	1,258.0

* LULUCF: Land Use, Land-Use Change and Forestry

3.2. KP-LULUCF Activities

See section 2.2 of executive summary.

E.S.4. Other Information (Indirect GHGs and SO₂)

Under the UNFCCC, it is required to report emissions not only 6 types of GHGs (CO₂, CH₄, N₂O, HFCs, PFCs and SF₆) that are controlled by the Kyoto Protocol, but also emissions of indirect GHGs (NO_x, CO and NMVOC) as well as SO₂. Their emission trends are indicated below.

Nitrogen oxide (NO_x) emissions in FY2010 were 1,744 thousand tonnes. They decreased by 14.6% since FY1990 and decreased by 1.9% compared to the previous year.

Carbon monoxide (CO) emissions in FY2010 were 2,577 thousand tonnes. They decreased by 42.6% since FY1990 and increased by 2.1% compared to the previous year.

Non-methane volatile organic compounds (NMVOC) emissions in FY2010 were 1,569 thousand tonnes. They decrease by 19.2% since FY1990 and increased by 0.4% compared to the previous year.

Sulfur dioxide (SO₂) emissions in FY2010 were 955 thousand tonnes. They decreased by 23.9% since FY1990 and decreased by 0.3% compared to the previous year.

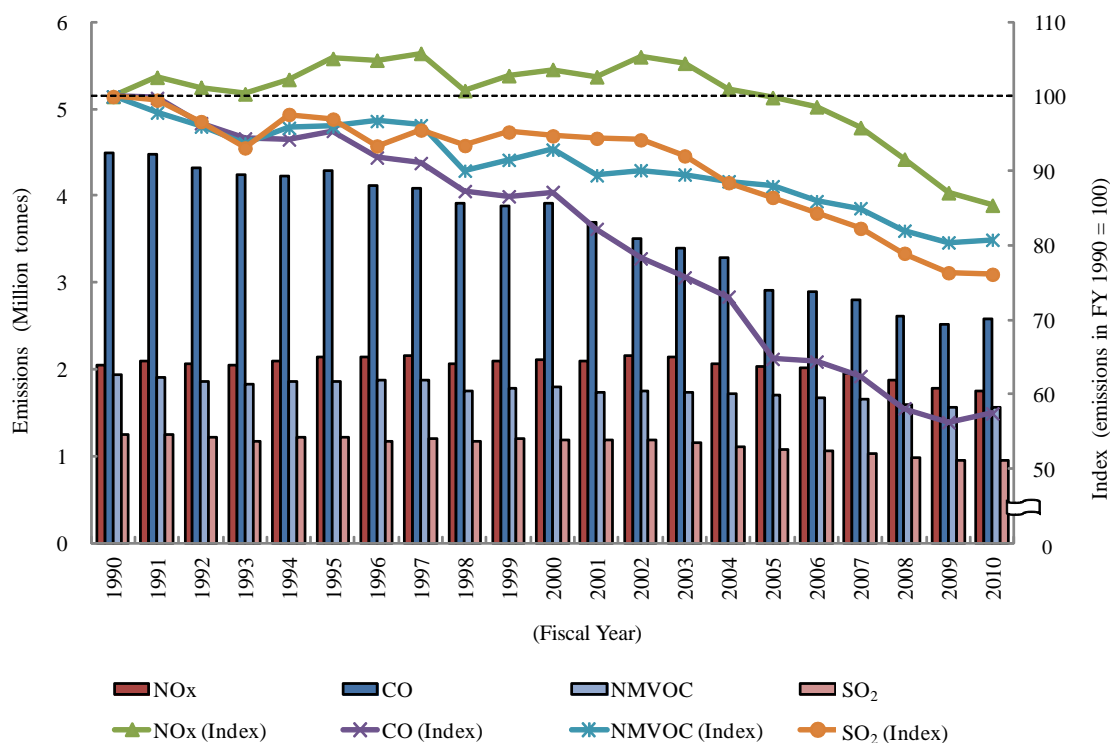


Figure 3 Trends in Emissions of Indirect GHGs and SO₂

⁶ It implies “Category” indicated in the Revised 1996 IPCC Guidelines and CRF.

Chapter 1. Introduction

1.1. Background Information on Japan's Greenhouse Gas Inventory

Japan reports the greenhouse gas (GHG) inventories, which contain the information on emissions and removals of GHGs, including indirect GHGs and SO₂ in Japan from FY1990 to FY2010¹, on the basis of Article 4 and 12 of the United Nations Framework Convention on Climate Change (UNFCCC) and of Article 7 of the Kyoto Protocol.

Estimation methodologies for the GHG inventories are required to be in line with the *Revised 1996 IPCC Guidelines for National Greenhouse Gas Inventories (Revised 1996 IPCC Guidelines)*, which was made by the Intergovernmental Panel on Climate Change (IPCC), and Japan's estimation methodologies are basically in line with these guidelines. In order to enhance transparency, consistency, comparability, completeness and accuracy of inventory, Japan also applies the *Good Practice Guidance and Uncertainty Management in National Greenhouse Gas Inventories* published in 2000 (GPG (2000)) and the *Good Practice Guidance for Land Use, Land-Use Change and Forestry* published in 2003 (GPG-LULUCF).

Japan's national inventory is reported in accordance with the *UNFCCC Reporting Guidelines on Annual Inventories (FCCC/SBSTA/2006/9)*. For the reporting of supplementary information required under Article 7, paragraph 1 of the Kyoto Protocol, the reporting guidelines (*Annotated outline of the National Inventory Report including reporting elements under the Kyoto Protocol*), the use of which is encouraged by the UNFCCC Secretariat, are applied.

1.2. A Description of Japan's Institutional Arrangement for the Inventory Preparation

The Ministry of the Environment (MOE), with the cooperation of relevant ministries, agencies and organizations, prepares Japan's national inventory and compiles supplementary information required under Article 7.1, and the MOE submits the inventory to the UNFCCC Secretariat in accordance with the UNFCCC and the Kyoto Protocol.

The MOE takes overall responsibilities for the national inventory and therefore makes every effort on improving the quality of inventory. The MOE organizes the "Committee for the Greenhouse Gas Emission Estimation Methods (Committee)" in order to integrate the latest scientific knowledge into the inventory and to modify it based on more recent international provisions. The estimation of GHG emissions and removals, the key category analysis and the uncertainty assessment are then carried out by taking the decisions of the Committee into consideration. Substantial activities, such as the estimation of emissions and removals and the preparation of Common Reporting Format (CRF) and National Inventory Report (NIR), are done by the Greenhouse Gas Inventory Office of Japan (GIO), which belongs to the Center for Global Environmental Research of the National Institute for Environmental Studies. The relevant ministries, agencies and organizations provide the GIO the appropriate data (e.g., activity data, emission factors, GHG emissions and removals) through compiling various statistics and also provide relevant information on supplementary information required under Article 7.1. The relevant ministries check and verify the inventories (i.e., CRF, NIR) including the spreadsheets that are actually utilized for the estimation, as a part of the Quality Control (QC) activities.

¹ "FY (fiscal year)" is used because the major part of CO₂ emission estimate is on the fiscal year basis (April to March).

The checked and verified inventories are Japan’s official values. The inventories are then made public by the MOE and are submitted to the UNFCCC Secretariat by the Ministry of Foreign Affairs.

Figure 1-1 shows the overall institutional arrangement for the inventory preparation within Japan. More detailed information on the role and responsibility of each relevant ministry, agency and organization in the inventory preparation process is described in Annex 6.

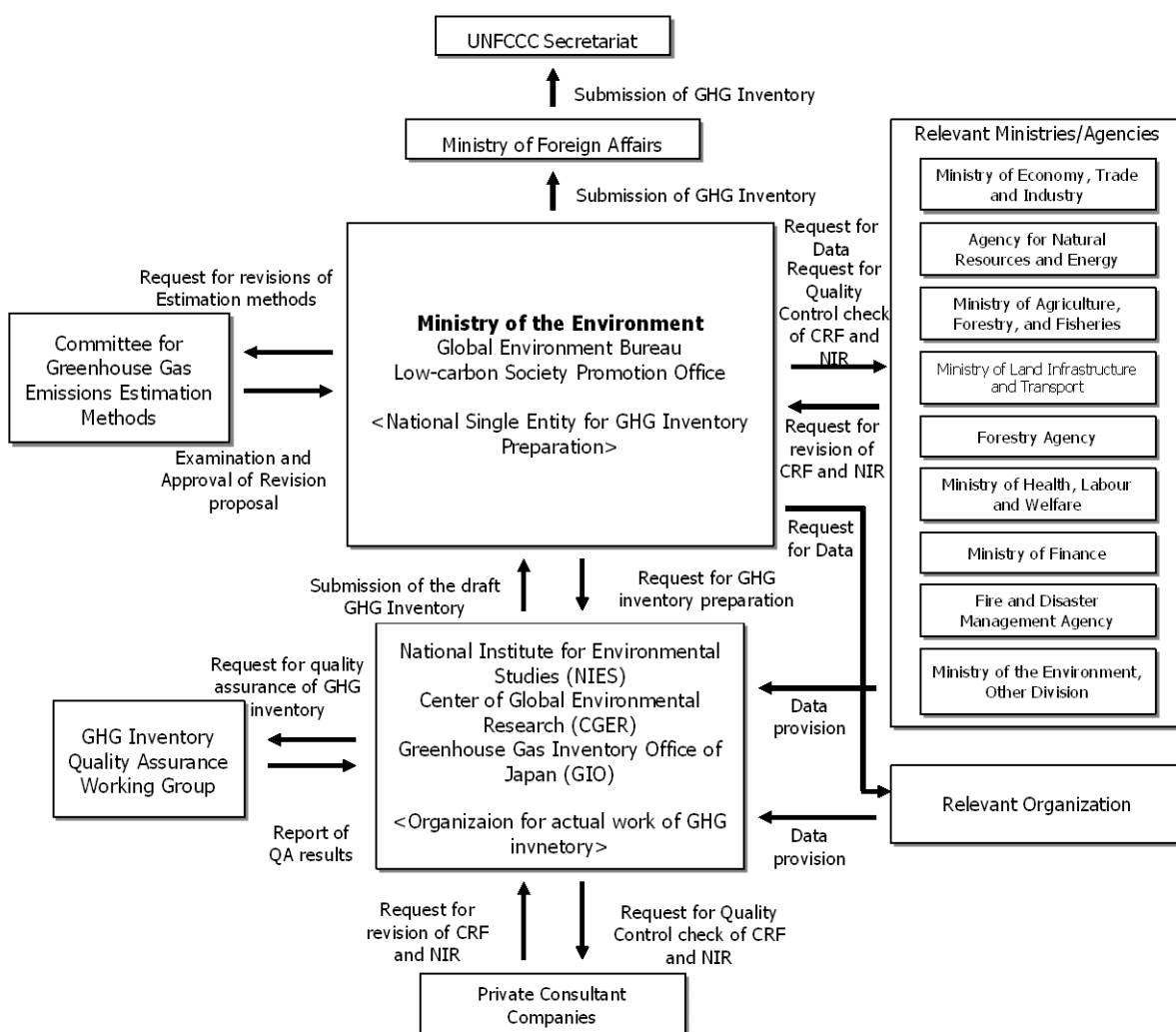


Figure 1-1 Japan’s institutional arrangement for the national inventory preparation

1.3. Brief Description of the Inventory Preparation Process

1.3.1. Annual cycle of the inventory preparation

Table 1-1 shows the annual cycle of the inventory preparation. In Japan, in advance of the estimation of national inventory submitted to the UNFCCC (submission deadline: 15th April), preliminary figures are estimated and published as a document for an official announcement. (In preliminary figures, only GHG emissions excluding removals are estimated.)

Table 1-1 Annual cycle of the inventory preparation

		*Inventory preparation in fiscal year "n"												
		Calendar Year n+1						CY n+2						
Process	Relevant Entities	Fiscal Year n+1												
		May	Jun	Jul	Aug	Sep	Oct	Nov	Dec	Jan	Feb	Mar	Apr	
1	Discussion on the inventory improvement	MOE, GIO	→	→	→	→	→	→	→	→	→	→	→	→
2	Holding the meeting of the Committee	MOE, (GIO, Private consultant)	→	→	→	→	→	→	→	→	→	→	→	→
3	Collection of data for the national inventory	MOE, GIO, Relevant Ministries/Agencies, Relevant organization, Private consultant									→	→	→	→
4	Preparation of a draft of CRF	GIO, Private consultant									→	→	→	→
5	Preparation of a draft of NIR	GIO, Private consultant									→	→	→	→
6	Implementation of the exterior QC and the coordination with the relevant ministries and agencies	MOE, GIO, Relevant Ministries/Agencies, Private consultant										→	→	→
7	Correction of the drafts of CRF and NIR	MOE, GIO, Private consultant											→	→
8	Submission and official announcement of the national inventory	MOE, Ministry of Foreign Affairs, GIO												★
9	Holding the meeting of the QA-WG	MOE, GIO	→	→	→	→								

★: Inventory submission and official announcement must be implemented within 6 weeks after April 15.

MOE: Ministry of the Environment

GIO: Greenhouse Gas Inventory Office of Japan

Committee: Committee for the Greenhouse Gas Emission Estimation Methods

QAWG: Inventory Quality Assurance Working Group

1.3.2. Process of the inventory preparation

1) Discussion on the inventory improvement (Step 1)

The MOE and the GIO identify the items, which need to be addressed by the Committee, based on the results of the previous inventory review of the UNFCCC, the recommendations of the “Inventory Quality Assurance Working Group (QAWG)”, the items needing improvement as identified at former Committee’s meetings, as well as any other items, requiring revision, as determined during previous inventory preparations. The schedule for the expert evaluation (step 2) is developed by taking the above mentioned information into account.

2) Holding the meeting of the Committee for the Greenhouse Gas Emission Estimation Methods [evaluation and examination of estimation methods by experts] (Step 2)

The MOE holds the meeting of the Committee, in which estimation methodologies for an annual inventory and the issues that require technical reviews are discussed by experts with different scientific backgrounds (refer to Annex 6).

3) Collection of data for the national inventory (Step 3)

The data required for preparing the national inventory and the supplementary information required under Article 7.1 of the Kyoto Protocol are collected.

4) Preparation of a draft of CRF [including the implementation of the key category analysis and the uncertainty assessment] (Step 4)

The data input and estimation of emissions and removals are carried out simultaneously by utilizing files containing spreadsheets (JNGI: Japan National GHG Inventory files), which have inter-connecting links among themselves based on the calculation formulas for emissions and removals. Subsequently, the key category analysis and the uncertainty assessment are also carried out.

5) Preparation of a draft of NIR (Step 5)

The draft of NIR is prepared by following the general guidelines made by the MOE and the GIO. The MOE and the GIO identify the points, which need to be revised or require an additional description by

taking the discussion at step 1 into account. The GIO and the selected private consulting companies prepare new NIR by updating data, and by adding and revising descriptions in the previous NIR.

6) Implementation of the exterior QC and the coordination with the relevant ministries and agencies (Step 6)

As a QC activity, the selected private consulting companies check the JNGI files and the initial draft of CRF (the 0th draft) prepared by the GIO (exterior QC). The companies not only check the input data and the calculation formulas in the files, but also verify the estimations by re-calculating the total amounts of GHG emissions determined by utilizing the same files. Because of this cross-check, any possible data input and emission estimation mistakes are avoided. They also check the content and descriptions of the initial draft of NIR (the 0th draft) prepared by the GIO. JNGI files draft CRF and draft NIR, which have been checked by the private consulting companies, are regarded as the primary drafts of inventories.

Subsequently, the GIO sends out the primary drafts of the inventories as well as of official announcements as electronic computer files to the MOE and the relevant ministries and agencies, and possible revisions are carried out. The data, which are estimated based on confidential data, are only sent out for confirmation to the ministry and/or the agency which provided these confidential data.

7) Correction of the drafts of CRF and NIR (Step 7)

When revisions are requested at step 6, the possible corrections are discussed among the MOE, the GIO and the relevant ministries and/or agencies. The corrected drafts are then the secondary drafts. The secondary drafts are sent out again to the relevant ministries and/or the agencies for conclusive confirmation. If there is no additional request for revision, they are considered to be the final versions.

8) Submission and official announcement of the national inventory (Step 8)

The completed inventory is submitted by the MOE via the Ministry of Foreign Affairs to the UNFCCC Secretariat. Information on the estimated GHG emissions and removals are officially announced and published on the MOE's homepage (<http://www.env.go.jp/>) with additional relevant information. The inventory is also published on the GIO's homepage (<http://www-gio.nies.go.jp/index-j.html>).

9) Holding the meeting of the Greenhouse Gas Inventory Quality Assurance Working Group (Step 9)

The QAWG, which is composed of experts who are not directly involved in or related to the inventory preparation process, is organized in order to guarantee the inventory's quality and to find out possible improvements.

This QAWG reviews the appropriateness of the estimation methodologies, activity data, emission factors, and the contents of CRF and NIR. GIO integrates the items, which were suggested for improvement by the QAWG, into the inventory improvement program, and utilizes them in discussions on the inventory estimation methods and in subsequent inventory preparation.

1.4. Brief General Description of Methodologies and Data Sources Used

The methodology used in estimation of GHG emissions or removals is basically in accordance with the *Revised 1996 IPCC Guidelines*, the *GPG (2000)* and the *GPG-LULUCF*. The country-specific

methodologies are also used for some categories (e.g., “4.C. methane emissions from rice cultivation”) in order to reflect the actual emission status in Japan.

Results of the actual measurements or estimates based on research conducted in Japan are used to determine the emissions factors (country-specific emissions factors). The default values given in the *Revised 1996 IPCC Guidelines*, the *GPG (2000)* and the *GPG-LULUCF* are used for: emissions, which are assumed to be quite low (e.g., “1.B.2.a.ii fugitive emissions from fuel (oil and natural gas”)), and where the possibility of emission from a given source is uncertain (e.g., “4.D.3. Indirect emissions from soil in agricultural land”).

1.5. Brief Description of Key Categories

Key category analysis is carried out in accordance with the *GPG (2000)* and the *GPG-LULUCF* (Tier 1, Tier 2 level assessment and trend assessment, qualitative analysis).

1.5.1. GHG Inventory

In FY2010, 39 sources and sinks were identified as Japan’s key categories (Table 1-2). For the base year of the UNFCCC (FY1990), 35 sources and sinks were identified as key categories (Table 1-3). More detailed information is described in Annex 1.

Table 1-2 Japan's key categories in FY2010

	A IPCC Category		B Direct GHGs	L1	T1	L2	T2
#1	1A Stationary Combustion	Solid Fuels	CO2	#1	#2	#4	#7
#2	1A Stationary Combustion	Liquid Fuels	CO2	#2	#1	#10	#8
#3	1A Stationary Combustion	Gaseous Fuels	CO2	#3	#3		
#4	1A3 Mobile Combustion	b. Road Transportation	CO2	#4	#6	#6	#24
#5	5A Forest Land	1. Forest Land remaining Forest Land	CO2	#5		#2	
#6	2A Mineral Product	1. Cement Production	CO2	#6	#7	#11	#10
#7	2F(a) Consumption of Halocarbons and SF6	1. Refrigeration and Air Conditioning Equipment	HFCs	#7	#5	#3	#1
#8	1A Stationary Combustion	Other Fuels	CO2	#8	#11	#7	#11
#9	6C Waste Incineration		CO2	#9		#5	
#10	1A3 Mobile Combustion	d. Navigation	CO2	#10	#14		
#11	1A3 Mobile Combustion	a. Civil Aviation	CO2	#11			
#12	2A Mineral Product	3. Limestone and Dolomite Use	CO2	#12		#20	
#13	4A Enteric Fermentation		CH4			#25	
#14	2A Mineral Product	2. Lime Production	CO2			#23	
#15	4B Manure Management		N2O			#8	
#16	4C Rice Cultivation		CH4			#17	
#17	1A Stationary Combustion		N2O			#16	#19
#18	5E Settlements	2. Land converted to Settlements	CO2			#22	#22
#19	6A Solid Waste Disposal on Land		CH4		#13	#19	#9
#20	4D Agricultural Soils	1. Direct Soil Emissions	N2O			#9	#13
#21	4D Agricultural Soils	3. Indirect Emissions	N2O			#13	#16
#22	1A3 Mobile Combustion	b. Road Transportation	N2O			#14	#12
#23	4B Manure Management		CH4			#15	#18
#24	2F(a) Consumption of Halocarbons and SF6	7. Semiconductor Manufacture	PFCs			#18	#15
#25	6C Waste Incineration		N2O			#12	
#26	2F(a) Consumption of Halocarbons and SF6	5. Solvents	PFCs		#9		#4
#27	6B Wastewater Handling		CH4				#26
#28	6B Wastewater Handling		N2O			#21	
#29	2F(a) Consumption of Halocarbons and SF6	8. Electrical Equipment	SF6		#8		#2
#30	2B Chemical Industry	3. Adipic Acid	N2O		#10		#17
#31	5B Cropland	2. Land converted to Cropland	CO2				#20
#32	5A Forest Land	2. Land converted to Forest Land	CO2				#25
#33	2E Production of Halocarbons and SF6	2. Fugitive Emissions	PFCs				#21
#34	2E Production of Halocarbons and SF6	2. Fugitive Emissions	SF6		#12		#3
#35	1A3 Mobile Combustion	a. Civil Aviation	N2O			#1	#6
#36	1A3 Mobile Combustion	d. Navigation	N2O			#24	
#37	2E Production of Halocarbons and SF6	2. Fugitive Emissions	HFCs				#23
#38	2E Production of Halocarbons and SF6	1. By-product Emissions (Production of HCFC-22)	HFCs		#4		#14
#39	1B Fugitive Emission	1a i. Coal Mining and Handling (under gr.)	CH4				#5

N.B. Figures recorded in the Level and Trend columns indicate the ranking of individual level and trend assessments.

Table 1-3 Japan's key categories in FY1990

	A IPCC Category		B Direct GHGs	L1	L2
#1	1A Stationary Combustion	Liquid Fuels	CO2	#1	#7
#2	1A Stationary Combustion	Solid Fuels	CO2	#2	#6
#3	1A3 Mobile Combustion	b. Road Transportation	CO2	#3	#8
#4	1A Stationary Combustion	Gaseous Fuels	CO2	#4	
#5	5A Forest Land	1. Forest Land remaining Forest Land	CO2	#5	#1
#6	2A Mineral Product	1. Cement Production	CO2	#6	#10
#7	2E Production of Halocarbons and SF6	1. By-product Emissions (Production of HCFC-22)	HFCs	#7	#27
#8	1A3 Mobile Combustion	d. Navigation	CO2	#8	
#9	6C Waste Incineration		CO2	#9	#3
#10	2F(a) Consumption of Halocarbons and SF6	8. Electrical Equipment	SF6	#10	#4
#11	2A Mineral Product	3. Limestone and Dolomite Use	CO2	#11	#23
#12	2F(a) Consumption of Halocarbons and SF6	5. Solvents	PFCs	#12	#9
#13	1A Stationary Combustion	Other Fuels	CO2	#13	#17
#14	4A Enteric Fermentation		CH4	#14	#28
#15	6A Solid Waste Disposal on Land		CH4	#15	#15
#16	2B Chemical Industry	3. Adipic Acid	N2O	#16	
#17	1A3 Mobile Combustion	a. Civil Aviation	CO2	#17	
#18	4C Rice Cultivation		CH4	#18	#22
#19	2A Mineral Product	2. Lime Production	CO2		#26
#20	4B Manure Management		N2O		#14
#21	5E Settlements	2. Land converted to Settlements	CO2		#21
#22	2E Production of Halocarbons and SF6	2. Fugitive Emissions	SF6		#5
#23	4D Agricultural Soils	1. Direct Soil Emissions	N2O		#11
#24	1A3 Mobile Combustion	b. Road Transportation	N2O		#13
#25	4D Agricultural Soils	3. Indirect Emissions	N2O		#16
#26	2B Chemical Industry	1. Ammonia Production	CO2		#30
#27	2F(a) Consumption of Halocarbons and SF6	7. Semiconductor Manufacture	PFCs		#19
#28	4B Manure Management		CH4		#18
#29	1B Fugitive Emission	1a i. Coal Mining and Handling (under gr.)	CH4		#12
#30	6B Wastewater Handling		CH4		#29
#31	6C Waste Incineration		N2O		#20
#32	6B Wastewater Handling		N2O		#24
#33	5E Settlements	1. Settlements remaining Settlements	CO2		#32
#34	1A3 Mobile Combustion	d. Navigation	N2O		#25
#35	1A3 Mobile Combustion	a. Civil Aviation	N2O		#2

N.B. Figures recorded in the column L (Level) indicate the ranking of level assessments.

The data of HFCs, PFCs and SF₆ utilized for this analysis are the 1995 values.

1.5.2. KP-LULUCF Activity

As a result of analysis implemented in accordance with GPG-LULUCF, Chapter 5, "Afforestation/Reforestation", "Deforestation", "Forest management" and "Revegetation" activities (CO₂) were identified as key categories for Japan's KP-LULUCF activities in FY2010. More detailed information is described in section 11.7 of chapter 11.

1.6. Information on the QA/QC Plan including Verification and Treatment of Confidentiality Issues

The role and the responsibility for each entity in the inventory preparation process are clarified in Japan's national system. The relevant entities are: MOE, GIO, relevant ministries, relevant agencies, relevant organizations, the Committee, selected private consulting companies and the QAWG. The QC activities (e.g., checking estimation accuracy, archiving documents) are carried out in each step of the inventory preparation process in accordance with the GPG (2000) and the GPG-LULUCF in order to

control the inventory's quality.

As a QA activity, the QAWG is established in order to implement a detailed review of each source or sink. The QAWG is composed of experts who are not directly involved in or related to the inventory preparation process. The QAWG reviews several sectors/categories annually with the aim of reviewing the entire inventory within a few years. The QAWG review was implemented for the Energy sector in FY2010.

For further information on the national system and process for inventory preparation, see sections 1.2 and 1.3 of this chapter. Detailed information on the QA/QC plan is described in Annex 6.1.

1.7. General Uncertainty Assessment, including Data on the Overall Uncertainty for the Inventory Totals

1.7.1. GHG Inventory

Total net GHG emissions in Japan for FY2010 were approximately 1,185 million tonnes (CO₂ equivalents). The total net emissions uncertainty was 2% and the uncertainty introduced into the trend in the total emissions was 1%. More detailed information on the uncertainty assessment is described in Annex 7.

Table 1-4 Uncertainty of Japan's GHG inventory

IPCC Category	GHGs	Emissions / Removals [Gg-CO ₂ eq.]		Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions ¹⁾	rank
		A	[%]				
1A. Fuel Combustion (CO ₂)	CO ₂	1,137,550.9	90.4%	1%	10	0.74%	2
1A. Fuel Combustion (Stationary:CH ₄ ,N ₂ O)	CH ₄ , N ₂ O	4,957.6	0.4%	27%	4	0.11%	8
1A. Fuel Combustion (Transport:CH ₄ ,N ₂ O)	CH ₄ , N ₂ O	2,694.0	0.2%	351%	1	0.80%	1
1B. Fugitive Emissions from Fuels	CO ₂ , CH ₄ , N ₂ O	409.0	0.0%	19%	5	0.01%	9
2. Industrial Processes (CO ₂ ,CH ₄ ,N ₂ O)	CO ₂ , CH ₄ , N ₂ O	42,373.9	3.4%	7%	8	0.25%	7
2. Industrial Processes (HFCs,PFCs,SF ₆)	HFCs, PFCs, SF ₆	23,524.2	1.9%	33%	2	0.66%	4
3. Solvent & other Product Use	N ₂ O	99.0	0.0%	5%	9	0.00%	10
4. Agriculture	CH ₄ , N ₂ O	25,499.6	2.0%	18%	6	0.39%	6
5. LULUCF	CO ₂ , CH ₄ , N ₂ O	-73,179.1	-5.8%	12%	7	0.71%	3
6. Waste	CO ₂ , CH ₄ , N ₂ O	20,873.8	1.7%	32%	3	0.57%	5
Total Net Emissions	(D)	1,184,802.8	(E) ²⁾	2%			

$$1) C = A \times B / D$$

$$2) E = \sqrt{C_1^2 + C_2^2 + \dots}$$

1.7.2. KP-LULUCF Activity

Japan's net removals in FY2010 were 50 million tonnes (CO₂ equivalents) and the uncertainty was 12%. More detailed information on the uncertainty assessment is described in section 11.4.1.5 of chapter 11 and Annex 7.

Table 1-5 Uncertainty of Japan's KP-LULUCF activities

Greenhouse gas source and sink activities	GHGs	Emissions/Removals [Gg CO ₂ eq.]		Emissions/Removals Uncertainty [%]	Rank	Emissions/Removals Uncertainty as % of total national emissions [%]	Rank
			%				
Article 3.3 activities Afforestation and Reforestation	CO ₂ , N ₂ O, CH ₄	-426	-1%	36%	1	0%	3
Article 3.3 activities Deforestation	CO ₂ , N ₂ O, CH ₄	4,823	10%	26%	2	-2%	4
Article 3.4 activities Forest management	CO ₂ , N ₂ O, CH ₄	-53,252	-107%	11%	4	12%	1
Article 3.4 activities Revegetation	CO ₂ , N ₂ O, CH ₄	-1,130	-2%	17%	3	0%	2
Total		-49,985	-100%	12%			

1.8. General Assessment of the Completeness

In this inventory report, emissions from some categories are not estimated and reported as “NE”. In FY2006, GHG emissions and removals from categories that were previously reported as NE were newly estimated by analyzing categories such as those, which possibly result in the emission of considerable amount of GHGs, as well as those, which require substantial improvement in their estimation methodology.

Source categories reported as NE in this year's report include those whose emissions are thought to be very small, those whose emissions are unknown, and those for which emission estimation methods have not been developed. For these categories, further investigation on their emission possibility and the development of estimation methodologies will be carried out in accordance with Japan's QA/QC plan. See Annex 5 for a list of not-estimated emission source categories.

For some categories, dealing with the emission sources of HFCs, PFCs and SF₆, activity data are not available from CY 1990 to 1994. Those categories are therefore reported as “NE” during that period.

Chapter 2. Trends in GHGs Emissions and Removals

2.1. Description and Interpretation of Emission and Removal Trends for Aggregate GHGs

2.1.1. GHGs Emissions and Removals

Total GHGs emissions in FY2010^{1,2} (excluding LULUCF³) were 1,258 million tonnes (in CO₂ eq.). They increased by 4.4% compared to the emissions in FY1990⁴ (excluding LULUCF). Compared to the emissions in the base year under the Kyoto Protocol⁵, they decreased by 0.3%.

It should be noted that actual emissions of HFCs, PFCs, and SF₆ in the period from CY1990 to 1994 are not estimated (NE)⁶.

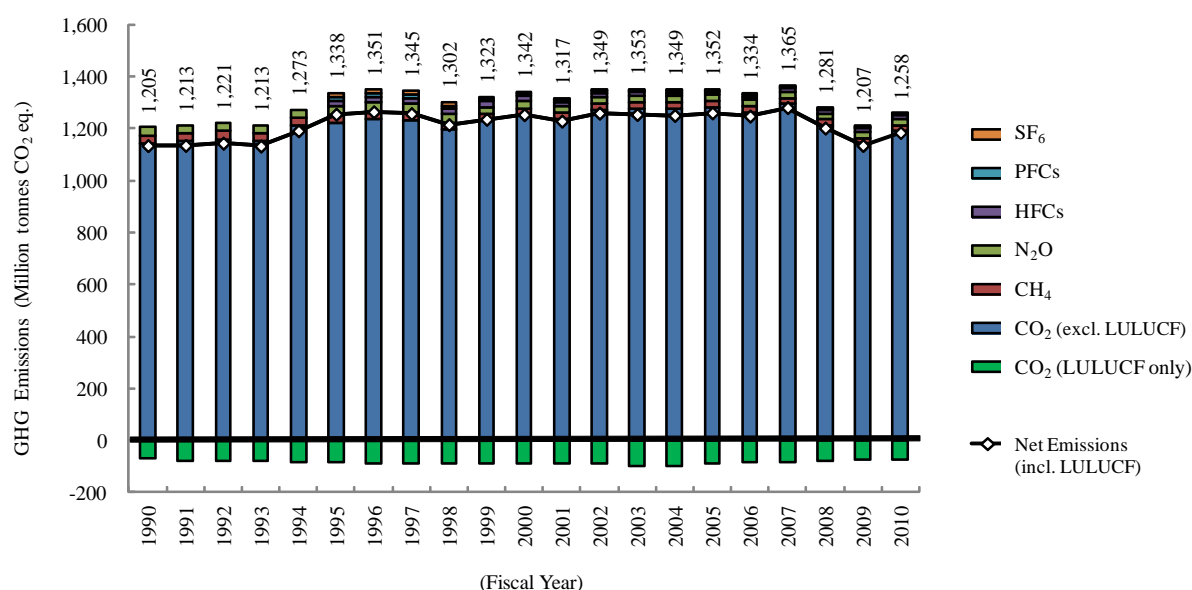


Figure 2-1 Trends in greenhouse gas emissions and removals in Japan

Carbon dioxide emissions in FY2010 were 1,192 million tonnes (excluding LULUCF), accounting for 94.8% of total GHGs emissions. They increased by 4.4% since FY1990 and increased by 4.4% compared to the previous year. Carbon dioxide removals⁷ in FY2010 were 73.2 million tonnes and were equivalent to 5.8% of total GHGs emissions. They increased by 4.3% since FY 1990 and increased by 1.8% compared to the previous year. Methane emissions in FY2010 (excluding

¹ “FY” (Fiscal Year), from April of the reporting year through March of the next year, is used because CO₂ is the primary GHGs emissions and estimated on a fiscal year basis. “CY” stands for “Calendar Year”.

² The sum of CO₂, CH₄, N₂O, HFCs, PFCs and SF₆ emissions converted to CO₂ equivalents, multiplied by their respective global warming potential (GWP). The GWP is a coefficient by means of which greenhouse gas effects of a given gas are made relative to those of an equivalent amount of CO₂. The coefficients are subjected to the *Second Assessment Report* (1995) issued by the Intergovernmental Panel on Climate Change (IPCC).

³ Abbreviation of “Land Use, Land-Use Change and Forestry”

⁴ The sum of CO₂, CH₄ and N₂O emissions converted to CO₂ equivalents multiplied by their respective GWP.

⁵ Japan’s base year under the Kyoto Protocol for CO₂, CH₄, N₂O emissions is FY 1990, while FY 1995 is the base year for HFCs, PFCs, and SF₆ emissions.

⁶ Potential emissions are reported in Common Reporting Format (CRF) for CY 1990 to 1994.

⁷ Since the inventory to be submitted under the UNFCCC reports all GHG emissions and removals from the LULUCF Sector, these values do not correspond to emissions and removals which can be accounted for compliance under the Kyoto Protocol (for ‘forest management’, 13 million carbon tonnes as the upper limit for Japan is given in the Appendix to the Annex to Decision 16/CMP.1.)

LULUCF) were 20.4 million tonnes (in CO₂ eq.), accounting for 1.6% of total GHGs emissions. They decreased by 36.2% since FY1990 and decreased by 2.1% compared to the previous year. Nitrous oxide emissions in FY2010 (excluding LULUCF) were 22.1 million tonnes (in CO₂ eq.), accounting for 1.8% of total GHGs emissions. They decreased by 30.3% since FY1990 and decreased by 2.2% compared to the previous year.

Hydrofluorocarbons emissions in CY2010 were 18.3 million tonnes (in CO₂ eq.), accounting for 1.5% of total GHGs emissions. They decreased by 9.9% since CY1995 and increased by 10.3% compared to the previous year. Perfluorocarbons emissions in CY2010 were 3.4 million tonnes (in CO₂ eq.), accounting for 0.3% of total GHGs emissions. They decreased by 76.1% since CY1995 and increased by 4.2% compared to the previous year. Sulphur hexafluoride emissions in CY2010 were 1.9 million tonnes (in CO₂ eq.), accounting for 0.1% of total GHGs emissions. They decreased by 89.0% since CY1995 and increased by 0.6% compared to the previous year.

Table 2-1 Trends in greenhouse gas emissions and removals in Japan

[Million tonnes CO ₂ eq.]	GWP	Base year of KP	1990	1991	1992	1993	1994	1995	1996	1997	1998	1999	2000	2001
CO ₂ (excl. LULUCF)	1	1,144.1	1,141.2	1,150.1	1,158.6	1,150.9	1,210.7	1,223.7	1,236.6	1,231.5	1,195.9	1,230.9	1,251.6	1,236.4
CO ₂ (incl. LULUCF)	1	NA	1,071.0	1,072.7	1,081.4	1,070.9	1,129.0	1,141.6	1,149.9	1,144.5	1,109.1	1,143.8	1,163.8	1,148.5
CO ₂ (LULUCF only)	1	NA	-70.2	-77.4	-77.1	-80.0	-81.7	-82.1	-86.8	-87.1	-86.9	-87.1	-87.8	-87.9
CH ₄ (excl. LULUCF)	21	33.4	32.0	31.8	31.5	31.2	30.6	29.7	29.0	27.9	27.1	26.5	25.9	25.1
CH ₄ (incl. LULUCF)	21	NA	32.0	31.8	31.5	31.3	30.6	29.7	29.0	27.9	27.1	26.5	25.9	25.1
N ₂ O (excl. LULUCF)	310	32.6	31.6	31.1	31.3	31.0	32.2	32.7	33.6	34.3	32.8	26.4	29.0	25.5
N ₂ O (incl. LULUCF)	310	NA	31.7	31.2	31.4	31.1	32.3	32.7	33.7	34.4	32.8	26.4	29.0	25.6
HFCs	HFC-134a: 1,300 etc.	20.2	NE	NE	NE	NE	NE	NE	20.3	19.9	19.4	19.9	18.8	16.2
PFCs	PFC-14: 6,500 etc.	14.0	NE	NE	NE	NE	NE	NE	14.2	14.8	16.2	13.4	10.4	7.9
SF ₆	23,900	16.9	NE	NE	NE	NE	NE	NE	17.0	17.5	15.0	13.6	9.3	6.0
Gross Total (excl. LULUCF)		1,261.3	1,204.9	1,213.0	1,221.4	1,213.2	1,273.5	1,337.5	1,351.4	1,344.8	1,302.3	1,323.4	1,341.9	1,317.1
Net Total (incl. LULUCF)		NA	1,134.8	1,135.7	1,144.3	1,133.3	1,191.9	1,255.6	1,264.8	1,257.8	1,215.5	1,236.3	1,254.2	1,229.2

[Million tonnes CO ₂ eq.]	GWP	2002	2003	2004	2005	2006	2007	2008	2009	2010	Emission increase from the base year of KP	Emission increase from 1990 (2010)	Emission increase from 1995 (2010)	Emission increase from previous year (2010)
CO ₂ (excl. LULUCF)	1	1,273.5	1,278.6	1,278.0	1,282.3	1,263.1	1,296.3	1,213.2	1,142.3	1,191.9	4.2%	4.4%	-	4.4%
CO ₂ (incl. LULUCF)	1	1,184.4	1,180.4	1,180.3	1,191.5	1,178.1	1,212.2	1,134.5	1,070.4	1,118.8	-	4.5%	-	4.5%
CO ₂ (LULUCF only)	1	-89.1	-98.2	-97.7	-90.7	-85.0	-84.2	-78.7	-71.9	-73.2	-	4.3%	-	1.8%
CH ₄ (excl. LULUCF)	21	24.2	23.7	23.2	22.9	22.5	22.1	21.5	20.9	20.4	-38.8%	-36.2%	-	-2.1%
CH ₄ (incl. LULUCF)	21	24.2	23.7	23.2	22.9	22.5	22.1	21.5	20.9	20.4	-	-36.2%	-	-2.1%
N ₂ O (excl. LULUCF)	310	24.8	24.5	24.5	24.1	24.1	22.8	22.8	22.6	22.1	-32.4%	-30.3%	-	-2.2%
N ₂ O (incl. LULUCF)	310	24.8	24.5	24.6	24.1	24.1	22.8	22.8	22.6	22.1	-	-30.5%	-	-2.2%
HFCs	HFC-134a: 1,300 etc.	13.7	13.8	10.6	10.5	11.7	13.3	15.3	16.6	18.3	-9.7%	-	-9.9%	10.3%
PFCs	PFC-14: 6,500 etc.	7.4	7.2	7.5	7.0	7.3	6.4	4.6	3.3	3.4	-75.8%	-	-76.1%	4.2%
SF ₆	23,900	5.6	5.3	5.1	4.8	4.9	4.4	3.8	1.9	1.9	-89.0%	-	-89.0%	0.6%
Gross Total (excl. LULUCF)		1,349.1	1,353.0	1,348.9	1,351.5	1,333.6	1,365.3	1,281.3	1,207.4	1,258.0	-0.3%	4.4%	-5.9%	4.2%
Net Total (incl. LULUCF)		1,260.1	1,254.8	1,251.2	1,260.8	1,248.6	1,281.1	1,202.6	1,135.5	1,184.8	-	4.4%	-	4.3%

* NA: Not Applicable

* NE: Not Estimated

* LULUCF: Land Use, Land-Use Change and Forestry

2.1.2. CO₂ Emissions per Capita

Total CO₂ emissions in FY 2010 (excluding LULUCF) were 1,192 million tonnes, and on a per capita basis, they were 9.31 tonnes. Compared to FY1990, they increased by 4.4% in total emissions, and increased by 0.8% in per capita emissions. Compared to the previous year, they increased by 4.4% in total emissions, and increased by 3.9% in per capita emissions.

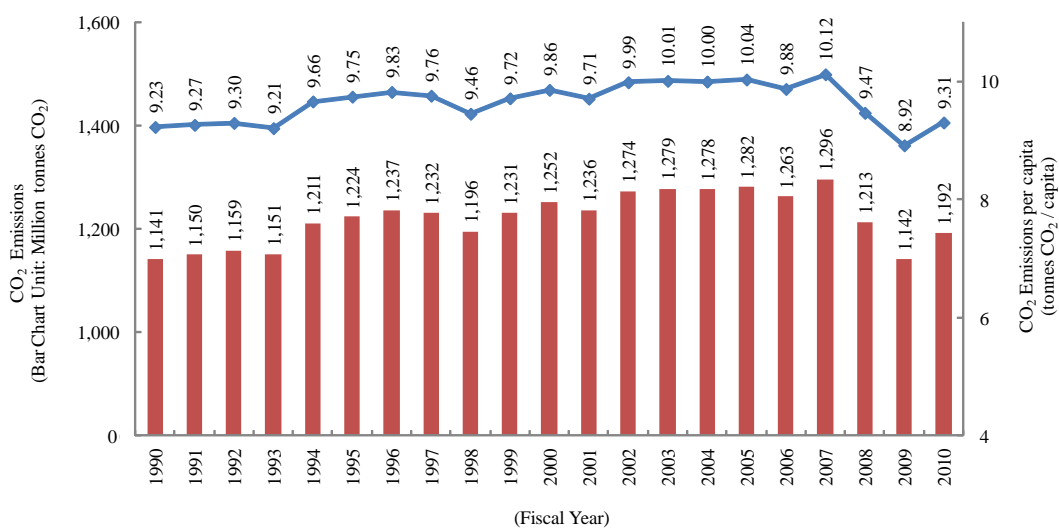


Figure 2-2 Trends in total CO₂ emissions and CO₂ emissions per capita

Source of population data: Ministry of Public Management, Home Affairs, Posts and Telecommunications Japan, *Population Census and Annual Report on Current Population Estimates*

2.1.3. CO₂ Emissions per Unit of GDP

Carbon dioxide emissions per unit of GDP (million yen) in FY2010 were 2.33 tonnes. They decreased by 7.3% since FY1990 and increased by 1.2% compared to the previous year.

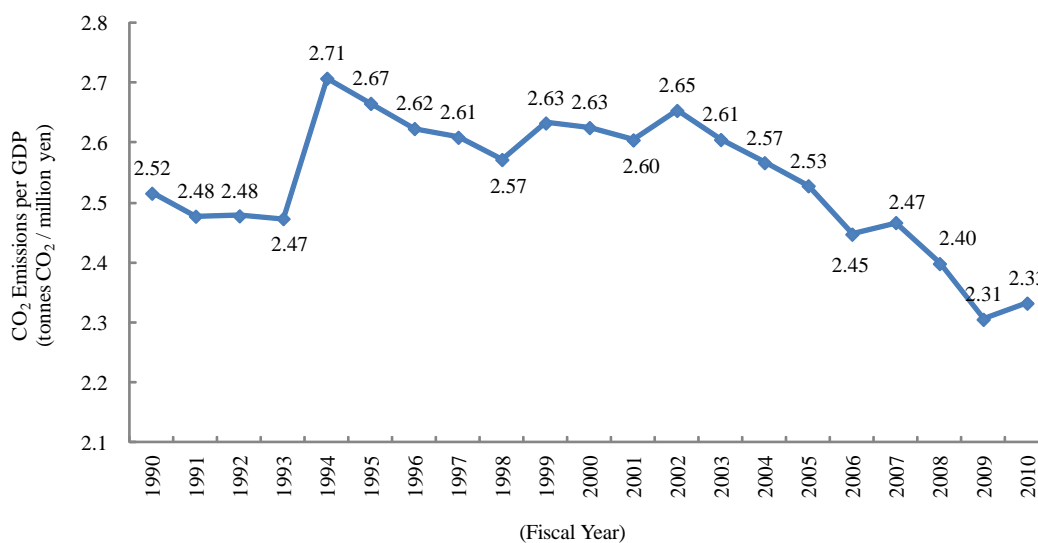


Figure 2-3 Trends in CO₂ emissions per unit of GDP

Source of GDP data: 1990-1993: EDMC / 1994-2010: Cabinet Office, Government of Japan, *Annual Report on National Accounts*

2.2. Description and Interpretation of Emission and Removal Trends by Gas

2.2.1. CO₂

Carbon dioxide emissions in FY2010 were 1,192 million tonnes (excluding LULUCF), accounting for 94.8% of total GHGs emissions. They increased by 4.4% since FY 1990 and increased by 4.4% compared to the previous year.

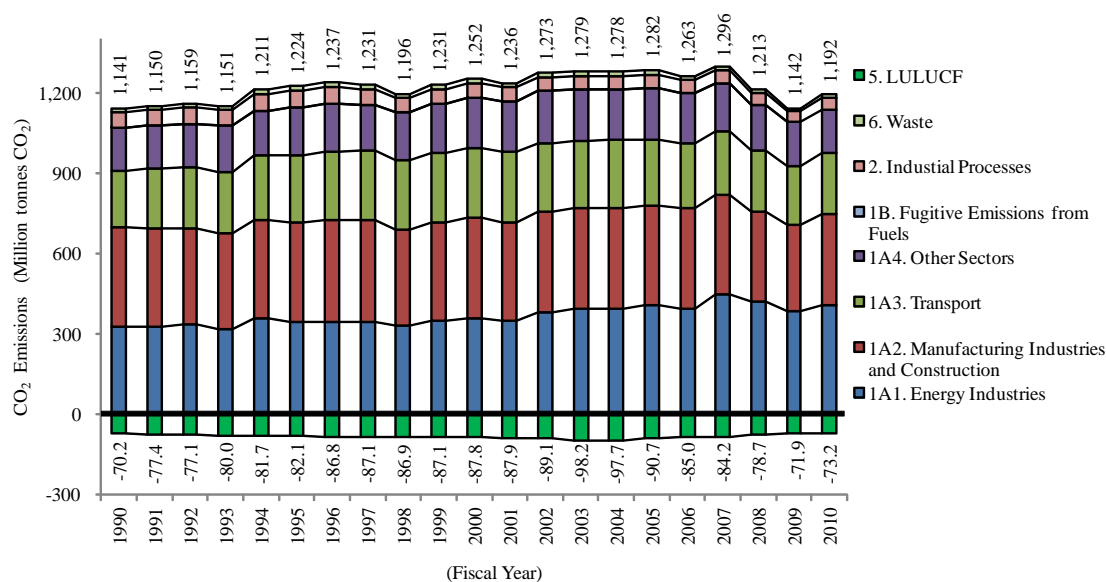


Figure 2-4 Trends in CO₂ emissions

The breakdown of CO₂ emissions in FY2010 shows that the Fuel Combustion, accounting for 95.4%. It is followed by the Industrial Processes (3.5%) and the Waste sectors (1.1%). As for the breakdown of CO₂ emissions within the Fuel Combustion, the Energy Industries accounts for 35.7% and is followed by the Industries at 30.1%, the Transport at 19.8%, and the Other Sectors⁸ at 14.4%.

The main driving factor for the increase in CO₂ emissions compared to the previous year is the recovery from the economic recession induced by the Global Financial Crisis of 2008; CO₂ emissions from the Industries sector increased because of the higher levels of manufacturing. In addition, electric power demand increased due to the relatively high number of days on which extremes of hot or cold were experienced.

By looking at the changes in emissions by sector, emissions from the Fuel Combustion in the Energy Industries increased by 25.2% since FY1990 and increased by 5.2% compared to the previous year. The main driving factor for the increase compared to the base year is the increase in electricity consumption.

Emissions from the Industries decreased by 7.7% since FY1990 and increased by 7.4% compared to the previous year.

Emissions from the Transport increased by 6.6% compared to FY1990 and increased by 1.0%

⁸ It covers emissions from Commercial/Institutional, Residential and Agriculture/Forestry/Fisheries.

compared to the previous year. The main driving factor for the increase compared to the base year is increase in demand for passenger transportation, compensating decrease in volume of freight transportation.

Emissions from the Other Sectors increased by 1.4% since FY1990 and increased by 1.5% compared to the previous year.

Carbon dioxide removals in FY2010 were 73.2 million tonnes, and they were equivalent to 5.8% of total GHGs emissions. They increased by 4.3% since FY 1990 and increased by 1.8% compared to the previous year.

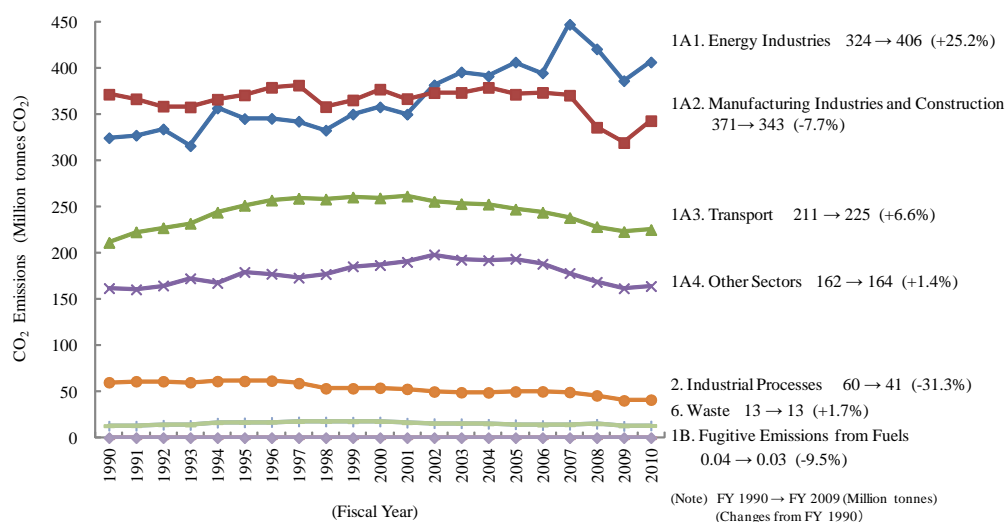


Figure 2-5 Trends in CO₂ emissions in each sector

(Figures in brackets indicate relative increase or decrease to the FY 1990 values)

Table 2-2 Trends in CO₂ emissions and removals in each sector

Category	1990	1995	2000	2005	2008	2009	2010
IA. Fuel Combustion	1,068,260	1,145,769	1,180,044	1,217,696	1,152,418	1,089,142	1,137,551
IA1. Energy Industries	324,253	344,948	357,574	406,039	420,263	385,896	406,096
Public Electricity and Heat Production	297,074	315,399	330,863	378,921	394,714	357,104	380,094
Petroleum Refining	15,893	16,956	17,285	16,441	14,324	14,564	15,001
Manufacture of Solid Fuels and Other Energy Industries	11,286	12,592	9,426	10,677	11,225	14,228	11,001
IA2. Manufacturing Industries and Construction	371,311	370,539	376,778	371,229	335,619	318,978	342,609
Iron and Steel	149,600	141,862	150,776	152,741	143,269	134,610	151,872
Non-Ferrous Metals	6,092	4,770	3,042	2,634	2,333	2,120	2,096
Chemicals	64,736	74,806	67,216	58,650	53,325	52,549	53,617
Pulp, Paper and Print	25,825	29,449	29,035	26,552	22,843	21,239	20,323
Food Processing, Beverages and Tobacco	13,129	14,407	13,161	11,326	8,862	8,761	8,817
Other Manufacturing	111,929	105,245	113,547	119,326	104,987	99,698	105,884
IA3. Transport	211,054	251,167	259,076	247,010	228,099	222,768	224,943
Civil Aviation	7,162	10,278	10,677	10,799	10,277	9,781	9,193
Road Transportation	189,228	225,381	232,827	222,652	205,933	202,018	204,277
Railways	932	819	707	644	600	586	588
Navigation	13,731	14,687	14,865	12,915	11,288	10,383	10,885
IA4. Other Sectors	161,641	179,115	186,615	193,419	168,436	161,500	163,902
Commercial/Institutional	83,593	93,269	101,450	110,678	98,756	93,283	92,336
Residential	56,668	66,320	68,958	67,583	59,023	57,792	61,095
Agriculture/Forestry/Fisheries	21,380	19,526	16,207	15,158	10,657	10,425	10,472
1B. Fugitive Emissions from Fuels	37	51	36	38	38	35	33
2. Industrial Processes	59,934	61,338	53,983	50,031	45,739	40,314	41,177
Mineral Products	55,369	56,761	49,842	46,903	43,009	37,714	38,280
Chemical Industry	4,209	4,220	3,893	2,887	2,574	2,488	2,737
Metal Production	356	357	248	242	156	112	160
5. LULUCF	-70,175	-82,056	-87,780	-90,742	-78,707	-71,873	-73,188
6. Waste	12,966	16,534	17,494	14,491	15,012	12,763	13,186
Total (including LULUCF)	1,071,021	1,141,636	1,163,777	1,191,515	1,134,500	1,070,381	1,118,760
Total (excluding LULUCF)	1,141,196	1,223,693	1,251,557	1,282,257	1,213,206	1,142,254	1,191,947

* LULUCF: Land Use, Land-Use Change and Forestry

2.2.2. CH₄

Methane emissions in FY2010 were 20.4 million tonnes (in CO₂ eq., including LULUCF), accounting for 1.6% of total GHGs emissions. They decreased by 36.2% since FY1990 and decreased by 2.1% compared to the previous year.

The breakdown of CH₄ emissions in FY2010 shows that the largest source is the Enteric Fermentation, which accounts for 33%. It is followed by the Rice Cultivation (27%) and the SWDS (16%).

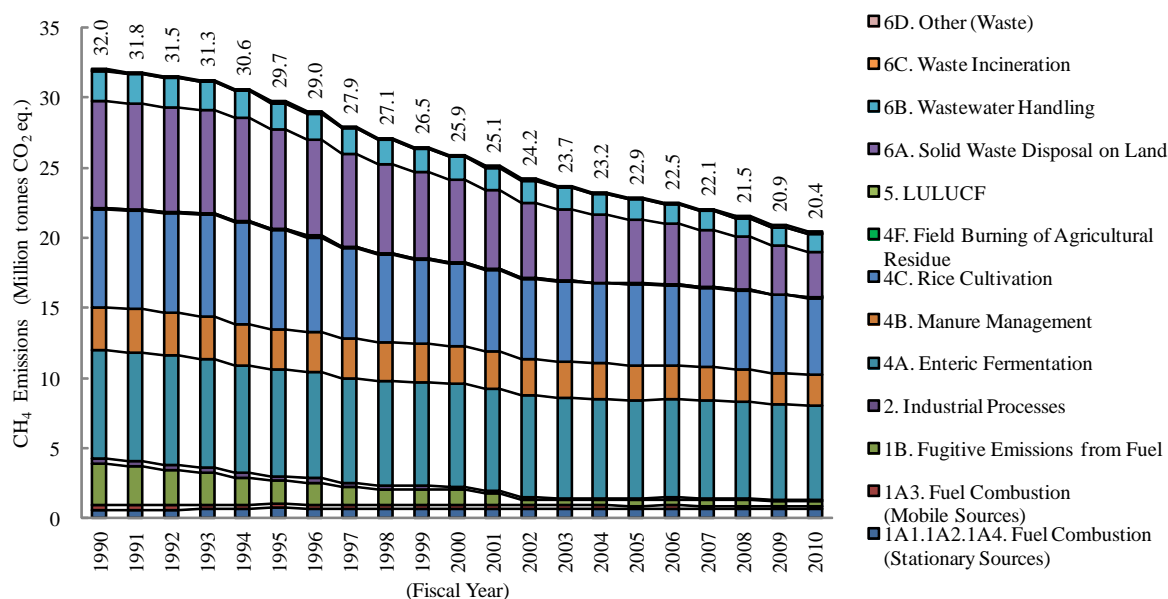


Figure 2-6 Trends in CH₄ emissions

Table 2-3 Trends in CH₄ emissions

[Thousand tonnes CO₂ eq.]

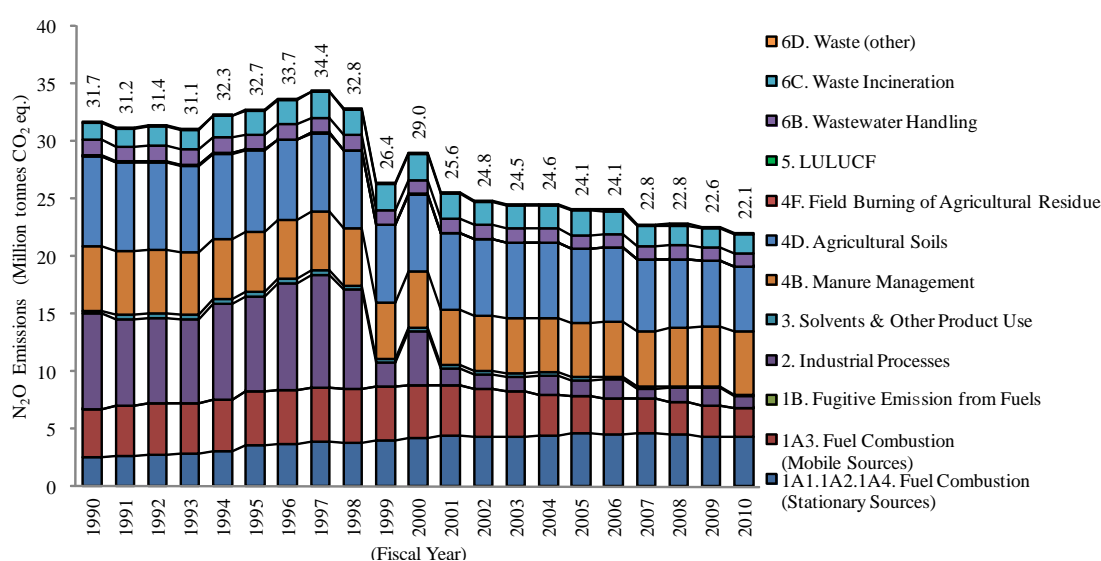
Category	1990	1995	2000	2005	2008	2009	2010
1A. Fuel Combustion	890	1,038	963	888	866	833	842
1A1. Energy Industries	30	34	44	37	44	42	46
1A2. Industries	355	438	352	351	367	364	373
1A3. Transport	298	309	298	237	191	179	168
1A4. Other Sectors	207	257	270	262	263	248	256
1B. Fugitive Emissions from Fuels	3,037	1,610	1,043	396	408	394	376
1B1. Solid Fuels	2,806	1,345	769	74	46	46	44
1B2. Oil & Natural Gas	231	265	274	322	362	348	331
2. Industrial Processes	358	322	196	134	121	110	119
4. Agriculture	17,831	17,676	16,045	15,309	14,876	14,625	14,387
4A. Enteric Fermentation	7,677	7,606	7,370	7,002	6,913	6,773	6,673
4B. Manure Management	3,094	2,893	2,678	2,503	2,302	2,247	2,205
4C. Rice Cultivation	6,960	7,083	5,920	5,739	5,599	5,545	5,452
4F. Field Burning of Agricultural Residue	101	94	77	65	62	59	57
5. LULUCF	9	9	8	9	22	9	2
6. Waste	9,914	9,081	7,645	6,128	5,250	4,919	4,719
6A. Solid Waste Disposal on Land	7,645	7,076	5,878	4,569	3,759	3,517	3,270
6B. Wastewater Handling	2,144	1,884	1,657	1,419	1,322	1,273	1,270
6C. Waste Incineration	13	15	13	14	12	11	10
6D. Other (Waste)	112	106	96	126	157	118	169
Total (including LULUCF)	32,039	29,736	25,900	22,864	21,543	20,889	20,445
Total (excluding LULUCF)	32,030	29,728	25,892	22,855	21,521	20,881	20,443

* LULUCF: Land Use, Land-Use Change and Forestry

2.2.3. N₂O

Nitrous oxide emissions in FY 2010 were 22.1 million tonnes (in CO₂ eq., including LULUCF), accounting for 1.8% of total GHGs emissions. They decreased by 30.5% since FY 1990 and decreased by 2.2% compared to the previous year. Their decrease since FY 1990 is mainly a result of a decrease in emissions from Industrial Processes (e.g. adipic acid production (87%)). There is a sharp decline in emissions from the Industrial Processes from FY 1998 to 1999, as N₂O abatement equipment came on stream in the adipic acid production plant in March 1999. However the N₂O emissions increased in FY 2000 because of a decrease in the equipment's efficiency; the emissions decreased again in FY 2001 with the resumption of normal operation.

The breakdown of N₂O emissions in FY 2010 shows that the largest source is the Agricultural Soils accounting for 25%. It is followed by the Manure Management (25%) and the Fuel Combustion (Stationary Sources) (19%).

Figure 2-7 Trends in N₂O emissionsTable 2-4 Trends in N₂O emissions

Category	1990	1995	2000	2005	2008	2009	2010
1A. Fuel Combustion	6,752	8,285	8,788	7,913	7,355	7,011	6,809
1A1. Energy Industries	922	1,413	1,709	2,119	2,118	2,032	2,005
1A2. Industries	1,350	1,871	2,126	2,093	2,051	1,978	1,942
1A3. Transport	4,206	4,652	4,589	3,320	2,843	2,672	2,526
1A4. Other Sectors	273	348	363	380	342	330	336
1B. Fugitive Emissions from Fuels	0.1	0.2	0.1	0.1	0.1	0	0
2. Industrial Processes	8,267	8,213	4,690	1,300	1,262	1,559	1,078
3. Solvent & Other Product Use	287	438	341	266	129	120	99
4. Agriculture	13,430	12,363	11,585	11,212	11,034	10,960	11,112
4B. Manure Management	5,533	5,152	4,885	4,748	5,019	5,247	5,475
4D. Agricultural Soils	7,864	7,179	6,674	6,443	5,996	5,694	5,619
4F. Field Burning of Agricultural Residue	33	32	25	21	20	19	18
5. LULUCF	91	62	33	16	11	8	6
6. Waste	2,914	3,358	3,561	3,373	3,039	2,921	2,969
6B. Wastewater Handling	1,295	1,252	1,216	1,166	1,161	1,133	1,132
6C. Waste Incineration	1,519	2,012	2,260	2,096	1,739	1,684	1,688
6D. Waste (other)	99	94	85	112	139	105	150
Total (including LULUCF)	31,740	32,718	28,997	24,081	22,829	22,580	22,074
Total (excluding LULUCF)	31,649	32,656	28,965	24,065	22,819	22,572	22,067

* LULUCF: Land Use, Land-Use Change and Forestry

2.2.4. HFCs

Hydrofluorocarbons emissions in CY 2010⁹ were 18.3 million tonnes (in CO₂ eq.), accounting for 1.5% of total GHGs emissions. They decreased by 9.9% since CY 1995, and increased by 10.3% compared to the previous year. Their decrease since CY 1995 (-99.8%) is mainly a result of a decrease in HFC-23, a by-product of HCFC-22 production that have been regulated under Act on the Protection of the Ozone Layer Through the Control of Specified Substances and Other Measures.

The breakdown of HFCs emissions in CY 2010 shows that the largest source is refrigerants of the Refrigeration and Air Conditioning Equipment accounting for 94%, and is followed by the Aerosols / MDI (4%).

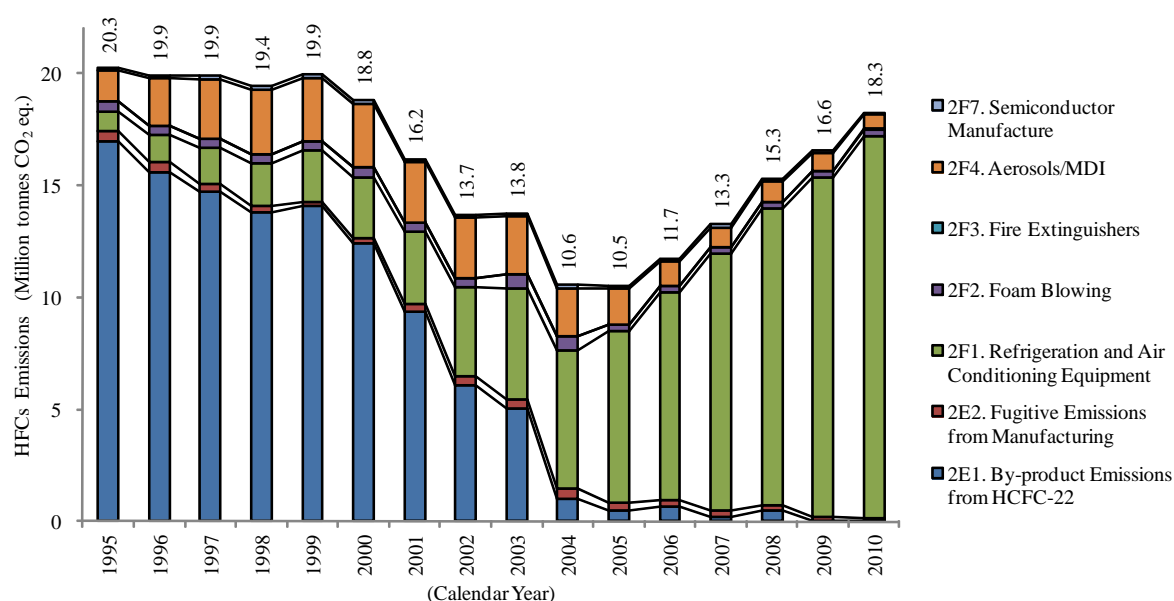


Figure 2-8 Trends in HFCs emissions

Table 2-5 Trends in HFCs emissions

Category	1995	2000	2005	2008	2009	2010
2E. Productions of F-gas	17,445	12,660	816	701	222	128
2E1. By-product Emissions from Production of HCFC-22	16,965	12,402	463	469	40	42
2E2. Fugitive Emissions	480	258	353	232	182	86
2F. Consumption of F-gases	2,815	6,141	9,702	14,597	16,332	18,128
2F1. Refrigeration and Air Conditioning Equipment	840	2,689	7,667	13,269	15,134	17,088
2F2. Foam Blowing	452	440	316	286	290	291
2F3. Fire Extinguishers	NO	3.7	5.9	6.3	7	7
2F4. Aerosols/MDI	1,365	2,834	1,572	890	809	640
2F7. Semiconductor Manufacture	158	174	141	146	92	102
Total	20,260	18,800	10,518	15,298	16,554	18,257

⁹ Emissions of HFCs, PFCs and SF₆ are estimated on a calendar year (CY) basis.

2.2.5. PFCs

Perfluorocarbons emissions in CY 2010 were 3.4 million tonnes (in CO₂ eq.), accounting for 0.3% of total GHGs emissions. They decreased by 76.1% since CY 1995, and increased by 4.2% compared to the previous year. Their decrease since CY 1995 (-87%) is mainly a result of a decrease in emissions from the Solvents.

The breakdown of PFCs emissions in CY 2010 shows that the largest source is the Semiconductor for Manufacture accounting for 53%. It is followed by the Solvents such as the ones for washing metals (40%) and the Fugitive Emissions from manufacturing (6%).

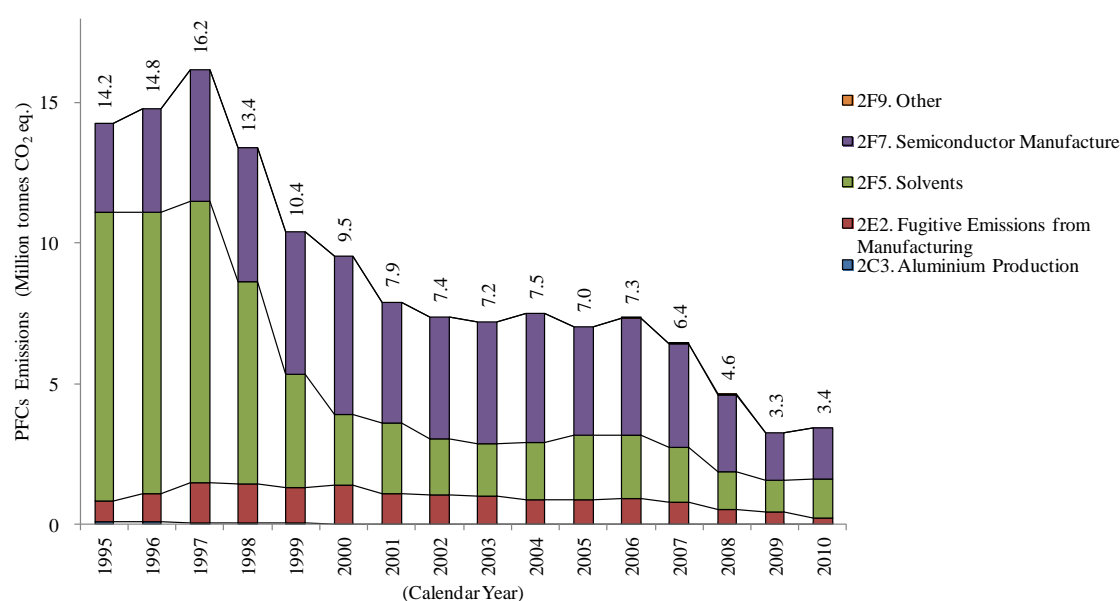


Figure 2-9 Trends in PFCs emissions

Table 2-6 Trends in PFCs emissions

[Thousand tonnes CO ₂ eq.]						
Category	1995	2000	2005	2008	2009	2010
2C3. Aluminium Production	70	18	15	15	11	10
2E2. Fugitive Emissions	763	1,359	837	524	399	200
2F. Consumption of F-gases	13,408	8,143	6,150	4,079	2,857	3,195
2F5. Solvents	10,264	2,506	2,289	1,318	1,142	1,376
2F7. Semiconductor Manufacture	3,144	5,637	3,861	2,756	1,715	1,819
2F9. Other	NE,NO	NE,NO	NE,NO	4.7	NA,NE,NO	NA,NE,NO
Total	14,240	9,519	7,002	4,618	3,268	3,405

2.2.6. SF₆

Sulphur hexafluoride emissions in CY 2010 were 1.9 million tonnes (in CO₂ eq.), accounting for 0.1% of total GHGs emissions. They decreased by 89.0% since CY 1995, and increased by 0.6% compared to the previous year. Their decrease since CY 1995 (-94%) is mainly a result of a decrease from the Electrical Equipment, due to the strengthening of the management of gases largely in electric power companies.

The breakdown of SF₆ emissions in CY 2010 shows that the largest source is Semiconductor Manufacture accounting for 38%. It is followed by Electrical Equipment (35%) and Aluminium and Magnesium Foundries (17%).

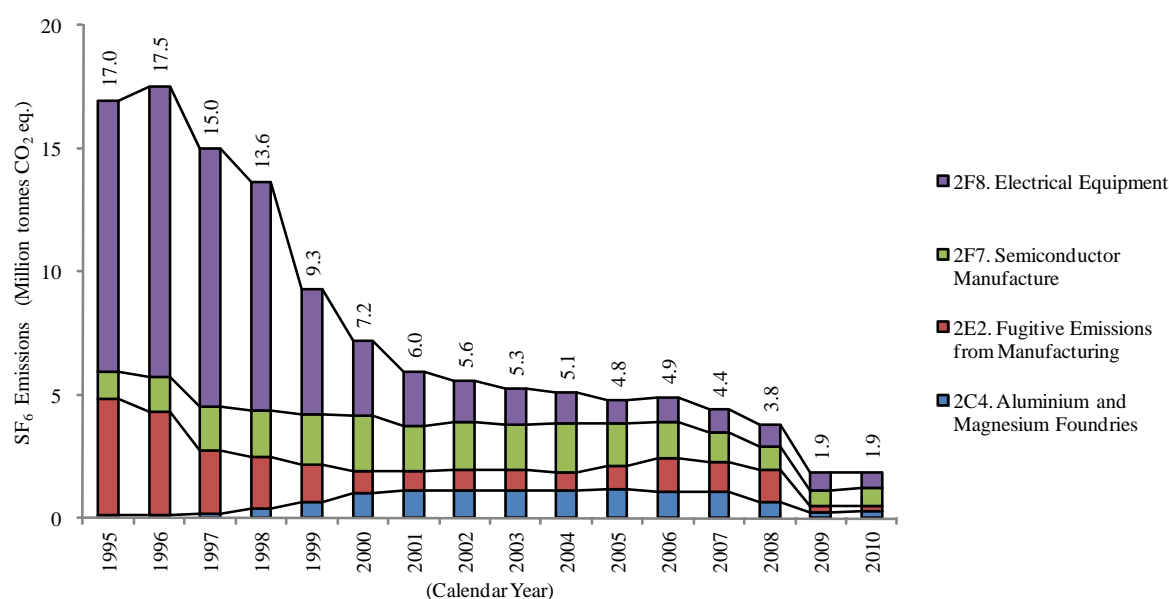


Figure 2-10 Trends in SF₆ emissions

Table 2-7 Trends in SF₆ emissions

[Thousand tonnes CO₂ eq.]

Category	1995	2000	2005	2008	2009	2010
2C4. SF ₆ Used in Aluminium and Magnesium Foundries	120	1,028	1,157	652	239	308
2E2. Fugitive Emissions	4,708	860	975	1,288	261	198
2F. Consumption of F-gases	12,134	5,300	2,676	1,855	1,352	1,356
2F7. Semiconductor Manufacture	1,129	2,250	1,733	952	606	704
2F8. Electrical Equipment	11,005	3,050	943	902	745	652
Total	16,961	7,188	4,808	3,795	1,851	1,862

2.3. Description and Interpretation of Emission and Removal Trends by Categories

The breakdown of GHGs emissions and removals in FY 2010 by sector¹⁰ shows that the Energy accounts for 91.1% of total GHGs emissions. It is followed by the Industrial Processes (5.2%), the Agriculture (2.0%), the Waste (1.7%) and the Solvents and Other Product Use (0.01%).

Removals by the LULUCF in FY 2010 were equivalent to 5.8% of total GHGs emissions.

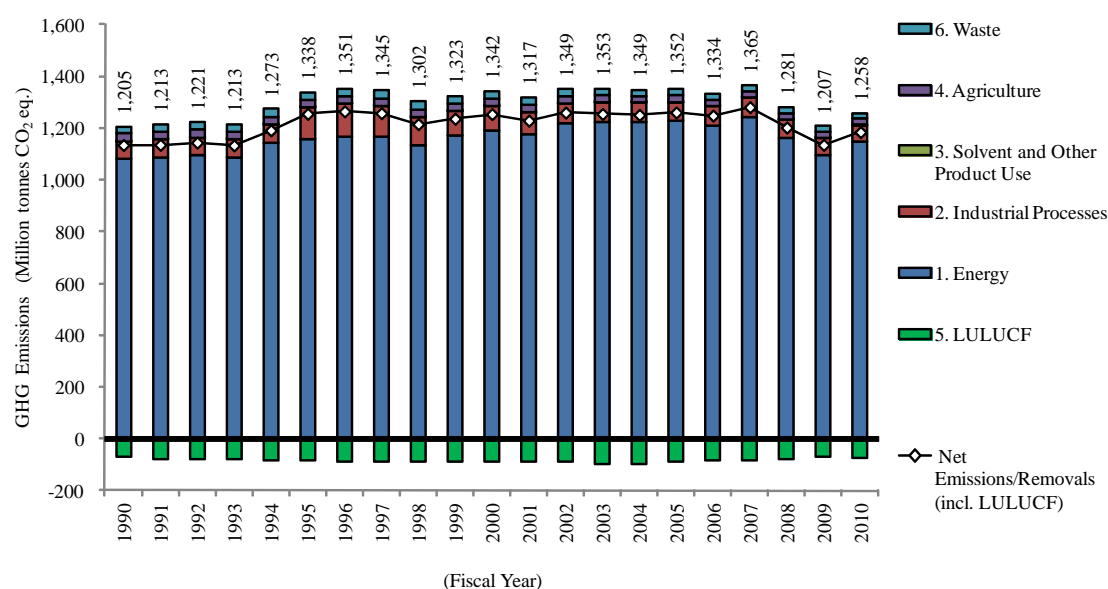


Figure 2-11 Trends in greenhouse gas emissions and removals in each sector

Table 2-8 Trends in greenhouse gas emissions and removals in each sector

[Million tonnes CO ₂ eq.]	1990	1991	1992	1993	1994	1995	1996	1997	1998	1999	2000
1. Energy	1,079.0	1,086.8	1,094.2	1,087.7	1,143.7	1,156.8	1,168.9	1,165.8	1,135.6	1,171.0	1,190.9
2. Industrial Processes	68.6	68.9	68.8	67.6	69.8	121.3	123.5	120.1	108.6	95.3	94.4
3. Solvent and Other Product Use	0.3	0.4	0.4	0.4	0.4	0.4	0.4	0.4	0.4	0.4	0.3
4. Agriculture	31.3	31.2	31.2	31.1	30.7	30.0	29.4	28.7	28.3	27.9	27.6
5. LULUCF	-70.1	-77.3	-77.1	-79.9	-81.6	-82.0	-86.7	-87.0	-86.8	-87.1	-87.7
6. Waste	25.8	25.7	26.8	26.4	28.9	29.0	29.3	29.7	29.3	28.9	28.7
Net Emissions/Removals (incl. LULUCF)	1,134.8	1,135.7	1,144.3	1,133.3	1,191.9	1,255.6	1,264.8	1,257.8	1,215.5	1,236.3	1,254.2
Emissions (excl. LULUCF)	1,204.9	1,213.0	1,221.4	1,213.2	1,273.5	1,337.5	1,351.4	1,344.8	1,302.3	1,323.4	1,341.9

[Million tonnes CO ₂ eq.]	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010
1. Energy	1,178.0	1,217.8	1,223.5	1,223.3	1,226.9	1,208.3	1,241.9	1,161.1	1,097.4	1,145.6
2. Industrial Processes	84.4	78.0	76.7	73.9	73.8	75.8	74.4	70.8	63.7	65.9
3. Solvent and Other Product Use	0.3	0.3	0.3	0.3	0.3	0.2	0.2	0.1	0.1	0.1
4. Agriculture	27.4	27.1	26.9	26.7	26.5	26.4	26.1	25.9	25.6	25.5
5. LULUCF	-87.9	-89.0	-98.2	-97.7	-90.7	-85.0	-84.1	-78.7	-71.9	-73.2
6. Waste	27.1	25.9	25.6	24.7	24.0	22.7	22.7	23.3	20.6	20.9
Net Emissions/Removals (incl. LULUCF)	1,229.2	1,260.1	1,254.8	1,251.2	1,260.8	1,248.6	1,281.1	1,202.6	1,135.5	1,184.8
Emissions (excl. LULUCF)	1,317.1	1,349.1	1,353.0	1,348.9	1,351.5	1,333.6	1,365.3	1,281.3	1,207.4	1,258.0

* LULUCF: Land Use, Land-Use Change and Forestry

¹⁰ It implies "Category" indicated in the Revised 1996 IPCC Guidelines and CRF.

2.3.1. Energy

Emissions from the Energy sector in FY 2010 were 1,146 million tonnes (in CO₂ equivalents). They increased by 6.2% since FY 1990 and increased by 4.4% compared to the previous year.

The breakdown of GHGs emissions from this sector in FY 2010 shows that CO₂ from Fuel Combustion accounts for 99.3%. The largest source within the Fuel Combustion is the Liquid Fuel CO₂, which accounted for 42%, and is then followed by the Solid Fuel CO₂ (38%) and the Gaseous Fuel CO₂ (18%).

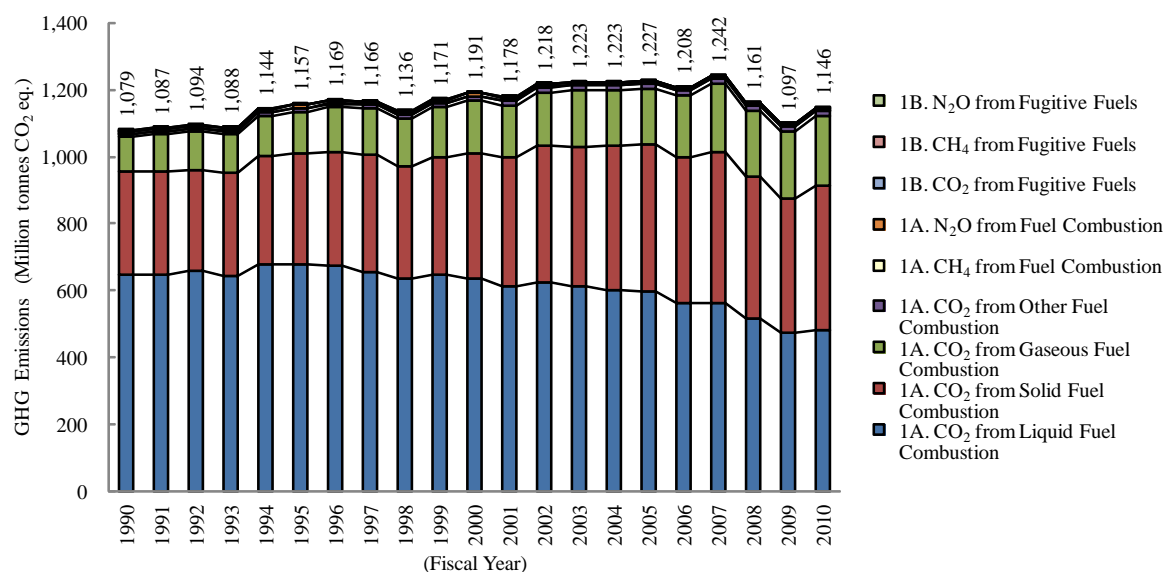


Figure 2-12 Trends in greenhouse gas emissions from the Energy sector

Table 2-9 Trends in greenhouse gas emissions from the Energy sector

[Thousand tonnes CO₂ eq.]

Source Category	1990	1995	2000	2005	2008	2009	2010
1A. Fuel Combustion	1,075,901	1,155,092	1,189,795	1,226,497	1,160,638	1,096,986	1,145,203
Liquid Fuel CO ₂	646,223	677,349	635,121	597,813	518,395	474,999	481,120
Solid Fuel CO ₂	308,620	331,720	376,521	437,937	420,521	401,560	431,476
Gaseous Fuel CO ₂	104,301	126,198	155,261	166,823	199,525	198,684	210,774
Other Fuels CO ₂ (Waste)	9,116	10,503	13,142	15,123	13,976	13,899	14,180
CH ₄	890	1,038	963	888	866	833	842
N ₂ O	6,752	8,285	8,788	7,913	7,355	7,011	6,809
1B. Fugitive Emissions from Fuel	3,074	1,661	1,079	433	446	430	409
CO ₂	37	51	36	38	38	35	33
CH ₄	3,037	1,610	1,043	396	408	394	376
N ₂ O	0.1	0.2	0.1	0.1	0.1	0	0
Total	1,078,975	1,156,753	1,190,874	1,226,930	1,161,084	1,097,416	1,145,612

2.3.2. Industrial Processes

Emissions from the Industrial Processes sector in FY 2010 were 65.9 million tonnes (in CO₂ eq.). They decreased by 3.9% since FY 1990, and increased by 3.5% compared to the previous year.

It should be noted that actual emissions of HFCs, PFCs, and SF₆ are not estimated (NE) for CY 1990 to 1994.

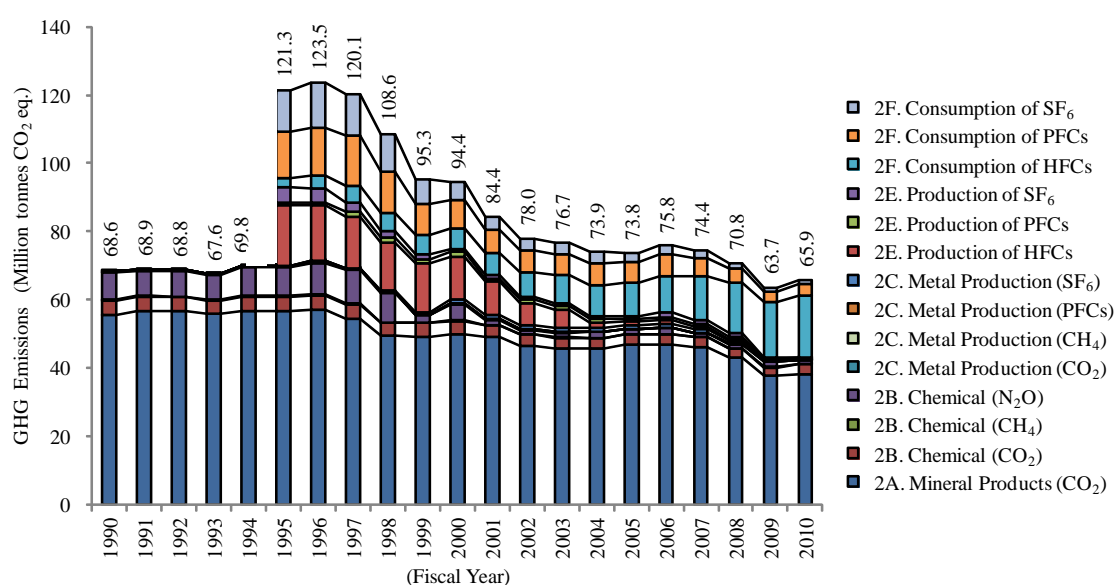


Figure 2-13 Trends in greenhouse gas emissions from the Industrial Processes sector

The breakdown of GHGs emissions from this sector in FY 2010 shows that the largest source is the Mineral Products such as CO₂ emissions from limestone in the cement production, accounting for 58%. It is followed by the Consumption of HFCs (28%) and the Consumption of PFCs (5%).

The main driving factors for decreases in CO₂, CH₄ and N₂O emissions since FY 1990 are the decrease in CO₂ emissions from cement production as the clinker production declined, and the decrease in N₂O emissions from adipic acid production as the N₂O abatement equipment came on stream. The main driving factors for decreases in HFCs, PFCs and SF₆ emissions since CY 1995 are the promotion of substitute materials use and of the capture and destruction of these gases.

Table 2-10 Trends in greenhouse gas emissions from the Industrial Processes sector

[Thousand tonnes CO₂ eq.]

Category	1990	1995	2000	2005	2008	2009	2010
2A. Mineral Products (CO ₂)	55,369	56,761	49,842	46,903	43,009	37,714	38,280
2B. Chemical Industry	12,814	12,737	8,762	4,304	3,943	4,144	3,919
CO ₂	4,209	4,220	3,893	2,887	2,574	2,488	2,737
CH ₄	338	304	179	117	106	97	104
N ₂ O	8,267	8,213	4,690	1,300	1,262	1,559	1,078
2C. Metal Production	375	564	1,311	1,431	838	375	493
CO ₂	356	357	248	242	156	112	160
CH ₄	19	18	17	17	15	13	15
PFCs	NE	70	18	15	15	11	10
SF ₆	NE	120	1,028	1,157	652	239	308
2E. Production of F-gas	NE	22,916	14,879	2,629	2,513	882	527
HFCs	NE	17,445	12,660	816	701	222	128
PFCs	NE	763	1,359	837	524	399	200
SF ₆	NE	4,708	860	975	1,288	261	198
2F. Consumption of F-gas	NE	28,356	19,584	18,528	20,531	20,541	22,679
HFCs	NE	2,815	6,141	9,702	14,597	16,332	18,128
PFCs	NE	13,408	8,143	6,150	4,079	2,857	3,195
SF ₆	NE	12,134	5,300	2,676	1,855	1,352	1,356
Total	68,559	121,335	94,377	73,793	70,834	63,656	65,898

2.3.3. Solvent and Other Product Use

Emissions from the Solvents and Other Product Use sector in FY 2010 were 100 thousand tonnes (in CO₂ eq.). They decreased by 65.5% since FY 1990, and decreased by 17.9% compared to the previous year. The only substance subject for estimation in this sector is laughing gas (N₂O) used as a general anesthetic in hospitals.

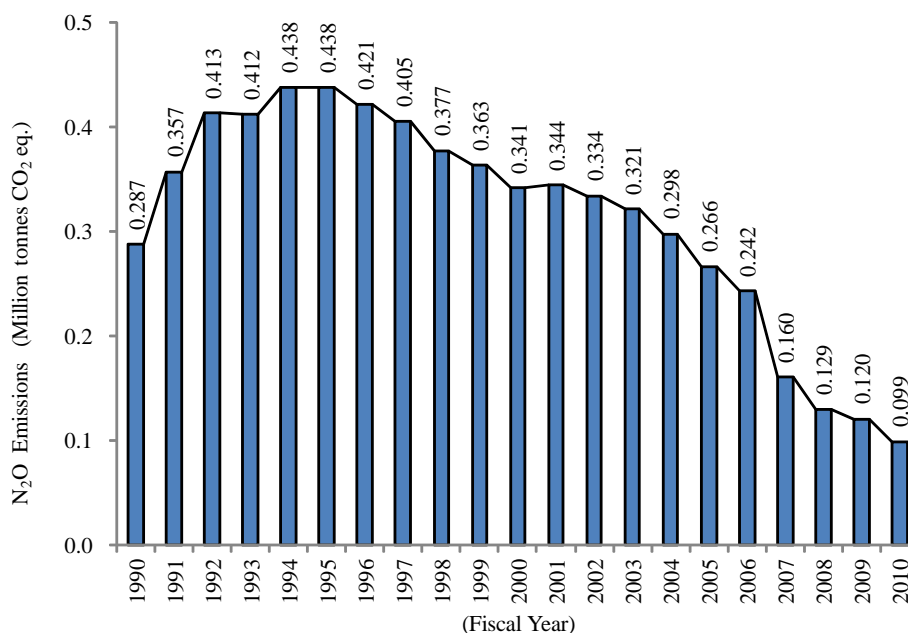


Figure 2-14 Trends in greenhouse gas emissions from the Solvent and Other Product Use sector

2.3.4. Agriculture

Emissions from the Agriculture sector in FY 2010 were 25.5 million tonnes (in CO₂ eq.). They decreased by 18.4% since FY 1990 and decreased by 0.3% compared to the previous year.

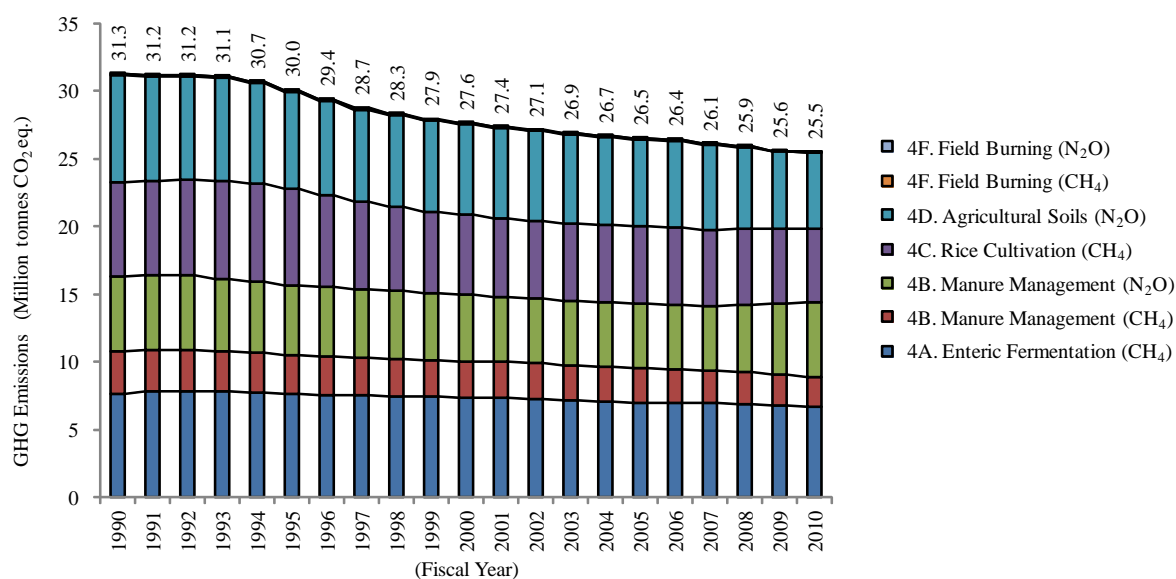


Figure 2-15 Trends in greenhouse gas emissions from the Agriculture sector

The breakdown of GHGs emissions from this sector in FY 2010 shows that the largest source is the Enteric Fermentation accounting for 26%. It is followed by the Agricultural Soils (22%) as a result of the nitrogen-based fertilizer applications, and the Rice Cultivation (21%).

The main driving factor for decrease in emissions since FY 1990 is the decrease in CH₄ emissions from the Rice Cultivation as a result of crop acreage decline, and the decrease in N₂O emissions from the Agricultural Soils, because the amount of nitrogen fertilizers applied to cropland had decreased.

Table 2-11 Trends in greenhouse gas emissions from the Agriculture sector

[Thousand tonnes CO₂ eq.]

Category	1990	1995	2000	2005	2008	2009	2010
4A. Enteric Fermentation(CH ₄)	7,677	7,606	7,370	7,002	6,913	6,773	6,673
4B. Manure Management	8,627	8,045	7,563	7,251	7,321	7,495	7,680
CH ₄	3,094	2,893	2,678	2,503	2,302	2,247	2,205
N ₂ O	5,533	5,152	4,885	4,748	5,019	5,247	5,475
4C. Rice Cultivation(CH ₄)	6,960	7,083	5,920	5,739	5,599	5,545	5,452
4D. Agricultural Soils (N ₂ O)	7,864	7,179	6,674	6,443	5,996	5,694	5,619
4F. Field Burning of Agricultural Res	133	126	103	87	82	78	76
CH ₄	101	94	77	65	62	59	57
N ₂ O	33	32	25	21	20	19	18
Total	31,261	30,039	27,629	26,521	25,910	25,585	25,500

2.3.5. Land Use, Land Use Change and Forestry (LULUCF)

Net Removals (including CO₂, CH₄ and N₂O emissions) from the LULUCF sector in FY 2010 was 73.2 million tonnes (in CO₂ eq.). They increased by 4.4% since FY 1990 and increased by 1.8% compared to the previous year. The decline trend in removals in recent years is largely due to maturity of Japanese forest.

The breakdown of GHGs emissions and removals from this sector in FY 2010 shows that the largest sink is the Forest land and its removals were 76.7 million tonnes accounting for 105% of this sector's net total emissions / removals.

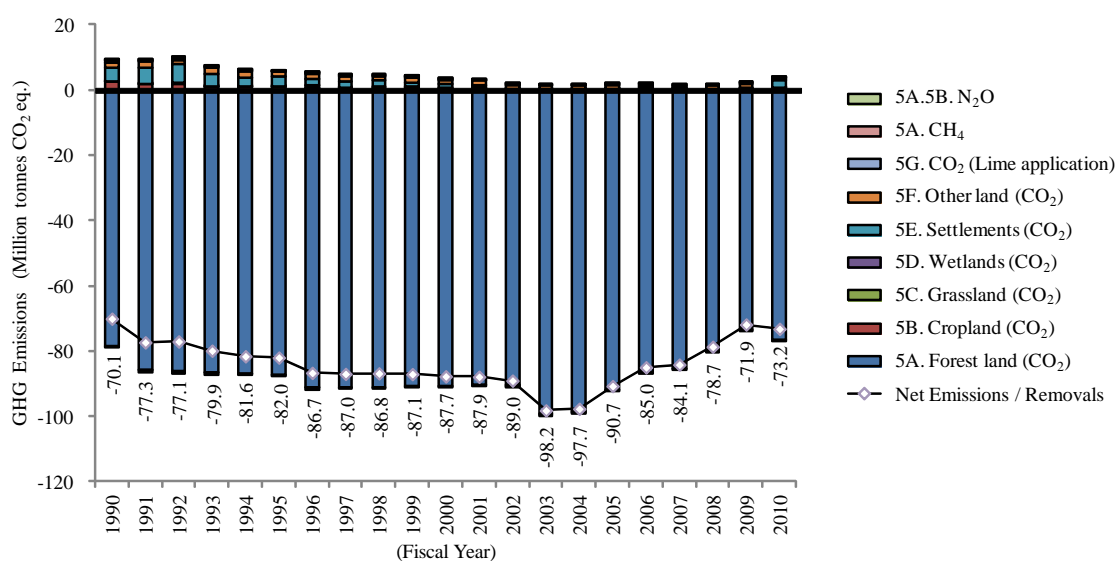


Figure 2-16 Trends in greenhouse gas emissions and removals from the LULUCF sector

Table 2-12 Trends in greenhouse gas emissions and removals from the LULUCF sector

[Thousand tonnes CO₂ eq.]

Category	1990	1995	2000	2005	2008	2009	2010
5A. Forest land	-78,583	-87,331	-90,681	-92,001	-79,904	-73,663	-76,675
CO ₂	-78,592	-87,341	-90,689	-92,011	-79,928	-73,673	-76,677
CH ₄	9	9	8	9	22	9	2
N ₂ O	0.9	0.9	0.8	0.9	2.2	0.9	0.2
5B. Cropland	2,603	883	388	292	233	265	459
CO ₂	2,513	823	356	277	224	258	452
CH ₄	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO
N ₂ O	90	61	32	15	8	8	6
5C. Grassland	-444	-481	-406	-336	-303	-276	-216
CO ₂	-444	-481	-406	-336	-303	-276	-216
CH ₄	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO
N ₂ O	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO
5D. Wetlands	86	360	451	16	16	23	82
CO ₂	86	360	451	16	16	23	82
CH ₄	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO
N ₂ O	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO
5E. Settlements	4,158	2,800	947	126	144	477	2,518
CO ₂	4,158	2,800	947	126	144	477	2,518
CH ₄	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO
N ₂ O	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO
5F. Other land	1,554	1,479	1,228	955	834	1,049	382
CO ₂	1,554	1,479	1,228	955	834	1,049	382
CH ₄	NO	NO	NO	NO	NO	NO	NO
N ₂ O	NO	NO	NO	NO	NO	NO	NO
5G. Other	550	303	333	231	306	270	270
CO ₂	550	303	333	231	306	270	270
Total	-70,075	-81,986	-87,739	-90,717	-78,674	-71,856	-73,179

2.3.6. Waste

Emissions from the Waste sector in FY 2010 were 20.9 million tonnes (in CO₂ eq.). They decreased by 19.1% since FY 1990 and increased by 1.3% compared to the previous year.

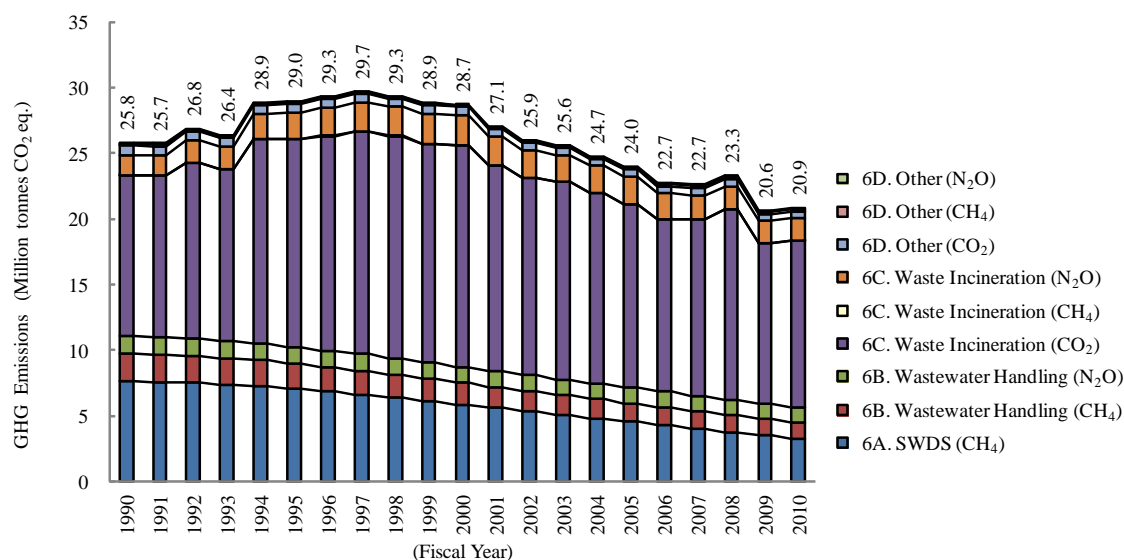


Figure 2-17 Trends in greenhouse gas emissions from the Waste sector

The breakdown of GHGs emissions from this sector in FY 2010 shows that the largest source is the Waste Incineration (CO₂), associated with waste derived from fossil fuels such as waste plastic and waste oil, accounting for 61%. It is followed by the SWDS (CH₄) (16%) and the Waste Incineration (N₂O) (8%), associated with waste substances including those that do not have a fossil fuel origin.

The main driving factor for decrease in emissions since FY 1990 is the decrease in CH₄ emissions from the SWDS as a result of decrease in the amount of disposal of biodegradable waste due to improvement of volume reduction ratio by intermediate treatment under Waste Management and Public Cleansing Act and other acts.

Table 2-13 Trends in greenhouse gas emissions from the Waste sector

[Thousand tonnes CO₂ eq.]

Category	1990	1995	2000	2005	2008	2009	2010
6A. Solid Waste Disposal on Land (CH ₄)	7,645	7,076	5,878	4,569	3,759	3,517	3,270
6B. Wastewater Handling	3,439	3,136	2,874	2,585	2,483	2,405	2,401
CH ₄	2,144	1,884	1,657	1,419	1,322	1,273	1,270
N ₂ O	1,295	1,252	1,216	1,166	1,161	1,133	1,132
6C. Waste Incineration	13,796	17,894	19,111	16,095	16,232	13,943	14,356
CO ₂	12,263	15,867	16,838	13,984	14,481	12,249	12,658
CH ₄	13	15	13	14	12	11	10
N ₂ O	1,519	2,012	2,260	2,096	1,739	1,684	1,688
6D. Other	914	868	837	744	826	737	847
CO ₂	703	668	656	507	530	514	528
CH ₄	112	106	96	126	157	118	169
N ₂ O	99	94	85	112	139	105	150
Total	25,794	28,974	28,700	23,993	23,300	20,603	20,874

2.4. Description and Interpretation of Emission Trends for Indirect GHGs and SO₂

Under the UNFCCC, it is required to report emissions not only 6 types of GHGs (CO₂, CH₄, N₂O, HFCs, PFCs and SF₆) that are controlled by the Kyoto Protocol, but also emissions of indirect GHGs (NO_x, CO and NMVOC) as well as SO₂. Their emission trends are indicated below.

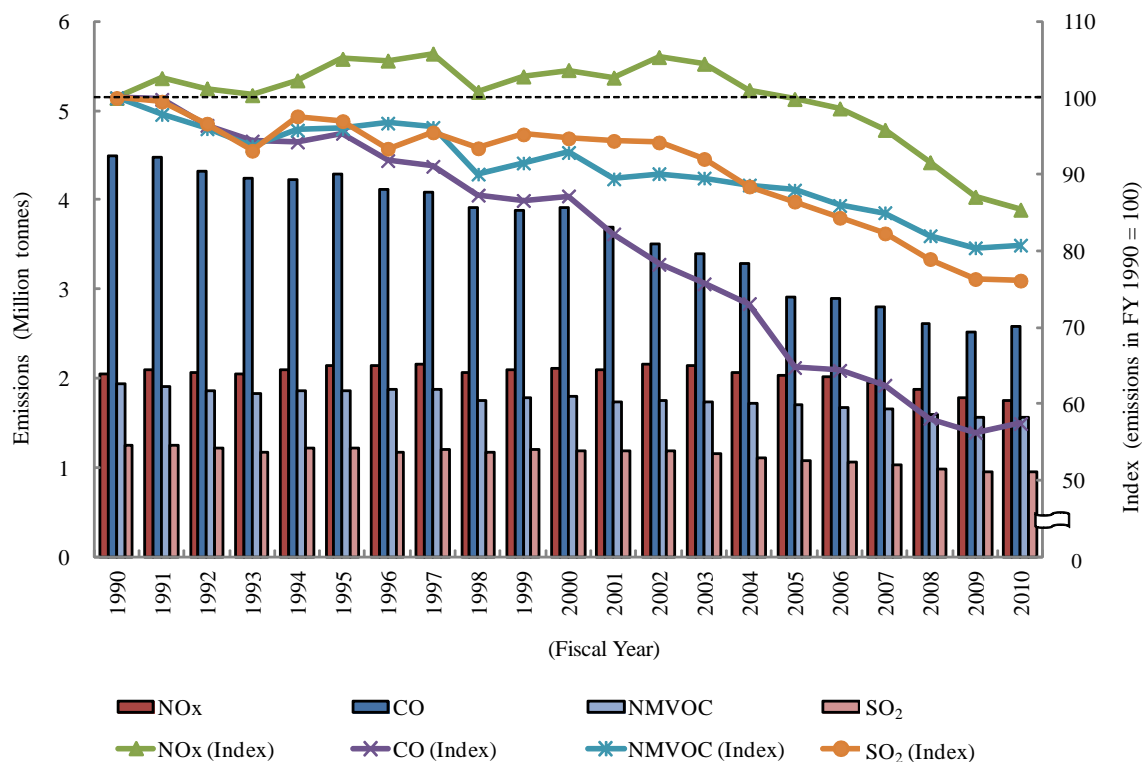


Figure 2-18 Trends in emissions of indirect greenhouse gases and SO₂

Nitrogen oxide (NO_x) emissions in FY 2010 were 1,744 thousand tonnes. They decreased by 14.6% since FY 1990 and decreased by 1.9% compared to the previous year.

Carbon monoxide (CO) emissions in FY 2010 were 2,577 thousand tonnes. They decreased by 42.6% since FY 1990 and increased by 2.1% compared to the previous year.

Non-methane volatile organic compounds (NMVOC) emissions in FY 2010 were 1,569 thousand tonnes. They decrease by 19.2% since FY 1990 and increased by 0.4% compared to the previous year.

Sulfur dioxide (SO₂) emissions in FY 2010 were 955 thousand tonnes. They decreased by 23.9% since FY 1990 and decreased by 0.3% compared to the previous year.

2.5. Emissions and removals from KP-LULUCF activities

The net removals from KP-LULUCF activities in FY2010 were 50.0 million tonnes (in CO₂ eq.). The breakdown of emissions and removals to each activity in the first commitment period of the Kyoto Protocol is shown in Table 2-14. For detailed information, see Chapter 11.

Table 2-14 Accounting summary for activities under Articles 3.3 and 3.4 of the Kyoto Protocol (CRF Information Table)

GREENHOUSE GAS SOURCE AND SINK ACTIVITIES	BY	Net emissions/removals				Accounting Parameters	Accounting Quantity
		2008	2009	2010	Total		
		(Gg CO ₂ equivalent)					
A. Article 3.3 activities							
A.1. Afforestation and Reforestation							-1230.68
A.1.1. Units of land not harvested since the beginning of the commitment period		-389.54	-415.03	-426.11	-1,230.68		-1230.68
A.1.2. Units of land harvested since the beginning of the commitment period							
A.2. Deforestation		2,456.72	3,115.09	4,822.89	10,394.70		10394.70
B. Article 3.4 activities							
B.1. Forest Management (if elected)		-45,388.77	-49,005.55	-53,251.78	-147,646.10		-147646.10
3.3 offset						9,164.02	-9164.02
FM cap						238,333.33	-138482.08
B.2. Cropland Management (if elected)	NA	NA	NA	NA	NA	NA	NA
B.3. Grazing Land Management (if elected)	NA	NA	NA	NA	NA	NA	NA
B.4. Revegetation (if elected)	-77.78	-1081.76	-1112.34	-1130.14	-3324.24	-233.34	-3090.90

- ※ The net removals by FM after application of 3.3 offset are lower than the upper limit (13 Mt-C times 5 (238,333 Gg-CO₂)) given in the Appendix to decision 16/CMP.1.
- ※ Since the total anthropogenic GHG emissions by sources and removals by sinks in managed forests since 1990 are larger than the net source of emissions incurred under Article 3.3, the offset rule according to paragraph 10 of the Annex to decision 16/CMP.1 is applied to Japan.
- ※ Methodologies for estimation and accounting of Article 3.3 and 3.4 activities are continuously reviewed. The values in Table 2-14 are estimated by using the current methodologies, and are only reported but not accounted for in the 2012 submission since Japan elected accounting for the entire commitment period. The issuance of removal units from LULUCF activities under the Kyoto Protocol is to be performed at the end of the first commitment period.
- ※ The total values and results of summing up each figure are not always the same because of the difference in display digit.

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Chapter 3. Energy (CRF sector 1)

3.1. Overview of Sector

Emissions from the energy sector consist of two main categories: fuel combustion and fugitive emissions from fuels. Fuel combustion includes emissions released into the atmosphere when fossil fuels (e.g., coal, oil products, and natural gas) are combusted. Fugitive emissions are intentional or unintentional releases of gases from fossil fuels by anthropogenic activities.

In Japan, fossil fuels are used to produce energy for a wide variety of purposes (e.g., production, transportation, and consumption of energy products) and CO₂ (Carbon dioxide), CH₄ (Methane), N₂O (Nitrous Oxide), NO_x (Nitrogen Oxide), CO (Carbon Monoxide), and NMVOC (Non-Methane Volatile Organic Compounds) are emitted in the process.

In 2010, GHG emissions (CO₂, CH₄ and N₂O) from energy sector accounted to 1,145,612 Gg-CO₂ eq., and represented 91.1% of the Japan's total GHG emissions (excluding LULUCF). The emissions from energy sector had increased by 6.2% compare to 1990.

3.2. Fuel Combustion (1.A.)

This category covers GHG emissions from combustion of fossil fuels such as coal, oil, and natural gas, and incineration of waste for energy purposes and with energy recovery.¹

This section includes GHG emissions from five sources: Energy Industries (1.A.1)—emissions from power generation and heat supply; Manufacturing Industries and Construction (1.A.2)—emissions from manufacturing industry and construction; Transport (1.A.3)—emissions from aviation, railways, road transport and shipping; Other Sectors (1.A.4)—emissions from commercial/institutional, residential, and agriculture/forestry/fishing sources; and Other (1.A.5)—emissions from the other sector.

In FY 2010, emissions from fuel combustion were 1,145,203 Gg-CO₂ eq., and represented 91.0% of GHG of the Japan's total GHG emissions (excluding LULUCF). The emissions had increased by 6.4% compared to 1990.

GHG emissions from fuel combustion in FY 2010 had increased by 4.4% compared to FY 2009. The primary reason for the emission increase in FY 2010 as compared to FY 2009 was the recovery from the economic recession induced by the Global Financial Crisis of 2008. CO₂ emissions from the industries sector increased because of the higher levels of manufacturing. In addition, electric power demand increased due to the relatively high number of days on which extremes of hot or cold were experienced.

¹ These emissions from waste incineration had been reported in the waste sector in 2008 submissions, regardless of use as energy or energy recovery. However, to comply with ERT observations and the requirements of IPCC Guidelines, the emissions are reported in the energy sector since 2009 submissions.

Table 3-1 Trends in GHGs emissions from fuel combustion (1.A)

Gas	Item	Unit	1990	1995	2000	2005	2008	2009	2010		
CO ₂	1.A.1. Energy Industries	a. Public Electricity and Heat Production	Gg-CO ₂	297,074	315,399	330,863	378,921	394,714	357,104	380,094	
		b. Petroleum Refining	Gg-CO ₂	15,893	16,956	17,285	16,441	14,324	14,564	15,001	
		c. Manufacture of Solid Fuels and Other Energy Industries	Gg-CO ₂	11,286	12,592	9,426	10,677	11,225	14,228	11,001	
	1.A.2. Manufacturing Industries and Construction	a. Iron and Steel	Gg-CO ₂	149,600	141,862	150,776	152,741	143,269	134,610	151,872	
		b. Non-Ferrous Metals	Gg-CO ₂	6,092	4,770	3,042	2,634	2,333	2,120	2,096	
		c. Chemicals	Gg-CO ₂	64,736	74,806	67,216	58,650	53,325	52,549	53,617	
		d. Pulp, Paper and Print	Gg-CO ₂	25,825	29,449	29,035	26,552	22,843	21,239	20,323	
		e. Food Processing, Beverages and Tobacco	Gg-CO ₂	13,129	14,407	13,161	11,326	8,862	8,761	8,817	
		f. Other	Gg-CO ₂	111,929	105,245	113,547	119,326	104,987	99,698	105,884	
	1.A.3. Transport	a. Civil Aviation	Gg-CO ₂	7,162	10,278	10,677	10,799	10,277	9,781	9,193	
		b. Road Transportation	Gg-CO ₂	189,228	225,381	232,827	222,652	205,933	202,018	204,277	
		c. Railways	Gg-CO ₂	932	819	707	644	600	586	588	
		d. Navigation	Gg-CO ₂	13,731	14,687	14,865	12,915	11,288	10,383	10,885	
	1.A.4. Other Sectors	a. Commercial/Institutional	Gg-CO ₂	83,593	93,269	101,450	110,678	98,756	93,283	92,336	
		b. Residential	Gg-CO ₂	56,668	66,320	68,958	67,583	59,023	57,792	61,095	
		c. Agriculture/Forestry/Fisheries	Gg-CO ₂	21,380	19,526	16,207	15,158	10,657	10,425	10,472	
	1.A.5 Other	a. Stationary	Gg-CO ₂	NO	NO	NO	NO	NO	NO	NO	
		b. Mobile	Gg-CO ₂	NO	NO	NO	NO	NO	NO	NO	
		Total	Gg-CO ₂	1,068,260	1,145,769	1,180,044	1,217,696	1,152,418	1,089,142	1,137,551	
	CH ₄	1.A.1. Energy Industries	a. Public Electricity and Heat Production	Gg-CH ₄	1.35	1.55	1.95	1.66	1.84	1.76	1.90
			b. Petroleum Refining	Gg-CH ₄	0.05	0.06	0.07	0.07	0.06	0.06	0.07
c. Manufacture of Solid Fuels and Other Energy Industries			Gg-CH ₄	0.02	0.03	0.06	0.05	0.17	0.18	0.20	
1.A.2. Manufacturing Industries and Construction		a. Iron and Steel	Gg-CH ₄	4.59	4.22	4.49	3.95	3.88	4.09	4.55	
		b. Non-Ferrous Metals	Gg-CH ₄	0.29	0.25	0.20	0.16	0.15	0.13	0.13	
		c. Chemicals	Gg-CH ₄	0.23	0.28	0.25	0.24	0.22	0.22	0.23	
		d. Pulp, Paper and Print	Gg-CH ₄	1.10	1.08	1.11	1.39	1.70	1.68	1.79	
		e. Food Processing, Beverages and Tobacco	Gg-CH ₄	0.11	0.14	0.13	0.13	0.12	0.12	0.13	
		f. Other	Gg-CH ₄	10.60	14.88	10.57	10.83	11.41	11.07	10.93	
1.A.3. Transport		a. Civil Aviation	Gg-CH ₄	0.14	0.17	0.21	0.23	0.22	0.22	0.22	
		b. Road Transportation	Gg-CH ₄	12.70	13.11	12.54	9.81	7.79	7.31	6.70	
		c. Railways	Gg-CH ₄	0.06	0.05	0.05	0.04	0.03	0.03	0.03	
		d. Navigation	Gg-CH ₄	1.27	1.37	1.41	1.23	1.07	0.98	1.03	
1.A.4. Other Sectors		a. Commercial/Institutional	Gg-CH ₄	1.02	3.19	4.38	4.46	5.69	5.11	5.09	
		b. Residential	Gg-CH ₄	8.23	8.61	8.15	7.76	6.64	6.49	6.89	
		c. Agriculture/Forestry/Fisheries	Gg-CH ₄	0.63	0.45	0.32	0.28	0.22	0.21	0.21	
1.A.5 Other		a. Stationary	Gg-CH ₄	NO	NO	NO	NO	NO	NO	NO	
		b. Mobile	Gg-CH ₄	NO	NO	NO	NO	NO	NO	NO	
		Total	Gg-CH ₄	42.39	49.43	45.87	42.28	41.22	39.67	40.10	
			Gg-CO ₂ eq.	890	1,038	963	888	866	833	842	
N ₂ O		1.A.1. Energy Industries	a. Public Electricity and Heat Production	Gg-N ₂ O	2.88	4.40	5.30	6.62	6.61	6.33	6.24
	b. Petroleum Refining		Gg-N ₂ O	0.08	0.14	0.20	0.19	0.18	0.18	0.19	
	c. Manufacture of Solid Fuels and Other Energy Industries		Gg-N ₂ O	0.02	0.02	0.02	0.02	0.04	0.04	0.04	
	1.A.2. Manufacturing Industries and Construction	a. Iron and Steel	Gg-N ₂ O	1.08	1.31	1.32	1.16	1.14	1.08	1.04	
		b. Non-Ferrous Metals	Gg-N ₂ O	0.19	0.18	0.15	0.03	0.03	0.03	0.02	
		c. Chemicals	Gg-N ₂ O	0.58	1.06	1.05	0.91	0.86	0.84	0.87	
		d. Pulp, Paper and Print	Gg-N ₂ O	0.48	0.90	0.93	0.94	1.11	1.15	1.15	
		e. Food Processing, Beverages and Tobacco	Gg-N ₂ O	0.24	0.26	0.26	0.25	0.24	0.23	0.23	
		f. Other	Gg-N ₂ O	1.78	2.34	3.15	3.45	3.24	3.05	2.95	
	1.A.3. Transport	a. Civil Aviation	Gg-N ₂ O	0.23	0.30	0.34	0.35	0.33	0.32	0.30	
		b. Road Transportation	Gg-N ₂ O	12.59	13.96	13.76	9.74	8.29	7.78	7.31	
		c. Railways	Gg-N ₂ O	0.39	0.34	0.29	0.27	0.25	0.24	0.24	
		d. Navigation	Gg-N ₂ O	0.36	0.39	0.40	0.35	0.30	0.28	0.29	
	1.A.4. Other Sectors	a. Commercial/Institutional	Gg-N ₂ O	0.38	0.59	0.69	0.77	0.74	0.71	0.71	
		b. Residential	Gg-N ₂ O	0.29	0.33	0.34	0.33	0.27	0.26	0.28	
		c. Agriculture/Forestry/Fisheries	Gg-N ₂ O	0.21	0.20	0.14	0.13	0.10	0.09	0.10	
	1.A.5 Other	a. Stationary	Gg-N ₂ O	NO	NO	NO	NO	NO	NO	NO	
		b. Mobile	Gg-N ₂ O	NO	NO	NO	NO	NO	NO	NO	
		Total	Gg-N ₂ O	21.78	26.72	28.35	25.53	23.72	22.62	21.97	
			Gg-CO ₂ eq.	6,752	8,285	8,788	7,913	7,355	7,011	6,809	
	Total of all gases		Gg-CO ₂ eq.	1,075,901	1,155,092	1,189,795	1,226,497	1,160,638	1,096,986	1,145,203	

3.2.1. Energy Industries (1.A.1.)

a) Source/Sink Category Description

This source category provides methods estimating CO₂ emissions from Public Electricity and Heat Production (1.A.1.a), Petroleum Refining (1.A.1.b), and Manufacture of Solid Fuels and Other Energy Industries (1.A.1.c).

b) Methodological Issues

The estimation methods, activity data, emission factors, and other parameters used in the Energy Industry (1.A.1), Manufacturing Industry and Construction (1.A.2) and Other Sectors (1.A.4) are basically common. Therefore, the estimation method and data used for all of them is summarized in this section.

The estimation method for waste incineration with energy use and energy recovery is described in Chapter.8.

【CO₂】

● Estimation Method

Tier 1 Sectoral Approach has been used in accordance with the decision tree of the *GPG (2000)* (Page 2.10, Fig. 2.1) to calculate emissions. Country-specific emission factors are used for all types of fuel.

$$E = \sum_{ij} [(A_{ij} - N_{ij}) \times GCV_i \times 10^{-3} \times EF_i \times OF_i] \times 44/12$$

E	: CO ₂ emissions from fossil fuel combustion [t-CO ₂]
A	: Energy consumption [t, kl, 10 ³ ×m ³]
N	: Non-energy product use of fossil fuels [t, kl, 10 ³ ×m ³]
GCV	: Gross calorific value [MJ/kg, MJ/l, MJ/m ³]
EF	: Carbon content of the fuel [t-C/TJ]
OF	: Oxidation factor
i	: Type of energy
j	: Sector

The calories and emissions from waste incineration with energy recovery are reported in Fuel Combustion (1.A.) in accordance with the *1996 Revised IPCC Guidelines* and the *GPG (2000)*. The fuel type is classified as “Other fuels”.

Estimation method, emission factors and activity data for emission from waste incineration with energy recovery is same as those used in the waste incineration (6.C.) in accordance with the *1996 Revised IPCC Guidelines*. Please refer to Chapter 8 for further details on estimation methods.

● Emission Factors

➤ Carbon emission factors

The carbon content of fuels expressed as the unit of calorific value (Gross Calorific Value) was used for carbon emission factors. The emission factors are country-specific values except a part of fuels that applied the default value provided in the *2006 IPCC Guidelines*.

Table 3-2 Emission factors for fuel combustion in gross calorific value

Fuel		Unit	1990	1995	2000	2005	2008	2009	2010	References	
Coal	Steel Making Coal	tC/TJ	24.5	24.5	24.5	24.5	24.5	24.5	24.5	-	
	Coking Coal	tC/TJ	24.5	24.5	24.5	24.5	24.5	24.5	24.5	2006 IPCC Guidelines for National Greenhouse Gas Inventories	
	PCI Coal	tC/TJ	24.5	24.5	24.5	24.5	24.5	24.5	24.5	same as Coking Coal	
	Imported Steam Coal	tC/TJ	24.7	24.7	24.7	24.7	24.7	24.7	24.7	-	
	Imported Coal : for general use	tC/TJ	24.7	24.7	24.7	24.7	24.7	24.7	24.7	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Imported Coal : for power generation	tC/TJ	24.7	24.7	24.7	24.7	24.7	24.7	24.7	same as Imported Coal : for general use	
	Indigenous Steam Coal	tC/TJ	24.9	24.9	24.9	24.9	24.9	24.9	24.9	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Underground	tC/TJ	24.9	24.9	24.9	24.9	24.9	24.9	24.9	same as Indigenous Steam Coal	
	Open Pit	tC/TJ	24.9	24.9	24.9	24.9	24.9	24.9	24.9	same as Indigenous Steam Coal	
	Hard Coal, Anthracite & Lignite	tC/TJ	25.5	25.5	25.5	25.5	25.5	25.5	25.5	2006 IPCC Guidelines for National Greenhouse Gas Inventories	
Coal Products	Coke	tC/TJ	29.4	29.4	29.4	29.4	29.4	29.4	29.4	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Coal Tar	tC/TJ	20.9	20.9	20.9	20.9	20.9	20.9	20.9	2006 IPCC Guidelines for National Greenhouse Gas Inventories	
	Coal Briquette	tC/TJ	29.4	29.4	29.4	29.4	29.4	29.4	29.4	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Coke Oven Gas	tC/TJ	11.0	11.0	11.0	11.0	11.0	11.0	11.0	2006 IPCC Guidelines for National Greenhouse Gas Inventories	
	Blast Furnace Gas	tC/TJ	27.3	26.9	26.6	26.5	26.4	26.5	26.3	established with annually calculated value in order to keep carbon balance in blast furnace and L.D. converter	
	Converter Furnace Gas	tC/TJ	38.4	38.4	38.4	38.4	38.4	38.4	38.4	2006 IPCC Guidelines for National Greenhouse Gas Inventories	
	Crude Oil for Refinery	tC/TJ	18.7	18.7	18.7	18.7	18.7	18.7	18.7	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Crude Oil for Power Generation	tC/TJ	18.7	18.7	18.7	18.7	18.7	18.7	18.7	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Bituminous Mixture Fuel	tC/TJ	20.0	20.0	20.0	20.0	20.0	20.0	20.0	2006 IPCC Guidelines for National Greenhouse Gas Inventories	
	Natural Gas Liquid & Condensate	tC/TJ	18.4	18.4	18.4	18.4	18.4	18.4	18.4	GHGs Estimation Methods Committee Report (Ministry of the Environment, Committee for the Greenhouse Gases Emissions Estimation Methods)	
Oil	Slack Gasoline	tC/TJ	18.2	18.2	18.2	18.2	18.2	18.2	18.2	adopted the value of Naphtha	
	Slack Kerosene	tC/TJ	18.5	18.5	18.5	18.5	18.5	18.5	18.5	adopted the value of Kerosene	
	Slack Diesel Oil or Gas Oil	tC/TJ	18.7	18.7	18.7	18.7	18.7	18.7	18.7	adopted the value of Diesel Oil or Gas Oil	
	Slack Fuel Oil	tC/TJ	19.5	19.5	19.5	19.5	19.5	19.5	19.5	adopted the value of Heating Oil C	
	Cracked Gasoline	tC/TJ	18.2	18.2	18.2	18.2	18.2	18.2	18.2	adopted the value of Naphtha	
	Cracked Diesel Oil or Gas Oil	tC/TJ	18.7	18.7	18.7	18.7	18.7	18.7	18.7	adopted the value of Diesel Oil or Gas Oil	
	Feedstock Oil for Refinery and Mixing	tC/TJ	18.7	18.7	18.7	18.7	18.7	18.7	18.7	adopted the value of Crude Oil for Refinery	
	Naphtha	tC/TJ	18.2	18.2	18.2	18.2	18.2	18.2	18.2	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Reformed Material Oil	tC/TJ	18.3	18.3	18.3	18.3	18.3	18.3	18.3	adopted the value of Gasoline	
	Gasoline	tC/TJ	18.3	18.3	18.3	18.3	18.3	18.3	18.3	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Premium Gasoline	tC/TJ	18.3	18.3	18.3	18.3	18.3	18.3	18.3	same as Gasoline	
	Regular Gasoline	tC/TJ	18.3	18.3	18.3	18.3	18.3	18.3	18.3	same as Gasoline	
	Jet Fuel	tC/TJ	18.3	18.3	18.3	18.3	18.3	18.3	18.3	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Kerosene	tC/TJ	18.5	18.5	18.5	18.5	18.5	18.5	18.5	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Gas Oil or Diesel Oil	tC/TJ	18.7	18.7	18.7	18.7	18.7	18.7	18.7	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Fuel Oil A	tC/TJ	18.9	18.9	18.9	18.9	18.9	18.9	18.9	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Fuel Oil C	tC/TJ	19.5	19.5	19.5	19.5	19.5	19.5	19.5	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Fuel Oil B	tC/TJ	19.2	19.2	19.2	19.2	19.2	19.2	19.2	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Fuel Oil C	tC/TJ	19.5	19.5	19.5	19.5	19.5	19.5	19.5	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Fuel Oil C for Power Generation	tC/TJ	19.5	19.5	19.5	19.5	19.5	19.5	19.5	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Lubricating Oil	tC/TJ	19.2	19.2	19.2	19.2	19.2	19.2	19.2	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Asphalt	tC/TJ	20.8	20.8	20.8	20.8	20.8	20.8	20.8	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Non Asphalt Heavy Oil Products	tC/TJ	20.8	20.8	20.8	20.8	20.8	20.8	20.8	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Oil Coke	tC/TJ	25.4	25.4	25.4	25.4	25.4	25.4	25.4	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Galvanic Furnace Gas	tC/TJ	38.4	38.4	38.4	38.4	38.4	38.4	38.4	adopted the value of Converter Furnace Gas	
	Refinery Gas	tC/TJ	14.2	14.2	14.2	14.2	14.2	14.2	14.2	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
	Liquefied Petroleum Gas	tC/TJ	16.3	16.3	16.3	16.1	16.1	16.1	16.1	GHGs Estimation Methods Committee Report (Ministry of the Environment, Committee for the Greenhouse Gases Emissions Estimation Methods)	
	Natural Gas	Liquefied Natural Gas	tC/TJ	13.5	13.5	13.5	13.5	13.5	13.5	13.5	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan
		Indigenous Natural Gas	tC/TJ	13.9	13.9	13.9	13.9	13.9	13.9	13.9	2006 IPCC Guidelines for National Greenhouse Gas Inventories
		Indigenous Natural Gas	tC/TJ	13.9	13.9	13.9	13.9	13.9	13.9	13.9	adopted the value of Indigenous Natural Gas
Coal Mining Gas		tC/TJ	13.5	13.5	13.5	13.5	13.5	13.5	13.5	Environmental Agency, The Report on Estimation of CO ₂ Emissions in Japan	
Off-gas from Crude Oil		tC/TJ	13.9	13.9	13.9	13.9	13.9	13.9	13.9	adopted the value of Indigenous Natural Gas	
Town Gas	Town Gas	tC/TJ	14.0	14.0	13.8	13.6	13.7	13.6	13.7	same as Town Gas	
	Town Gas	tC/TJ	14.0	14.0	13.8	13.6	13.7	13.6	13.7	established with annually calculated value in order to keep carbon balance in prodeced town gas	
	Small Scale Town Gas	tC/TJ	16.3	16.3	16.3	16.1	16.1	16.1	16.1	adopted the value of LPG	

Emission factors were developed based on three different concepts; (a) Energy sources other than Blast Furnace Gas (BFG) and Town gas, (b) BFG, and (c) Town gas.

Table 3-2 provides the emission factors for CO₂ by fuel types.

(a) Energy sources other than Blast Furnace Gas (BFG) and Town gas

Carbon emission factors of energy sources other than Blast Furnace Gas (BFG) and Town gas were used values provided in “The Report on Estimation of CO₂ Emissions in Japan (Environmental Agency, 1992)”, “GHGs Estimation Methods Committee Report (Committee for the Greenhouse Gases Emissions Estimation Methods, The Ministry of Environment)” and “2006 IPCC Guidelines”.

The result of evaluation in *Evaluating and Analyzing the Validity of Carbon Emission Factors for Different Fuels* (Kainou, 2005) were adopted for setting emission factors. In the choice of carbon emission factors, adequacy assessment was conducted for emission factors in *the Report on Estimation of CO₂ Emissions in Japan* (Environmental Agency, 1992), which were used in the inventories submitted up to 2005. These were assessed based on the following three criteria, and the values assessed as adequate continue to be used in this inventory.

- 1) Evaluation and analysis by comparison of theoretical upper and lower limits
- 2) Evaluation and analysis by comparison with the *Revised 1996 IPCC Guidelines* default values
- 3) Group evaluation and analysis by carbon balance using the *General Energy Statistics*

Summaries of evaluations were indicated below.

1) Evaluation and analysis by comparison of theoretical upper and lower limits

The validity of carbon emission factors is evaluated to compare intended emission factor and emission factor calculated by theoretical from standard enthalpy change of formation of pure matter, such as hydrogen, methane and carbon monoxide, because most of the fuels required to evaluate carbon emission factors are hydrocarbons containing a few impurities, and because a physicochemical correspondence exists between the standard gross calorific values of pure hydrocarbons and carbon emission factors.

2) Evaluation and analysis by comparison with the *Revised 1996 IPCC Guidelines* default values

The validity of carbon emission factors is judged by using the *Revised 1996 IPCC Guidelines* default values or the *2006 IPCC Guidelines* reference values² and their statistical reliability (uncertainty) information. However, because the average properties of fuels envisaged in the IPCC Guidelines and those of fuels used in Japan are not necessarily the same, carbon emission factors can be appropriately judged based on statistical examination of group evaluation and analysis mentioned below even when figures deviate, as long as a valid reason for the deviation exists.

² When *Evaluating and Analyzing the Validity of Carbon Emission Factors for Different Fuels* was submitted, the *2006 IPCC Guidelines* was not submitted. These values were reference values, some of these reference values were revised.

3) Group evaluation and analysis by carbon balance using the *General Energy Statistics*

The validity of fuel-specific carbon emission factors for some petroleum product and coal product factor groups can be evaluated using the *General Energy Statistics* to analyze carbon balance in coal products and oil products.

The values assessed as inadequate were substituted by the values given in the *GHGs Estimation Methods Committee Report* (Committee for the Greenhouse gases Emissions Estimation Methods, Ministry of the Environment) and the *2006 IPCC Guidelines*.

(b) Blast Furnace Gas (BFG)

During iron and steel production process, in the blast furnace and converter furnace, the amount of energy and carbon contained in coke and PCI coal which are injected to the processes and these contained in BFG and CFG which are calculated should be theoretically balanced. Since the composition of BFG is unstable, emission factors for BFG was established with annually calculated value in order to keep carbon balance in blast furnace and converter furnace during the iron and steel production process.

Emission factor for BFG was established with annually calculated value in order to keep carbon balance in blast furnace and converter furnace during iron and steel production process. The amount of carbon excluded carbon contained in CFG from carbon (contained in 'Coke' and 'PCI coal') injected to blast furnace indicated under 'Steel process gas' is considered to be carbon contained in BFG. Emission factor for BFG was established as carbon described above divided by calorific values of BFG generated. The equation for emission factor and the overview of carbon flow for iron & steel and calculation process are shown below.

Calculation to establish emission factor for BFG is conducted every year.

$$EF_{BFG} = [(A_{coal} \times EF_{coal} + A_{coke} \times EF_{coke}) - A_{CFG} \times EF_{CFG}] / A_{BFG}$$

EF	: Carbon emission factor [tC/TJ]
A	: Fuel consumption [TJ]
BFG	: Blast Furnace Gas
coal	: PCI coal
coke	: coke
CFG	: Converter Furnace Gas

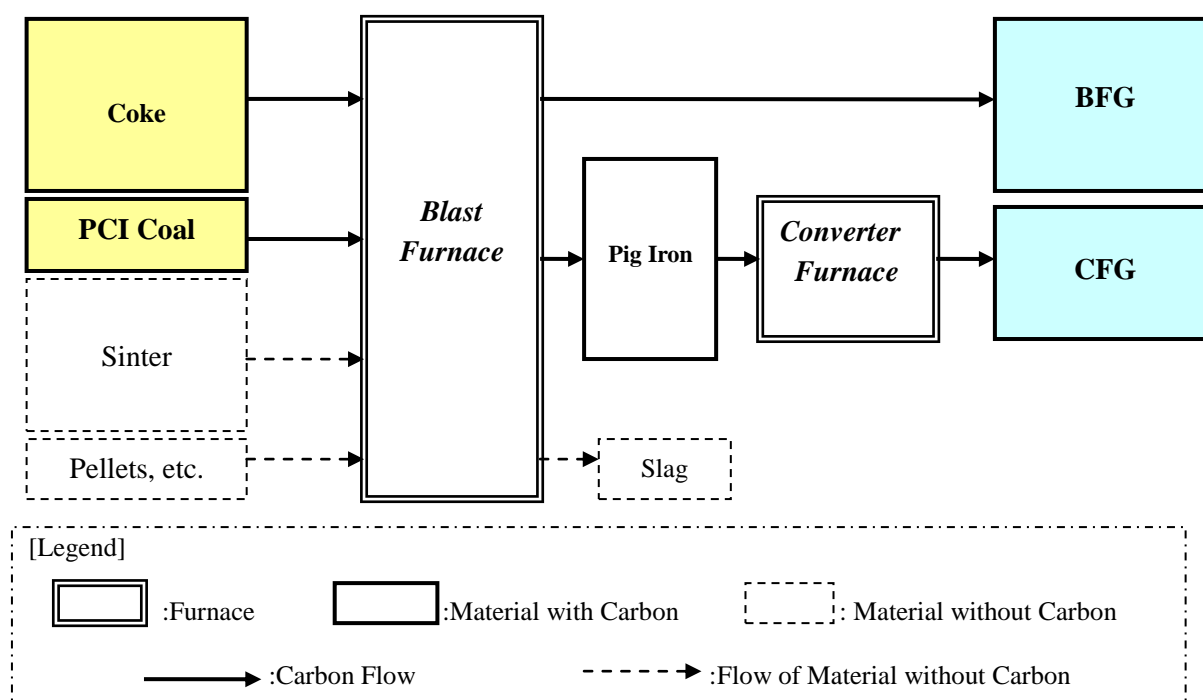


Figure 3-1 Manufacturing Flow for Coke, Coke Oven Gas and Blast Furnace Gas
(Overview of carbon flow for iron & steel)

Table 3-3 Calculation of Emission Factors for BFG

Steel Process Gas		1990	1995	2000	2005	2008	2009	2010	Note
Input									
PCI Coal	Gg-C	1,574	2,593	3,518	3,111	2,950	2,659	3,550	A
Coke	Gg-C	12,830	11,432	12,021	11,382	10,818	10,358	11,067	B
Input Total	Gg-C	14,404	14,024	15,539	14,492	13,768	13,017	14,616	C: A + B
Output									
CFG(LDG)	Gg-C	2,541	2,359	2,726	2,804	2,727	2,589	2,798	D
Difference	Gg-C	11,863	11,665	12,813	11,688	11,041	10,428	11,818	E: C - D
Output									
BFG	TJ	434,801	433,504	481,768	441,357	417,636	393,685	448,708	F
EF BFG	t-C/TJ	27.3	26.9	26.6	26.5	26.4	26.5	26.3	E / F

(c) Town gas

'Town gas' consists of 'Town gas' provided by town gas supplier and 'Small scale town gas' provided by small scale town gas supplier.

In the case of small scale town gas supplier:

Because most part of small scale town gas is LPG, the same emission factor for LPG was adopted for small scale town gas

In the case of town gas supplier:

Town gas is produced from the mixture of raw materials and air dilution. In order to calculate town gas emission factors, total carbon contained in fossil fuel used as raw materials was divided by the total calorific value of produced town gas. Emission factors for town gas were established based on carbon balance in 'Town gas production'. To calculate town gas emission factors, the

total carbon in fossil fuel inputs used as raw materials (COG, Kerosene, Refinery gas, LPG, LNG and Indigenous natural gas) was divided by the total calorific value of the town gas production. Calculation to establish emission factor for town gas is conducted every year.

$$EF_{TG} = \sum (A_i \times EF_i) / P_{TG}$$

- EF : Carbon emission factor [tC/TJ]
 A : Fuel consumption [TJ]
 P : Calorific value of the town gas production [TJ]
 TG : Town gas
 i : Feedstocks (COG, Kerosene, Refinery gas, LPG, LNG, Indigenous natural gas)

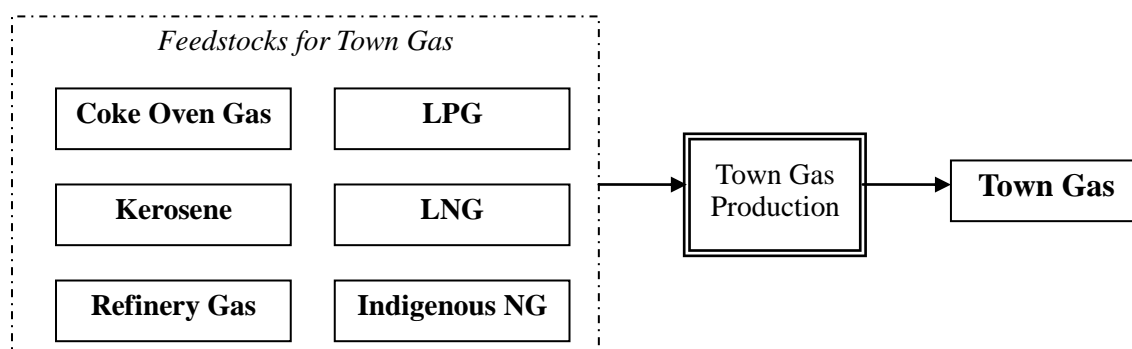


Figure 3-2 Manufacturing Flow for Town Gas

Table 3-4 Calculation of Emission Factors for Town Gas

Town Gas Production		1990	1995	2000	2005	2008	2009	2010	Note
Input									
COG	Gg-C	211	134	105	22	0	0	0	a1
Kerosene	Gg-C	200	275	69	6	0	0	0	a2
Refinery Gas	Gg-C	186	199	186	145	88	13	0	a3
LPG	Gg-C	1,931	2,104	1,791	1,069	679	700	782	a4
LNG	Gg-C	6,253	9,107	11,642	16,563	19,378	19,181	20,943	a5
Indigenous NG	Gg-C	551	661	848	1,190	1,822	1,768	1,603	a6
Input Total	Gg-C	9,331	12,480	14,641	18,994	21,967	21,663	23,328	A: Σ a
Output									
Town Gas	TJ	664,661	892,307	1,061,122	1,391,962	1,607,991	1,593,032	1,697,063	B
EF Town Gas	t-C/TJ	14.0	14.0	13.8	13.6	13.7	13.6	13.7	A/B

➤ Oxidation factor

For each type of energy, country-specific oxidation factors were established considering the actual conditions of fuel combustion in Japan based on survey on related industrial groups, manufacturing corporations and experts.

Gaseous Fuels

Every result of measurement of soot concentration of boiler to generate powers in 2004 for gaseous fuels combustion shows that no soot was emitted; therefore, it is considered that gaseous fuels are completely combusted. The results of questionnaires also show that gaseous fuels are completely combusted. Hence, oxidation factor for gaseous fuels combustion was set to 1.0.

Table 3-5 Data of gaseous fuel combustion

Fired condition	Provider	Survey
Complete combustion	The Federation for Electric Power Companies Japan (FEPC)	measurement of soot concentration of boiler to generate powers in 2004

Liquid Fuels (Petroleum Fuels)

Carbon contained in liquid fuel is considered to be almost completely combusted; however, unburned fuel loss, about 0.5%, may occur depending on its fired condition. Because the data of actual measurement was not available, considering meticulous combustion management and smoke treatment in Japan, oxidation factor for liquid fuels combustion was set to 1.0.

Solid Fuels

Oxidation factor for solid fuels varies depending on fired condition, type of furnace, and coal property; therefore, it is quite difficult to obtain representational data set of actual measurement of unburned fuel loss. Meanwhile, almost all the unburned carbon generated during combustion in furnace is considered to be contained in coal ash. Coal ash is effectively utilized or landfilled. Carbon contained in coal ash which is used as raw material of cement is oxidized to CO₂ and emitted into the atmosphere during calcinations processes.

Average oxidation factor from 1990 to 2003 considering unburned carbon oxidized in firing process of coal ash eventually is 0.996, expressed as 3 significant digits. 2 significant digits are considered to be adequate in the view of other coefficients' accuracy; therefore, oxidation factor for solid fuels is set to 1.0 rounding off to two significant digits.

● *Activity Data*

The data given in the *General Energy Statistics* compiled by the Agency for Natural Resources and Energy were used for the activity data. The *General Energy Statistics* (Energy Balance Table) provides a comprehensive overview of domestic energy supply and demand to grasp what are converted from energy sources, such as coal, oil, natural gas and others, provided in Japan and what are consumed in what sectors. The objective of this *General Energy Statistics* is to help to quantitatively understand energy supply and demand and to make judgments about the situation, in addition to helping with planning for energy and environmental policy, and with measuring, assessing, and otherwise gauging policy effectiveness.

General Energy Statistics (Energy Balance Table) indicates an overview of domestic energy supply and demand, shows the main energy sources used in Japan as “Columns” and the supply, conversion and consumption sectors as “Rows”, in a matrix. Specifically, columns comprise 11 major categories (coal, coal products, oil, oil products, natural gas, town gas, new and renewable energy, large-scale hydropower, nuclear power, electricity, and heat) and the necessary sub-categories and a more detailed breakdown of the sub-categories. Rows comprise 3 major sectors — primary energy supply (primary supply), energy conversion (conversion), and final energy consumption (final consumption) — plus the necessary sub-categories and a more detailed breakdown of the sub-categories.

In calculating the energy supply and demand amounts for *General Energy Statistics*, it is assumed that each energy source, such as gasoline or electricity, is homogeneous in terms of gross calorific value per original unit (MJ/kg, MJ/L, MJ/m³), and that homogeneous energy sources are supplied, converted, and consumed. Values for supply, conversion, and consumption in original units as determined from

official statistical sources are multiplied by gross calorific value per original unit to obtain energy supply and demand amounts.

The calculation process in the *General Energy Statistics* is as follows:

- (1) Set calorific values and carbon emission factors.
- (2) Build energy supply and demand modules.
- (3) Prepare original unit tables (integrate modules and prepare main table and summary table) (units in t, kl, m³, etc).
- (4) Prepare energy unit tables (Units are J).
- (5) Prepare energy-derived carbon tables (given are carbon content).

General Energy Statistics adopts “actual calorific values” based on calculation based on annual official statistics for some fuel types which can be recalculated. For other fuel types which cannot be recalculated and whose composition is stable, “standard calorific values” based on relevant official statistics and document are adopted.

The complete Energy Balance Tables for the years since FY 1990 are available on the following internet site:

<http://www.enecho.meti.go.jp/info/statistics/jukyu/result-2.htm> (Japanese version only)

Please refer to the simplified energy balance tables provided in Annex 2.

For the activity data for energy industries, the data reported in the following sectors in the *General Energy Statistics* were used: “Power Generation, General Electric Utilities” [#2110, codes in bracket indicate column and row number indicated in the *Interpretation of General Energy Statistics*] which reports energy consumption associated with electric power generation by electric power suppliers, and “Power Generation, Independent Power Producing” [#2150]; “District Heat Supply” [#2350] which provides energy consumption associated with heat energy and cold energy by thermal energy suppliers; “Own use, General Electric Utilities” [#2911] which reports energy consumption associated with captive (own) use of energy industries; “Own use, Independent Power Producing” [#2912]; “Own use, District Heat Supply” [#2913]; “Own use, Oil Refinery” [#2916]; “Own use, Town Gas” [#2914]; “Own Use, Steel Coke” [#2915]; and “Own use, Other Conversion” [#2917] (Numbers in parentheses indicate corresponding sector numbers in the *General Energy Statistics*).

Table 3-6 shows the correspondence between sectors of Japan’s Energy Balance Table from the *General Energy Statistics* and those of the CRF.

Table 3-6 Correspondence between sectors of Japan's Energy Balance Table and of the CRF (1.A.1)

CRF		Japan's Energy Balance Table	
1A1	Energy Industries		
1A1a	Public Electricity and Heat Production	Power Generation, General Electric Utilities	#2110
		Own use, General Electric Utilities	#2911
		Power Generation, Independent Power Producing	#2150
		Own use, Independent Power Producing	#2912
		District Heat Supply	#2350
		Own use, District Heat Supply	#2913
1A1b	Petroleum Refining	Own use, Oil Refinery	#2916
1A1c	Manufacture of Solid Fuels and Other Energy Industries	Coal Products	#2500
		Own use, Town Gas	#2914
		Own use, Steel Coke	#2915
		Own use, Other Conversion	#2917

➤ *Gross calorific value*

Gross calorific values used in Japan's Energy Balance Table (*General Energy Statistics*) are adopted. Table 3-7 shows trends in gross calorific value for each fuel type. Japan's Energy Balance Table (*General Energy Statistics*) is adopting actual calorific values based on calculation based on annual official statistics for some fuel types which can be recalculated. For other fuel types which cannot be recalculated and whose composition is stable, "standard calorific values" based on relevant official statistics and documents are adopted. The "standard calorific value" is revised approximately once in every 5 years.

Table 3-7 Trends in gross calorific value of each fuel type

Fuel		Code	Unit	1990	1995	2000	2005	2008	2009	2010	
Coal	Steel Making Coal	\$110	MJ/kg	31.8	31.8	28.9	29.0	29.0	29.0	29.0	
	Coking Coal	\$111	MJ/kg	31.8	30.5	29.1	29.1	29.1	29.1	29.1	
	PCI Coal	\$112	MJ/kg	31.8	30.5	28.2	28.2	28.2	28.2	28.2	
	Imported Steam Coal	\$130	MJ/kg	26.0	26.0	26.6	25.7	25.7	25.7	25.7	
	Imported Coal : for general use	\$131	MJ/kg	26.0	26.0	26.6	25.7	25.7	25.7	25.7	
	Imported Coal : for power generation	\$132	MJ/kg	24.9	26.1	26.4	25.5	25.3	25.4	25.3	
	Indigenous Steam Coal	\$135	MJ/kg	24.3	24.3	22.5	22.5	22.5	22.5	22.5	
	Underground	\$136	MJ/kg	24.3	24.3	23.2	23.2	23.2	23.2	23.2	
	Open Pit	\$137	MJ/kg	18.7	18.7	18.7	18.7	18.7	18.7	18.7	
Hard Coal, Anthracite & Lignite	\$140	MJ/kg	27.2	27.2	27.2	26.9	26.9	26.9	26.9		
Coal Products	Coke	\$161	MJ/kg	30.1	30.1	30.1	29.4	29.4	29.4	29.4	
	Coal Tar	\$162	MJ/kg	37.3	37.3	37.3	37.3	37.3	37.3	37.3	
	Coal Briquette	\$163	MJ/kg	23.9	23.9	23.9	23.9	23.9	23.9	23.9	
	Coke Oven Gas	\$171	MJ/m ³ N	21.5	21.6	21.3	21.4	21.2	21.1	21.3	
	Blast Furnace Gas	\$172	MJ/m ³ N	3.5	3.6	3.6	3.4	3.4	3.4	3.4	
	Converter Furnace Gas	\$173	MJ/m ³ N	8.4	8.4	8.4	8.4	8.4	8.4	8.4	
Oil	Crude Oil for Refinery	\$210	MJ/l	38.3	38.3	38.2	38.1	38.2	38.1	38.2	
	Crude Oil for Power Generation	\$220	MJ/l	39.1	39.2	39.6	38.5	39.5	39.7	39.7	
	Bituminous Mixture Fuel	\$221	MJ/kg	30.1	30.3	29.9	22.4	22.4	22.4	22.4	
	Natural Gas Liquid & Condensate	\$230	MJ/l	35.7	35.5	35.4	35.0	32.9	34.8	34.8	
Oil Products	Slack Gasoline	\$271	MJ/l	33.6	33.6	33.6	33.5	33.5	33.5	33.5	
	Slack Kerosene	\$272	MJ/l	36.8	36.8	36.8	36.7	36.7	36.7	36.7	
	Slack Diesel Oil or Gas Oil	\$273	MJ/l	38.6	38.6	38.6	38.6	38.6	38.6	38.6	
	Slack Fuel Oil	\$274	MJ/l	41.8	41.8	41.8	41.8	41.8	41.8	41.8	
	Cracked Gasoline	\$275	MJ/l	33.6	33.6	33.6	33.5	33.5	33.5	33.5	
	Cracked Diesel Oil or Gas Oil	\$276	MJ/l	38.6	38.6	38.6	38.6	38.6	38.6	38.6	
	Feedstock Oil for Refinery and Mixing	\$277	MJ/l	38.3	38.3	38.2	38.1	38.2	38.1	38.2	
	Naphtha	\$281	MJ/l	33.6	33.6	33.6	33.5	33.5	33.5	33.5	
	Reformed Material Oil	\$282	MJ/l	35.1	35.1	35.1	35.1	35.1	35.1	35.1	
	Gasoline	\$310	MJ/l	34.6	34.6	34.6	34.6	34.6	34.6	34.6	
	Premium Gasoline	\$311	MJ/l	35.1	35.1	35.1	35.1	35.1	35.1	35.1	
	Regular Gasoline	\$312	MJ/l	34.5	34.5	34.5	34.5	34.5	34.5	34.5	
	Jet Fuel	\$320	MJ/l	36.4	36.4	36.7	36.7	36.7	36.7	36.7	
	Kerosene	\$330	MJ/l	36.8	36.8	36.8	36.7	36.7	36.7	36.7	
	Gas Oil or Diesel Oil	\$340	MJ/l	38.1	38.1	38.2	37.8	37.9	37.9	38.1	
	Fuel Oil A	\$351	MJ/l	39.7	39.6	39.3	39.1	39.9	39.9	39.9	
	Fuel Oil C	\$355	MJ/l	42.7	42.2	42.0	42.0	42.2	42.0	42.1	
	Fuel Oil B	\$356	MJ/l	40.2	40.2	40.4	40.4	40.4	40.4	40.4	
	Fuel Oil C	\$357	MJ/l	42.7	42.2	42.0	42.0	42.2	42.0	42.1	
	Fuel Oil C for Power Generation	\$358	MJ/l	41.1	41.1	41.3	41.2	41.2	41.2	41.3	
	Lubricating Oil	\$365	MJ/l	40.2	40.2	40.2	40.2	40.2	40.2	40.2	
	Asphalt	\$371	MJ/kg	41.6	41.2	40.9	41.0	41.1	41.0	41.0	
	Non Asphalt Heavy Oil Products	\$372	MJ/kg	41.6	41.2	40.9	41.0	41.1	41.0	41.0	
	Oil Coke	\$375	MJ/kg	35.6	35.6	35.6	29.9	29.9	29.9	29.9	
	Galvanic Furnace Gas	\$376	MJ/m ³ N	8.4	8.4	8.4	8.4	8.4	8.4	8.4	
	Refinery Gas	\$380	MJ/m ³ N	39.3	39.3	44.9	44.9	44.9	44.9	44.9	
	Liquified Petroleum Gas	\$390	MJ/kg	50.2	50.2	50.2	50.8	50.8	50.8	50.8	
	Natural Gas	Liquefied Natural Gas	\$410	MJ/kg	54.6	54.6	54.6	54.6	54.6	54.6	54.6
		Indigenous Natural Gas	\$420	MJ/m ³ N	42.1	42.4	42.6	42.9	44.7	44.8	44.7
Indigenous Natural Gas		\$421	MJ/m ³ N	42.1	42.4	42.6	42.9	44.7	44.8	44.7	
Coal Mining Gas		\$422	MJ/m ³ N	36.0	36.0	16.7	16.7	16.7	16.7	16.7	
Off-gas from Crude Oil		\$423	MJ/m ³ N	42.1	42.4	42.6	42.9	44.7	44.8	44.7	
Town Gas	Town Gas	\$450	MJ/m ³ N	41.9	41.9	41.1	44.8	44.8	44.8	44.8	
	Town Gas	\$460	MJ/m ³ N	41.9	41.9	41.1	44.8	44.8	44.8	44.8	
	Small Scale Town Gas	\$470	MJ/m ³ N	100.5	100.5	100.5	100.5	100.5	100.5	100.5	

【CH₄, N₂O】**● Estimation Method**

Because it is possible to use fuel-specific, sector-specific and furnace-specific activity data, and also to set country-specific emission factors, CH₄ and N₂O emissions from fuel combustion in this category is calculated by using Tier 2 country-specific emission factors in accordance with the *1996 Revised IPCC Guidelines* and *GPG (2000)*. However, in residential and other sectors in which activity data for different furnace types cannot be used, Tier 1 IPCC default emission factors were used.

Estimation equation is as follows. Emissions were calculated by multiplying fuel-specific, furnace-specific and sector-specific activity data by fuel-specific and furnace-specific emission factors.

$$E = \sum (EF_{ij} \times A_{ijk})$$

E	: Emissions from combustion of fuel by stationary sources (kgCH ₄ , kgN ₂ O)
EF _{ij}	: Emission factor for fuel type i, furnace type j (kgCH ₄ /TJ, kgN ₂ O/TJ)
A _{ijk}	: Fuel consumption for fuel type i, furnace type j, sector k (TJ)
i	: Fuel type
j	: Furnace type
k	: Sector

● Emission Factors

Based on data obtained from surveys conducted in Japan (Table 3-9), chimney flue CH₄, N₂O and O₂ concentrations, and the theoretical (dry) exhaust gas volumes, theoretical air volumes, and higher heating values (gross calorific values) shown in Table 3-8 were employed to establish emission factors for each kind of facility using the following combustion calculation formula.

$$EF = C_{CH_4, N_2O} \times \{G_0' + (m-1) \times A_0\} \times MW \div V_m \div GCV$$

EF	: emission factor [kgCH ₄ /TJ, kgN ₂ O/TJ]
C _{CH₄ or N₂O}	: CH ₄ or N ₂ O concentration in exhaust gas [ppm]
G ₀ '	: theoretical exhaust gas volume for each fuel combustion (dry) [m ³ N/ original unit]
A ₀	: theoretical air volume for each fuel combustion [m ³ N/ original unit]
m	: air ratio ≡ actual air volume/ theoretical air volume (-)
MW	: molecular weight of CH ₄ (constant)=16 [g/mol] molecular weight of N ₂ O(constant)=44 [g/mol]
V _m	: one mole ideal gas volume in standardized condition (constant)=22.4 [10 ⁻³ m ³ /mol]
GCV	: gross calorific value for each fuel combustion [MJ/ original unit]

However, air ratio “m” is approximately provided with O₂ concentration in exhaust gas, as the equation below.

$$m = \frac{21}{21 - C_{O_2}}$$

C_{O₂} : O₂ concentration in exhaust gas (%)

CH₄ and N₂O emission factors by each fuel and furnace types were averaged after dividing emission factor of each kind of facilities according to fuel and furnace types (Table 3-10, Table 3-11). Anomalous values were excluded according to t-testing or expert opinion when calculating average values.

For CH₄ and N₂O emissions from electric arc furnaces, combustion calculation was carried out using measurement results for CH₄ and N₂O concentrations in exhaust gas, dry exhaust gas volume per unit time, and calorific value per unit time.

Table 3-8 Theoretical exhaust gas and air volumes, higher heating value for different fuels

Fuel type	Original unit	Theoretical exhaust gas volume (dry)	Higher heating value	Theoretical air volume	Remarks
		m ³ N/l,kg,m ³ N	kJ/l,kg,m ³ N,kWh	m ³ N/l,kg,m ³ N	
Fuel oil A	l	8.900	39,100	9.500	1
Fuel oil B	l	9.300	40,400	9.900	1
Fuel oil C	l	9.500	41,700	10.100	1
Diesel oil	l	8.800	38,200	9.400	1
Kerosene	l	8.400	36,700	9.100	1
Crude oil	l	8.747	38,200	9.340	1
Naphtha	l	7.550	34,100	8.400	1
Other liquid fuels	l	9.288	37,850	9.687	2
Other liquid fuels (heavy)	l	9.064	37,674	9.453	2
Other liquid fuels (light)	l	9.419	35,761	9.824	2
Steam coal	kg	7.210	26,600	7.800	1
Coke	kg	7.220	30,100	7.300	1
Harvested wood	kg	3.450	14,367	3.720	2
Charcoal	kg	7.600	30,500	7.730	3
Other solid fuels	kg	7.000	33,141	7.000	2
Town gas	m ³	9.850	46,047	10.949	2
Coke oven gas (COG)	m ³	4.500	21,100	4.800	1
Blast furnace gas (BFG)	m ³	1.460	3,410	0.626	1
Liquefied natural gas (LNG)	kg	11.766	54,500	13.093	1
Liquefied petroleum gas (LPG)	kg	11.051	50,200	12.045	1
Converter furnace gas (CFG) (Linz-Donawitz gas : LDG)	m ³	2.200	8,410	1.500	1
Refinery gas (offgas)	m ³	11.200	44,900	12.400	1
Other gaseous fuels	m ³	4.587	28,465	4.096	2
Other gaseous fuels (petroleum)	m ³	7.889	40,307	7.045	2
Other gaseous fuels (steel)	m ³	2.812	19,097	2.511	2
Other gaseous fuels (mining)	m ³	3.396	38,177	3.032	2
Other gaseous fuels (other)	m ³	4.839	23,400	4.321	2
Pulping waste liquor	kg	3.245	13,898	3.499	2
Electricity	kWh		3,600		1

Note 1: Theoretical exhaust gas and air volumes are the standard values given in the Ministry of the Environment's *General Survey of the Emissions of Air Pollutants*, except for town gas, LNG, and LPG, for which values calculated from constituent data were used. For town gas, the constituents of town gas 13A were considered to be representative. Regarding higher heating value, standard calorific values given in the *General Energy Statistics* were used for items marked 1, and standard values given in the *General Survey of the Emissions of Air Pollutants* (based on the 1992 survey) for items marked 2 in the Remarks column. The higher heating value for steam coal (imported) was used for the higher heating value of steam coal. The item marked 3 in the Remarks column was set by the 2005 Committee based on reference materials etc.

Table 3-9 References for measurement data used in establishment of emission factors

	References
1	Hokkaido Prefecture, <i>Report of GHG Emissions Intensity from Stationary Combustion</i> , 1991
2	Hyogo Prefecture, <i>Report of GHG Emissions Intensity from Stationary Combustion</i> , 1991
3	Osaka Prefecture, <i>Study of GHG Emissions Intensity from Stationary Combustion</i> , 1991
4	Hokkaido Prefecture, <i>Report of GHG Emissions Intensity from Stationary Combustion</i> , 1992
5	Hyogo Prefecture, <i>Report of GHG Emissions Intensity from Stationary Combustion</i> , 1992
6	City of Kitakyusyu, <i>Report of GHG Emissions Intensity from Stationary Combustion</i> , 1992
7	Hyogo Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1993
8	Hyogo Prefecture, <i>Report of GHG Emissions Intensity from Stationary Combustion</i> , 1994
9	Kanagawa Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1995
10	Niigata Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1995
11	Osaka Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1995
12	Hiroshima Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1995
13	Fukuoka Prefecture, <i>Report of GHG Emission Factors from Stationary Combustion</i> , 1995
14	City of Osaka, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1995
15	City of Kobe, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1995
16	Hokkaido Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1996
17	Ishikawa Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1996
18	Kyoto Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1996
19	Osaka Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1996
20	Hyogo Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1996
21	Hiroshima Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1996
22	Fukuoka Prefecture, <i>Report of GHG Emission Factors from Stationary Combustion</i> , 1996
23	Kyoto Prefecture, <i>Report of GHG Emission Factors from Stationary Combustion</i> , 1997
24	Hyogo Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1997
25	Fukuoka Prefecture, <i>Report of GHG Emission Factors from Stationary Combustion</i> , 1997
26	Japan Sociality Atmospheric Environment, <i>Reports on Greenhouse gas emissions estimation methodology</i> , 1996
27	Osaka Prefecture, <i>Study of GHG Emission Factors from Stationary Combustion</i> , 1999
28	Hyogo Prefecture, <i>Report of GHG Emission Factors from Stationary Combustion</i> , 2000
29	The Institute of Applied Energy, <i>Report for Trend of Fuel Quality in Lowering Environmental Atmospheric Quality</i> , 2000
30	Measurement Data prepared by Committee for the Greenhouse Gases Emissions Estimation Methods in FY1999
31	Data prepared by the Federation of Electric Power Companies of Japan
32	IPCC, <i>Revised 1996 IPCC Guidelines (Reference Manual)</i> , 1997

Table 3-10 CH₄ emission factors for different fuels and furnaces (unit: kg-CH₄/TJ)

Furnace type	Fuel type	Emission factor	Remarks
Boiler	Fuel oils B and C, crude oil	0.10	Average of 9 facilities
Boiler	Fuel oil A, diesel oil, kerosene, naphtha, other liquid fuels	0.26	Average of 2 facilities
Boiler	Gaseous fuel	0.23	Average of 5 facilities
Boiler	Steam coal, coke, other solid fuels	0.13	Average of 7 facilities
Boiler	Harvested wood, charcoal	75	Average of 4 facilities
Boiler	Pulping waste liquor	4.3	Average of 2 facilities
Sintering furnace for smelting of metals (except copper, lead, zinc)	Solid fuel, liquid fuel, gaseous fuel	31	Average of 6 facilities
Pelletizing furnace (steel and non-ferrous metal)	Solid fuel, liquid fuel, gaseous fuel	1.7	Average of 2 facilities
Metal rolling furnace, metal treating furnace, metal forging furnace	Liquid fuel, gaseous fuel	0.43	Average of 11 facilities
Petroleum and gas furnaces	Liquid fuel, gaseous fuel	0.16	Average of 27 facilities
Catalytic regenerator	Coke, carbon	0.054	Average of 11 facilities
Brick kiln, ceramic kiln, and other kiln	Solid fuel, liquid fuel, gaseous fuel	1.5	Average of 2 facilities
Aggregate drying kiln, cement raw material drying kiln, brick raw material drying kiln	Solid fuel, liquid fuel, gaseous fuel	29	Average of 6 facilities
Other drying kilns	Solid fuel, liquid fuel, gaseous fuel	6.6	Average of 8 facilities
Electric arc furnace	Electricity	13	Average of 6 facilities
Other industrial furnaces	Solid fuel	13	Average of 14 facilities
Other industrial furnaces	Liquid fuel	0.83	Average of 14 facilities
Other industrial furnaces	Gaseous fuel	2.3	Average of 6 facilities
Gas turbine	Liquid fuel, gaseous fuel	0.81	Average of 11 facilities
Diesel engine	Liquid fuel, gaseous fuel	0.70	Average of 8 facilities
Gas engine, petrol engine	Liquid fuel, gaseous fuel	54	Average of 6 facilities
Household equipment	Solid fuel	290	IPCC default value converted to higher heating value
Household equipment	Liquid fuel	9.5	IPCC default value converted to higher heating value
Household equipment	Gaseous fuel	4.5	IPCC default value converted to higher heating value
Household equipment	Biomass fuel	290	IPCC default value converted to higher heating value

Table 3-11 N₂O emission factors for different fuels and furnaces (unit: kg-N₂O/TJ)

Furnace type	Fuel type	Emission factor	Remarks
Boiler	Fuel oils B and C, crude oil	0.22	Average of 10 facilities
Boiler	Fuel oil A, diesel oil, kerosene, naphtha, other liquid fuels	0.19	Average of 2 facilities
Boiler	Gaseous fuel	0.17	Average of 5 facilities
Boiler (other than fluidized-bed boilers)	Solid fuel	0.85	Average of 9 facilities
Normal pressure fluidized-bed boiler	Solid fuel	54	Average of 11 facilities
Pressurized fluidized-bed boiler	Steam coal	5.2	Data from 1 facility
Boiler	Pulping waste liquor	0.17	Average of 2 facilities
Blast furnace	Coke oven gas, blast furnace gas, other gaseous fuel	0.047	Average of 2 facilities
Petroleum furnace, gas furnace	Liquid fuel, gaseous fuel	0.21	Average of 27 facilities
Catalytic regenerator	Coke, carbon	7.3	Average of 12 facilities
Electric arc furnace	Electricity	3.3	Average of 6 facilities
Coke oven	Town gas, coke oven gas, blast furnace gas, converter gas, offgas, other gaseous fuels	0.14	Average of 3 facilities
Other industrial furnace	Solid fuel	1.1	Average of 20 facilities
Other industrial furnace	Liquid fuel	1.8	Average of 31 facilities
Other industrial furnace	Gaseous fuel	1.2	Average of 18 facilities
Gas turbine	Liquid fuel, gaseous fuel	0.58	Average of 12 facilities
Diesel engine	Liquid fuel, gaseous fuel	2.2	Average of 9 facilities
Gas engine, petrol engine	Liquid fuel, gaseous fuel	0.85	Average of 7 facilities
Household equipment	Solid fuel	1.3	IPCC default value converted to higher heating value
Household equipment	Liquid fuel	0.57	IPCC default value converted to higher heating value
Household equipment	Gaseous fuel	0.090	IPCC default value converted to higher heating value
Household equipment	Biomass fuel	3.8	IPCC default value converted to higher heating value

● Activity Data

The data are estimated in the General Survey of the Emissions of Air Pollutants which provides details on fuel consumption for each type of furnaces and fuels, because stationary combustion fuel consumption data for the each type of furnaces are not available in the *General Energy Statistics*.

Fuel consumption by each sector (Energy Conversion, Industry, Commercial & Others, and Residential) for each type of fuels as presented in the *General Energy Statistics* was further divided among each furnace types proportionally to fuel consumption data in the General Survey of the Emissions of Air Pollutants to obtain the activity data for each sector, each fuel type and each furnace type. However, because the data in the General Survey of the Emissions of Air Pollutants does not differentiate between the pressurized fluidized-bed boiler, normal pressure fluidized-bed boiler, and other boilers, the fuel consumptions of these fluidized-bed boilers are calculated separately. Fuel consumption data of pressurized fluidized-bed furnace were provided by Federation of Electric Power Companies. Fuel consumption data of normal pressure fluidized-bed furnace were provided from companies which had past operation records of normal pressure fluidized-bed furnaces since 1990.

The data of solid fuel boilers excepted for fluidized-bed furnaces are estimated by subtracting the data of fluidized-bed furnace from the data of whole solid fuel boiler.

The exhaustive General Survey of the Emissions of Air Pollutants for all facilities emitting soot and smoke were carried out in fiscal 1992, 1995, 1996, and 1999. For years in which exhaustive General Survey of the Emissions of Air Pollutants were not carried out, the percentages of fuel consumption accounted for by each furnace type were interpolated using the data obtained in the years exhaustive survey carried out.

The procedure for calculating activity data is as follows:

- 1) Fuel consumption data from the General Survey of the Emissions of Air Pollutants is collated respectively for each fuel type, furnace type and sector.
- 2) The percentage of fuel consumption accounted for by each furnace type is calculated for each fuel type and sector.
- 3) Fuel consumption for different fuel types and sectors provided in the *General Energy Statistics* is multiplied by the percentage calculated in (2) to obtain fuel-specific, furnace-specific, and sector-specific activity data.

$$A_{ijk} = A_{EBik} \times W_{ijk}$$

- A_{ijk} : Activity data for fuel type i, furnace type j, sector k (TJ)
 A_{EBik} : Fuel consumption for fuel type i, sector k from *General Energy Statistics* (TJ)
 W_{ijk} : Ratio of furnace type j associated with consumption of fuel type i in sector k
 i : Fuel type
 j : Furnace type
 k : Sector

$$W_{ijk} = A_{MAPijk} / \sum_m A_{MAPimk}$$

- A_{MAPijk} : Fuel consumption for fuel type i, furnace type j, sector k according to General Survey of the Emissions of Air Pollutants (TJ)

- 4) The fuel-specific, furnace-specific, and sector-specific fuel consumption in the General Survey of the Emissions of Air Pollutants is used as activity data for the consumption of fuels (such as charcoal) not included in the *General Energy Statistics*, and furnaces for which *General Energy Statistics* fuel consumption data cannot be used (in specific terms, electricity consumption of electric arc furnaces and carbon fuels of catalytic regenerators).
- 5) In the residential sector, fuel consumption for different fuel types provided in the *General Energy Statistics* is used as activity data.

The N₂O emissions from solid fuel in 1.A.1.a (Public Electricity and Heat Production) increased between 1994 and 1995. The reason for the increase is that a new large sized fluidized-bed boiler for power generation went on line in 1995. As a result, the solid fuel consumption of fluidized-bed boiler for public power generation increased in 1995, resulting in an increase of N₂O emissions from solid fuel in this category.

➤ *Outline of the General Survey of the Emissions of Air Pollutants*

The General Survey of the Emissions of Air Pollutants is a statistical survey conducted to (1) promote reasonable and effective atmospheric environmental policy, (2) obtain information on current

activities within the context of the Air Pollutant Control Law (e.g., the current status of regulation of stationary sources that emit soot and smoke in facilities that are registered to a local government and in facilities that emit ordinary soot or particular soot, and the current status of air pollutant control), (3) develop the submitted data on facilities emitting soot and smoke, and (4) estimate the amounts of air pollutant emissions from facilities that emit soot and smoke. This survey is conducted with survey questionnaires. The response sheets and this survey's explanations are distributed to target facilities mentioned above.

c) Uncertainties and Time-series Consistency

● *Uncertainties*

【CO₂】

Carbon-Hydrogen ratio of hydrocarbons is strongly correlating with calorific value in theory, then, standard deviation of sample data of each fuel's calorific value are used for uncertainty assessment of emission factors based on assumption that deviation of carbon content and that of calorific value is equal. The uncertainty of energy consumption in TJ given in the *General Energy Statistics* was assessed based on the given statistical error of solid fuels, liquid fuels, and gaseous fuels. As a result, the uncertainty for emissions was determined to be 1% for CO₂ emissions from fuel combustion. A summary of uncertainty assessment methods is provided in Annex 7.

【CH₄, N₂O】

The uncertainties for emission factors were evaluated on the basis of applied statistical procedures, expert judgment, and default data for each energy type. The uncertainties of activity data were estimated by using standard deviation and the percentage of data collection indicated in General Survey of the Emissions of Air Pollutants. The uncertainties for emissions from fuel combustion were estimated to be 47% for CH₄ emissions and 33% for N₂O emissions. A summary of uncertainty assessment methods are provided in Annex 7.

● *Time-series Consistency*

The emissions were calculated in a consistent manner in all time series.

The carbon emission factors of all energy sources have been calculated by a consistent estimation method in all time series.

The emission factors for CH₄ and N₂O have been calculated by a consistent estimation method since FY 1990.

The activity data was used from data in *General Energy Statistics* in all time series, and the statistics are made by a consistent estimation method in all time series.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

GHG emissions in FY 2008 and FY 2009 were recalculated with the revision of the fuel consumption for FY 2008 and FY 2009 in the *General Energy Statistics*. Also, N₂O emissions from FY 1990 to FY 2009 were recalculated with the correction of the gross calorific value of steam coal burned in normal

pressure fluidized-bed boilers.

Updating the amount of incinerated municipal and industrial incinerated waste, the emission estimates for the period FY2005 and FY2007-2009 were recalculated. For details, see Section 8.4.2 of Chapter 8.

f) Source-specific Planned Improvements

Over 15 years have passed since the investigation for “The Report on Estimation of CO₂ Emissions in Japan” (Environment Agency) which is used as a data source of carbon emission factors of fuels was implemented. Then, the consideration on development of revised carbon emission factors of each fuel by direct measurement of carbon content and calorific value has been started with the cooperation of the Agency of Natural Resources and Energy.

The use of fuel consumption data in the General Survey of the Emissions of Air Pollutants for FY 2002 onward was prohibited for any purposes other than the original one specified for the General Survey of the Emissions of Air Pollutants, while that is not the case with the data in the General Survey of the Emissions of Air Pollutants for FY 1999 and earlier years. The use of General Survey of the Emissions of Air Pollutants in the GHG inventory was added to the purpose of the General Survey of the Emissions of Air Pollutants by the current examination toward the reuse of the General Survey of the Emissions of Air Pollutants and was recently officially accepted. Japan will keep consider applying the latest the General Survey of the Emissions of Air Pollutants data in the future inventory.

3.2.2. Manufacturing Industries and Construction (1.A.2)

a) Source/Sink Category Description

This category provides the estimation methods for determining CO₂ emissions from Iron and Steel (1.A.2.a); Non-ferrous Metals (1.A.2.b); Chemicals (1.A.2.c); Pulp, Paper, and Print (1.A.2.d); Food Processing, Beverages, and Tobacco (1.A.2.e); and Other (1.A.2.f).

b) Methodological Issues

● Estimation Method

See Section 3.2.1 b) (1.A.1).

● Emission Factors

See Section 3.2.1 b) (1.A.1).

● Activity Data

The data presented in *General Energy Statistics* were used for activity data, as was the case for the Energy Industry (1.A.1).

Activity data for manufacturing industry sectors were calculated by totaling energy consumption from production activities in factories and offices (final energy consumption), energy consumption related to non-utility power generation for use in one’s own factories and offices (non-utility power generation), and energy consumption related to steam production for use in own factories and offices (industrial steam) shown in *General Energy Statistics*. Because the energy consumption for production activities in factories and offices contained a certain amount used as raw materials (non-energy use), this amount was subtracted.

The non-utility power generation and industrial steam generation sectors are included in the energy conversion sector in *General Energy Statistics*. However, the *Revised 1996 IPCC Guidelines for National Greenhouse Gas Inventories* allocates CO₂ emissions from energy consumption for power or steam generation to the sectors generating that power or steam. As such, these CO₂ emissions are added to those from each industry in the final energy consumption sector and are provided in 1.A.2.

The IEF of CO₂ emissions from liquid fuels in 1.A.2.f (Other) decreases between 1997 and 1998, and increases between 1998 and 1999 because of revisions made to statistics on the manufacturing sector. The manufacturing sector data in Japan's Energy Balance Table (*General Energy Statistics*), the activity data, are based on the Ministry of Economy, Trade and Industry's *Yearbook of the Current Survey of Energy Consumption*. Subjects to be surveyed to obtain the data for the *Yearbook of the Current Survey of Energy Consumption* were changed in December, 1997. The survey for the industries of Dyeing, Rubber Product and Non-ferrous metal Product has been discontinued since 1998. Also, since 1998, business institutions or designated items to be surveyed for the industries of Chemicals, Cement & Ceramics, Glass Wares, Iron and Steel, Non-ferrous Metals and Machinery has been changed. For these reasons, and the IEF of CO₂ emissions from liquid fuels in 1.A.2.f (Other) changed. The details are documented and described in Annex.2.

Table 3-12 shows correspondence between sectors of Japan's Energy Balance Table and of the CRF (1.A.2).

Table 3-12 Correspondence between sectors of Japan's Energy Balance Table and of the CRF (1.A.2)

CRF		Japan's Energy Balance Table		
1A2	Manufacturing Industries and Construction			
	1A2a	Iron and Steel	Auto: Iron & Steel	#2217
			Steam Generation: Iron & Steel	#2307
			Final Energy Consumption, Iron & Steel	#6580
			Non-Energy, Iron & Steel	#9680
	1A2b	Non-Ferrous Metals	Auto: Non-Ferrous Metal	#2218
			Steam Generation: Non-Ferrous Metal	#2308
			Final Energy Consumption, Non-Ferrous Metal	#6590
			Non-Energy, Non-Ferrous Metal	#9690
	1A2c	Chemicals	Auto: Chemical Textiles	#2212
			Steam Generation: Chemical Textiles	#2302
			Final Energy Consumption, Chemical Textiles	#6530
			Non-Energy, Chemical Textiles	#9630
			Auto: Chemical	#2214
			Steam Generation: Chemical	#2304
			Final Energy Consumption, Chemical	#6550
			Non-Energy, Chemical	#9650
	1A2d	Pulp, Paper and Print	Auto: Pulp & Paper	#2211
			Steam Generation: Pulp & Paper	#2301
			Final Energy Consumption, Pulp & Paper	#6520
			Non-Energy, Pulp & Paper	#9620
	1A2e	Food Processing, Beverages and Tobacco	Final Energy Consumption, Food	#6510
			Non-Energy, Non-Manufacturing Industry (Food)	#9610
1A2f	Other			
	Mining	Final Energy Consumption, Mining	#6120	
		Non-Energy, Non-Manufacturing Industry (Mining)	#9610	
	Construction	Final Energy Consumption, Construction	#6150	
		Non-Energy, Non-Manufacturing Industry (Construction)	#9610	
	Oil Products	Auto: Oil products	#2213	
		Steam Generation: Oil products	#2303	
		Final Energy Consumption, Oil products	#6540	
		Non-Energy, Oil products	#9640	
	Glass Wares	Auto: Glass Wares	#2215	
		Steam Generation: Glass Wares	#2305	
		Final Energy Consumption, Glass Wares	#6560	
		Non-Energy, Glass Wares	#9660	
	Cement&Ceramics	Auto: Cement & Ceramics	#2216	
		Steam Generation: Cement & Ceramics	#2306	
		Final Energy Consumption, Cement & Ceramics	#6570	
		Non-Energy, Cement & Ceramics	#9670	
	Machinery	Auto: Machinery & Others	#2219	
		Steam Generation: Machinery & Others	#2309	
		Final Energy Consumption, Machinery	#6600	
		Non-Energy, Machinery	#9700	
	Duplication Adjustment	Auto: Duplication Adjustment	#2220	
		Steam Generation: Duplication Adjustment	#2310	
Final Energy Consumption, Duplication Adjustment		#6700		
Non-Energy, Duplication Adjustment		#9710		
Other Industries & SMEs	Auto: Others	#2250		
	Final Energy Consumption, Other Industries & SMEs	#6900		
	Non-Energy, Other Industries & SMEs	#9720		

- Auto: Non-utility power generation
- #9xxx items are not energy use activity.

c) Uncertainties and Time-series Consistency

See Section 3.2.1 c).

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

GHG emissions in FY 2009 were recalculated with the revision of the fuel consumption in FY 2009 in the *General Energy Statistics*. Also, N₂O emissions from FY 1990 to FY 2009 were recalculated with the correction of the gross calorific value of steam coal burned in normal pressure fluidized-bed boilers.

Updating the amount of biomass-based plastic products consumed, the emission estimates for the period FY2007-2009 were recalculated. Also, updating the amount of industrial waste used as raw material or fuel, the emission estimates for FY2008 were recalculated. For details, see Section 8.4.3 of Chapter 8.

f) Source-specific Planned Improvements

See Section 3.2.1 f)

3.2.3. Transport (Mobile Combustion) (1.A.3.:CO₂)**a) Source/Sink Category Description**

This category provides the methods used to estimate CO₂ emissions from Civil Aviation (1.A.3.a), Road Transportation (1.A.3.b), Railways (1.A.3.c), and Navigation (1.A.3.d).

b) Methodological Issues**● Estimation Method**

See Section 3.2.1 b).

Because CO₂ emissions from natural gas-powered vehicles and steam locomotives are included in Commercial /Institutional section of Other Sectors (1.A.4), CO₂ emissions from these source are reported as "IE."

● Emission Factors

See Section 3.2.1 b).

The carbon emission factor for liquid fuels (diesel oil) in 1.A.3.b (Road Transportation) is the lowest in Annex I Parties for two reasons. One is because the quality standard for diesel oil in Japan is different from other countries. Crude oil with high sulphur content imported from Middle East must be decomposed and go through ultradeep desulfurization to be low-sulphur diesel oil (<10ppm) according to Japanese automobile exhaust gas regulations. The other reasons is because gas oil used for purposes other than road transport is called "Fuel oil A" to distinguish it from diesel oil. The carbon balance of Japanese petroleum refineries including diesel oil and Fuel oil A nearly matches according to statistics, so these carbon emission factors are not irregular.

● Activity Data

The data given in the *General Energy Statistics* were used for activity data.

Values subtracting final energy consumption reported under 'Non-energy' [#9850] from energy consumption reported under 'Civil Aviation' [#8140] [#8540], 'Road Transportation' [#8110] [#8510] [#8115] [#8190] [#8590], 'Railways' [#8120] [#8520] and 'Navigation' [#8130] [#8530] in Japan's Energy Balance Table (*General Energy Statistics*) are used for activity data. Because energy consumption reported under 'Non-energy' was used for the purposes other than combustion and was considered not emitting CO₂, these values were deducted. (see Table 3-13)

Table 3-13 Correspondence between sectors of Japan's Energy Balance Table and of the CRF (1.A.3)

CRF		Japan's Energy Balance Table	
1A3	Transport		
1A3a	Civil Aviation	Final Energy Consumption, Passenger Air	#8140
		Final Energy Consumption, Freight Air	#8540
		Non-Energy, Transportation (Air)	#9850
1A3b	Road Transportation	Final Energy Consumption, Passenger Car	#8110
		Final Energy Consumption, Freight, Freight Truck & Lorry	#8510
		Final Energy Consumption, Passenger Bus	#8115
		Final Energy Consumption, Passenger, Transportation fraction estimation error	#8190
		Final Energy Consumption, Freight, Transportation fraction estimation error.	#8590
		Non-Energy, Transportation (Car, Truck & Lorry, Bus)	#9850
1A3c	Railways	Final Energy Consumption, Passenger Rail	#8120
		Final Energy Consumption, Freight Rail	#8520
		Non-Energy, Transportation (Rail)	#9850
1A3d	Navigation	Final Energy Consumption, Passenger Ship	#8130
		Final Energy Consumption, Freight Ship	#8530
		Non-Energy, Transportation (Ship)	#9850
1A3e	Other Transportation	-	-

• #9xxx items are not energy use activity.

c) Uncertainties and Time-series Consistency

See Section 3.2.1 c).

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

GHG emissions in FY 2007 and FY 2009 were recalculated due to the revision of the fuel consumption in FY 2007 and FY 2009 in the *General Energy Statistics*.

f) Source-specific Planned Improvements

There are no major planned improvements in this source category.

3.2.4. Transport (Mobile Combustion) (1.A.3.:CH₄, N₂O)

This section provides the estimation methods for CH₄ and N₂O emissions from Civil Aviation (1.A.3.a), Road Transportation (1.A.3.b), Railways (1.A.3.c), and Navigation (1.A.3.d).

3.2.4.1. Civil Aviation (1.A.3.a.)

a) Source/Sink Category Description

This section provides the estimation methods for CH₄ and N₂O emissions from energy consumption in civil aviation. Greenhouse gases associated with the domestic operation of Japanese airliners are mainly emitted from jet fuels. In addition, a small amount of aviation gasoline used by light aircraft and helicopters is also a source of CH₄ and N₂O emission.

b) Methodological Issues

● Estimation Method

Emissions have been calculated using the Tier 2a method for jet fuel and the Tier 1 for aviation gasoline, in accordance with Decision Tree of the *GPG (2000)* (Page 2.58, Fig. 2.7).

CH₄ and N₂O emissions associated with landing and take-off (LTO) of domestic airliners using jet fuel
= Emission factor per LTO 1 cycle per domestic airliner × Number of LTO cycles of aircraft in domestic routes

CH₄ and N₂O emissions from domestic airliner during cruising using jet fuel
= Emission factor associated with jet fuel consumption × Jet fuel consumption by aircraft during cruising in domestic routes

CH₄ and N₂O emission associated with flight of gasoline-powered domestic aircraft
= Emission factor associated with consumption of aviation gasoline × Consumption of aviation gasoline by aircraft in domestic routes

● Emission Factors

➤ Jet fuel

The default values given in the *Revised 1996 IPCC Guidelines* are used for emission factors for CH₄ and N₂O for LTO. The values used for emission factors for CH₄ and N₂O for cruising were calculated by converting the default values given in the *Revised 1996 IPCC Guidelines* into kg-CH₄/l using the specific gravity of jet fuel (0.78 t/kl). (See the following table)

➤ Aviation gasoline

The default values given in the *Revised 1996 IPCC Guidelines* are used for emission factors for CH₄ and N₂O (See the following table).

Table 3-14 CH₄ and N₂O emission factors for aircraft

		CH ₄	N ₂ O
jet aircraft (Jet fuel)	During takeoff and landing*	0.3 [kg-CH ₄ /LTO]	0.1 [kg-N ₂ O/LTO]
	During cruise	0 [kg-CH ₄ /kl]	0.078 [kg-N ₂ O/kl]
Other than jet aircraft (Aviation gasoline)	-	0.06 [g-CH ₄ /MJ]	0.0009 [g-N ₂ O/MJ]

* LTO=Landing/takeoff cycle

Source: Ministry of the Environment, *Results of Review of Greenhouse Gases Emissions Estimations Part 3* (August 2002). *Revised 1996 IPCC Guidelines*, Volume 3, Table 1-47

● Activity Data

➤ Jet fuel

The number of takeoffs and landings given in the *Statistical Yearbook of Air Transport* of the Ministry of Land, Infrastructure, Transport and Tourism is used as activity data at takeoff and landing. Fuel Consumption for takeoff and landing was calculated by multiplying fuel consumption for one takeoff or landing given in the IPCC guidelines, by the number of takeoffs and landings given above.

Fuel consumption for cruising was estimated by subtracting the amount of jet fuel consumed at takeoff and landing, from total jet fuel consumption calculated from the *Statistical Yearbook of Air Transport* of Ministry of Land, Infrastructure, Transport and Tourism.

➤ *Aviation gasoline*

Consumption of gasoline in airplane sector taken from the *General Energy Statistics* of the Agency for Natural Resources and Energy was used for activity data.

Table 3-15 Activity Data used for emission estimates of aircraft

Item	Unit	1990	1995	2000	2005	2008	2009	2010
number of LTO cycle	LTO	430,654	532,279	667,559	715,767	726,415	716,804	714,671
Jet fuel consumption of Cruise	kl	2,330,514	3,223,547	3,537,205	3,543,856	3,334,851	3,146,174	2,923,113
Gasoline consumption	kl	5,345	6,029	4,287	7,662	2,773	2,358	1,882

c) Uncertainties and Time-series Consistency

● **Uncertainties**

As the uncertainty of emission factors, default values given in the *GPG (2000)* (200% for CH₄ and 10,000% for N₂O) were applied. The uncertainty of activity data was 10%; determined as a standard value by the Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties for the emissions were determined to be 200% for CH₄ and 10,000% for N₂O. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

Emission factors were used same values since FY 1990. Activity data for jet fuel from the *Statistical Yearbook of Air Transport* and aviation gasoline from the *General Energy Statistics* have been used consistently since FY 1990.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.

e) Source-specific Recalculations

CH₄ and N₂O emissions in FY 2009 were recalculated due to the revision of the aviation gasoline consumption in FY 2009 in the *General Energy Statistics*.

f) Source-specific Planned Improvements

No improvements are planned.

3.2.4.2. Road Transportation (1.A.3.b.)

Emissions from automobiles in Japan are calculated for the following vehicle categories:

Table 3-16 Reporting categories and definitions of emissions from automobiles

Vehicle Type	Definition	Fuel type for emission reporting			
		Gasoline	Diesel	LPG	LNG
Light passenger vehicle	Light vehicle used for transportation of people.	○	—	—	—
Light cargo truck	Light vehicle used for transportation of cargo	○	—	—	—
Passenger vehicle	Regular passenger vehicle or small vehicle used for transportation of people, with a capacity of 10 persons or less.	○	○	○	—
Bus	Regular passenger vehicle or small vehicle used for transportation of people, with a capacity of 11 persons or more.	○	○	—	—
Small cargo truck	Small vehicle used for transportation of cargo.	○	○	—	—
Regular cargo truck	Regular vehicle used for transportation of cargo.	○	○	—	—
Special-purpose vehicle	Regular, small or light vehicle used for special purposes, including flushers, advertising vans, hearses, and others.	○	○	—	—
NPG vehicle	Any of the above vehicles that use natural gas as fuel.	—	—	—	○
Motorcycle	Two-wheeled vehicle	○	—	—	—

Different estimation methods are used for the categories of Light Passenger Vehicles, Light Cargo Trucks, Passenger Vehicles, Buses, Small Cargo Trucks, Regular Cargo Trucks, and Special-purpose Vehicles (3.2.4.2.a), Natural gas-powered Vehicles (3.2.4.2.b), and Motorcycles (3.2.4.2.c). Thus, they are described in the following sections.

3.2.4.2.a. Light Passenger Vehicles, Light Cargo Trucks, Passenger Vehicles, Buses, Small Cargo Trucks, Regular Cargo Trucks, and Special-purpose Vehicles

a) Source/Sink Category Description

This section provides the estimation methods for CH₄ and N₂O emissions from light passenger vehicles, light cargo trucks, passenger vehicles, buses, small cargo trucks, regular cargo trucks, and special-purpose vehicles.

b) Methodological Issues

● Estimation Method

Emissions have been calculated distance traveled per vehicle type by emission factors using the Tier 3 method, in accordance with Decision Tree of the *GPG (2000)* (Page 2.45, Fig. 2.5). The country-specific emission factors were used for some vehicle type, and the default emission factors were used for the other vehicle type. The activity data was estimated by using distance traveled and fuel efficiency which were provided from the Ministry of Land, Infrastructure, Transport and Tourism's *Statistical Yearbook of Motor Vehicle Transport*.

● Emission Factors

Emission factors for CH₄ and N₂O have been established for each type of fuel in each vehicle type, using the data shown in Table 3-17. "JAMA data" means that the raw emission factors of Japan Automobile Manufacturers Association are arranged as combine mode emission factors³ etc. by car

³ JAMA data were provided by test mode. The emission factors were calculated using "combined driving mode" mainly. "Combined driving mode" = "10.15 driving mode" × 0.88 + "11 driving mode" × 0.12. "10.15 driving mode" is

regulation year. The emission factors are estimated by multiplying arranged emission factors of JAMA by number of vehicles per car regulation year of each car classification (see Table 3-18, Table 3-19). “Measured data” means that the emission factor is based on actual Japanese data. The emission factors were weighted averages of actual Japanese data estimated per each class of running speed, by proportion of distance traveled per each class of running speed given in the Ministry of Land, Infrastructure, Transport and Tourism’s *Road Transport Census*. The emission factors reflect the actual motor vehicle operation in Japan because the proportion of distance traveled by each class of running speed during congestion was applied. “1996GL” and “GPG(2000)” mean the emission factors were established using the default values in IPCC guidelines.

Detailed method for the determination of the emission factors are described in the *Greenhouse Gases Estimation Methods Committee Report – Transportation* (Ministry of Environment; February, 2006).

Table 3-17 Data source of the emission factors of vehicle

Vehicle Type	Gasoline engine		Diesel engine	
	CH ₄	N ₂ O	CH ₄	N ₂ O
Light passenger vehicle	JAMA data	JAMA data		
Light cargo truck	JAMA data	JAMA data		
Passenger vehicle	JAMA data	JAMA data	JAMA data	JAMA data
Bus	1996GL	GPG(2000) +	Measured data	1996GL
Small cargo truck	JAMA data	JAMA data	JAMA data	JAMA data
Regular cargo truck	1996GL	GPG(2000) +	JAMA data	JAMA data
Special-purpose vehicle	1996GL	GPG(2000) +	Measured data	1996GL

JAMA data: Calculated by using driving mode test data provided by Japan Automobile Manufacturers Association

Measured data: Using actual Japanese data other than above JAMA data

1996GL: Using the default values in 1996 revised IPCC guidelines.

GPG(2000)+ : Calculated by using default data indicated in GPG (2000) in consideration of the fuel consumption by car type indicated in the *Statistical Yearbook of Motor Vehicle Transport* and calorific value indicated in the *General Energy Statistics*.

Table 3-18 CH₄ emission factors for road transportation

Fuel	Vehicle Type	Unit	1990	1995	2000	2005	2008	2009	2010
Gasoline	Light Passenger Vehicle	g-CH ₄ /km	0.008	0.008	0.008	0.007	0.006	0.006	0.005
	Passenger Vehicle (including LPG)	g-CH ₄ /km	0.015	0.015	0.014	0.011	0.009	0.009	0.008
	Light Cargo Truck	g-CH ₄ /km	0.020	0.020	0.019	0.013	0.009	0.009	0.008
	Small Cargo Truck	g-CH ₄ /km	0.022	0.021	0.021	0.015	0.011	0.011	0.010
	Regular Cargo Truck	g-CH ₄ /km	0.035	0.035	0.035	0.035	0.035	0.035	0.035
	Bus	g-CH ₄ /km	0.035	0.035	0.035	0.035	0.035	0.035	0.035
	Special Vehicle	g-CH ₄ /km	0.035	0.035	0.035	0.035	0.035	0.035	0.035
Diesel	Passenger Vehicle	g-CH ₄ /km	0.011	0.012	0.012	0.013	0.013	0.013	0.013
	Small Cargo Truck	g-CH ₄ /km	0.010	0.011	0.010	0.009	0.009	0.009	0.008
	Regular Cargo Truck	g-CH ₄ /km	0.017	0.016	0.015	0.014	0.013	0.012	0.012
	Bus	g-CH ₄ /km	0.019	0.018	0.017	0.017	0.017	0.017	0.017
	Special Vehicle	g-CH ₄ /km	0.017	0.015	0.013	0.013	0.013	0.013	0.013

a hot start driving mode and “11 driving mode” is a cold start driving mode.

Table 3-19 N₂O emission factors for road transportation

Fuel	Vehicle Type	Unit	1990	1995	2000	2005	2008	2009	2010
Gasoline	Light Passenger Vehicle	g-N ₂ O/km	0.015	0.015	0.014	0.009	0.007	0.006	0.005
	Passenger Vehicle (including LPG)	g-N ₂ O/km	0.024	0.024	0.020	0.012	0.009	0.007	0.006
	Light Cargo Truck	g-N ₂ O/km	0.024	0.024	0.022	0.013	0.009	0.008	0.008
	Small Cargo Truck	g-N ₂ O/km	0.020	0.021	0.021	0.013	0.009	0.008	0.008
	Regular Cargo Truck	g-N ₂ O/km	0.039	0.041	0.038	0.037	0.035	0.035	0.036
	Bus	g-N ₂ O/km	0.045	0.046	0.044	0.041	0.042	0.040	0.041
	Special Vehicle	g-N ₂ O/km	0.039	0.042	0.037	0.031	0.030	0.028	0.027
Diesel	Passenger Vehicle	g-N ₂ O/km	0.006	0.005	0.004	0.004	0.004	0.005	0.005
	Small Cargo Truck	g-N ₂ O/km	0.009	0.010	0.011	0.012	0.012	0.012	0.012
	Regular Cargo Truck	g-N ₂ O/km	0.015	0.015	0.015	0.017	0.028	0.030	0.032
	Bus	g-N ₂ O/km	0.025	0.025	0.025	0.025	0.025	0.025	0.025
	Special Vehicle	g-N ₂ O/km	0.025	0.025	0.025	0.025	0.025	0.025	0.025

● Activity Data

Estimates of annual distance traveled by each vehicle type and by each type of fuel have been used as activity data. The method of estimating activity data was to multiply the proportion of distance traveled for each fuel, which was calculated from fuel consumption and fuel efficiency, by the distance traveled for each vehicle type given in the Ministry of Land, Infrastructure, Transport and Tourism's *Statistical Yearbook of Motor Vehicle Transport*.

Table 3-20 Distance traveled per vehicle type

vehicle type	fuel type	Unit	1990	1995	2000	2005	2008	2009	2010
Light vehicle	Gasoline	10 ⁶ vehicle-km	15,281	39,386	70,055	102,601	121,327	128,585	129,695
Passenger vehicle	Gasoline	10 ⁶ vehicle-km	289,697	323,022	363,991	372,663	351,943	355,499	347,593
	Diesel Oil	10 ⁶ vehicle-km	42,252	66,787	58,832	30,902	17,692	14,879	12,791
	LPG	10 ⁶ vehicle-km	18,368	17,192	15,382	13,971	12,864	12,362	11,937
Bus	Gasoline	10 ⁶ vehicle-km	95	32	21	46	73	85	99
	Diesel Oil	10 ⁶ vehicle-km	7,016	6,736	6,598	6,605	6,503	6,464	6,492
Light cargo truck	Gasoline	10 ⁶ vehicle-km	85,336	84,534	74,914	73,789	73,312	72,382	70,690
Small cargo truck + Cargo passenger truck	Gasoline	10 ⁶ vehicle-km	36,981	25,892	24,988	26,597	26,345	26,054	26,410
	Diesel Oil	10 ⁶ vehicle-km	55,428	62,032	57,221	41,674	36,295	33,281	30,824
Regular cargo truck	Gasoline	10 ⁶ vehicle-km	447	361	331	741	1,059	1,088	1,156
	Diesel Oil	10 ⁶ vehicle-km	66,434	78,086	82,693	78,866	77,887	74,146	73,587
Special vehicle	Gasoline	10 ⁶ vehicle-km	827	851	1,584	1,556	1,726	1,822	1,822
	Diesel Oil	10 ⁶ vehicle-km	10,420	15,373	19,115	18,869	19,851	19,361	19,779

● N₂O emissions from gasoline vehicle in Japan

With the stipulation of the "1978 Emissions Regulation," the Three-way Catalyst started to be installed in gasoline automobiles in Japan. Then N₂O emissions per distance traveled increased. Until around 1986 when The Three-way Catalyst became widely used, N₂O emissions per distance traveled kept on increasing. New emission regulation on automobile were not stipulated until 1997, therefore, N₂O emissions per distance traveled were stable from 1986 to 1997. From 1997, Low Emission Vehicles were introduced. From 2000, with the stipulation of the "2000 Emission Regulation," N₂O emissions per distance traveled started to decrease in response to the introduction of the Close-coupled Catalytic Converter. Since 1997, the trend of N₂O emissions per distance traveled is on the decrease.

● Completeness

➤ Biomass fuels

Currently, since very little ethanol fuel exists in Japan, there are very few ethanol-powered vehicles. For that reason, the emissions of CH₄ and N₂O associated with the use of vehicles using biomass as fuel has been reported as "NO".

➤ Other (Methanol)

The number of methanol vehicles owned in Japan was only 19 at the end of March 2007 (data surveyed by the Ministry of Land, Infrastructure, Transport and Tourism). Therefore activity data is negligible, and has not been reported, as it is assumed that the emissions are also negligible.

c) Uncertainties and Time-series Consistency

● *Uncertainties*

As the uncertainty of emission factors for the CH₄ and N₂O emissions from all types of vehicles, default values given in the *GPG (2000)* (40% for CH₄ and 50% for N₂O) were applied. For the uncertainty for activity data, 50% for standard values determined by the Committee for the Greenhouse Gas Emission Estimation Methods was applied. As a result, the uncertainties of the emission from all road transportation including natural gas-powered vehicles and motorcycles were determined to be 64% for CH₄ and 71% for N₂O. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

Emission factors were developed by using same method since FY 1990. Activity data have been estimated using the data in the *Statistical Yearbook of Motor Vehicle Transport*, in a consistent estimation method from FY 1990 onward.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.

e) Source-specific Recalculations

For gasoline light vehicle, gasoline passenger vehicle, gasoline small cargo truck, gasoline light cargo truck, diesel small cargo truck, diesel regular cargo truck, and LPG passenger vehicle, new emission factors for CH₄ and N₂O in response to the enforcement of the New Long-term Regulation for exhaust gas (from FY 2005) were provided by JAMA, and emission factors for CH₄ and N₂O were revised. As a result, emissions for CH₄ and N₂O from FY 2005 to FY 2009 were recalculated.

f) Source-specific Planned Improvements

For some types of vehicles, it is necessary to consider whether more suitable emission factors, representing Japan's circumstances should be established on the basis of actual measurements, because the default values presented in the *Revised 1996 IPCC Guidelines* and *GPG (2000)* are currently used.

3.2.4.2.b. Natural gas-powered vehicles

a) Source/Sink Category Description

This section provides the estimation methods for CH₄ and N₂O emissions from natural gas-powered vehicles.

b) Methodological Issues

● *Estimation Method*

Emissions were calculated by multiplying the distance traveled per type of natural gas-powered vehicle by the emission factor for the vehicle type.

● Emission Factors

CH₄ emission factors for natural gas-powered small cargo trucks, passenger vehicle, light vehicle, light cargo trucks, regular cargo trucks and bus were determined using JAMA data and the same method used for the same type of gasoline or diesel powered vehicles.

N₂O emission factors for small cargo trucks and regular cargo trucks were determined using the emission factors established for each travel speed category, based on the actual measurements taken in Japan, and weighted by the percentage of distance traveled for each travel speed category reported in the *Road Transport Census* (Ministry of Land, Infrastructure, Transport and Tourism).

In the absence of actual measurement data in Japan, N₂O emission factors for passenger vehicle, light passenger vehicle, light cargo trucks, Special-purpose vehicles and bus and CH₄ emission factor for special-purpose vehicles were determined by the method indicated in the following Table 3-21.

Table 3-21 CH₄ and N₂O emission factors for natural gas-powered vehicles

Vehicle Type	Calculation Method for Emission Factor		Average Emission Factor	
	CH ₄	N ₂ O	CH ₄ [g-CH ₄ /km]	N ₂ O [g-N ₂ O/km]
Small cargo truck	JAMA data	Determined based on actual measurements	0.020	0.0002
Passenger vehicle	JAMA data	Used the emission factors for small cargo truck, taking the specifications of each vehicle type into account.	0.019	0.0002
light passenger vehicle, light cargo truck	JAMA data		0.013	
Regular cargo truck	JAMA data	Determined based on actual measurements	0.082	0.0128
Special-purpose vehicle	Determined from the percentage of distance traveled per travel speed category which was adjusted by the emission factor per travel speed category for regular cargo trucks, taking travel patterns of natural gas-powered special-purpose vehicles into consideration.		0.093	0.0145
Bus	JAMA data	Determined from the emission factor for regular cargo truck which was adjusted by the ratio of equivalent inertia weight, taking vehicle weight into consideration.	0.050	0.0384

● Activity Data

Annual distance traveled per vehicle type was determined by multiplying the number of natural gas-powered vehicles by the annual distance traveled per vehicle. The number of these vehicles was taken from the number of registered natural gas-powered vehicles per type in data compiled by the Japan Gas Association. For the annual distance traveled per vehicle type, the value specific to the natural gas-powered vehicles could not be determined. As a result, the calculation of activity data used the annual distance traveled per vehicle for all fuel types which had been determined from the distance traveled per vehicle type and the number of registered vehicles per type reported in the *Statistical Yearbook of Motor Vehicle Transport*.

Table 3-22 Annual distance traveled by natural gas-powered vehicles per vehicle type

vehicle type	Unit	1990	1995	2000	2005	2008	2009	2010
Passenger vehicle	1,000 vehicle-km	54	104	6,516	13,528	14,016	14,271	14,008
Bus	1,000 vehicle-km	0	1,860	18,743	53,936	64,005	65,079	65,956
Truck	1,000 vehicle-km	91	2,459	77,394	384,460	565,364	572,016	591,048
Small cargo truck	1,000 vehicle-km	184	8,088	32,426	57,045	72,550	75,529	78,680
Light vehicle	1,000 vehicle-km	0	301	12,934	49,543	69,299	74,951	79,106
Garbage vehicle	1,000 vehicle-km	0	300	6,955	38,816	50,304	52,287	54,472

c) *Uncertainties and Time-series Consistency*

● *Uncertainties*

The uncertainty of emission factors for both CH₄ and N₂O were determined as 1000% by expert judgment. The uncertainty of activity data was 50%; determined as a standard value by the 2002 Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties of the emissions were determined to be 1001% for both CH₄ and N₂O. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

The same emission factors were used since FY 1990. Activity data were estimated by using the data in the *Statistical Yearbook of Motor Vehicle Transport* and the *Natural Gas Mining Association Data*, in the same estimation method consistently since FY 1990.

d) *Source-specific QA/QC and Verification*

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.

e) *Source-specific Recalculations*

No recalculation was performed.

f) *Source-specific Planned Improvements*

To set more precise emission factors that better reflect actual conditions, it is needed to stock much more data on the annual distance traveled per vehicle type and improve the estimation methods used.

3.2.4.2.c. Motorcycles

a) *Source/Sink Category Description*

This section provides the estimation methods for CH₄ and N₂O emissions from motorcycles.

b) *Methodological Issues*

● *Estimation Method*

Emissions from motorcycles were estimated based on the method developed in Japan by the Ministry of Environment for the estimation of emissions from vehicles not subject to the PRTR⁴.System. The emissions were calculated for two emission sources of “Hot start” and “Increment at cold start”, using the equations below. For details of the calculation method, see the *Greenhouse Gases Estimation Methods Committee Report – Transportation* (February, 2006).

CH₄ and N₂O emissions from hot-starting of motorcycles

= Emission factor for vehicle-km per type of motorcycle × Total annual distance traveled by motorcycles per type

CH₄ emissions from increment at cold starting of motorcycles

= Emission factor per start per type × Number of engine start-ups per year by each type of motorcycle

⁴ Pollutant Release and Transfer Register

● Emission Factors

➤ Hot start

The THC (Total Hydro Carbon) emission factor for hot starts, derived from the actual measurement data in Japan, was multiplied by the ratio of the CH₄ emission factor to the THC emission factor, obtained from actual measurements. The THC emission factors for motorcycles were established for each vehicle type, stroke, and unregulated/regulated status. Accordingly, the emission factor per travel speed was determined for each type of motorcycle by apportioning the number of motorcycles in operation to these categories based on the estimated component ratio. For N₂O, the default emission factor for *US Motorcycles/European Motorcycles* given in the *Revised 1996 IPCC Guidelines* [0.002(gN₂O/km)] is used for unregulated status and JAMA data are used for regulated status (Integrated EFs are estimated from the ratio of each number of motorcycles for unregulated/regulated status).

➤ Increment at cold start

The emission factor was determined for each type of motorcycle by multiplying the THC emission factor for cold-start increment, derived from the actual measurement data in Japan, by the CH₄ and THC emission factors for hot start, and apportioning the results based on the ownership component ratio. No emission factor is set for N₂O because the increment at cold start for N₂O is assumed to be included in the default emission factor for hot start

Table 3-23 CH₄ and N₂O emission factors for motorcycles

Emission Source	Vehicle Type	Unit	1990	1995	2000	2005	2008	2009	2010
Two-wheel vehicle (hot start, CH ₄)	Small motor vehicle: first kind	g-CH ₄ /km	0.124	0.118	0.101	0.066	0.042	0.036	0.033
	Small motor vehicle: second kind	g-CH ₄ /km	0.088	0.090	0.082	0.051	0.030	0.028	0.025
	Light two-wheel vehicle	g-CH ₄ /km	0.155	0.159	0.137	0.069	0.043	0.037	0.033
	Small two-wheel vehicle	g-CH ₄ /km	0.117	0.119	0.112	0.069	0.046	0.041	0.034
Two-wheel vehicle (cold start, CH ₄)	Small motor vehicle: first kind	g-CH ₄ /number	0.039	0.039	0.033	0.022	0.018	0.017	0.016
	Small motor vehicle: second kind	g-CH ₄ /number	0.012	0.012	0.013	0.016	0.018	0.018	0.018
	Light two-wheel vehicle	g-CH ₄ /number	0.016	0.016	0.018	0.024	0.026	0.027	0.027
	Small two-wheel vehicle	g-CH ₄ /number	0.043	0.043	0.042	0.035	0.032	0.032	0.031
Two-wheel vehicle (hot start, N ₂ O)	Small motor vehicle: first kind	g-N ₂ O/km	0.002	0.002	0.002	0.002	0.002	0.002	0.002
	Small motor vehicle: second kind	g-N ₂ O/km	0.002	0.002	0.002	0.002	0.002	0.002	0.002
	Light two-wheel vehicle	g-N ₂ O/km	0.002	0.002	0.002	0.001	0.001	0.001	0.001
	Small two-wheel vehicle	g-N ₂ O/km	0.002	0.002	0.002	0.002	0.001	0.001	0.001

● Activity Data

➤ Hot start

Based on the motorcycle operation data in the *Road Transport Census*, annual distance traveled was determined for each type of motorcycle and travel speed category using the ratio of total distance traveled per type, obtained from sources including the *Survey of Motorcycle Market Trends* and the ratio of distance traveled per travel speed category, estimated from the *Road Transport Census*. In the determination of the activity data for this source, the rate of reduction of motorcycle operation due to rain or snow as well as increases in the ownership and the distance traveled during the years outside the survey were taken into consideration.

➤ Increment at cold start:

The annual number of engine startups (times/year) per type of motorcycle was determined by the following formula:

$$\begin{aligned} & \text{Number of engine startups} \\ & = (\text{Expected operation of new motorcycle in number of days in year})_{\text{type}} \times (\text{Operation factor})_{\text{elapsed years}} \times \\ & (\text{Reduction rate of operation due to rain and snow})_{\text{prefecture}} \times (\text{Average number of startups per day})_{\text{type}} \times (\text{Number} \\ & \text{of motorcycles owned})_{\text{type, prefecture, elapsed years}} \end{aligned}$$

c) *Uncertainties and Time-series Consistency*

● *Uncertainties*

As the uncertainty of emission factors, default values given in the *GPG (2000)* (40% for CH₄ and 50% for N₂O) were applied. The uncertainty of activity data was 50%; this was determined as a standard value by the 2002 Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties of the emissions were determined to be 64% for CH₄ and 71% for N₂O. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

Same Estimation Methods were used since FY 1990. Activity data were estimated using the data in the *Statistical Yearbook of Motor Vehicle Transport* in a consistent estimation method since FY 1990.

d) *Source-specific QA/QC and Verification*

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6. In addition, as a part of QC check, estimation files for motorcycles were improved.

e) *Source-specific Recalculations*

CH₄ and N₂O emissions of FY 2009 were recalculated due to the fact that statistics of number of motorcycles owned (The number of vehicle owned by Automobile Inspection and Registration Association and JAMA data) were revised. In addition, due to the improvement of the estimation files for motorcycles, the emissions for all years were recalculated.

f) *Source-specific Planned Improvements*

No improvements are planned.

3.2.4.3. Railways (1.A.3.c.)

a) *Source/Sink Category Description*

This section provides the estimation methods for CH₄ and N₂O emissions from railways. Emissions from railways come mainly from diesel-engine railway cars and locomotives that use diesel oil. In addition, there are small amounts of emissions from coal-fired steam locomotives.

b) *Methodological Issues*

● *Estimation Method*

Emissions were calculated by multiplying emission factor by fuel consumption on a calorific basis. The *GPG (2000)* does not provide a decision tree for a calculation method for this source.

CH₄ and N₂O emissions from diesel engines in railways

= Emission factor for diesel engines in railways × Annual consumption of diesel oil by diesel engines in railways

CH₄ and N₂O emissions from steam locomotives

= Emission factor for coal in rail transportation × Annual consumption of coal by steam locomotives

● *Emission Factors*

For emission factors for diesel-powered railway cars and , the default value shown in the *Revised 1996 IPCC Guidelines* under *Diesel engines – Railways* was used after the conversion to a per-liter value using the calorific value of diesel oil.

For emission factors for steam locomotives, the default value shown in the *Revised 1996 IPCC Guidelines* under *Coal – Railways* was used after the conversion to a per-weight value using the calorific value of imported steam coal.

Table 3-24 Default values for railway emission factors

	Diesel Engines	Steam Locomotives
CH ₄ emission factor	0.004 [g-CH ₄ /MJ]	10 [kg-CH ₄ /TJ]
N ₂ O emission factor	0.03 [g-N ₂ O/MJ]	1.4 [kg-N ₂ O/TJ]

Source: *Revised 1996 IPCC Guidelines*, Vol. 3, p. 1.91, Table 1-49; p. 1.35, Table 1-7; and p. 1.36, Table 1-8

● Activity Data

For the consumption of diesel oil by diesel engines in railways, diesel oil consumption in the railway sector shown in the *General Energy Statistics* compiled by the Agency for National Resources and Energy was used as the activity data.

Coal consumption by steam locomotives was considered to be the value shown in the *Statistical Yearbook of Railway Transport* (Ministry of Land, Infrastructure, Transport and Tourism) in the table “*Cost of Consumption of Operating Electricity, Fuel and Oil*” under *Cost under the Other fuel – Cost*. The cost-based value was divided by the coal price for each year (for imported steam coal) shown in the *Directory of Energy and Economic Statistics* to estimate the coal consumption.

Table 3-25 Activity Data used for estimation of emissions from railways

Fuel type	Unit	1990	1995	2000	2005	2008	2009	2010
Diesel oil	kl	356,224	313,235	269,711	248,211	230,381	224,972	224,972
Coal	kt	17	19	28	13	7	7	7

c) Uncertainties and Time-series Consistency

● Uncertainties

The uncertainties for emission factors were determined to be 5.0% for CH₄ and 5.0% for N₂O in accordance with the Committee for the Greenhouse Gas Emission Estimation Methods. For the uncertainty of activity data of diesel-engine locomotive, 10% given in the *Statistical Yearbook of Railway Transport*, was applied. For the uncertainty of activity data of coal-fired steam locomotives, 105% aggregated by the values given in the *Statistical Yearbook of Railway Transport* and the *Directory of Energy and Economics Statistics*, was applied. As a result, the uncertainties of the emissions were determined to be 11% for CH₄ and N₂O from diesel-engine locomotives and 101% for CH₄ and N₂O from coal-fired steam locomotives. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

The same emission factors were used since FY 1990. The data given in the *General Energy Statistics* for diesel-engine locomotives were used as activity data consistently since FY 1990. Activity data for coal-fired steam locomotives were calculated using the data in the *Statistical Yearbook of Railway Transport* and the *Directory of Energy and Economics Statistics*, in a consistent estimation method in all time-series.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.

e) Source-specific Recalculations

GHG emissions in FY 2007 and FY 2009 were recalculated with the revision of the diesel fuel consumption described in the *General Energy Statistics* in FY 2007 and FY 2009. Also, GHG emissions from FY 1990 to FY 2009 were recalculated with the correction of the GCV to NCV conversion rate of coal.

f) Source-specific Planned Improvements

For the emission factor for diesel engine-railways, it is needed to discuss whether more suitable emission factors (i.e., those that better reflect Japan's circumstances) should be established on the basis of actual measurements, because the default values presented in the *Revised 1996 IPCC Guidelines* and *GPG (2000)* are currently used.

3.2.4.4. Navigation (1.A.3.d.)**a) Source/Sink Category Description**

This section provides the estimation methods for CH₄ and N₂O emissions from navigation. Ships emit CH₄ and N₂O through the use of diesel oil and fuel oils A, B and C during their navigation.

b) Methodological Issues**● Estimation Method**

Emissions were calculated using the default values for CH₄ and N₂O given in the *Revised 1996 IPCC Guidelines*, in accordance with Decision Tree of the *GPG (2000)* (Page 2.52, Fig. 2.6).

<i>CH₄ and N₂O emissions associated with navigation of domestic vessels</i> = Emission factors for diesel oil and fuel oils A, B and C relating to domestic vessels × Consumption of each type of fuel by domestic vessels

● Emission Factors

The default values for Ocean-Going Ships (diesel engines) given in the *Revised 1996 IPCC Guidelines* (See the following table) were converted to emission factor per liter using the calorific value for each type of fuel (diesel oil, fuel oil A, B and C).

Table 3-26 Default emission factors for navigation

	Value
CH ₄ Emission Factor	0.007 [g-CH ₄ /MJ]
N ₂ O Emission Factor	0.002 [g-N ₂ O/MJ]

Source: *Revised 1996 IPCC Guidelines* Vol. 3, page 1.90, Table 1-48

● Activity Data

Consumption of each fuel type in internal navigation sector taken from the *General Energy Statistics* of the Agency for Natural Resources and Energy was used for activity data.

Table 3-27 Activity Data used for estimation of emissions from ships

Fuel type	Unit	1990	1995	2000	2005	2008	2009	2010
Diesel oil	1,000 kl	133	208	204	195	189	163	163
Fuel oil (A)	1,000 kl	1,602	1,625	1,728	1,324	1,046	946	1,021
Fuel oil (B)	1,000 kl	526	215	152	63	25	20	18
Fuel oil (C)	1,000 kl	2,446	3,002	3,055	2,873	2,592	2,420	2,516

c) *Uncertainties and Time-series Consistency*

● *Uncertainties*

As the uncertainty of emission factors, default values given in the *GPG (2000)* (200% for CH₄ and 1,000% for N₂O) were applied. The uncertainty of activity data was 13%. This was a precision value (95% confidence interval) provided in the *Statistical Yearbook of Coastwise Vessel Transport* that was an original statistic of the *General Energy Statistics*. As a result, the uncertainties of the emissions were determined to be 64% for CH₄ and 71% for N₂O. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

Emission factors were used same values since FY 1990. The activity data given in the *General Energy Statistics* were used as the activity data for navigation consistently since FY 1990.

d) *Source-specific QA/QC and Verification*

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.

e) *Source-specific Recalculations*

GHG emissions in FY 2009 were recalculated with the revision of the fuel consumption described in the *General Energy Statistics* in FY 2009. Also, GHG emissions from FY 1990 to FY 2009 were recalculated with the correction of the GCV to NCV conversion rate of fuel oil C.

f) *Source-specific Planned Improvements*

For the emission factor for navigation, it is needed to discuss to set more suitable factors (i.e., those that better reflect Japan's circumstances) that are based on actual measurements, because the default values presented in the *Revised 1996 IPCC Guidelines* are currently used.

3.2.5. Other Sectors (1.A.4)

a) *Source/Sink Category Description*

This category provides the estimation methods for CO₂ emissions from Commercial /Institutional (1.A.4.a), Residential (1.A.4.b) and Agriculture / Forestry / Fisheries (1.A.4.c).

b) *Methodological Issues*

● *Estimation Method*

See Section 3.2.1 b).

● *Emission Factors*

See Section 3.2.1 b).

● Activity Data

The data given in the *General Energy Statistics* compiled by the Agency for Natural Resources and Energy were used for activity data as well energy industry (1.A.1).

Activity data for each sub-category are the values for final energy consumption in Commercial/Institutional (#7500), Residential (#7100), and Agriculture/Forestry/Fisheries (#6110) sector in *General Energy Statistics*. Because the energy consumption above includes the amount of Non-energy use which was used for purposes other than combustion, these values were deducted from the energy consumption in each category.

Table 3-28 Correspondence between sectors of Japan's Energy Balance Table and of the CRF (1.A.4)

CRF		Japan's Energy Balance Table	
1A4	Other Sectors		
1A4a	Commercial/Institutional	Final Energy Consumption, Commercial & Others	#7500
		Non-Energy, ResCom & others (Commercial & Others)	#9800
1A4b	Residential	Final Energy Consumption, Residential	#7100
		Non-Energy, ResCom & others (Residential)	#9800
1A4c	Agriculture/Forestry/Fisheries	Final Energy Consumption, Agriculture, Forestry & Fishery	#6110
		Non-Energy, Non-Manufacturing Industry (Agriculture, Forestry & Fishery)	#9610

• #9xxx items are not energy use activity.

c) Uncertainties and Time-series Consistency

See Section 3.2.1 c).

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6

e) Source-specific Recalculations

GHG emissions in FY 2009 were recalculated due to the revision of the fuel consumption in FY 2009 in *General Energy Statistics*.

f) Source-specific Planned Improvements

There are no major planned improvements in this source category.

3.2.6. Distinctive trend

ERT recommend that Japan provide clear explanations of trends of such as GHG emissions, activities, and emission factors. In this section, these explanations were described.

The N₂O emissions from solid fuel in 1.A.1.a (Public Electricity and Heat Production) increased between 1994 and 1995. The reason for the increase is that a new large sized fluidized-bed boiler for power generation went on line in 1995. As a result, the solid fuel consumption of fluidized-bed boiler for public power generation increased in 1995, resulting in an increase of N₂O emissions from solid fuel in this category.

The IEFs (Implied Emission Factor) of CO₂ emissions from solid fuels in 1.A.1.c (Manufacture of Solid Fuels and Other Energy Industries) have been pulled up and down by “emission from carbon balances” derived from transformation of solid fuel by manufacture of solid fuels. The apparent annual change of this category is caused by the difference of mass-balance between Coking coal and Coke and other coal products, may be caused by statistical error, unobserved stockpiles in the process and/or spontaneous input-output unbalance.

Furthermore, the gross calorific value (GCV) trends for solid fuel are declining since 1990. In 1970 to 1990, Japanese steel manufacturers used conventional coking coal for feedstock for Coke, but due to the shortage of coking coal and price increase, they developed new Coke making technology to use steam coal with pre-treatment as feedstock for Coke instead. Similarly, they changed PCI coal from coking coal and steam coal mixture to steam coal with pre-treatment. Japanese steel manufacturers have been trying to make high-quality coke from cheap coal for economic reasons. Because conventional coking coal has a high carbon content and GCV than steam coal, and the new technology has introduced step-by-step, so apparent GCV gradually decreased in these years. For emission factors for fuel combustion in gross calorific value, refer to Table 3-2.

3.2.7. Comparison of Sectoral and Reference Approaches

This comparison is documented and described in Annex 4.

3.2.8. International Bunker Fuels

a) Source/Sink Category Description

This sector provides the estimation methods for determining CO₂, CH₄, and N₂O emissions from the fuel consumed for international navigation and aviation.

Exclusion of emissions from bunker fuels used for international navigation and aviation from the national totals has been reported as a memo item.

b) Methodological Issues

● *Estimation Method*

Emissions of CO₂, CH₄ and N₂O from this source are derived by multiplying the consumption of each type of fuel handled by bonds by the emission factor.

● *Emission Factors*

[CO₂]

The emission factors used for CO₂ are the same as those for the energy sectors, fuel combustion (CO₂) in energy sectors (Refer to Section 3.2.1).

[CH₄, N₂O]

Default values given in the *Revised 1996 IPCC Guidelines* are used for CH₄ and N₂O emission factors.

Table 3-29 Emission factors for CH₄ and N₂O from international bunkers

Transport mode	Type of fuel	CH ₄ emission factor	N ₂ O emission factor
Aircraft	Jet fuel	0.002 [g-CH ₄ /MJ] ^a	0.1 [kg-N ₂ O/t] ^b
Shipping	Fuel oil A, Fuel oil B, Fuel oil C, Diesel oil, Kerosene	0.007 [g-CH ₄ /MJ] ^c	0.002 [g-N ₂ O/MJ] ^c

a. Revised 1996 IPCC Guidelines Vol. 3, Table 1-47

b. Revised 1996 IPCC Guidelines Vol. 3, Table 1-52

c. Revised 1996 IPCC Guidelines Vol. 3, Table 1-48

● Activity Data

Totals for bonded imports and bonded exports given in the Ministry of Economy, Trade and Industry's Yearbook of Mineral Resources and Petroleum Products Statistics (former Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke) are used for emissions of CO₂, CH₄, and N₂O from the relevant source.

A and B in the diagram below correspond to the items under bonded exports and bonded imports, respectively, in the Yearbook of Mineral Resources and Petroleum Products Statistics (former Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke). C equals to the sum of A and B and it is used as the activity data for this source of emissions. This is considered to be approximately equivalent to the amount of the fuels sold in Japan for the international aviation and navigation.

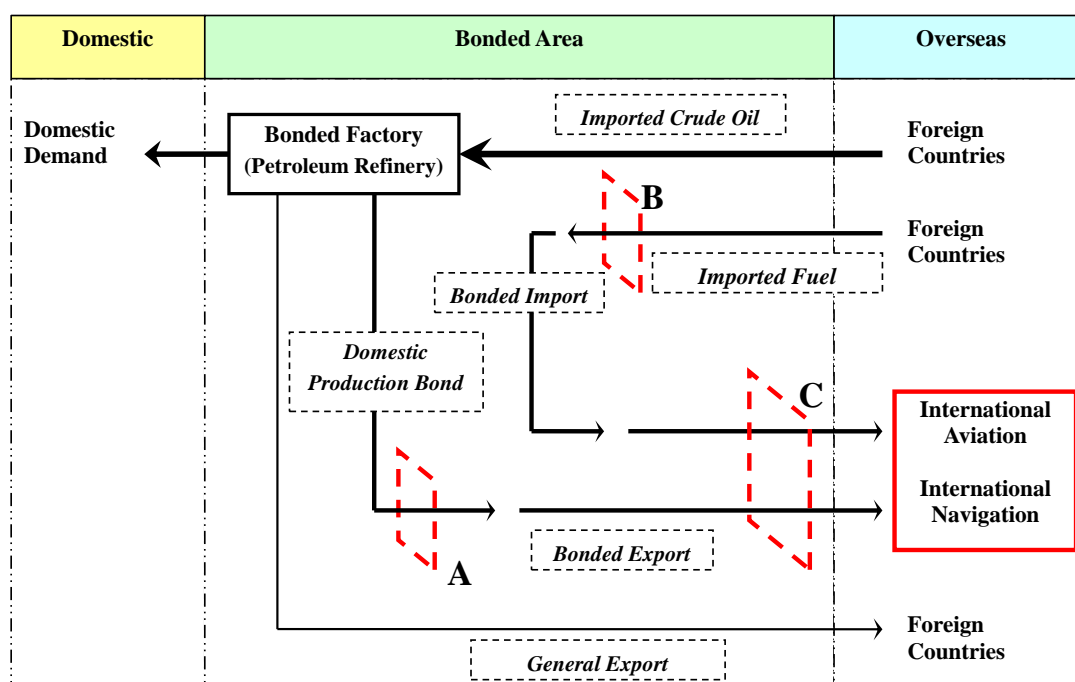


Figure 3-3 Activity data for international bunkers

It is assumed that jet fuel is used by aircraft, while fuel oil A, B, C, diesel oil and kerosene are used by vessels. Fuel oil A, B, and C are used for propulsion of international water-borne vessels. Diesel oil and kerosene are used only for fuels of private power generator (eg. Air heating).

【CO₂】

The kiloliter-based consumption data given in the Ministry of Economy, Trade and Industry's

Yearbook of Mineral Resources and Petroleum Products Statistics (former Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke) is converted to a Joule-based data using the standard calorific values given in the *General Energy Statistics* by the Agency for Natural Resources and Energy.

【CH₄, N₂O】

The *Revised 1996 IPCC Guidelines* provide a default emission factor that is based on net calorific values. Therefore, activity data in gross calorific values are converted to net calorific values by multiplying them by the conversion rate of 0.95, except 0.975 for fuel oil C.

In addition, regarding activity data of N₂O from an international aviation, the *Revised 1996 IPCC Guidelines* provide a default emission factor in weight units. In order to adapt the activity data to this unit, the kiloliter-based consumption data is multiplied by the density identified by the Petroleum Association of Japan for N₂O from aircraft (0.78 [g/cm³]).

c) Other issues

The desk review report in 2004 indicated that there was a significant difference between bunker active data reported in the CRF (table 1.C) and bunker consumption data reported to the International Energy Agency (IEA). The followings explain the causes for the difference.

➤ *Data Update*

The ERT in 2004 used the following IEA energy balances for analysis.

- Data for 2000-2001: ” ENERGY BALANCES OF OECD COUNTRIES 2000-2001」 II 94-95”
- Data for 2002-2003: “ ENERGY BALANCES OF OECD COUNTRIES 2002-2003」 II 94-95”

After the publication of the data, it was found out that there were some errors in data of 2000 and 2001 submitted to IEA, including omission of full counting of imported bunker fuel and errors in the values of exported diesel oil. In March 2006, Japan reported the revision of these errors and the errors have been corrected since then.

➤ *Difference of fuel types reported as “bunker”*

Up to Japan’s national greenhouse gas inventories submitted in May 2004, Japan reported the bonded imports and exports of fuel oil A, B, and C as navigation bunker. In IEA energy balance, navigation bunker reported includes bonded diesel oil, kerosene and lubricant, other than bonded fuel oil A, B and C. This difference causes the variation between inventory data and IEA data.

Japan revised the estimation method in the inventory submitted in August 2004 and has reported bonded diesel oil and kerosene consumption as navigation bunker since then⁵.

➤ *Errors of density and conversion factor*

Data for the IEA energy balance need to be reported in the metric-ton unit. Japan calculates and reports to IEA values in metric-ton by multiplying the volume of fuel combustion given in the *Yearbook of Mineral Resources and Petroleum Products Statistics* by the density of each fuel type given in the *information of petroleum*, Sekiyu –Tsushin. IEA converts the values in metric ton into tons of oil equivalent (TOE) by using conversion factors. Given that the values are expressed in net

⁵ Lubricant is not included because lubricant is not combusted by use.

calorific-based value equivalent, and the conversion factors used in IEA are net calorific value.

Conversion of a unit to TOE by using information given in the inventory can be conducted by multiplying the volume of fuel consumption by gross calorific-based values.

This difference in the conversion process causes the variation between IEA energy balance and Japan's energy statistics for inventory preparation.

Glossary

Bonded Jet Fuel

Under the Tariff Law, aircrafts (Japanese and non-Japanese) flying international routes are deemed to be “aircraft for international use”, and the fuel they consume is tariff-free, subject to the completion of the required procedures. The application of this legislation means that if fuel is refined from crude oil imported to Japanese refinery, both the crude oil import tariff and the petroleum tax are waived. Similarly, if fuel has been imported as a product, the product import tariff is waived. The foregoing is termed as “bonded jet fuel”.

Bonded Fuel Oil

Vessels that ply voyages between Japan and other countries are deemed to be “foreign trade vessels”, under the Tariff Law. The majority of their fuel is consumed outside Japanese territorial waters, and, therefore both tariffs and the petroleum tax are waived. The foregoing is termed as “bonded fuel oil”.

Bonded Export

The demand for fuel supplied to aircrafts (Japanese and non-Japanese) flying international routes and ships (Japanese and non-Japanese) that ply foreign ocean routes is termed as “bonded demand”. Jet fuel is supplied to aircrafts while fuel oil is supplied to ships. Of these bonded demand, the fuel supplied from products that was produced from crude oil is counted as bonded exports by the Ministry of Economy, Trade and Industry.

Bonded Import (Bond to Bond)

Fuel products that are imported from foreign countries, landed in a bonded area and supplied from the bonded area to bonded demand without going through domestic customs, is counted as bonded imports by the Ministry of Economy, Trade and Industry.

3.2.9. Feedstocks and Non-Energy Use of Fuels

In the method used to estimate GHG emissions from fuel combustion (1.A.), the energy consumption in the category of Non-energy use (#9500) in the *General Energy Statistics* was deducted from the total energy consumption, because these amounts of fuel was used as feedstocks without combustion and oxidation process.

The Non-energy category consists of the following two requirements: (1) Consumption which can be confirmed as clearly being employed for non-energy uses by official statistics, such as surveys of feedstocks inputs according to *Current Survey of Energy Consumption* which is the data source of *General Energy Statistics*; and (2) Products which are from the outset produced for the purpose of

non-energy use.

(However, that portion which is confirmed from official statistics such as *Current Survey of Energy Consumption* as having been employed for energy uses is treated as energy consumption and excluded from non-energy use.)

CO₂ emissions from combustion and oxidation in the process of production, use and abandonment of the amount of feedstocks and non-energy use which were deducted from 1.A are separately reported in the following sectors.

- ◆ Ammonia Production (2.B.1)
- ◆ Silicon Carbide Production (2.B.4)
- ◆ Calcium Carbide Production (2.B.4)
- ◆ Ethylene Production (2.B.5)
- ◆ Use of Electric Arc Furnaces in Steel Production (2.C.1)
- ◆ Wastes Incineration (Simple Incineration) (waste oil and waste plastics) (6.C)
- ◆ Emissions from the Decomposition of Petroleum-Derived Surfactants (6.D)

3.2.10. CO₂ capture from flue gases and subsequent CO₂ storage

The amount of CO₂ capture from flue gases and subsequent CO₂ storage was not estimated in Japan.

3.2.11. Emission from waste incineration with energy recovery

Below three cases that utilize waste as crude material meets definition of the emission from waste incineration with energy recovery.

- Waste incineration with energy recovery
- Direct use of waste as fuel
- Use of waste processed as fuel

Estimation method for emission from these sources is applied waste incineration (6.C.) method in accordance with the *1996 Revised IPCC Guidelines*. The value of emission is included in fuel combustion (1.A.1. and 1.A.2.) in accordance with the *1996 Revised IPCC Guidelines* and the *GPG (2000)*. Please refer to Chapter 8 for the details of the estimation methods.

The reporting category of the emissions for each type of waste is, according to its use as fuel or raw material, classified to either “Energy Industry (Category 1.A.1.)” or “Manufacturing and Construction (1.A.2)”. The fuel type is classified as “Other fuels”.

Greenhouse gas emissions during the direct use of waste as a raw material, such as plastics used as reducing agents in blast furnaces or as a chemical material in coking furnaces, or use of intermediate products manufactured using the waste as a raw material, are estimated in this category.

Refuse-derived solid fuels (RDF: Refuse Derived Fuel, RPF: Refuse Paper and Plastic Fuel) are used for the estimation of emissions from fuels produced from waste. The reporting categories of the above emissions are included in “Energy Industry (Category 1.A.1.)” or “Manufacturing/Construction (1.A.2)” according to the use of waste as fuels. The fuel type is classified as “Other fuels”.

Table 3-30 Categories for the calculation of emissions from waste incineration with energy recovery

Incineration	Waste category	Estimation classification	Category of estimation	CO ₂	CH ₄	N ₂ O
Waste incineration with energy recovery	Municipal solid waste	Plastic	1.A.1	○	○ Estimated in bulk	○ Estimated in bulk
		Synthetic textile	1.A.1	○		
		Other (biogenic)	1.A.1	/		
	Industrial solid waste	Waste oil	1.A.1	○	○	○
		Waste plastic	1.A.1	○	○	○
		Other (biogenic)	1.A.1	/	○	○
Direct use of waste as fuel	Municipal solid waste	Plastic	1.A.1/2	○	○	○
	Industrial solid waste	Waste oil	1.A.2	○	○	○
		Waste plastic	1.A.2	○	○	○
		Waste wood	1.A.2	/	○	○
	Waste tire	Fossil origin	1.A.1/2	○	○	○
		Biogenic origin	1.A.1/2	/		
Use of waste processed as fuel	Refuse derived fuel (RDF·RPF)	Fossil origin	1.A.1/2	○	○	○
		Biogenic origin	1.A.1/2	/		

* CO₂ emissions from the incineration of biomass-derived waste (including biomass-based plastics and waste animal and vegetable oil) is not included in the total emissions in accordance with the Revised 1996 IPCC Guidelines; instead it is estimated as a reference value and reported under “Biogenic” in Table 6.A,C of the CRF.

The greenhouse gas emissions from waste incineration for energy purpose and with energy recovery are shown in Table 3-31.

Table 3-31 GHG Emission from waste incineration with energy recovery and use of waste as raw material or fuel

Gas	Item	Unit	1990	1995	2000	2005	2008	2009	2010			
CO ₂	1.A.1 Energy Industries	a. Public Electricity and Heat Production	Gg-CO ₂	6,493	7,080	9,075	7,965	6,707	6,795	6,684		
		b. Petroleum Refining	Gg-CO ₂	NO	NO	1	6	4	5	6		
		c. Manufacture of Solid Fuels and Other Energy Industries	Gg-CO ₂	NO	NO	15	239	193	204	242		
	1.A.2 Manufacturing Industries and Construction	a. Iron and Steel	Gg-CO ₂	NO	NO	308	634	377	444	549		
		b. Non-Ferrous Metals	Gg-CO ₂	118	63	51	17	3	2	2		
		c. Chemicals	Gg-CO ₂	14	64	89	66	66	67	72		
		d. Pulp, Paper and Print	Gg-CO ₂	NO	55	113	993	1,604	1,651	1,711		
		e. Food Processing, Beverages and Tobacco	Gg-CO ₂	IE	IE	IE	IE	IE	IE	IE	IE	
			Mining	Gg-CO ₂	IE	IE	IE	IE	IE	IE	IE	
			Construction	Gg-CO ₂	IE	IE	IE	IE	IE	IE	IE	
			Oil Products	Gg-CO ₂	IE	IE	IE	IE	IE	IE	IE	
			Glass Wares	Gg-CO ₂	IE	IE	IE	IE	IE	IE	IE	
			f. Other	Cement & Ceramics	Gg-CO ₂	597	1,122	1,876	2,317	2,467	2,428	2,510
				Machinery	Gg-CO ₂	41	26	20	10	NO	NO	NO
				Duplication Adjustment	Gg-CO ₂	NO	NO	NO	NO	NO	NO	NO
				Other Industries & Small & Medium Enterprises	Gg-CO ₂	1,854	2,092	1,595	2,877	2,556	2,305	2,403
			Total		Gg-CO ₂	9,116	10,503	13,142	15,123	13,976	13,899	14,180
CH ₄	1.A.1 Energy Industries	a. Public Electricity and Heat Production	Gg-CH ₄	0.54	0.54	0.60	0.15	0.14	0.13	0.13		
		b. Petroleum Refining	Gg-CH ₄	NO	NO	0.000002	0.000018	0.000010	0.000013	0.000016		
		c. Manufacture of Solid Fuels and Other Energy Industries	Gg-CH ₄	NO	NO	IE	IE	IE	IE	IE		
	1.A.2 Manufacturing Industries and Construction	a. Iron and Steel	Gg-CH ₄	NO	NO	NA	0.00036	0.00065	0.00065	0.00066		
		b. Non-Ferrous Metals	Gg-CH ₄	0.00032	0.00018	0.00014	0.00008	0.00002	0.00001	0.00001		
		c. Chemicals	Gg-CH ₄	0.00006	0.00013	0.00019	0.00019	0.00019	0.00019	0.00020		
		d. Pulp, Paper and Print	Gg-CH ₄	NO	0.0001	0.0002	0.0027	0.0045	0.0046	0.0047		
		e. Food Processing, Beverages and Tobacco	Gg-CH ₄	IE	IE	IE	IE	IE	IE	IE	IE	
			Mining	Gg-CH ₄	IE	IE	IE	IE	IE	IE	IE	
			Construction	Gg-CH ₄	IE	IE	IE	IE	IE	IE	IE	
			Oil Products	Gg-CH ₄	IE	IE	IE	IE	IE	IE	IE	
			Glass Wares	Gg-CH ₄	IE	IE	IE	IE	IE	IE	IE	
			f. Other	Cement & Ceramics	Gg-CH ₄	0.04	0.08	0.15	0.21	0.25	0.24	0.22
				Machinery	Gg-CH ₄	0.00018	0.00012	0.00009	0.00005	NO	NO	NO
				Duplication Adjustment	Gg-CH ₄	NO	NO	NO	NO	NO	NO	NO
				Other Industries & Small & Medium Enterprises	Gg-CH ₄	1.77	1.77	2.22	2.90	4.02	4.02	3.73
			Total		Gg-CH ₄	2.34	2.39	2.98	3.26	4.41	4.39	4.09
		Gg-CO ₂ eq	49.20	50.28	62.52	68.53	92.53	92.17	85.80			
N ₂ O	1.A.1 Energy Industries	a. Public Electricity and Heat Production	Gg-N ₂ O	1.20	1.33	1.56	1.14	1.04	0.96	0.96		
		b. Petroleum Refining	Gg-N ₂ O	NO	NO	0.00001	0.00012	0.00006	0.00008	0.00010		
		c. Manufacture of Solid Fuels and Other Energy Industries	Gg-N ₂ O	NO	NO	IE	IE	IE	IE	IE		
	1.A.2 Manufacturing Industries and Construction	a. Iron and Steel	Gg-N ₂ O	NO	NO	NA	0.0007	0.0013	0.0013	0.0013		
		b. Non-Ferrous Metals	Gg-N ₂ O	0.00024	0.00013	0.00011	0.00006	0.00001	0.00001	0.00001		
		c. Chemicals	Gg-N ₂ O	0.00004	0.00060	0.00092	0.00107	0.00110	0.00113	0.00121		
		d. Pulp, Paper and Print	Gg-N ₂ O	NO	0.0007	0.0014	0.0175	0.0279	0.0286	0.0295		
		e. Food Processing, Beverages and Tobacco	Gg-N ₂ O	IE	IE	IE	IE	IE	IE	IE	IE	
			Mining	Gg-N ₂ O	IE	IE	IE	IE	IE	IE	IE	
			Construction	Gg-N ₂ O	IE	IE	IE	IE	IE	IE	IE	
			Oil Products	Gg-N ₂ O	IE	IE	IE	IE	IE	IE	IE	
			Glass Wares	Gg-N ₂ O	IE	IE	IE	IE	IE	IE	IE	
			f. Other	Cement & Ceramics	Gg-N ₂ O	0.01	0.02	0.04	0.05	0.05	0.05	0.05
				Machinery	Gg-N ₂ O	0.00013	0.00008	0.00007	0.00003	NO	NO	NO
				Duplication Adjustment	Gg-N ₂ O	NO	NO	NO	NO	NO	NO	NO
				Other Industries & Small & Medium Enterprises	Gg-N ₂ O	0.03	0.03	0.03	0.05	0.06	0.06	0.05
			Total		Gg-N ₂ O	1.24	1.38	1.63	1.26	1.18	1.10	1.10
		Gg-CO ₂ eq	385.39	428.89	506.38	391.20	364.52	339.81	340.36			

3.3. Fugitive Emissions from Fuels (1.B.)

The Fugitive Emissions subsector consists of intentional and unintentional emissions of CO₂, CH₄, and N₂O from unburned fossil fuels during their mining, production, processing, refining, transportation, storage, and distribution.

There are two main source categories in this sector: Solid Fuels (1.B.1), emissions from coal mining and handling, and Oil and Natural Gas (1.B.2), emissions from the oil and natural gas industries. The main source of emissions from solid fuels is CH₄ contained in coal bed, whereas fugitive emissions, venting, flaring, volatilization, and accidents are the main emission sources in the oil and natural gas industries.

In 2010, GHG emissions from fugitive emission from fuels were 409 Gg-CO₂ eq. and accounted for 0.03 % of the Japan's total GHG emissions (excluding LULUCF). The emissions have decreased by 87 % compared to 1990.

Table 3-32 Emission trends of the fugitive emissions subsector (1.B)

Gas	IPCC Category			Unit	1990	1995	2000	2005	2008	2009	2010
CH ₄	1.B.1 Solid Fuels	a. Coal Mining	i. Underground Mines	Gg-CH ₄	132.63	63.45	36.11	3.07	1.55	1.67	1.65
			ii. Surface Mines	Gg-CH ₄	1.01	0.58	0.51	0.43	0.63	0.53	0.47
	1.B.2 Oil and Natural Gas	b. Natural Gas		Gg-CH ₄	1.35	1.75	1.42	1.41	1.30	1.21	1.17
				Gg-CH ₄	8.95	9.87	10.98	13.30	15.35	14.81	14.07
			c. Venting and Flaring	- Venting	Gg-CH ₄	0.58	0.86	0.53	0.51	0.47	0.43
	- Flaring	Gg-CH ₄		0.11	0.14	0.11	0.13	0.14	0.13	0.12	
	Total				Gg-CH ₄	144.63	76.66	49.67	18.84	19.44	18.77
				Gg-CO ₂ eq.	3,037.14	1,609.87	1,043.15	395.74	408.29	394.26	375.73
CO ₂	1.B.1 Solid Fuels	a. Coal Mining	i. Underground Mines	Gg-CO ₂	NE	NE	NE	NE	NE	NE	NE
			ii. Surface Mines	Gg-CO ₂	NE	NE	NE	NE	NE	NE	NE
	1.B.2 Oil and Natural Gas	b. Natural Gas		Gg-CO ₂	0.14	0.20	0.14	0.15	0.12	0.11	0.10
				Gg-CO ₂	0.25	0.27	0.31	0.38	0.45	0.43	0.41
			c. Venting and Flaring	- Venting	Gg-CO ₂	0.005	0.007	0.005	0.004	0.004	0.004
	- Flaring	Gg-CO ₂		36.22	50.44	35.58	37.06	37.27	34.60	32.64	
	Total				Gg-CO ₂	36.62	50.92	36.03	37.60	37.85	35.15
N ₂ O	1.B.1 Solid Fuels	a. Coal Mining	i. Underground Mines	Gg-N ₂ O	NE	NE	NE	NE	NE	NE	NE
			ii. Surface Mines	Gg-N ₂ O	NE	NE	NE	NE	NE	NE	NE
	1.B.2 Oil and Natural Gas	b. Natural Gas		Gg-N ₂ O	3.06E-07	3.40E-07	3.74E-07	5.10E-07	2.38E-07	2.04E-07	2.04E-07
				Gg-N ₂ O							
			c. Venting and Flaring	- Venting	Gg-N ₂ O						
	- Flaring	Gg-N ₂ O		0.00036	0.00050	0.00036	0.00038	0.00039	0.00036	0.00034	
	Total				Gg-N ₂ O	0.00036	0.00050	0.00036	0.00038	0.00039	0.00036
				Gg-CO ₂ eq.	0.11	0.16	0.11	0.12	0.12	0.11	0.11
Total of all gas				Gg-CO ₂ eq.	3,073.88	1,660.95	1,079.29	433.46	446.26	429.52	408.98

3.3.1. Solid Fuels (1.B.1.)

3.3.1.1. Coal Mining and Handling (1.B.1.a.)

3.3.1.1.a. Underground Mines (1.B.1.a.i.)

a) Source/Sink Category Description

Coal contains CH₄ that forms during the coalification process. Most will have been naturally released from the ground surface before mine development, but mining releases the CH₄ remaining in coal beds into the atmosphere.

The number of operational coal mines in Japan has decreased and coal production has decreased greatly as well. As a result, the amount of the CH₄ emissions from coal mining has shown a yearly decrease.

Furthermore, the coal mining practices have changed recently, resulting in the decreasing trend of CH₄ IEF. Specifically, coal is now mined in more shallow areas, therefore emitting less CH₄. This is because deep areas are costly to mine compared to coal in shallow areas. Additionally, areas which have been previously mined, therefore already releasing CH₄, are re-mined for coal, using the latest technology. This contributes to low CH₄ emission per amount of coal mined also if compared with other countries.

Although a reporting column is provided for CO₂ emissions associated with coal mining, in the absence of a default emission factor, emissions from this source were reported as “NE”. Coal mining exists in Japan, and, depending on the CO₂ concentration in the coal being mined, the CO₂ may be released into the atmosphere during mining activity. Although it is believed that coal beds in Japan do not contain CO₂ at a concentration level that is higher than that in the atmosphere, emissions cannot be calculated because of the absence of actual measurements. Because of the absence as well of a default value for CO₂ emissions associated with coal mining, emissions from this source are not reported.

b) Methodological Issues

● Estimation Method

➤ Mining Activities

Emissions from mining activities were drawn from actual measurements obtained from individual coal mines using the Tier 3 method, in accordance with Decision Tree of the *GPG (2000)* (Page 2.72, Fig. 2.10).

➤ Post-Mining Activities

Emissions from post-mining activities were estimated using the Tier 1 method, which uses default emission factors in accordance with Decision Tree of the *GPG (2000)* (Page 2.73, Fig. 2.11). It was estimated by multiplying the amount of coal mined from underground mining by the emission factor.

● Emission Factors

➤ Mining Activities

The emission factor for mining activities was established by dividing the emissions of CH₄ gas identified in a survey by Japan Coal Energy Center (J-COAL), by the production volume of coal from underground mines.

Table 3-33 Emission factors for mining activities – Underground mines

Item	Unit	1990	1995	2000	2005	2008	2009	2010	Reference
Coal Production of Underground Mines	kt	6,775	5,622	2,364	738	536	575	588	Surveyed by J-COAL
CH ₄ Total Emissions	1000 m ³	181,358	80,928	48,110	2,781	1,001	1,089	1,025	Surveyed by J-COAL
CH ₄ Total Emissions	Gg-CH ₄	121.5	54.2	32.2	1.9	0.7	0.7	0.7	=CH ₄ [1000m ³] / 1000 X 0.67 [Gg/10 ⁶ m ³]
Emission Factor	kg-CH ₄ /t	17.9	9.6	13.6	2.5	1.3	1.3	1.2	CH ₄ Total Emissions / Coal Production of Underground Mines

➤ Post-Mining Activities

Due to the lack of data for emissions from post-mining activities in Japan, emission factors were calculated (1.64 [kg CH₄/t]) by converting the median value (2.45 m³/t) of the default values (0.9 – 4.0 m³/t) given in the *Revised 1996 IPCC Guidelines* by the density of CH₄, 0.67 (1,000 t/10⁶ m³) at 20°C

and 1 atmosphere.

● **Activity Data**

➤ *Mining Activities, Post-Mining Activities*

The value used for activity data for underground mining and post-mining activities was derived by subtracting the surface mining production from the total coal production as given in the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* prepared by the Ministry of Economy, Trade and Industry and the data provided by Japan Coal Energy Center (J-COAL).

Table 3-34 Trends in coal production

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Total Coal Production	kt	7,980	6,317	2,974	1,249	1,290	1,206	1,145
Surface Mines	kt	1,205	695	610	511	754	631	557
Underground Mines	kt	6,775	5,622	2,364	738	536	575	588

c) *Uncertainties and Time-series Consistency*

● **Uncertainties**

Uncertainty for CH₄ emissions from mining activities was calculated to be 5% based on the values of measurement error and error of gas flow velocity fluctuation.

Uncertainty for CH₄ emissions from post-mining activities was 200%, which is the value of the default data in *GPG (2000)*. A summary of uncertainty assessment methods is provided in Annex 7.

● **Time-series Consistency**

The CH₄ emissions data for mining activities in underground mines have been derived from *Japan Coal Energy Center (J-COAL)* statistics consistently since FY 1990.

Total coal production and coal production on surface mines were provided by the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* prepared by the Ministry of Economy, Trade and Industry from FY 1990 to FY 2000. Thereafter, they have been provided by the Japan Coal Energy Center (J-COAL), because categories of surface mining production and total coal production in the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* is no longer conducted. The data from the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* prepared by the Ministry of Economy, Trade and Industry until 2000 are provided by Japan Coal Energy Center (J-COAL). Therefore, total coal production data from both of these sources are same and have been used in a consistent manner since FY 1990.

d) *Source-specific QA/QC and Verification*

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

In order to ensure safety of coal mine workers in Japan, monitoring the concentration of CH₄ and CO in coal mines is ordained by law. Under the law, mining companies must set rules on monitoring management. Mining companies monitor accurately under strict management and checks, and compile relevant reports. Furthermore, national authorities regularly check monitoring measurements and safety reports.

e) Source-specific Recalculations

There have been no recalculations to emissions from this source category.

f) Source-specific Planned Improvements

There are no major planned improvements in this source category.

3.3.1.1.b. Surface Mines (1.B.1.a.ii.)**a) Source/Sink Category Description**

This category provides the estimation methods for fugitive emissions of CH₄ occur during the coal mining and post-mining activities on surface mines.

b) Methodological Issues● **Estimation Method**➤ **Mining Activities**

CH₄ emissions were calculated using the Tier 1 method and the default emission factor in accordance with Decision Tree of the *GPG (2000)* (Page 2.71, Fig. 2.9).

➤ **Post-Mining Activities**

CH₄ emissions were calculated using the Tier 1 method and the default emission factor in accordance with Decision Tree of the *GPG (2000)* (Page 2.73, Fig. 2.11).

Both were calculated by multiplying the amount of coal mined from surface mining by the relevant emission factors.

● **Emission Factors**➤ **Mining Activities**

A value (0.77 [kg-CH₄/t-coal]) was used as the emission factor for mining activities. It was derived by converting the median (1.15 [m³/t]) of the default values given in the *Revised 1996 IPCC Guidelines* (0.3–2.0 [m³/t]), using the concentration of CH₄ at one atmospheric pressure and 20°C (0.67 [Gg/10⁶m³]).

➤ **Post-Mining Activities**

A value (0.067 [kg-CH₄/t-coal]) was used as emission factor for post-mining activities. It was derived by converting the median (0.1 [m³/t]) of the default values given in the *Revised 1996 IPCC Guidelines* (0–0.2 [m³/t]), using the concentration of CH₄ at one atmospheric pressure and 20°C (0.67 [Gg/10⁶m³]).

● **Activity Data**

The figure for the surface production given in the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* prepared by the Ministry of Economy, Trade and Industry and the data provided by the Japan Coal Energy Center (J-COAL) were used as the activity data for mining and post-mining activities (see Table 3-34).

c) Uncertainties and Time-series Consistency● **Uncertainties**

The uncertainties for emission factors were applied 200% of default data indicated in the *GPG (2000)*. The uncertainty of activity data was 10%; this was determined as a standard value by the Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties of CH₄ emissions

from surface mines were estimated to 200% for both mining and post-mining activities. Summary of uncertainty assessment methods are provided in Annex 7.

● **Time-series Consistency**

Total coal production and coal production on surface mines were provided by the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* prepared by the Ministry of Economy, Trade and Industry from FY 1990 to FY 2000. Thereafter, they have been provided by the Japan Coal Energy Center (J-COAL), because categories of surface mining production and total coal production in the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* is no longer conducted. The data from the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* prepared by the Ministry of Economy, Trade and Industry until 2000 are provided by Japan Coal Energy Center (J-COAL). Therefore, total coal production data from both of these sources are same and have been used in a consistent manner since FY 1990.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

There have been no recalculations to emissions from this source category.

f) Source-specific Planned Improvements

There are no major planned improvements in this source category.

3.3.1.2. Solid Fuel Transformation (1.B.1.b.)

In Japan, the production of briquettes is believed to meet the description of the activity of conversion to solid fuel. The process of coal briquette production includes introducing water to coal, and squeeze-drying it. Therefore, the process is not thought to involve any chemical reactions, but the emission of CO₂, CH₄ or N₂O cannot be denied. However, as no actual measurements have been taken, however, it is not presently possible to calculate emissions. CO₂, CH₄ and N₂O emissions associated with the conversion to solid were reported as “NE” in the absence of default values.

3.3.2. Oil and Natural Gas (1.B.2.)

3.3.2.1. Oil (1.B.2.a.)

3.3.2.1.a. Exploration (1.B.2.a.i.)

a) Source/Sink Category Description

This category provides the estimation methods for fugitive emissions of CO₂, CH₄ and N₂O occur during the exploratory drilling of oil and gas fields and pre-production tests.

b) Methodological Issues

● **Estimation Method**

CO₂, CH₄ and N₂O emissions associated with oil exploration drilling and pre-production testing was

calculated using the Tier 1 Method in accordance with the Decision Tree of *GPG (2000)*. Emissions were calculated by multiplying the number of exploratory drilling wells, and the number of wells tested for oil and gas during pre-production testing, by their respective emission factors.

● **Emission Factors**

The emission factors from the *GPG (2000)* for drilling and testing wells were used.

Table 3-35 Emission factors for exploratory drilling and testing wells [Gg/number of wells]

	CH ₄	CO ₂	N ₂ O
Drilling	4.3×10^{-7}	2.8×10^{-8}	0
Testing	2.7×10^{-4}	5.7×10^{-3}	6.8×10^{-8}

Source: GPG (2000), p. 2.86, Table 2.16

● **Activity Data**

➤ *Drilling*

The data given in the *Natural Gas Data Year Book* compiled by the Natural Gas Mining Association were used for exploratory drilling wells.

➤ *Testing*

It was not possible to readily ascertain statistically the number of wells in which oil and gas testing had been carried out, and even where such tests are conducted, not all wells are successful. For that reason, the number of wells tested for oil and gas used the median values of the number of exploratory drilling wells and the number of successful wells shown in the *Natural Gas Data Year Book*. As for the most recent year, the data of the previous year were provisionally used.

Table 3-36 Trends in the number of exploratory drilling wells and those tested for oil and gas

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Number of Wells Drilled	wells	8	7	7	10	6	4	4
Number of Wells Succeeded	wells	1	3	4	5	1	2	2
Number of Wells Tested	wells	5	5	6	8	4	3	3

c) **Uncertainties and Time-series Consistency**

● **Uncertainties**

Because all emission factors for exploration of oil and natural gas were the default values in *GPG (2000)*, the uncertainties for emission factors were assessed based on default values (25%) described in *GPG (2000)*. The uncertainty of activity data was 10%; this was determined as a standard value by the Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties for emissions were estimated to be 27% each for the fugitive emissions of CO₂, CH₄, and N₂O that occur during the exploration of oil and natural gas. A summary of uncertainty assessment methods are provided in Annex 7.

● **Time-series Consistency**

Emission factors have used consistent values since FY 1990. Activity data have been calculated by using annual data from the *Natural Gas Data Year Book* and a consistent estimation method since FY 1990.

d) **Source-specific QA/QC and Verification**

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

Since the activity data of FY 2009 was obtained, the GHG emissions of FY 2009 were recalculated. Also the activity data of FY 2003 was revised, the GHG emissions of FY 2003 were recalculated.

f) Source-specific Planned Improvements

There have been no major planned improvements in this source category.

3.3.2.1.b. Production (1.B.2.a.ii.)

a) Source/Sink Category Description

This category provides the estimation methods for fugitive emissions of CO₂ and CH₄ occur during production of crude oil, as well as when measuring instruments are lowered into oil wells during inspection of operating oil fields.

b) Methodological Issues

● Estimation Method

Emissions relating to fugitive emissions from petroleum production and servicing of oilfield production wells were calculated using the Tier 1 method in accordance with Decision Tree of the *GPG (2000)* (Page 2.81, Fig. 2.13). Emissions were calculated by multiplying the amount of crude oil production by the emission factor.

● Emission Factors

➤ Production

The default value for conventional crude oil given in the *GPG (2000)* was used for the emission factor of fugitive emissions from petroleum production. (The median of the default values was used for CH₄).

Table 3-37 EF for fugitive emissions from petroleum production [Gg/10³kl]

		CH ₄ ¹⁾	CO ₂	N ₂ O ²⁾
Conventional Oil	Fugitive emissions	1.45×10 ⁻³	2.7×10 ⁻⁴	0

Source: GPG (2000) Table 2.16

1) The default value is 1.4×10⁻³ – 1.5×10⁻³

2) Excluded from calculations, as the default value is 0 (zero)

➤ Servicing

The default value given in the *GPG (2000)* was used as the emission factor for fugitive emissions from servicing of petroleum production wells.

Table 3-38 Emission factors for fugitive emissions from servicing of petroleum production wells [Gg/number of wells]

	CH ₄	CO ₂	N ₂ O ¹⁾
Production Well (Servicing)	6.4×10 ⁻⁵	4.8×10 ⁻⁷	0

Source: GPG (2000) Table 2.16

1) Excluded from calculations, as the default value is 0 (zero)

- **Activity Data**

- **Production**

The values for production of crude oil in Japan given in the Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke and the Yearbook of Mineral Resources and Petroleum Products Statistics prepared by the Ministry of Economy, Trade and Industry were used as the activity data for fugitive emissions from production. However, condensates were not included.

- **Servicing**

Because the number of oil wells and natural gas wells cannot be separated, the total fugitive emissions from servicing of oil and natural gas wells are reported in the subcategory *1.B.2.b.ii. Exploration*. The oil is reported as “IE” here.

c) Uncertainties and Time-series Consistency

- **Uncertainties**

As the uncertainty of emission factors, default values given in the *GPG (2000)* (25% for CO₂ and 25% for CH₄) were applied. The uncertainty of activity data was 5%; this was determined as a standard value by the Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties for the emissions were determined to be 25% for CO₂ and for CH₄. The uncertainty assessment methods are summarized in Annex 7.

- **Time-series Consistency**

Emission factors have been used consistent values since FY 1990. Activity data have been calculated using annual data from the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* and the *Yearbook of Mineral Resources and Petroleum Products Statistics* prepared by the Ministry of Economy, Trade and Industry, in a consistent manner since FY 1990.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

There have been no recalculations to emissions from this source category.

f) Source-specific Planned Improvements

There are no major planned improvements in this source category.

3.3.2.1.c. Transport (1.B.2.a.iii.)

a) Source/Sink Category Description

This category provides the estimation methods for fugitive emissions of CO₂ and CH₄ occur during the transportation of crude oil and condensate through pipelines, tank trucks, and tank cars to refineries.

b) Methodological Issues

- **Estimation Method**

Emissions relating to fugitive emissions associated with transport were calculated using the Tier 1 method in accordance with Decision Tree of the *GPG (2000)* (Page 2.81, Fig. 2.13). Emissions were

calculated by multiplying the amount of crude oil or condensate production by the emission factors.

Fugitive emissions from transporting oil from domestic oilfield at sea to land and fugitive emissions from land transport were estimated. Crude oil for sea transport is carried out entirely by pipeline, and is not expected to generate any fugitive emissions from other transportation mode. Land transport includes a number of methods, including pipeline, tank trucks, and tank cars, but it is difficult to differentiate them statistically. For that reason, it has been assumed that all of the produced oil is transported by tank trucks or tank cars in estimations.

● *Emission Factors*

The default values given in the *GPG (2000)* were used as the emission factors.

Table 3-39 Emission factors for transportation of crude oil and condensate [Gg/10³kl]

	CH ₄	CO ₂	N ₂ O ¹⁾
Transportation of crude oil	2.5×10 ⁻⁵	2.3×10 ⁻⁶	0
Transportation of condensate	1.1×10 ⁻⁴	7.2×10 ⁻⁶	0

Source: GPG (2000) Table 2.16

1) Excluded from calculations, as the default value is 0 (zero)

● *Activity Data*

The values for production of oil in Japan given in the Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke and the Yearbook of Mineral Resources and Petroleum Products Statistics prepared by the Ministry of Economy, Trade and Industry, were used as the activity data for fugitive emissions from transport.

Table 3-40 Production of crude oil and condensate in Japan

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Oil Production Excluding Condensate	kl	420,415	622,679	385,565	370,423	340,593	309,526	292,539
Condensate Production	kl	234,111	242,859	375,488	540,507	632,654	607,672	560,106
Oil Production (Total)	kl	654,526	865,538	761,053	910,930	973,247	917,198	852,645

c) *Uncertainties and Time-series Consistency*

● *Uncertainties*

As the uncertainty of emission factors, default values given in the *GPG (2000)* (25% for CO₂ and 25% for CH₄) were applied. The uncertainty of activity data was 5%; this was determined as a standard value by the Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties for the emissions were determined to be 25% for CO₂ and for CH₄. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

Emission factors have been used consistent values since FY 1990. Activity data have been calculated using annual data from the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* and the *Yearbook of Mineral Resources and Petroleum Products Statistics* prepared by the Ministry of Economy, Trade and Industry, in a consistent estimation method since FY 1990.

d) *Source-specific QA/QC and Verification*

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference

materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

There have been no recalculations to emissions from this source category.

f) Source-specific Planned Improvements

There are no major planned improvements in this source category.

3.3.2.1.d. Refining / Storage (1.B.2.a.iv)

a) Source/Sink Category Description

This category provides the estimation methods for fugitive emissions of CH₄ occur when crude oil is refined or stored at oil refineries.

CO₂ emissions from this source were reported as “NE”. Refining / Storage activities exist in Japan and extremely small amount of CO₂ may be released into the atmosphere from the activities if CO₂ is included in crude oil. Because there is no examples of actual measurements of the CO₂ content of crude oil as well as a default value, CO₂ emissions from this source were not estimated.

b) Methodological Issues

● **Estimation Method**

➤ **Oil Refining**

Emissions relating to fugitive emissions from refining were calculated using the Tier 1 method in accordance with Decision Tree of the *GPG (2000)* (Page 2.82, Fig. 2.14).

➤ **Oil Storage**

Emissions relating to fugitive emissions from storage should be calculated using the Tier 1 method in accordance with Decision Tree of the *GPG (2000)* (Page 2.82, Fig.2.14), but as the country-specific emission factor is available for this emissions source, it was applied to the inventories instead.

● **Emission Factors**

➤ **Oil Refining**

With respect to the emissions factors for the fugitive emissions during the refining processes, the amount of CH₄ emitted during crude oil refining processes was considered to be negligible because fugitive emission of CH₄ was unlikely to occur in Japan during crude oil refining at normal operation. For that reason, the lower limit of the default values shown in the *Revised 1996 IPCC Guidelines* was adopted.

Table 3-41 Emission factor during refining of crude oil

Emission Factor [kg-CH ₄ /PJ]	
Oil Refining	90 ¹⁾

Source: Revised 1996 IPCC Guidelines, Volume 3 Table 1-58

1) The default value is 90–1,400

➤ **Oil Storage**

Oil is stored in either corn-roof tanks or floating-roof tanks. All oil storage in Japan adopts floating-roof tanks, which means that fugitive CH₄ emissions are considered to be very small. If fugitive CH₄ emissions were to occur, they could only occur by vaporization of oil left on the exposed wall wet with oil when the floating roof descends as the stored oil is removed; thus, the amount of

fugitive CH₄ emissions would be small.

The Petroleum Association of Japan has conducted experiments relating to the evaporation of CH₄ from tank walls by modeling the floating-roof tank to calculate estimates of CH₄ emissions.

The emission factor associated with storage of crude oil is a value derived by converting the estimates of the Petroleum Association (0.007 Gg/year as at 1998) to a net calorific value and dividing it by the relevant activity data.

Table 3-42 Assumptions for calculation of emission factor during oil storage

CH ₄ Emissions [kg-CH ₄ /year]	Input of Crude Oil to Oil Refining Industry		Emission Factor [kg-CH ₄ /PJ]
	[PJ: Gross Calorific Value] ¹⁾	[PJ: Net Calorific Value] ²⁾	
7,000	9,921	9,424.95	0.7427

1) Agency for Natural Resources and Energy, *General Energy Statistics*

2) Net Calorific Value = Gross Calorific Value × 0.95

● Activity Data

The value used for activity data during refining and storing was the converted net calorific values of NGL and refined crude oil in petroleum refining industry taken from the *General Energy Statistics* compiled by the Agency for Natural Resources and Energy.

Table 3-43 Amount of crude and NGL refined in Japan

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Oil and NGL Refined	PJ:NCV	7,732	8,907	8,898	8,820	8,054	7,542	7,497

c) Uncertainties and Time-series Consistency

● Uncertainties

For the uncertainty of emission factors for fugitive emissions of CH₄ occurring when crude oil is refined or stored at oil refineries, values shown in the *Revised 1996 IPCC Guidelines* are applied. The uncertainties for emission factors were applied 25% of default data indicated in the *GPG (2000)* in accordance with Decision Tree of uncertainty assessment of emission factor. The uncertainty for activity data was evaluated to be 0.9% by combining the uncertainty of crude oil and NGL indicated in the *General Energy Statistics*. As a result, the uncertainties for emissions were determined to 25% for CH₄ emissions from the source. Summary of uncertainty assessment methods are provided in Annex 7.

● Time-series Consistency

Emission factors have been used consistent values since FY 1990. Activity data have been calculated using annual data from the *General Energy Statistics*, in a consistent estimation method since FY 1990.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

GHG emissions in FY 2009 were recalculated because of the revision of the fuel consumption in FY 2009 in the *General Energy Statistics*.

f) Source-specific Planned Improvements

There are no major planned improvements in this source category.

3.3.2.1.e. Distribution of Oil Products (1.B.2.a.v.)

Petroleum products are distributed in Japan, and where CO₂ and CH₄ are dissolved, it is conceivable that either or both will be emitted as a result of the relevant activity. The level of CO₂ or CH₄ emitted by the activity is probably negligible, in light of the composition of the petroleum products, but because there are no examples of measurement of the CO₂ or CH₄ content of petroleum products, it is not currently possible to calculate emissions. Emissions were reported as “NE” in the absence of the default emission factors.

3.3.2.2. Natural Gas (1.B.2.b.)

3.3.2.2.a. Exploration (1.B.2.b.i.)

There are test drillings of oil and gas fields in Japan, and it is conceivable that the activity could give rise to emissions of CO₂, CH₄, or N₂O. It is difficult, however, to distinguish between oilfields and gas fields prior to test drilling, therefore the emissions were reported as “IE” because the calculation was combined with the subcategory of *1.B.2.a.i. Fugitive Emissions Associated with Oil Exploration*.

3.3.2.2.b. Production / Processing (1.B.2.b.ii.)

a) Source/Sink Category Description

This category provides the estimation methods for CO₂ and CH₄ emissions of fugitive emissions from the production of natural gas, processing through the adjusting of its constituent elements, and through the lowering of measurement instruments during servicing of natural gas production wells.

b) Methodological Issues

● *Estimation Method*

Fugitive emissions from the production of natural gas, processing through the adjusting of its constituent elements, and through the lowering of measurement instruments during servicing of natural gas production wells was calculated using the Tier 1 method, and in accordance with Decision Tree of the *GPG (2000)* (Page 2.80, Fig. 2.12).

Fugitive emissions during natural gas production and conditioning processes were estimated by multiplying the amount of natural gas production by their respective emission factors. Fugitive emissions during gas field inspections were calculated by multiplying the number of production wells by the emission factor.

● *Emission Factors*

➤ *Production*

The default values given in the *GPG (2000)* were used for the emission factors of fugitive emissions during the production of natural gas. (The median of the default values was used for CH₄).

Table 3-44 Emission factors of fugitive emissions during production of natural gas [Gg/10⁶ m³]

		CH ₄ ¹⁾	CO ₂	N ₂ O ²⁾
Natural Gas Production	Fugitive Emissions	2.75×10 ⁻³	9.5×10 ⁻⁵	0

Source: GPG (2000) Table 2.16

1) The default values are 2.6×10⁻³ – 2.9×10⁻³

2) Excluded from calculations, as the default value is 0 (zero)

➤ Processing

The default values given in the *GPG (2000)* for the emission factors of fugitive emissions during processing of natural gas were used. (The median of the default values was used for CH₄).

Table 3-45 Emission factors during processing of natural gas [Gg/10⁶ m³]

		CH ₄ ¹⁾	CO ₂	N ₂ O ²⁾
Processing of Natural Gas	Processing in general (General treatment plant, Sweet Gas Plants)	8.8×10 ⁻⁴	2.7×10 ⁻⁵	0

Source: GPG (2000) Table 2.16

1) The default values are 6.9×10⁻⁴ – 10.7×10⁻⁴

2) Excluded from calculations, as the default value is 0 (zero)

➤ Servicing

The default values for fugitive emissions during servicing of natural gas production wells given in the *GPG (2000)* were used.

Table 3-46 Emission factors during servicing of natural gas production wells [Gg/number of wells]

	CH ₄	CO ₂	N ₂ O ¹⁾
Production Well (Servicing)	6.4×10 ⁻⁵	4.8×10 ⁻⁷	0

Source: GPG (2000) Table 2.16

1) Excluded from calculations, as the default value is 0 (zero)

● Activity Data

➤ Production and Processing

The production volume of natural gas in Japan given in the Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke and the Yearbook of Mineral Resources and Petroleum Products Statistics prepared by the Ministry of Economy, Trade and Industry, was used as the activity data during its production and processing.

➤ Servicing

Because the number of oil wells and natural gas wells cannot be separated for the entire time series, the total fugitive emissions from servicing of oil and natural gas wells are reported here. The number of oil/natural gas wells shown in the *Natural Gas Data Year Book* published by the Japan Natural Gas Association was used. As for the most recent year, the data of the previous year was provisionally used.

Table 3-47 Natural gas production and the number of producing and capable wells

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Natural Gas Production	10 ⁶ m ³	2,066	2,237	2,499	3,140	3,706	3,555	3,343
Number of Producing and Capable Wells	wells	1,230	1,205	1,137	1,115	1,065	1,049	1,049

c) Uncertainties and Time-series Consistency

● Uncertainties

As the uncertainty of emission factors for the CO₂ and CH₄ emissions from fugitive emissions of the production and processing of natural gas, default values given in the *GPG (2000)* (25% for CO₂ and 25% for CH₄) were applied. The uncertainty of activity data was 5%; this was determined as a standard value by the Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties for the emissions were determined to be 25% for CO₂ and for CH₄.

As the uncertainty of emission factors for the CO₂ and CH₄ emissions from fugitive emissions from servicing of oil and natural gas wells, default values given in the *GPG (2000)* (25% for CO₂ and 25% for CH₄) were applied. The uncertainty of activity data was 10%; this was determined as a standard value by the Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties for the emissions were determined to be 27% for CO₂ and for CH₄.

The uncertainty assessment methods are summarized in Annex 7.

● ***Time-series Consistency***

Emission factors have used consistent values since FY 1990. Activity data have been calculated by using annual data on the production volume of natural gas from the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* and the *Yearbook of Mineral Resources and Petroleum Products Statistics* prepared by the Ministry of Economy, Trade and Industry, and on the number of oil/natural gas wells from the *Natural Gas Data Year Book*. A consistent estimation method has been used since FY 1990.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

Since the activity data of FY 2009 was obtained, the GHG emission of FY 2009 was recalculated.

f) Source-specific Planned Improvements

There are no major planned improvements in this source category.

3.3.2.2.c. Transportation (1.B.2.b.iii.)

a) Source/Sink Category Description

This category provides the estimation methods for CH₄ emissions in conjunction with transportation of domestically produced natural gas, such as the release of gas when relocating and building pipelines, and the release of gas used to operate pressure regulators.

Emissions from CO₂ in this source are reported as “NA”. Approximately 90% of town gas is based on LNG and is free of CO₂. However, domestically produced natural gas from some of Japan’s natural gas formations contains CO₂. Because nearly all of this CO₂ is removed at natural gas production plants before the gas is sent to pipelines, the natural gas provided by town gas suppliers likely contains hardly any CO₂. Emission of CO₂ removed at natural gas production plants is assigned to natural gas production and processing (1.B.2.b.ii).

b) Methodological Issues

● Estimation Method

Total natural gas pipeline length is multiplied by a Japan-specific emission factor to calculate CH₄ emissions occurring in conjunction with releases by pipeline construction and relocation, and releases of gas used to operate pressure regulators.

● Emission Factors

The amount of CH₄ emitted from a 1-km length of domestic natural gas pipeline over a 1-year period is defined as the emission factor, and is set by dividing the CH₄ emission amount by pipeline length. Due to the insufficiency of past data, it was decided to use a uniform emission factor that was set using FY2004 data for 1990 and subsequent years. Data were provided by the Japan Natural Gas Association.

1) Gas Releases Due To Pipeline Relocation

The equation below was used as the basis for calculating the CH₄ amount released when in-pipe pressure is reduced for relocating gas pipelines. Further, after relocation work is complete it is necessary to flush the pipeline with natural gas, which is released before introduction into the pipeline. The amount of CH₄ is determined by measuring with a gas meter or calculating it using means such as pipeline pressure when introducing the gas. These were calculated for each pipeline relocation and the annual cumulative total determined.

$$\text{CH}_4 \text{ emission amount} = \text{volume of pipe section with reduced pressure} \times \text{pressure before reduction (absolute pressure)} / \text{atmospheric pressure (absolute pressure)} \times \text{CH}_4 \text{ content (CH}_4 \text{ per Nm}^3\text{)}$$

2) Gas Releases Due To Pipeline Installation

After installation work is complete, it is necessary to flush the pipeline with natural gas, which is released before introduction into the pipeline. The amount of CH₄ is determined by measuring with a gas meter or calculating it using means such as pipeline pressure when gas is introduced, and their annual cumulative total determined.

3) Release of Gas for Operating Pressure Regulators

The amount of natural gas used in accordance with specifications of pressure regulators for reducing gas supply pressure is calculated as follows.

$$\text{CH}_4 \text{ emission amount} = \text{amount used according to pressure regulator specifications} \times \text{number of regulators installed} \times \text{CH}_4 \text{ content (CH}_4 \text{ per Nm}^3\text{)}$$

Table 3-48 FY2004 CH₄ emissions as a concomitant of natural gas transportation

	Amount of gas used (Nm ³ /day)	Number of work	Number of establishment	Amount of gas releases (k-Nm ³)	CH ₄ content (t-CH ₄ /kNm ³)	CH ₄ releases (t-CH ₄)
Pipeline Relocation & Installation	---	77	---	843	0.645	544
Gas for Operating Pressure Regulators	19	---	48	333	0.643	215
Total	---	---	---	---	---	759

➤ *Total Pipeline Length*

We used 2,090 km as the total length of natural gas pipeline of the main association members covered by an FY2004 study by the Japan Natural Gas Association, which is the pipeline whose emissions are of concern here.

$\begin{aligned} \text{Emission factor} &= \text{CH}_4 \text{ release amount} / \text{total pipeline length} \\ &= 759 \text{ t-CH}_4 / 2090 \text{ km} \\ &= 0.363 \text{ t-CH}_4/\text{km} \end{aligned}$

● *Activity Data*

The length of natural gas pipeline laid in Japan given by the Japan Natural Gas Association in its *Natural Gas Data Year Book* was used as the activity data of the length of natural gas pipeline laid. As for the most recent year, the data of the previous year was provisionally used.

Table 3-49 Length of natural gas pipeline installation

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Natural Gas Pipeline length	km	1,984	2,195	2,434	2,721	3,016	3,027	3,027

c) *Uncertainties and Time-series Consistency*

● *Uncertainties*

A country-specific emission factor is used for CH₄ in conjunction with transportation. As the uncertainty of emission factors, default values given in the *GPG (2000)* (25% for CH₄) were applied because according to the Decision Tree, either expert judgement or the default value given in the *GPG (2000)* is to be adopted. The uncertainty of activity data was 10%; this was determined as a standard value by the Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties for the emissions were determined to be 27% for CH₄. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

Emission factors have been used consistent values since FY 1990. Activity data have been calculated using annual data from the *Natural Gas Data Year Book*, in a consistent estimation method since FY 1990.

d) *Source-specific QA/QC and Verification*

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) *Source-specific Recalculations*

Since the activity data of FY 2009 was obtained, the GHG emission of FY 2009 was recalculated.

f) *Source-specific Planned Improvements*

The CH₄ emissions in conjunction with transportation of domestically produced natural gas are estimated on the premise that the full transportation of natural gas is sent to pipelines(1.B.2.b.iii.), however, recently there are some cases of the transportation of LNG by tank trucks or tank cars. LNG transported by tank trucks and tank cars is basically sealed. There is no research on the actual situation for whole in Japan, and no default value, so this current estimation method is continuously adopted. If sufficient data on CH₄ emissions from transportation of natural gas by the tank trucks or tank cars is obtained in the future, the possibilities of estimation methods for this category should be considered.

3.3.2.2.d. Distribution (1.B.2.b.iv.-)

a) Source/Sink Category Description

This category provides the estimation methods for CH₄ emitted from the normal operation of LNG receiving terminals, town gas production facilities, and satellite terminals, as well as during regular maintenance or construction, and for CH₄ emitted from town gas supply networks.

In Japan, liquefied petroleum gas, coal, coke, naphtha, crude oil, and natural gas are refined and blended at gas plants into gas, which, after being conditioned to produce a certain calorific value, is supplied to urban areas through gas lines. Such gas fuel is called “town gas”, of which more than 90% is LNG-based.

Japan reports the emissions associated with the production of town gas (Natural Gas Supplies) in the category of *1.B.2.b. Natural Gas Distribution*. The town gas production is accounted for in this category, even though it may not meet the definition in the *Revised 1996 IPCC Guidelines* exactly, because of the lack of a category more appropriate for reporting of emissions from town gas production.

Emissions from CO₂ in this source are reported as “NA”. More than 90% of town gas is based on LNG and is free of CO₂. However, domestically produced natural gas from some of Japan’s natural gas formations contains CO₂. Because nearly all of this CO₂ is removed at natural gas production plants before the gas is sent to pipelines, the natural gas provided by town gas suppliers likely contains hardly any CO₂. Emission of CO₂ removed at natural gas production plants is assigned to natural gas production and processing (1.B.2.b.ii).

b) Methodological Issues

● Estimation Method

➤ LNG Receiving Terminals, Town Gas Production Facilities, and Satellite Terminals (Natural Gas Supplies)

Some of the main emission sources are gas samples taken for analysis and residual gas emitted at times such as regular maintenance of manufacturing facilities. The Tier 1 method is employed in accordance with the *GPG (2000)* decision tree (page 2.82, Fig. 2.14). However, because it is possible to use a Japan-specific emission factor, the amounts of liquefied natural gas and natural gas used as town gas feedstock were multiplied by a Japan-specific emission factor to obtain emissions.

➤ Town Gas Supply Networks

CH₄ emissions from high-pressure pipelines and from medium- and low-pressure pipelines and holders are calculated by multiplying the total length of city gas pipeline by the emission factor. CH₄ emissions from service pipes are calculated by multiplying the number of users by the emission factor.

● Emission Factors

➤ LNG Receiving Terminals, Town Gas Production Facilities, and Satellite Terminals (Natural Gas Supplies)

The emission factor was calculated by dividing emission of CH₄ during the normal operation of LNG

receiving terminals, town gas production facilities, and satellite terminals in Japan, as well as during regular maintenance or construction, by the calorific value of the raw material input (LNG, natural gas). The emission factor calculated using FY1998 data was 905.41 (kgCH₄/PJ), while that calculated using FY2007 data was 264.07 (kgCH₄/PJ). The main reason for the emission factor change was the reduction in CH₄ emissions, which was due to progress in reduction measures such as the installation of new sampling and recovery lines used for gas analyses (changes to lines that recover gas from atmospheric dispersion) in LNG receiving terminals and town gas production facilities. Because measures to reduce CH₄ emissions have been gradually implemented, emission factors for the period from FY1999 to FY2006 were set by linear interpolation. At this time, measures to reduce CH₄ emissions have been generally implemented, thereby affording little expectation of major change in the emission factor for the time being. Therefore, the FY2007 emission factor value will be kept the same for FY2008 and subsequent years.

➤ *Town Gas Supply Networks*

Emission sources in the supply of domestically produced town gas are (i) high-pressure pipelines, (ii) medium- and low-pressure pipelines and holders, and (iii) service pipes. FY2004 data were used to calculate CH₄ emissions for each of the minor categories of each of the emission sources shown in Table 3-50. The emission factor for high-pressure pipelines and for medium- and low-pressure pipelines and holders was set using the CH₄ amount emitted from 1 km of the town gas pipeline length during 1 y, while that for service pipes was set using the CH₄ amount emitted from 1000 users' homes during 1 y.

Table 3-50 CH₄ emissions from town gas pipelines and emission factors (Established by FY2004 data)

Emission Sources		CH ₄ emissions (t/yr)	Source sizes	Emission factors
High-pressure pipelines	New pipeline installation Pipeline relocation	180	Total high-pressure pipeline 1799 km	0.100 t-CH ₄ /km
Medium- and low-pressure pipelines and holders	Construction and demolition Fugitive emissions Burner and other inspections Holder construction and overhauling	93	Total medium- and low-pressure pipeline 226,016 km	0.411 kg-CH ₄ /km
Service pipes	Installing service pipes Post-installation purging Removal Changing meters Fugitive emissions, etc. Rounds for opening valves and regular maintenance Equipment repairs (Especially high emissions when doing work at user sites (homes))	19	User homes 27,298,000	0.696 kg-CH ₄ /1000 homes

● *Activity Data*

➤ *LNG Receiving Terminals, Town Gas Production Facilities, and Satellite Terminals (Natural Gas Supplies)*

The amounts of LNG and natural gas shown in the *General Energy Statistics* (Agency for Natural

Resources and Energy) as used as raw material for town gas.

Table 3-51 Liquefied natural gas and natural gas used as material for town gas

Item	Unit	1990	1995	2000	2005	2008	2009	2010
LNG Consumption with Town Gas Production	PJ	464	676	864	1,230	1,439	1,424	1,555
Natural Gas Consumption with Town Gas Production	PJ	40	48	61	86	131	127	115

➤ *Town gas supply networks*

Estimates use the high-pressure pipeline length, total medium- and low-pressure pipeline length, and number of users given in the *Gas Industry Yearbook* of the Agency for Natural Resources and Energy Gas Market Division.

Table 3-52 High-pressure pipeline length, total medium- and low-pressure pipeline length, and number of users

Item	Unit	1990	1995	2000	2005	2008	2009	2010
High-pressure Pipeline Length	km	1,067	1,281	1,443	1,898	2,029	2,066	2,124
Total Medium- and Low-pressure Pipeline Length	km	180,239	197,474	214,312	230,430	239,336	241,675	244,022
Number of Users	10 ³ houses	21,334	23,580	25,858	27,762	28,599	28,774	28,902

c) *Uncertainties and Time-series Consistency*

● *Uncertainties*

Although CH₄ emission factor of natural gas supplies is country-specific, the uncertainty of emission factor is the default value (25%) given in the *GPG (2000)* because the application of statistical treatment was considered to be unsuitable. The uncertainty of activity data was determined to be 8.7% by combining of the uncertainty of LNG and natural gas presented in the *General Energy Statistics*. As a result, the uncertainties for emissions were estimated to be 26% for CH₄ emissions from natural gas supplies.

A country-specific emission factor is used for CH₄ emissions from town gas supply networks. The uncertainties for emission factors of town gas supply network were the default values presented in *GPG (2000)* (25% for CH₄) were applied because default value of expert opinion or *GPG (2000)* is adopted in accordance with Decision Tree of uncertainty assessment of emission factor. For the uncertainty for activity data, the value set by the Committee for Greenhouse Gas Emission Estimation Methods (10%) was applied. As a result, the uncertainties for emissions were estimated to be 27% for CH₄ emissions from town gas supply network. A summary of uncertainty assessment methods are provided in Annex 7.

● *Time-series Consistency*

Emission factors have used consistent values as described above since FY 1990. Activity data have been calculated using annual data on LNG and natural gas consumption and town gas production from *General Energy Statistics* and data on the town gas supply network from the *Gas Industry Yearbook*. A consistent estimation method has been used since FY 1990.

d) *Source-specific QA/QC and Verification*

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

There have been no recalculations to emissions from this source category.

f) Source-specific Planned Improvements

There are no major planned improvements in this source category.

3.3.2.2.e. At industrial plants and power station / in residential and commercial sectors (1.B.2.b.v.)

Conceivable sources of these CH₄ emissions include gas pipe work in buildings, but because these emissions are included in those of “Natural Gas Distribution” (distribution through the town gas network) (1.B.2.b.iv), CH₄ emissions from this source are reported as “IE.” Additionally, because CO₂ is basically not included among town gas constituents, CO₂ emissions from this source are reported as “NA.”

3.3.2.3. Venting and Flaring (1.B.2.c.)

This section includes fugitive emissions of CO₂ and CH₄ occur from venting during oil field development, crude oil transportation, refining processes, and product transportation in the petroleum industry and as well as during gas field development, natural gas production, transportation, and processing in natural gas industry.

It also includes CO₂, CH₄, N₂O emissions from flaring during the above processes.

3.3.2.3.a. Venting (Oil) (1.B.2.c.-venting i.)**a) Source/Sink Category Description**

This category provides the estimation methods for CO₂ and CH₄ from venting in the petroleum industry.

b) Methodological Issues**● Estimation Method**

Emissions from venting in the petroleum industry were calculated using the Tier 1 Method in accordance with the Decision Tree of *GPG (2000)* (Page 2.81, Fig. 2.13) by multiplying the amount of crude oil production by the default emission factors.

● Emission Factors

The default values for conventional oil given in the *GPG (2000)* were used for the emission factors of oilfield venting. (The median of the default values was used for CH₄).

Table 3-53 Emission factors of oilfield venting

		CH ₄ ¹⁾	CO ₂	N ₂ O ²⁾
Conventional Oil	Venting valves [Gg/1000 m ³]	1.38×10 ⁻³	1.2×10 ⁻⁵	0

Source: GPG (2000) Table 2.16

1) The default values are 6.2×10⁻⁵ - 270×10⁻⁵

2) Excluded from calculations, as the default value is 0 (zero)

- **Activity Data**

The production volume of oil in Japan given by the Ministry of Economy, Trade and Industry in its Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke and the Yearbook of Mineral Resources and Petroleum Products Statistics was used as the activity data of fugitive emissions from oilfield venting. The production of condensate was excluded from the calculation (see Table 3-40).

c) Uncertainties and Time-series Consistency

- **Uncertainties**

As the uncertainty of emission factors, default values given in the *GPG (2000)* (25% for CO₂ and CH₄) were applied. The uncertainty of activity data was 5%; this was determined as a standard value by the Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties for the emissions were determined to be 25% for CO₂ and CH₄. The uncertainty assessment methods are summarized in Annex 7.

- **Time-series Consistency**

Emission factors have been used consistent values as described above since FY 1990. Activity data have been calculated using annual data from the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* and the *Yearbook of Mineral Resources and Petroleum Products Statistics*, in a consistent estimation method since FY 1990.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

There have been no recalculations to emissions from this source category.

f) Source-specific Planned Improvements

There have been no major planned improvements in this source category.

3.3.2.3.b. Venting (Gas) (1.B.2.c.-venting ii.)

CO₂ and CH₄ emissions from venting in the natural gas industry were considered only for the amount during transportation because *GPG (2000)* provides emissions factors only for transportation. Intentional CO₂ emissions from natural gas pipelines are reported as “NA” because CO₂ emissions during Transportation of natural gas are considered as “NA” (1.B.2.b.iii). Intentional CH₄ emissions from natural gas pipelines are reported as “IE” because they are included in emissions during natural gas transportation (1.B.2.b.iii).

3.3.2.3.c. Venting (Oil and Gas) (1.B.2.c.-venting iii.)

Statistical data are reported for two categories of petroleum and natural gas in Japan. As a result, fugitive emissions from venting in the combined petroleum and natural gas industries were reported as “IE” since they were accounted for respectively in the emissions from venting in the petroleum industry (1.B.2.c.i) and the natural gas industry (1.B.2.c.ii).

3.3.2.3.d. Flaring (Oil) (1.B.2.c.-flaring i.)

a) Source/Sink Category Description

This category provides the estimation methods for CO₂, CH₄, and N₂O from flaring in the petroleum industry.

b) Methodological Issues

● Estimation Method

CO₂, CH₄, and N₂O emissions from flaring in the petroleum industry were calculated using the Tier 1 Method in accordance with the Decision Tree of *GPG (2000)*, by multiplying the amount of crude oil production in Japan by the default emissions factors.

● Emission Factors

In the absence of actual measurement data or country-specific emission factors in Japan, the default values shown in *GPG (2000)* were used. It should be noted that the median values were used for CH₄ emissions.

Table 3-54 Emission factors for flaring in the oil industry

	Unit	CH ₄ ¹⁾	CO ₂	N ₂ O
Flaring (Conventional Oil)	Gg/10 ³ m ³	1.38×10 ⁻⁴	6.7×10 ⁻²	6.4×10 ⁻⁷

Source: GPG (2000), Table 2.16

1) Default value: 0.05×10⁻⁴ to 2.7×10⁻⁴

● Activity Data

For the calculation of activity data for this emission source, the amounts of crude oil production shown in the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* and the *Yearbook of Natural Resources and Petroleum Products*, both published by Ministry of Economy, Trade and Industry, were used. The production of condensate was excluded from the calculation (see Table 3-40).

c) Uncertainties and Time-series Consistency

● Uncertainties

As the uncertainty of emission factors, default values given in the *GPG (2000)* (25% for CO₂, CH₄, and N₂O) were applied. The uncertainty of activity data was 5%; this was determined as a standard value by the Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties for the emissions were determined to be 25% for CO₂, CH₄, and N₂O. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

Emission factors have been used consistent values as described above since FY 1990. Activity data have been calculated using annual data from the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* and the *Yearbook of Mineral Resources and Petroleum Products Statistics*, in a consistent estimation method since FY 1990.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

There have been no recalculations to emissions from this source category.

f) Source-specific Planned Improvements

There have been no major planned improvements in this source category.

3.3.2.3.e. Flaring (Natural Gas) (1.B.2.c.-flaring ii.)**a) Source/Sink Category Description**

This category provides the estimation methods for CO₂, CH₄, and N₂O from flaring in the natural gas industry.

b) Methodological Issues**● Estimation Method**

CO₂, CH₄, and N₂O emissions associated with flaring in the natural gas industry were calculated using the Tier 1 Method in accordance with the Decision Tree of *GPG (2000)*. Emissions were calculated by multiplying the amount of production of natural gas by the emission factors. The total emissions associated with flaring both during gas production and processing were reported as the emissions from flaring in the natural gas industry.

● Emission Factors

The default values for fugitive emissions from flaring (Natural Gas) given in the *GPG (2000)* were used.

Table 3-55 Emission factors for flaring in the natural gas industry

		Unit	CH ₄	CO ₂	N ₂ O
Flaring in the natural gas industry	Gas production	Gg/10 ⁶ m ³	1.1 × 10 ⁻⁵	1.8 × 10 ⁻³	2.1 × 10 ⁻⁸
	Gas processing	Gg/10 ⁶ m ³	1.3 × 10 ⁻⁵	2.1 × 10 ⁻³	2.5 × 10 ⁻⁸

Source: *GPG (2000)*, Table 2.16

● Activity Data

For the calculation of activity data for this emission source, the amounts of domestic production of natural gas shown in the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* and the *Yearbook of Natural Resources and Petroleum Products*, both published by Ministry of Economy, Trade and Industry, were used (see Table 3-47).

c) Uncertainties and Time-series Consistency**● Uncertainties**

As the uncertainty of emission factors, default values given in the *GPG (2000)* (25% for CO₂, CH₄, and N₂O) were applied. The uncertainty of activity data was 5%; this was determined as a standard value by the Committee for the Greenhouse Gas Emission Estimation Methods. As a result, the uncertainties for the emissions were determined to be 25% for CO₂, CH₄, and N₂O. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

Emission factors have been used consistent values as described above since FY 1990. Activity data have been calculated using annual data from the *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke* and the *Yearbook of Mineral Resources and Petroleum Products Statistics*,

in a consistent estimation method since FY 1990.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. The QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. Details of the QA/QC activities are provided in Annex 6.

e) Source-specific Recalculations

There have been no recalculations to emissions from this source category.

f) Source-specific Planned Improvements

There have been no major planned improvements in this source category.

3.3.2.3.f. Flaring (Oil and Gas) (1.B.2.c.-flaring iii.)

Statistical data are reported for two categories of petroleum and natural gas in Japan. As a result, fugitive emissions from flaring in the combined petroleum and natural gas industries were reported as “IE” since they were accounted for respectively in the emissions from flaring in the petroleum industry (1.B.2.c.i) and the natural gas industry (1.B.2.c.ii.)

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Chapter 4. Industrial Processes (CRF sector 2)

4.1. Overview of Sector

Chemical and physical transformation in industrial processes produce atmospheric GHG emissions. This chapter describes the methodologies of estimating industrial process emissions shown in Table 4-1.

In 2010, total GHG emissions from the industrial processes sector amounted to approximately 65,898Gg-CO₂ eq., accounting for 5.2% of national total emissions (excluding LULUCF) in Japan. The emissions (excluding F-gases) from this sector has decreased by 38.2% compared to 1990. The emissions of halocarbons and SF₆ from this sector has decreased by 54.3% compared to 1995.

Table 4-1 Emission source categories in the industrial processes sector

Emission source categories			CO ₂	CH ₄	N ₂ O	HFCs	PFCs	SF ₆	
2.A Mineral Products	2.A.1	Cement Production	○						
	2.A.2	Lime Production	○						
	2.A.3	Limestone and Dolomite Use	○						
	2.A.4	Soda Ash Production and Use	○						
	2.A.5	Asphalt Roofing	NE						
	2.A.6	Road Paving with Asphalt	NE						
	2.A.7	Other	IE, NO	NA, NO	NA, NO				
2.B Chemical Industry	2.B.1	Ammonia Production	○	NE	NA				
	2.B.2	Nitric Acid Production			○				
	2.B.3	Adipic Acid Production	NA		○				
	2.B.4	Carbide Production	Silicon Carbide	○	○				
			Calcium Carbide	○	NA				
	2.B.5	Other	Carbon Black		○				
			Ethylene	○	○	NA			
			1,2-Dichloroethane		○				
Styrene				○					
Methanol									
		Coke	IE	○	NA				
2.C Metal Production	2.C.1	Iron and Steel Production	Steel	IE	NA				
			Pig Iron	IE	NA				
			Sinter	IE	IE				
			Coke	IE	IE				
			Use of Electric Arc Furnaces in Steel Production	○	○				
	2.C.2	Ferroalloys Production	IE	○					
	2.C.3	Aluminium Production	IE	NE			○		
2.C.4	SF ₆ Used in Aluminium and Magnesium Foundries	Aluminium						NO	
		Magnesium						○	
2.C.5	Other	NO	NO	NO					
2.D Other Production	2.D.1	Pulp and Paper							
	2.D.2	Food and Drink	IE						
2.E Production of Halocarbons and SF ₆	2.E.1	By-product emissions: Production of HCFC-22				○			
	2.E.2	Fugitive emissions				○	○	○	

(continued on next page)

Emission source categories				CO ₂	CH ₄	N ₂ O	HFCs	PFCs	SF ₆		
2.F Consumption of Halocarbons and SF ₆	2.F.1	Refrigeration and Air Conditioning Equipment	Domestic Refrigeration	manufacturing				○	NO	NO	
				stocks				IE	NO	NO	
				disposal				IE	NO	NO	
			Commercial Refrigeration	Commercial Refrigeration	manufacturing				○	NO	NO
					stocks				IE	NO	NO
				disposal				IE	NO	NO	
				Automatic Vending Machine	manufacturing				○	NO	NO
					stocks				IE	NO	NO
					disposal				IE	NO	NO
			Transport Refrigeration	manufacturing				IE	NO	NO	
				stocks				IE	NO	NO	
				disposal				IE	NO	NO	
			Industrial Refrigeration	manufacturing				IE	NO	NO	
				stocks				IE	NO	NO	
				disposal				IE	NO	NO	
			Stationary Air-Conditioning (Household)	manufacturing				○	NO	NO	
				stocks				IE	NO	NO	
				disposal				IE	NO	NO	
	Mobile Air-Conditioning (Car Air Conditioners)	manufacturing				○	NO	NO			
		stocks				IE	NO	NO			
		disposal				IE	NO	NO			
	2.F.2	Foam Blowing	Hard Foam	Urethane Foam	manufacturing				○	NO	NO
					stocks				○	NO	NO
					disposal				IE	NO	NO
				High Expanded Polyethylene Foam	manufacturing				○	NO	NO
					stocks				NO	NO	NO
					disposal				NO	NO	NO
				Extruded Polystyrene Foam	manufacturing				○	NO	NO
					stocks				○	NO	NO
					disposal				IE	NO	NO
				Soft Foam							NO
	2.F.3	Fire Extinguishers			manufacturing				NO	NO	NO
					stocks				○	NO	NO
					disposal				NO	NO	NO
	2.F.4	Aerosols/Metered Dose Inhalers	Aerosols	manufacturing				○	NO	NO	
				stocks				○	NO	NO	
				disposal				IE	NO	NO	
			Metered Dose Inhalers	manufacturing				○	NO	NO	
				stocks				○	NO	NO	
				disposal				IE	NO	NO	
	2.F.5	Solvents			manufacturing				NO	NO	NO
					stocks				IE	○	NO
					disposal				IE	IE	NO
	2.F.6	Other Applications Using ODS Substitutes						IE	NA	NA	
	2.F.7	Semiconductors	Semiconductors	manufacturing				IE	IE	IE	
				stocks				○	○	○	
				disposal				NA	NA	NA	
Liquid Crystals			manufacturing				IE	IE	IE		
			stocks				○	○	○		
			disposal				NA	NA	NA		
2.F.8	Electrical Equipment			manufacturing					○		
				stocks					○		
				disposal					IE		
2.F.9	Other						NA	NE, ○	IE		

Emissions reported indicated as ○, and refer to Abbreviations list for notation keys.

4.2. Mineral Products (2.A.)

This category covers CO₂ emissions from the calcination of mineral raw material such as CaCO₃, MgCO₃, Na₂CO₃, etc. This section includes GHG emissions from Cement Production (2.A.1), Lime Production (2.A.2.), Limestone and Dolomite Use (2.A.3.) and Soda Ash Production and Use (2.A.4.). In 2010, emissions from Mineral Products were 38,280 Gg-CO₂ eq. and represented 3.0% of total

GHG emissions (excluding LULUCF). The emissions decreased by 30.9% compared to 1990.

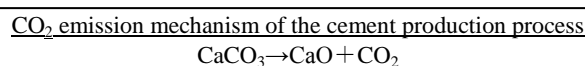
Table 4-2 CO₂ Emissions from 2.A. Mineral Products

Gas	Emission sub-category		Units	1990	1995	2000	2005	2008	2009	2010
CO ₂	2.A Mineral Products	2.A.1 Cement Production	Gg-CO ₂	37,905	41,275	34,394	31,579	27,925	24,755	23,784
		2.A.2 Lime	Gg-CO ₂	6,674	5,795	5,900	6,646	6,594	5,371	6,285
		2.A.3 Limestone and Dolomite Use	Gg-CO ₂	10,522	9,441	9,339	8,480	8,332	7,450	8,073
		2.A.4 Soda Ash Production and Use	Gg-CO ₂	267	250	209	197	159	138	138
	Total			Gg-CO ₂	55,369	56,761	49,842	46,903	43,009	37,714

4.2.1. Cement Production (2.A.1.)

a) Source/Sink Category Description

CO₂ is emitted by the calcination of limestone, the main component of which is calcium carbonate, during the production of clinker, an intermediate product of cement and the main component of which is calcium oxide.



b) Methodological Issues

● Estimation Method

Following the *GPG (2000)* decision tree, the CO₂ emissions from this source was estimated by multiplying the amount of clinker produced by an emission factor.

<u>CO₂ emissions (t-CO₂) from cement production</u> = emission factor (t-CO ₂ /t-clinker) × clinker production (t) × cement kiln dust correction coefficient
--

● Emission Factors

Multiplying the CaO content of clinker by the molecular weight ratio of CaO and CO₂ (0.785) yields the emission factor. Because Japan's cement industry takes in large amounts of waste and byproducts from other industries and recycles them as substitute raw materials for cement production, clinker contains CaO from sources other than carbonates. This CaO does not go through the limestone calcination stage and therefore does not emit CO₂ during the clinker production process. For that reason, emission factors were determined by estimating the CaO content of clinker from carbonates, by subtracting CaO originating from waste and other sources from the total CaO content of clinker. Japan applies 1.00 for the cement kiln dust (CKD) correction coefficient, because normally almost all CKD is recovered and used again in the production process, as confirmed by the Cement Association. The emission factors for CO₂ emitted from cement production were calculated using the following procedure.

- 1 Estimate dry weight of waste and other materials input in raw material processing.
- 2 Estimate the amount and content of CaO from waste and other materials in clinker.

- 3 Estimate the CaO content of clinker, excluding the CaO from waste and other materials.
- 4 Determine the clinker emission factor.

Emission factors of CO₂ emissions from cement production

$$= [(CaO \text{ content of clinker}) - (CaO \text{ content of clinker from waste and other materials})] \times 0.785$$

CaO content of clinker from waste and other materials

$$= \frac{\text{dry weight of inputs of waste and other materials} \times \text{CaO content of waste and other materials}}{\text{clinker production volume}}$$

➤ **Estimating dry weight of waste and other materials input in raw material processing**

The following 13 types of waste and other materials were chosen for this calculation: coal ash (incineration residue), sewage sludge incineration ash, municipal solid waste incineration ash, glass refuse/ceramics refuse, concrete refuse, blast furnace slag (water granulated), blast furnace slag (slow-cooled), steelmaking slag, nonferrous slag, casting sand, particulates/dust, coal ash (fluidized bed furnace ash), and coal ash (from dust collectors) (these waste account for over 90% of the CaO from waste and other materials). Waste amounts (emission-based) and the water content of each waste and other material were determined from studies by the Cement Association of Japan (only for 2000 and thereafter).

➤ **Estimating the amount and content of CaO from waste and other materials in clinker**

The dry weights of each type of waste and other materials found above are multiplied by the CaO content for each type as found by the Cement Association, thereby calculating the total CaO amount in clinker derived from waste and other materials. This is divided by clinker production amount to find the CaO content from waste and other materials in clinker. Because data for 1990 to 1999 are unavailable, averages for 2000 through 2003 were used.

➤ **Estimating the CaO content of clinker, excluding the CaO from waste and other materials**

CaO content in waste and other materials is subtracted from the average CaO content of clinker as determined by the Cement Association, which yields the proportion of CaO in clinker that is used to set emission factors.

Table 4-3 Composition of Waste Origin Material

Group	Types of waste	Water content	CaO content
Incineration residue	Coal ash	7.2 - 14.5%	5.0 - 5.8%
	Sewage sludge incineration ash *	11.6 - 14.9%	7.4 - 12.5%
	Municipal solid waste incineration ash *	20.3 - 24.4%	10.0 - 26.5%
Glass refuse, Concrete refuse, and Ceramics refuse	Glass refuse, Ceramics refuse *	16.8 - 32.7%	17.5 - 31.1%
	Concrete refuse *	10.0 - 22.2%	6.4 - 43.9%
Slag	Blast furnace slag (water granulated)	5.0 - 8.7%	40.0 - 42.4%
	Blast furnace slag (slow-cooled)	5.7 - 6.5%	40.8 - 41.5%
	Steelmaking slag	7.7 - 11.4%	34.8 - 40.5%
	Nonferrous slag	5.6 - 8.4%	6.4 - 10.0%
	Casting sand *	9.8%	6.5%
Particulates (dust collector dust)	Particulates/dust	8.9 - 14.3%	9.0 - 13.4%
	Coal ash (fluidized bed furnace ash) *	0.1 - 1.7%	14.5 - 20.7%
	Coal ash	1.4 - 3.9%	4.6 - 5.0%

* Newly added from FY2009.

Table 4-4 Emission factors of CO₂ from cement production

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Average CaO content in clinker	%	65.9	65.9	66.0	65.9	65.9	65.8	65.8
Waste Origin CaO content in clinker	%	2.6	2.6	2.9	2.0	1.9	1.7	1.7
CaO content in clinker excluding waste origin CaO	%	63.3	63.3	63.0	63.9	63.9	64.1	64.1
CO ₂ /CaO		0.785	0.785	0.785	0.785	0.785	0.785	0.785
EF	t-CO ₂ /t	0.497	0.497	0.495	0.501	0.502	0.503	0.503

● Activity Data

Cement Association provides the data on the amount of clinker produced. Because there is no statistics on clinker production from 1990 to 1999, an estimation is made for past (1990–1999) clinker production using the average values of the 2000–2003 ratios of clinker production (Cement Association data) to limestone consumption (Ministry of Economy, Trade and Industry, Yearbook of Ceramics and Building Materials Statistics).

Table 4-5 Clinker production

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Consumption of Limestone (actual)	kt (dry)	89,366	97,311	81,376	-	-	-	-
Clinker Production (actual)	kt	-	-	69,528	63,003	55,647	49,195	47,279
Clinker Production (actual) / Consumption of Limestone (actual)*		0.853	0.853					
Estimated Clinker Production after correction**	kt	76,253	83,032	69,528	63,003	55,647	49,195	47,279

* Clinker Production (actual) / Consumption of Limestone (actual) for 1990-1999 is the average value of 2000-2003.

** Values for FY 1990-1999 are corrected using estimation, and values for FY2000 and on are actual.

c) Uncertainties and Time-series Consistency

● Uncertainty

For the uncertainty of the CO₂ emission factor from cement production, the standard value given in the *GPG (2000)* was applied. For the uncertainty of activity data, the value of 10% given by the Committee for the Greenhouse Gas Emission Estimation Methods was used. As a result, the uncertainty of emissions was estimated to be 10%. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

CO₂ emissions from cement production from 1990 to 1999 is estimated using estimated activity data and emission factors based on values provided by the Cement Association. For years from 2000 and onward, the methodology described in the sections above is consistently applied using the data provided by Cement Association.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)*. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.

e) Source-specific Recalculations

There have been no source-specific recalculations.

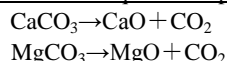
f) Source-specific Planned Improvements

No improvements are planned.

4.2.2. Lime Production (2.A.2.)**a) Source/Sink Category Description**

CO₂ is emitted during the calcination of CaCO₃, MgCO₃ in limestone used as raw material to produce quicklime.

CO₂ generation mechanism of quicklime production process

**b) Methodological Issues**

- **Estimation Method**

CO₂ emissions are calculated by multiplying limestone consumption by the country-specific emission factor.

CO₂ emissions (t-CO₂) generated by use of raw materials in quicklime production

= raw material-specific emission factor (t-CO₂/t-raw material) × amount of limestone consumption (t-product)

- **Emission Factors**

An emission factor per unit raw material (limestone) (0.428 t-CO₂/t-raw material) provided by the Japan Lime Association was used.

The Emission factor per unit raw material was calculated by finding the CO₂ emissions per unit raw material estimated from the amounts of carbon and other substances in raw material constituents and quicklime products, and then finding the weighted average using production amounts of each district. The emission factor for lime production is the same for all years because annual change is thought to be small.

- **Activity Data**

Limestone consumption data for quicklime and slaked lime use, categorized under 'Ceramic industry - other ceramics and quarry products' in the Adjusted Price Transaction Table is used. It is converted to dry weight using the water content from limestone used for cement.

The Adjusted Price Transaction Table (RIETI, 2010):

The Adjusted Price Transaction Table is a table created from the monetary input table in the Input-Output Table and the consumption data provided in industrial statistics, and is an application of similar estimation methods as in the General Energy Statistics.

In the existing transaction table attached to the Input-Output Table, although expressing the domestic supply and demand of products without any omission/duplication, there exists the possibility of over/under evaluation of transaction depending on the sector if the actual price differs, since transaction in each sector is based on the input from the average price across all industries. In contrast, the Adjusted Price Transaction Table attempts to eliminate differences between sectors, by taking into consideration the uneven transaction prices based on the differences in product quality/form in each sector, and through using statistical values in industrial statistics etc to the extent which possible.

By using consumption data in the Adjusted Price Transaction Table as activity data, it is considered

possible to capture activity data for all industries without omission/duplication, and to achieve a correct categorization of emission/non-emission related use, based on its detailed breakdown of sectors.

In the inventory, limestone/dolomite consumption data by sector in the Adjusted Price Transaction Table will be used as activity data for each limestone related source, excluding that for 'Cement Production (2.A.1.)'.

As for the dolomite consumed in dolomitic lime production, it is accounted for under 'Limestone and Dolomite Use (2.A.3.)', therefore it will not be included under 'Lime Production (2.A.2.)'. As for the re-absorption of CO₂ by the production of light calcium carbonate, it is already deducted by accounting for limestone consumption equivalent to the amount of light calcium carbonate production subtracted from lime production, under the lime production sector,

Table 4-6 Limestone Consumption

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Limestone consumption (dry)	kt	15,595	13,540	13,785	15,527	15,406	12,548	14,684

c) Uncertainties and Time-series Consistency

● *Uncertainty*

The uncertainty for CO₂ emissions from quicklime lime production was estimated. The uncertainty of 15% as given in the *GPG (2000)* was used for emission factors for both types of lime. For the uncertainty of activity data, 5% was used. As a result, the uncertainty of emissions was estimated to be 16%. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series consistency*

Limestone consumption data provided in the Adjusted Price Transaction Table is used as lime production activity data for all years from FY1990. The emission factors are constant for all years from FY1990. Therefore, CO₂ emission from lime production has been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

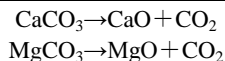
No improvements are planned.

4.2.3. Limestone and Dolomite Use (2.A.3.)

a) Source/Sink Category Description

Limestone contains CaCO_3 and minute amounts of MgCO_3 , and dolomite contains CaCO_3 and MgCO_3 . The heating of limestone and dolomite releases CO_2 derived from CaCO_3 and MgCO_3 .

CO₂ generating mechanism of limestone and dolomite use



b) Methodological Issues

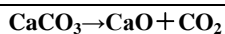
● Estimation Method

The amounts of limestone and dolomite used are multiplied by the emission factors to calculate emissions.

● Emission Factors

➤ Limestone

The emission factor is calculated by adding the value obtained when multiplying the molecular weight ratio of CO_2 and CaCO_3 by the percentage of CaO that can be extracted from limestone (55.4%, the median value of the “54.8% to 56.0%” given in The Story of Lime [Japan Lime Association]) and the value obtained when multiplying the molecular weight ratio of CO_2 and MgCO_3 by the percentage of MgO that can be extracted from limestone (0.5%, the median value of the “0.0% to 1.0%” given in The Story of Lime [Japan Lime Association]).



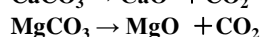
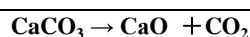
- Proportion of CaO extractable from limestone: 55.4 %
(Median of 54.8% to 56.0%: Japan Lime Association, The Story of Lime)
- Proportion of MgO extractable from limestone: 0.5 %^b
(Median of 0.0% to 1.0%: Japan Lime Association, The Story of Lime)
- Molecular weight of CaCO_3 (primary constituent of limestone) : 100.0869^a
- Molecular weight of MgCO_3 : 84.3139^a
- Molecular weight of CaO : 56.0774^a
- Molecular weight of MgO : 40.3044^a
- Molecular weight of CO_2 : 44.0095^a
- CaCO_3 content = proportion of CaO extractable from limestone × molecular weight of CaCO_3 / molecular weight of CaO
= (55.4% × 100.0869) / 56.0774 × 100 = 98.88%
- MgCO_3 content = proportion of MgO extractable from limestone × molecular weight of MgCO_3 / molecular weight of MgO
= 0.5% × 84.3139 / 40.3044 = 1.05%
- Emission factor = (molecular weight of CO_2 / molecular weight of CaCO_3 × CaCO_3 content)
+ (molecular weight of CO_2 / molecular weight of MgCO_3 × MgCO_3 content)
= 44.0095 / 100.0869 × 98.88 + 44.0095 / 84.3139 × 1.05
= 0.4348 + 0.0055 = 0.4402 [t- CO_2 /t]
= 440 [kg- CO_2 /t]

Sources)

- a. IUPAC “Atomic Weights of the Elements 1999”
(<http://www.chem.qmul.ac.uk/iupac/AtWt/AtWt99.html>)
- b. Japan Lime Association “The Story of Lime”

➤ Dolomite

The emission factor is calculated by adding the value obtained when multiplying the molecular weight ratio of CO₂ and CaCO₃ by the percentage of CaO that can be extracted from dolomite (34.5%, the median value of the 33.1% to 35.85% range given in *The Story of Lime* [Japan Lime Association]) and the value obtained when multiplying the molecular weight ratio of CO₂ and MgCO₃ by the percentage of MgO that can be extracted from dolomite (18.3%, the median value of the 17.2% to 19.5% range given in *The Story of Lime* [Japan Lime Association]).



- Proportion of CaO extractable from dolomite: 34.5%
(Median value of the 33.1% to 35.85% range given in *The Story of Lime* [Japan Lime Association])
- Proportion of MgO extractable from dolomite: 18.3%
(Median value of the 17.2% to 19.5% range given in *The Story of Lime* [Japan Lime Association])
- Molecular weight of CaCO₃ (major constituent of dolomite): 100.0869
- Molecular weight of MgCO₃ (major constituent of dolomite): 84.3142
- Molecular weight of CaO: 56.0774
- Molecular weight of MgO: 40.3044
- Molecular weight of CO₂: 44.0098

- CaCO₃ content = proportion of CaO extractable from dolomite × molecular weight of CaCO₃ / molecular weight of CaO

$$= 34.5\% \times 100.0869 / 56.0774$$

$$= 61.53\%$$
- MgCO₃ content = proportion of MgO extractable from dolomite × molecular weight of MgCO₃ / molecular weight of MgO

$$= 18.3\% \times 84.3142 / 40.3044$$

$$= 38.39\%$$

- Emission factor = molecular weight of CO₂ / molecular weight of CaCO₃ × CaCO₃ content
+ molecular weight of CO₂ / molecular weight of MgCO₃ × MgCO₃ content

$$= 44.0098 / 100.0869 \times 0.6153 + 44.0098 / 84.3142 \times 0.3839$$

$$= 0.2706 + 0.2004$$

$$= 0.4709 \quad [\text{t-CO}_2/\text{t}]$$

$$= 471[\text{kg-CO}_2/\text{t}]$$

● Activity Data

Of the limestone and dolomite consumption data in the Adjusted Price Transaction Table, all limestone and dolomite consumption categorized under 'emissive use,' excluding sectors that correspond to 'Cement Production (2.A.1.)' and 'Lime Production (2.A.2),' i.e., 'Ceramic industry – cement' and 'Ceramic industry - other ceramic, stone, and clay products - quicklime and slaked lime,' will be accounted for. (For dolomite, all sectors excluding 'Ceramic industry – cement'). Activity data is in dry weight, converted using the water content from limestone used for cement.

The sectors in the Adjusted Price Transaction Table corresponding to the five main uses are as follows:

Table 4-7 Main uses and corresponding sectors in the Adjusted Price Transaction Table

Main uses	Corresponding sectors in the Adjusted Price Transaction Table (Limestone)	Corresponding sectors in the Adjusted Price Transaction Table (Dolomite)
Steel/Refining	2611-01 Steel - pig iron to 2611-04 Steel - crude ore (electric furnace)	2611-01 Steel - pig iron to 2631-03 Steel - cast and forged materials (iron)
	2631-02 Steel - cast iron pipe, -03 cast and forged materials (iron)	
	2711-01 Non-ferrous metal - copper, -02 lead and zinc	2711-02 Non-ferrous metal - lead and zinc
	2722-03 Non-ferrous metal - non-ferrous metal cast and forged products	
Glass products	2511-01 Ceramic industry - sheet glass to 2519-09 Ceramic industry - other glass products	2511-01 Ceramic industry - sheet glass/safety glass
Desulfurization of exhaust gas	0621-01 Mining industry - materials for ceramics	
Ceramics products		0621-01 Mining industry - raw minerals for ceramics
		0621-09 Mining industry - other non-metal ore
	2531-01 Ceramic industry - pottery, china and earthenware	2531-01 Ceramic industry - ceramics
	2599-01 Ceramic industry - clay refractories	2599-01 Ceramic industry - refractory, -03 carbon graphite
		2599-09 Ceramic industry - other ceramic, stone, and clay products
		2811-01 Metal Products - metal products for construction use to 2899-09 Metal Products - other metal products
		8611-09 Private services - other amusement and recreation services
Chemical products	2011-02 Chemical Products - chemical fertilizers	2011-02 Chemical Products - chemical fertilizers
	2022-09 Chemical Products - other inorganic chemical industry products	2022-09 Chemical Products - other inorganic chemical industry products
		2039-02 Chemical Products - processed oil and fat products
	2039-09 Chemical Products - other organic chemical industry products	2039-09 Chemical Products - other organic chemical industry products
		2061-01 Chemical Products - medicaments
		2079-09 Chemical Products - other chemical end products

Note: The numbers before the sector names are categorization numbers in the Adjusted Price Transaction Table.

Table 4-8 Amounts of limestone and dolomite consumption

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Limestone consumption								
For Steel and Refinement (dry)	kt	14,415	13,588	13,593	12,542	12,164	10,987	11,814
For Glass Products (dry)	kt	66	42	26	31	16	12	17
For Flue Gas Desulfurization (dry)	kt	2,048	2,157	2,134	2,503	2,334	2,092	2,143
For Ceramic Products (dry)	kt	435	1,108	1,108	424	490	336	320
For Chemical Products (dry)	kt	3,614	1,714	1,725	624	695	471	442
Dolomite consumption								
For Steel and Refinement (dry)	kt	1,144	1,089	1,160	1,530	1,534	1,096	1,575
For Glass Products (dry)	kt	264	250	203	230	160	126	151
For Ceramic Products (dry)	kt	1,561	1,227	1,020	1,130	1,295	1,577	1,615
For Chemical Products (dry)	kt	147	96	84	53	35	36	34

c) Uncertainties and Time-series Consistency● *Uncertainty*

The uncertainty of emission factors for limestone and dolomite were estimated using expert judgment. The uncertainty of emission factors for limestone and dolomite were determined to be 16.4%, 3.5% respectively. The uncertainty for activity data were estimated as 4.8% and 3.9% for limestone and dolomite, respectively, and the uncertainty for emissions were estimated as 17% and 5%, respectively. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series consistency*

Limestone and dolomite consumption data provided in the Adjusted Price Transaction Table is used as limestone and dolomite use activity data for all years from FY1990. The emission factors are constant for all years from FY1990. Therefore, CO₂ emission from limestone and dolomite use has been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

Recalculations have been done for 2008 and 2009 based on updating the activity data for limestone use.

f) Source-specific Planned Improvements

The further improvement of the accuracy of the Adjusted Price Transaction Table will be considered.

4.2.4. Soda Ash Production and Use (2.A.4.)**4.2.4.1. Soda Ash Production (2.A.4.-)**

In Japan, the ammonium chloride soda process is used to produce soda ash (Na₂CO₃). The soda ash production process involves calcinating limestone and coke in a lime kiln, which emits CO₂. Almost all lime-derived CO₂ is stored in the product.

In the soda ash production process, purchased CO₂ is sometimes input through a pipeline, but because these CO₂ emissions are from the ammonia industry, they are already included in “Ammonia Production (2.B.1)”. Also, the coke consumed is listed as that for heating in the Yearbook of the Current Survey of Energy Consumption, and thus CO₂ emissions from coke are already counted under “Fuel Combustion (1.A)”. Therefore all emissions from this source are already included in other categories, and are reported as “IE”. Coke is input as a heat-source and CO₂ source.

The *Revised 1996 IPCC Guidelines* offer a method to calculate CO₂ emissions from calcinating trona (Na₂CO₃-NaHCO₃-2H₂O), but these emissions are not estimated because in Japan soda ash has never been manufactured by trona calcination.

4.2.4.2. Soda Ash Use (2.A.4.-)*a) Source/Sink Category Description*

CO₂ is released during the use of soda ash (Na₂CO₃).

b) Methodological Issues● **Estimation Method**

CO₂ emissions from soda ash use are calculated by multiplying soda ash consumption by the country-specific emission factor.

● **Emission Factors**

Soda ash consumption data categorized under 'for emission purpose' in the Adjusted Price Transaction Table does not differentiate between domestic products and imported products, therefore the emission factor is established by taking a weighted average of the below emission factors for domestic soda ash and imports, by total domestic shipment and total import amounts.

For domestic soda ash, the emission factor is set as follows using data on the purity of soda ash. (The annual fluctuation in purity of soda ash is small, therefore the emission factor will be set constant over the time-series.)

$$\begin{aligned}
 & \text{Emission factor for domestic soda ash} \\
 & = \text{purity of soda ash (arithmetic mean between the 2 domestic companies)} \\
 & \quad \times \text{molecular weight of CO}_2 / \text{molecular weight of Na}_2\text{CO}_3 \\
 & = 0.995 \times 44.01 / 105.99 \\
 & = 0.413
 \end{aligned}$$

For soda ash imported, and other disodium carbonate imported, there is not enough information to set representative emission factors, therefore the default value (0.415 t-CO₂/t-Na₂CO₃) specified in the *Revised 1996 IPCC Guidelines* (vol. 3 p. 2.13) is used continuously.

● **Activity Data**

Soda ash consumption data categorized under 'for emission purpose' in the Adjusted Price Transaction Table is used.

Table 4-9 Soda ash consumption (limestone)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Soda ash consumption (soda ash)	kt	647	605	504	476	384	333	333

c) Uncertainties and Time-series Consistency● **Uncertainty**

For the uncertainty of the emission factor from soda ash use, the lime production value was applied since it is a similar source category to soda ash. For the uncertainty of activity data, 6.3% uncertainty was applied. The uncertainty of CO₂ emissions from soda ash use was estimated as 16%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series consistency**

Soda ash consumption data provided in the Adjusted Price Transaction Table is used as soda ash use activity data for all years from FY1990. The emission factor is constant for all years from FY1990. Therefore, CO₂ emission from soda ash use has been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

Recalculations have been done for 2009 based on updating the activity data.

f) Source-specific Planned Improvements

No improvements are planned.

4.2.5. Asphalt Roofing (2.A.5.)

Asphalt roofing is manufactured in Japan, but information on the manufacturing process and activity data is inadequate, and it is not possible to definitively conclude that CO₂ is not emitted from the manufacturing of asphalt roofing. Emissions have also never been actually measured, and as no default emission value is available, it is not currently possible to calculate emissions. Therefore, it has been reported as “NE”.

4.2.6. Road Paving with Asphalt (2.A.6.)

Roads in Japan are paved with asphalt, but almost no CO₂ are thought to be emitted in the process. It is not possible, however, to be completely definitive about the absence of emissions. Emissions have also never been actually measured, and as no default emission value is available, it is not currently possible to calculate emissions. Therefore, it has been reported as “NE”.

4.3. Chemical Industry (2.B.)

This category covers CO₂, CH₄, and N₂O emissions from the processes of chemical productions.

This section includes GHG emissions from five sources: Ammonia Production (2.B.2), Nitric Acid Production (2.B.2.), Adipic Acid Production (2.B.3.), Carbide Production (2.B.4.), Other (2.B.5.).

In 2010, emissions from Chemical Industry were 3,919 Gg-CO₂ eq. and represented 0.3% of GHG of the Japan's total GHG emissions (excluding LULUCF). The emissions had decreased by 69.4% compared to 1990.

Table 4-10 Emissions from 2.B. Chemical Industry

Gas	Emission sub-category			Units	1990	1995	2000	2005	2008	2009	2010	
CO ₂	2.B Chemical Industry	2.B.1	Ammonia	Gg-CO ₂	3,385	3,436	3,188	2,155	1,990	1,909	2,106	
		2.B.4	Carbide Production	Silicon Carbide	Gg-CO ₂	C	C	C	C	C	C	C
				Calcium Carbide	Gg-CO ₂	C	C	C	C	C	C	C
		2.B.5	Other	Ethylene	Gg-CO ₂	C	C	C	C	C	C	C
	Total				Gg-CO ₂	4,209	4,220	3,893	2,887	2,574	2,488	2,737
CH ₄	2.B Chemical Industry	2.B.4	Carbide Production	Silicon Carbide	Gg-CH ₄	0.02	0.05	0.03	0.03	0.03	0.03	0.03
				Carbon Black	Gg-CH ₄	0.28	0.27	0.27	0.28	0.25	0.22	0.26
		2.B.5	Other	Ethylene	Gg-CH ₄	0.09	0.10	0.11	0.11	0.10	0.11	0.10
				1,2-Dichloroethane	Gg-CH ₄	0.01	0.02	0.02	0.02	0.02	0.02	0.02
				Styrene	Gg-CH ₄	0.07	0.09	0.09	0.10	0.08	0.09	0.09
				Methanol	Gg-CH ₄	0.17	0.15	NO	NO	NO	NO	NO
	Coke				Gg-CH ₄	15.47	13.82	8.00	5.02	4.59	4.13	4.45
	Total				Gg-CH ₄	16.11	14.50	8.52	5.57	5.07	4.60	4.95
Total				Gg-CO ₂ eq.	338	304	179	117	106	97	104	
N ₂ O	2.B Chemical Industry	2.B.2	Nitric Acid	Gg-N ₂ O	2.47	2.46	2.57	2.52	1.62	1.54	1.81	
		2.B.3	Adipic Acid	Gg-N ₂ O	24.20	24.03	12.56	1.68	2.45	3.49	1.66	
	Total				Gg-N ₂ O	26.67	26.49	15.13	4.19	4.07	5.03	3.48
	Total				Gg-CO ₂ eq.	8,267	8,213	4,690	1,300	1,262	1,559	1,078
Total of All Gases				Gg-CO ₂ eq.	12,814	12,737	8,762	4,304	3,943	4,144	3,919	

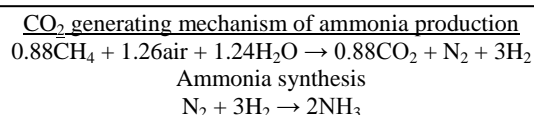
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4.3.1. Ammonia Production (2.B.1.)

a) Source/Sink Category Description

1) CO₂

In ammonia production, CO₂ is emitted when hydrocarbon feedstock is broken down to make H₂.



2) CH₄

Emission of CH₄ from the ammonia production has been confirmed by actual measurements. As there are not enough sufficient examples to enable the establishment of an emission factor, it is not currently possible to calculate emissions. The *Revised 1996 IPCC Guidelines* also do not give a default emission factor. Therefore, CH₄ was reported as “NE”.

3) N₂O

Emission of N₂O from ammonia production is theoretically impossible, and given that even in actual measurements the emission factor for N₂O is below the limits of measurement, N₂O was reported as “NA”.

b) Methodological Issues

● Estimation Method

CO₂ emissions are calculated by multiplying the amount of fuels consumed as ammonia feedstock by emission factors.

● Emission Factors

The same emission factors that are used to calculate CO₂ emissions from the fuel combustion sector (Chapter 3) are used for each feedstock listed in Table 4-11. It should be noted that the implied

emission factor changes every year, since the composition of the feedstocks consumed for ammonia production varies annually.

Table 4-11 Emission factors and calorific values of feedstocks used when producing ammonia

Feedstock	Emission Factors (tC/TJ)	Calorific value		(Units)
		1990	2005	
Naphtha	18.17	33.5	33.6	MJ/l
Liquefied petroleum gas (LPG)	16.13	50.2	50.8	MJ/kg
Petroleum-derived hydrocarbon gases (petrochemical offgases)	14.15	39.3	44.9	MJ/m ³
Natural gas	13.90	41.0	43.5	MJ/m ³
Coal (thermal coal, imports)	24.71	26.0	25.7	MJ/kg
Petroleum coke	25.35	35.6	29.9	MJ/kg
Liquefied natural gas (LNG)	13.47	54.4	54.6	MJ/kg
Coke oven gas (COG)	10.99	20.1	21.1	MJ/m ³

(Reference) General Energy Statistics, Agency for Natural Resources and Energy

● Activity Data

The fixed units (including weight and volume) for the fuel types in Table 4-12 below, which are from the Ministry of Economy, Trade and Industry's Yearbook of the Current Survey of Energy Consumption, were converted using the calorific values in the Agency for Natural Resources and Energy's General Energy Statistics, and results were used as activity data. Consumption data on some fuel types are confidential.

Table 4-12 Amount of feedstocks used for ammonia production

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Naphtha	kl	189,714	477,539	406,958	92,453	67,062	72,045	70,067
LPG	t	226,593	45,932	5,991	0	0	0	0
Off gas	10 ³ m ³	C	230,972	240,200	147,502	151,553	140,783	143,634
Natural Gas	10 ³ m ³	C	100,468	86,873	77,299	50,260	21,773	41,640
Coal	t	C	209,839	726	1,239	802	522	629
Oil Coke	t	C	273,125	420,862	353,983	336,633	351,594	394,116
LNG	t	C	46,501	23,395	165,606	162,342	145,699	157,918
COG	10 ³ m ³	C	35,860	55,333	0	0	0	0

C: Confidential

● Point to Note

Fuel consumption in this category has been deducted from energy sector activity data (see Chapter 3).

c) Uncertainties and Time-series Consistency

● Uncertainty

The uncertainty of each fuel was estimated. For the uncertainty of emission factors, the values given in Chapter 3 were applied. The standard value, 5%, given by the Committee for the Greenhouse Gas Emission Estimation Methods was used. As a result, the uncertainty of emissions from the fuels are of the following: naphtha 7%; LPG 6%; hydrocarbon gas 22%; natural gas 7%; coal (steam coal, imported coal) 7%; petroleum coke 23%; LNG 10%; and COG 25%. The uncertainty assessment methods are summarized in Annex 7.

- **Time-series Consistency**

For activity data, the same sources are used throughout the time series, from the Current Survey of Energy Consumption. The emission factor is constantly based on the General Energy Statistics throughout the time series. Therefore, CO₂ emission from ammonia production has been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

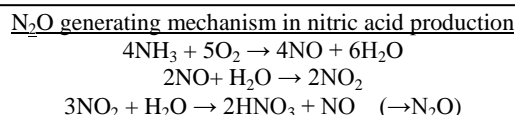
f) Source-specific Planned Improvements

No improvements are planned.

4.3.2. Nitric Acid Production (2.B.2.)

a) Source/Sink Category Description

N₂O is emitted when nitric acid (HNO₃) is produced from ammonia.



In Japan, the main processes used in nitric acid production are the New Fauser Process (medium pressure) and Chemico Process (high pressure), both based on the Ostwald chemical process. With regard to N₂O decomposition, there are catalytic decomposition units in operation.

b) Methodological Issues

- **Estimation Method**

N₂O emissions were estimated by multiplying the nitric acid production amount by an emission factor, based on the method given in *GPG (2000)* (page 3.31, Equation 3.9). Since emissions data for individual factories is confidential, the nitric acid production amount and the emission factor were set for Japan's total production. Due to the current lack of data on the amount of N₂O destroyed, the equation has no term for destruction.

<p><u>N₂O emissions (kg-N₂O) from nitric acid production</u></p> <p>= emission factor [kg-N₂O/t] × nitric acid production volume [t]</p>

- **Emission Factors**

Because data for individual factories are confidential, the emission factor was set by using each factory's nitric acid production amount to find the weighted average of Japan's 10 nitric acid producing factories' emission factors (measurement data). These emission factors take N₂O recovery and destruction into account.

Table 4-13 N₂O emission factors for nitric acid production

Item	Unit	1990	1995	2000	2005	2008	2009	2010
EF for Nitric Acid Production	kg-N ₂ O/t	3.50	3.51	3.92	4.18	3.35	3.34	3.58

- **Activity Data**

Production amounts of nitric acid are directly provided by the Ministry of Economy, Trade and Industry.

Table 4-14 Amount of Nitric acid production

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Nitric Acid Production	t	705,600	701,460	655,645	602,348	484,070	460,600	506,071

c) Uncertainties and Time-series Consistency

- **Uncertainty**

The uncertainty of the emission factor was estimated using a 95% confidence interval for emission factors. For the uncertainty of activity data, the standard value of 5% given by the Committee for the Greenhouse Gas Emission Estimation Methods was used. As a result, the uncertainty of emissions was estimated as 46%. The uncertainty assessment methods are summarized in Annex 7.

- **Time-series Consistency**

Emissions throughout the time series are consistently estimated using the activity data and emission factors provided by the Ministry of Economy, Trade and Industry.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.3.3. Adipic Acid Production (2.B.3.)

a) Source/Sink Category Description

N₂O is emitted in the adipic acid (C₆H₁₀O₄) production process through the reaction of cyclohexanone, cyclohexanol, and nitric acid.

b) Methodological Issues

- **Estimation Method**

Emissions were estimated using the N₂O generation rates, N₂O decomposition amount, and adipic acid production amount of the relevant operating sites, in accordance with the *GPG (2000)* decision tree (Page 3.32, Fig. 3.4).

N₂O emissions from adipic acid production

$$= [\text{N}_2\text{O generation rate} \times (1 - \text{N}_2\text{O generation rate} \times \text{decomposition unit operation rate})] \times \text{adipic acid production rate}$$

- **Emission Factors**

Values calculated using the above equation has been used as the emission factors. Parameters were established by the following methods. Relevant data used in estimation is confidential.

- **Rate of generation of nitrous oxide**

Actual measurement data provided from the sole producer of adipic acid as an end product in Japan.

- **Rate of decomposition of nitrous oxide**

The figure used is the result of measurement of the rate of decomposition of nitrous oxide in the operating site.

- **Operating rate of decomposition unit**

A full-scale survey on the number of operation hours is conducted annually for N₂O decomposition units and adipic acid production plants. The operating rate is based on this survey.

Calculation of operating ratio of decomposition unit

$$\begin{aligned} & \text{Operating ratio of decomposition unit (\%)} \\ &= \text{Number of hours of decomposition unit in operation} \\ & / \text{Number of hours of adipic acid production plants in operation} \times 100 (\%) \end{aligned}$$

Number of hours of decomposition unit in operation:

Hours starting from the beginning of feeding the entire volume of N₂O gases until the end of feeding

Number of hours of adipic acid production plants in operation:

Hours starting from the beginning of feeding materials until the end of feeding

- **Activity Data**

The activity data for nitrous oxide emissions associated with the manufacturing of adipic acid is the amount of adipic acid produced provided to the Ministry of Economy, Trade and Industry by the manufacturer. Relevant data used in estimation is confidential.

- **Point to Note**

From 1990 to 1997, N₂O emissions from adipic acid production increased gradually. However, N₂O decomposition units were installed in adipic acid production plants in March 1999, and emissions since then have decreased dramatically. There was a temporary growth in the emissions in 2000 due to the low operating ratio of N₂O decomposition units caused by a breakdown of the decomposition units.

c) **Uncertainties and Time-series Consistency**

- **Uncertainty**

The uncertainty of the emission factor for adipic acid was estimated by combining the uncertainty of the N₂O generation rate, N₂O decomposition rate, and the operating rate of the decomposition unit. As a result, the uncertainty of the emission factor was estimated as 9%. A 2% uncertainty given by

the *GPG (2000)* was applied for activity data. As a result, the uncertainty for adipic acid was estimated as 9%. The uncertainty assessment methods are summarized in Annex 7.

- ***Time-series Consistency***

Activity data and emission factors consistently provided by the producer of adipic acid are used to estimate emissions throughout the time series.

- d) ***Source-specific QA/QC and Verification***

See section 4.2.1. d) .

- e) ***Source-specific Recalculations***

There have been no source-specific recalculations.

- f) ***Source-specific Planned Improvements***

No improvements are planned.

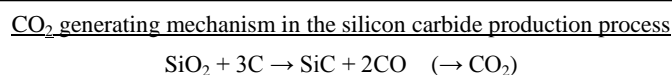
4.3.4. Carbide Production (2.B.4.)

4.3.4.1. Silicon Carbide Production (2.B.4.-)

- a) ***Source/Sink Category Description***

- 1) ***CO₂***

CO₂ is emitted by the reaction of petroleum coke with silica as raw materials in the production of silicon carbide.



- 2) ***CH₄***

In Japan, silicon carbide is produced in electric arc furnaces, and it is believed that CH₄ is generated from the oxidation of coke, which is used as a reducing agent in silicon carbide production.

- b) ***Methodological Issues***

- 1) ***CO₂***

- ***Estimation Method***

Emissions are calculated by multiplying the amount of petroleum coke used as silicon carbide feedstock by an emission factor.

- ***Emission Factors***

Because Japan does not have measurement data or emission factor data, the default value 2.3 [t-CO₂/t] for silicon carbide production in the *Revised 1996 IPCC Guidelines* (vol. 3 p. 2.21) is used.

- **Activity Data**

The activity data for CO₂ emissions from silicon carbide production is the amount of petroleum coke consumed, which is provided by Japan's only silicon carbide production facility. The data is confidential.

2) CH₄

- **Estimation Method**

Emissions were calculated by multiplying an emission factor based on actual figures obtained in Japan by the energy consumption of electric arc furnaces. This is the same method used for calculating CH₄ emissions in the Fuel Combustion Sector (1.A. Solid Fuels).

- **Emission Factors**

The emission factor of energy consumption in electric arc furnaces (12.8 kg-CH₄/TJ) was determined by using the formula for calculating fuel combustion and actual data from Japanese measurement surveys of CH₄ concentrations in gas ducts, concentrations of O₂ and theoretical flue gas amounts (dry), theoretical air demand, and high calorific values. See Chapter 3 3.2.1 Stationary Combustion (1.A.1., 1.A.2., 1.A.4.: CH₄ and N₂O)

- **Activity Data**

Energy consumption amounts included in the "electric furnace" category for the iron and steel industries of the General Survey of the Emissions of Air Pollutants were used. (From 2000 and onward, 1999 values are used.)

Table 4-15 Energy consumption from electric arc furnaces (for carbide)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Furnaces (for Carbide)	TJ	1,576	4,277	2,454	2,454	2,454	2,454	2,454

c) Uncertainties and Time-series Consistency

- **Uncertainty**

1) CO₂

For the uncertainty of the CO₂ emission factor, 100% was applied as provided by the *GPG (2000)* for a similar category. For the uncertainty of activity data, the standard value of 10% given by the Committee for the Greenhouse Gas Emission Estimation Methods was used. The uncertainty assessment methods are summarized in Annex 7.

2) CH₄

The uncertainty of the CH₄ emission factor and activity data were estimated as 163% and 5%, respectively, as estimated in Chapter 3. The uncertainty for emissions is estimated as 163%. The uncertainty assessment methods are summarized in Annex 7.

- **Time-series Consistency**

For CO₂ and CH₄ activity data, the same sources are consistently used throughout the time series-the former from the manufacturing facility, and the latter from the General Survey of the Emissions of Air

Pollutants. The emission factors for both gases are constant throughout the time series. Therefore, CO₂ and CH₄ emissions from silicon carbide have been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

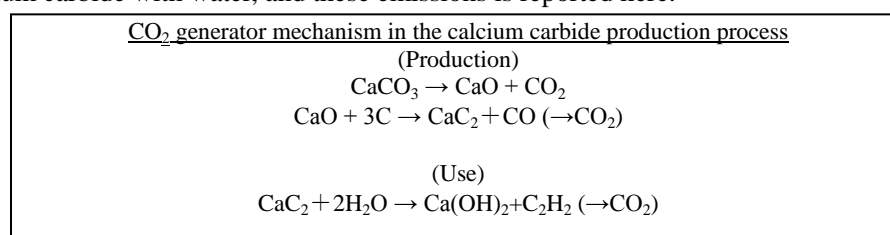
The use of fuel consumption data in the General Survey of the Emissions of Air Pollutants for FY 2002 onward was prohibited for any purposes other than the original one specified for the General Survey of the Emissions of Air Pollutants, while that is not the case with the data in the General Survey of the Emissions of Air Pollutants for FY 1999 and earlier years. The use of General Survey of the Emissions of Air Pollutants in the GHG inventory was added to the purpose of the General Survey of the Emissions of Air Pollutants by the current examination toward the reuse of the General Survey of the Emissions of Air Pollutants and was recently officially accepted. Japan will continue to consider applying the latest the General Survey of the Emissions of Air Pollutants data in the future inventory.

4.3.4.2. Calcium Carbide Production and Use (2.B.4.-)

a) Source/Sink Category Description

1) CO₂

CO₂ is generated in the process of making the quicklime, and is also emitted by the combustion of CO occurring from calcium carbide production. However, the former is included in emissions from Chemical Products in “Limestone and Dolomite Use (2.A.3.),” therefore only reducing agent-origin emissions are accounted for here. Further, CO₂ is generated by the combustion of acetylene, which is generated by reacting calcium carbide with water, and these emissions is reported here.



2) CH₄

Byproduct gases (mainly CO) generated in carbide production include a small amount of CH₄, all of which is recovered and burned as fuel, with none being emitted outside the system. Therefore emissions from this source are reported as “NA”.

b) Methodological Issues

● **Estimation Method**

CO₂ emissions are calculated by multiplying calcium carbide production by the following emission

factor, based on the *Revised 1996 IPCC Guidelines*.

- **Emission Factors**

For years FY1990 to 2007, because Japan does not have measurement data or emission factor data, the default value in the *Revised 1996 IPCC Guidelines* is used.

Table 4-16 CO₂ Emission factors for calcium carbide production and consumption (FY1990-2007)

Units	From reducing agent in production	From use
t-CO ₂ /t	1.09	1.10

Source: *Revised 1996 IPCC Guidelines*, vol. 3, p. 2.22.

For years after FY2008, country-specific emission factors from reducing agents during production (changes annually) are used, which are based on measurement data from the two calcium carbide producing companies in Japan. These emission factors are confidential.

The default emission factor (1.10 t-CO₂/t) for calcium carbide use is also used for FY2008 and onwards.

- **Activity Data**

Calcium carbide production data provided by the Carbide Industry Association are used as the calcium carbide production amount. The data are confidential.

c) Uncertainties and Time-series Consistency

- **Uncertainty**

For the uncertainty of the CO₂ emission factor, 100% was applied as provided by the *GPG (2000)* for a similar category. For the uncertainty of activity data, the standard value of 10% given by the Committee for the Greenhouse Gas Emission Estimation Methods was used. As a result, the uncertainty for CO₂ emissions from calcium carbide was estimated as 100%. The uncertainty assessment methods are summarized in Annex 7.

- **Time-series Consistency**

For activity data, the same sources are used throughout the time series. The emission factor is constant from 1990 to 2007 and for years after 2008, the country-specific emission factor will be used. This is because there is no data available on emission factors for previous years, and because emission factors may fluctuate over time due to changes in scale of production or improvements in manufacturing technology, therefore the default emission factors will be used for FY1990 to FY2007.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations

f) Source-specific Planned Improvements

No improvements are planned.

4.3.5. Other (2.B.5.)

4.3.5.1. Carbon Black Production (2.B.5.-)

a) Source/Sink Category Description

Carbon black is made by breaking down acetylene, natural gas, oil mist, and other feedstocks by incomplete combustion at 1,300°C or higher. The CH₄ in the tail gas (offgas) emitted from the carbon black production process is released into the atmosphere.

b) Methodological Issues

● Estimation Method

CH₄ emissions from carbon black production are calculated by multiplying the carbon black production amount by Japan's country-specific emission factor, in accordance with the *Revised 1996 IPCC Guidelines*.

● Emission Factors

Five major companies, providing 96% of domestic production, recover CH₄ generated in the carbon black production processes and use it in recovery furnaces and flare stacks. Therefore, there are no emissions during normal operation. The emission factor was established by estimating emissions of CH₄ during routine inspections and the boiler inspection carried out by the five major domestic producers, and taking a weighted average by using production amounts of carbon black. The emission factor is 0.35 [kg-CH₄/t].

Table 4-17 CH₄ emissions and carbon black production by five main domestic producers

	Carbon black production [t/year]	CH ₄ emissions [kg-CH ₄ /year]	Emission factor [kg-CH ₄ /t]
Total from five main companies	701,079	246,067	0.35

Source: Data provided by the Carbon Black Association (1999 actual results)

● Activity Data

Carbon black production amounts given in the Yearbook of Chemical Industries Statistics compiled by the Ministry of Economy, Trade and Industry were used for activity data for CH₄ emissions associated with the manufacturing of carbon black.

Table 4-18 Carbon black production amount

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Carbon Black Production	t	792,722	758,536	771,875	805,461	725,113	634,733	730,352

c) Uncertainties and Time-series Consistency

● Uncertainty

The uncertainty for the emission factor for carbon black was calculated by finding the 95% confidence interval of emission factors. The estimated uncertainty was 54.8%. For the uncertainty of activity data, the standard value of 5% given by the Committee for the Greenhouse Gas Emission Estimation Methods was used. As a result, the uncertainty of carbon black production emissions was estimated at 55%. The uncertainty assessment methods are summarized in Annex 7.

- **Time-series Consistency**

For activity data, the same source-the Yearbook of Chemical Industries Statistics are used throughout the time series. The emission factor is constant throughout the time series. Therefore, CH₄ emissions from carbon black production have been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

The possibility of double counting of CH₄ from furnaces in the Energy sector should be investigated.

4.3.5.2. Ethylene Production (2.B.5.-)

a) Source/Sink Category Description

1) CO₂, CH₄

CO₂ is emitted when it is separated in the ethylene production process. CH₄ is emitted by naphtha cracking through steam cracking in the ethylene production process.

2) N₂O

There is almost no nitrogen contained in naphtha, the raw material of ethylene, and the ethylene production process takes place under conditions that are almost completely devoid of oxygen. Emissions are reported as “NA” in accordance with the judgment of experts that theoretically there are no N₂O emissions.

b) Methodological Issues

- **Estimation Method**

CH₄ and CO₂ emissions from ethylene production were calculated by multiplying ethylene production by Japan’s country-specific emission factor, in accordance with the *Revised 1996 IPCC Guidelines*.

- **Emission Factors**

- **CO₂**

The emission factor was set, based on a survey conducted by the Japan Petrochemical Industry Association in 2009 on the CO₂ emission factor from ethylene production. This emission factor is confidential.

- **CH₄**

Estimates of amount of exhaust gas from flare stacks at a normal operation and an unsteady operation at operating sites in Japan (assuming that 98% of the amount that enters is combusted), and measured amount of exhaust gas from naphtha cracking furnaces and furnaces heated by re-cycled gas, were

divided by the production amount to calculate emission factors for each company. The weighted average based on production from each company was then applied to establish the emission factor of 0.015 [kg-CH₄/t]. (Surveyed by the Japan Petrochemical Industry Association)

● **Activity Data**

Ethylene production amounts from the Yearbook of Chemical Industries Statistics compiled by the Ministry of Economy, Trade and Industry were used as activity data for emissions of CH₄ and CO₂ from ethylene production.

Table 4-19 Ethylene production amount

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Ethylene Production	kt	5,966	6,951	7,566	7,549	6,520	7,219	6,999

c) Uncertainties and Time-series Consistency

● **Uncertainty**

The uncertainty for both CO₂ and CH₄ emission factors for ethylene were calculated by finding the 95% confidence interval of emission factors, based on the decision tree for uncertainty assessment. The estimated uncertainty for both CO₂ and CH₄ were 77.2%. For the uncertainty of activity data, the standard value of 5% given by the Committee for the Greenhouse Gas Emission Estimation Methods was used. As a result, the uncertainty for both CO₂ and CH₄ were estimated as 77%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

For activity data, the same sources are used throughout the time series. The emission factor is constant throughout the time series. Therefore, CO₂ and CH₄ emissions from ethylene production have been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.3.5.3. 1,2-Dichloroethane (2.B.5.-)

a) Source/Sink Category Description

1,2-dichloroethane (Ethylene Dichloride) is manufactured by reacting ethylene (C₂H₄) and chlorine (Cl₂). The product then passes through washing, refining, and thermolysis processes to become a vinyl chloride monomer (C₂H₃Cl). A very small amount of CH₄ is contained in the exhaust gases of the reaction, and of the washing and refining processes.

b) Methodological Issues● **Estimation Method**

CH₄ emissions from 1,2-dichloroethane production are calculated by multiplying production amount by Japan's country-specific emission factor, in accordance with the *Revised 1996 IPCC Guidelines*.

● **Emission Factors**

The concentration of CH₄ in waste gas from three member companies of the Vinyl Environmental Council (representing approximately 70% of total 1,2-dichloroethane production in Japan) was measured, and a weighted average was calculated to establish the emission factor. The emission factor is 0.0050 [kg-CH₄/t]. (Surveyed by the Vinyl Environmental Council)

● **Activity Data**

1,2-Dichloroethane production amounts from the Yearbook of Chemical Industries Statistics compiled by the Ministry of Economy, Trade and Industry were used as activity data for CH₄ emissions from 1,2-dichloroethane production.

Table 4-20 1,2-Dichloroethane production amount

Item	Unit	1990	1995	2000	2005	2008	2009	2010
1,2-Dichloroethane Production	kt	2,683	3,014	3,346	3,639	3,243	3,213	3,155

c) Uncertainties and Time-series Consistency● **Uncertainty**

The uncertainty of the CH₄ emission factor for 1,2-dichloroethane production were estimated by finding the 95% confidence interval, based on expert judgment. The uncertainty was estimated as 100.7%. For the uncertainty of activity data, the standard value of 5% given by the Committee for the Greenhouse Gas Emission Estimation Methods was used. As a result, the uncertainty of 1,2-dichloroethane production was estimated as 101%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

For activity data, the same sources are used throughout the time series. The emission factor is constant throughout the time series. Therefore, CH₄ emissions from 1,2-Dichloroethane production have been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.3.5.4. Styrene Production (2.B.5.-)

a) Source/Sink Category Description

CH₄ is emitted in the styrene production process.

b) Methodological Issues

● Estimation Method

CH₄ emissions from styrene production were calculated by multiplying styrene production amount by Japan's country-specific emission factor, based on the method given in the *Revised 1996 IPCC Guidelines*.

● Emission Factors

Estimates of amount of exhaust gas from flare stacks at a normal operation and an unsteady operation at operating sites in Japan (assuming that 98% of the amount that enters is combusted), and measured amount of waste gas from heating furnaces, were divided by the production amount to calculate emission factors for each company. The weighted average by production from each company was then applied to establish the emission factor. The emission factor is 0.031 [kg-CH₄/t]. (Surveyed by the Japan Petrochemical Industry Association)

● Activity Data

Styrene monomer production amounts from the Yearbook of Chemical Industries Statistics compiled by the Ministry of Economy, Trade and Industry were used as activity data for CH₄ emissions from styrene production.

Table 4-21 Styrene production amount

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Styrene Production	kt	2,227	2,952	3,020	3,375	2,699	3,043	3,019

c) Uncertainties and Time-series Consistency

● Uncertainty

The uncertainty for the CH₄ emission factor for styrene production was estimated by finding the 95% confidence interval of emission factors, based on the decision tree for uncertainty assessment. The estimated uncertainty was 113.2%. For the uncertainty of activity data, the standard value of 5% given by the Committee for the Greenhouse Gas Emission Estimation Methods was used. As a result, the uncertainty of emissions was estimated as 113%. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

For activity data, the same sources are used throughout the time series. The emission factor is constant throughout the time series. Therefore, CH₄ emissions from styrene production have been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.3.5.5. Methanol Production (2.B.5.-)**a) Source/Sink Category Description**

CH₄ is emitted in the production of methanol.

b) Methodological Issues● **Estimation Method**

CH₄ emissions from methanol production are calculated using the method given in the *Revised 1996 IPCC Guidelines*.

According to industry organizations, the production (synthesis) of methanol stopped in Japan in 1995 due to the price difference with overseas methanol. Since then all methanol has been imported, and methanol production plants disappeared from Japan in about 1995. According to the Yearbook of Chemical Industries Statistics, beginning in 1997 there is also no production of refined methanol. The methanol refining process merely dewateres the synthesized methanol, therefore, theoretically no CH₄ is generated.

Accordingly, from 1990 to 1995, emissions are reported using the production amounts in industry organization statistics. For 1996 and thereafter, emissions are reported as “NO” because it is assumed that methanol has not been produced (synthesized) since 1995.

● **Emission Factors**

The default value for methanol given in the *Revised 1996 IPCC Guidelines* was used. The emission factor is 2 [kg-CH₄/t] (Refer to *Revised 1996 IPCC Guidelines* Vol. 2 p 2.22, Table 2-9).

● **Activity Data**

Production amounts of methanol (on calendar year basis) given by the Methanol and Formalin Association were used as activity data for CH₄ emissions from methanol production.

Table 4-22 Methanol production amount

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Methanol Production	t	83,851	75,498	NO	NO	NO	NO	NO

c) Uncertainties and Time-series Consistency● **Uncertainty**

The uncertainty is not estimated.

● **Time-series Consistency**

For activity data, the same sources are used throughout the time series. The emission factor is constant throughout the time series. Therefore, CH₄ emissions from methanol production have been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.3.5.6. Coke Production (2.B.5.-)**a) Source/Sink Category Description****1) CO₂**

This category is reported as “IE” because the emissions of CO₂ from coke production are included in the coal products and production section of the Fuel Combustion Sector (1.A.).

2) CH₄

CH₄ is emitted in coke production.

3) N₂O

We have no measurements of the concentration of N₂O in the gas leaking from coking furnace lids, but N₂O emissions from this source are reported as “NA,” the reason being that experts say that N₂O is likely not produced because the atmosphere in a coke oven is normally at least 1,000°C, and is reducing.

b) Methodological Issues● **Estimation Method**

CH₄ emissions from coke production were calculated by multiplying coke production amount by Japan’s country-specific emission factor, based on the method given in the *Revised 1996 IPCC Guidelines*.

● **Emission Factors**

CH₄ emissions from coke production come from two sources: CH₄ in combustion exhaust gas from gas leakage from the carbonization chamber to the combustion chamber, and CH₄ emitted from the coking furnace lid, the desulfurization tower, or the desulfurization recycling tower, in the carbonization process of coal.

➤ **Combustion exhaust gas**

The concentration of CH₄ in the exhaust gas from coking furnaces operated by five companies at seven operating sites (surveyed by the Japan Iron and Steel Federation, actual results for FY1999) was weighted by the production amount of coke to derive a weighted average, which was established as the emission factor. The emission factor is 0.089 [kg-CH₄/t].

➤ **Coking furnace lid, desulfurization tower, and desulfurization recycling tower**

The Japan Iron and Steel Federation has had a voluntary plan in place since fiscal year 1997 to manage noxious atmospheric pollutants, and CH₄ emissions have been estimated from emissions of other substances from the lid of coking furnaces. The emission factor has been established by taking a weighted average using this data and the amount of production of coke.

Table 4-23 Emission factor of CH₄ from coking furnace lids, desulfurization towers, and desulfurization recycling towers

Item	Unit	1990-1996	1997-1999	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010
CH ₄ EFs	[kg-CH ₄ /t]	0.238	0.180	0.119	0.062	0.052	0.042	0.055	0.043	0.039	0.040	0.037	0.032	0.031

* Emission factor change is assumed to be small for FY1990-1996, therefore actual data values for FY1995 is used for other years with no data. For FY1997-1999, it is assumed that values for 1998 and 1999 are the same as those of 1997. For FY2000 and on, actual data values are adopted.

Source: Japan Iron and Steel Federation data

➤ CH₄ emission factor for coke production

The aforementioned Combustion Exhaust Gas and Coking Furnace Lids, Desulfurization Towers, and Desulfurization Recycling Towers have been added, and the resulting figure has been used as the emission factor.

● Activity Data

As the activity of CH₄ emissions from coke production, the inventory used the coke production amount given in the Yearbook of Mineral Resources and Petroleum Products Statistics (previously the Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke) compiled by the Ministry of Economy, Industry and Trade.

Table 4-24 Coke production amount

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Coke Production	kt	47,338	42,279	38,511	38,009	36,551	34,140	37,036

● Completeness

The SBDT¹ (Table 2(I).A-Gs2) in the CRF requires emissions of CO₂ and CH₄ from coke production to be reported as a sub-category of 2.C.1. Steel Manufacture, but coke is also manufactured in Japan in industries other than the steel industry. The emissions have therefore been counted in this category.

c) Uncertainties and Time-series Consistency

● Uncertainty

For the uncertainty of the emission factor for coke production, the uncertainty of fuel combustion emissions from the coking furnace and coking furnace lids were estimated separately. The uncertainty of fuel combustion emissions from the coking furnace and coking furnace lids was estimated as 98.5% and 61.8%, respectively. For the uncertainty of activity data, the standard value of 5% given by the Committee for the Greenhouse Gas Emission Estimation Methods was used. The uncertainty assessment methods are summarized in Annex 7.

¹ SBDT: Sectoral Background Data Table

● **Time-series Consistency**

For activity data, the same sources are used throughout the time series. The emission factor is based on the information provided by the Japan Iron and Steel Federation estimated using a consistent methodology throughout the time series. Therefore, CH₄ emissions from coke production have been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.4. Metal Production (2.C.)

This category covers CO₂, CH₄, PFCs and SF₆ emissions from the manufacturing processes of metal products.

This section includes GHG emissions from three sources: Iron and Steel Production (2.C.1), Ferroalloys Production (2.C.2.), Aluminium Production (2.C.3.), and SF₆ Used in Aluminium and Magnesium Foundries (2.C.4.).

In 2010, emissions from Metal Production were 493 Gg-CO₂ eq. and represented 0.04% of GHG of the Japan's total GHG emissions (excluding LULUCF). The total emissions of CO₂ and CH₄ from this category had decreased by 53.5% compared to 1990. The total of halocarbons and SF₆ had increased by 68.2% compared to 1995.

Table 4-25 Emissions from 2.C. Metal Production

Gas	Emission sub-category			Units	1990	1995	2000	2005	2008	2009	2010	
CO ₂	2.C Metal Production	2.C.1	Iron and Steel Production	Use of Electric Arc Furnaces in Steel Production	Gg-CO ₂	356.09	357.22	248.42	241.93	155.77	111.99	159.86
CH ₄	2.C Metal Production	2.C.1	Iron and Steel Production	Use of Electric Arc Furnaces in Steel Production	Gg-CH ₄	0.74	0.72	0.67	0.68	0.61	0.51	0.59
		2.C.2	Ferroalloys Production		Gg-CH ₄	0.19	0.14	0.13	0.13	0.11	0.11	0.12
	Total				Gg-CH ₄	0.92	0.85	0.80	0.80	0.72	0.62	0.71
	Total				Gg-CO ₂ eq.	19.36	17.92	16.84	16.89	15.02	12.96	14.87
Total of Gases					Gg-CO ₂ eq.	375.45	375.15	265.26	258.81	170.80	124.95	174.73
Gas	Emission sub-category			Units	1990	1995	2000	2005	2008	2009	2010	
PFCs	2.C Metal Production	2.C.3	Aluminium Production		Gg-CO ₂ eq.		69.74	17.78	14.80	14.67	11.02	10.38
SF ₆	2.C Metal Production	2.C.4	SF ₆ Used in Aluminium and Magnesium Foundries		t		5.00	43.00	48.42	27.30	10.00	12.88
					Gg-CO ₂ eq.		119.50	1,027.70	1,157.31	652.47	239.00	307.90
Total of Gases					Gg-CO ₂ eq.		189.24	1,045.48	1,172.11	667.14	250.02	318.28

4.4.1. Iron and Steel Production (2.C.1.)

4.4.1.1. Steel Production (2.C.1.-)

1) CO₂

Coke oxidizes when it is used as a reduction agent in steel production, and CO₂ is generated. The amount of coke used has been included under consumption of fuel in the Fuel Combustion Sector (1.A.), and the CO₂ generated through the oxidization of coke used as a reducing agent has already been calculated under Fuel Combustion Sector (1.A.). Therefore, it has been reported as “IE”.

4.4.1.2. Pig Iron Production (2.C.1.-)

1) CO₂

CO₂ generated from pig iron production is emitted when coke is used as a reduction agent. The amount of coke used has been included under consumption of fuel in the Fuel Combustion Sector (1.A.), and the CO₂ generated through the oxidization of coke used as a reducing agent has already been calculated under Fuel Combustion Sector (1.A.). Therefore, it has been reported as “IE”.

2) CH₄

It is theoretically impossible for CH₄ generation in association with pig iron production, and it has been confirmed that CH₄ is not emitted from actual measurements. Therefore, emissions have been reported as “NA”.

4.4.1.3. Sinter Production (2.C.1.-)

1) CO₂

CO₂ generated when making sinter is all generated by the combustion of coke fines; these emissions come under the Fuel Combustion Sector (1.A.). As they are already calculated in this 1.A. sector, they are reported as “IE”.

CO₂ emissions from limestone and dolomite used when making sinter are counted under “4.2.3. Limestone and Dolomite Use”.

2) CH₄

CH₄ generated when making sinter is all generated by the combustion of coke fines; these emissions come under the Fuel Combustion Sector (1.A.). As they are already calculated in this sector, they are reported as “IE”.

4.4.1.4. Coke Production in Iron and Steel Production (2.C.1.-)

1) CO₂

Coke is mainly produced in iron and steel production in Japan. This category is reported as “IE” because the emissions of CO₂ from coke production are included in the coal products and production section of the Fuel Combustion Sector (1.A.).

2) CH₄

Emissions of CH₄ were calculated at 4.3.5.6. Coke (2.B.5.-), and have been reported as “IE”.

4.4.1.5. Use of Electric Arc Furnaces in Steel Production (2.C.1.-)

a) Source/Sink Category Description

CO₂ is emitted from carbon electrodes when using electric arc furnaces to make steel. CH₄ is also emitted from electric arc furnaces during steel production.

b) Methodological Issues

1) CO₂

● Estimation Method

CO₂ emissions from arc furnaces for steel production are estimated by amount of carbon calculated by weight of production and import of carbon electrodes minus weight of export of carbon electrodes. This difference of the carbon is assumed to be diffused to the atmosphere as CO₂. The carbon included in electric furnaces gas given in the General Energy Statistics are subtracted from the CO₂ emission in this source since these emissions are included in category 1.A fuel combustion.

● Activity Data

Production of carbon electrodes given in Yearbook of Ceramics and Building Materials Statistics compiled by the Ministry of Economy, Trade and Industry, and import and export of carbon electrodes given in Trade Statistics of Japan, Ministry of Finance are used.

Table 4-26 CO₂ emission from carbon electrodes of furnaces

	Unit	1990	1995	2000	2005	2008	2009	2010
#A Import	t	12,341	18,463	11,363	15,075	15,116	11,218	17,321
#B Domestic production	t	211,933	186,143	184,728	216,061	201,256	169,545	205,081
#C Export	t	87,108	92,812	107,998	138,409	134,509	116,489	139,757
#D Electric furnaces gas	t	39,983	14,300	20,293	26,700	39,349	33,709	39,017
Domestic consumptions (#A + #B - #C - #D)	t	97,184	97,493	67,800	66,028	42,514	30,564	43,629
CO ₂ emissions	Gg-CO ₂ eq.	356	357	248	242	156	112	160

2) CH₄

● Estimation Method

Emissions were calculated by multiplying an emission factor based on actual measurements obtained in Japan by the energy consumption of electric arc furnaces. This is the same method used for calculating CH₄ emissions in the Fuel Combustion Sector (1.A. Solid Fuels).

● Emission Factors

The emission factor of energy consumption of electric arc furnaces (12.8 kg-CH₄/TJ) was determined by using the data from actual measurement surveys. (See Chapter 3, 3.2.1 and Chapter 4, 4.3.4.1)

● **Activity Data**

Energy consumption amounts included in the "electric furnace" category for the iron and steel industries of the General Energy Statistics were used.

Table 4-27 Energy consumption from electric arc furnaces

Consumption	Unit	1990	1995	2000	2005	2008	2009	2010
Furnaces	TJ	57,564	55,986	52,457	52,747	47,316	39,753	45,800

c) Uncertainties and Time-series Consistency

1) *CO₂*

● **Uncertainty**

Because all CO₂ from electric arc furnaces are assumed to escape into the atmosphere, no emission factor has been set. Therefore, by assessing the uncertainty for activity data the uncertainty for emissions is assessed. As a result of combining the uncertainties of the parameters for activity data, the uncertainty was estimated as 4.5%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

For activity data (emissions), the same sources are used throughout the time series. Therefore, CO₂ emissions from electric arc furnaces have been estimated in a consistent manner throughout the time-series.

2) *CH₄*

● **Uncertainty**

The uncertainty for the emission factor has been estimated as 163% and the uncertainty for activity data has been estimated as 5% (see chapter 3). As a result, the uncertainty for CH₄ emissions has been estimated as 163%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

For activity data, the same sources are used throughout the time series. The emission factor is constant throughout the time series. Therefore, CH₄ emissions from electric arc furnaces in steel production have been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

Recalculations were conducted for FY2009, since the renewed value of the energy consumption in electric arc furnaces was provided.

f) Source-specific Planned Improvements

No improvements are planned.

4.4.2. Ferroalloys Production (2.C.2.)

a) Source/Sink Category Description

1) CO₂

Ferroalloys are produced in Japan, and the CO₂ that is generated in association with the ferroalloys production is emitted as a result of the oxidization of coke used as a reducing agent. Consumption of coke is included in consumption of fuel under the Fuel Combustion Sector (1.A.), and CO₂ generated as a consequence of the oxidization of coke used as a reduction agent has already been calculated under the Fuel Combustion Sector (1.A.). Residual carbon in the ferroalloys is oxidized when the ferroalloys are used in the production of steel, and are released into the atmosphere as CO₂. Therefore, it has been reported as "IE".

2) CH₄

Ferroalloys are manufactured in Japan in electric arc furnaces, small-scale blast furnaces, and Thermit furnaces. CH₄ generated in association with ferroalloy production is thought to be generated when the oxidization of coke, a reduction agent, takes place.

b) Methodological Issues

● Estimation Method

CH₄ emissions from ferroalloy production were calculated by multiplying an emission factor based on actual measurements obtained in Japan by the energy consumption of electric arc furnaces. This is the same method used for calculating CH₄ emissions in the Fuel Combustion Sector (1.A.1 Energy Industries).

● Emission Factors

The value for the emission factor of electric arc furnaces (12.8 kg-CH₄/TJ) was used because these furnaces produce ferroalloys.

● Activity Data

Energy consumption amounts included in the "ferroalloy" category for the iron and steel industries of the General Energy Statistics were used.

Table 4-28 Energy consumption from ferroalloy production

Consumption	Unit	1990	1995	2000	2005	2008	2009	2010
Furnaces (for Ferroalloys)	TJ	14,456	10,699	10,181	10,072	8,578	8,458	9,510

c) Uncertainties and Time-series Consistency

● Uncertainty

The uncertainty for the emission factor has been estimated as 163% and the uncertainty for activity data has been estimated as 5% (see chapter 3). As a result, the uncertainty for CH₄ emissions has been estimated as 163%. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

For activity data, the same sources are used throughout the time series. The emission factor is constant throughout the time series. Therefore, CH₄ emissions from furnaces for ferroalloy have

been estimated in a consistent manner throughout the time-series.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.4.3. Aluminium Production (2.C.3.)

a) Source/Sink Category Description

1) CO₂

Aluminum refining is conducted in Japan. CO₂ generated in association with aluminum smelting is emitted in conjunction with the oxidization of the anode paste used as a reducing agent. Consumption of coke, the main ingredient in the anode paste has been included in fuel consumption under the Fuel Combustion Sector (1.A.), and the CO₂ that is generated by the oxidization of coke used as a reducing agent has already been calculated under the Fuel Combustion Sector (1.A.). Therefore, it has been reported as “IE”.

2) CH₄

Aluminum refining is conducted in Japan. There is a small amount of hydrogen in the pitch that acts as a raw material for the anode paste used in aluminum smelting. Theoretically, therefore, it is possible that CH₄ could be generated. As there is no actual data on emissions, however, it is not possible to calculate emissions. There is also no emission factor offered in the *Revised 1996 IPCC Guidelines*, and no data on the hydrogen content of pitch can be obtained. As it is not possible to estimate an emission factor, emissions have been reported as “NE”.

3) PFCs

PFCs are emitted during aluminum refining, due to the use of a fluoride melt consisting mainly of cryolite during electrolysis.

b) Methodological Issues

● **Estimation Method**

Estimating emissions involved multiplying the production amount of primary aluminum refining by Japan’s country-specific emission factors calculated using the equation prescribed in the *Revised 1996 IPCC Guidelines*.

● **Emission Factors**

The equation prescribed in the Tier 1b method of the *Revised 1996 IPCC Guidelines* was used to determine emission factors, as shown in the table below.

Table 4-29 PFCs emission factor of aluminum production

Item	Unit	1995	2000	2005	2008	2009	2010
PFC-14 (CF ₄)	kgPFC-14/t	0.542	0.369	0.307	0.300	0.301	0.300
PFC-116 (C ₂ F ₆)	kgPFC-116/t	0.0542	0.0369	0.0307	0.0300	0.0301	0.0300

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy, Trade and Industry

● **Activity Data**

As the activity data for PFC emissions in conjunction with aluminum refining, we used the aluminum production amounts given in the Yearbook of Minerals and Non-Ferrous Metals Statistics compiled by the Ministry of Economy, Trade and Industry. Japan's primary aluminum production is small, at about 0.03% of world production.

c) **Uncertainties and Time-series Consistency**

● **Uncertainty**

For the uncertainty of the emission factor, 33% was applied, according to the *GPG (2000)* default value. For the uncertainty of the activity data, 5%, the value set by the Committee for Greenhouse Gas Estimation Methods was applied. As a result, the uncertainty of the emissions was determined to be 33%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

Emissions from 1990 to 1994 have not been estimated due to the lack of data. For years after 1995, The Chemical and Bio Sub-Group, Ministry of Economy, Trade and Industry annually collects and estimates F gas emissions.

d) **Source-specific QA/QC and Verification**

The data collected and estimated by the Chemical and Bio Sub-Group, Ministry of Economy, Trade and Industry is verified by the Committee for Greenhouse Gas Estimation Methods and is used in the inventory.

e) **Source-specific Recalculations**

There have been no source-specific recalculations.

f) **Source-specific Planned Improvements**

No improvements are planned.

4.4.4. SF₆ Used in Aluminium and Magnesium Foundries (2.C.4.)

4.4.4.1. Aluminium Foundry

Emission from this source was reported as "NO" as it was been confirmed that Japan had no record of the use of SF₆ in aluminum forging processes.

4.4.4.2. Magnesium Foundry

a) Source/Sink Category Description

SF₆ is emitted in magnesium foundries, due to its use as cover gas to prevent oxidation of molten magnesium.

b) Methodological Issues

Emissions are an aggregation of all SF₆ used by magnesium foundries. The data that has been reported is given in documentation prepared by the Chemical and Bio Sub-Group of the Ministry of Economy, Trade and Industry's Industrial Structure Council, for emissions of SF₆ used in magnesium foundries. The associated indices are given in the table below.

Table 4-30 Indices related to SF₆ emitted from magnesium foundries

Item	Unit	1995	2000	2005	2008	2009	2010
Consumption of SF ₆	t	5	43	48	27	10	13

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy, Trade and Industry

c) Uncertainties and Time-series Consistency

● Uncertainty

For the uncertainty of the emission factor, 0% was applied, due to the fact that the amount of emissions is equal to the amount of magnesium used. For the uncertainty of the activity data, 5% was applied, according to the value set by the Committee for Greenhouse Gas Estimation Methods. As a result, the uncertainty of the emissions was determined to be 5%. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

See section 4.4.3. c) .

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.5. Other Production (2.D.)

4.5.1. Pulp and Paper (2.D.1.)

Pulp and Paper production possibly emit nitrogen oxides (NO_x), carbon monoxide (CO), non-CH₄ volatile organic compounds (NMVOC), and sulfur dioxide (SO₂). These emissions are reported in Annex 3.

4.5.2. Food and Drink (2.D.2.)

Foods and drinks are manufactured in Japan, and because CO₂ is used in the manufacturing process (frozen CO₂ and raw material for carbonated drinks, etc.), it is conceivable that CO₂ is emitted into the atmosphere in the course of manufacturing. The CO₂ used in the process of manufacturing foods and drinks, however, is a by-product gas of petrochemical products, and as such emissions have already been incorporated into the Fuel Combustion Sector (1.A.), they have been reported as “IE”.

4.6. Production of Halocarbons and SF₆ (2.E.)

This category covers HFCs, PFCs and SF₆ emissions from the manufacturing processes of Halocarbons and SF₆.

This section includes GHG emissions from two sources: By-product Emissions: Production of HCFC-22 (2.E.1) and Fugitive Emissions (2.E.2).

In 2010, emissions from Production of Halocarbons and SF₆ were 527 Gg-CO₂ eq. and represented 0.04% of GHG of Japan’s total GHG emissions (excluding LULUCF). The emissions had decreased by 97.7% compared to 1995.

Table 4-31 Emissions from 2.E. Production of Halocarbons and SF₆

Gas	Emission sub-category		Units	1995	2000	2005	2008	2009	2010	
HFCs	2.E Production of Halocarbons and SF ₆	2.E.1	By-product emissions: Production of HCFC-22	Gg-CO ₂ eq.	16,965.00	12,402.00	463.32	469.17	39.78	42.12
		2.E.2	Fugitive emissions	Gg-CO ₂ eq.	480.12	257.84	352.69	232.24	182.36	86.22
	Total			Gg-CO ₂ eq.	17,445.12	12,659.84	816.01	701.41	222.14	128.34
PFCs	2.E Production of Halocarbons and SF ₆	2.E.2	Fugitive emissions	Gg-CO ₂ eq.	762.85	1,359.00	837.49	523.80	399.48	200.24
SF ₆		2.E.2	Fugitive emissions	t	197.00	36.00	40.80	53.90	10.90	8.30
	Gg-CO ₂ eq.			4,708.30	860.40	975.12	1,288.21	260.51	198.37	
Total of All Gases				Gg-CO ₂ eq.	22,916.27	14,879.24	2,628.62	2,513.42	882.13	526.96

4.6.1. By-product Emissions: Production of HCFC-22 (2.E.1.)

a) Source/Sink Category Description

HFC-23 is generated as a by-product of HCFC-22 production.

b) Methodological Issues

● Estimation Method

Estimating emissions involved subtracting the recovery and destruction amount of by-product HFC-23 (measured data) from the amount of by-product HFC-23 generated at HCFC-22 production plants in Japan. The amount of by-product HFC-23 was estimated by multiplying the production of HCFC-22 by the generation rate of HFC-23 (obtained from the results of composition analysis of the interior of a reactor).

The recovery/destruction units are constantly running when the plants are in operation. If any trouble

arises in the units, management practices are to stop the plant operation.

Emissions of by-product HFC-23 associated with the production of HCFC-22

$$\text{Emissions of HFC-23} = \text{Production of HCFC-22 (t)} \times \text{Rate of generation of HFC-23 (\%)} \\ - \text{Amount of recovery and destruction (t)}$$

Table 4-32 Indices related to By-product Emissions of HFC-23: Production of HCFC-22

Item	Unit	1995	2000	2005	2008	2009	2010
Production of HCFC-22	t	81,000	95,271	65,715	60,401	26,682	46,149
Rate of generation of HFC-23	%	2.13%	1.70%	1.90%	2.00%	2.34%	2.01%
Emission rate to production	%	1.79%	1.11%	0.06%	0.07%	0.01%	0.01%
Emissions	t	1,450	1,060	40	40	3	4
	Mt-CO ₂ eq.	16.97	12.40	0.46	0.47	0.04	0.04

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy, Trade and Industry

*Emissions decreased because all manufacturing facilities were equipped with recovery/destruction units in 2004. The low emission rate to production is due to efforts made in preventing the fall of the operating rates through the improvement in techniques of operation management of destruction facilities and maintenance.

c) *Uncertainties and Time-series Consistency*

● *Uncertainty*

For the uncertainty of the emission factor, 2% was applied, according to the *IPCC 2006 Guidelines* default value. For the uncertainty of the activity data, 5% was applied, according to the value set by the Committee for Greenhouse Gas Estimation Methods. As a result, the uncertainty of the emissions was determined to be 5%. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

See section 4.4.3. c) .

d) *Source-specific QA/QC and Verification*

See section 4.4.3. d) . Emissions are surveyed for all production plants in Japan. Composition analysis is carried out frequently, as in the case where one plant takes measurements every day. Concentration measurements are implemented at the vent of the plant.

e) *Source-specific Recalculations*

There have been no source-specific recalculations.

f) *Source-specific Planned Improvements*

No improvements are planned.

4.6.2. Fugitive Emissions (2.E.2.)

a) *Source/Sink Category Description*

HFCs, PFCs, SF₆ are emitted as fugitive emissions during their manufacturing. For decomposition of the residual gases and cleansing of the containment shell, or releasement into the atmosphere, these emissions are reported under this subcategory.

b) Methodological Issues

● Estimation Method

Emissions were reported based on measurement data at each of HFCs, PFCs, SF₆ manufacturing plant in Japan. Recovery etc is hereby taken into account. The recovery/destruction units are constantly running when the plants are in operation. If any trouble arises in the units, management practices are to stop the plant operation.

Fugitive emissions in production from this source category were reported by subtracting the amount of production from the amount of HFCs, PFCs, SF₆ generated at each gas manufacturing facility. Emissions of HFCs for each year were given by the Japan Fluorocarbon Manufactures Association, and emissions of PFCs and SF₆ were given by the Japan Chemical Industry Association.

The associated indices are given in the table below.

Table 4-33 Indices related to fugitive emissions from HFCs production

Item	Unit	1995	2000	2005	2008	2009	2010
Emissions	Mt-CO ₂ eq.	0.480	0.258	0.353	0.232	0.182	0.086

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy, Trade and Industry

Table 4-34 Indices related to fugitive emissions from PFCs production

Item	Unit	1995	2000	2005	2008	2009	2010
Production of PFCs	t	1,207	2,336	2,726	2,802	2,028	2,800
Emissions	t	107	181	107	67	50	25
	Mt-CO ₂ eq.	0.763	1.359	0.837	0.524	0.399	0.200

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy, Trade and Industry

Table 4-35 Indices related to fugitive emissions from SF₆ production

Item	Unit	1995	2000	2005	2008	2009	2010
Production of SF ₆	t	2,392	1,556	2,313	2,647	2,562	2,201
Emissions	t	197.0	36.0	40.8	53.9	10.9	8.3
	Mt-CO ₂ eq.	4.708	0.860	0.975	1.288	0.261	0.198

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy, Trade and Industry

* Emissions decreased because all manufacturing facilities were equipped with recovery/destruction units in 2009.

c) Uncertainties and Time-series Consistency

● Uncertainty

For the uncertainties of the emission factors, 100% was applied for all HFCs, PFCs and SF₆, according to the *GPG (2000)* default value. For the uncertainties of the activity data, 10% was applied for all HFCs, PFCs and SF₆, according to the value set by the Committee for Greenhouse Gas Estimation Methods. As a result, the uncertainties of the emissions for all HFCs, PFCs and SF₆ were determined to be 100%. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

See section 4.4.3. c) .

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.7. Consumption of Halocarbons and SF₆ (2.F.)

This category covers HFCs, PFCs and SF₆ emissions from the manufacturing, utilization and disposal processes of the products of Halocarbons and SF₆ used. This section includes GHG emissions from nine sources: Refrigeration and Air Conditioning Equipment (2.F.1), Foam Blowing (2.F.2.), Fire Extinguishers (2.F.3.), Aerosols (2.F.4.) Solvents (2.F.5.), Other applications using ODS substitutes (2.F.6.), Semiconductors (2.F.7.), Electrical Equipment (2.F.8.) and Other (2.F.9.).

In 2010, emissions from Consumption of Halocarbons and SF₆ were 22,679 Gg-CO₂ eq., and represented 1.8% of GHG of Japan's total GHG emissions (excluding LULUCF). The emissions had decreased by 20.0% compared to 1995.

Table 4-36 Emissions from 2.F. Consumption of Halocarbons and SF₆

Gas	Emission sub-category		Units	1995	2000	2005	2008	2009	2010	
HFCs	2.F Consumption of Halocarbons and SF ₆	2.F.1	Refrigeration and Air Conditioning Equipment	Gg-CO ₂ eq.	840.40	2,688.60	7,667.03	13,268.94	15,133.69	17,088.19
		2.F.2	Foam Blowing	Gg-CO ₂ eq.	451.76	440.31	316.30	286.38	290.18	290.97
		2.F.3	Fire Extinguishers	Gg-CO ₂ eq.	NO	3.73	5.92	6.35	6.55	6.72
		2.F.4	Aerosols/Metere d Dose Inhalers	Gg-CO ₂ eq.	1,365.00	2,834.35	1,571.89	889.55	809.25	640.09
		2.F.7	Semiconductors	Gg-CO ₂ eq.	157.89	173.60	141.06	145.68	92.36	102.19
	Total			Gg-CO ₂ eq.	2,815.05	6,140.59	9,702.21	14,596.89	16,332.03	18,128.16
PFCs	2.F Consumption of Halocarbons and SF ₆	2.F.5	Solvents	Gg-CO ₂ eq.	10,263.55	2,505.63	2,289.26	1,318.27	1,142.15	1,375.99
		2.F.7	Semiconductors	Gg-CO ₂ eq.	3,144.23	5,637.07	3,860.52	2,756.49	1,715.19	1,818.65
		2.F.9	Other-Railway Silicon Rectifiers	Gg-CO ₂ eq.	NO	NO	NO	4.67	NO	NO
	Total			Gg-CO ₂ eq.	13,407.78	8,142.70	6,149.78	4,079.42	2,857.34	3,194.63
SF ₆	2.F Consumption of Halocarbons and SF ₆	2.F.7	Semiconductors	t	47.22	94.16	72.50	39.85	25.37	29.45
		2.F.8	Electrical Equipment	t	460.46	127.62	39.45	37.74	31.19	27.29
	Total			t	507.68	221.77	111.95	77.60	56.56	56.74
	Total			Gg-CO ₂ eq.	12,133.65	5,300.39	2,675.51	1,854.54	1,351.76	1,356.15
Total of All Gases				Gg-CO ₂ eq.	28,356.48	19,583.69	18,527.50	20,530.85	20,541.13	22,678.94

4.7.1. Refrigeration and Air Conditioning Equipment (2.F.1.)

4.7.1.1. Domestic Refrigeration Production, Use and Disposal (2.F.1.-)

a) Source/Sink Category Description

1) HFCs

HFCs are emitted from the production, use (including failure of devices), and disposal of domestic refrigeration.

2) PFCs

Emission from this source in the “production” category was reported as “NO” as Japan had no record of their use in the production of the products. The emission was also reported as “NO” in the “use” and “disposal” categories, because it was unlikely that PFCs were used in imported products, or refrigerants were refilled.

b) Methodological Issues

● Estimation Method

The collected amount of HFC under regulation was subtracted from 1) fugitive refrigerant ratio from production, 2) fugitive refrigerant ratio from use (including failure of devices), and 3) refrigerant contained at the time of disposal, separately, based on production and shipment amounts and refrigerant contained. Then, all there were combined.

Emissions from use and disposal were estimated by summing up the values calculated for each year of the production of devices.

Emissions of HFCs from Domestic Refrigeration

$$\begin{aligned} \text{HFC emissions} = & \text{total refrigerant contained at production} \times \text{fugitive refrigerant ratio at production} \\ & + \sum (\text{number of operated devices containing HFC} \times \text{refrigerant contained per operated device} \\ & \times \text{fugitive refrigerant ratio from use}) \\ & + \sum (\text{number of disposed devices containing HFC} \times \text{refrigerant contained per disposed device}) \\ & - \text{collected volume of HFC} \end{aligned}$$

The associated indices are given in the table below.

Table 4-37 Indices related to emissions of HFCs from domestic refrigeration

Item	Unit	1995	2000	2005	2008	2009	2010
Total HFC charged in the year of production	t	520	590	0.3	0	0	0
Fugitive refrigerant ratio at production	%	1.00%	1.00%	0.17%	0%	0%	0%
Number of operated HFC devices	1,000 devices	7,829	33,213	41,796	34,509	31,471	27,925
Refrigerant charged per device at production	g	150	125	125	125	125	125
Operational fugitive ratio (including failure)	%	0.3%	0.3%	0.3%	0.3%	0.3%	0.3%
Number of HFC devices disposed	1,000 devices	0	177	1,839	3,154	3,445	3,588
Amount of HFC collected under law	t/year	—	—	52	111	139	167
Emissions	t	8.7	40.1	187.8	283.9	289.0	276.3
	Mt-CO ₂ eq.	0.011	0.052	0.244	0.369	0.376	0.359

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

c) Uncertainties and Time-series Consistency● **Uncertainty**

For the uncertainties of the emission factors, 50% was applied for all production, use, and disposal, according to the values used in a similar category. For the uncertainties of the activity data, 40% was applied for all production, use, and disposal, according to the value set by the Committee for Greenhouse Gas Estimation Methods. As a result, the uncertainties of the emissions for all production, use, and disposal were determined to be 64%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

See section 4.4.3. c) .

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

More accurate information became available on the amount of HFC collected for 2009, resulting in recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.7.1.2. Commercial Refrigeration Production, Use and Disposal (2.F.1.-)**4.7.1.2.a. Commercial Refrigeration****a) Source/Sink Category Description****1) HFCs**

HFCs are emitted from the manufacturing, operation, maintenance, accidents, and disposal of commercial refrigeration.

2) PFCs

Emissions from this source in the “production” category were reported as “NO” as Japan had no record of their use in the production of the products. The emission was also reported as “NO” in the “use” and “disposal” categories, because it was unlikely that PFCs were used in imported products, or refrigerants were refilled.

b) Methodological Issues● **Estimation Method**

In accordance with the IPCC Guidelines, emissions of each species of F-gases from 1) manufacturing, 2) installation, 3) operation and 4) disposal are estimated for the devices below.

centrifugal refrigerating machine, screw refrigerating machine, refrigerator-freezer unit, transport refrigerator-freezer unit, separately placed showcase, built-in showcase, ice making machinery, water fountain, commercial refrigerator-freezer, all-in-one air conditioning system, gas heat pump, chilling unit

Emissions of HFCs from Commercial Refrigeration

Methods below are applied to each type of device and refrigerant

1) manufacturing

Emissions from manufacturing = Σ (number of device produced \times amount of refrigerant contained
 \times fugitive refrigerant ratio from manufacturing)

2) installation

Emissions from operation = Σ (number of device charged refrigerant in place produced
 \times amount of refrigerant contained \times fugitive refrigerant ratio from installation)

3) operation

Emissions from maintenance = Σ (number of devices operated \times amount of refrigerant contained
 \times fugitive refrigerant ratio from operation) - amount collected

4) disposal

Emissions from disposal = Σ (number of devices disposed \times average amount of refrigerant contained)
- amount collected

* In the estimation of emissions from maintenance, the yearly decrease is reflected in the “amount of refrigerant contained.” The “number of devices operated” and “number of devices disposed” are estimated from the amount of shipment and lifetime of device.

The associated indices are given in the table below.

Table 4-38 Indices related to emissions of HFCs from commercial refrigeration

Item	Unit	1995	2000	2005	2008	2009	2010
Number of HFC devices produced	1,000 devices	222	380	1,413	1,445	987	1,122
Average amount of refrigerant charged at production	g/device	358	587	3,377	3,532	3,276	3,280
Fugitive refrigerant ratio at production	%	0.2%	0.2%	0.2%	0.1%	0.1%	0.2%
Number of devices charged in production place	1,000 devices	9	32	138	199	175	171
Average amount of refrigerant during installation	g/device	17,806	9,221	23,914	26,529	25,361	23,766
Fugitive refrigerant ratio during installation	%	1.2%	1.4%	1.8%	1.7%	1.6%	1.6%
Number of devices operated	1,000 devices	375	1,957	6,770	10,027	10,847	11,743
Amount of refrigerant during operation	g/device	1,012	1,043	4,549	5,629	5,791	5,961
Fugitive refrigerant ratio during use	%	2-17% (depending on the kind of device)					
Number of devices disposed	1,000 devices	1	23	127	248	260	397
Amount of HFC collected under law during maintenance	t	0	0	0	436	503	548
Amount of HFC collected under law at disposal	t	0	0	183	200	230	269
Emissions from manufacturing	Mt-CO ₂ eq.	0.003	0.008	0.126	0.195	0.168	0.164
Emissions from stocks	Mt-CO ₂ eq.	0.036	0.229	2.900	6.432	7.582	8.814
Emissions from disposal	Mt-CO ₂ eq.	0.003	0.046	0.501	1.630	1.995	2.333
Emissions	Mt-CO ₂ eq.	0.042	0.283	3.527	8.258	9.746	11.311

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

* From 2002 onward, “amount of refrigerant” and “fugitive refrigerant ratio from operation” increased because devices became larger with the increase of commercial package AC devices.

Table 4-39 Type of HFC and emission factors during use, by type of commercial refrigeration

Type of commercial refrigeration	Type of HFC	Amount of refrigerant	Emission factor *	Share in the number of devices operated (2010)
Small-size refrigerators (built-ins etc)	R-404A, HFC-134a etc	0.1 - 3 kg	2%	40%
Separately installed showcases	R-404A, R-407C etc	20 - 41 kg	16%	3%
Mid-size refrigerators (excluding Separately installed showcases)	R-404A, R-407C etc	2 - 30 kg	13 - 17%	6%
Large-size refrigerators	HFC-134a, R404A etc	300 - 2,300 kg	7 - 12%	0.05%
All-in-one air conditioning systems for buildings	R-410A, R-407C etc	37 kg	3.5%	7%
Other commercial air conditioning devices (excluding All-in-one air conditioning systems for buildings)	R-410A, R-407C etc	3 - 43 kg	3 - 5%	44%

Source: Documents of the Refrigerant Policy Working Group, Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

* Includes for emissions during servicing, accidents, and breakdowns

c) *Uncertainties and Time-series Consistency*

● *Uncertainty*

See section 4.7.1.1. c) .

● *Time-series Consistency*

See section 4.4.3. c) .

d) *Source-specific QA/QC and Verification*

See section 4.4.3. d) .

e) *Source-specific Recalculations*

There have been no source-specific recalculations.

f) *Source-specific Planned Improvements*

No improvements are planned.

4.7.1.2.b. Automatic Vending machine Production, Use and Disposal

a) *Source/Sink Category Description*

1) *HFCs*

HFCs are emitted from manufacturing, accidents, and disposals of automatic vending machines.

2) *PFCs*

Emission from this source in the “production” category was reported as “NO” as Japan had no record of their use in production. The emissions were also reported as “NO” in the “use” and “disposal” categories, because it was unlikely that PFCs were used in imported products or refrigerants were refilled..

b) *Methodological Issues*

● *Estimation Method*

Emissions of F-gases from 1) manufacturing, 2) accidents and 3) disposals are estimated, based on

production and shipment amounts and amounts of refrigerants charged.

<u>Emissions of HFCs from Automatic Vender machine</u>	
1) manufacturing	Emissions from manufacturing = Σ (number of device produced \times amount of refrigerant contained \times fugitive refrigerant ratio from manufacturing)
2) accident	Emissions from accident = Σ (number of devices operated \times amount of refrigerant contained \times incidence rate \times average fugitive rate in accident)
3) disposal	
(a) until 2001	Emissions from disposal = Σ {number of devices disposed \times amount of refrigerant contained \times (1 - collection rate) }
(b) from 2002 onward	Emissions from disposal = Σ (number of devices disposed \times average amount of refrigerant contained) - amount collected

The associated indices are given in the table below.

Table 4-40 Indices related to emissions of HFCs from automatic vender machines

Item	Unit	1995	2000	2005	2008	2009	2010
Number of HFC devices produced	1,000 devices	0	272	355	270	173	173
Refrigerant charged per device	g	0	300	220	219	219	219
Fugitive refrigerant ratio at production	%	0.4%	0.4%	0.3%	0.3%	0.3%	0.3%
Number of devices operated	1,000 devices	0	284	1,999	2,384	2,368	2,279
Incidence rate	%	0.4%	0.4%	0.3%	0.3%	0.3%	0.3%
Fugitive refrigerant ratio (failure)	%	20.0%	20.0%	20.0%	20.0%	20.0%	20.0%
Fugitive refrigerant ratio (fixing)	%	0.9%	0.9%	0.5%	0.4%	0.4%	0.4%
Number of devices disposed	1,000 devices	0	0	0	213	293	286
Emissions	t	0.00	0.39	0.57	12.44	16.83	16.41
	Mt-CO ₂ eq.	0.000	0.001	0.001	0.019	0.026	0.025

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

* Accidents of devices charged with HFCs almost never occurred in 1999 and 2000, therefore, were reported as 0. After 2001 onward, the number of accidents are reflected in the estimation.

c) Uncertainties and Time-series Consistency

● Uncertainty

See section 4.7.1.1. c) .

● Time-series Consistency

See section 4.4.3. c) .

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

Recalculation was conducted for 2009, due to error correction.

f) Source-specific Planned Improvements

No improvements are planned.

4.7.1.3. Transport Refrigeration Production, Use and Disposal (2.F.1.-)

1) HFCs

Emission was reported as “IE” since HFCs in this category had been included in the total reported in 4.7.1.2. Commercial Refrigeration (2.F.1.-).

2) PFCs

Emission from this source in the “production” category was reported as “NO” since Japan had no record of their use in the production. The emission was also reported as “NO” in the “use” and “disposal” categories, because it was unlikely that PFCs were used in imported products or refrigerants were refilled.

4.7.1.4. Industrial Refrigeration Production, Use and Disposal (2.F.1.-)

1) HFCs

HFCs emissions have been reported as “IE”, as they are included in 4.7.1.2. Commercial Refrigeration (2.F.1.-).

2) PFCs

Emission from this source in the “production” category was reported as “NO” since Japan had no record of their use in the production of the products. The emission was also reported as “NO” in the “use” and “disposal” categories, because it was unlikely that PFCs were used in imported products or refrigerants were refilled.

4.7.1.5. Stationary Air-Conditioning (Household) Production, Use and Disposal (2.F.1.-)

a) Source/Sink Category Description

1) HFCs

HFCs are emitted from the manufacturing, operation, and disposals of household stationary air-conditioning devices.

2) PFCs

Emission from this source in the “production” category was reported as “NO” since Japan had no record of their use in production. The emission was also reported as “NO” in the “use” and “disposal” categories, because it was unlikely that PFCs were used in imported products or refrigerants were refilled..

b) Methodological Issues

● Estimation Method

In accordance with the IPCC Guidelines, emissions of each species of F-gases from 1) manufacturing, 2) operation, 3) disposals are estimated, based on production and shipment amounts and amounts of refrigerants charged.

Emissions of HFCs from Stationary Air-Conditioning (Household)

1) manufacturing

Emissions from manufacturing = Σ (number of devices produced \times volume of refrigerant contained \times fugitive refrigerant ratio from manufacturing)

2) operation

Emissions from operation = Σ (number of devices for shipment \times average volume of refrigerant contained \times fugitive refrigerant ratio from operation)

3) disposals

Emissions from disposal = Σ (number of devices disposed \times average volume of refrigerant contained) - volume collected

* In the estimation of emissions from operation, the yearly decrease is reflected in the “average volume of refrigerant contained.” The “number of devices for shipment” and “number of devices disposed” are estimated from volume of shipment and lifetime of device.

The associated indices are given in the table below.

Table 4-41 Indices related to emissions of HFCs (R-410a) from stationary air-conditioning (household)

Item	Unit	1995	2000	2005	2008	2009	2010
Number of HFC devices produced	1,000 devices	0	1,077	3,981	3,970	2,618	3,169
Refrigerant charged per device	g	1,000	1,000	1,000	1,000	1,000	1,000
Fugitive refrigerant ratio at production	%	0.2%	0.2%	0.2%	0.2%	0.2%	0.2%
Number of devices operated	1,000 devices	0	1,726	26,091	47,584	53,966	61,540
Average refrigerant charged during use	g/device	0	1,000	1,000	1,000	1,000	1,000
Fugitive refrigerant ratio during use	%	2%	2%	2%	2%	2%	2%
Number of devices disposed	1,000	0	2	83	351	524	764
Average refrigerant stock in device disposed	g/device	0	954	911	870	856	841
Amount of HFC collected under law	t/year	-	-	10	67	122	231
Emissions	t	0	38	596	1,206	1,426	1,675
	Mt-CO ₂ eq.	0.000	0.066	1.029	2.080	2.460	2.890

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

c) Uncertainties and Time-series Consistency

● *Uncertainty*

See section 4.7.1.1. c) .

● *Time-series Consistency*

See section 4.4.3. c) .

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

More accurate information became available on the amount of HFC collected for 2009, resulting in recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.7.1.6. Mobile Air-Conditioning (Car Air Conditioners) Production, Use and Disposal (2.F.1.-)

a) Source/Sink Category Description

1) HFCs

HFCs are emitted from manufacturing, operation, breakdowns, accidents, and disposals of mobile air-conditioning devices.

2) PFCs

Emission from this source in the “production” category was reported as “NO” since Japan had no record of their use in production. The emission was also reported as “NO” in the “use” and “disposal” categories, because it was unlikely that PFCs were used in imported products or refrigerants were refilled.

b) Methodological Issues

● Estimation Method

In accordance with the IPCC Guidelines, emissions of each species of F-gases from 1) manufacturing, 2) operation, 3) breakdowns, 4) accidents and 5) disposals are estimated.

<p>Emissions of HFCs from Mobile Air-Conditioning (Car Air Conditioners)</p> <p>Methods below are applied for each type of car</p> <p>1) manufacturing Emissions from manufacturing = Σ (number of devices produced \times amount of refrigerant contained \times fugitive refrigerant ratio from manufacturing)</p> <p>2) operation Emissions from operation = Σ (number of cars operated \times amount of refrigerant contained \times fugitive refrigerant ratio from operation)</p> <p>3) breakdowns Emissions from maintenance = Σ (number of cars operated \times amount of refrigerant contained \times rate of breakdowns \times fugitive refrigerant ratio from breakdowns)</p> <p>4) accidents Emissions from accident = Σ (number of cars in completely destroyed \times amount of refrigerant contained at time of accident)</p> <p>5) disposal (a) until 2001 Emissions from disposal = Σ {number of cars disposed \times amount of refrigerant contained \times (1 - collection rate) }</p> <p>(b) from 2002 onward Emissions from disposal = Σ (number of cars disposed \times average amount of refrigerant contained) - amount collected</p> <p>* In the estimation of emissions from operation, the yearly decrease is reflected in the “amount of refrigerant contained.”</p>

Table 4-42 Indices related to emissions of HFC-134a from car air conditioners

Item	Unit	1995	2000	2005	2008	2009	2010
Number of cars produced	1,000 devices	9,745	9,761	10,407	11,163	7,653	9,292
Fugitive refrigerant during production	g	4	4	3	3	1	1
Number of cars operated with HFC air conditioners	1,000 devices	15,655	42,374	60,364	64,543	64,407	65,091
Average refrigerant charged per device	g	700	615	548	520	497	497
Fugitive refrigerant ratio during use per year per device (normal car)	g	15	15	10	10	10	10
Breakdown incidence	%	4%	4%	4%	4%	4%	4%
Fugitive refrigerant ratio from breakdown cars	%	50%	50%	50%	50%	50%	50%
Number of cars completely destroyed	1,000 devices	50	136	193	207	210	208
Average refrigerant charged in completely destroyed car	g	681	610	522	475	460	446
Number of cars disposed	1,000 devices	116	789	2,058	2,176	2,498	2,895
Average refrigerant charged upon disposal	g	676	593	522	466	456	443
Amount of HFC collected (under law from FY2002 and beyond)	t/year	-	-	531	686	787	898
Emissions	t	605	1,759	2,205	1,956	1,944	1,925
	Mt-CO ₂ eq.	0.787	2.287	2.866	2.543	2.527	2.502

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

Table 4-43 Type of HFC in car air conditioners and emission factor during use

Type of HFC	Emission factor during use
HFC-134a	5.2%

Source: Documents of the Refrigerant Policy Working Group, Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

c) *Uncertainties and Time-series Consistency*

● *Uncertainty*

See section 4.7.1.1. c) .

● *Time-series Consistency*

See section 4.4.3. c) .

d) *Source-specific QA/QC and Verification*

See section 4.4.3. d) .

e) *Source-specific Recalculations*

More accurate information became available on the number of cars completely destroyed and amount of HFC collected for 2009, resulting in recalculations.

f) *Source-specific Planned Improvements*

No improvements are planned.

4.7.2. Foam Blowing (2.F.2.)

4.7.2.1. Hard Foam Production (2.F.2.-)

4.7.2.1.a. Urethane Foam

a) *Source/Sink Category Description*

HFC-134a is emitted as a result of foam blowing agent use.

b) Methodological Issues

● Estimation Method

In accordance with the IPCC Guidelines (closed-cell foams), emissions were calculated assuming that 10% of the emission from foam blowing agents used each year occurred within the first year after production, with the remainder emitted over 20 years at the rate of 4.5% per year. The data on the amount of foam blowing agents used each year was provided by the Japan Urethane Foam Association, Japan Urethane Raw Materials Association.

It is difficult to separate the “use” emission from that at the time of “disposal” because urethane foams were disposed of at various times. Accordingly, the emissions in the “use” and “disposal” categories were combined and reported under the “use” category, while the emission in the “disposal” category was reported as “IE”.

Urethane-related HFC-134a emissions

HFC-134a emissions

= Amount of HFC-134a used [t] × Leakage during foam blowing [%]

+ Total amount used upto the previous year [t] × Percentage of annual emissions during use [%]

= (Emission during production) + (Emission during use)

Table 4-44 Indices related to emissions of HFC-134a from urethane foam

Item	Unit	1995	2000	2005	2008	2009	2010
HFC-134a Use	t	0	167	224	145	109	66
Leakage during foam blowing	%	10%	10%	10%	10%	10%	10%
Annual emissions rate during use	%	4.5%	4.5%	4.5%	4.5%	4.5%	4.5%
Emissions within the first year after production	t	0	17	35	15	11	7
Emissions during use	t	0	0	44	75	82	86
Emissions	t	0.0	16.7	78.8	89.5	92.4	93.0
Emissions during production	Mt-CO ₂ eq.	0.000	0.022	0.046	0.019	0.014	0.009
Emissions during use	Mt-CO ₂ eq.	0.000	0.000	0.057	0.098	0.106	0.112
Emissions	Mt-CO ₂ eq.	0.000	0.022	0.102	0.116	0.120	0.121

Source: For HFC-134a Use, leakage during foam blowing, and annual emissions rate during use, Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

* The amount of HFC-134a used in 1995-1999 was zero.

c) Uncertainties and Time-series Consistency

● Uncertainty

For the uncertainties of the emission factors, 50% was applied for both production and use, according to the values used in a similar category. For the uncertainties of the activity data, 50% was applied for both production and use, according to *GPG (2000)*'s default value. As a result, the uncertainties of the emissions for both production and use were determined to be 71%. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

See section 4.4.3. c) .

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.7.2.1.b. High Expanded Polyethylene Foam (2.F.2.-)*a) Source/Sink Category Description*

HFC-134a is emitted as a result of foam blowing agent use.

b) Methodological Issues● *Estimation Method*

In accordance with the IPCC Guidelines (open-cell foams), emissions were calculated assuming that all of the emissions from foam blowing agents used occurred at the time of production. The amount of the emissions from foam blowing agents used each year was provided by the High Expanded Polyethylene Foam Industry Association.

Table 4-45 Indices related to emissions of HFC-134a from high expanded polyethylene foam

Item	Unit	1995	2000	2005	2008	2009	2010
HFC-134a Use	t	346	322	128	100	100	100
Emissions	t	346	322	128	100	100	100
	Mt-CO ₂ eq.	0.450	0.419	0.166	0.130	0.130	0.130

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

Table 4-46 Indices related to emissions of HFC-152a from high expanded polyethylene foam

Item	Unit	1995	2000	2005	2008	2009	2010
HFC-152a Use	t	14	NO	NO	NO	NO	NO
Emissions	t	14	NO	NO	NO	NO	NO
	Mt-CO ₂ eq.	0.002	0	0	0	0	0

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

c) Uncertainties and Time-series Consistency● *Uncertainty*

See section 4.7.2.1.a. c).

● *Time-series Consistency*

See section 4.4.3. c) .

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.7.2.1.c. Extruded Polystyrene Foam Production (2.F.2.-)*a) Source/Sink Category Description*

HFC-134a is emitted as a result of foam blowing agent use.

b) Methodological Issues● *Estimation Method*

Emissions were calculated assuming that 25% of the emission of foam blowing agents occurs within the first year after production, with the remainder emitted over 30 years at the rate of 2.5% per year. The amount of the emissions from foam blowing agents used each year was provided by the Extruded Polystyrene Foam Industry Association. This assumption is consistent with the IPCC Good Practice Guidance and the estimation method under PRTR for the amount of transferred HCFC at polystyrene foam production sites.

It is difficult to separate the “use” emission from that at the time of “disposal” because heat insulation material is disposed of at various times such as the renovation and dismantling of buildings, and in times of disaster. Since disposed polystyrene foam is considered to be emitting HFCs as same as that in use, these emissions are combined and reported under “use”, while the emissions from “disposal” were reported as “IE”.

Extruded polystyrene foam-related HFC-134a emissions

HFC-134a emissions =

Amount of HFC-134a used in particular year [t] × Leakage during foam blowing 25%

+ Total amount used in the past up to the previous year [t] × Annual emission rate during use [%]

Table 4-47 Indices related to emissions of HFC-134a from extruded polystyrene foam

Item	Unit	1995	2000	2005	2008	2009	2010
HFC-134a Use	t	0	0	26	0	0	0
Foam productization rate	%	75%	75%	75%	75%	75%	75%
Annual emission rate during use	%	-	-	2.5%	2.5%	2.5%	2.5%
Emissions during production	t	0	0	7	0	0	0
Emissions during use	t	0	0	30	31	31	31
Emissions	t	0	0	37	31	31	31
Emissions during production	Mt-CO ₂ eq.	0.00	0.00	0.01	0.00	0.00	0.00
Emission during use	Mt-CO ₂ eq.	0.00	0.00	0.04	0.04	0.04	0.04
Emissions	Mt-CO ₂ eq.	0.00	0.00	0.05	0.04	0.04	0.04

Source: For HFC-134a Use, foam productization rate, and annual emissions rate during use, Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

* The amount of HFC-134a used in 1995-2000 was zero.

c) Uncertainties and Time-series Consistency● **Uncertainty**

See section 4.7.2.1.a. c).

● **Time-series Consistency**

See section 4.4.3. c) .

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

Recalculation was conducted for 2005, due to error correction.

f) Source-specific Planned Improvements

No improvements are planned.

4.7.2.2. Soft Foam (2.F.2.-)

All foam using HFCs for forming is hard foam. Emissions have therefore been reported as “NO”.

4.7.3. Fire Extinguishers (2.F.3.)**a) Source/Sink Category Description**

HFCs are emitted by the use of halogen fire extinguishers.

b) Methodological Issues● **Estimation Method**

HFC-23 and HFC-227ea are used for the productions of fire extinguishers. However, as of 2004, only HFC-227ea is filled in the bottles for fire extinguishing equipments, and each company purchases pre-filled HFC-23 fire extinguisher bottles.

HFCs emission from this category was reported as “NO” by expert judgment since HFC-227ea was a very small amount, 0.0007(t) (= 700g) when emission from production in FY2004 was estimated. For use, at the time around 1995, almost no HFC filled fire extinguishers existed on the market, therefore it is assumed that there was not any use, resulting in NO for 1995 emissions.

For 1996 and following years, calculations were performed using the following equation and based on the HFC extinguishing agent installations and stocks.

HFC emissions from use of fire extinguishers

$$\text{HFC emissions [t]} = \text{HFC extinguishing agent installations and stocks [t]} \times \text{Emission factor during use}$$

Concerning the emission at the time of disposal of fire extinguishers, it is reported as “NO” because the use of HFC for fire extinguishers has just started, and also the expected lifetime of buildings is 30-40 years, therefore they are unlikely to be disposed of as of present.

● Emission Factors

There are still no findings on the emission factor of HFC extinguishing agents when using them. The emission rate (0.00088) determined from refills of halons (provided by the Fire and Disaster Management Agency), which are similar extinguishing agents, was adopted as the emission factor for this category.

Table 4-48 References for the Emission factor of fire extinguishers
(The emission ratio of halon fire extinguishers)

	Unit	2002	2003	2004	2005	2006	2007	Average
Installations of halon 1301 (A)	t	17,094	17,090	17,060	16,994	17,075	16,889	17,034
Refills of halon 1301 (B)	t	13	13	22	13	14	15	15
(B) / (A)		0.00076	0.00076	0.00129	0.00076	0.00082	0.00089	0.00088

● Activity Data

HFC stock amounts provided by the Fire Defense Agency were used as activity data for HFC emissions from fire extinguishing agents use.

Table 4-49 The amounts of the HFC extinguishing agent installations and stocks

Item	Unit	1995	2000	2005	2008	2009	2010
Installations and stocks of HFC-23	t	NO	306	478	501	512	523
HFC-23 emissions	t	NO	0.27	0.42	0.44	0.45	0.46
	Gg-CO ₂ eq.	NO	3.15	4.92	5.16	5.27	5.39
Installations and stocks of HFC-227ea	t	NO	225	392	467	498	522
HFC-227ea emissions	t	NO	0.20	0.34	0.41	0.44	0.46
	Gg-CO ₂ eq.	NO	0.57	1.00	1.19	1.27	1.33
Total emissions	Gg-CO ₂ eq.	NO	3.73	5.92	6.35	6.55	6.72

c) Uncertainties and Time-series Consistency

● Uncertainty

For the uncertainties of the emission factor for fire extinguisher use, 50% was applied, according to the values used in a similar category. For the uncertainties of the activity data, 40% was applied according to the value set by the Committee for Greenhouse Gas Estimation Methods. As a result, the uncertainties of the emissions during use for the category were determined to be 64%. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

Calculations are performed with a method consistently used from FY1995, based on an emission factor and activity data received from the Fire Defense Agency.

d) Source-specific QA/QC and Verification

The data received from the Fire Defense Agency is compiled by the Chemical and Bio Sub-Group, Ministry of Economy, Trade and Industry. It is verified by the Committee for Greenhouse Gas Estimation Methods and is used in the inventory.

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.7.4. Aerosols (2.F.4.)

4.7.4.1. Aerosols (2.F.4.-)

a) Source/Sink Category Description

HFCs are emitted from the manufacturing and use of aerosols.

b) Methodological Issues

● *Estimation Method*

In accordance with the IPCC Guidelines, emissions were calculated on the assumption that 50% of the emission from the amount of aerosol filled in the products (potential emissions) occurred in the year of production, with the remaining 50% emitted in the following year. Fugitive emissions from manufacturing is considered as the balance between the amount used for production and the actual measurement amount filled in the products, and it is included in the emissions. The data on the amount used for production and the amount filled in the products were provided by the Aerosol Industry Association of Japan. HFC is considered to be actually remaining in disposed aerosols at some level. However, the amount of emission at the time of “disposal” was reported as “IE” since it is included in the calculation for the “use” category.

F-gas (HFC-134a, HFC-152a) emissions associated with the manufacturing of Aerosol

$$\begin{aligned} \text{F-gas emissions in year } n &= \text{Fugitive emissions during manufacturing (t)} \\ &+ \text{F-gas potential emissions in year } (n-1) \times 50 (\%) \\ &+ \text{F-gas potential emissions in year } n \times 50 (\%) \end{aligned}$$

$$\begin{aligned} \text{Fugitive emissions during manufacturing} &= \text{F-gas consumed during manufacturing in year } n \\ &- \text{F-gas potential emissions} \end{aligned}$$

The associated indices are given in the table below.

Table 4-50 Indices related to emissions of HFC-134a from aerosols

Item	Unit	1994	1995	2000	2005	2008	2009	2010
Potential Emissions	t	800	1,300	2,044	604	343	230	200
Fugitive emissions during production*	t	-	-	80.2	24.9	12.8	10.0	8.1
Emissions in the year produced, during use	t	400	650	1,022	302	172	115	100
Remaining (emissions in the next year)	t	400	650	1,022	302	172	115	100
Emissions	t	-	1,050	2,137	908	338	297	223
	Mt-CO ₂ eq.	-	1.365	2.778	1.181	0.439	0.386	0.290

Source: Potential Emissions: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

* Fugitive emissions from 1994 to 1997 are included in potential emissions.

Table 4-51 Indices related to emissions of HFC-152a from aerosols

Item	Unit	1995	2000	2005	2008	2009	2010
Potential Emissions	t	NO	34	1,300	1,416	764	558
Fugitive emissions during production	t	NO	1.1	28.9	380.5	494.0	638.0
Emissions in the year produced, during use	t	NO	17	650	708	382	279
Remaining (emissions in the next year)	t	0	17	650	708	382	279
Emissions	t	0	18	1,217	1,685	1,584	1,299
	Mt-CO ₂ eq.	0.000	0.003	0.170	0.236	0.222	0.182

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

c) *Uncertainties and Time-series Consistency*

● *Uncertainty*

For the uncertainties of the emission factors, 0% was applied for all production, use and disposal, due to the fact that the amount of emissions is equal to the amount of aerosols used. For the uncertainties of the activity data, 40% was applied for all production, use, and disposal, according to the value set by the Committee for Greenhouse Gas Estimation Methods. As a result, the uncertainties of the emissions for all production, use and disposal were determined to be 40%. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

See section 4.4.3. c) .

d) *Source-specific QA/QC and Verification*

See section 4.4.3. d) .

e) *Source-specific Recalculations*

There have been no source-specific recalculations.

f) *Source-specific Planned Improvements*

No improvements are planned.

4.7.4.2. Metered Dose Inhalers (2.F.4.-)

a) *Source/Sink Category Description*

HFCs are emitted from the use and disposal of metered dose inhaler (MDI) s.

b) *Methodological Issues*

● *Estimation Method*

In accordance with the IPCC Guidelines, emissions were calculated on the assumption that from the amount used each year, 50% of the emission occurred in the year of production, with the remaining 50% emitted in the following year.

The amount of purchased gas, the amount of the use of domestically produced MDI, and the use of imported MDI, and the amount of disposal of MDI were provided by the Federation of Pharmaceutical Manufacturers' Associations of Japan (FPMAJ). FPMAJ estimates the amount of HFC disposal by mainly including destructed MDI that were defective products.

F-gas (HFC-134a, HFC-227ea) emissions associated with the manufacturing of MDI

F-gas emissions in year n = Fugitive emissions during manufacturing (t)
 + F-gas potential emissions in year (n - 1) × 50 (%)
 + F-gas potential emissions in year n × 50 (%)
 - amount of disposal of F-gas contained in MDI

Potential emissions of F-gas = F-gas contained in domestic produced MDI + F-gas contained in imported MDI

The associated indices are given in the table below.

Table 4-52 Indices related to emissions of HFC-134a from MDI

Item	Unit	1995	2000	2005	2008	2009	2010
Purchases of F-gas	t	0.0	1.4	1.1	1.1	0.9	1.1
Usage of domestic MDI	t	0.0	1.4	0.9	0.9	0.9	1.1
Usage of imported MDI	t	0	42	71	62	57	57
Amount collected and destroyed	t	0.0	0.1	1.9	0.5	0.4	2.5
Emissions	t	NO	37	63	61	60	56
	Mt-CO ₂ eq.	0.000	0.048	0.082	0.080	0.078	0.072

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

Table 4-53 Indices related to emissions of HFC-227ea from MDI

Item	Unit	1995	2000	2005	2008	2009	2010
Purchases of F-gas	t	0.0	0.0	42.8	48.0	29.3	37.0
Usage of domestic MDI	t	0.0	0.0	41.0	45.9	27.8	36.0
Usage of imported MDI	t	0.0	3.6	2.1	9.0	1.6	0.4
Amount collected and destroyed	t	0.0	0.0	1.2	1.6	0.9	0.8
Emissions	t	NO	1.8	48.1	46.4	42.8	33.1
	Mt-CO ₂ eq.	0.000	0.005	0.139	0.135	0.124	0.096

For the Usage of domestic MDI, Usage of imported MDI, and Amount collected and destroyed:

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

c) Uncertainties and Time-series Consistency

● Uncertainty

For the uncertainties of the emission factors, 0% was applied for all production, use and disposal, due to the fact that the amount of emissions is equal to the amount of MDI used. For the uncertainties of the activity data, 40% was applied for all production, use and disposal, according to the value set by the Committee for Greenhouse Gas Estimation Methods. As a result, the uncertainties of the emissions for all production, use and disposal were determined to be 40%. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

See section 4.4.3. c) .

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.7.5. Solvents (2.F.5.)*a) Source/Sink Category Description*

HFCs and PFCs are emitted from the use of solvents. The liquids PFCs used were C₅F₁₂ (PFC-41-12) and C₆F₁₄ (PFC-51-14). HFCs used as solvents correspond to confidential data; therefore, these data are reported as included numbers in the total of PFCs.

b) Methodological Issues● *Estimation Method*

Assuming that almost all of the total amount of liquid PFC shipment was used in cleaners and for cleaning purposes each year, the entire amount was reported in the "use" category as the amount of emissions. Emission from manufacturing was reported as "NO" since there is no practice to blend before use. Emission at the time of disposal was reported as "IE" on the assumption, from the point of view of conservativeness, that the entire amount including that was disposed of, was emitted during use, because of the difficulty in determining the status of the disposal of PFCs. It is confirmed that no disposals were identified in 1995. The associated indices are given in the table below. Emissions from PFCs contained in railway rectifiers are subtracted from liquid PFC emissions to yield the total PFC emissions.

Table 4-54 Indices related to emissions of PFCs etc. from solvents use

Item	Unit	1995	2000	2005	2008	2009	2010
Liquid PFC emissions	Gg-CO ₂ eq.	10356.1	2624.0	2289.3	1318.3	1142.1	1376.0
Liquid PFC contained in Railway rectifiers	Gg-CO ₂ eq.	92.5	118.4	0.0	0.0	0.0	0.0
PFC emissions from solvents	Gg-CO ₂ eq.	10263.6	2505.6	2289.3	1318.3	1142.1	1376.0

Source for liquid PFC: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy Trade and Industry

c) Uncertainties and Time-series Consistency● *Uncertainty*

For the uncertainties of the emission factors, 0% was applied for solvent use, due to the fact that the amount of emissions is equal to the amount of solvent used. For the uncertainties of the activity data, 40% was applied for solvent using according to the value set by the Committee for Greenhouse Gas Estimation Methods. As a result, the uncertainties of the emissions were determined to be 40%. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

Emissions are estimated in a manner consistent over the time-series methodologically and from the point of view of data source.

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.7.6. Other applications using ODS substitutes (2.F.6.)

Refrigerants filled in research and medical equipment are captured and included in other refrigerant categories, therefore the emissions from this category is reported as "IE", based on expert judgment.

4.7.7. Semiconductors Manufacture (2.F.7.)**4.7.7.1. Semiconductors (2.F.7.-)****a) Source/Sink Category Description**

HFCs, PFCs and SF₆, are emitted from the manufacturing of semiconductors.

b) Methodological Issues**● Estimation Method**

Methods of emissions from semiconductors are in line with the IPCC *GPG (2000)*. These emissions are estimated with purchase of F-gases, process supply rate, use rate of F-gas, removal rate, by-product generation ratio and removal ratio for by-products.

In addition, regarding the treatment of 10% as residue of process supply rate, these emissions are reported in this category in case of a 90% recharging rate and subsequent shipment. In cases of decomposing the residual 10% and cleansing the containment shell, or releasement into the atmosphere, these emissions are reported in "2.E.2. Production of Halocarbons and SF₆".

Japan Electronics and Information Technology Industries Association data are used for F-gases purchased.

Emissions from manufacturing (during F-gas charging to containment shell for shipment) are already reported in "2.E.2. Production of Halocarbons and SF₆", therefore, are reported as "IE" for this category. Theoretically, emissions from disposal can not be generated, therefore are reported as "NA".

F-gas emissions in Semiconductor Manufacturing

Methods below are applied for each F-gas:

(i) HFC-23, PFCs (PFC-14, PFC-116, PFC-218, PFC-c318), SF₆ emissions

Emissions
= Total CO₂ equivalent emissions from all production lines
- Total CO₂ equivalent amount destroyed in all production lines

Total CO₂ equivalent emissions from all production lines
= \sum each production line \sum {amount purchased per F-gas \times process supply rate
 \times (1 - use rate of F-gas) \times GWP}

Total CO₂ equivalent amount destroyed in all production lines
= \sum each production line \sum {amount purchased per F-gas \times process supply rate
 \times (1 - use rate of F-gas) \times fraction of F-gas destroyed \times GWP}

(For production lines without destruction facilities: fraction of F-gas destroyed = 0)

(ii) By-produced PFC-14 emissions

Emissions
= Total CO₂ equivalent emissions from all production lines
- Total CO₂ equivalent amount destroyed in all production lines

Total CO₂ equivalent emissions from all production lines
= \sum each production line \sum (purchases of PFCs \times process supply rate
 \times by production rate \times GWP)

Total CO₂ equivalent amount destroyed in all production lines
= \sum each production line \sum (purchases of PFCs \times process supply rate
 \times by production rate \times fraction of F-gas destroyed \times GWP)

(For production lines without destruction facilities: fraction of F-gas destroyed = 0)

Relevant indices are shown in Table below.

Table 4-55 Indices related to emissions of F-gases from manufacturing of semiconductors

Item	Unit	1995	2000	2005	2008	2009	2010
PFC-14 purchased	t	313.0	299.9	231.5	276.9	208.9	265.3
PFC-116 purchased	t	209.5	561.2	393.2	284.9	171.5	194.3
PFC-218 purchased	t	0.0	9.9	181.8	181.0	129.5	167.0
PFC-c318 purchased	t	0.6	38.6	24.8	40.2	33.3	35.8
HFC-23 purchased	t	47.8	49.4	42.1	73.7	53.8	67.1
SF ₆ purchased	t	90.8	131.9	96.8	79.1	60.2	76.7
Process supply rate	%	90%	90%	90%	90%	90%	90%
Use rate of PFC etc	%	20 - 70% (depending on the substance)					
Fraction of F-gas destroyed	%	90%	90%	90%	90%	90%	90%
CF ₄ by-production rate	%	C ₂ F ₆ (PFC-116): 10%, C ₃ F ₈ (PFC-218): 20%					
By-production CF ₄ removal rate	%	90%	90%	90%	90%	90%	90%
HFCs emissions	Mt-CO ₂ eq.	0.158	0.172	0.138	0.142	0.090	0.099
PFCs emissions	Mt-CO ₂ eq.	3.046	5.409	3.712	2.665	1.672	1.765
SF ₆ emissions	Mt-CO ₂ eq.	1.005	1.484	1.111	0.694	0.433	0.469

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy, Trade and Industry

Table 4-56 Use rate of HFCs, PFCs, and SF₆ during semiconductor manufacturing

Item	Unit	1995 - 2010
Use rate of PFC-14	%	20
Use rate of PFC-116	%	30
Use rate of PFC-218	%	60
Use rate of PFC-c318	%	70
Use rate of HFC-23	%	70
Use rate of SF ₆	%	50

*: use rates are default values from the IPCC guidelines.

c) Uncertainties and Time-series Consistency

● *Uncertainty*

For the uncertainties of the emission factors, 50% was applied for all HFCs, PFCs and SF₆, according to the values used in a similar category. For the uncertainties of the activity data, 40% was applied for all HFCs, PFCs and SF₆, according to the value set by the Committee for Greenhouse Gas Estimation Methods. As a result, the uncertainties of the emissions for all HFCs, PFCs and SF₆ were determined to be 64%. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

See section 4.4.3. c) .

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.7.7.2. Liquid Crystals (2.F.7.-)

a) Source/Sink Category Description

HFCs, PFCs and SF₆, are emitted from the manufacturing of liquid crystals.

b) Methodological Issues

● *Estimation Method*

Same methods applied to semiconductors are also applied to emissions from manufacturing of liquid crystals. World LCD Industry Cooperation Committee has established a voluntary action plan to reduce PFCs emissions and has engaged in reducing PFC emissions. In these activities, it should be applied IPCC methods.

Table 4-57 Indices related to emissions of F-gases from manufacturing of liquid crystals

Item	Unit	1995	2000	2005	2008	2009	2010
PFC-14 purchased	t	20.7	47.3	77.8	69.3	51.9	93.7
PFC-116 purchased	t	0.4	2.7	9.9	4.1	2.3	0.0
PFC-c318 purchased	t	0.0	0.0	0.8	1.9	1.7	1.6
HFC-23 purchased	t	0.1	0.7	1.6	1.5	1.1	1.1
SF ₆ purchased	t	11.5	85.3	101.4	146.8	127.1	176.9
Use rate of PFC	%	90%	90%	90%	90%	90%	90%
Fraction of F-gas destroyed	%	20 - 70% (depending on the substance)					
CF ₄ by-production rate	%	90%	90%	90%	90%	90%	90%
By-production CF ₄ removal rate	%	C ₂ F ₆ (PFC-116): 10%					
Desellection Efficiency of CF ₄	%	90%	90%	90%	90%	90%	90%
HFCs emissions	Mt-CO ₂ eq.	0.000	0.002	0.003	0.003	0.003	0.003
PFCs emissions	Mt-CO ₂ eq.	0.099	0.228	0.149	0.092	0.043	0.053
SF ₆ emissions	Mt-CO ₂ eq.	0.124	0.766	0.622	0.259	0.174	0.235

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy, Trade and Industry

*: use rate of PFC etc is a default value from the IPCC guidelines.

c) *Uncertainties and Time-series Consistency*

● *Uncertainty*

See section 4.7.7.1. c).

● *Time-series Consistency*

See section 4.4.3. c) .

d) *Source-specific QA/QC and Verification*

See section 4.4.3. d) .

e) *Source-specific Recalculations*

There have been no source-specific recalculations.

f) *Source-specific Planned Improvements*

No improvements are planned.

4.7.8. Electrical Equipment (2.F.8.)

a) *Source/Sink Category Description*

SF₆ are emitted during the manufacturing and use of electrical equipment.

b) *Methodological Issues*

● *Estimation Method*

Emissions from producing electrical equipment were calculated by multiplying the amount of SF₆ purchased by assembly fugitive rate. Emissions from the use of electrical equipment were calculated based on the fugitive rate during the use of electrical equipment. Emissions from the inspection and disposal of electrical equipment were obtained by actual measurements of SF₆.

In CRF, the emission was reported as "IE" after including the emission from disposal into the use of electrical equipment.

SF₆ emissions from the production of electrical equipment

$$\text{SF}_6 \text{ Emissions from the production} = \text{SF}_6 \text{ purchased (t)} \times \text{assembly fugitive rate (\%)}$$
SF₆ emission from the use of electrical equipment

SF₆ emission from the use

$$= \text{Stocks of SF}_6 \times \text{rate of emitted SF}_6 \text{ into the environment during the use of electrical equipments (0.1\%)}$$
SF₆ emission from the inspection of electrical equipment

$$\text{SF}_6 \text{ emission from the inspection} = \text{actual measurements of SF}_6$$
SF₆ emission from the disposal of electrical equipment

$$\text{SF}_6 \text{ emission from the disposal} = \text{actual measurements of SF}_6$$

The associated indices are given in the table below.

Table 4-58 Indices related to emissions of SF₆ from electrical equipment assembly

Item	Unit	1995	2000	2005	2008	2009	2010
SF ₆ purchased	t	1,380	649	629	784	459	315
SF ₆ charged to electrical equipment	t	1,464	450	582	726	410	282
Stocks (other than in electrical equipment)	t	-	105	29	40	38	26
Assembly fugitive rate	%	29%	15%	3%	2%	2%	2%
Emissions	t	400	100	23	19	11	7
	Mt-CO ₂ eq.	9.560	2.402	0.548	0.444	0.263	0.165

For SF₆ purchased, SF₆ charged to electrical equipment, Stocks in other than electrical equipment, Assembly fugitive rate:

Source: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy, Trade and Industry

Table 4-59 Indices related to emissions of SF₆ during the use of electrical equipment

Item	Unit	1995	2000	2005	2008	2009	2010
Stocks of SF ₆	t	6,300	8,000	8,700	9,000	9,000	9,100
Operational fugitive rate	%	0.1%	0.1%	0.1%	0.1%	0.1%	0.1%
SF ₆ emissions during use *	t	6.3	8.0	8.7	9.0	9.0	9.1
SF ₆ emissions during maintenance and disposal *	t	54.00	14.00	2.50	5.10	3.40	3.50
SF ₆ emissions during use, maintenance, and disposal	t	60.46	27.13	16.51	19.17	20.19	20.39
	Gg-CO ₂ eq.	1444.99	648.36	394.48	458.19	482.56	487.34

* excluding data from the Greenhouse Gas Accounting and Reporting System

Source: For Stocks of SF₆, Operational fugitive rate, SF₆ emissions during use, maintenance, and disposal: Documents of Group for Prevention of Global Warming, Chemical and Bio Sub-Group, Industrial Structure Council, Ministry of Economy, Trade and Industry

c) Uncertainties and Time-series Consistency

● Uncertainty

For the uncertainties of the emission factors, 30% was applied for production, and 50% was applied for use and disposal, according to the *GPG (2000)*'s default value. For the uncertainties of the activity data, 40% was applied for all production, use and disposal, according to the value set by the Committee for Greenhouse Gas Estimation Methods. As a result, the uncertainty of the emissions for production was determined to be 50%, and 64% for use and disposal. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

See section 4.4.3. c) .

d) Source-specific QA/QC and Verification

See section 4.4.3. d) .

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

4.7.9. Other - Railway Silicon Rectifiers (2.F.9.)**a) Source/Sink Category Description**

PFCs are emitted at disposal of railway silicon rectifiers.

b) Methodological Issues● **Estimation Method**

Based on the number of devices containing PFC-51-14, the amount of PFC-51-14 contained, and lifetime of the devices, given in the Survey on Management Methods of Halons/Liquid PFCs etc, the amount of PFC-51-14 disposed after use in railway silicon rectifiers in each fiscal year was estimated. This was done by multiplying the number of railway silicon rectifiers disposed per year, by the amount of PFC contained in each device. PFC emissions are calculated by subtracting the amount of PFC-51-14 destroyed in a specific fiscal year from the PFC disposed after use in railway silicon rectifiers in the same fiscal year.

PFC emissions at disposal of railway silicon rectifiers

$$= \text{PFC disposed after use in railway silicon rectifiers} - \text{PFC destroyed}$$

Table 4-60 Amounts of PFC Disposed from Railway Silicon Rectifiers

Item	Unit	1995	2000	2005	2008	2009	2010
Amount of PFC disposed	Gg-CO ₂ eq.	NO	NO	NO	0.63	NO	NO

c) Uncertainties and Time-series Consistency● **Uncertainty**

For the uncertainty of the emission factor from railway silicon rectifiers, the 0% value for solvents was applied since it is a similar source category. For the uncertainties of the activity data, 40% was applied. As a result, the uncertainties of the emissions were determined to be 40%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

Emissions are estimated in a manner consistent over the time-series methodologically and from the point of view of data source.

d) Source-specific QA/QC and Verification

See section 4.2.1. d) .

e) Source-specific Recalculations

Recalculation was conducted for 2008 and 2009, since the actual amount of PFC disposed became available.

f) Source-specific Planned Improvements

No improvements are planned.

References

1. IPCC, *Revised 1996 IPCC Guidelines for National Greenhouse Gas Inventories*, 1997
2. IPCC, *Good Practice Guidance and Uncertainty Management in National Greenhouse Gas Inventories*, 2000
3. IUPAC website “Atomic Weights of the Elements 1999”
(<http://www.chem.qmul.ac.uk/iupac/AtWt/AtWt99.html>)
4. Ministry of the Environment Committee for the Greenhouse Gases Emissions Estimation Methods, GHGs Estimation Methods Committee Report Part 1, August 2006
5. Ministry of Economy, Trade and Industry, *Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke*
6. Ministry of Economy, Trade and Industry, *Yearbook of Chemical Industries Statistics*
7. Ministry of Economy, Trade and Industry, Documents of Group for prevention of global warming, Chemical and Bio Sub-Group, Industrial Structure Council
8. Agency for Natural Resources and Energy, *General Energy Statistics*
9. Ministry of Economy, Trade and Industry, *Yearbook of Mineral Resources and Petroleum Products Statistics*
10. Ministry of Economy, Trade and Industry, *Yearbook of Minerals and Non-Ferrous Metals Statistics*
11. Ministry of Economy, Trade and Industry, *Yearbook of Current Survey of Energy Consumption*
12. Ministry of Economy, Trade and Industry, *Yearbook of Ceramics and Building Materials Statistics*
13. Ministry of Economy, Trade and Industry, *Yearbook of Iron and Steel, Non-ferrous Metals, and Fabricated Metals Statistics*
14. Ministry of Economy, Trade and Industry, *Yearbook of Iron and Steel Statistics*
15. Ministry of Finance, *Trade Statistics of Japan*
16. Japan Lime Association, *The Story of Lime*
17. Adjusted Price Transaction Table, The Research Institute of Economy, Trade and Industry

Chapter 5. Solvent and Other Product Use (CRF sector 3)

5.1. Overview of Sector

CO₂, N₂O, and NMVOC are emitted from solvent and other product use. In this chapter, CO₂ and N₂O emissions due to the following product uses are discussed (see Annex 3 for NMVOC):

- Paint application
- Degreasing and dry-cleaning
- Chemical products, manufacture and processing
- Other (e.g. anesthesia)

In 2010, total GHG emissions from the solvent and other product use sector amounted to 99 Gg-CO₂ eq., accounting for 0.01% of total national emissions (excluding LULUCF) from Japan. “3.D.1. Use of Nitrous Oxide for Anesthesia” is the only greenhouse gas emission source in this sector.

5.2. Paint Application (3.A.)

Paint solvents are used in Japan, but their application is basically restricted to mixing, therefore are assumed not to entail chemical reactions. Therefore, they do not generate CO₂ or N₂O during use. They have been reported as “NA.”

5.3. Degreasing and Dry-Cleaning (3.B.)

1) CO₂

Degreasing and dry-cleaning are practiced in Japan.

Degreasing is defined as, “washing processes that do not involve chemical reactions”, and it is assumed that it does not generate CO₂. Although the CO₂ emissions may occur in association with washing methods involving dry ice or carbonic gas, such methods are not thought to be used in Japan. There are no processes in dry-cleaning in which chemical reactions may occur, and it is basically assumed that it does not generate CO₂. However washing methods using liquefied carbonic gas are being used experimentally in research facilities and it is not possible to completely negate the possibility of CO₂ emissions.

As a result, these activities have been reported as “NE” due to the fact that there are no sufficient data available on the actual condition of emissions from degreasing and dry-cleaning and the absence of a default emission factor prevents any calculations from being performed.

2) N₂O

Degreasing and dry-cleaning are practiced in Japan, but degreasing is defined as, ‘washing processes that do not involve chemical reactions’, and there are no processes in dry-cleaning in which chemical reactions may occur. Therefore, it is assumed that N₂O is not generated. In Japan, there are also no methods which have the potential to emit N₂O used for degreasing or dry-cleaning, and they have therefore been reported as “NA”.

5.4. Chemical Products, Manufacture and Processing (3.C.)

NMVOC emissions occur from production and use of chemical products. NMVOC is reported in Annex 3.

5.5. Other (3.D.)

5.5.1. Use of Nitrous Oxide for Anesthesia (3.D.1)

a) Source/Sink Category Description

Nitrous oxide is emitted during anesthetics (laughing gas) use. Since 2006, some hospitals have installed N₂O destruction units, and the reductions achieved are reflected in the total emissions. Only N₂O is used as an anesthetic in Japan, and CO₂ is not. Therefore, CO₂ emissions have been reported as “NA”.

In 2010, total GHG emissions from this category amounted to 99 Gg-CO₂ eq., accounting for 0.01% of total national emissions (excluding LULUCF) from Japan.

Table 5-1 N₂O emissions during anesthetics (laughing gas) use

Gas	Category		Units	1990	1995	2000	2005	2008	2009	2010
				Gg-N ₂ O	Gg-CO ₂ eq.	Gg-CO ₂ eq.	Gg-CO ₂ eq.	Gg-CO ₂ eq.	Gg-CO ₂ eq.	Gg-CO ₂ eq.
N ₂ O	3.D Other	3.D.1 Use of N ₂ O for Anesthesia	Gg-N ₂ O	0.93	1.41	1.10	0.86	0.42	0.39	0.32
			Gg-CO ₂ eq.	287.07	437.58	340.99	266.41	129.10	120.50	98.95

b) Methodological Issues

● Estimation Method

In relation to emissions of N₂O from use of anesthetics, the actual amount of N₂O shipped as an anesthetic by pharmaceutical manufacturers or importers has been reported for 2005 and preceding years. For 2006 and beyond, the amount of N₂O collected is calculated using the amount of Laughing Gas used in domestic hospitals equipped with N₂O destruction units for anesthesia, and a destruction rate of 99.9 %. This is subtracted from the N₂O shipped for medical use to yield the amount of N₂O emitted.

Amount of N ₂ O emitted during the use of laughing gas = N ₂ O shipped for medical use – Amount of laughing gas used in hospitals equipped with N ₂ O destruction units × destruction rate
--

● Emission Factors

It is assumed that all of the N₂O used as medical gas escapes into the atmosphere, unless collected. Therefore, no emission factor has been established.

● Activity Data

The volume of shipments of N₂O for anesthetics (on calendar year basis) is given in the Ministry of Health, Labour and Welfare's Statistics of Production by Pharmaceutical Industry. This is used for 2005 and preceding years, and for 2006 to 2009, the amount of N₂O collected in three, and after 2010 collected in four domestic hospitals equipped with N₂O destruction units is subtracted from the above-mentioned shipment.

Table 5-2 Laughing gas shipment amount and N₂O collected in domestic hospitals
(calendar year basis)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Laughing gas shipment amount	kg-N ₂ O	926,030	1,411,534	1,099,979	859,389	417,919	389,749	320,111
N ₂ O collected in domestic hospitals	kg-N ₂ O	-	-	-	-	1,454	1,049	914

c) Uncertainties and Time-series Consistency

● *Uncertainty*

Because all N₂O used for anesthetics are assumed to escape into the atmosphere, no emission factor has been set. Therefore, the uncertainty for activity data is also the uncertainty for emissions. As Statistics of Production by Pharmaceutical Industry is a fundamental statistic based on statistical law, a 5% uncertainty was given for this emission source.

● *Time-series Consistency*

The volumes of shipments are taken from the Statistics of Production by Pharmaceutical Industry in a consistent manner throughout the time series.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the GPG (2000). Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.

e) Source-specific Recalculations

There have been no source-specific recalculations.

f) Source-specific Planned Improvements

No improvements are planned.

5.5.2. N₂O from Fire Extinguishers (3.D.2)

1) CO₂

Many types of fire extinguishers in Japan are filled with CO₂, which is emitted into the atmosphere when a fire extinguisher is used. All of the CO₂ with which the fire extinguishers are filled, however, is the by-product gas generated from petrochemicals or petroleum refining. Such emissions are included in the calculation of Chapter 1, section 1.A.1.b. Petroleum Refining, and therefore, have been reported as "IE".

2) N₂O

N₂O is not used in the fire extinguishers in Japan. Therefore the N₂O emissions from this category are reported as "NO".

5.5.3. N₂O from Aerosol Cans (3.D.-)

1) CO₂

Aerosol products, which fill spray cans with CO₂, are manufactured in Japan. It is assumed that CO₂ could be emitted into the atmosphere when the aerosol products are used. However, because the CO₂ used in the aerosol industry is a by-product gas of petrochemical products, these emissions are counted in the Combustion of Fuel sector (1.A.), and have been reported as “IE” here.

2) N₂O

Aerosol products manufactured in Japan do not use N₂O. Theoretically, no N₂O is emitted, and it has been reported as “NA” here.

References

1. Ministry of the Environment Committee for the Greenhouse Gases Emissions Estimation Methods, *Review of Greenhouse Gases Emissions Estimation Methods Part 2*, August 2002.
2. Ministry of Health, Labour and Welfare's *Statistics of Production by Pharmaceutical Industry*.

Chapter 6. Agriculture (CRF sector 4)

6.1. Overview of Sector

Greenhouse gas emissions from the agricultural sector are calculated in five categories: 4A, 4B, 4C, 4D, and 4F. In 4A: Enteric Fermentation, CH₄ gas generated and emitted by cattle, buffalo, sheep, goats, horses, and swine as the result of enteric fermentation is reported. In 4B: Manure Management, CH₄ and N₂O generated by treatment of manure excreted by cattle, buffalo, sheep, goats, horses, swine and poultry are reported. In 4C: Rice Cultivation, CH₄ emissions from paddy fields (continuously flooded and intermittently flooded) cultivated for rice production are reported. In 4D: Agricultural Soils, CH₄ and N₂O emitted directly and indirectly from agricultural soil as well as pastures, ranges, and paddocks manure are reported. Emissions for 4E Prescribed Burning of Savannas are reported as NO, since Japan has no emission source in this category, while CH₄ and N₂O (as well as CO, which is described in Annex 3) emissions from field burning of grains, legumes, root crops, and sugar cane during agricultural activities are reported in 4F: Field Burning of Agricultural Residues.

The Revised 1996 IPCC Guidelines require emissions from the agricultural sector to be reported as a three-year average. The Japanese inventory uses the year before and the year after the relevant year to report a three-year average for emissions.

GHG emissions in the Agricultural Sector in FY 2010 were 25,500 Gg-CO₂ eq., comprising 2.0% of total emissions (excluding LULUCF). The value represents a reduction by 18.4% from FY 1990.

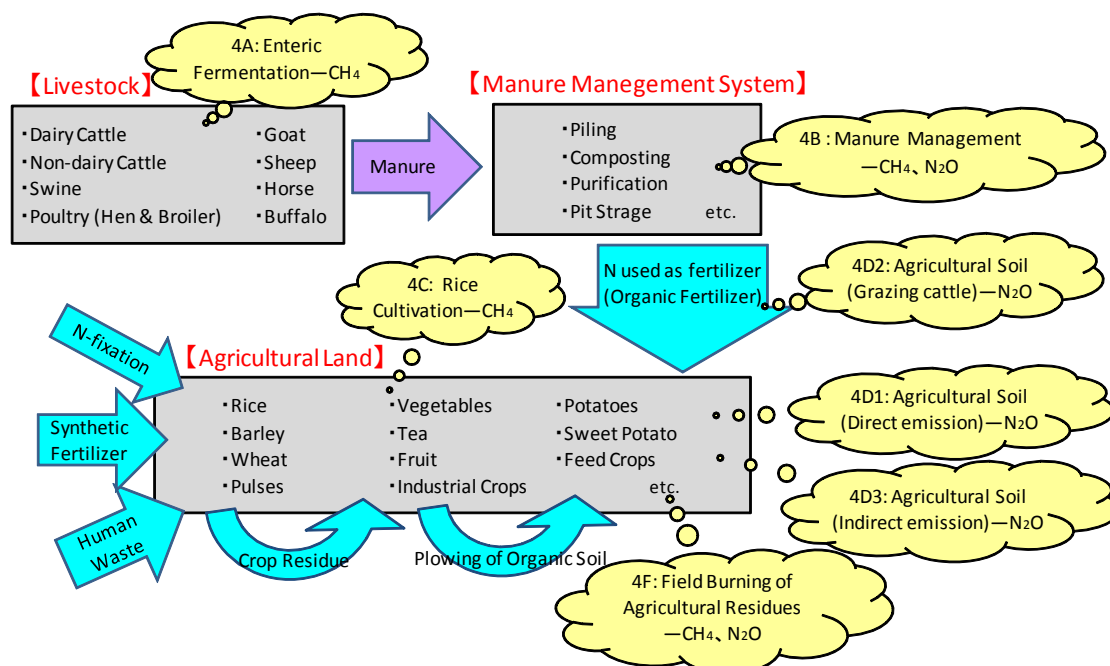


Figure 6-1 Relationships among the categories in the agricultural sector

6.2. Enteric Fermentation (4.A.)

Ruminants such as cattle, buffalo, sheep, and goats have multi-chamber stomachs. The rumen carries out anaerobic fermentation to break down cellulose and other substances, thereby releasing CH₄.

Horses and swine are not ruminants and have monogastric stomachs, but fermentation in their digestive tracts produces small amounts of CH₄, which is released into the atmosphere. These CH₄ emissions are calculated and reported in the *Enteric Fermentation (4.A.)* section.

GHG emissions from Enteric Fermentation in FY 2010 were 6,673 Gg-CO₂eq., comprising 0.5% of total emissions (excluding LULUCF). The Value represents a reduction by 13.1% from FY 1990.

Table 6-1 CH₄ emissions from enteric fermentation

Gas	Livestock species	Unit	1990	1995	2000	2005	2008	2009	2010
CH ₄	4.A.1.- Dairy Cattle	Gg-CH ₄	192.6	184.4	172.8	162.9	154.7	151.7	150.0
	4.A.1.- Non-Dairy Cattle	Gg-CH ₄	158.2	164.6	165.5	158.2	162.1	158.5	155.5
	4.A.2. Buffalo	Gg-CH ₄	0.012	0.007	0.005	0.004	0.004	0.004	0.004
	4.A.3. Sheep	Gg-CH ₄	0.09	0.06	0.05	0.04	0.05	0.06	0.06
	4.A.4. Goats	Gg-CH ₄	0.11	0.08	0.09	0.07	0.06	0.06	0.06
	4.A.6. Horses	Gg-CH ₄	2.1	2.1	1.9	1.6	1.5	1.5	1.5
	4.A.8. Swine	Gg-CH ₄	12.5	11.0	10.7	10.6	10.8	10.8	10.7
	Total	Gg-CH ₄	365.6	362.2	351.0	333.4	329.2	322.5	317.8
	Gg-CO ₂ eq	7,677	7,606	7,370	7,002	6,913	6,773	6,673	

6.2.1. Cattle (4.A.1.)

a) Source/Sink Category Description

This section provides the estimation methods for CH₄ emissions from enteric fermentation in Cattle.

b) Methodological Issues

●Estimation Method

In accordance with Decision Tree of the *GPG (2000)* (Page 4.24 Fig. 4.2), calculations for dairy and non-dairy cattle should be performed using the Tier 2 method. The Tier 2 method requires the total energy intake of livestock to be multiplied by the CH₄ conversion factor to derive the emission factor, but it has been in practice in Japan on livestock-related research to use amount of dry matter intake. It is considered that, by applying the results of previous researches, the estimation method using amount of dry matter intake provides more accurate data. For that reason, a technique similar to the Tier 2 Method but specific to Japan was used for the calculation of CH₄ emissions associated with enteric fermentation by cattle. The emissions were calculated by multiplying the cattle population (dairy and non-dairy) by the emission factors established based on their dry matter intake.

As cattle begin to eat normal feed at the age of five to six months, the calculation of the CH₄ emissions associated with enteric fermentation includes cattle aged five months or older (cattle under five months are excluded from estimation). To reflect the actual situation of emissions in Japan, categorization of cattle is defined as shown below, and the estimation of CH₄ emissions is conducted by type and age.

Table 6-2 Categorization and assumptions underlying calculation of CH₄ emissions associated with enteric fermentation in cattle

Animal type		Assumptions for Calculation of Emissions	
Dairy cattle	Lactating	—	
	Non-lactating	—	
	Heifers (under 2 years old, excluding 5- and 6-months old)	Calculation excludes 6/24 of the population which was assumed to be 6 months old or younger; therefore actually covering only 18/24 of the population under 2 years old.	
	Heifers (5 to 6 months old)	Calculation covers 5- and 6-month old comprising 2/24 of the population under 2 years old.	
	Heifers (under 5 months old)	Covering 4/24 of the population under 2 years old. Excluded from CH ₄ emission estimation.	
Non-dairy cattle	Breeding cows	(1 year and older)	—
		(under 1 year, excluding 5- and 6-months old)	Calculation excludes 6/12 of the population which was assumed to be 6 months old or younger; therefore covering 6/12 of the population under 1 year old.
		(5 and 6 months old)	Calculation covers 5- and 6-month old comprising 2/12 of the population under 1 year old.
		(under 5 months old)	Covering 4/12 of the population under 1 year old. Excluded from CH ₄ emission estimation.
	fattening cattle	Japanese cattle (1 year and older)	—
		Japanese cattle (under 1 year, excluding 5- and 6-months old)	Calculation excludes 6/12 of the population which was assumed to be 6 months old or younger; therefore covering 6/12 of the population under 1 year old.
		Japanese cattle (5 to 6 months old)	Calculation covers 5- and 6-month old comprising 2/12 of the population under 1 year old.
		Japanese cattle (under 5 months old)	Covering 4/12 of the population under 1 year old. Excluded from CH ₄ emission estimation.
		Dairy breeds (excluding 5- and 6-months old)	Calculation excludes 6/24 of the population which was assumed to be 6 months old or younger; therefore covering 18/24 of the population under 2 years old.
		Dairy breeds (5 to 6 months old)	Calculation covers 5- and 6-month old comprising 2/24 of the population under 2 years old.
		Dairy breeds (under 5 months old)	Covering 4/24 of the population under 2 years old. Excluded from CH ₄ emission estimation.

● Emission Factors

The emission factor for CH₄ associated with enteric fermentation in cattle has been established on the basis of breath testing of ruminant livestock in Japan; it is based on the measured data for volume of CH₄ generated from dry matter intake. Results of measurements have made it clear that it is possible to estimate CH₄ from enteric fermentation in ruminant livestock using the equation given below, which uses dry matter intake as the explanatory variable (Shibata et al. (1993), Reference 30).

<p><u>CH₄ Emission Factor for associated with enteric fermentation in cattle (kg-CH₄/head)</u> = (Volume of CH₄ generated [l/day/head]) / (Volume of 1 mol) × (molecular weight of CH₄) × (No. of days in year) = Y / 22.4 (l/mol/head) × 0.016 (kg/mol) × 365 or 366 (day)</p> <p><u>Volume of CH₄ generated per head per day (=Y) (l/mol/head)</u> = -17.766 + 42.793 DMI - 0.849 (DMI)²</p> <p>DMI : Dry matter intake [kg/day/head]</p>
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Average dry matter intake estimated from Japan Feed Standards compiled by the Japan Livestock Industry Association is applied to the above equation to establish emission factors. The dry matter intake was calculated by substituting fat corrected milk, body weight, and weight gain by daily growth into the equation established for each type of cattle. Data for the fat corrected milk was obtained from

the Statistics on Milk and Dairy Products (Ministry of Agriculture, Fisheries and Forestry; MAFF) and the Statistics on Livestock (MAFF), and those for the fat content in milk from the Statistics of Livestock Production Costs (MAFF). Both sets of the data are updated on a yearly basis. Data for body weight and weight gain by daily growth were obtained from the table of weight by age (months) for each type of cattle included at the back of the Japanese Feeding Standards (Japan Livestock Industry Association). Equations to estimate Dry Matter Intake were revised in 2006 for daily cattle (Lactating and Non-lactating) and in 2008 for non-dairy cattle (Japanese cattle(M)).

Table 6-3 Equation to estimate Dry Matter Intake (DMI) by cattle

Animal type		Equation
Dairy cattle	Lactating	After 2006: $DMI=1.3922+0.05839 \times W^{0.75}+0.40497 \times FCM$ $FCM=(15 \times FAT/100+0.4) \times MILK$ Before 2005: $DMI=2.98120+0.00905 \times W+0.41055 \times FCM$ $FCM=(15 \times FAT /100+0.4) \times MILK$
	Non-lactating	After 2006: $DMI=0.017 \times W$ Before 2005: $DMI=(0.1163 \times W^{0.75}/0.82)/4.41/0.52*1.1$
	Heifers	$DMI=0.49137+0.01768 \times W+0.91754 \times DG$
Non-dairy cattle	Breeding cows	$DMI= [0.1067 \times W^{0.75} + (0.0639 \times W^{0.75} \times DG) / (0.78 \times q + 0.006)] / (q \times 4.4)$ $q=0.4213+0.1491 \times DG$
	Japanese cattle (M)	After 2008: $DMI=-3.481+2.668 \times DG+4.548 \times 10^{-2} \times W-7.207 \times 10^{-5} \times W^2+3.867 \times 10^{-8} \times W^3$ Before 2007: $DMI= [0.1124 \times W^{0.75} + (0.0546 \times W^{0.75} \times DG) / (0.78 \times q + 0.006)] / \{q \times (1.653 - 0.00123 \times W)\}$ $q=0.5304+0.0748 \times DG$
	Japanese cattle (F)	$DMI=[0.1108 \times W^{0.75} + (0.0609 \times W^{0.75} \times DG) / (0.78 \times q + 0.006)] / (q \times 4.4)$ $q=0.5018+0.0956 \times DG$
	Dairy breeds (excluding 5- and 6-months old)	$DMI=[0.1291 \times W^{0.75} + (0.0510 \times W^{0.75} \times DG) / (0.78 \times q + 0.006)] / (q \times 4.4)$ $q=(0.933+0.00033 \times W) \times (0.498+0.0642 \times DG)$
	Dairy breeds (5 to 6 months old)	$DMI=[0.1291 \times W^{0.75} + \{ (1.00+0.030 \times W^{0.75}) \times DG \} / (0.78 \times q + 0.006)] / (q \times 4.4)$ $q=(0.859-0.00092 \times W) \times (0.790+0.0411 \times DG)$

W: Weight, FCM: Fat Corrected Milk, FAT: Fat content in milk, MILK: Milk Yield, DG: Daily Growth, q: Energy metabolic rate

Source: Japan Livestock Industry Association, *Japan Feed Standards*

Table 6-4 Fat content in milk (FAT) and Milk Yield (MILK) by cattle

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Milk yield (Lactating)	kg/head/day	24.9	26.8	28.1	30.1	30.8	30.6	30.6
Fat content in milk (Lactating)	%	3.7	3.8	3.9	4.0	3.9	3.9	3.9

Table 6-5 Weight by cattle (W)

Animal Type		Unit	1990	1995	2000	2005	2009	2010	2011	
Dairy Cattle	Lactating	kg/head	595.9	602.8	621.4	622.7	623.0	623.0	623.0	
	Non-lactating	kg/head	595.9	602.8	621.4	622.7	623.0	623.0	623.0	
	Heifer: Under Two Years, over six months	kg/head	342.4	349.3	364.9	374.2	376.1	376.1	376.1	
	Heifer: Five and six months	kg/head	140.0	140.6	146.3	162.8	166.1	166.1	166.1	
Non-Dairy Cattle	Breeding Cows	One Year and Over	kg/head	426.6	426.6	487.3	450.9	429.1	429.1	429.1
		Under One Year, over six months	kg/head	230.2	230.2	279.7	259.3	247.0	247.0	247.0
	fattening cattle	Five and six months	kg/head	141.0	141.0	157.1	146.8	140.7	140.7	140.7
		Japanese cattle (M): One Year and Over	kg/head	574.3	574.3	574.3	572.3	571.0	571.0	571.0
		Japanese cattle (M): Under One Year, over six months	kg/head	273.4	273.4	273.4	274.6	275.4	275.4	275.4
		Japanese cattle (M): Five and six months	kg/head	146.7	146.7	146.7	147.9	148.6	148.6	148.6
		Japanese cattle (F): One Year and Over	kg/head	388.0	388.0	462.5	427.7	406.8	406.8	406.8
		Japanese cattle (F): Under One Year, over six months	kg/head	230.2	230.2	279.7	259.3	247.0	247.0	247.0
		Japanese cattle (F): Five and six months	kg/head	141.0	141.0	157.1	146.8	140.7	140.7	140.7
		Dairy breed: Over six months	kg/head	479.8	479.8	479.8	479.8	479.8	479.8	479.8
Dairy breed: Five and six months	kg/head	194.8	194.8	194.8	194.8	194.8	194.8	194.8		

Table 6-6 Daily Growth by cattle (DG)

Animal Type		Unit	1990	1995	2000	2005	2009	2010	2011	
Dairy Cattle	Lactating	kg/head/day	—	—	—	—	—	—	—	
	Non-lactating	kg/head/day	—	—	—	—	—	—	—	
	Heifer: Under Two Years, over six months	kg/head/day	0.60	0.63	0.65	0.59	0.58	0.58	0.58	
	Heifer: Five and six months	kg/head/day	0.69	0.70	0.76	0.88	0.90	0.90	0.90	
Non-Dairy Cattle	Breeding Cows	One Year and Over	kg/head/day	0.17	0.17	0.14	0.13	0.13	0.13	0.13
		Under One Year, over six months	kg/head/day	0.70	0.70	0.94	0.86	0.81	0.81	0.81
		Five and six months	kg/head/day	0.74	0.74	1.04	0.96	0.91	0.91	0.91
	fattening cattle	Japanese cattle (M): One Year and Over	kg/head/day	0.60	0.60	0.60	0.59	0.58	0.58	0.58
		Japanese cattle (M): Under One Year, over six months	kg/head/day	1.07	1.07	1.07	1.07	1.07	1.07	1.07
		Japanese cattle (M): Five and six months	kg/head/day	0.94	0.94	0.94	0.95	0.95	0.95	0.95
		Japanese cattle (F): One Year and Over	kg/head/day	0.28	0.28	0.27	0.25	0.24	0.24	0.24
		Japanese cattle (F): Under One Year, over six months	kg/head/day	0.70	0.70	0.94	0.86	0.81	0.81	0.81
		Japanese cattle (F): Five and six months	kg/head/day	0.74	0.74	1.04	0.96	0.91	0.91	0.91
		Dairy breed: Over six months	kg/head/day	0.92	0.92	0.92	0.92	0.92	0.92	0.92
		Dairy breed: Five and six months	kg/head/day	1.10	1.10	1.10	1.10	1.10	1.10	1.10

Table 6-7 Dry matter intake by cattle (DMI)

Animal Type		Unit	1990	1995	2000	2005	2009	2010	2011	
Dairy Cattle	Lactating	kg/head/day	18.2	19.2	20.0	20.9	21.0	20.9	20.9	
	Non-lactating	kg/head/day	8.2	8.3	8.5	8.5	10.6	10.6	10.6	
	Heifer: Under Two Years, over six months	kg/head/day	7.1	7.2	7.5	7.7	7.7	7.7	7.7	
	Heifer: Five and six months	kg/head/day	3.6	3.6	3.8	4.2	4.3	4.3	4.3	
Non-Dairy Cattle	Breeding Cows	One Year and Over	kg/head/day	6.6	6.6	7.1	6.6	6.3	6.3	6.3
		Under One Year, over six months	kg/head/day	5.5	5.5	6.7	6.2	5.9	5.9	5.9
		Five and six months	kg/head/day	3.8	3.8	4.4	4.1	4.0	4.0	4.0
	fattening cattle	Japanese cattle (M): One Year and Over	kg/head/day	8.4	8.4	8.4	8.3	7.7	7.7	7.7
		Japanese cattle (M): Under One Year, over six months	kg/head/day	6.8	6.8	6.8	6.8	7.2	7.2	7.2
		Japanese cattle (M): Five and six months	kg/head/day	4.3	4.3	4.3	4.4	4.4	4.4	4.4
		Japanese cattle (F): One Year and Over	kg/head/day	5.7	5.7	6.4	6.0	5.7	5.7	5.7
		Japanese cattle (F): Under One Year, over six months	kg/head/day	4.9	4.9	6.1	5.6	5.3	5.3	5.3
		Japanese cattle (F): Five and six months	kg/head/day	3.4	3.4	4.1	3.8	3.6	3.6	3.6
		Dairy breed: Over six months	kg/head/day	8.7	8.7	8.7	8.7	8.7	8.7	8.7
		Dairy breed: Five and six months	kg/head/day	5.3	5.3	5.3	5.3	5.3	5.3	5.3

Table 6-8 Emission factor associated with enteric fermentation by cattle

Animal Type		Unit	1990	1995	2000	2005	2009	2010	2011	
Dairy Cattle	Lactating	kgCH ₄ /head/year	125.0	128.3	130.0	131.9	132.1	131.8	132.2	
	Non-lactating	kgCH ₄ /head/year	72.0	72.7	74.0	74.1	88.7	88.7	88.9	
	Heifer: Under Two Years, over six months	kgCH ₄ /head/year	63.4	64.7	66.9	67.8	68.0	68.0	68.1	
	Heifer: Five and six months	kgCH ₄ /head/year	32.7	32.9	34.4	38.1	38.8	38.8	38.9	
Non-Dairy Cattle	Breeding Cows	One Year and Over	kgCH ₄ /head/year	59.0	59.2	63.1	59.3	57.0	57.0	57.1
		Under One Year, over six months	kgCH ₄ /head/year	49.8	50.0	60.1	56.3	53.8	53.8	54.0
		Five and six months	kgCH ₄ /head/year	34.9	35.0	40.4	37.8	36.2	36.2	36.3
	fattening cattle	Japanese cattle (M): One Year and Over	kgCH ₄ /head/year	73.2	73.4	73.2	72.8	68.5	68.5	68.7
		Japanese cattle (M): Under One Year, over six months	kgCH ₄ /head/year	61.1	61.3	61.1	61.2	64.5	64.5	64.7
		Japanese cattle (M): Five and six months	kgCH ₄ /head/year	39.6	39.7	39.6	39.9	39.8	39.8	39.9
		Japanese cattle (F): One Year and Over	kgCH ₄ /head/year	51.8	51.9	58.1	54.2	51.9	51.9	52.0
		Japanese cattle (F): Under One Year, over six months	kgCH ₄ /head/year	44.3	44.5	55.3	51.2	48.7	48.7	48.8
		Japanese cattle (F): Five and six months	kgCH ₄ /head/year	31.0	31.0	37.4	34.6	32.9	32.9	33.0
		Dairy breed: Over six months	kgCH ₄ /head/year	75.6	75.8	75.6	75.6	75.6	75.6	75.8
		Dairy breed: Five and six months	kgCH ₄ /head/year	48.0	48.1	48.0	48.0	48.0	48.0	48.1

●Activity Data

For activity data of this source, the herd size for each type of livestock at 1 February in each year, recorded by the Ministry of Agriculture, Forestry and Fisheries in its *Livestock Statistics* is used.

Table 6-9 Livestock population for cattle (Single year)

Animal Type		Unit	1990	1995	2000	2005	2009	2010	2011		
Dairy Cattle	Lactating	1000 head	1,082	1,035	971	900	830	805	805		
	Non-lactating	1000 head	332	299	249	231	200	195	195		
	Heifer: Under two years, over six months	1000 head	491	445	379	379	341	351	351		
	Heifer: Five and six months	1000 head	55	49	42	42	38	39	39		
	Heifer: Under five months	1000 head	109	99	84	84	76	78	78		
Dairy Cattle Total		1000 head	2,068	1,927	1,725	1,636	1,484	1,467	1,467		
Non-Dairy Cattle	Breeding Cows	One Year and Over	1000 head	679	646	612	594	651	636	636	
		Under One Year, over six months	1000 head	17	13	12	14	17	16	16	
		Five and six months	1000 head	6	4	4	5	6	5	5	
		Under five months	1000 head	12	9	8	9	11	11	11	
	fattening cattle	Japanese cattle (M): One Year and Over	1000 head	368	412	385	374	425	409	409	
		Japanese cattle (M): Under One Year, over six months	1000 head	125	133	114	119	132	127	127	
		Japanese cattle (M): Five and six months	1000 head	42	44	38	40	44	42	42	
		Japanese cattle (M): Under five months	1000 head	83	89	76	80	88	85	85	
		Japanese cattle (F): One Year and Over	1000 head	197	265	246	290	339	336	336	
		Japanese cattle (F): Under One Year, over six months	1000 head	102	105	93	89	106	101	101	
		Japanese cattle (F): Five and six months	1000 head	34	35	31	30	35	34	34	
		Japanese cattle (F): Under five months	1000 head	68	70	62	59	70	67	67	
		Dairy breed: Over six months	1000 head	805	808	845	789	726	671	671	
		Dairy breed: Five and six months	1000 head	89	90	94	88	81	75	75	
		Dairy breed: Under five months	1000 head	179	180	188	175	161	149	149	
		Non-Dairy Cattle Total		1000 head	2,805	2,901	2,805	2,755	2,892	2,763	2,763

* Data for 2011 are substituted by data for 2010

* three-year average data are reported in the CRF.

c) Uncertainties and Time-series Consistency

● Uncertainties

An uncertainty assessment was conducted for the categories indicated in Table 6-2, there were 4 categories for dairy cattle and 11 categories for non-dairy cattle. The uncertainties for emission factors were calculated by finding the 95% confidence interval in accordance with the equation indicated in the section *Emission Factors*. Populations of cattle (Activity data) are decided by survey of total population in the *Livestock Statistics*, but standard error for cattle is not described. Therefore, the uncertainties for activity data were determined to be 5% in accordance with decision tree indicated in Annex 7. As a result, the uncertainties of the emissions were determined to be 15% for dairy cattle and 19% for non-dairy cattle. The uncertainty assessment methods are summarized in Annex 7.

● Time-series Consistency

Emission factors were calculated consistently from FY 1990 onward by the method mentioned in the section on Emission Factors. Activity data were used consistently from FY 1989 onward from the data in *Livestock Statistics*.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1.

Comparison between results of Japan's estimation method and IPCC Tier 2 method was conducted. As a result, for dairy cattle, considering the error of CH₄ conversion factor ($Y_m = 0.60 \pm 0.05$), the emissions based on Japan's method were in the range calculated by IPCC Tier 2 method. Therefore, it is considered that there were no significant differences between emissions of Japan's method and IPCC Tier 2 method. On the other hand, for non-dairy cattle, it became clear that the emissions of Japan's method are about 10% higher than IPCC Tier 2 method. Analysis of the factor about this

difference will be continued.

e) *Source-specific Recalculations*

Since the fat content in milk (FAT) and the amount of milk production per lactating cow were updated, the emissions from dairy cattle were revised from FY2007 to FY2009. In the Agriculture sector, a 3-year average has been used. Thus, the emissions for FY2009 were revised in accordance with the revision and/or update of the activity data for FY2010.

f) *Source-specific Planned Improvements*

- It is planned to discuss the development of the estimation method, which reflects the emissions reduction with technologies that suppress methane fermentation by controlling the rumen fermentation (such as by the addition of fatty acid calcium to feed) and by improving the feed efficiency with the Total Mixed Ration (TMR) feeding.

6.2.2. Buffalo, Sheep, Goats, Horses & Swine (4.A.2., 4.A.3., 4.A.4., 4.A.6., 4.A.8.)

a) *Source/Sink Category Description*

This section provides the estimation methods for CH₄ emissions from enteric fermentation in Buffalo, Sheep, Goats, Horses and Swine.

b) *Methodological Issues*

● *Estimation Method*

CH₄ emissions were calculated using the Tier 1 Method in accordance with the Decision Tree of the *GPG (2000)*.

● *Emission Factors*

The emission factor for CH₄ associated with sheep and goats has been established in the same way as for cattle, based on the emissions of CH₄ estimated from dry matter intake.

In Japan, most of sheep are farmed for meat and they are smaller than sheep for wool production assumed in the *Revised 1996 IPCC Guidelines* and *GPG (2000)* as default. Therefore, it is considered that emission factor for sheep in Japan is lower than default in IPCC guidelines. As for goats, research findings in this regard do not exist in Japan. However, the emission factor for goat was regarded as equivalent to the one for sheep by the experts (the expert judgment). Therefore, the emission factor for sheep is also used for goats.

The emission factor for swine has been established on the basis of results of research conducted in Japan. The emission factor used for horses and buffalo is the default value given in the *Revised 1996 IPCC Guidelines*.

Table 6-10 Emission factors for CH₄ associated with enteric fermentation in sheep, goats, horses, swine and buffalo

Animal type	Dry Matter Intake [kg]	CH ₄ Emission factor [kg/year/head]
Sheep, goats ^a	0.8	4.1
Swine ^b	—	1.1
Horses ^c	—	18.0
Buffalo ^c	—	55.0

a: Calculated by the formula: (CH₄ generated [l/day/head]) / (Volume of 1 mol) × (molecular weight of CH₄) × (no. of days in year)

b: Mamoru Saito, *Methane emissions from fattening swine and expectant swine* (1988) (Reference 29)

c: *Revised 1996 IPCC Guidelines*

●Activity Data

The values used for activity data are used for sheep and goats given in the *Statistical Document of Livestock Breeding* offered by the Japan Livestock Industry Association. The values used for activity data for swine are the herd size at February 1st in each year, as recorded by the Ministry of Agriculture, Forestry and Fisheries in its *Livestock Statistics*. The values used for activity data for horses given in the *Statistical Document of Horse* offered by the Ministry of Agriculture, Forestry and Fisheries, for buffalo given *Statistics on Livestock in Okinawa Prefecture*.

Table 6-11 Activity data associated with enteric fermentation by buffalo, sheep, goats, swine, and horses

Type of animal	Unit	1990	1995	2000	2005	2009	2010	2011
Sheep	1000 head	21	14	12	9	14	14	14
Goats	1000 head	26	19	22	16	14	14	14
Swine	1000 head	11,336	9,900	9,788	9,621	9,834	9,768	9,768
Horses	1000 head	116	118	105	87	82	82	82
Buffalo	1000 head	0.21	0.12	0.10	0.08	0.08	0.08	0.08

* Data for 2011 are substituted by data for 2010

c) Uncertainties and Time-series Consistency

●Uncertainties

An uncertainty assessment was conducted by each livestock category. The uncertainties for emission factors were applied 50% of default data given in the *GPG (2000)*. As the uncertainty for activity data, 0.85% of standard error for swine given in the *Livestock Statistic* was applied to swine. Since sample standard deviation can't be obtained and expert judgment is impossible, and non-fundamental statistics, 100% was applied to other livestock in accordance with the decision tree of uncertainty assessment. As a result, the uncertainties of the emissions were determined to be 50% for swine and 112% for buffalo, sheep and goats. The uncertainty assessment methods are summarized in Annex 7.

●Time-series Consistency

For emission factors, same values were used consistently from FY 1990 to FY 2010. Activity data for sheep and goats applied the data given in the *Statistical Document of Livestock Breeding*, those for swine applied the data given in the *Livestock Statistics*; those for horses applied the data given in *Statistical Document of Horse*, and those for buffalo applied the data given in the *Livestock Statistics of Okinawa*, consistently since FY 1989.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1.

With regard to the significant difference between Japan's EF and the default EF in the IPCC Guidelines for sheep, the reason is described in the 'Emissions factors' section above.

e) Source-specific Recalculations

In the Agriculture sector, a 3-year average has been used. Thus, the emissions from FY2006 to FY2009 were revised in accordance with the revision and/or update of the activity data from FY2007 to FY2010.

f) Source-specific Planned Improvements

Although the default emission factor in the *Revised 1996 IPCC Guidelines* or the *GPG (2000)* has been used for some livestock categories, there is a need to discuss whether it is possible to establish country-specific emission factors for Japan.

6.2.3. Poultry (4.A.9.)

It is conceivable that CH₄ is emitted from enteric fermentation in poultry, but the Japanese literature offers no data on emission factors, and neither the *Revised 1996 IPCC Guidelines* nor the *GPG (2000)* offer default emission factors. Therefore, this category has been reported as “NE”. In addition, poultry other than hens and broiler are not covered by official statistics, suggesting that they may be assumed to be negligible.

6.2.4. Camels and Llamas, Mules and Asses (4.A.5., 4.A.7.)

Japan reported “NO” in this subcategory as it was unlikely that these animals were raised for agricultural purposes.

6.2.5. Other (4.A.10.)

The only livestock that are bred in Japan are cattle, buffalo, sheep, goats, horses, swine and poultry. Therefore, this category has been reported as “NO”.

6.3. Manure Management (4.B.)

Livestock manure generates CH₄ when its organic content is converted to CH₄ gas through CH₄ fermentation, or when CH₄ from enteric fermentation dissolved in manure is released by aeration or agitation. In manure management, N₂O is produced mainly by microorganism via nitrification and denitrification processes.

CH₄ and N₂O emissions from manure management in FY 2010 are 2,205 Gg-CO₂ eq. and 5,475 Gg-CO₂ eq., comprising 0.2% and 0.4% of total emissions (excluding LULUCF), respectively. The value represents a reduction by 28.7% and 1.0% from FY 1990, respectively.

Table 6-12 CH₄ and N₂O emissions from livestock manure management

Gas	Livestock species	Unit	1990	1995	2000	2005	2008	2009	2010
CH ₄	4.B.1.- Dairy Cattle	Gg-CH ₄	123.2	115.7	106.2	98.2	90.8	90.9	91.7
	4.B.1.- Non-Dairy Cattle	Gg-CH ₄	4.5	4.6	4.5	4.4	4.5	4.4	4.3
	4.B.2. Buffalo	Gg-CH ₄	0.0004	0.0002	0.0002	0.0002	0.0002	0.0002	0.0002
	4.B.3. Sheep	Gg-CH ₄	0.006	0.004	0.003	0.003	0.003	0.004	0.004
	4.B.4. Goats	Gg-CH ₄	0.005	0.003	0.004	0.003	0.003	0.002	0.002
	4.B.6. Horses	Gg-CH ₄	0.2	0.2	0.2	0.2	0.2	0.2	0.2
	4.B.8. Swine	Gg-CH ₄	15.9	13.9	13.6	13.5	11.0	8.4	5.7
	4.B.9. Poultry	Gg-CH ₄	3.5	3.2	3.0	2.9	3.0	3.1	3.2
	Total	Gg-CH ₄	147.3	137.8	127.5	119.2	109.6	107.0	105.0
Gg-CO ₂ eq		3,094	2,893	2,678	2,503	2,302	2,247	2,205	
N ₂ O	4.B.1.- Dairy Cattle	Gg-N ₂ O	2.7	2.6	2.3	2.2	2.4	2.7	3.1
	4.B.1.- Non-Dairy Cattle	Gg-N ₂ O	2.8	2.9	2.8	2.8	3.0	3.2	3.3
	4.B.2. Buffalo	Gg-N ₂ O	0.00012	0.00007	0.00005	0.00004	0.00004	0.00004	0.00004
	4.B.3. Sheep	Gg-N ₂ O	0.007	0.005	0.004	0.003	0.004	0.004	0.005
	4.B.4. Goats	Gg-N ₂ O	0.03	0.02	0.03	0.02	0.02	0.02	0.02
	4.B.6. Horses	Gg-N ₂ O	0.1	0.1	0.1	0.1	0.1	0.1	0.1
	4.B.8. Swine	Gg-N ₂ O	4.8	4.2	4.1	4.1	4.9	5.6	6.3
	4.B.9. Poultry	Gg-N ₂ O	7.4	6.8	6.4	6.2	5.8	5.3	4.9
	Total	Gg-N ₂ O	17.8	16.6	15.8	15.3	16.2	16.9	17.7
Gg-CO ₂ eq		5,533	5,152	4,885	4,748	5,019	5,247	5,475	
Total of all gases		Gg-CO ₂ eq	8,627	8,045	7,563	7,251	7,321	7,495	7,680

6.3.1. Cattle, Swine and Poultry (4.B.1., 4.B.8., 4.B.9.)

a) Source/Sink Category Description

This section provides the estimation methods for CH₄ and N₂O emissions for manure management from cattle (Daily cattle and Non-daily cattle), swine and poultry (Hen and Broilers). The estimations for cattle were conducted separately for “shed” and “pastured” cattle. CH₄ emissions were reported in this category and N₂O emissions for “pastured” were reported in “4.D.2 Pasture, Range and Paddock Manure”.

b) Methodological Issues

1) Cattle, Swine and Poultry in shed and barn

● Estimation Method

CH₄ emissions in shed associated with the manure management were calculated by multiplying the amount of organic matter contained in manure from each type of livestock by the emission factor for each type of treatment method.

$$E = \sum (EF_n \times A_n)$$

E: CH₄ emissions associated with the management of manure excreted by cattle, swine and poultry (g-CH₄)

EF_n: Emission factor for treatment method *n* (g-CH₄/g-Organic matter);

A_n: Amount of organic matter contained in manure treated by method *n* (g-Organic matter).

N₂O emissions were calculated by multiplying the amount of nitrogen contained in manure of each type of animal by the emission factor for each type of treatment method.

$$E = \sum (EF_n \times A_n) \times 44/28$$

E : N₂O emission associated with management of manure excreted by cattle, swine and poultry (g-N₂O)

EF_n : Emission factor for treatment method n (g-N₂O/g-N);

A_n : Amount of nitrogen contained in manure treated by method n (g-N)

● Emission Factors

Emission factors for CH₄ and N₂O associated with Animal Waste Management System (hereafter, AWMS) have been established for each treating method of for each type of livestock, on the basis of the results of research carried out in Japan after reviewing its validity in accordance with the decision tree shown in Figure 6-2.

Moisture for dairy cattle feces is high, and they easily make anaerobic condition. It is considered to be the reason for high CH₄ emission factor of piling.

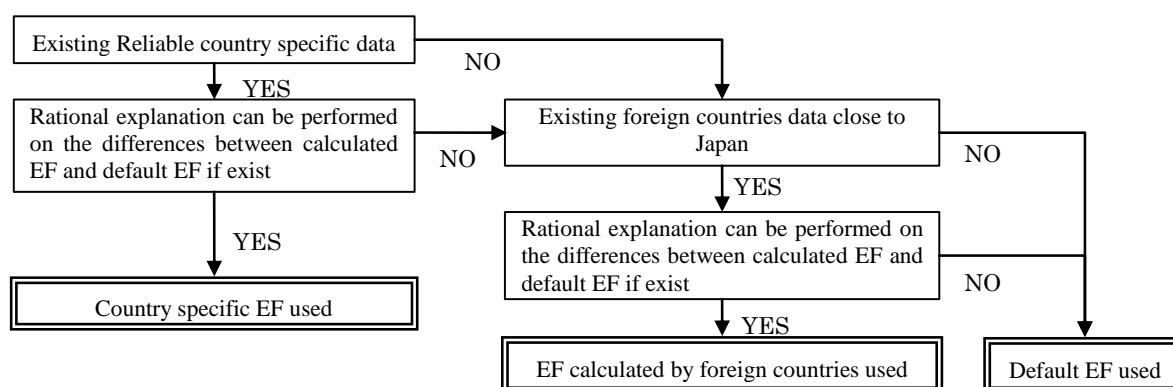


Figure 6-2 Decision tree for determination of EF

Table 6-13 CH₄ Emission factors for each method of treating manure from cattle, Swine, Hen & Broiler (g-CH₄/g-Organic matter)

treating method		Daily Cattle		Non-daily cattle		Swine		Hen, Broiler	
12. Pit storage		3.90 %	D ¹	3.00 %	D ¹	8.7 %	D ¹	—	
13. Sun drying		0.20 %	J ³	0.20 %	J ³	0.20 %	J ³	0.20 %	J ³
14. Other	14a. Thermal drying	0 %							
	14b. Composting (feces)	0.044 %	D ¹	0.034 %	D ¹	0.080 %	J ⁹	0.080 %	J ⁹
	14c. Piling	3.80 %	J ⁵	0.13 %	J ⁵	0.16 %	J ⁵	0.14 %	J ⁵
	14d. Incineration	0.4 %							
	14e. Composting (urine)	0.044 %	D ¹	0.034 %	D ¹	0.097 %	D ¹	—	
	14e. Composting (feces and urine mixed)					0.080 %	J ⁹		
	14f. Purification	0.0087 %	D ¹	0.0067 %	D ¹	0.019 %	D ¹	—	
	14g. Methane fermentation (feces)	3.80 %	P	0.13 %	P	0.16 %	P	0.14 %	P
	14g. Methane fermentation (feces and urine mixed)	3.90 %	S	3.0 %	S	8.7 %	S	—	
	14k. Other (feces)	3.80 %	M	0.4 %	M	0.4 %	M	0.4 %	M
14k. Other (feces and urine mixed)	3.90 %	M	3.0 %	M	8.7 %	M	—		

Table 6-14 N₂O Emission factors for each method of treating manure from cattle, Swine Hen & Broiler (g-N₂O-N/g-N)

treating method		Daily Cattle		Non-daily cattle		Swine		Hen, Broiler	
12. Pit storage				0.10 %				D ¹	
13. Sun drying				2.0 %				D ¹	
14. Other	14a. Thermal drying			2.0 %				D ¹	
	14b. Composting (feces)	0.25 %		J ⁷	0.16 %		J ⁹		
	14c. Piling	2.40 %	J ⁵	1.60 %	J ⁵	2.50 %	J ⁵	2.0 %	D ¹
	14d. Incineration			0.1 %				O ⁴	
	14e. Composting (urine)			2.0%				D ¹	
	14e. Composting (feces and urine mixed)	2.0%	D ¹	0.25%	J ⁷	0.16%	J ⁹	—	
	14f. Purification			5.0 %				J ⁸	
	14g. Methane fermentation (feces)	2.40 %	P	1.60 %	P	2.50 %	P	2.0 %	P
	14g. Methane fermentation (feces and urine mixed)			0.1 %				S	
	14k. Other (feces)	2.4%	M	2.0%	M	2.5%	M	2.0%	M
14k. Other (feces and urine mixed)	5.0%	M	5.0%	M	5.0%	M	—		

D: Default value of Revised 1996 IPCC Guidelines

J: Established by data of Japan

O: Established by data of other countries

Z: Emission can not occur because of mechanism

P: Application of the value of "Piling"

S: Application of the value of "Pit storage"

M: Application of the maximum values of the treating methods for "feces" or "feces and urine mixed"

* Manure excreted by hen and broiler was categorized as feces since it contains a very small amount of urine.

Sources for Table 6-13 and Table 6-14

1: GPG (2000) (Reference 4)

2: IPCC, *Revised 1996 IPCC Guidelines* (Reference 3)

3: Makoto Ishibashi et al., "Development of technology of reducing GHG on the livestock industry(second report)" (2003) (Reference 34)

4: Japan Livestock Technology Association, *GHGs emissions control in livestock Summary*, (2002) (Reference 22)

5: Takashi Osada et al., *Greenhouse gas generation from livestock waste composting* (2005) (Reference 38)

6: IPCC(1995): IPCC 1995 Report (Reference 2)

7: Takashi Osada et al., *Determination of nitrous oxide, methane, and ammonia emissions from a swine waste composting process* (2000) (Reference 36)

8: Takashi Osada, *Nitrous Oxide Emission from Purification of Liquid Portion of Swine Wastewater* (2003) (Reference 37)

9: Project Report of Survey on Prevention of Global Warming in the Agriculture, Forest and Fisheries Sector within the Environment and Biomass Comprehensive Strategy Promotion Project in FY 2008 (Nationwide Survey) (Reference 47)

●Activity Data

The values used for the activity data are estimates of the amount of organic matter and the amount of nitrogen excreted annually by various types of livestock, respectively.

Total annual amount of organic matter by domestic livestock was calculated by multiplying the population of each type of animal by the amount of manure per head by the proportion of organic matter in feces or urine. Total nitrogen amount was calculated by multiplying the population of each type of animal by the nitrogen content amount of feces or urine excreted per head. The amount of organic matter and nitrogen amount was allocated to each category of manure management by multiplying the total amount by the percentage of manure treated separately and the percentage per

treatment method. For livestock population, same references indicated in ‘4.A. Enteric Fermentation’ are used.

In order to avoid duplication with the cattle under grazing, the cattle population was calculated by subtracting activity data for grazing cattle determined by the formula, “Grazing population × Number of grazing days (190 days) / Number of days in year (365 or 366 days)”, from the total population of dairy and non-dairy cattle.

Estimating activity data for CH₄ (amount of organic matter excreted)

Amount of organic matter excreted [Gg] = Livestock population [1000 head]
 × Amount of feces and urine excreted [kg/head/day] × days per year [day] × organic matter content in feces and urine [%] × proportions of feces and urine separated [%] × share of each treating method [%] / 1000

Estimating activity data for N₂O (amount of nitrogen excreted by each type of livestock)

Amount of nitrogen excreted [Gg-N] = Livestock population [1000 head]
 × Nitrogen content amount of feces and urine excreted [kg-N/head/day] × Days per year [day]
 × Proportion of feces and urine separated [%] × Share of each treating method [%] / 1000

Table 6-15 Livestock population for Hen and Broiler

Type of livestock	Unit	1990	1995	2000	2005	2009	2010	2011
Hen	1000 head	188,786	190,634	186,202	180,697	179,849	178,703	178,703
Broiler	1000 head	142,740	118,123	106,311	103,687	107,141	107,141	107,141

* Data for 2011 are substituted by data for 2010

Table 6-16 Amount of feces and urine excreted and Nitrogen content amount by type of livestock

Type of livestock		Amount of feces or urine excreted [kg/head/day]		Nitrogen content amount in feces or urine excreted [g-N/head/day]	
		feces	urine	Feces	urine
Dairy Cattle	Lactating	45.5	13.4	152.8	152.7
	Non-lactating and Inexperienced Birthing	29.7	6.1	38.5	57.8
	Heifer: Under Two Years	17.9	6.7	85.3	73.3
Non-Dairy Cattle	Under Two years	17.8	6.5	67.8	62.0
	Over Two Years	20.0	6.7	62.7	83.3
	Dairy breed	18.0	7.2	64.7	76.4
Swine	Growing-Finishing	2.1	3.8	8.3	25.9
	Breeding	3.3	7.0	11.0	40.0
Hen	poult	0.059	-	1.54	-
	adult	0.136	-	3.28	-
Broiler		0.130	-	2.62	-

Source: M, Tsuiki et al., A Computer Program for Estimating the Amount of Livestock Wastes. (Reference 44)

Table 6-17 Organic matter content in feces and urine, by type of livestock (wet base)

Type of livestock	Organic matter content	
	Feces	Urine
Dairy Cattle	16%	0.5%
Non-Dairy Cattle	18%	0.5%
Swine	20%	0.5%
Hen	15%	—
Broiler	15%	—

Source: Japan Livestock Technology Association, *GHGs emissions control in livestock Summary*. (2002) (Reference 22)

Table 6-18 Proportion of separated and mixed treatment of manure, by type of livestock

Type of livestock	Separated		Mixed	
	~2008	2009~	~2008	2009~
Dairy Cattle	60%	45.5%	40%	54.5%
Non-Dairy Cattle	7%	4.8%	93%	95.2%
Swine	70%	73.9%	30%	26.1%
Hen	100%	100%	—	—
Broiler	100%	100%	—	—

Source: Until 2008: Japan Livestock Technology Association, *GHGs emissions control in livestock Summary*. (2002) (Reference 22),

From 2009 onward:MAFF, *Survey of current status for livestock manure management system* (2009) (Reference 57)

Table 6-19 Percentage of manure management by type of animal

State of Manure (Separated or Mixed)		Treating method	Dairy Cattle		Non-Dairy Cattle		Swine		Hen		Broiler	
			1989~ 2008	2009~	1989~ 2008	2009~	1989~ 2008	2009~	1989~ 2008	2009~	1989~ 2008	2009~
Separated	Feces	Sun drying	2.8%	2.0%	1.5%	0.9%	7.0%	0.7%	30.0%	8.2%	15.0%	2.5%
		Thermal drying	0.0%	0.0%	0.0%	0.0%	0.7%	0.1%	3.0%	2.2%	0.0%	1.1%
		Composting	9.0%	6.6%	11.0%	8.1%	62.0%	48.2%	42.0%	49.6%	5.1%	19.3%
		Piling	88.0%	90.1%	87.0%	89.8%	29.6%	49.3%	23.0%	36.8%	66.9%	36.7%
		Incineration	0.2%	0.0%	0.5%	—	0.7%	0.6%	2.0%	1.6%	13.0%	30.5%
		Methane Fermentation	—	—	—	—	—	0.1%	—	—	—	0.1%
		public sewage	—	0.0%	—	—	—	—	—	—	—	—
	Pasturage	—	0.0%	—	—	—	—	—	0.0%	—	0.1%	
	Other	—	1.3%	—	1.2%	—	1.0%	—	1.6%	—	9.9%	
	Urine	Sun drying	—	0.0%	—	0.0%	—	0.0%	—	—	—	—
		Composting (urine)	1.5%	1.7%	9.0%	1.2%	10.0%	5.4%	—	—	—	—
		Purification	2.5%	5.1%	2.0%	4.4%	45.0%	76.3%	—	—	—	—
		Pit storage	96.0%	89.6%	89.0%	91.5%	45.0%	15.3%	—	—	—	—
		Methane Fermentation	—	1.9%	—	0.0%	—	0.5%	—	—	—	—
Public sewage		—	0.8%	—	0.6%	—	0.4%	—	—	—	—	
Other		—	0.9%	—	2.4%	—	2.1%	—	—	—	—	
Mixed	Sun drying	4.7%	1.1%	3.4%	0.7%	6.0%	0.2%	—	—	—	—	
	Thermal drying	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	—	—	—	—	
	Composting (urine)	20.0%	22.9%	22.0%	10.8%	29.0%	21.3%	—	—	—	—	
	Piling	14.0%	50.9%	74.0%	85.6%	20.0%	51.3%	—	—	—	—	
	Purification	0.3%	0.2%	0.0%	0.0%	22.0%	18.5%	—	—	—	—	
	Pit storage	61.0%	15.4%	0.6%	0.1%	23.0%	4.0%	—	—	—	—	
	Incineration	—	0.1%	—	0.0%	—	0.0%	—	—	—	—	
	Methane Fermentation	—	1.7%	—	0.0%	—	2.0%	—	—	—	—	
	Public sewage	—	0.1%	—	0.0%	—	0.7%	—	—	—	—	
	Pasturage	—	6.5%	—	1.1%	—	0.0%	—	—	—	—	
Other	—	1.2%	—	1.6%	—	1.9%	—	—	—	—		

Source: ~2008: Japan Livestock Technology Association, *GHGs emissions control in livestock Part4*. (1999) (Reference 23)
2009~: MAFF, *Survey of current status for livestock manure management system* (2009) (Reference 57)

●Completeness

Poultry other than hens and broiler are not covered by official statistics, and they are assumed to be negligible. Therefore, only hens and broiler are considered as estimation target from poultry.

●Climate Regions

In the Tier 1 method, the *GPG (2000)* requires that emissions be calculated using herd size by climate regions.

In accordance with the climate categories given in the *Revised 1996 IPCC Guidelines*, Japan should be divided into temperate and cool zones. The average temperature over all prefectures in Japan is around 15 °C. This figure is almost the same as the threshold given in the *Revised 1996 IPCC*

Guidelines. Therefore, emissions have been calculated on the assumption that all of Japan falls into the temperate zone, without a need to categorize regions into temperate or cool zone.

2) Cattle under grazing

Organic matter contained in manure excreted by livestock during grazing (i.e. dung and urine deposited onto grazing and watering grounds by the grazing livestock) is converted to CH₄ through the CH₄ fermentation process, and emitted into the atmosphere. The nitrogen-containing manure also generates ammonium ions, which in turn generates N₂O in the process of oxidation to nitrate from ammonium ions under aerobic conditions.

Emissions in this category are reported for cattle grazing owing to the unavailability of statistics and other information regarding the grazing of other animals. CH₄ emissions are reported in this category and N₂O emissions from grazing cattle are reported in 4D2.

● Estimation Method

For CH₄ and N₂O emitted from pasture, range, and paddock manure, the amount of emissions was calculated for cattle by multiplying the Japan-specific emission factors by the total grazing population in accordance with the Decision Tree in the *GPG (2000)* (page 4.55, Fig. 4.7).

● Emission Factors

Data for the amounts (g) of CH₄ and N₂O emitted from manure excreted per head of cattle per day were used as the emission factors. The data were established by multiplying the model output value of carbon content in manure excreted by grazing cattle during the grazing period by the actual measurement values of CH₄ and N₂O generated per amount of carbon contained in the manure of the grazing cattle.

The amount of carbon contained in the manure of the grazing cattle was calculated by a growth model of grazing cattle based on grass production, quality of grass, climatic conditions, and age in days of grazing cattle.

Table 6-20 Emission factors for grazing cattle

GHGs	Emission Factors	Unit
CH ₄	3.67	[g-CH ₄ /head/day]
N ₂ O	0.32	[g-N ₂ O-N/head/day]

Source: Japan Livestock Technology Association, *GHGs emissions control in livestock Part6*. (2001) (Reference 24)

● Activity Data

Activity data was determined by multiplying the grazing population by the duration of the grazing period. The grazing population was derived from the total grazing population in both public and private pastures reported in the *Livestock Statistics*. For the grazing population in prior years, the percentage of the average grazing population (= Grazing population reported in the *Livestock Statistics* / Total population raised) as in FY 2003 and FY 2004 was determined first, and then the grazing population for each fiscal year was calculated on the assumption that the percentage was the same in all fiscal years.

The duration of 190 days was established for the grazing period, using the values for seasonal grazing (average grazing period: 172.8 days; the number of pastures 623) and year-round grazing (assumed

grazing period: 365 days; the number of pastures 61) indicated in the *Report on National Factual Survey of Cattle Pastures (2000)*, and averaging the grazing days weighted by the number of pastures.

Table 6-21 Trends in the population of grazing cattle

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Amount of grazing daily cattle	head	302,219	281,603	252,088	245,100	290,600	287,291	287,291
Amount of grazing non-daily cattle	head	99,734	103,162	99,759	116,300	113,400	108,319	108,319

* Data for 2011 are substituted by data for 2010

3) Reporting in Common Reporting Format (CRF)

In the CRF, with regard to CH₄ emissions from this category, it is required to report emissions by each livestock. However, for N₂O emissions from this category, it is required to report emissions by AWMS (11. Anaerobic Lagoons, 12. Liquid Systems, 13. Solid Storage and Dry Lot, 14. Other).

For cattle, swine, and poultry, Japan's country-specific manure management categories and the implementation rates of the management categories have been established for each type of animal. For details, see Table 6-22 below.

The current CRF divides the reporting categories into Anaerobic Lagoons, Liquid Systems, Solid Storage and Dry Lots, and Other. In Japan, however, composting is widely practiced, particularly with respect to domestic livestock feces. Consequently the composting-related subcategories of "Piling" and "Composting" have been established under the Other category. Additional subcategories of "Thermal drying" and "Incineration", which are practiced for the purposes of amount reduction and easier handling of dung, have been also included in the Other category. Urine undergoes purification treatment as sewage with high concentrations of pollutants. Accordingly, a subcategory of "Purification" has been added to the CRF category of Other.

Composting is widely practiced in Japan because, among other things: (1) it is essential for Japanese livestock farmers to facilitate transportation and handling, because the lack of space required for the on-site reduction of manure makes it necessary to direct the manure for uses outside their farms; and (2) compost is in considerably higher demand as a fertilizer for various crops than slurry or liquid manure in Japan where fertilizers tend to be lost by heavy rain and the expectations of the protection of water quality, prevention of odor, and sanitary management are high.

"11. Anaerobic Lagoons" have been reported as "NO". Because there are quite small number of livestock farmers who has enough area of field to spread manure, and it is assumed that there are no livestock farmers who use anaerobic lagoons. There are cases when manure is spread to fields in Japan, but even in these cases, stirring is conducted before the spreading. Therefore, there are no anaerobic manure management systems.

Table 6-22 Correspondence between the Japanese and CRF manure management categories

Sub-categories in Japan		CRF	Description of Treatment	
Manure treatment	Manure management category			
Separate treatment	Feces	Sun drying	13. Solid Storage and Dry Lot	Dried under sunlight to facilitate handling (for storage and odor prevention).
		Thermal drying	14. Other (a. Thermal drying)	Dried by heat to facilitate handling.
		Composting	14. Other (b. Composting)	Fermented for several days to several weeks with forced aeration and agitation in lidded or closed tanks.
		Piling	14. Other (c. Piling)	Piling system is a method of composting. Piled about 1.5-2m height on compost bed or in shed to ferment for several months with occasional turning.
		Incineration	14. Other (d. Incineration)	For amount reduction or disposal, and use as an energy source (e.g. chicken manure boiler).
		Methane fermentation	14. Other (g. Methane Fermentation (feces))	Slurry livestock manure is fermented under anaerobic conditions. Generated methane gas is used as an energy source.
		Public sewage	-	Released into public sewage without purification or aeration management. Emissions are included in the Waste sector.
	Urine	Pasture, Range and Paddock	14. Other (i. Pasture, Range and Paddock)	Livestock are fed on a land with vegetation to eat. N ₂ O Emissions are reported in the 'Pasture, Range and Paddock (4.D.2.)'
		Other	14. Other (k. other (feces))	Treated with the method not mentioned above.
		Liquid Composting	14. Other (e. Composting (liquid))	Treated in an aeration storage tank.
		Purification	14. Other (f. Purification)	Separate pollutants using aerobic microbes, such as activated sludge.
		Pit storage	12. Liquid systems	Stored in a storage tank.
		Methane fermentation	14. Other (g. Methane Fermentation (mixed))	Same as above (Methane fermentation)
		Public sewage	-	Same as above (Public sewage)
Mixed treatment	Other	14. Other (k. other (mixed))	Treated with the method not mentioned above.	
	Sun drying	13. Solid Storage and Dry Lot	Dried under sunlight to facilitate handling.	
	Thermal drying	14. Other (a. Thermal drying)	Same as above, Thermal drying.	
	Liquid Composting	14. Other (e. Composting (liquid))	Treated in an aeration storage tank.	
	Piling	14. Other (c. Piling)	Same as above, Piling.	
	Purification	14. Other (f. Purification)	Same as above, Purification.	
	Pit storage	12. Liquid systems	Stored in a storage tank (e.g. slurry storage).	
	Methane fermentation	14. Other (g. methane fermentation (mixed))	Same as above (Methane fermentation).	
	Public sewage	-	Same as above (Public sewage)	
	Pasture, Range and Paddock	14. Other (i. Pasture, Range and Paddock)	Same as above (Pasture, Range and Paddock)	
Other	14. Other (k. other (mixed))	Treated with method not mentioned above		

4) Nitrogen in Livestock Manure Applied to Agricultural Soil

At present, the amount of manure-derived organic fertilizer application in 4.D.3 *Indirect Emissions* is calculated by subtracting the amount of volatilization into the atmosphere, the amount treated by "Incineration" and "Purification", and the amount disposed in landfill as waste from the total nitrogen content of livestock manure. Buffalo, sheep, goats, and horses are excluded from the calculation in 4.D.3 because they produce very small amounts of manure and details of their management in Japan are unknown.

● *Estimation Method*

The percentage of application of manure-derived organic fertilizers was calculated by subtracting the nitrogen contents in the livestock manure disposed of in the "direct final disposal", the nitrogen volatilized as N₂O, the nitrogen volatilized as ammonia and nitrogen oxides, and the nitrogen

eliminated by the “incineration” and “purification”, from the total nitrogen contained in livestock manure excreted in a shed and barn.

$$N_D = N_{all} - N_{N_2O} - N_{NH_3+NO_x} - N_{inc+waa} - N_{waste}$$

N_D :	Amount of nitrogen in manure-derived fertilizer applied to agricultural soil (kg-N)
N_{all} :	Total amount of nitrogen excreted by livestock (deposited in shed and barn) (kg-N)
N_{N_2O} :	Nitrogen in livestock manure volatilized as N_2O (deposited in shed and barn) (kg-N)
$N_{NH_3+NO_x}$:	Nitrogen in manure volatilized as NH_3 and NO_x (deposited in shed and barn) (kg- NH_3 -N + NO_x -N)
$N_{inc+waa}$:	Nitrogen eliminated by “incineration” and “purification”(deposited in shed and barn) (kg-N)
N_{waste} :	Amount of nitrogen in manure that is disposed of in the “final direct disposal” (kg-N)

➤ **Amount of N_2O volatilized into the atmosphere**

The amount of N_2O volatilized into the atmosphere was determined from the calculation results of N_2O emissions from livestock manure.

➤ **Amount volatilized as ammonia and nitrogen oxides**

The amount of nitrogen that was volatilized as ammonia and nitrogen oxides from livestock manure was calculated by multiplying the amount of nitrogen excreted by each type of animal by the percentage of nitrogen that was volatilized as ammonia and nitrogen oxides from manure of each type of animal. Because the percentage of nitrogen that is volatilized as nitrogen oxides is unknown, the percentages of the volatilization of ammonia and nitrogen oxides from manure were determined together with the percentage volatilized as ammonia based on the data in the *Estimated Volatilization of Ammonia from Livestock Manure in the Control of Greenhouse Gas Emissions in Livestock: Summary* (Japan Livestock Technology Association).

Table 6-23 Estimated percentage of volatilized ammonia from livestock manure

Type of Animal	Value
Dairy and non-dairy cattle	10%
Swine	20%
Hen and broiler	30%

Source: Japan Livestock Technology Association, *GHGs emissions control in livestock Summary*. (2002) (Reference 22)

➤ **Nitrogen eliminated by incineration or purification**

The amount was determined from the values of nitrogen disposed of through incineration and purification processes in manure management.

➤ **Nitrogen in manure disposed of in direct final disposal**

Livestock manure disposed of in landfill as waste is either treated before disposal (“treated disposal”) or sent directly to landfill untreated (“direct final disposal”).

Because the manure that was disposed of in “direct final disposal” was detained as a mixture of dung and urine prior to the disposal in landfill, a portion of manure held under the Storage subcategory in the Mixed Treatment category was deemed to have been disposed of in “direct final disposal” (note: manure of hen and broiler was deemed to have been treated under the “Feces - Piling” subcategory). The amount of manure that was disposed of in “treated disposal” is negligible and its treatment method is unknown; therefore, manure that was treated before final disposal was included in the calculation of the manure disposed in the “direct final disposal”.

For the amount of nitrogen in manure disposed of in “direct final disposal,” the total amounts of manure disposed in the “direct final disposal” and “treated disposal” shown in the *Report on the Survey for Research on the Wide-range Movement of Wastes and the State of Cyclical Use of Wastes* were apportioned to the amount of dung and urine of cattle and swine that was treated under the Storage subcategory of the Mixed Treatment category and the amount of manure of hen and broilers that was treated under the “Feces - Piling” of feces subcategory. The amounts that had been apportioned to the cattle and swine were further apportioned to dung and urine. Finally, the amounts of nitrogen content were calculated by multiplying the apportioned amounts by the nitrogen content calculated by dividing nitrogen amount in manure treated in storage system by manure amount treated in storage system in each of dung and urine of each type of animal.

Nitrogen content in livestock manure disposed in the direct final disposal

= Total amount of direct final disposal and treated final disposal × Average nitrogen contents in manure treated by storage system

= Total amount of direct final disposal and treated final disposal × Nitrogen amount in manure treated by storage system / Manure amount treated by storage system

Table 6-24 Nitrogen in livestock manure applied to agricultural soil

Item	Unit	1990	1995	2000	2005	2009	2010	2011
the amount of N in animal manure (N _{all})	tN	789,405	748,584	708,663	683,651	698,037	686,867	688,737
the amount of N ₂ O-N released from animal (except Incineration method and Wastewater manage method) (N _{N₂O})	tN	8,934	8,485	7,981	7,690	8,151	8,000	8,022
the amount of NH ₃ -N and NO _x -N released from animal manure (N _{NH₃+Nox})	tN	144,935	137,392	130,075	125,673	118,173	116,772	117,092
the amount of N vanished by Incineration method and Wastewater manage method (N _{inc+waa})	tN	69,056	60,313	57,938	56,691	97,666	97,151	97,417
the amount of N vanished by burying in the ground. (N _{waste})	tN	489	464	429	417	593	769	769
the amount of N used as fertilizer (N _D)	tN	565,991	541,931	512,239	493,180	473,453	464,175	465,438

c) Uncertainties and Time-series Consistency

● Uncertainties

An uncertainty assessment was conducted for individual livestock categories. For cattle, uncertainty assessments were conducted separately for “shed” and “pastured” cattle and both uncertainties combined.

For the uncertainties of the emission factors for livestock, excluding pastured cattle, the values given in the *GPG (2000)* and the values calculated by expert judgment in accordance with the decision tree for uncertainty assessment, were applied. For the uncertainties of emission factors for pastured cattle, the values calculated by expert judgment were applied in accordance with the decision tree for uncertainty assessment.

For the uncertainties of the activity data, 0.85% (the standard error for swine given in the *Livestock Statistics*) was applied to swine, and 7.29% (the standard error for hens given in the *Livestock Statistics*) was applied to hens, and broilers. For cattle (total population), 5% is adopted, same as “6.2.1. Enteric Fermentation, Cattle”. Activity data for pastured cattle is indicated in the *Livestock Statistics*, but standard error is not indicated and it is difficult to judge applying above precision for

cattle (total). Therefore, 50% was applied for pastured cattle in accordance with the decision tree of uncertainty.

As a result, the uncertainties of the emissions for CH₄ and N₂O were determined to be 78% and 91% for dairy cattle, 73% and 125% for non-dairy cattle, 106% and 92% for Swine, 54% and 80% for Poultry, respectively. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

Emission factors were used consistently from FY 1989 onward by the method. Activity data were calculated consistently from FY 1989 onward from the data in *Livestock Statistics*.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. For some country specific emission factors, there were significant differences between the default emission factors. In the case, the factors of differences were analyzed. QA/QC activities are summarized in Annex 6.1.

e) Source-specific Recalculations

Since new ‘proportion of separated and mixed treatment of manure, by type of livestock’ and ‘percentage of manure management by type of animal’ were reported, emissions for this category FY2008 and FY2009 were revised. In the Agriculture sector, a 3-year average has been used. Thus, the emissions for FY2009 were revised in accordance with the revision and/or update of the activity data for FY2010.

f) Source-specific Planned Improvements

As research on actual emissions has been continuously conducted by the organizations and agencies concerned, a review of emission factors and parameters will be implemented when the new data are obtained.

In addition, since the estimation of the amount of nitrogen fertilized in agricultural soil from livestock manure has a possibility of overestimate, the nitrogen flow in the whole Agriculture sector has been continuously investigated in the Committee for Greenhouse Gas Emission Estimation Methods.

6.3.2. Buffalo, Sheep, Goats & Horses (4.B.2., 4.B.3., 4.B.4., 4.B.6.)

a) Source/Sink Category Description

This section provides the estimation methods for CH₄ and N₂O emissions for manure management from Buffalo, Sheep, Goats and Horses.

b) Methodological Issues

● **Estimation Method**

CH₄ and N₂O emissions were calculated by using the Tier 1 method in accordance with the Decision Tree of the *GPG (2000)* (Page 4.33, Fig. 4.3 and Fig. 4.4).

CH₄ emissions associated with manure management (kg-CH₄)= Emission factor for animal (kg-CH₄/year/head) × Population of the animal (head)*N₂O emission associated with livestock manure (kg-N₂O)*= Emission factor per manure management category of each type of animal [kg-N₂O-N/kg-N] × Nitrogen content of manure [kg-N/head] × Percentage of manure management category × Population of livestock [head]● **Emission Factors**

For the emission factors for CH₄, the default values for temperate zones in industrialized nations, given in the *Revised 1996 IPCC Guidelines* were used. For buffalo, the default value given for the temperate zone in Asia was used.

For the emission factors for N₂O, the default values of “Other animals” for temperate zones in Asia & Far East, given in the *Revised 1996 IPCC Guidelines* were used.

Table 6-25 Emission factors for sheep, goats and horses

Type of livestock	Emission Factors [kg-CH ₄ /head/year]	reference
Sheep	0.28	<i>Revised 1996 IPCC Guidelines</i> Vol. 2 p. 4.6 Table 4-4
Goats	0.18	
Horses	2.08	
Buffalo	2	<i>Revised 1996 IPCC Guidelines</i> , Vol. 3, p. 4.13, Table 4-6

Table 6-26 Emission factors for buffalo, sheep, goats and horses

Manure Management Category		[kg-N ₂ O-N/ kg-N]
11.	Anaerobic Lagoons	0.1%
12.	Liquid Systems (Pit storage)	0.1%
13.	Solid Storage and Dry Lot (Sun drying)	2.0%
14. Other	h. Daily Spread	0.0%
	i. Pasture Range and Paddock	2.0%
	j. Used Fuel	0.0%
	k. Other system	0.5%

Source: *Revised 1996 IPCC Guidelines*, Vol. 3, page 4.121, Table B-1 (Reference 3)● **Activity Data**

For CH₄, same as ‘4.A. Enteric Fermentation’, Calculation of activity data for sheep and goats used the values listed in the *Statistical Document of Livestock Breeding* offered by the Japan Livestock Industry Association and horses used the values listed in the *Statistical Document of Horse* offered by the MAFF. Data for buffalo in the calculation used the population of buffalo listed in the *Statistics on Livestock in Okinawa Prefecture* (Table 6-11).

For N₂O, in order to determine the activity data for buffalo, sheep, goats, and horses, first, the total nitrogen was calculated by multiplying the population of each type of animal by the nitrogen content of manure per head of animal. Then, the amount of nitrogen per manure management category was calculated by multiplying the total nitrogen by the percentage of each management category. For the nitrogen contents of manure and the percentage of each manure management category, the default values given in the *Revised 1996 IPCC Guidelines* were used. For the population size per type of livestock, the same values used in the calculation of CH₄ emissions were used.

Table 6-27 Amounts of nitrogen in manure excreted by buffalo, sheep, goats, and horses

Type of Animal	[kg-N/head/year]
Buffalo*	40
Sheep	12
Goats*	40
Horses*	40

Source: *Revised 1996 IPCC Guidelines*, Vol. 3, page 4.99, Table 4-20 (Reference 3)

* Value for “Other animals” was used.

Table 6-28 Percentage of each manure management category for buffalo, sheep, goats, and horses

Treatment Category		Percentage of Treatment			
		Buffalo	Sheep	Goats	Horses
11.	Anaerobic Lagoons	0%	0%	0%	0%
12.	Liquid Systems (Pit storage)	0%	0%	0%	0%
13.	Solid Storage and Dry Lot (Sun drying)	14%	0%	0%	0%
14. Other	h. Daily Spread	16%	0%	0%	0%
	i. Pasture, Range and Paddock	29%	83%	95%	95%
	j. Used as Fuel	40%	0%	0%	0%
	k. Other system	0%	17%	5%	5%

Source: *Revised 1996 IPCC Guidelines*

c) *Uncertainties and Time-series Consistency*

● *Uncertainties*

An uncertainty assessment was conducted for individual livestock categories. With respect to the uncertainties for emission factors for CH₄ and N₂O from each livestock, 100%—the concerned or similar sources given in the *GPG (2000)*—were applied in accordance with the decision tree for uncertainty assessment. For the uncertainty of the activity data in each livestock, 100% was applied in accordance with decision tree. As a result, the uncertainties of the emissions were determined to be 141% for each livestock. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

For emission factors, same values were used consistently from FY 1989 to FY 2008. Activity data were calculated consistently from FY 1989 onward from the data in the *Statistical Document of Livestock Breeding*, the *Statistical Document of Horse* and the *Livestock Statistics of Okinawa*.

d) *Source-specific QA/QC and Verification*

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1.

e) *Source-specific Recalculations*

In the Agriculture sector, a 3-year average has been used. Thus, the emissions from FY2006 to FY2009 were revised in accordance with the revision and/or update of the activity data from FY2007 to FY2010.

f) *Source-specific Planned Improvements*

There is a need to discuss whether Japan’s country-specific emission factors will be established on the basis of actual measurements.

6.3.3. Camels and Llamas, Mules and Asses (4.B.5., 4.B.7.)

Japan reported “NO” in this section as these animals were not likely to be raised for agricultural purposes.

6.3.4. Other (4.B.10.)

The only livestock that are bred in Japan are cattle, buffalo, sheep, goats, horses, swine and poultry. Therefore, this category has been reported as “NO”.

6.4. Rice Cultivation (4.C.)

CH₄ is generated under anaerobic conditions by the action of microbes. Therefore, paddy fields provide favorable conditions for CH₄ generation.

Intermittently and continuously flooded paddy fields are targeted in this category. In Japan, Rice cultivation is practiced mainly on intermittently flooded paddy field.

CH₄ emissions from Rice Cultivation in FY 2010 are 5,452 Gg-CO₂ eq., comprising 0.4% of total emissions (excluding LULUCF). The value represents a reduction by 21.7% from FY 1990.

Table 6-29 CH₄ emissions from rice cultivation

Gas	Item	Unit	1990	1995	2000	2005	2008	2009	2010
CH ₄	4.C.1.- Intermittently Flooded	Gg-CH ₄	319.9	325.5	272.1	263.8	257.3	254.8	250.6
	4.C.1.- Continuously Flooded	Gg-CH ₄	11.6	11.8	9.8	9.5	9.3	9.2	9.1
	Total	Gg-CH ₄	331.4	337.3	281.9	273.3	266.6	264.0	259.6
		Gg-CO ₂ eq	6,960	7,083	5,920	5,739	5,599	5,545	5,452

6.4.1. Intermittently Flooded (Single Aeration) (4.C.1.-)

a) Source/Sink Category Description

This section provides the estimation methods for CH₄ emissions from intermittently flooded rice cultivation.

● Water management regime in Japanese paddy fields

The general practice of intermittent flooding (single aeration) by paddy farmers in Japan is different in nature from the intermittently flooded paddy field (multi aeration) concept in the *Revised 1996 IPCC Guidelines*. The diagram below presents the outline.

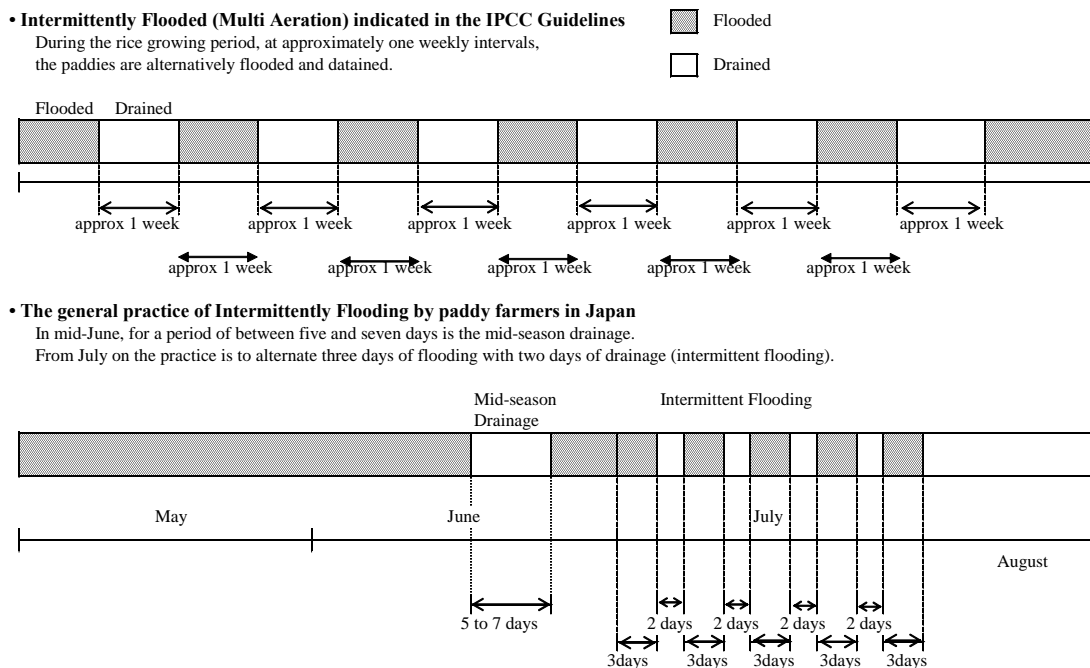


Figure 6-3 Comparison of water management regime in Japan and intermittent flooding (Multi aeration) indicated in the *IPCC Guidelines*

b) Methodological Issues

● **Estimation Method**

CH₄ emissions from intermittently flooded paddy fields (single aeration) were calculated by taking the overall usage of organic matter management into account, since the actual measurements of emission factors per soil type for each type of organic matter management (type of applied organic matter) existed.

The amount of CH₄ generated per type of soil for each method of organic matter management was calculated by multiplying the area of intermittently flooded paddy fields by the “amount of CH₄ generated per type of soil per unit area for each management method”, “proportion of the area of each type of soil”, and “proportion of each organic management method”.

$$\begin{aligned}
 & \underline{CH_4 \text{ emission from intermittently flooded paddy fields (single aeration) (kg-CH}_4)} \\
 & = \sum (\text{Emission factor for organic matter management method } n \text{ for soil type } m \text{ [kg-CH}_4\text{/m}^2] \times \\
 & \quad \text{Area of paddy fields [m}^2] \times \text{Percentage of intermittently flooded paddy field} \times \text{Proportion of} \\
 & \quad \text{soil type } m \times \text{Proportion of organic matter management method } n)
 \end{aligned}$$

● **Emission Factors**

The following table summarizes the emission factors established for each category of this source.

The established emission factors are based on actual measurements of five soil types, with and without straw amendment. Actual data on soil types subject to composting is not available, but the CH₄ emission of composted soil is 1.2 to 1.3 times of un-composted soil. Therefore, the emission factor for composted soil, by soil type, was established as 1.25 times larger than the value for un-composted soil.

Table 6-30 CH₄ emission factor for intermittently flooded paddy fields (single aeration)

Type of soil	Straw amendment [g-CH ₄ /m ² /year]	Various compost amendment [g-CH ₄ /m ² /year]	No-amendment [g-CH ₄ /m ² /year]
Andosol	8.50	7.59	6.07
Yellow soil	21.4	14.6	11.7
Lowland soil	19.1	15.3	12.2
Gley soil	17.8	13.8	11.0
Peat soil	26.8	20.5	16.4

Source: Haruo Tsuruta (2000) (Reference 31)

●Activity Data

It is assumed that intermittently flooded paddy fields (single aeration) comprise some 98% of planted paddy area and continuously flooded paddies comprise the remaining 2%¹.

The method of establishing activity data for emissions of CH₄ from intermittently flooded paddy fields (single aeration) was to multiply the planted paddy area given in the Ministry of Agriculture, Forestry and Fisheries in *Statistics of Cultivated and Planted area*, by the proportion of area by each soil types (Takata et al. (2009)), and then by the proportion subject to organic matter management. Since the survey for proportion of organic matter management has been conducted every year since FY2008, their data has been reflected to the estimation.

Table 6-31 Proportion of Japan's surface area represented by specific soil types

Soil type		~1991	1992	1997	2001	2002~
Andosol	Andosol, moist andosol, andosol gley soil	13.06%	13.06%	13.14%	13.20%	13.20%
Yellow soil	Brown forest soil, gray ground soil, gley ground soil, yellow soil, dark red soil, red soil, lithosol	11.31%	11.31%	11.03%	10.80%	10.80%
Lowland soil	Brown lowland soil, grey lowland soil, regosol	40.82%	40.82%	40.62%	40.46%	40.46%
Gley soil	Gley soil, strong gley soil	28.94%	28.94%	29.20%	29.40%	29.40%
Peat soil	Black peat, peat soil	5.85%	5.85%	6.02%	6.15%	6.15%

*1992 data and 2001 data were original data indicated in Takata et al.(2009). 1993-2000 data were calculated by using interpolation between 1992 and 2001. 1992 data was used for data before FY1991 and 2001 data was used for data after FY2002.

Source: Calculated from Takata et al.(2009) (Reference 48)

Table 6-32 Proportion of organic matter management in Japan

Organic matter	1990~2007	2008	2009	2010
Straw amendment	60%	65%	61%	57%
Various compost amendment	20%	18%	23%	26%
No-amendment	20%	17%	16%	17%

Source : 1990~2007: MAFF, "Basis Survey of Soil Environment" (Reference 49)

After 2008: MAFF, "Survey of Greenhouse Gas Emissions from Soils and Soil Carbon Sequestration" (Reference 50)

Table 6-33 Area of paddy fields

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Area of paddy field	kha	2,055	2,106	1,763	1,702	1,621	1,625	1,574

Source: *Statistics of Cultivated and Planted Area* (MAFF) (Reference 13)

c) Uncertainties and Time-series Consistency

●Uncertainties

¹ Revised 1996 IPCC Guidelines, vol.2 Workbook, p4.18, Table 4.9

The uncertainties for CH₄ emissions from intermittently flooded (single aeration) paddy fields are assessed with respect to each organic matter management method (straw amendment, various compost amendment and no-amendment), because the uncertainty assessment methods differ for each management regime.

For the uncertainties of the emission factors the values given in the *GPG (2000)* or the values calculated by expert judgment were applied in accordance with the decision tree for uncertainty assessment. For the uncertainty of the activity data, 0.31% for area of paddy fields given in the *Statistics of Cultivated and Planted Area* was applied.

As a result, the uncertainties of the emissions were determined to be 32% for straw amendment, 32% for various compost amendment and 46% for no-amendment. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

Emissions are estimated by using consistent estimation methods and data sources.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1.

e) Source-specific Recalculations

In the Agriculture sector, a 3-year average has been used. Thus, the emissions for FY2009 were revised in accordance with the revision for proportion of organic matter management for FY 2010 and the revision and/or update of the activity data for FY2010.

f) Source-specific Planned Improvements

The Ministry of Agriculture, Forestry and Fisheries is currently conducting a comprehensive study aimed at agricultural land. A part of results of this study were reflected for estimation in this year. There will be a review to be conducted on the estimation methods and parameter when new results of the study become available.

Work is progressing on developing an estimation method that uses the DNDC model, and the application of Tier 3 will be discussed in the future.

6.4.2. Continuously Flooded (4.C.1.-)

a) Source/Sink Category Description

This section provides the estimation methods for CH₄ emissions from continuously flooded rice cultivation.

b) Methodological Issues

● **Estimation Method**

CH₄ emissions have been calculated by using country-specific emission factors for different soil types and for different organic amendments, in accordance with Decision Tree of the *GPG (2000)* (Page

4.79, Fig. 4.9).

● **Emission Factors**

Research results in Japan (Association for Advancement of Agricultural Science (2000), Reference 28) indicate that emissions of CH₄ from intermittently flooded paddy fields are 42% to 45% less than those from continuously flooded paddy fields. This knowledge formed the basis for the establishment of an emission factor for CH₄ from continuously flooded paddy fields: divide the implied emission factor, which is gotten by divided emissions by crop field area, for intermittently flooded paddy fields by 0.565 (1-0.435). Since proportion of area by soil types and proportion of organic matter management change every year, the implied emission factor for intermittently flooded paddy fields changes every year. Therefore, the emission factor for continuously flooded paddy fields changed annually.

Table 6-34 Emission factor for CH₄ from continuously flooded paddy fields

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Continuously flooded paddy fields	gCH ₄ /m ² /year	28.12	28.12	28.12	28.12	28.38	28.05	28.05
Intermittently flooded paddy fields (single aeration)*	gCH ₄ /m ² /year	15.89	15.89	15.89	15.89	16.04	15.85	15.85

* Implied emission factor

● **Activity Data**

It is assumed that intermittently flooded paddy fields (single aeration) comprise some 98% of planted paddy area and continuously flooded paddies comprise the remaining 2%.

The method of establishing activity data for emissions of CH₄ from continuously flooded paddy fields was to multiply the planted paddy area given in the Ministry of Agriculture, Forestry and Fisheries in *Statistics of Cultivated and Planted area*, by 2%.

c) **Uncertainties and Time-series Consistency**

● **Uncertainties**

The uncertainties for emission factors were calculated from the uncertainties of each parameter decided by expert judgment. For the uncertainty for activity data, 0.31% of standard error for area of paddy field given in the *Statistics of Cultivated and Planted Area* was applied. As a result, the uncertainty of the emissions was determined to be 116%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

Refer to section 6.4.1. *Intermittently Flooded*.

d) **Source-specific QA/QC and Verification**

Refer to section 6.4.1. *Intermittently Flooded*.

e) **Source-specific Recalculations**

In the Agriculture sector, a 3-year average has been used. Thus, the emissions for FY2009 were revised in accordance with the revision and/or update of the activity data for FY2010.

f) **Source-specific Planned Improvements**

Japan's CH₄ emission ratio of "Intermittently Flooded / Continuously Flooded" are measured on only one site; therefore, further data collection is regarded as necessary.

6.4.3. Rainfed & Deep Water (4.C.2., 4.C.3.)

As indicated in the IRRI (International Rice Research Institute) *World Rice Statistics 1993–94*, rainfed and deep water paddy fields do not exist in Japan. Therefore, this category has been reported as “NO”.

6.4.4. Other (4.C.4.)

Just as indicated in the IRRI (International Rice Research Institute) *World Rice Statistics 1993-94*, a possible source of emissions in this category is upland rice field, but since upland rice field are not flooded, like the soil of fields, they are aerobic and do not become anaerobic. The bacteria that generate CH₄ are obligatory anaerobic bacterium, and unless the soil is maintained in an anaerobic state, there will be no generation of CH₄. As generation of CH₄ is not feasible, this category was reported as “NA”.

6.5. Agricultural Soils (4.D.)

This section provides the estimation methods for N₂O direct emissions from soils (by applied synthetic fertilizers, organic fertilizers, nitrogen fixation by N-fixing crops, crop residue and plowing of organic soil), and for N₂O indirect emissions (by atmospheric deposition and nitrogen leaching and run-off).

●Direct Emissions (N₂O)

Application of synthetic fertilizers, organic fertilizers, nitrogen fixation by N-fixing crops or use of crop residues for soil amendment generates ammonium ions in the soil. The soil emits N₂O in the process of oxidizing the ammonium ions into nitrate-nitrogen under aerobic conditions. N₂O is emitted via denitrification of nitrate. N₂O is generated when organic soil containing nitrogen is plowed.

●Indirect Emissions (N₂O)

Nitrogen compounds such as ammonia, that volatilize and are released into the atmosphere from synthetic fertilizers applied to agricultural soils and organic fertilizers derived from livestock manure are deposited on soil as the results of various actions, including turbulent diffusion, molecular diffusion, effect of electrostatic forces, chemical reactions, plant respiration, and being washed out of the air by rain. In this section, the amount of N₂O generated by microbe activity on the deposited nitrogen compounds was calculated.

N₂O is generated by the action of microbes on nitrogen that leaches or runs off as nitrate from synthetic fertilizers and manure-derived fertilizers applied to agricultural soil.

N₂O emissions from agricultural soils in FY 2010 are 5,619 Gg-CO₂ eq., comprising 0.4% of total emissions (excluding LULUCF). The value represents a reduction by 28.6% from FY 1990.

Table 6-35 N₂O emissions from agricultural soils

Gas	Item	Unit	1990	1995	2000	2005	2008	2009	2010	
N ₂ O	4.D.1. Direct Emission	Synthetic Fertilizers	Gg-N ₂ O	6.2	5.4	4.9	4.8	4.1	3.7	3.7
		Organic Fertilizers	Gg-N ₂ O	4.3	3.9	3.6	3.5	3.4	3.4	3.4
		N-fixing Crops	Gg-N ₂ O	0.3	0.2	0.3	0.3	0.3	0.3	0.3
		Crop Residue	Gg-N ₂ O	2.1	2.1	2.0	1.9	1.9	1.8	1.8
		Plowing of Organic Soil	Gg-N ₂ O	0.4	0.4	0.4	0.4	0.4	0.4	0.4
	4.D.2. Pasture, Range and Paddock Manure	Gg-N ₂ O	0.04	0.04	0.03	0.03	0.04	0.04	0.04	
	4.D.3. Indirect Emission	Atmospheric Deposition	Gg-N ₂ O	5.1	4.8	4.4	4.3	4.1	4.0	3.9
		Nitrogen Leaching and Run-off	Gg-N ₂ O	6.9	6.4	5.8	5.6	5.2	4.9	4.8
	Total		Gg-N ₂ O	25.4	23.2	21.5	20.8	19.3	18.4	18.1
			Gg-CO ₂ eq	7,864	7,179	6,674	6,443	5,996	5,694	5,619

6.5.1. Direct Soil Emissions (4.D.1.)

6.5.1.1. Synthetic Fertilizers (4.D.1.-)

a) Source/Sink Category Description

This section provides the estimation methods for N₂O emissions by the application of synthetic fertilizers.

b) Methodological Issues

● Methodology for Estimating Emissions / Removals of GHGs

N₂O emissions were calculated, using country-specific emission factors in accordance with Decision Tree of the *GPG (2000)* (Page. 4.55 Fig. 4.7).

$$\frac{N_2O \text{ emissions associated with the application of synthetic fertilizer in agricultural soil (crop field)} (kg-N_2O)}{=} \text{Emission factor [kg-N}_2\text{O-N/kg-N]} \times \text{Amount of nitrogen contained in synthetic fertilizer applied in crop field [kg-N]} \times 44/28$$

● Emission Factors

Emission factors were established based on actual data measurement conducted in Japan.

Emission factors for N₂O associated with the application of synthetic fertilizers and organic fertilizers were defined as the same value, because there was no significant difference between emission factors of synthetic fertilizers and organic fertilizers, analyzing data for N₂O emissions from agricultural fields in Japan.

Comparing emission factors among various crops, it was identified that emission factor of tea was significantly higher and emission factor of rice was significantly lower than those of other crops. As there were not significant differences among the other crops, three emission factors were defined (for rice, tea and other crops). Emission factor of Japan is lower than that of default value in the *Revised 1996 IPCC Guidelines*. It is the reason that the volcanic ash soil that is widely distributed in Japan releases little N₂O emissions. The emission factor of rice is adopted as a default value within the 2006 IPCC Guidelines and its validity has been internationally confirmed.

Table 6-36 N₂O emission factor for synthetic fertilizer to agricultural soil

Crop species	Emission Factor (kg-N ₂ O-N/kgN)
Paddy rice	0.31 %
Tea	2.9 %
Other species	0.62 %

Source: Akiyama et al., Direct N₂O emissions and estimate of N₂O emission factors from Japanese agricultural soils. (2006) (Reference 39)

Akiyama et al., Estimations of emission factors for fertilizer-induced direct N₂O emissions from agricultural soils in Japan: Summary of available data (2006) (Reference 40)

●Activity Data

For coordination with the way emission factors have been set, the amount of synthetic fertilizer used by crop type is used as the activity data. The amount of synthetic fertilizer used can be ascertained from statistical information on the total amount used, but because there are no data enabling one to determine the annual amounts applied by crop type, values corresponding to the amounts of nitrogen applied for each crop type are found by taking the area of land planted with each crop type that can be found using statistical information and multiplying by the results of studies on the amounts of synthetic fertilizers applied per unit area for each crop type in Japan. Total synthetic fertilizer demand is apportioned to each crop type in accordance with the corresponding application amount for each crop type.

Activity data for N₂O emissions from the application of synthetic fertilizers to crop field

Amount of nitrogen-based fertilizer applied to agricultural soil of each crop field [t]
 = Demand for synthetic fertilizer [tN] × (Area of each crop field [ha] × Amount of synthetic fertilizer used in each crop field [kgN/10a]) / (Σ Area of each crop field [ha] × Amount of synthetic fertilizer used in each crop field [kgN/10a])

The amounts of fertilizer applied by crop type are known because the amounts of synthetic and organic fertilizers applied for each crop type were determined by a farming study conducted in 2000 (*A report on an Investigation of how to quantify the amount of Greenhouse Gases Emissions reduced in 2000 F.Y.* (Reference 28)). Because experts reason that there is likely little year-on-year change in application amounts to crops except for paddy rice and tea, data on the amounts of synthetic fertilizer applied per unit area according to the 2000 study (Reference 28) were applied uniformly for these crops in all years.

Because of regulations and other factors, fertilizer application amounts for tea change from year to year. Nonaka (2005) (Reference 45) has found the amounts of nitrogen applied to tea fields (the total of synthetic and organic) in 1993, 1998, and 2002. For these application amounts, the ratio of synthetic fertilizer to organic fertilizer applied to tea according to the 2000 study (Reference 28) was used to estimate the amounts of synthetic and organic fertilizer applied, which were then used in calculations. Time-series data were prepared by interpolating from 1993 to 2002, using the 1993 data for previous years, and using the 2002 data for subsequent years (see Table 6-36). For paddy rice, the report uses application amount data for years that can be determined using Statistical Survey on Farm Management and Economy (Ministry of Agriculture, Forestry and Fisheries). The value of paddy rice was substituted for upland rice.

Table 6-37 Demand for synthetic fertilizer

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Demand for Synthetic Fertilizer	tN	611,955	527,517	487,406	471,190	350,135	350,135	350,135

* The data for 2010 and 2011 are substituted by the data for 2009

Source: Yearbook of Fertilizer Statistics (Pocket Edition) (References 17)

Table 6-38 Amount of synthetic fertilizers application per area by each type of crop (other than rice and tea)

Type of crop	Amount of application [kg N/10a]
Vegetables	21.27
Fruit	14.70
Potatoes	12.70
Pulse	3.10
Feed crops	10.00
Sweet potato	6.20
Wheat	10.00
Coarse cereal (including Buckwheat)	4.12
Mulberries	16.20
Industrial crops	22.90
Tobacco	15.40

Source: Association for Advancement of Agricultural Science, *Establishment of GHGs reduction model, Incorporated foundation* (References 28)

Table 6-39 Amount of synthetic fertilizers application per area (rice and tea)

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Amount of synthetic fertilizers application per area (rice)	kg-N/10a	9.65	8.71	7.34	6.62	5.80	5.80	5.80
Amount of synthetic fertilizers application per area (tea)	kg-N/10a	57.23	54.88	48.06	44.76	44.76	44.76	44.76

* The data of rice for 2010 and 2011 are substituted by the data for 2009

Source: Rice: MAFF, "Research of agricultural management"

Tea: Kunihiko Nonaka (2005) (References 45)

Table 6-40 Area of cropping by each type of crop

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Vegetables *	ha	620,100	564,400	524,900	476,300	468,700	465,400	465,400
Rice	ha	2,055,000	2,106,000	1,763,000	1,702,000	1,621,000	1,625,000	1,574,000
Fruit *	ha	346,300	314,900	286,200	265,400	250,700	246,900	246,900
Tea	ha	58,500	53,700	50,400	48,700	47,300	46,800	46,200
Potatoes *	ha	115,800	104,400	94,600	86,900	83,100	82,500	82,500
Pulse *	ha	256,600	155,500	191,800	193,900	197,500	189,000	189,000
Feed crops	ha	1,096,000	1,013,000	1,026,000	1,030,000	1,008,000	1,012,000	1,030,000
Sweet potato	ha	60,600	49,400	43,400	40,800	40,500	39,700	38,900
Wheat	ha	366,400	210,200	236,600	268,300	266,200	265,700	271,700
Coarse cereal (including Buckwheat)	ha	29,600	23,400	38,400	45,900	47,500	49,700	49,700
Mulberries *	ha	59,500	26,300	5,880	2,998	2,011	2,011	2,011
Industrial crops	ha	142,900	124,500	116,300	110,300	106,430	104,680	105,280
Tobacco	ha	30,000	26,400	24,000	19,100	15,770	15,120	15,120
Upland rice	ha	18,900	11,600	7,060	4,470	3,000	2,890	2,370

* Data for 2011 are substituted by data for 2010

Source: Potatoes: MAFF, *Vegetable Production and Shipment Statistics*, Tobacco: *Japan Tobacco Survey*, Mulberries: MAFF *Survey*, Other crops: MAFF, *Statistics of Cultivated and Planted Area* (Note: The values of "Potatoes" is excluded in "Vegetable", and "Tea" and "Tobacco" is excluded in "Industrial crops".)

c) *Uncertainties and Time-series Consistency*

● *Uncertainties*

N₂O emissions by the application of synthetic fertilizers were estimated for each crop species. Thus, the uncertainties of N₂O emissions by the application of synthetic fertilizers were also calculated for each crop species and then finally combined as total uncertainties. The uncertainties for the emission factors were calculated by combining the uncertainties of parameters, estimated by expert judgment or using sample standard deviations. As a result, the uncertainties for emission factors were determined to be 220.0% for paddy rice, 211.7% for tea, 181.7% for other crops. For the uncertainty for activity data, 0.31% for paddy rice and 0.26% for other crops (the value for area of upland fields), which is standard error given in the Statistics of Cultivated and Planted Area, was applied. As a result, the uncertainties of the emissions were determined to be 139%. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

Emissions are estimated by using consistent estimation methods and data sources.

d) *Source-specific QA/QC and Verification*

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1. Comparison with Japan's EF and the default EF in the IPCC Guidelines is described in the section 'Emissions factors' above.

e) *Source-specific Recalculations*

In the Agriculture sector, a 3-year average has been used. Thus, the emissions for FY 2008 and FY2009 were revised in accordance with the revision and/or update of the activity data for FY 2009 and FY2010.

f) *Source-specific Planned Improvements*

The same emission factor has been used for synthetic and organic fertilizers. Thus, it is needed to discuss whether it is possible to obtain separate emission factors for these two types of fertilizer.

6.5.1.2. Organic Fertilizer (Application of Animal Waste) (4.D.1.-)

a) *Source/Sink Category Description*

This section provides the estimation methods for N₂O emissions by application of organic fertilizer (livestock and other compost and barnyard manure).

b) *Methodological Issues*

● *Estimation Method*

Emissions of N₂O have been calculated in accordance with Decision Tree of the *GPG (2000)* (Page 4.55, Fig. 4.7).

$$\begin{aligned}
 & \underline{N_2O \text{ emissions from the application of organic fertilizers to agricultural soils (kg-N}_2\text{O)}} \\
 & = \sum_{\text{Type of crop}} \{ \text{Emission factor by type of crop (kg-N}_2\text{O-N/kg-N)} \\
 & \quad \times \text{amount of nitrogen applied, by type of crop (kg-N)} \} \times 44/28
 \end{aligned}$$

●Emission Factors

The same country specific emission factor used for synthetic fertilizer is used. (Table 6-36)

●Activity Data

Activity data was derived by multiplying the area of cultivation for each type of crop, by the amount of nitrogen applied per unit area for each type of crop (excluding tea). Because of regulations and other factors, fertilizer application amounts for tea change from year to year, same as the synthetic fertilizers. Nonaka (2005) (Reference 45) has found the amounts of nitrogen applied to tea fields (the total of synthetic and organic) in 1993, 1998, and 2002. For these application amounts, the ratio of synthetic fertilizer to organic fertilizer applied to tea according to the 2000 study (Reference 28) was used to estimate the amounts of synthetic and organic fertilizer applied, which were then used in calculations. Time-series data were prepared by interpolating from 1993 to 2002, using the 1993 data for previous years, and using the 2002 data for subsequent years (see Table 6-39). Area of cultivated land by type of crop is same as synthetic fertilizers.

$\begin{aligned} &\text{Amount of nitrogen applied, by type of crop (kg-N)} \\ &= \text{Area of cultivated land by type of crop (ha)} \\ &\quad \times \text{Amount of nitrogen as organic fertilizer applied per unit area, by type of crop (kg-N/10a)} \times 10 \end{aligned}$

Table 6-41 Amount of nitrogen as organic fertilizers application per area by each type of crop (excluding tea)

Type of crop	Amount of application [kg-N/10a]
Vegetables	23.62
Paddy rice *	3.2
Fruit	10.90
Potatoes	7.94
Pulse	6.24
Feed crops	10.00
Sweet potato	8.85
Wheat	5.70
Coarse cereal (including Buckwheat)	1.81
Mulberries	0.00
Industrial crops	3.96
Tobacco	11.41

*the value of paddy rice was substituted for upland rice.

Source: Association for Advancement of Agricultural Science, A report on an Investigation of how to quantify the amount of Greenhouse Gases Emissions reduced in 2000 F.Y.. (Reference 28)

Table 6-42 Amount of nitrogen as organic fertilizers application per area for tea

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Amount of organic fertilizers application per area (tea)	kg-N/10a	20.77	19.92	17.44	16.24	16.24	16.24	16.24

Source: Total amount of synthetic and organic fertilizers : Nonaka (2005) (Referenace 45)

c) Uncertainties and Time-series Consistency

●Uncertainties

An uncertainty assessment was conducted by the same method as in 6.5.1.1. Synthetic Fertilizers. As a result, the uncertainty of the emissions was determined to be 152%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

Emissions are estimated by using consistent estimation methods and data sources.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1.

e) Source-specific Recalculations

In the Agriculture sector, a 3-year average has been used. Thus, the emissions for FY2009 were revised in accordance with the revision and/or update of the activity data for FY2010.

f) Source-specific Planned Improvements

The same emission factor has been used for synthetic and organic fertilizers. Thus, it is needed to discuss whether it is possible to obtain separate emission factors for these two types of fertilizer.

6.5.1.3. N-fixing Crops (4.D.1.-)

a) Source/Sink Category Description

This section provides the estimation methods for N₂O emissions from nitrogen fixed by N-fixing crops.

b) Methodological Issues

● **Estimation Method**

Emissions are calculated by taking the amount of nitrogen fixed by nitrogen-fixing crops, which is estimated using Japan's observation data, and multiplying by country-specific emission factor.

$$E = EF * F_{BN} * 44 / 28$$

- E : N₂O emission associated with N-fixation by N-fixing crops (kg-N₂O)
 EF : Emission factor (kg-N₂O- N/kg-N)
 F_{BN} : Amount of nitrogen fixed by N-fixing crops (kg-N)

● **Emission Factors**

The N₂O emission factor for emissions from application of synthetic fertilizer, which is set using Japan's measurement results, is set on the basis of emissions from both nitrogen from fertilizer application and the amount of nitrogen fixed by nitrogen-fixing crops. Therefore, it is set as the emission factor of N₂O emissions from nitrogen fixed by N fixing crops. Although there are three kinds of emission factors for synthetic fertilizers, such as for "rice", "tea", and "other crops", (see Table 6-36), the EF of "other crops" (0.0062[kgN₂O-N/kg-N]) is applied in view of the target crops.

● **Activity Data**

The amount of nitrogen in the above-ground part biomass of N fixing crops is considered to be reasonably substituted for the amount of annual nitrogen fixation by the N fixing crops cultivated in one year. The nitrogen content data in the harvest in the crops and a harvest residue of our country in Owa (1996) was used, and the nitrogen amounts fixed by N fixing crops are calculated by the

following methods. The target crops are broadly classified into "pulse (dried grain) and vegetables", and "feed crops."

➤ *Pulse (dried grain) and Vegetables*

Included in calculations for nitrogen-fixing crops are the pulses (dried seeds) of soybeans, adzuki beans, kidney beans, and peanuts, and the vegetables of string beans, snow peas, broad beans, and green soybeans.

The amount of nitrogen fixed by nitrogen-fixing crops (F_{BN}) was set by transforming Tier 1b Equation 4.26 of GPG (2000) and multiplying the crop yield for N-fixing crops ($Crop_{BFi}$) by the amount of nitrogen per crop yield and crop residue, which was determined by Japanese research data.

$$F_{BN} = \sum_i [Crop_{BFi} \cdot (Frac_{NCRBFi} + Frac_{NRESBFi})]$$

F_{BN}	: The amount of nitrogen fixed by N-fixing crops (kg-N)
$Crop_{BFi}$: Actual crop yield for N-fixing crops i (t)
$Frac_{NCRBFi}$: Amount of nitrogen per crop yield for N-fixing crops i (kg-N/t)
$Frac_{NRESBFi}$: Amount of nitrogen per crop residue for N-fixing crops i (kg-N/t)

➤ *Feed crops*

In Japan, grass and legume feed crops are sown together. Statistical information enables one to ascertain only the crop yield and planted areas of grass-only feed crops and mixed grass–legume feed crops. Because that makes it impossible to directly find the harvest amount and planted area of legume-only feed crops, for the sake of convenience, it is used 10% for the proportion of legume feed crops in mixed-sown in accordance with the judgments of experts based on a Japanese study² and other sources, and estimated the crop yield of legume feed crops.

Japanese research data include those on the nutrient content in the stubble and roots of grass–legume mixed feed crops, and taking into account that calculations for nitrogen-fixing crops in the 2006 IPCC Guidelines cover the amount of aboveground biomass residue and underground biomass plowed into soil, it was decided that calculation of the nitrogen amount fixed by legume feed crops would directly use the amount of nitrogen in stubble and root residue instead of the amount of nitrogen in harvested aboveground biomass, and estimates were made with the following equation, obtained by transforming GPG (2000) Equation 4.27.

$$F_{BN} = \sum_i [Crop_{BF} \cdot Frac_{NCBGF}]$$

F_{BN}	: Amount of nitrogen fixed by leguminous feed crops (kg-N)
$Crop_{BF}$: Actual crop yield for leguminous feed crop (t)
$Frac_{NCBGF}$: Amount of nitrogen contained in the underground part per crop yield for leguminous feed crop (kg-N/t)

² Research results of Hokkaido prefectural Agricultural Experiment Stations” Current status and issues of feed crop production in meadow in Hokkaido I. Current status of crop yield and nutrient value” <http://www.agri.pref.hokkaido.jp/center/kenkyuseika/gaiyosho/h12gaiyo/20003161.htm>

Table 6-43 Parameters used in estimating for N-fixing crops

Type of crop	Amount of fixed nitrogen per unit crop yield (kg-N/t)	Proportion of dry matter
Soybeans	69.17	1.000
Adzuki beans	40.68	1.000
Kidney beans	50.13	1.000
Peanuts	63.00	1.000
Strings beans	1.98* ²	0.302* ¹
Snow pea	2.65* ²	0.302* ¹
Broad beans	9.57* ¹	0.302* ¹
Green soybeans	9.57	0.302
Leguminous feed crop	2.74	0.200

*1 The value for green soybeans is substituted.

*2 Each crop value are calculated by using nitrogen ratio included in harvest for each crop and green soybeans and by using the amount of fixed nitrogen per unit crop yield for green soybeans .

c) *Uncertainties and Time-series Consistency*

● *Uncertainties*

N₂O emissions for nitrogen fixed by N fixing crops were estimated for each crop species. Thus, the uncertainties of N₂O emissions for nitrogen fixed by N fixing crops were also calculated for each crop species and then finally combined as total uncertainties. The uncertainties for the emission factors were calculated by combining the uncertainties of parameters decided by expert judgment and indicated in GPG (2000). The uncertainties for activity data were determined to be 0.26% of standard error for the area of crop field indicated in the Statistics of Cultivated and Planted Area. As a result, the uncertainties for emission from nitrogen fixed by N fixing crops were determined to be 99%.

● *Time-series Consistency*

Emissions are estimated by using consistent estimation methods and data sources.

d) *Source-specific QA/QC and Verification*

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1.

e) *Source-specific Recalculations*

In the Agriculture sector, a 3-year average has been used. Thus, the emissions for FY2009 were revised in accordance with the revision and/or update of the activity data for FY2010.

f) *Source-specific Planned Improvements*

The percentage of legume feed crops in mixed-sown pastures is needed to be elaborated. To make the shift to the estimation method in accordance with the *2006 IPCC Guidelines* in the future, it is a need to obtain data of underground crop residue plowed into soil. But sufficient their data don't exist now. These improvement are investigation issues in the future.

6.5.1.4. Crop Residue (4.D.1.-)

a) *Source/Sink Category Description*

This section provides the estimation methods for N₂O emissions by crop residue plowed into soil.

b) Methodological Issues

● Estimation Method

N₂O emissions associated with the application of crop residues to agricultural soils were calculated by multiplying the default emissions factors given in the Revised 1996 IPCC Guidelines by the nitrogen input through the use of crop residues plowed into soil.

$$\begin{aligned} & \underline{N_2O \text{ emission associated with the use of crop residues plowed into soil (kg-N}_2\text{O)} \\ & = \text{Default emission factor [kg-N}_2\text{O-N/kg-N]} \times \text{Nitrogen input through the use of crop residues} \\ & \text{plowed into soil [kg-N]} \times 44/28 \end{aligned}$$

● Emission Factors

The default emission factor, 0.0125 [kg-N₂O-N/kg-N], shown in the *Revised 1996 IPCC Guidelines* and the *GPG (2000)* was used.

● Activity Data

[Rice]

For the amount of rice crop residue plowed into soil, the data for rice straw and rice chaff indicated in the survey of MAFF was used. The nitrogen content of this crop was calculated by multiplying the aforementioned data by nitrogen content in crop residue (kgN/t) calculated from Matsumoto (2000).

[Wheat, Barley]

The total amount of nitrogen in residue for wheat and barley was calculated by multiplying the amount of crop production (by MAFF, ‘*Statistics of Cultivated and Planted Area*’ or ‘*Vegetable Production and Shipment Statistics*’) by nitrogen content in residue per crop production calculated from Matsumoto (2000). Amount of nitrogen in residue plowed into soil was calculated by multiplying the total amount of nitrogen in residue by the proportion of the amount of nitrogen in residue plowed into soil estimated from crop area of each treatment for wheat straw surveyed by MAFF.

[Crops other than rye (for grain), oats (for grain) and Tea]

The amount of nitrogen in each crop residue plowed into soil were calculated by multiplying nitrogen content in residue per crop production calculated from Matsumoto (2000) by annual crop production (by MAFF, *Statistics of Cultivated and Planted Area*) by the ratio other than burned in the field (burned in the field: 0.1, the default value in the *Revised 1996 IPCC Guidelines*).

For the amount of nitrogen in crop residue plowed into soil, the data of the *Document of Kagoshima prefectural Institute for Agricultural Development* was used for sweet potato and sugarcane, and the data of *Hokkaido Fertiliser Recommendations 2010* was used for sugar beets, potato, Japanese radish and onion, and the data of Owa (1996) was used for Chinese cabbage and Lettuce.

When any crop has no available data with respect to nitrogen content included in crop residue per crop production, the value for a similar type of crop was used. The same values were adopted for all fiscal years. For feed crops, the area not plowed into soil was excluded. For the crops which were assumed that field burning is not practiced in Japan, and which were not included in the calculation for the Field Burning of Crop Residues (4.F), “Proportion burned in field” were considered as “zero”.

Table 6-44 Residue/Crop production ratio, Nitrogen content in residue and Nitrogen content in residue per crop production for main crops

Crop type	Residue /Crop production ratio [t (residue)/ t (crop yield)] (A)	Nitrogen content in residue [kg-N/t (residue)] (B)	Nitrogen content in residue per crop production [kg-N/t (crop yield)] (A)×(B)	Note
Rice	-	6.88 ^a	-	Wet weight
Barley	1.39 ^a	3.68 ^a	0.511	
Wheat	1.39 ^a	3.68 ^a	0.511	
Soy	1.40 ^a	10.9 ^a	15.19	
Potatoes	0.0321 ^d	2.22 ^b	0.71	
Sweet potatoes	0.808 ^c	2.29 ^c	1.85	
Sugarbeets	0.0617 ^d	15.4 ^b	0.95	
Sugarcane	0.102 ^c	5.48 ^c	0.56	
Maize	1.20 ^a	3.52 ^a	4.22	
Japanese radish	0.033 ^d	2.84 ^b	0.93	
Chinese cabbage	0.018 ^d	4.03 ^d	0.71	
Cabbage	0.672 ^a	2.72 ^a	1.83	
Lettuce	0.040 ^d	4.08 ^d	1.64	
Onion	0.015 ^d	1.24 ^b	0.019	

a: Matsumoto N., Development of Estimation Method and Evaluation of Nitrogen Flow in Regional Areas (2000) (Reference 55)

b: Hokkaido Government, Hokkaido Fertiliser Recommendations 2010. (2010) (Reference 56)

c: Document of Kagoshima prefectural Institute for Agricultural Development

d: Owa N., *New Trends in Technology for Efficient Use of Nutrients – Nutritional Balance of Crops in Japan* (1996) (Reference 33)

Amount of nitrogen in crop residue plowed into soil (kg-N) (rice)

= Annual amount of residue plowed into soil [t] × Nitrogen content in crop residue [kgN/t]

Amount of nitrogen in crop residue plowed into soil (kg-N) (wheat and barley)

= $\sum_{\text{crop}} \{ \text{Annual crop production [t]} \times \text{Proportion crop residue plowed into soil per crop production [\%]} \times \text{Nitrogen content in residue per crop production [kgN/t]} \}$

Amount of nitrogen in crop residue plowed into soil (kg-N) (crops other than rye, oats, tea, rice, wheat and barley)

= $\sum_{\text{crop}} \{ \text{Annual crop production [t]} \times \text{Nitrogen content in residue per crop production [kgN/t]} \times (1 - \text{Proportion burned in field}) \}$

[Rye and Oats (for grain)]

In accordance with the default technique described in the *Revised 1996 IPCC Guidelines* and the *GPG (2000)*, the amount of nitrogen applied to soil by plowing in crop residues was determined by multiplying the annual production of each type of crop by the default value of each of the percentage of residues in the production of each crop, the average percentage of dry matter in the residues, the percentage less the percentage burned in the field, and the nitrogen content in the residues.

Nitrogen plowed into soil with crop residues (kg-N) (rye and oats)

= Annual crop yield (t) × Proportion of residue to crop yield × Average proportion of dry matter in crop residue (t-dm/t) × (1 – Proportion burned in field) × Nitrogen content (t-N/t-dm) × 10³

The production amount of rye and oats were calculated by multiplying the planted area by the yield per unit area. The planted area was divided into the area used for grain, for green crops and for others. However, the available statistics were not reported the category of rye for grain, (the survey has been discontinued since 1992 production) and therefore the value of the “total planted area” less the “area planted for green crops” taken from the available statistics was used as the area cultivated for grain

expediently, even though the planted area in this report covers the planting for grain only.

Table 6-45 Planted area of rye and oats (for grain)

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Rye	ha	50	119	110	120	170	170	100
Oats	ha	4,000	2,517	1,600	800	500	500	500

Source: The data are calculated by using the *Statistics of Cultivated and Planted Area* (MAFF) (Reference 13)

Table 6-46 Yields of rye and oats per unit area

Crop	Yield per unit area	Note
Rye	424 [kg/10a]	Data determined by specialists based on the results of rye cultivation tests in Japan
Oats	223 [kg/10a]	Data available only up to FY 1994. The 1994 figures were used for all fiscal years since most of the data before 1994 were available for major prefectures only.

Table 6-47 Proportion of residue to crop production, average proportion of dry matter in crop residues, nitrogen content

Crop	Proportion of residue	Average proportion of dry matter in residue	Nitrogen content	Proportion burned in field
Rye	2.84	0.90	0.0048	0.10
Oats	2.23	0.92	0.0070	0.10
Source	Determined by specialists	<i>GPG (2000)</i> , p. 4.58, Table 4.16		<i>Revised 1996 Guidelines</i> , Vol. 3, p. 4.83

[Tea]

For tea, "Leaf fall" and "Autumn pruning" were targeted as the residues which return into soils annually. In addition, as residues return into soil once in several years, "Medium pruning", which prunes the part of 30-50 cm from the ground and carried out once in about five years, was targeted. For the "Medium pruning", it assumed that it carried out by one fifth in every year in all area of tea field, and all of tea field will be renewal in five years. The residues' nitrogen contents were calculated by multiplying by nitrogen contents per unit area of "Leaf fall", "Autumn pruning" and "Medium pruning" by crop field areas. The crop field areas used for this were the data indicated in the *Statistics of Cultivated and Planted Area* by MAFF.

Nitrogen plowed into soil with crop residues (kgN) (Tea)

= (Nitrogen amount included in residue by autumn pruning [kgN/10a] + Nitrogen amount included in residue by leaf fall [kgN/10a]) × 10 × Cultivated area of tea [ha] + Nitrogen amount included in the residue by medium pruning [kgN/10a] × 10 × 1/5 × Cultivated area of tea [ha]

Table 6-48 Amount of nitrogen content included in tea residue of branch pruning

Kind of branch pruning		Amount of Nitrogen content (kg-N/10a)	Reference
Autumn pruning	Annual	7.7	Hoshina et al..(1982) (Reference 51), Kinoshita et al.. (2005) (Reference 52), Tachibana et al.. (1996) (Reference 53)
Medium pruning	Once in five years	19.4	Ohta et al.. (1996) (Reference 54)
Leaf fall	Annual	11.5	Hoshina et al..(1982) (Reference 51)

c) Uncertainties and Time-series Consistency

● Uncertainties

Because the estimation methods differ from one crop to the other, their uncertainties were calculated for respective crops.

The uncertainties of emission factors for crops other than rye and oats were assessed for each crop by combining the uncertainties for each parameter calculated by expert judgment and given for standard values in the *GPG (2000)*. The uncertainties of emission factors for rye and oats were calculated to combine each parameter determined by expert judgment or standard values in the *GPG (2000)*, and were determined to be 388% for rye and 392% for oats.

The uncertainties for activity data were assessed as 0.26% for tea and 0.32% for other crops by applying the standard errors in the *Crop statistics* and the *Statistics of Cultivated and Planted Area*, respectively.

As a result, the uncertainty of the emission combined from each crop uncertainty was determined to be 211%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

Emissions are estimated by using consistent estimation methods and data sources.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1.

e) Source-specific Recalculations

Because of the revision of nitrogen content in crop residue for some crops, the amount of nitrogen put into soils as crop residues were revised, and the emissions from FY1990 to FY2009 were revised. In the Agriculture sector, a 3-year average has been used. Thus, the emissions for FY2009 were revised in accordance with the revision and/or update of the activity data for FY2010.

f) Source-specific Planned Improvements

It is needed to discuss whether it will be possible to establish country-specific emission factors for Japan.

6.5.1.5. Plowing of Organic Soil (4.D.1.-)

a) Source/Sink Category Description

In Japan, there are organic soils in Hokkaido. Two types, “muck soil” and “peat soil”, are treated as organic soils. In Japan, the creation of farmland on organic soils was mostly completed by the 1970s, and in general farmers till land that has had soil dressing.

b) Methodological Issues

● **Estimation Method**

Emissions of N₂O from the plowing of organic soil were calculated by multiplying the area of the plowed organic soil of paddy field and upland field by the emission factor in accordance with the *Revised 1996 IPCC Guidelines* and the *GPG (2000)*.

$\begin{aligned} & \underline{N_2O \text{ emission associated with the plowing of organic soil (kg-N}_2\text{O)}} \\ & = \text{Emission factor for plowing of organic soil [kg-N}_2\text{O/ha]} \times \text{Area of plowed organic soil [ha]} \times \\ & 44/28 \end{aligned}$

●Emission Factors

For paddy cultivation in organic soils, it is known that N₂O emission in paddy field is lower than the one in upland field. In Japan, Nagata (2006) (Reference 43) observed N₂O emissions for paddy of organic soil in Hokkaido, but the observations included emissions from applied nitrogen. Therefore, country-specific emission factor is determined to be 0.30 [kgN₂O-N/ha/year] by deducting country-specific emission factor of fertilizers indicated in Akiyama (2006). For the upland field of organic soil, some observation results exists (Nagata 2006, Nagata 2009 (Reference 46)), but there is not much difference from the default of temperate region (8[kgN₂O-N/ha/year]) indicated in GPG(2000) p4.60 Table4.17. Therefore, default value is used for upland field.

●Activity Data

The area of plowed organic soil was established by multiplying the cultivated areas of paddy fields and common upland fields, obtained from the *Statistics of Cultivated and Planted Area* (MAFF), by the percentage of organic soils (peat soil and muck soil) in paddy fields and common upland fields in Japan. The percentage of organic soils was used data made from Takata et al.(2009)

Table 6-49 Percentage of organic soil

Soil type	~1991	1992	1997	2001	2002~
Paddy field	5.85%	5.85%	6.02%	6.15%	6.15%
Upland field	1.94%	1.94%	2.01%	2.07%	2.07%

*1992 data and 2001 data were original data. 1993-2000 data were calculated by using interpolation between 1992 and 2001. 1992 data was used for data before 1991 and 2001 data was used for data after 2002.

Source: Calculated from Takata et al.(2009) (Reference 48)

Table 6-50 Areas of organic soil

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Area of organic soil (paddy field)	ha	166,491	163,328	161,541	157,194	154,119	153,504	152,151
Area of organic soil (field)	ha	24,735	24,296	24,420	24,281	24,198	24,198	24,116

c) Uncertainties and Time-series Consistency

●Uncertainties

N₂O emissions by plowing of organic soil were calculated in two category, paddy field and upland field. Therefore, the uncertainties were also calculated separately, and finally two uncertainties were combined as total uncertainty.

The uncertainties for emission factors were calculated aggregating the uncertainties of each parameter given in the *GPG (2000)* and references or calculated from the data of references. The combined uncertainties for emission factor were determined to be 248% for paddy field and 900% for upland field. For the uncertainty for activity data, 0.11% of the standard error for paddy rice and 0.26% of the standard error for upland field crops given in the *Statistics of Cultivated and Planted Area* were applied. As a result, the uncertainties of the emissions were determined to be 712%. The uncertainty assessment methods are summarized in Annex 7.

●Time-series Consistency

Emissions are estimated by using consistent estimation methods and data sources.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1.

e) Source-specific Recalculations

No recalculations.

f) Source-specific Planned Improvements

For paddy fields, country-specific emission factors are used. However, there are issues such as the exclusion of influence for stubble which remains in the ground surface after harvest and the influence of the plowing of straw residue, to avoid double counting of emissions. It is necessary to promote further elaborate check to reflect more suitable national condition to the emission factor, including upland field which use default emission factor.

6.5.1.6. Direct Emissions (CH₄)

CH₄-generating bacteria are absolutely anaerobic, and if soil is not maintained in an anaerobic state, CH₄ generation is not possible. Upland soils are normally oxidative and in aerobic condition. Therefore, CH₄ is not produced by these soils. For that reason, direct emission of CH₄ from soil has been reported as “NA”.

6.5.2. Pasture, Range and Paddock Manure (4.D.2.)

The method for calculating CH₄ and N₂O emissions from pasture, range, and paddock cattle manure is described in 6.3.1 “Livestock Waste Management: Cattle, Swine and Poultry (4.B.1., 4.B.8., 4.B.9.)” (see 6.3.1). N₂O emissions are counted in 4.D.2.

6.5.3. Indirect Emissions (4.D.3.)**6.5.3.1. Atmospheric Deposition (4.D.3.-)****a) Source/Sink Category Description**

This section provides the estimation methods for N₂O indirect emissions caused by atmospheric deposition of nitrogen compounds volatilized as NH₃ and NO_x from synthetic fertilizer or domestic livestock manure.

b) Methodological Issues**●Estimation Method**

N₂O emissions have been calculated in accordance with Decision Tree of the *GPG (2000)* (Page 4.69, Fig. 4.8).

Calculation of N₂O emissions associated with atmospheric deposition

Emissions of N₂O from atmospheric deposition [kg-N₂O]
 = emission factor [kg-N₂O-N/kg-NH₃-N+NO_x-N]
 × Amount of nitrogen volatilized from ammonia and nitrogen oxides from livestock manure
 and synthetic fertilizers [kg-NH₃-N+NO_x-N] × 44/28

● Emission Factors

The default value given in the Revised 1996 IPCC Guidelines has been used as the emission factor for this source.

Table 6-51 Emission factor for N₂O emissions associated with atmospheric deposition

	Emission Factor [kg-N ₂ O-N/kg-NH ₃ -N & NO _x -N deposited]
N ₂ O emissions associated with atmospheric deposition	0.01

Source: Revised 1996 IPCC Guidelines Vol.2 Table 4-18 (GPG (2000) Table4.18) (Reference 3)

● Activity Data

The amounts of nitrogen (kg) contained in ammonia and nitrogen oxides that volatilize from synthetic fertilizers and livestock manure applied to agricultural soil were calculated for activity data. For the amount of manure-derived nitrogen applied to agricultural soil, the portion of nitrogen content in the livestock manure in Japan which was returned to agricultural soil, calculated in the 4.B. *Manure Management* section, was used to maintain consistency in the nitrogen cycle. Also, the portion of human waste which was returned to agricultural soil as fertilizer was added to the activity data reported in this section.

$$A = N_{FERT} * Frac_{GASF} + N_{ANI}$$

$$= N_{FERT} * Frac_{GASF} + N_B * Frac_{GASM1} + (N_D + N_{FU}) * Frac_{GASM2}$$

- A: Amount of nitrogen that volatilizes as ammonia and nitrogen oxides from synthetic fertilizers, livestock manure, and human waste (kg-NH₃-N+NO_x-N)
- N_{FERT}: Demand for synthetic nitrogen fertilizers (kg-N)
- Frac_{GASF}: Percentage of volatilization as ammonia and nitrogen oxides from synthetic fertilizers (kg-NH₃-N + NO_x-N/kg-N)
- N_{ANI}: Amount of nitrogen that volatilizes as ammonia and nitrogen oxides from livestock manure and human waste (kg-NH₃-N + NO_x-N/kg-N)
- N_B: Amount of nitrogen included in livestock manure (kg-N)
- Frac_{GASM1}: Percentage of volatilization as ammonia and nitrogen oxides from livestock manure during treatment (kg NH₃-N + NO_x-N/kgN)
- N_D: Amount of manure-derived fertilizer applied to agricultural soil (kg-N)
- N_{FU}: Amount of human waste-derived fertilizer applied to agricultural soil (kg-N)
- Frac_{GASM2}: Percentage of volatilization as ammonia and nitrogen oxides from nitrogen contained in livestock manure and human waste applied to agricultural soils(kg-NH₃-N + NO_x-N/kg-N)

For synthetic fertilizers, “demand for nitrogen-based fertilizers” given in the Ministry of Agriculture, Forestry and Fisheries *Yearbook of Fertilizer Statistics (Pocket Edition)* was used for the amount of fertilized nitrogen (N_{FERT}), and the default value given in the Revised 1996 IPCC Guidelines was used for the percentage of volatilization (Frac_{GASF}).

For livestock manure, the values determined in the *Manure Management (4.B.)* section (excluding the amount dispersed in the atmosphere as N₂O as well as the amount treated by the “Incineration” or “Purification” in the *Manure Management (4.B.)*) (Table 6-24) was used for livestock manure applied to farmland (N_D), and the default value given in the Revised 1996 IPCC Guidelines was used for the percentage of volatilization (Frac_{GASM}).

Activity data of human waste was calculated by multiplying the amount of human waste-derived nitrogen calculated with *Waste Treatment in Japan* by the value indicated in Table 6-24.

The amount of nitrogen volatilized in the process of treating livestock manure as NH_3 and NO_x was calculated by multiplying the nitrogen amount of livestock manure excreted in shed and pasture (N_B) by the figures indicated in Table 6-24.

Table 6-52 Proportion of nitrogen volatilized from synthetic fertilizers and livestock manure as ammonia or nitrogen oxides

	Value	Unit
Frac _{GASF}	0.1	[kg- NH_3 -N + NO_x -N/kg of synthetic fertilizer nitrogen applied]
Frac _{GASM}	0.2	[kg- NH_3 -N + NO_x -N/kg of nitrogen excreted by livestock]

Source: Revised 1996 Guidelines Vol. 2, Table 4-17 (Reference 3)

Table 6-53 Nitrogen returned to agricultural soil

Item	Unit	1990	1995	2000	2005	2009	2010	2011
N applied to agricultural soil from livestock waste	tN	565,991	541,931	512,239	493,180	473,453	464,175	465,438
N applied to agricultural soil from human waste	tN	10,394	4,747	2,116	874	457	457	458

c) Uncertainties and Time-series Consistency

●Uncertainties

N_2O emissions volatilized from atmospheric deposition were calculated in two categories, nitrogen compounds derived from synthetic fertilizer and from livestock manure (including human waste). Therefore, the uncertainties were also calculated separately, and finally two uncertainties were combined as total uncertainty.

The uncertainties for emission factors were calculated by aggregating the uncertainty of each parameter, estimated by expert judgment or given as the standard values in the *GPG (2000)*. The aggregated uncertainty of emission factor was 107% for the application of synthetic fertilizer, and 71% for the application of livestock manure. For the uncertainties of the activity data for applied synthetic fertilizers, the same values as in 6.5.1.1. [*Direct Soil Emission:*] *Synthetic Fertilizers* were applied. For applied livestock manure, the uncertainties of the activity data were calculated from 6.3.1. [*Manure Management:*] *Cattle, Swine, and Poultry*. The total emissions uncertainty aggregated from all the uncertainties was 75%. The uncertainty assessment methods are summarized in Annex 7.

●Time-series Consistency

Emissions are estimated by using consistent estimation methods and data sources.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1.

e) Source-specific Recalculations

In responding to the revision of livestock population in the Manure Management (4.B.), the amount of livestock-origin nitrogen returns into crop field soil was changed; therefore, emissions for this category from FY2006 to FY 2009 were revised.

f) Source-specific Planned Improvements

It is needed to discuss the establishment of country-specific emission factors and the ratios of volatile nitrogen compounds in synthetic fertilizers.

6.5.3.2. Nitrogen Leaching and Run-off (4.D.3.-)

a) Source/Sink Category Description

This section provides the estimation methods for N₂O emissions from Nitrogen Leaching and Run-off.

b) Methodological Issues

● Estimation Method

N₂O emissions were calculated according to the Decision Tree in the *GPG (2000)* (Page 4.69, Fig. 4.8), by multiplying Japan's country-specific emission factors by the amount of nitrogen that leached and run-off.

$\frac{N_2O \text{ emission associated with nitrogen that leached and run-off (kg-N}_2\text{O)}{=} \text{Emission factor associated with nitrogen leaching and run-off [kg-N}_2\text{O-N/kg-N]} \times \text{Nitrogen that leached and run-off [kg-N]} \times 44/28$
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● Emission Factors

The N₂O emission from this source was calculated using the Japan-specific emission factor that had been established by various studies.

Table 6-54 Emission factor for N₂O emissions associated with nitrogen leaching and run-off

	[kg-N ₂ O-N/kg-N]
N ₂ O emission from nitrogen that leaches or runs off	0.0124

Source: Takuji Sawamoto et al., Evaluation of emission factors for indirect N₂O emission due to nitrogen leaching in agro-ecosystems. (Reference 35)

● Activity Data

Activity data was calculated by multiplying the default value of proportion of leaching and run-off given in the *Revised 1996 IPCC Guidelines* by the amount of nitrogen in livestock manure applied to agricultural soil and synthetic fertilizer derived from atmospheric deposition.

Table 6-55 Frac_{LEACH}: Proportion of nitrogen applied subject to leaching and run-off

Value	Unit
0.3	[kg-N/kg nitrogen of fertilizer or manure]

Source: *Revised 1996 IPCC Guidelines* Vol. 2, Table 4-17 (Reference 3)

c) Uncertainties and Time-series Consistency

● Uncertainties

N₂O emissions for nitrogen leaching and run-off were calculated in two category, synthetic fertilizer and livestock manure (including human waste). Therefore, the uncertainties were also calculated separately, and finally two uncertainties were combined as total uncertainty.

The uncertainties for emission factors were calculated aggregating the uncertainties of each parameter, estimated by expert judgments or given for standard values in the *GPG (2000)*. The aggregated uncertainty for emission factor was determined to be 113% for both synthetic fertilizers and livestock

manure. For the uncertainty of activity data, the same method used at “6.5.3.1. Atmospheric Deposition” was applied. As a result, the uncertainty of the emissions was determined to be 97%. The uncertainty assessment methods are summarized in Annex 7.

● **Time-series Consistency**

Emissions are estimated by using consistent estimation methods and data sources.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1.

e) Source-specific Recalculations

In responding to the revision of livestock population in the Manure Management (4.B.), the amount of livestock-origin nitrogen returns into crop field soil was changed; therefore, emissions for this category from FY2006 to FY 2009 were revised.

f) Source-specific Planned Improvements

Refer to the section “6.5.3.1. Atmospheric Deposition”.

6.5.3.3. Indirect Emissions (CH₄) (4.D.3.-)

Direct CH₄ emissions were zero, and indirect CH₄ emissions from crop fields were also taken as zero. Therefore, these sources have been reported as “NA”.

Except for atmospheric deposition or nitrogen leaching and run-off, there is no conceivable source of CH₄ emissions from cultivated farmland soil other than direct emissions from soil, animal production, and indirect emissions. Therefore, they have been reported as “NO”.

6.5.4. Other (4.D.4)

Because it is not likely that agricultural sources of CH₄ and N₂O emissions exist in Japan other than the direct soil emissions, and indirect emissions, these sources were reported as “NO” as was the case in previous years.

6.6. Prescribed Burning of Savannas (4.E.)

This source is given in the *Revised 1996 IPCC Guidelines* as “being for the purpose of managing pastureland in sub-tropical zones”. There is no equivalent activity in Japan, and this source has been reported as “NO”.

6.7. Field Burning of Agricultural Residues (4.F.)

Incomplete burning of crop residues in field releases CH₄ and N₂O into the atmosphere. CH₄ and N₂O emissions from this source are calculated and reported in this category.

CH₄ and N₂O emissions from Field Burning of Agricultural Residues in FY 2010 are 57 Gg-CO₂ eq. and 18 Gg-CO₂ eq., comprising 0.005% and 0.001% of total emissions (excluding LULUCF),

respectively. The value represents a reduction by 43.0% and 44.5% for CH₄ and N₂O from FY 1990, respectively.

Table 6-56 CH₄ and N₂O emissions from field burning of agriculture residues

Gas	Item	Unit	1990	1995	2000	2005	2008	2009	2010	
CH ₄	4.F.1. Cereals	Wheat	Gg-CH ₄	0.42	0.23	0.30	0.40	0.36	0.29	0.25
		Barley	Gg-CH ₄	0.15	0.10	0.09	0.08	0.08	0.07	0.06
		Maize	Gg-CH ₄	1.89	1.66	1.48	1.32	1.37	1.38	1.35
		Oats	Gg-CH ₄	0.02	0.02	0.04	0.04	0.03	0.03	0.03
		Rye	Gg-CH ₄	0.00	0.00	0.00	0.00	0.00	0.00	0.00
		Rice	Gg-CH ₄	2.06	2.27	1.53	1.06	0.87	0.83	0.83
	4.F.2. Pulses	Peas	Gg-CH ₄	0.01	0.01	0.00	0.00	0.00	0.00	0.00
		Soybeans	Gg-CH ₄	0.08	0.04	0.08	0.07	0.08	0.08	0.08
		Adzuki beans	Gg-CH ₄	0.02	0.02	0.02	0.02	0.01	0.01	0.01
		Kidney beans	Gg-CH ₄	0.01	0.00	0.00	0.00	0.00	0.00	0.00
		Peanuts	Gg-CH ₄	0.01	0.01	0.01	0.00	0.00	0.00	0.00
	4.F.3. Tubers and Roots	Potatoes	Gg-CH ₄	0.03	0.03	0.02	0.02	0.02	0.02	0.02
		Sugarbeat	Gg-CH ₄	0.06	0.06	0.06	0.06	0.06	0.06	0.05
	4.F.4. Sugarcane	Gg-CH ₄	0.06	0.04	0.04	0.03	0.04	0.04	0.04	
Total	Gg-CH ₄		4.8	4.5	3.7	3.1	2.9	2.8	2.7	
	Gg-CO ₂ eq		101	94	77	65	62	59	57	
N ₂ O	4.F.1. Cereals	Wheat	Gg-N ₂ O	0.006	0.003	0.005	0.006	0.005	0.004	0.004
		Barley	Gg-N ₂ O	0.002	0.002	0.001	0.001	0.001	0.001	0.001
		Maize	Gg-N ₂ O	0.027	0.024	0.021	0.019	0.020	0.020	0.019
		Oats	Gg-N ₂ O	0.001	0.001	0.002	0.002	0.002	0.002	0.002
		Rye	Gg-N ₂ O	0.000	0.000	0.000	0.000	0.000	0.000	0.000
		Rice	Gg-N ₂ O	0.056	0.062	0.042	0.029	0.024	0.023	0.023
	4.F.2. Pulses	Peas	Gg-N ₂ O	0.000	0.000	0.000	0.000	0.000	0.000	0.000
		Soybeans	Gg-N ₂ O	0.003	0.002	0.003	0.003	0.004	0.004	0.003
		Adzuki beans	Gg-N ₂ O	0.001	0.001	0.001	0.001	0.001	0.001	0.000
		Kidney beans	Gg-N ₂ O	0.000	0.000	0.000	0.000	0.000	0.000	0.000
		Peanuts	Gg-N ₂ O	0.000	0.000	0.000	0.000	0.000	0.000	0.000
	4.F.3. Tubers and Roots	Potatoes	Gg-N ₂ O	0.003	0.002	0.002	0.002	0.002	0.002	0.002
		Sugarbeat	Gg-N ₂ O	0.004	0.003	0.004	0.004	0.004	0.003	0.003
	4.F.4. Sugarcane	Gg-N ₂ O	0.001	0.001	0.001	0.001	0.001	0.001	0.001	
Total	Gg-N ₂ O		0.11	0.10	0.08	0.07	0.06	0.06	0.06	
	Gg-CO ₂ eq		33	32	25	21	20	19	18	
Total of all gases			Gg-CO ₂ eq	133	126	103	87	82	78	76

6.7.1. Rice, Wheat, Barley, Rye, and Oats (4.F.1.)

a) Source/Sink Category Description

This section provides the estimation methods for CH₄ and N₂O emissions from field burning of agricultural residues of rice, wheat, barley, rye, and oats.

b) Methodological Issues

● Estimation Method

CH₄ and N₂O emissions were calculated, using the default method indicated in the *Revised 1996 IPCC Guidelines* and the *GPG (2000)*, by multiplying the amounts of carbon and nitrogen released by field burning by the CH₄ emission factor and N₂O emission factor, respectively.

Wheat, barley, rye, and oats were cultivated either as grain or green crops. The portions of the green crops which were cultivated for use of the entire aboveground mass for cattle feed were excluded from

the calculation of emissions.

$$\begin{aligned} & \underline{CH_4 \text{ emission associated with field burning of agricultural residues (kg-CH}_4\text{)}} \\ & = \text{CH}_4 \text{ emission factor (kg-CH}_4\text{-C/kgC)} \times \text{Total carbon released (kg-C)} \times 16/12 \end{aligned}$$

$$\begin{aligned} & \underline{N_2O \text{ emission associated with field burning of agricultural residues (kg-N}_2\text{O)}} \\ & = \text{N}_2\text{O emission factor (kg-N}_2\text{O-N/kgN)} \times \text{total nitrogen released (kg-N)} \times 44/28 \end{aligned}$$

● Emission Factors

The default values shown in the Revised 1996 IPCC Guidelines and the GPG (2000) were used.

Table 6-57 Emission factors for CH₄ and N₂O emissions associated with field burning of rice, wheat, barley, rye, and oats residues

	Value	Unit
CH ₄	0.005	[kg-CH ₄ /kg-C]
N ₂ O	0.007	[kg-N ₂ O/kg-N]

Source: Revised IPCC Guidelines Vol.2 Table 4-16 (Reference 3)

● Activity Data

[Crops other than rice]

Activity data was calculated in accordance with the default method technique shown in the *Revised 1996 IPCC Guidelines* and the *GPG (2000)*, by multiplying by the crop yield by “Proportion of residue to crop yield”, “Proportion of dry matter in residue”, “Proportion burned in field”, “Oxidation rate” and “Carbon/nitrogen content of residues”.

$$\begin{aligned} & \underline{\text{Total carbon/total nitrogen released by field burning of agricultural residues (kg-C, kg-N)}} \\ & = \text{Annual crop yield (t)} \times \text{Proportion of residue to crop yield} \times \text{Proportion of dry matter in residue} \\ & \quad (\text{t-dm/t}) \times \text{Proportion burned in field} \times \text{Oxidation rate} \times \text{Carbon/nitrogen content of} \\ & \quad \text{residues (t-C/t-dm, t-N/t-dm)} \times 10^3 \end{aligned}$$

[Rice]

For rice, Amount of burning rice straw and rice chaff on crop field is surveyed by MAFF. The residues' nitrogen content was calculated by multiplying by the aforementioned data by nitrogen content (kgN/t) indicated in Japan's country-specific data of nutrient balance for each crop (Matsumoto, 2000). Therefore, emission was calculated by multiplying by the crop yield by “Amount of burning rice straw and rice chaff”, “Proportion of dry matter in residue”, “Oxidation rate” and “Carbon/nitrogen content of residues”.

$$\begin{aligned} & \underline{\text{Total carbon/total nitrogen released by field burning of agricultural residues (kgC, kgN) (Rice)}} \\ & = \text{Amount of burning rice straw and rice chaff [t]} \times \text{proportion of dry matter in residue [t-dm/t]} \times \\ & \quad \text{Oxidation rate} \times \text{Carbon/nitrogen content of residues [t-C/t-dm, t-N/t-dm]} \times 10^3 \end{aligned}$$

Table 6-58 Amount of burning rice straw and rice chaff on crop field

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Rice straw	t	438,197	536,908	429,091	276,619	183,904	183,904	183,904
Rice chaff	t	581,302	528,290	291,260	260,289	209,927	209,927	209,927
Total	t	1,019,499	1,065,198	720,350	536,908	393,831	393,831	393,831

Reference: Survey by MAFF

➤ **Annual crop yield**

[Wheat (grain), and barley (grain)]

The values reported in the *Crop Statistics* were used for the yield of wheat, and barley (grain).

- **Wheat and barley (green crops)**

Because data of the yields of green crop wheat and barley (excluding those for fodder) were not directly available, the annual yields were calculated by multiplying the area planted with wheat for green crops and other purposes, as shown in the *Statistics of Cultivated and Planted Area*, by the yield per unit area established for green crop rye and oats (excluding those for fodder) and proportionally divided by the yield of wheat and barley (grain)..

- **Rye and oats**

Because data of the yields of rye and oats were not directly available, the total annual yields were calculated by multiplying the area planted with rye or oats, as indicated based on the *Statistics of Cultivated and Planted Area*, by the yield per unit area

Table 6-59 Yield of rye and oats per unit area (kg/10a)

Crop	Yield per unit area	Data Source
Rye (grain)	424	Determined by specialists (based on rye crop tests in Japan)
Oats (grain)	223	MAFF, <i>Crop Statistics</i> (Reference 14)
Rye and Oats (green crops)	1,100	Determined by specialists (based on literature)

➤ **Residue/ Crop production ratio, dry matter fraction in residue, carbon content, proportion burned in field, and oxidation rate.**

The proportion burned in field for Wheat, Barley, Rye and Oats were determined on the basis of data of crop area by treating method for wheat straw surveyed by MAFF as shown in Table 6-60. Since the survey data don't exist for before FY2006, the value for FY2007 is applied to these years. Table 6-61 shows other parameters for each crop.

Table 6-60 Proportion burned in field for Wheat, Barley, Rye and Oats

Item	Unit	1990	1995	2000	2005	2009	2010	2011
Proportion burned in field	%	13.5	13.5	13.5	13.5	11.6	10.6	10.6

*Estimated from the survey by MAFF

➤ **Nitrogen content**

The specific nitrogen content value was determined for each of rice, wheat, barley, and oats (green crop), based on the results of various studies carried out in Japan. The nitrogen content of green crop wheat/barley was calculated using the average of nitrogen contents in wheat and barley weighted by yield. The default nitrogen content values in the *GPG (2000)* were used for rye and oats (grain). The nitrogen content for rye (green crop) was calculated by multiplying Japan's country-specific value for oats (green crop) by the value resulting from "rye (grain) / oats (grain)".

Table 6-61 Proportions of residue to crop yield, dry matter in residue, carbon content, and oxidation rate

Crop	Residue/ Crop product ratio	Dry matter fraction in residue ^{a)}	Carbon content	Nitrogen content	Oxidation rate
Rice	---	0.85 ^a	0.4144 ^a	0.00688 ⁱ	0.90 ^b
Wheat (grain)	1.39 ¹	0.85 ^a	0.4853 ^a	0.00368 ⁱ	
Barley (grain)	1.39 ¹	0.85 ^a	0.4567 ^a	0.00368 ⁱ	
Wheat/barley (green crop)	---	0.17 ^c	0.48 ^{d,g}	0.017 ^{h,g}	
Rye	2.84 ^e	0.90 ^c	0.4710 ⁱ	0.0048 ^f	
Oats	2.23 ^e	0.92 ^c	0.4710 ⁱ	0.007 ^f	
Rye (green crop)	---	0.17 ^c	0.4710 ⁱ	0.0116 ^h	
Oats (green crop)	---	0.17 ^c	0.4710 ⁱ	0.0169 ^h	

a: *GPG (2000)*, p. 4.58, Table 4.16 (Reference 4)

b: Survey by MAFF

c: Determined based on the percentage of dry matter in green crop wheat indicated in the *Standard Table of Feed Composition in Japan* (National Agriculture Research Organization, pub. by Japan Livestock Association)

d: Determined based on the values shown in the *GPG (2000)* for wheat (grain) and barley (grain) by apportioning for yields

e: Determined based on the results of crop tests for rye and oats in Japan

f: Used the average of the values shown for “wheat” and “barley” in the *Good Practice Guidance (2000)*.

g: Values change over the years

h: Owa, *New Trends in Technology for Efficient Use of Nutrients – Nutritional Balance of Crops in Japan* (1996) (Reference 33)

i: Matsumoto N., Development of Estimation Method and Evaluation of Nitrogen Flow in Regional Areas (2000) (Reference 55)

c) Uncertainties and Time-series Consistency

●Uncertainties

The uncertainty assessment was conducted by each crop. The uncertainties for emission factors were calculated to combine the uncertainty of each parameter determined by expert judgment or given in the *GPG (2000)* as the default values. The uncertainties for activity data applied the standard error in each statistics (the *Crop Statistics* and the *Statistics of Cultivated and Planted Area*) or the value decided by the 2002 Committee for Greenhouse Gas Emission Estimation Methods. The uncertainty assessment results of the emissions by each crop were provided in Annex 7 Table 11. The uncertainty assessment methods are summarized in Annex 7.

●Time-series Consistency

Emissions are estimated by using consistent estimation methods and data sources.

d) Source-specific QA/QC and Verification

Tier 1 QC activities have been conducted in accordance with the *GPG (2000)* methods. Tier 1 QC activities focus on the verification of the parameters for activity data and emission factors and the archive of reference materials. QA/QC activities are summarized in Annex 6.1.

e) Source-specific Recalculations

The amount of rice straw and rice chaff burned on crop field and proportion burned in field for wheat, barley, rye, and oats from FY2008 to FY2010 were revised. Therefore, emissions from FY2007 to FY2010 were revised.

f) Source-specific Planned Improvements

For the use of the default parameter in the *Revised 1996 IPCC Guidelines* or the *GPG (2000)*, it is needed to discuss whether country-specific parameter can be established for Japan.

6.7.2. Maize, Peas, Soybeans, Adzuki beans, Kidney beans, Peanuts, Potatoes, Sugarbeets & Sugar cane (4.F.1., 4.F.2., 4.F.3., 4.F.4.)

a) Source/Sink Category Description

This section provides the estimation methods for CH₄ and N₂O emissions from field burning of agricultural residues by Maize, Peas, Soybeans, Adzuki beans, Kidney beans, Peanuts, Potatoes, Sugarbeet & Sugar cane.

b) Methodological Issues

● **Estimation Method**

CH₄ and N₂O emissions were calculated in accordance with the relevant Decision Tree in the *GPG (2000)* (page 4.52, Fig. 4.6), by multiplying the total carbon released or total nitrogen released, as calculated by the default method, by the emission factors.

● **Emission Factors**

Same emission factors used for rice, wheat, and barley residues were used (Table 6-57)

● **Activity Data**

Activity data was calculated by multiplying the yield of each crop shown in the *Crop Statistics* and the *Vegetable Production and Shipment Statistics* published by MAFF by the parameters shown in the following calculation formula.

<p><i>Total carbon released by field burning of agricultural residues [kg-C] (Potatoes, Sugarbeets, Sugarcane)</i> = Annual crop production [t] × Residue/Crop product ratio × Dry matter fraction in residue [t-dm/t] × Proportion burned in field × Oxidation rate × Carbon content of residues [t-C/t-dm] × 10³</p>
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<p><i>Total carbon released by field burning of agricultural residues [kg-C] (Crops other than Potatoes, Sugarbeets, Sugarcane)</i> = Annual crop production [t] × Residue/Crop product ratio [t-dm/t] × Proportion burned in field × Oxidation rate × Carbon content of residues [t-C/t-dm] × 10³</p>
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<p><i>Total nitrogen released by field burning of agricultural residues [kg-N]</i> = Annual crop production [t] × Residue/Crop product ratio × Proportion burned in field × Oxidation rate × Carbon/nitrogen content of residues [t-N/t-dm or t-N/t] × 10³</p>
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Table 6-62 Residue/ crop product ration, dry matter, carbon content, nitrogen content, proportion burned in field , and oxidation rate

Crop	Residue/ Crop product ratio	Dry matter fraction in residue	Carbon content	Nitrogen content	Proportion burned in field	Oxidation rate
Maize	1.20 ^{eA}	0.86 ^h	0.4709 ^{hD}	0.0035 ^{eE}	0.10 ^c	0.90 ^c
Peas	0.60 ^{eA}	0.87 ^h	0.45 ^{aD}	0.0101 ^{eE}		
Soybean	1.40 ^{eA}	0.89 ^h	0.45 ^{aD}	0.0109 ^{eE}		
Adzuki beans	0.89 ^{eA}	0.89 ^h	0.45 ^{aD}	0.0098 ^{eE}		
Kidney beans	0.60 ^{eA}	0.89 ^h	0.45 ^{aD}	0.0101 ^{eE}		
Peanuts	0.94 ^{eA}	0.86 ^h	0.45 ^{aD}	0.0054 ^{eE}		
Potatoes	0.032 ^{bB}	-	0.4226 ^{hD}	0.0222 ^{fD}		
Sugarbeets	0.062 ^{bB}	-	0.4072 ^{hD}	0.0154 ^{fD}		
Sugar cane	0.102 ^{gB}	-	0.4235 ^{hD}	0.0055 ^{gD}		

A: Residue (wet) / Crop production (wet)

B: Residue (dry) / Crop production (wet)

D: N content (or C content) / Residue (dry)

E: N content (or C content) / Residue (wet)

Source:

a. In the absence of default values, the values for dicotyledonous and monocotyledonous plants were used. Murayama, N., et al., *Alimentation of Crops and Fertilizer*, Buneido, p. 26 (Bowen: Trace Elements in Biochemistry, 1966)

b. Owa, *New Trends in Technology for Efficient Use of Nutrients – Nutritional Balance of Crops in Japan* (1996) (Reference 33)

c: *Revised 1996 IPCC Guidelines*

d: Although default values are not available, the median value of the values indicated in the *Revised 1996 IPCC Guidelines*, Vol. 2, p. 4.30 (0.001 – 0.02) were used.

e: Matsumoto N., *Development of Estimation Method and Evaluation of Nitrogen Flow in Regional Areas* (2000) (Reference 55)

f: Hokkaido Government, *Hokkaido Fertiliser Recommendations 2010*. (2010) (Reference 56)

g: Document of Kagoshima prefectural Institute for Agricultural Development

h: *GPG (2000)*, p. 4.58, Table 4.16 (Reference 4)

c) *Uncertainties and Time-series Consistency*

● *Uncertainties*

The uncertainty assessment was conducted by each crop. The uncertainties for emission factors were calculated to aggregate the uncertainty of each parameter determined by expert judgment and given for default values in *the GPG (2000)*. For the uncertainties of the activity data, the value decided by the Committee for Greenhouse Gas Emission Estimation Methods in 2002 was applied. The uncertainty assessment results of the emissions by each crops were provided in Annex 7 Table 11. The uncertainty assessment methods are summarized in Annex 7.

● *Time-series Consistency*

Emissions are estimated by using consistent estimation methods and data sources.

d) *Source-specific QA/QC and Verification*

Refer to section” 6.7.1. Rice, Wheat, Barley, Rye, and Oats”.

e) *Source-specific Recalculations*

In the Agriculture sector, a 3-year average has been used. Thus, the emissions for FY2009 were revised in accordance with the revision and/or update of the activity data for FY2010.

f) Source-specific Planned Improvements

For the use of the default parameter in the *Revised 1996 IPCC Guidelines* or the *GPG (2000)*, it is needed to discuss whether country-specific parameter can be established for Japan.

6.7.3. Dry bean (4.F.2.-)

Dry beans are a type of kidney beans, and the term refers to the mature, husked vegetable. Kidney beans in Japan are eaten before ripening, however, which means there is little of this type of product. Kidney beans are included in Beans (4.F.2.), under ‘Other crops’ and, therefore, the dry beans have been reported as “IE”.

6.7.4. Other (4.F.5.)

It is possible that agricultural residue other than cereals, pulse, root vegetables and sugar canes are burnt in the fields. However, data on actual activity is not available and it is not possible to establish the emission factor. Therefore, these sources have been reported as “NE”.

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Chapter 7. Land Use, Land-Use Change and Forestry (CRF sector 5)

7.1. Overview of Sector

The land use, land-use change, and forestry (LULUCF) sector deals with greenhouse gas (GHG) emissions and removals resulting from land use such as forestry activities and land-use change. Japan classifies its national land into six categories—Forest land, Cropland, Grassland, Wetlands, Settlements, and Other land—and subdivides each of them into two subcategories by distinguishing them on the basis of whether or not land conversion has been occurred, in accordance with the *GPG-LULUCF*. It also uses 20 years, a default value in the *GPG-LULUCF*, when distinguishing the land conversion. GHG emissions and removals in this sector consist of carbon stock changes in five carbon pools (aboveground biomass, belowground biomass, dead wood, litter, and soil), direct N₂O emissions from N fertilization, N₂O emissions from drainage of soils, N₂O emissions from disturbance associated with land-use conversion to cropland, CO₂ emissions from agricultural lime application, and non-CO₂ emissions from biomass burning. In this chapter, above- and below ground biomass are referred to collectively as “living biomass”, and dead wood and litter collectively as “dead organic matter”.

Japan’s total land area as of FY2010 is about 37.8 million ha. The largest portion of the national land is Forest land, which covers about 25.0 million ha. The second-largest portion is Cropland, which covers about 3.98 million ha. In addition, Grassland, Wetlands, Settlements, and Other land cover about 0.99 million ha, 1.33 million ha, 3.76 million ha, and 2.77 million ha, respectively.

Japan’s national land is an archipelago consisting of Hokkaido, Honshu, Shikoku, Kyushu and other islands, and lies off the east coast of the Eurasian Continent. The archipelago has the general shape of a crescent and extends from northeast to southwest. Its northernmost point is located at about 45 degrees north latitude, and its southernmost point is located at about 20 degrees north latitude. Most of Japan’s national land is located in a temperate, humid climate zone. Some islands in the southern part of Japan belong to a subtropical climate zone, and the northern part of Japan is located in a cool-temperate climate zone. The average annual temperature and precipitation in Tokyo, the capital city of Japan located in the temperate, humid climate zone, are 16.3 degrees centigrade and 1,528.8mm; those in Sapporo, Hokkaido prefecture, located in the cool-temperate climate zone, are 8.9 degrees centigrade and 1,106.5 mm; and those in Naha, Okinawa prefecture, located in the subtropical climate zone, are 23.1 degrees centigrade and 2,040.8 mm, respectively.¹

The LULUCF sector contains both sources and sinks; however, in Japan, it has been a net sink continuously since FY1990. Net removals in FY2010 were 72,909 Gg-CO₂; this accounts for 5.9% of the total national emissions (excluding LULUCF). The net removals in FY2010 also represent an increase of 4.0% over the FY1990 value and an increase of 1.5% over the FY2009 value.

This chapter is divided into 14 sections. Section 7.2 describes the method of determining land-use categories. Section 7.3 describes general parameters for estimating carbon stock changes from land-use conversion. Sections 7.4 to 7.9 explain the estimation methods of carbon stock changes in each land-use category. GHG emissions by the LULUCF sector resulting from other than carbon stock

¹ The average annual temperatures and precipitation are the average of the years between FY 1981 and 2010. See National Astronomical Observatory, *2012 Chronological Scientific Tables* (Tokyo: Maruzen Inc., 2011) pp.182-183 and pp.194-195. With respect to the degrees of latitude, see Geographical Survey Institute, *Degrees of Latitudes and Longitudes of Japan’s Northernmost, Southernmost, Easternmost and Westernmost Points* <<http://www.gsi.go.jp/KOKUJYOHO/center.htm>>.

changes are described in sections 7.10 to 7.14.

7.2. Method of determining land-use categories

7.2.1. Basic approach

In accordance with the 6 land-use categories in the *GPG-LULUCF*, land is classified on the basis of the definitions in existing statistics and other sources. As for Forest land and Cropland, country-specific subcategories are determined (Forest land: forests with standing trees (intensively managed forests / semi-natural forests) / forests with less standing trees / bamboo; Cropland: rice fields / upland fields / orchard).

“Land remaining Land” and “Land converted to Land” in each land-use category are determined based on existing statistics. Land-use categories that cannot be directly determined from existing statistics are determined using estimation measures such as allocation of areas of land conversion by means of the ratio of actual land areas for each land-use category.

Table 7-1 Land-use transition matrix for Japan in FY1990

		(kha)					
Before Conversion	Forest land	Cropland	Grassland	Wetlands	Settlements	Other land	Total
After Conversion							
Forest Land	24,946.8	2.7	0.7	IE	IE	0.1	24,950.3
Cropland	7.0	4,587.8	0.002	0.3	IE	1.3	4,596.4
Grassland	1.0	0.9	1,017.6	0.1	IE	2.0	1,021.6
Wetlands	0.3	0.02	0.01	1,319.6	0.002	0.1	1,320.0
Settlements	19.3	21.4	3.2	IE	3,175.2	IE	3,219.0
Other land	4.8	15.4	3.9	IE	IE	2,638.8	2,662.8
Total	24,979.1	4,628.2	1,025.3	1,320.0	3,175.2	2,642.2	37,770.0

Table 7-2 Land-use transition matrix for Japan in FY2010

		(kha)					
Before Conversion	Forest land	Cropland	Grassland	Wetlands	Settlements	Other land	Total
After Conversion							
Forest Land	24,965.8	0.3	0.1	IE	IE	0.01	24,966.2
Cropland	1.2	3,975.0	0.000	0	IE	0.0	3,976.3
Grassland	0.2	0.7	991.9	0	IE	0.3	993.1
Wetlands	0.2	0.02	0.003	1,329.7	0.001	0.1	1,330.0
Settlements	10.8	7.2	1.3	IE	3,739.7	IE	3,759.0
Other land	0.8	7.2	3.2	IE	IE	2,754.2	2,765.4
Total	24,979.1	3,990.4	996.6	1,329.7	3,739.7	2,754.6	37,790.0

(Note) The conversion areas indicated as “IE” are included in the following categories.

- Wetlands, Settlements converted to Forest land → Other land converted to Forest land
- Settlements converted to Cropland → Other land remaining Other land
- Settlements converted to Grassland → Other land remaining Other land
- Wetlands, Other land converted to Settlements → Other land remaining Other land
- Wetlands, Settlements converted to Other land → Other land remaining Other land

7.2.2. Methods of determining land-use categories and areas

Japan determines land-use categories and areas on the basis of existing statistics (Table 7-3). Among them, the areas of “Land converted to Forest land” are estimated based on data of the areas of afforestation and reforestation under Article 3, paragraph 3, of the Kyoto Protocol, which are determined by utilizing orthophotos taken at the end of 1989 and recent satellite images, in addition to

existing statistics. The areas of “Forest land converted to other land-use categories” are estimated based on data of the areas of deforestation determined in the same way as afforestation and reforestation, in addition to data of the *World Census of Agriculture and Forestry* and the Forestry Agency’s records. For detailed information on the methods of determining the areas of afforestation, reforestation and deforestation, see section 11.3.2.3 in Chapter 11.

Table 7-3 Method of determining land use categories and areas

Land use category	Method of determining land use category	Method of determining area
Forest	Forests under Forest Law Article 5 and 7.2.	Forests with standing trees (intensively managed forests, semi-natural forests), forests with less standing trees and bamboo in the forests which are included in the regional forests plan according to the <i>Forestry Status Survey</i> [-2004] and the <i>National Forest Resources Database</i> [2005-] (Forestry Agency). ² Definitions of forest subcategories are given in Table 7-4.
Cropland	Rice fields, upland fields and orchard.	Rice fields, upland fields and orchard according to <i>Statistics of Cultivated and Planted Area</i> by the MAFF.
Grassland	Pasture land, grazed meadow land and grassland other than pasture land and grazed meadow land ³ .	Pasture land according to <i>Statistics of Cultivated and Planted Area</i> by the MAFF, grazed meadow land according to <i>World Census of Agriculture and Forestry</i> by the MAFF, and grassland other than pasture land and grazed meadow land identified in <i>Land Use Status Survey</i> by the MLITT.
Wetlands	Lands covered with water (such as dams), rivers, and waterways.	Lands covered with water, rivers, and waterways according to <i>Land Use Status Survey</i> , <i>Survey of Forestry regions</i> by the MLITT. Among them, the lands that are subject to revegetation activities (e.g. green areas along rivers and erosion control sites, a part of urban parks) are allocated to Settlements.
Settlements	Urban areas that do not constitute Forest land, Cropland, Grassland or Wetlands. Urban green areas are all wooded and planted areas that do not constitute Forest land.	Settlements are roads, residential land, school reservations, park and green areas, road sites, environmental facility sites, golf courses, ski courses and other recreation sites identified in <i>Land Use Status Survey</i> and other surveys by the MLITT. The included figures for urban green areas are taken from the surveys on urban green facilities conducted by the MLITT. (Details are shown at Table 11-10).
Other land	Any land that does not belong to the above land-use categories.	Determined by subtracting the total area belonging to the other land-use categories from the total area of national land according to <i>Land Use Status Survey</i> by the MLITT.

MAFF: Ministry of Agriculture, Forestry and Fisheries; MLITT: Ministry of Land, Infrastructure, Transport and Tourism

² The *Forestry Status Survey* and the *National Forest Resources Database* use the same definitions and survey methods for forests, and these two data bases have time-series consistency.

³ Grassland other than pasture land and grazed meadow land is the land that remains after subtracting grazed meadow land and national land under the jurisdiction of the Forestry Agency from “Grassland other than forests” in the *World Census of Agriculture and Forestry*. Its present status is mainly wild grassland (including perennial pasture land, degenerated pasture land, and areas abandoned after cultivation and becoming wild).

Table 7-4 Definitions of forest subcategories

<u>Forest with standing trees:</u> Forest that does not fall under "Forest with less standing trees" and has a tree crown cover of standing trees 30% or higher (including young stands with the degree of stocking ⁴ of 3 or higher). Even if the tree crown cover of standing trees is less than 30%, forest in which the sum of the crown covers of both standing trees and bamboo is 30% or higher, while dominated by standing trees, is also included.	<u>Intensively managed forest:</u> Forest land that is subject to artificial regeneration such as tree planting and seeding, and in which no less than 50% of the volume (or the number) of standing trees are of tree species subject to artificial regeneration.
<u>Forest with less standing trees:</u> Forest in which the sum of the tree crown covers of both standing trees and bamboo is less than 30 percent.	<u>Semi-natural forest:</u> Forest with standing trees which is not classified as intensively managed forests
<u>Bamboo:</u> Forest that does not fall under "Forest with standing trees" and has a tree crown cover of bamboo (excluding "sasa" (a genus of running bamboo)) 30% or higher. Even if the tree crown cover of bamboo is less than 30%, forest in which the sum of the crown covers of both standing trees and bamboo is 30% or higher, while dominated by bamboo is also included.	

Reference: Forestry Agency of Japan, *Forest Status Survey* (March, 2007)

7.2.3. Survey methods and due dates of major land area statistics

Table 7-5 shows the survey methods and due dates of major land area statistics.

Table 7-5 Survey methods and due dates of major land area statistics

Name of the statistics / census	Survey method	Survey due date	Frequency	Presiding ministry
<i>Forest Status Survey</i>	Complete count survey	March, 31 st	Approximately 5 years	MAFF (Forestry Agency)
<i>National Forest Resources Database</i>	Complete count survey	April, 1 st	Every year (Since 2005)	MAFF (Forestry Agency)
<i>Statistics of Cultivated and Planted Area (Survey of cropland area)</i>	[Cropland area] Ground measurement survey (sample) [Expansion area and converted area of cropland] Tabular survey (using documents from relevant agencies and aerial photographs, etc.)	[Cropland area] July, 15th [Expansion area and converted area of cropland] July, 15th in the previous year until July, 14th	Every year	MAFF
<i>World Census of Agriculture and Forestry (Survey of Forestry Regions~2000)</i>	Complete count survey	August, 1st	Every 10 years	MAFF
<i>Land Use Status Survey</i>	Complete count Survey	March, 31st	Every year	MLITT

Details for urban green facilities are shown at Table 11-10.

MAFF: Ministry of Agriculture, Forestry and Fisheries; MLITT: Ministry of Land, Infrastructure, Transport and Tourism

7.2.4. Land area estimation methods

Some land areas cannot be directly determined from existing statistics; therefore, they are estimated using the following methods:

- Interpolation or trend extrapolation
- Allocation of areas of land conversion by means of the ratio of actual land areas for each

⁴ The degree of stocking is the ratio of actual volume to the expected volume of the forest stand, multiplied by 10.

land-use category

- Allocation of areas of land conversion by means of the ratio of converted land areas for a certain year

- ***Interpolation and trend extrapolation***

- ***Method***

The areas of forest land before 2004 were surveyed at an interval of approximately five years, and it was difficult to directly determine the areas of forest land in the unsurveyed years. Therefore, they were estimated by interpolation or extrapolation by means of linear expressions based on the areas in the surveyed years.

- ***Land-use category***

5.A.2. Land converted to Forest land (FY1991- 1994, 1996- 2001 and 2003- 2004).

- ***Allocation of areas of land conversion by means of the ratio of actual land areas for each land-use category***

- ***Method***

In Japan, it is difficult to obtain the areas of “Upland field converted to Forest land”, “Orchard converted to Forest land” and “Pasture land converted to Forest land” directly from existing statistics, since those are collectively reported as “Arable land”. Therefore, these land areas were estimated by multiplying the “Arable land converted to Forest land” by the ratios of actual land areas for each of the land-use categories (Upland field, Orchard and Pasture land).

- ***Land-use category***

- 5.A.2. Land (Cropland and Grassland) converted to Forest land
- 5.B.2. Land (Forest land, Grassland, Wetlands and Other land) converted to Cropland
- 5.C.2. Land (Forest land, Cropland, Wetlands and Other land) converted to Grassland
- 5.E.2. Land (Cropland and Grassland) converted to Settlements
- 5.F.2. Land (Cropland and Grassland) converted to Other land

- ***Allocation of areas of land conversion by means of the ratio of converted land area for a certain year***

- ***Method***

In Japan, it is difficult to directly obtain annual land areas of Cropland, Grassland, Settlements and Other land converted to Wetlands, respectively. Therefore, the annual land ratios of Cropland, Grassland, Settlements and Other land converted to Wetlands to “Land converted to Wetlands” in FY1998, which are assumed to be the same as the land ratio in each year, are multiplied by the areas of “Land converted to Wetlands” in each year to obtain the area of respective land use category converted to Wetlands.

- ***Land use category***

5.D.2. Land (Cropland, Grassland, Settlements and Other land) converted to Wetlands

7.3. Parameters for estimating carbon stock changes from land use conversions

Prior to the sections describing detailed methods for each land-use category, basic parameters used for estimating carbon stock changes due to land use conversions are shown here (Table 7-6 to Table 7-9) to prevent the reiteration of indicating these parameters in each subsequent section. For some parameters and estimation methods, details are given in the section indicated in “Note”.

Table 7-6 Living biomass stocks for each land-use category before and after conversion

Land use category		Biomass stock [t-d.m./ha]	Note	
Before conversion	Forest land	137.28 (FY2010)	Calculated by utilizing the values of biomass stocks in land of deforestation under Article 3, paragraph 3, of the Kyoto Protocol, which are provided from the NFRDB. In addition, the values before 2004 are extrapolated by means of trend from 2005 to the latest year. (Reference values [t-d.m./ha]: FY1990: 101.48, FY2005: 129.02, FY2008: 133.17, FY2009: 135.23)	
	Cropland	Rice field	0	Biomass stocks are assumed to be "0".
		Upland field	0	Biomass stocks are assumed to be "0".
		Orchard	30.63	Calculated by multiplying the average age and growth rate given in Ito <i>et al.</i> "Estimating the Annual Carbon Balance in Warm-Temperature Deciduous Orchards in Japan"
	Grassland	13.50	GPG-LULUCF Table 3.4.2 and Table 3.4.3 (warm temperate wet)	
	Wetlands, Settlements and Other land	0	Biomass stocks are assumed to be "0".	
Immediately after conversion	All land uses	0	Biomass stocks immediately after conversion are assumed to be "0".	
After conversion	Forest land	-	Removals in this land are directly estimated based on the implied removal factor of AR activity under the Kyoto Protocol. See section 7.4.2.b)1)	
	Cropland	Rice field	0	Biomass stocks are assumed to be "0".
		Upland field	0	Biomass stocks are assumed to be "0".
		Orchard	0	Biomass stocks are assumed to be "0".
	Grassland	2.70	One fifth of the value of GPG-LULUCF Table 3.4.2 and Table 3.4.3 (warm temperate wet)	
	Settlements	-	See section 7.8.2.b)1)	
	Wetlands and Other land	0	Biomass stocks are assumed to be "0".	

Table 7-7 Carbon stocks of dead wood for each land-use category before and after conversion

Land-use Category		Carbon Stock [t-C/ha]	Note
Before Conversion	Forest land	15.02 (FY2010)	Calculated from carbon stocks in dead wood in all forests. (Reference values [t-C/ha]: FY1990: 16.35, FY2005: 16.35, FY2008: 15.96, FY2009: 15.05)
	Cropland, Grassland, Wetlands, Settlements, Other land	0*	Assumed as zero (Section 4.3.2 in Volume 4 of the 2006 IPCC Guidelines, Tier.1)
Immediately after conversion	All land uses	0	Biomass stocks immediately after conversion are assumed to be "0".
After conversion	Forest land	13.01	Average carbon stocks per unit area in 20-year-old forests obtained by the CENTURY-jfos model
	Cropland, Grassland, Wetlands, Other land	0*	Assumed as zero (Section 4.3.2 in Volume 4 of the 2006 IPCC Guidelines, Tier.1)
	Settlements	0	Assumed as zero

* For some subcategories, stock changes are estimated as zero despite the fact that carbon stock values exist. See each section for details.

Table 7-8 Carbon stocks of litter for each land-use category before and after conversion

Land-use Category		Carbon Stock [t-C/ha]	Note
Before Conversion	Forest land	7.29 (FY2010)	Calculated from carbon stocks in litter in all forests. (Reference values [t-C/ha]: FY1990: 7.18, FY2005: 7.18, FY2008: 7.03, FY2009: 7.28)
	Cropland, Grassland, Wetlands, Settlements, Other land	0*	Assumed as zero (Section 4.3.2 in Volume 4 of the 2006 IPCC Guidelines, Tier.1)
Immediately after conversion	All land-uses	0	Biomass stocks immediately after conversion are assumed to be "0".
After conversion	Forest land	5.644	Average carbon stocks per unit area in 20-year-old forests obtained by the CENTURY-jfos model
	Cropland, Grassland, Wetlands, Other land	0*	Assumed as zero (Section 4.3.2 in Volume 4 of the 2006 IPCC Guidelines, Tier.1)
	Settlements	—	See section 7.8.2.b)2)

* For some subcategories, stock changes are estimated as zero despite the fact that carbon stock values exist. See each section for detail.

Table 7-9 Carbon stocks of soil for each land-use category before and after conversion

Land-use Category		Carbon Stock [t-C/ha]	Note
Before conversion	Forest land	85.31 (FY2010)	Value of soil carbon stocks at 0-30 cm depth one year before the inventory year National average value calculated by the CENTURY-jfos model. In addition, the value in FY2006 is applied to the years before 2005. (Reference values [t-C/ha]: FY1990: 85.74, FY2005: 85.74, FY2008: 84.21, FY2009: 85.31)
	Wetlands	88.00	Default value (GPG-LULUCF Table 3.3.3, Wetland soils/ Warm temperate).
After conversion	Forest land	82.954	Value of soil carbon stocks at 0-30 cm depth. Average carbon stocks per unit area in 20-year-old forests obtained by the CENTURY-jfos model.
	Wetlands	-	Under investigation
Commonly used before and after conversion	Rice field	71.38	Value of soil carbon stocks at 0-30 cm depth. Data provided from Dr. Makoto Nakai, National Institute for Agro-Environmental Sciences (Undisclosed) Cropland: see section 7.5.2.b)3) Grassland (pasture land): see section 7.6.2.b)2)
	Upland field	86.97	
	Orchard	77.46	
	Cropland (average)	76.40	
	Grassland	134.91	
	Settlements	-	Under investigation
Other land	-	Under investigation	

7.4. Forest land (5.A.)

Forests absorb CO₂ from the atmosphere by photosynthesis, fix carbon as organic substances, and store these substances for a given period. In contrast, forests emit CO₂ due to the effects of events such as logging and natural disturbances.

All forests in Japan are managed forests, and they consist of intensively managed forests, semi-natural forests, bamboo, and forests with less standing trees. Japan's forest land area in FY2010 was about 25.0 million ha—about 66.1% of the total national land area. The net removal by this category in

FY2010 was 76,677 Gg-CO₂ (excluding 2.33 Gg-CO₂ eq. of CH₄ and N₂O emissions resulting from biomass burning); this represents a decrease of 2.4% below the FY1990 value, and an increase of 4.1% over the FY2009 value.

In this section, Forest land is divided into two subcategories, “Forest land remaining Forest land (5.A.1.)” and “Land converted to Forest land (5.A.2.)”, and they are described separately in the following subsections.

Table 7-10 Emissions and removals in Forest land resulting from carbon stock changes

Gas	Category	Carbon pool	Unit	1990	1995	2000	2005	2008	2009	2010
CO ₂	5.A. Forest land	Total	Gg-CO ₂	-78,592.3	-87,340.7	-90,689.2	-92,010.9	-79,927.7	-73,673.0	-76,676.9
		Living Biomass	Gg-CO ₂	-72,530.6	-79,560.1	-83,695.2	-86,939.9	-76,498.9	-70,940.9	-74,553.8
		Dead Wood	Gg-CO ₂	-2,853.2	-3,798.1	-2,848.3	-1,094.6	188.1	703.8	1,157.6
		Litter	Gg-CO ₂	-2,694.5	-2,355.2	-1,779.1	-1,039.1	-720.6	-604.0	-517.5
		Soil	Gg-CO ₂	-514.1	-1,627.2	-2,366.6	-2,937.3	-2,896.2	-2,831.9	-2,763.2
	5.A.1. Forest land remaining Forest land	Total	Gg-CO ₂	-76,762.1	-86,456.0	-90,066.4	-91,548.0	-79,535.4	-73,331.6	-76,372.1
		Living Biomass	Gg-CO ₂	-71,107.2	-78,851.5	-83,188.6	-86,556.8	-76,171.6	-70,657.7	-74,301.1
		Dead Wood	Gg-CO ₂	-2,512.3	-3,629.3	-2,727.0	-1,003.5	265.6	770.8	1,217.3
		Litter	Gg-CO ₂	-2,546.7	-2,282.0	-1,726.4	-999.5	-687.0	-575.0	-491.6
		Soil	Gg-CO ₂	-595.9	-1,693.3	-2,424.3	-2,988.1	-2,942.5	-2,869.7	-2,796.7
	5.A.2. Land converted to Forest land	Total	Gg-CO ₂	-1,830.3	-884.7	-622.8	-463.0	-392.3	-341.4	-304.8
		Living Biomass	Gg-CO ₂	-1,423.3	-708.6	-506.6	-383.1	-327.3	-283.3	-252.8
		Dead Wood	Gg-CO ₂	-340.9	-168.9	-121.3	-91.1	-77.6	-67.0	-59.7
		Litter	Gg-CO ₂	-147.9	-73.3	-52.6	-39.5	-33.7	-29.0	-25.9
		Soil	Gg-CO ₂	81.8	66.1	57.8	50.8	46.2	37.9	33.6

7.4.1. Forest land remaining Forest land (5.A.1.)

a) Source/Sink Category Description

This subcategory deals with carbon stock changes in “Forest land remaining Forest land”, which has remained forested without conversion for the past 20 years as of FY2010. The net removal by this subcategory in FY2010 was 76,372 Gg-CO₂ (excluding 2.33 Gg-CO₂ eq. of CH₄ and N₂O emissions resulting from biomass burning); this represents a decrease of 0.5% below the FY1990 value and an increase of 4.1% over the FY2009 value.

b) Methodological Issues

1) Carbon stock changes in Living Biomass in “Forest land remaining Forest land”

● Estimation Method

In accordance with the decision tree provided in the *GPG-LULUCF*, carbon stock changes in living biomass in all Forest land are estimated by the Tier 2 stock change method using the country specific values of the amount of biomass accumulation. In this method, the carbon stock change in the living biomass pool is estimated by calculating the difference between the absolute amounts of carbon stocks in the pool at two points of time.

$$\Delta C_{LB} = \sum_k \{(C_{i2} - C_{i1}) / (t_2 - t_1)\}_k$$

ΔC_{LB} : annual change in carbon stocks in living biomass (t-C/yr)

t_1, t_2 : time points of carbon stock measurement

C_{i1} : total carbon in biomass calculated at time t_1 (t-C)

C_{i2} : total carbon in biomass calculated at time t_2 (t-C)

k : type of forest management

The carbon stocks in living biomass are calculated by multiplying the stand volume of each tree species by wood density, the biomass expansion factor, the root-to-shoot ratio and the carbon fraction of dry matter. These parameters except the carbon fraction are determined for each tree species.

$$C = \sum_j \{ [V_j \cdot D_j \cdot BEF_j] \cdot (1 + R_j) \cdot CF \}$$

- C : carbon stock in living biomass (t-C)
- V : merchantable volume (m³)
- D : wood density (t-d.m./m³)
- BEF : biomass expansion factor for conversion of merchantable volume
- R : root-to-shoot ratio
- CF : carbon fraction of dry matter (t-C/t-d.m.)
- j : tree species

Since Japan calculates the carbon stock change of living biomass in the total forest land in this manner, the carbon stock change of living biomass in “Forest land remaining Forest land” is obtained by subtracting the carbon stock change in “Land converted to Forest land” from the total change. For the method of estimating carbon stock change in “Land converted to Forest land”, see section 7.4.2.b)1).

● Parameters

➤ Volume

The Forestry Agency has developed the National Forest Resources Database (NFRDB) in order to estimate GHG emissions/removals from forests. The data in the NFRDB are based on the information on areas, tree species and forest ages, contained in the “Forest Registers”.

Merchantable volumes are estimated by multiplying the areas for each tree species and forest age in the NFRDB by merchantable volumes per area for each tree species and forest age in yield tables. Base data for the volumes per area are shown in Table 7-11 below. With respect to estimating the volumes of Japanese cedar, Hinoki cypress and Japanese larch in private forests, which are major tree species of intensively managed forests in Japan, the volumes per area reported in new yield tables, reflecting the newest survey results, are applied.

$$V = \sum_{m,j} (A_{m,j} \cdot v)$$

- V : merchantable volume (m³)
- A : area (ha)
- v : merchantable volume per area (m³/ha)
- m : age class or forest age
- j : tree species

Table 7-11 Yield tables used to estimate merchantable volume

Tree species			Yield tables	
			Private Forest	National Forest
Intensively managed forests	Conifer	Japanese cedar, Hinoki cypress, Japanese larch	New yield tables	Yield tables developed by Regional Forest Offices
		Other conifer	Yield tables developed by prefectures	
	Broad leaf			
Semi-natural forests				

- **Forest registers and yield tables developed by prefectures or Regional Forest Offices**

When forest plans are established for private and national forests (all forest lands are divided into 158 planning areas, and forest plans are established each year for 1/5 [about 30] of them), field surveys are implemented in these forests to develop a Forest Register which includes data on area, forest age, volume by tree species and so on. When forest plans are established (private forests: by each prefecture, national forests: by Regional Forest Offices of National Forests), the Forest Registers are updated to reflect the change in volume due to growth, cutting and disturbances. In general, the volume data described in the Forest Registers are estimated based on land area data and yield tables, which provide stand growth in the case that typical forest practices are implemented for each region, tree species and site class (yield tables show the relationship between forest age or age class and volume per area).

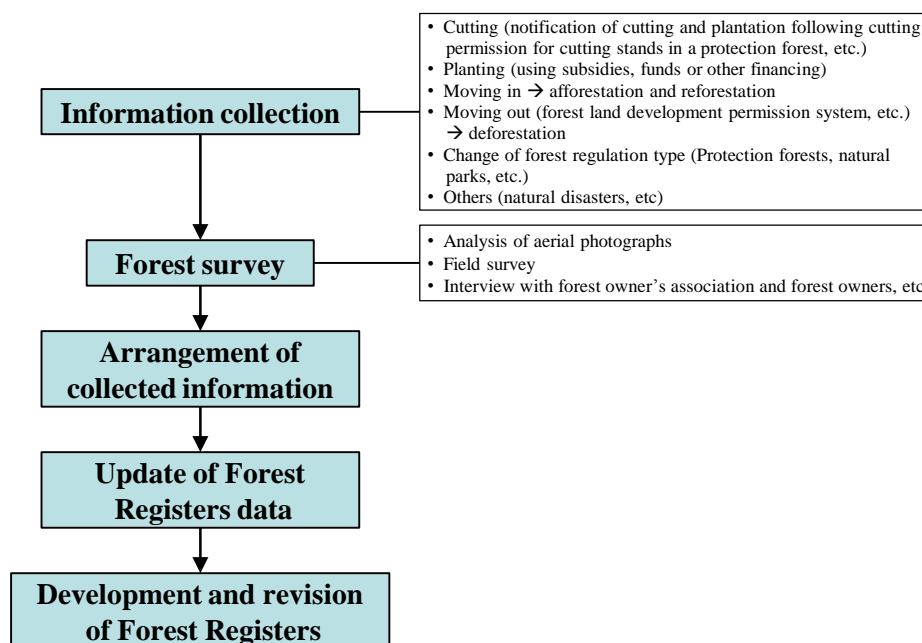


Figure 7-1 Procedures for developing Forest Registers

- **New yield tables (Japanese cedar, Hinoki cypress, Japanese larch)**

In 2006, the Forestry and Forest Products Research Institute developed new yield tables for Japanese cedar, Hinoki cypress and Japanese larch based on the results of a field survey over the country. The area of these three tree species covers 82% of intensively managed forests in private forests.

The new yield tables for Japanese cedar were established for 7 regions, those for Hinoki cypress for 4 regions and those for Japanese larch for 2 regions.

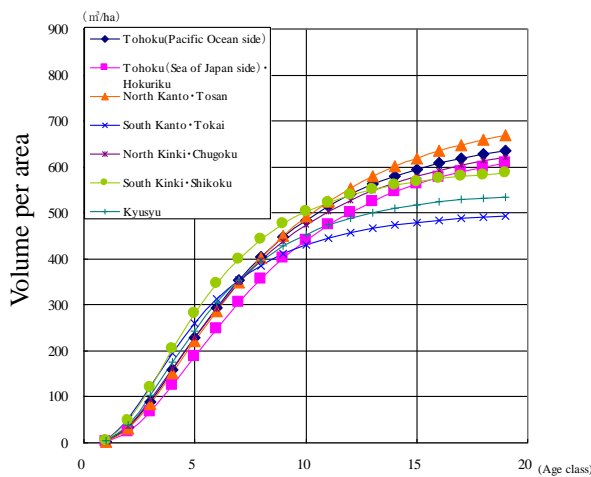


Figure 7-2 Yield tables made from forest resources monitoring survey data (Japanese cedar : 7 areas)

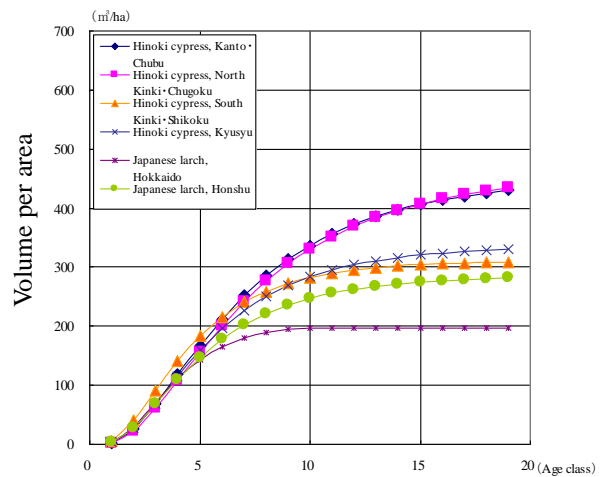


Figure 7-3 Yield tables made from forest resources monitoring survey data (Hinoki cypress : 4 areas, Japanese larch : 2 areas)

➤ **Biomass expansion factor and Root-to-shoot ratio**

The biomass expansion factors (BEF) and root-to-shoot ratios (R) were set based on the results from a biomass survey on dominant tree species, and existing research reports which were implemented by the Forestry and Forest Products Research Institute (Table 7-12).

BEFs were calculated for two age classes (20 years and below / 21 years and above) and for each tree species, because it was identified that BEFs differed between young forests and mature forests as apparent in Figure 7-4 below. On the other hand, R values were established only for tree species, because the root-to-shoot ratio was not correlated with forest age (Figure 7-5).

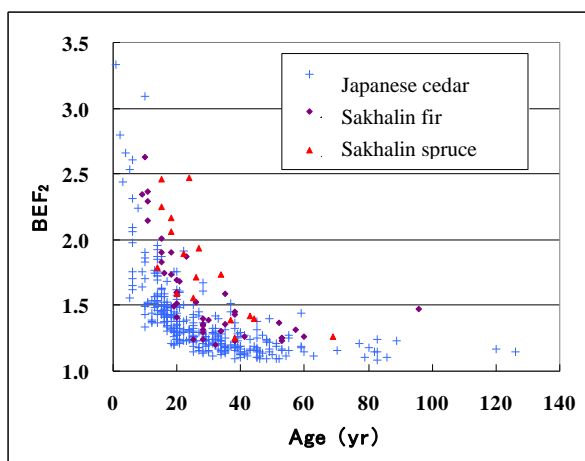


Figure 7-4 Relationship between biomass expansion factor (BEF) and forest age

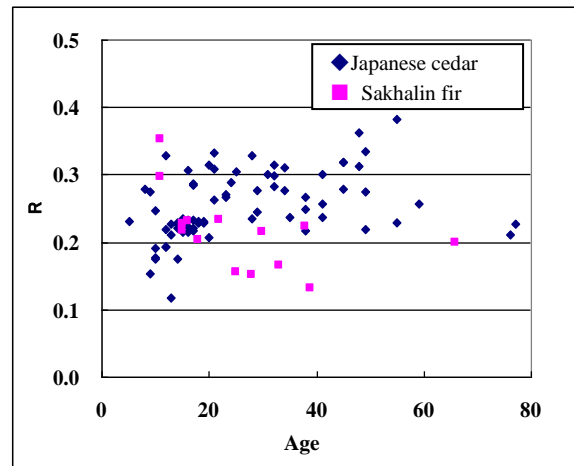


Figure 7-5 Relationship between root-to-shoot ratio (R) and forest age

➤ **Wood density**

Wood density (D) data were set based on the results from biomass survey on dominant tree species and existing research reports which were implemented by the Forestry and Forest Products Research

Institute (Table 7-12). These D values were established only for tree species, because wood density was not correlated with forest age.

➤ **Carbon fraction of dry matter**

The default value given in the *GPG-LULUCF* has been adopted as the carbon fraction (CF) of dry matter (Table 7-12).

Table 7-12 Biomass expansion factor, root-to-shoot ratio, wood density for tree species and carbon fraction

		BEF [-]		R [-]	D [t-d.m./m ³]	CF [t-C./t-d.m.]	Note	
		≤ 20	> 20					
Conifer trees	Japanese cedar	1.57	1.23	0.25	0.314	0.5		
	Hinoki cypress	1.55	1.24	0.26	0.407			
	Sawara cypress	1.55	1.24	0.26	0.287			
	Japanese red pine	1.63	1.23	0.26	0.451			
	Japanese black pine	1.39	1.36	0.34	0.464			
	Hiba arborvitae	2.38	1.41	0.20	0.412			
	Japanese larch	1.50	1.15	0.29	0.404			
	Momi fir	1.40	1.40	0.40	0.423			
	Sakhaline fir	1.88	1.38	0.21	0.318			
	Japanese hemlock	1.40	1.40	0.40	0.464			
	Yezo spruce	2.18	1.48	0.23	0.357			
	Sakhaline spruce	2.17	1.67	0.21	0.362			
	Japanese umbrella pine	1.39	1.23	0.20	0.455			
	Japanese yew	1.39	1.23	0.20	0.454			
	Ginkgo	1.50	1.15	0.20	0.450			
	Exotic conifer trees	1.41	1.41	0.17	0.320			
		Other conifer trees	2.55	1.32	0.34		0.352	Applied to Hokkaido, Aomori, Iwate, Miyagi, Akita, Yamagata, Fukushima, Tochigi, Gunma, Saitama, Niigata, Toyama, Yamanashi, Nagano, Gifu and Shizuoka prefectures
			1.39	1.36	0.34		0.464	Applied to Okinawa prefecture
		1.40	1.40	0.40	0.423	Applied to prefectures other than above		
Broad leaf trees	Japanese beech	1.58	1.32	0.26	0.573			
	Oak (evergreen tree)	1.52	1.33	0.26	0.646			
	Japanese chestnut	1.33	1.18	0.26	0.419			
	Japanese chestnut oak	1.36	1.32	0.26	0.668			
	Oak (deciduous tree)	1.40	1.26	0.26	0.624			
	Japanese popular	1.33	1.18	0.26	0.291			
	Alder	1.33	1.25	0.26	0.454			
	Japanese elm	1.33	1.18	0.26	0.494			
	Japanese zelkova	1.58	1.28	0.26	0.611			
	Cercidiphyllum	1.33	1.18	0.26	0.454			
	Japanese big-leaf	1.33	1.18	0.26	0.386			
	Maple tree	1.33	1.18	0.26	0.519			
	Amur cork	1.33	1.18	0.26	0.344			
	Linden	1.33	1.18	0.26	0.369			
	Kalopanax	1.33	1.18	0.26	0.398			
	Paulownia	1.33	1.18	0.26	0.234			
	Exotic broad leaf trees	1.41	1.41	0.16	0.660			
	Japanese birch	1.31	1.20	0.26	0.468			
	Other broad leaf trees	1.37	1.37	0.26	0.469	Applied to Chiba, Tokyo, Kochi, Fukuoka, Nagasaki, Kagoshima, and Okinawa prefectures		
		1.52	1.33	0.26	0.646	Applied to Mie, Wakayama, Oita, Kumamoto, Miyazaki, and Saga prefecture		
		1.40	1.26	0.26	0.624	Applied to prefectures other than above		

BEF: Biomass expansion factor (20 = age class)

R: Root-to-shoot ratio

D: Wood density

CF: Carbon Fraction

● **Activity Data (Area)**

➤ **Determining the forest area**

Forest areas of intensively managed forests, semi-natural forests, forests with less standing trees and bamboo under the forest planning system are obtained from the “Forest Status Survey” for years earlier than FY2004 and from the National Forest Resource Database (NFRDB) for FY2005 and onward. Data for FY1991 through FY1994, FY1996 through FY2001, and FY2003 through FY2004 are estimated by interpolation by means of linear expression. In addition, area data of Sakhalin fir, Yezo spruce, Japanese chestnut oak and Oak (deciduous tree) before FY1990 are not available individually; therefore, these data are estimated from “other conifer” and “other broad leaf” area divided by the area ratio in FY1995.

Table 7-13 Classifications in Forest Status Survey (before 2004) and National Forest Resource Database (after 2005)

Conifer trees		Broad leaf trees	
Before 2004	After 2005	Before 2004	After 2005
Japanese cedar	Japanese cedar	Japanese chestnut oak	Japanese chestnut oak
Hinoki cypress	Hinoki cypress	Oak (deciduous tree)	Oak (deciduous tree)
Pine	Japanese red pine	Other broad leaf	Japanese beech
	Japanese black pine		Oak (evergreen tree)
Japanese larch	Japanese larch		Japanese chestnut
Sakhalin fir	Sakhalin fir		Japanese poplar
Yezo spruce	Yezo spruce		Alder
	Sakhalin spruce		Japanese elm
Other conifer	Sawara cypress		Japanese zelkova
	Hiba arborvitae		Cercidiphyllum
	Momi fir		Japanese big-leaf magnolia
	Japanese hemlock		Maple tree
	Japanese umbrella pine		Amur cork
	Japanese yew		Japanese lime
	Ginkgo		Linden
	Exotic conifer trees		Kalopanax
Other needle leaf	Paulownia		
		Exotic broad leaf trees	
		Other broad leaf	

➤ **Obtaining the land area of “Forest land remaining Forest land”**

This land area is estimated by subtracting the cumulative total area of “Land converted to Forest land” during the past 20 years from the total area of “Forest land” in the year subject to estimation. All areas of “Land converted to Forest land” are assumed to be intensively managed forests. For the activity data of “Land converted to Forest land”, see section 7.4.2.b)1).

Table 7-14 Area of “Forest land remaining Forest land” within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Forest land remaining Forest land	kha	24,807.4	24,826.1	24,825.2	24,954.0	24,936.6	24,919.8	24,941.2
Intensively managed forests	kha	10,144.9	10,284.8	10,279.7	10,298.3	10,275.9	10,270.4	10,260.0
Semi-natural forests	kha	13,354.5	13,220.3	13,195.2	13,315.7	13,333.5	13,349.6	13,360.8
Cut-over forests and lesser stocked forests	kha	1,159.0	1,171.0	1,197.4	1,186.0	1,170.8	1,142.8	1,161.7
Bamboo	kha	149.0	150.0	152.9	154.0	156.4	157.1	158.6

Source: Forest Status Survey, National Forest Resources Database (Forest Agency)

2) Carbon Stock Changes in Dead Organic Matter and Soils in “Forest land remaining Forest land”

● Estimation Method

In accordance with the decision tree provided in the *GPG-LULUCF*, these pools are estimated by the Tier 3 method. Emissions and removals from organic soils are reported as “IE” because emissions from and removals by mineral and organic soils are estimated in the model in an integrated manner.

Carbon emissions/removals in each pool per unit area are estimated by using the CENTURY-jfos model and are multiplied by the land area of each forest management type. The sum of the emissions/removals of all forest management types are the annual changes in total carbon stocks in dead wood, litter and soil.

$$\Delta C_{dls} = \sum_{k,m,j} (A_{k,m,j} \times (d_{k,m,j} + l_{k,m,j} + s_{k,m,j}))$$

ΔC_{dls} : Annual change in carbon stocks in dead wood, litter and soil (t-C/yr)

A : Area (ha)

d : Average carbon stock change per unit area in dead wood (t-C/ha/yr)

l : Average carbon stock change per unit area in litter (t-C/ha/yr)

s : Average carbon stock change per unit area in soil (t-C/ha/yr)

k : Type of forest management

m : Age class or forest age

j : Tree species

● Parameters

Average carbon stock changes per unit area for dead wood, litter and soils are calculated by the CENTURY-jfos model, which was modified from the CENTURY model (Colorado State University) to be applicable to Japanese climate, soil, and vegetation conditions.

➤ Assumptions and Parameters as the Keys for the CENTURY-jfos Model

Since the amounts of tree growth and stable soil carbon stocks were thought to vary depending on climatic or locational conditions, the data of climatic values and soil carbon stocks were aggregated for each tree species in each prefecture (Table 7-15). It was assumed that forests have continually existed and been routinely utilized, and that their soil carbon stocks have been in a nearly steady state. Next, the parameters in the CENTURY-jfos model were adjusted. First, the growth parameters of above-ground biomass were adjusted so that the model be fitted to the growth in the yield tables in association with climatic values calculated per prefecture and per tree species. Second, the parameters were adjusted so that soil carbon stocks after the 60-year cutting age after a spinup of 3,000 years be fitted to the parameters for each of the prefectures and tree species calculated by Morisada *et al.* (2004). The methodologies of adjusting each parameter are in accordance with Sakai *et al.* (in preparation).

Tuning of the CENTURY-jfos Model

The Forestry and Forest Products Research Institute adjusted the CENTURY model in order to apply it to the Japanese forest environment. That is, forests were classified by predominant tree species (Japanese Cedar, Hinoki Cypress, Pine species, Japanese Larch, Sakhaline Fir, Sakhaline Spruce, broad leaf trees, and other conifer trees), and the geographical distribution of the tree species and soil types underneath were identified for each prefecture. The climate conditions to run the model were prepared from the mesh climate data provided by the Meteorological Agency of Japan (Japan Meteorological Agency, 2002). The model was adjusted with parameters on tree growth so that the tree growth in the model conformed to yield tables, and it was also tuned so that its output of carbon stocks in soil conformed to actual values based on field surveys for each prefecture and tree species (Table 7-15). The model after these modifications was named as the CENTURY-jfos model. After the tuning, carbon stocks in dead wood, litter and soil, and their stock changes were calculated by the CENTURY-jfos for different types of forest management such as management with thinning or without thinning.

Average annual carbon stock changes per unit area in dead wood, litter and soil are calculated for 1 – 19 age classes (for 100 years) for each type of forest management by means of CENTURY-jfos in order to estimate carbon stock changes in these carbon pools using the same activity data as for living biomass.

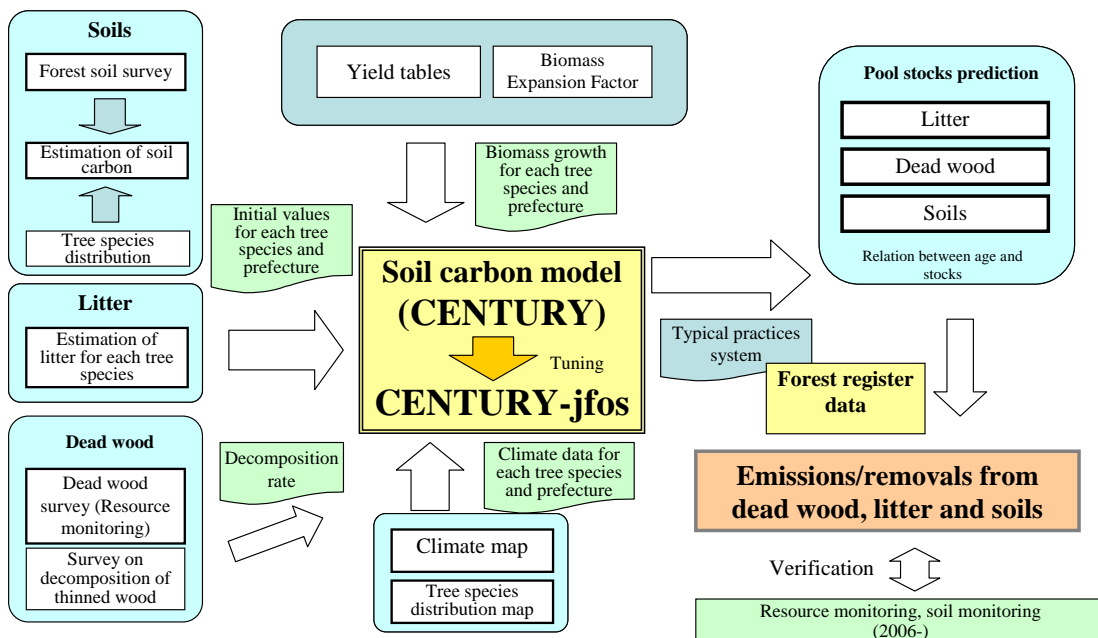


Figure 7-6 Estimation of emissions/removals in dead wood, litter and soils

Table 7-15 Standard soil carbon stocks used for the CENTURY-jfos model

(t-C/ha [30 cm depth])

Prefecture No.	Prefecture	Tree Species							
		Japanese Cedar	Hinoki Cypress	Pine species	Japanese Larch	Sakhaline Fir	Sakhaline Spruce	Broad Leaf Trees	Other Conifer Trees
1	Hokkaido	98.0	NA	100.6	91.0	88.0	93.7	91.0	89.4
2	Aomori	92.1	NA	94.3	83.3	109.1	NA	89.0	89.8
3	Iwate	89.5	93.6	92.7	93.9	98.1	NA	91.3	93.3
4	Miyagi	86.1	70.8	78.5	90.3	110.9	NA	82.8	80.5
5	Akita	81.1	NA	72.4	81.0	108.5	NA	82.6	79.6
6	Yamagata	83.2	79.7	68.0	81.0	97.4	NA	74.4	76.9
7	Fukushima	84.3	83.7	81.1	89.3	108.6	NA	81.4	85.0
8	Ibaraki	84.3	83.4	97.6	NA	NA	NA	91.2	90.8
9	Tochigi	83.0	86.1	91.6	100.6	133.4	NA	93.1	96.4
10	Gunma	88.7	88.3	93.9	95.1	98.1	NA	86.5	93.9
11	Saitama	81.3	82.4	96.2	106.8	NA	NA	85.8	94.7
12	Chiba	93.9	85.7	65.6	NA	NA	NA	84.6	76.4
13	Tokyo	79.2	81.6	85.7	94.7	NA	NA	63.9	84.3
14	Kanagawa	91.9	99.8	89.8	NA	NA	NA	94.9	99.1
15	Niigata	83.9	51.3	63.4	86.7	133.0	NA	85.3	86.9
16	Toyama	90.3	NA	72.5	88.5	106.0	NA	94.5	100.2
17	Ishikawa	82.7	80.2	70.2	NA	133.4	NA	86.6	74.3
18	Fukui	88.7	85.8	79.8	NA	NA	NA	90.1	80.6
19	Yamanashi	93.0	93.9	98.0	99.3	NA	NA	93.9	95.6
20	Nagano	102.1	100.5	96.0	108.4	106.0	NA	97.9	103.3
21	Gifu	100.5	94.8	79.1	99.6	107.8	NA	95.8	93.9
22	Shizuoka	94.6	96.7	69.1	90.7	NA	NA	90.0	93.7
23	Aichi	91.2	85.0	60.1	NA	NA	NA	78.5	77.2
24	Mie	92.1	84.4	63.8	97.1	NA	NA	78.7	80.5
25	Shiga	83.5	73.0	59.6	NA	NA	NA	79.5	65.8
26	Kyoto	74.0	67.4	63.3	NA	NA	NA	66.4	64.6
27	Osaka	78.9	74.0	60.9	NA	NA	NA	67.5	66.0
28	Hyogo	88.3	71.8	53.0	123.6	NA	NA	63.4	61.9
29	Nara	79.6	69.8	65.5	NA	NA	NA	73.4	69.4
30	Wakayama	72.1	70.5	58.2	NA	NA	NA	62.8	69.9
31	Tottori	73.8	74.9	75.6	121.2	NA	NA	72.3	75.4
32	Shimane	69.0	66.6	61.2	77.3	NA	NA	64.6	63.2
33	Okayama	80.3	73.7	51.4	121.2	NA	NA	65.2	63.6
34	Hiroshima	74.0	71.8	54.0	71.2	NA	NA	65.0	58.7
35	Yamaguchi	64.9	60.9	49.3	NA	NA	NA	55.2	54.8
36	Tokushima	72.9	63.7	63.6	NA	NA	NA	66.7	63.7
37	Kagawa	57.7	61.9	56.6	NA	NA	NA	57.2	57.7
38	Ehime	80.1	75.1	63.2	85.4	NA	NA	67.4	74.1
39	Kochi	81.4	76.1	73.8	NA	NA	NA	74.1	76.2
40	Fukuoka	97.3	88.9	77.5	NA	NA	NA	86.5	88.3
41	Saga	83.6	83.0	69.1	NA	NA	NA	79.6	82.9
42	Nagasaki	82.9	84.5	82.6	NA	NA	NA	78.9	84.5
43	Kumamoto	108.7	96.0	79.3	NA	NA	NA	93.5	95.6
44	Oita	109.9	100.5	108.3	130.3	NA	NA	99.1	101.4
45	Miyazaki	106.1	102.0	93.7	NA	NA	NA	98.0	99.6
46	Kagoshima	108.4	102.4	75.7	NA	NA	NA	90.8	97.0
47	Okinawa	58.5	NA	58.9	NA	NA	NA	58.0	58.5

● Activity Data (Area)

Forest area data provided by the NFRDB were used for the estimation.

c) Uncertainties and Time-series Consistency

● Uncertainty Assessment

The uncertainties of the parameters and activity data for living biomass were individually assessed on the basis of field study results, expert judgment, or the default values described in the *GPG-LULUCF*. The uncertainty estimates for dead organic matter and soil were assessed by calculating the variance of outputs from the CENTURY-jfos model. As a result, the uncertainty estimate was 11% for the total removals by “Forest land remaining Forest land”. The methodology used in the uncertainty assessment was in accordance with the guideline described in Annex 7. Uncertainty estimates regarding the major parameters in this category are shown in Table 7-16.

Table 7-16 Uncertainty estimates regarding major parameters in the Forest land category

		Uncertainty Estimates (%)	Country Specific (CS) or Default(D)	Remarks	
Forest land area		5.9	CS	Estimated based on uncertainty estimates of land areas in the NFRDB. Used 5.9% without distinguishing tree species.	
Volume of timber per area		22	CS	Estimated based on analysis of comparison between yield table and measured data	
Biomass Expansion Factor	Japanese cedar	≤20	3.5	CS	Estimated based on measured values
		>20	1.1	CS	
	Hinoki cypress	≤20	3.2	CS	
		>20	1.6	CS	
	Oak (deciduous tree)	≤20	8.6	CS	
		>20	2.1	CS	
Wood density	Japanese cedar		2.5	CS	
	Hinoki cypress		1.7	CS	
	Oak (deciduous tree)		1.6	CS	
Carbon fraction of dry matter	All tree species	2.0	D	Estimated based on the GPG-LULUCF default value. Used 2.0% without distinguishing tree species.	
Dead wood	All forests	22.1	CS	Result of uncertainty analysis of CENTURY-jfos model.	
Litter		51.0			
Soils		19.9			

● *Time-series Consistency*

There were no data for forest areas for FY1991 to FY1994, FY1996 to FY2001, and FY2003 to FY2004. Therefore, the time-series consistency was ensured by estimating these forest areas by means of interpolation.

d) *Source-/Sink-specific QA/QC and Verification*

QC is implemented in accordance with the Tier 1 approach described in the GPG (2000) and the GPG-LULUCF. The Tier 1 approach includes procedures for checking parameters and activity data, and for archiving references. The QA/QC activity procedures are described in section 6.1 of Annex 6.

e) *Source-/Sink-specific Recalculations*

● *Carbon stock changes in mineral soils on Intensively Managed Forests in FY2009*

Due to change in statistical value of the area of Land converted to Forest land, the carbon stock change is revised.

● *Carbon stock changes in living biomass on Forest with less standing trees*

Due to error correction on wood density parameter used for the estimation of carbon stock in Forest with less standing trees, carbon stock change on the land for FY1991 to FY2004 were recalculated.

● *Uncertainty assessment*

Estimation level of subcategories for uncertainty assessment was revised. In addition, uncertainties of

dead wood, litter and soils were updated due to recalculations made in response to the reexamination of the model.

f) Source-/Sink-specific Planned Improvements

None.

7.4.2. Land converted to Forest land (5.A.2)

a) Source/Sink Category Description

This subcategory deals with the carbon stock changes in Forest land converted from other land-use categories within 20 years. The net removal by this subcategory in FY2010 was 304.80 Gg-CO₂; this represents a decrease of 83.3% below the FY1990 value and a decrease of 10.7% below the FY2009 value.

b) Methodological Issues

1) Carbon stock change in Living Biomass in “Land converted to Forest land”

● *Estimation Method*

In the Tier 2 method, the annual carbon stock change in “Land converted to Forest land” (ΔC_{LF}) is to be estimated by summing the loss of carbon stock due to conversion (ΔC_L) and the change of carbon stock accumulated after conversion (ΔC_F). However, it is difficult to extract removals occurred on “Land converted to Forest land” directly from the data of the NFRDB because it deals with the stock change of living biomass of both "Forest land remaining Forest land" and "Land converted to Forest land" after land conversion collectively. On the other hand, it can be assumed that “Forest land subjected to AR activities under Article 3, paragraph 3 of the Kyoto Protocol” and “Land converted to Forest land” have similar nature. Therefore, ΔC_F was estimated by multiplying the area of converted land by the carbon stock change per unit area due to AR activities⁵. The ΔC_F value is reported all together under “Rice field converted to Forest land” in the CRF. ΔC_L was estimated and reported for each land-use category. For conversions from Rice field, Upland field and Other land, where carbon stocks of living biomass are assumed as 0, the carbon losses are reported as “NA”. For conversions from Wetlands and Settlements, where the areas of conversion are reported as “IE”, the carbon losses are reported as “NO”.

$$\Delta C_{LF} = \Delta C_L + \Delta C_F$$

$$\Delta C_L = \sum_i \{A_i \times (B_a - B_{b,i}) \times CF\}$$

$$\Delta C_F = A_{LF} \times IEF_{AR}$$

ΔC_{LF} : Annual carbon stock change in “Land converted to Forest land” (t-C/yr)

ΔC_L : Annual carbon stock change at the land conversion (t-C/yr)

ΔC_F : Carbon stock change in the converted land within 20 years since conversion (t-C/yr)

i : Land-use category before conversion

A_i : Annual increase of land area that has been converted from land-use type i to forest (ha/yr)

⁵ In the FY 2010 estimation, the average value of carbon stock change per unit area between FY 2005 and 2009 was adopted.

- B_a : Dry matter weight per unit area immediately after conversion to forest (t-C/yr)
 $B_{b,i}$: Dry matter weight per unit area before conversion from land-use type i to forest (t-C/yr)
 A_{LF} : Area of converted Forest land within 20 years (ha)
 IEF_{AR} : Average carbon stock change per unit area due to AR activities (equal to the implied removal factor) (t-C/ha/yr)
 CF : Carbon fraction of dry matter (t-C/t-d.m.)

● Parameters

➤ *Per unit area removals of Afforestation and Reforestation activities*

The average value of carbon stock change per unit area due to AR activities between FY2005 and FY2009 (2.8 t-C/ha) was applied to all reporting years.

➤ *Biomass stock in each Land-Use Category*

The parameters of Orchard and Grassland before conversion, shown in Table 7-6, are used.

● Activity Data (Area)

The areas of “Land converted to Forest land within 20 years” were calculated by summing the annually converted areas during the past 20 years. The estimation methods for annually converted areas from each land-use category are described below.

➤ *Total area of “Land converted to Forest land”*

It is presumed that the areas of “Land converted to Forest land” include AR areas, forest land restored from degraded land by natural succession, and land whose land-use categories are changed to “Forest land” due to other reasons. It is tentatively regarded that the areas of “Land converted to Forest land” are similar to the AR areas, and that the areas are determined in accordance with the concept of “overlap” described as a time series consistency and recalculation approach on section 6 of chapter 5 in the *GPG-LULUCF*, by using the AR areas and areas of forested cropland reported in the *Statistics of Cultivated and Planted Area*. In concrete terms, the AR areas are identified in detail by utilizing orthophotos taken at the end of 1989 and recent satellite images, but they are provided only from the FY2006 values. Therefore, the areas of “Land converted to Forest land” are estimated by setting an adjustment factor from the ratio between the AR areas since FY2006 and areas of forested cropland provided by the *Statistics of Cultivated and Planted Area*, and multiplying the areas of forested cropland since FY1990 by the adjustment factor. For further information on determining AR areas, see section 11.3.2.3 in Chapter 11.

➤ *Areas of “Cropland and Grassland converted to Forest Land”*

The areas of “Cropland converted to Forest land” are determined by utilizing the areas of forested cropland reported in the *Statistics of Cultivated and Planted Area*. As its subcategories, the areas of “Cropland converted to Forest land” are categorized to “Rice fields converted to Forest land”, “Upland fields converted to Forest land” and “Orchards converted to Forest land”. The areas of “Rice fields converted to Forest land” are determined by utilizing the areas of rice fields converted to forests provided by the *Statistics of Cultivated and Planted Area*. The areas of “Upland fields and Orchards converted to Forest land” are estimated by dividing the areas of arable land converted to forests, also provided by the *Statistics of Cultivated and Planted Area*, by means of the existing area ratios of upland fields, orchards and pasture land.

The areas of “Grassland converted to Forest land” are calculated by summing the areas of pasture land

converted to forests reported in the *Statistics of Cultivated and Planted Area* and those of grazed meadow converted to forests reported in *A Move and Conversion of Cropland*.

➤ **Areas of “Other land converted to Forest land”**

The areas of “Wetlands, Settlements, and Other land converted to Forest land” cannot be obtained directly from statistics. Therefore, they are estimated by subtracting the summed areas of “Cropland converted to Forest land” and “Grassland converted to Forest land” from the total area of “Land converted to Forest land”, and the areas of “Wetlands, Settlements, and Other land converted to Forest land” are reported collectively in “Other land converted to Forest land”.

Table 7-17 Area of land converted to Forest land (single year)

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Forest land	kha	3.5	1.6	1.4	0.8	0.6	0.5	0.4
Cropland converted to Forest land	kha	2.7	1.2	1.1	0.6	0.5	0.4	0.3
Rice field	kha	0.9	0.5	0.4	0.2	0.2	0.2	0.1
Upland field	kha	1.3	0.6	0.5	0.3	0.2	0.2	0.1
Orchard	kha	0.5	0.2	0.2	0.1	0.1	0.0	0.0
Grassland converted to Forest land	kha	0.7	0.3	0.3	0.2	0.1	0.1	0.1
Wetlands converted to Forest land	kha	IE	IE	IE	IE	IE	IE	IE
Settlements converted to Forest land	kha	IE	IE	IE	IE	IE	IE	IE
Other land converted to Forest land	kha	0.1	0.0	0.0	0.0	0.0	0.0	0.0

Table 7-18 Land converted to Forest land within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Forest land	kha	142.9	70.8	50.9	38.2	32.5	28.1	25.0
Cropland converted to Forest land	kha	121.9	57.7	40.6	30.0	25.3	21.9	19.6
Rice field	kha	53.8	23.7	15.9	11.0	9.0	8.4	7.6
Upland field	kha	46.8	23.7	17.7	14.0	12.2	10.2	9.1
Orchard	kha	21.4	10.3	6.9	4.9	4.1	3.3	2.9
Grassland converted to Forest land	kha	19.3	11.6	9.0	7.3	6.4	5.4	4.9
Wetlands converted to Forest land	kha	IE	IE	IE	IE	IE	IE	IE
Settlements converted to Forest land	kha	IE	IE	IE	IE	IE	IE	IE
Other land converted to Forest land	kha	1.7	1.5	1.2	0.9	0.8	0.7	0.6

2) Carbon Stock Changes in Dead Organic Matter and Soils in “Land converted to Forest land”

● Estimation Method

Carbon stock changes in dead wood, litter and soils were calculated under the assumption that these carbon stocks change linearly over 20 years from those in land-use categories “Other than Forest land” to those in Forest land. The calculation was implemented by applying the average carbon stocks obtained by the CENTURY-jfos model, in which mineral soils and organic soils are integrated. Therefore, emissions from organic soils were reported as “IE”.

$$\Delta C_{LF,i} = A_i \times (C_{after} - C_{before,i}) / 20$$

$\Delta C_{LF,i}$: annual change in carbon stocks in dead wood, litter or soils in land-use category i converted to Forest land (t-C/yr)

A_i : area of land-use category i being converted to Forest land within the past 20 years (ha)

C_{after} : average carbon stocks per unit area in land-use category after conversion (forests) (t-C/ha)

$C_{before,i}$: average carbon stocks per unit area in land-use category i before conversion (t-C/ha)

i : land-use category (Cropland, Grassland, Wetlands, Settlements, or Other land)

- **Parameters**

Parameters for each carbon pool in Table 7-7 (dead wood), Table 7-8 (litter) and Table 7-9 (soil) were used, in particular, for the categories Cropland, Grassland, Wetlands, Settlements and Other land before conversion and for the category Forest land after conversion.

- **Activity Data (Area)**

- **Total areas of “Land converted to Forest land”**

See Table 7-18.

c) Uncertainties and Time-series Consistency

- **Uncertainty Assessment**

The uncertainties of the parameters and activity data for living biomass, dead organic matter, and soil were individually assessed on the basis of field study results, expert judgment, or the default values described in the *GPG-LULUCF*. As a result, the uncertainty estimate was 24% for the entire removal by “Land converted to Forest land”. The methodology used in the uncertainty assessment is described in Annex 7. In addition, concrete uncertainty estimates for individual parameters in this category will be reported in future submissions after investigation is completed.

- **Time-series Consistency**

Time-series consistency for this subcategory is ensured.

d) Source-/Sink-specific QA/QC and Verification

QC is implemented in accordance with the Tier 1 approach described in the GPG (2000) and the *GPG-LULUCF*. The Tier 1 approach includes procedures for checking parameters and activity data, and for archiving references. The QA/QC activity procedures are described in section 6.1 of Annex 6.

e) Source-/Sink-specific Recalculations

- **Carbon stock changes in living biomass**

In the previous submission, implied removal factor (IEF_{AR}) of living biomass of AR was calculated by including both loss of biomass stocks on non-forest land immediately before conversion (the loss of biomass) and gain of biomass stock on AR land after conversion. However, as the calculated living biomass of total forest did not include loss of biomass stocks on non-forest land immediately before conversion, IEF_{AR} was recalculated by excluding “the loss of biomass”. Since the 2010 inventory submission, in which carbon stock changes on “FF land” and “LF land” were reported collectively, “the loss of biomass” has not been reported in the CRF tables. This submission again took into account the “the loss of biomass”. The gains of living biomass were jointly reported in the gain column for “Rice field converted to Forest land” on the CRF; while the reporting for the losses, which was reported as “IE” in the previous submission, were updated as follows: The losses by land conversion from Orchard and Grassland were reported at the relevant categories in the CRF tables.; notation keys of the loss on Rice field, Upland field and Other land were changed to “NA”; notation keys of Wetlands and Settlements were changed to “NO”.

f) Source-/Sink-specific Planned Improvements

- **Breakdown of the area of “Land converted to Forest Land”**

Although the area of “Land converted to Forest land” reported under the Convention and AR area under the Kyoto Protocol are nearly consistent, there is a discrepancy between them in estimating carbon stock changes. The reason for this is that the estimation of breakdown of the area of “Land

converted to Forest land” to its subcategories is based on statistics in the Convention report, while that for the Kyoto Protocol is based on the AR survey utilizing orthophotos and satellite images. Hence, the validity of the estimation method is presently being reviewed, including a method which does not subcategorize the conversion area.

● **Carbon Stock Changes in Soils in “Cropland and Grassland converted to Forest Land”**

The areas converted to Forest land from upland fields, orchards and pasture land are estimated by multiplying the total areas converted from Cropland to Forest land by each area ratio of upland fields, orchards and pasture land. However, this estimation method may not represent the true status of these areas. Hence, the validity of the estimation method is presently being reviewed.

● **Carbon Stock Changes in Soils in “Other land converted to Forest Land”**

Reporting carbon stock changes in soils in “Other land converted to Forest land” presently continues to be examined with respect to set values and setting methods of carbon stock changes in land before conversion.

7.5. Cropland (5.B)

Cropland is the land that produces annual and perennial crops; it includes temporarily fallow land. Cropland in Japan’s inventory consists of rice fields, upland fields and orchards.

In FY2010, Japan’s Cropland area was about 3.98 million ha, which is equivalent to about 10.5% of the national land. The area of organic soil in Cropland is about 0.18 million ha. The emissions from this category in FY2010 were 452 Gg-CO₂ (excluding 6.2 Gg-CO₂ eq. of N₂O emissions resulting from disturbance associated with land-use conversion to Cropland and 270 Gg-CO₂ of CO₂ emissions resulting from lime application to agricultural soils); this represents a decrease of 82.0% below the FY1990 value and an increase of 75.7% over the FY2009 value.

This section divides cropland into two subcategories, “Cropland remaining Cropland (5.B.1.)” and “Land converted to Cropland (5.B.2.)”, and describes them separately in the following subsections.

Table 7-19 Emissions and removals in Cropland resulting from carbon stock changes

Gas	Category	Carbon pool	Unit	1990	1995	2000	2005	2008	2009	2010	
CO ₂	5.B. Cropland	Total	Gg-CO ₂	2,513.2	822.8	355.7	277.1	224.2	257.5	452.4	
		Living Biomass	Gg-CO ₂	1,298.6	287.4	99.4	129.7	124.3	154.7	309.7	
		Dead Wood	Gg-CO ₂	418.4	85.0	27.2	32.7	29.8	34.4	67.8	
		Litter	Gg-CO ₂	183.7	37.3	11.9	14.4	13.1	16.7	32.9	
		Soil	Gg-CO ₂	612.5	413.1	217.1	100.3	57.0	51.7	42.1	
	5.B.1. Cropland remaining Cropland	Total	Gg-CO ₂	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE
		Living Biomass	Gg-CO ₂	NA	NA	NA	NA	NA	NA	NA	NA
		Dead Wood	Gg-CO ₂	NA	NA	NA	NA	NA	NA	NA	NA
		Litter	Gg-CO ₂	NA	NA	NA	NA	NA	NA	NA	NA
		Soil	Gg-CO ₂	NE	NE	NE	NE	NE	NE	NE	NE
	5.B.2. Land converted to Cropland	Total	Gg-CO ₂		2,513.2	822.8	355.7	277.1	224.2	257.5	452.4
		Living Biomass	Gg-CO ₂		1,298.6	287.4	99.4	129.7	124.3	154.7	309.7
		Dead Wood	Gg-CO ₂		418.4	85.0	27.2	32.7	29.8	34.4	67.8
		Litter	Gg-CO ₂		183.7	37.3	11.9	14.4	13.1	16.7	32.9
		Soil	Gg-CO ₂		612.5	413.1	217.1	100.3	57.0	51.7	42.1

7.5.1. Cropland remaining Cropland (5.B.1)

a) Source/Sink Category Description

This subcategory deals with carbon stock changes in Cropland, which has remained as Cropland during the past 20 years.

With respect to living biomass, the carbon stock change in perennial tree crops (fruit trees) is the subject of estimation according to the *GPG-LULUCF*. However, in Japan, tree growth is limited by trimming in order to have high productivity by keeping the tree height low, and by pruning lateral branches to improve tree shape. Therefore, carbon accumulation because of tree growth cannot be expected, and the annual carbon fixing volume of perennial tree crops in all orchards is stated as “NA.”

Carbon stock changes in dead organic matter are estimated as zero (0) by applying the Tier 1 method, which assumes that the carbon stocks are not changed, according to section 3.3.1.2.1 in the *GPG-LULUCF*. Thus, the carbon stock changes are reported as “NA”.

Carbon stock changes in and CO₂ emissions from soils are presently not estimated due to lack of data for estimation. Hence, this carbon pool is reported as “NE”.

Table 7-20 Areas of “Cropland remaining Cropland” within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Cropland remaining Cropland	kha	4,063.8	4,016.5	3,968.9	3,930.0	3,911.8	3,907.6	3,903.7

b) Source-/Sink-specific Recalculations

None.

c) Source-/Sink-specific Planned Improvements

● Carbon Stock Changes in Soils in “Cropland remaining Cropland”

Research and data collection activities for estimating carbon stock changes in cropland soils have been in progress. Japan is planning to report the carbon stock changes in its future submission when their estimation and reporting become possible.

● CO₂ Emissions from Cultivated Organic Soils in Cropland

The organic soil cropland in Japan is generally developed by admixture of soil and this formation usually makes CO₂ emissions lower than normal CO₂ emissions on cultivated organic soil in cropland. The calculation using the default emission factor in *GPG-LULUCF* is expected to lead overestimation of emissions; however, the actual conditions of CO₂ emissions from organic soils in Cropland subject to admixture of soil in Japan are presently under investigation. Japan is planning to report the relevant emissions in its future submission when their estimation and reporting become possible after the investigation is completed.

7.5.2. Land converted to Cropland (5.B.2)

a) Source/Sink Category Description

This subcategory deals with the carbon stock changes which occurred in the lands that were converted

from other land use categories to Cropland within the past 20 years. The emissions from this subcategory in FY2010 were 452 Gg-CO₂ (excluding 6.2 Gg-CO₂ eq. of N₂O emissions resulting from disturbance associated with land-use conversion to Cropland and 270 Gg-CO₂ of CO₂ emissions resulting from lime application to agricultural soils); this represents a decrease of 82.0% below the FY1990 value and an increase of 75.7% over the FY2009 value.

With respect to living biomass, its carbon stock change as a result of land-use conversion from other land use to Cropland is estimated. This process includes both temporary loss of living biomass in the land before and subsequent gain of living biomass after conversion.

With respect to dead organic matter, Japan used the CENTURY-jfos model to estimate carbon stocks of dead organic matter in Forest land, and then estimated carbon stock changes in “Forest land converted to Cropland”. Carbon stock changes in other subcategories converted to cropland were reported as “NA”, supposing that no carbon stock changes occurred, or “NE” where suitable knowledge for estimating carbon stocks for the land-use categories was not available.

With respect to soil, its carbon stock changes as a result of land-use conversion from other land uses to Cropland are estimated. CO₂ emissions in organic soils were reported as “NE” due to lack of data for estimation.

b) Methodological Issues

1) Carbon stock changes in Living Biomass in “Land converted to Cropland”

● Estimation Method

The Tier 2 method is applied for “Forest land converted to Cropland” using the country specific value of the amount of biomass accumulation. The Tier 1 method is applied for land uses other than “Forest land converted to Cropland” using provisional and default values.

$$\Delta C = \Delta C_i + \Delta C_j$$

$$\Delta C_i = A (CR_a - CR_i) \times CF$$

$$\Delta C_j = A \times CR_j \times CF$$

ΔC : annual carbon stock change in the converted land (t-C/yr)

ΔC_i : annual carbon stock change at the time of land conversion (t-C/yr)

ΔC_j : annual carbon stock change in the converted land after conversion (t-C/yr)

i : land use before conversion

j : land use after conversion

A : area of converted land for the current year (ha)

CR_a : dry matter biomass weight per unit area immediately following conversion (t-d.m./ha), default value=0

CR_i : dry matter biomass weight per unit area before land was converted from land-use type i (t-d.m./ha)

CR_j : change of dry matter biomass weight per unit area accumulated after conversion (t-d.m./ha/yr)

CF : carbon fraction of dry matter (t-C/t-d.m.)

● Parameters

➤ Biomass stock in each Land-Use Category

The values shown in Table 7-6 are used for the estimation of biomass stock changes upon land-use conversion and subsequent changes in biomass stock due to biomass growth in the converted land.

➤ **Carbon Fraction of Dry Matter**

0.5 (t-C/t-d.m.) (GPG-LULUCF, default value)

● **Activity Data (Area)**

Annually converted areas to Cropland were used for estimating carbon stock changes in living biomass in “Land converted to Cropland”.

It was assumed that the areas of “Forest land converted to other land use” (Cropland, Grassland, Wetlands, Settlement and Other land) were consistent with the area of deforestation (D area) reported under Article 3, paragraph 3, of the Kyoto Protocol. Thus, the area of “Forest land converted to Cropland” was estimated by allocating the D area. Since the D survey has been conducted since FY2005, the applied method to calculate the D area for FY1990 to FY2004 and for post FY2005 are as follows, respectively.

➤ **From FY1990 to FY2004**

The areas of “Forest land converted to Cropland, Grassland, Settlements and Other land” are estimated by multiplying the areas, which are calculated by subtracting the area of “Forest land converted to Wetlands” from the total areas converted from Forest land, by the land ratios of “Forest land converted to Cropland, Grassland, Settlements and Other land”, respectively.

The total area converted from Forest land was determined based on the areas provided by the *World Census of Agriculture and Forestry*, the Forestry Agency’s records, and D areas under Article 3, paragraph 3, of the Kyoto Protocol. In concrete terms, the D areas are identified in detail by utilizing orthophotos taken at the end of 1989 and recent satellite images, but they are provided only for the FY1990 values and onward. Therefore, the total areas converted from Forest land are estimated by setting an adjustment factor from the ratio between the D areas since FY1990 and the areas converted from forests provided by the *World Census of Agriculture and Forestry* and the Forestry Agency’s records, and multiplying the areas converted from forests since FY1970 by the adjustment factor. For further information on determining the D areas, see section 11.3.2.3 in Chapter 11.

The respective ratios of Forest land converted to other land-use categories except Wetlands are estimated from areas of private forests converted to other land-use categories resulting from forest land development, based on the Forestry Agency’s records, and the ratios are regarded as the same as those for national forests.

➤ **After FY2005**

The areas of “Forest land converted to Cropland, Grassland, Wetlands, Settlements and Other land” were estimated by multiplying the D area by the land ratios of Forest land converted to each land-use category. Both the ratio and the area were determined by the D survey.

The areas of land converted from land-use categories other than Forest land to Cropland are determined by applying expansion area values provided by the *Statistics of Cultivated and Planted Area*. The converted areas are divided into rice fields, upland fields, orchards, and pasture land proportionately by means of the current area ratios. The areas of rice fields, upland fields, and orchards are allocated to Cropland, while the area of pasture land is allocated to Grassland.

It should be noted that the area presented in the CRF “Table 5.B SECTORAL BACKGROUND DATA FOR LAND USE, LAND-USE CHANGE AND FORESTRY – Cropland” is not the annually converted area in FY2010 but the sum of annually converted areas during the past 20 years.

Table 7-21 Area of land converted to Cropland (single year)

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Cropland	kha	8.6	4.4	3.1	2.0	1.5	1.1	1.3
Forest land converted to Cropland	kha	7.0	1.4	0.5	0.5	0.5	0.6	1.2
Rice field	kha	0.011	0.018	0.002	0.000	0.045	0.023	0.120
Upland field	kha	7.0	1.4	0.5	0.5	0.5	0.6	1.1
Orchard	kha	IE	IE	IE	IE	IE	IE	IE
Grassland converted to Cropland	kha	0.002	0.022	0.012	0.027	0.005	0.004	0.0003
Wetlands converted to Cropland	kha	0.34	0.03	0.07	0	0.47	0	0
Settlements converted to Cropland	kha	IE	IE	IE	IE	IE	IE	IE
Other land converted to Cropland	kha	1.3	2.9	2.5	1.4	0.5	0.5	0.04
Rice field	kha	0.2	1.1	1.3	0.3	0.1	0.1	0.03
Upland field	kha	1.1	1.9	1.2	1.1	0.4	0.5	0.01
Orchard	kha	IE	IE	IE	IE	IE	IE	IE

2) Carbon Stock Change in Dead Organic Matter in “Land converted to Cropland”

● Estimation Method

Carbon stock changes in dead organic matter in “Forest land converted to Cropland” were estimated by applying the Tier 2 estimation method using the value of carbon stock in dead organic matter obtained by the CENTURY-jfos model. All carbon stocks in dead organic matter in the subcategory are assumed oxidized and emitted as CO₂ within the year of conversion in accordance with the description in section 3.4.2.2.1 in the *GPG-LULUCF*. In addition, as described in the Parameters section below, carbon stocks of dead organic matter in Cropland are assumed to be zero.

$$\Delta C_{DOM} = \sum ((C_{after,i} - C_{before,i}) \times A)$$

ΔC_{DOM} : Carbon stock changes in dead organic matter in the converted land (t-C/yr)

$C_{after,i}$: Average carbon stock per unit area in dead wood or litter after conversion (t-C/ha)

Note: carbon stocks after conversion are assumed as “0” (zero).

$C_{before,i}$: Average carbon stock per unit area in dead wood or litter before conversion (t-C/ha)

A : Area of converted land within the year of conversion (ha)

i : type of dead organic matter (dead wood or litter)

With regard to “Grassland converted to Cropland”, carbon stocks of dead wood and litter carbon pools were assumed to be minor and the stock changes could be ignored, and were thus reported as “NA”. With regard to “Wetlands and Settlements converted to Cropland”, these were also reported as “NA”, since carbon stock changes were assumed as zero to zero, supposing that basically no such carbon pools exist in reclaimed wetland and that carbon stocks of dead organic matter in Settlements before conversion were assumed as negligible. “Other land converted to Cropland”, which is estimated to be cropland restoration, was reported as “NE”, because suitable knowledge for estimating carbon stock changes for this land-use conversion was not available.

● Parameters

Average carbon stocks in dead wood and litter in Forest land before conversion are shown in Tables 7-6 and 7-7. In addition, it is assumed that they become zero immediately after conversion, and will not accumulate after conversion.

● Activity Data (Area)

Annually converted areas to Cropland are used for estimating carbon stock changes in dead organic matter in “Land converted to Cropland”.

3) Carbon Stock Changes in Soils in “Land converted to Cropland”

● Estimation Method

Carbon stock changes in soils were calculated by applying the Tier 2 estimation method in accordance with the estimation method for “Land converted to Cropland” (GPG-LULUCF, page 3-89) using the values of carbon stock in soils which are country specific or obtained by the CENTURY-jfos model.

$$\Delta C_i = A_i \times (C_{after,i} - C_{before,i}) / 20$$

ΔC_i : Annual change in carbon stocks in soils in converted land (t-C/yr)

A_i : Area converted to land-use category i within the past 20 years (ha)

$C_{after,i}$: Average carbon stocks per unit area in land-use category i after conversion (t-C/ha)

$C_{before,i}$: Average carbon stocks per unit area in land-use category i before conversion (t-C/ha)

i : Land-use category

● Parameters

Average soil carbon stocks in each land-use category shown in Table 7-9 were used. For reference, soil carbon stocks of Cropland (rice field, upland field and orchard) are described below in detail:

➤ Soil carbon stocks in Rice field, Upland field and Orchard

For the carbon stocks in rice fields, upland fields and orchard soils, the country-specific soil survey data were applied. As soil carbon stocks per unit area vary from one soil group to another (such as Andosols, Gray lowland soils and Gley soils), the average soil carbon stocks in rice field, upland field and orchard are calculated by averaging the soil carbon stock data per unit area at 0-30 cm depth weighted by the area for each soil group.

Table 7-22 Soil carbon stocks in Rice field

Soil Type	Area [ha]	Proportion	Carbon Stock / ha [t-C/ha]	Carbon Stock [t-C]
Lithosols	*	---	*	---
Sand-Dune Regosols	*	---	89.04	---
Andisols	17,169	0.6%	125.24	2,150,246
Wet Andosols	274,319	9.5%	113.68	31,184,584
Gleyed Andosols	50,760	1.8%	101.74	5,164,322
Cambisols	6,640	0.2%	59.48	394,947
Gray Upland Soils	79,236	2.7%	60.37	4,783,477
Gley Upland Soils	40,227	1.4%	60.71	2,442,181
Red Soils	*	---	*	---
Yellow Soils	144,304	5.0%	63.21	9,121,456
Dark Red Soils	1,770	0.1%	56.26	99,580
Fluvisols	141,813	4.9%	59.71	8,467,654
Gleysols	1,056,571	36.6%	61.59	65,074,208
Gleysols	889,199	30.8%	64.83	57,646,771
Muck Soils	75,944	2.6%	91.89	6,978,494
Histosols	109,465	3.8%	114.95	12,583,002
Total	2,887,417	100.0%		206,090,923
Average			80.19	
Weighted Average			71.38	Applied Value

*: Data difficult to obtain with high accuracy.

Table 7-23 Soil carbon stocks in Upland field

Soil Type	Area [ha]	Proportion	Carbon Stock / ha [t-C/ha]	Carbon Stock [t-C]
Lithosols	7,148	0.4%	69.25	494,999
Sand-Dune Regosols	22,297	1.2%	21.49	479,163
Andisols	851,061	46.5%	109.15	92,893,308
Wet Andosols	72,195	3.9%	149.51	10,793,874
Gleyed Andosols	1,850	0.1%	120.98	223,813
Cambisols	287,464	15.7%	65.16	18,731,154
Gray Upland Soils	71,855	3.9%	79.77	5,731,873
Gley Upland Soils	4,324	0.2%	*	---
Red Soils	25,243	1.4%	42.23	1,066,012
Yellow Soils	105,641	5.8%	47.13	4,978,860
Dark Red Soils	29,130	1.6%	45.15	1,315,220
Fluvisols	231,051	12.6%	50.05	11,564,103
Gleysols	75,095	4.1%	53.75	4,036,356
Gleysols	13,163	0.7%	65.94	867,968
Muck Soils	1,673	0.1%	78.72	131,699
Histosols	32,316	1.8%	184.91	5,975,552
Total	1,831,506	100.0%		159,283,954
Average			78.88	
Weighted Average			86.97	Applied Value

*: Data t difficult to obtain with high-accuracy.

Table 7-24 Soil carbon stocks in Orchard

Soil Type	Area [ha]	Proportion	Carbon Stock / ha [t-C/ha]	Carbon Stock [t-C]
Lithosols	7,682	1.9%	66.48	510,699
Sand-Dune Regosols	1,897	0.5%	27.77	52,680
Andisols	86,083	21.3%	119.03	10,246,459
Wet Andosols	2,530	0.6%	103.82	262,665
Gleyed Andosols	*	---	115.08	---
Cambisols	148,973	36.9%	68.35	10,182,305
Gray Upland Soils	6,424	1.6%	70.55	453,213
Gley Upland Soils	*	---	*	---
Red Soils	19,937	4.9%	63.68	1,269,588
Yellow Soils	75,973	18.8%	64.48	4,898,739
Dark Red Soils	6,141	1.5%	54.61	335,360
Fluvisols	35,261	8.7%	69.32	2,444,293
Gleysols	10,075	2.5%	57.35	577,801
Gleysols	2,065	0.5%	*	---
Muck Soils	135	0.0%	59.44	8,024
Histosols	130	0.0%	*	---
Total	403,306	100.0%		31,241,826
Average			72.30	
Weighted Average			77.46	Applied Value

*: Data difficult to obtain with high accuracy.

● Activity Data (Area)

Areas of “Land converted to Cropland” during the past 20 years are assumed as the sum of the areas of annually converted land to Cropland during the past 20 years. The assumed areas are applied to the estimation of the carbon stock changes in soils in “Land converted to Cropland”. The areas are shown in Table 7-25 below.

Table 7-25 Area of land converted to Cropland within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Cropland	kha	493.1	319.7	183.7	108.5	74.8	65.3	58.0
Forest land converted to Cropland	kha	279.3	203.0	120.6	54.6	35.5	28.0	22.2
Rice field	kha	279.3	203.0	120.6	54.6	35.5	28.0	22.2
Upland field	kha	IE	IE	IE	IE	IE	IE	IE
Orchard	kha	IE	IE	IE	IE	IE	IE	IE
Grassland converted to Cropland	kha	8.6	4.8	0.6	0.5	0.5	0.5	0.5
Wetlands converted to Cropland	kha	11.9	3.9	2.0	1.2	1.2	1.1	0.7
Settlements converted to Cropland	kha	IE	IE	IE	IE	IE	IE	IE
Other land converted to Cropland	kha	193.3	108.0	60.4	52.2	37.6	35.8	34.5
Rice field	kha	27.7	16.2	11.2	9.9	11.6	11.5	11.3
Upland field	kha	165.6	91.8	49.2	42.3	26.0	24.3	23.2
Orchard	kha	IE	IE	IE	IE	IE	IE	IE

c) Uncertainties and Time-series Consistency

● Uncertainty Assessment

Uncertainties of the parameters and the activity data for living biomass, dead organic matter, and soil were individually assessed on the basis of field study results, expert judgment, or the default values described in the *GPG-LULUCF*. The uncertainty was estimated as 28% for the entire emission from the “Land converted to Cropland”. More detailed information on the uncertainty assessment is described in Annex 7. Uncertainty estimates of some major parameters, which were used for the uncertainty assessment for this category, are shown in Table 7-26 as an example.

Table 7-26 Uncertainty estimates regarding major parameters in the Cropland category

Land-use category		Uncertainty (%)	Country Specific (CS) or Default (D)	Note
Cropland Area	Rice Field	0.15	CS	Original uncertainty of statistics
	Upland Field	0.27	CS	

● Time-series Consistency

Although the methods to estimate the area of “Forest land converted to other land-use” are different between FY1990-2004 and post FY2005 as described in section 7.5.2.b)1), time-series consistency for this subcategory is basically ensured.

d) Source-/Sink-specific QA/QC and Verification

QC is implemented in accordance with the Tier 1 approach described in the GPG (2000) and the *GPG-LULUCF*. The Tier 1 approach includes procedures for checking parameters and activity data, and for archiving references. More detailed information on the QA/QC activity procedures is described in section 6.1 of Annex 6.

e) Source-/Sink-specific Recalculations

● Carbon stock change in soils of “Other land converted to Cropland” in FY 2008

Due to error correction of the estimated area of Other land converted to Cropland in FY2008, carbon stock change in soils was recalculated.

● Carbon Stock Changes in Forest land converted to Cropland

Living biomass accumulation in Forest land subjected to conversion has been estimated by extrapolation of the trend of living biomass accumulation in the D area during the period between 2005 and the latest year. In line with this, the carbon stock changes were recalculated in accordance with the updated living biomass accumulation reflecting the value of FY2010.

f) Source-/Sink-specific Planned Improvements

● Methods of Obtaining Data of the area of “Grassland converted to Cropland”

Data on the area of land converted from Grassland to Cropland other than "Land converted from Grassland (pasture land) to Cropland (rice field)" cannot be obtained from currently available statistics, so the carbon stock changes in these areas have not been estimated. Therefore, the methods of obtaining the following area data of conversion need to be investigated.

- from pasture land to upland field
- from pasture land to orchard
- from grazed meadow to rice field
- from grazed meadow to upland field
- from grazed meadow to orchard

● Estimation Method of Soil Carbon Stock Change upon “Land converted to Cropland”

The estimation method will be considered when new data and information are obtained.

7.6. Grassland (5.C)

Grassland is generally covered with perennial pasture and is used mainly for harvesting fodder or grazing. In FY2010, Japan’s grassland area was about 0.99 million ha, which is equivalent to about 2.6% of the national land. The area of organic soil in the Grassland is about 0.04 million ha. The net removals from this category in FY2010 were 216 Gg-CO₂ (excluding 270 Gg-CO₂ of CO₂ emissions resulting from lime application to agricultural soils); this represents a decrease of 51.4% below the FY1990 value and a decrease of 21.9% below the FY2009 value.

This section divides grassland into two subcategories, “Grassland remaining Grassland (5.C.1.)” and “Land converted to Grassland (5.C.2.)”, and describes them separately in the following subsections.

Table 7-27 Emissions and removals from Grassland resulting from carbon stock changes

Gas	Category	Carbon pool	Unit	1990	1995	2000	2005	2008	2009	2010	
CO ₂	5.C. Grassland	Total	Gg-CO ₂	-444.0	-481.1	-405.8	-335.6	-302.6	-276.2	-215.9	
		Living Biomass	Gg-CO ₂	48.0	-17.2	-26.6	-28.4	-31.2	-21.4	17.9	
		Dead Wood	Gg-CO ₂	58.9	12.8	4.2	4.5	4.0	4.6	11.8	
		Litter	Gg-CO ₂	25.8	5.6	1.8	2.0	1.7	2.2	5.7	
		Soil	Gg-CO ₂	-576.7	-482.4	-385.3	-313.7	-277.1	-261.7	-251.3	
	5.C.1. Grassland remaining Grassland	Total	Gg-CO ₂	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE
		Living Biomass	Gg-CO ₂	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE
		Dead Wood	Gg-CO ₂	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE
		Litter	Gg-CO ₂	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE
		Soil	Gg-CO ₂	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE	NA,NE
	5.C.2. Land converted to Grassland	Total	Gg-CO ₂	-444.0	-481.1	-405.8	-335.6	-302.6	-276.2	-215.9	
		Living Biomass	Gg-CO ₂	48.0	-17.2	-26.6	-28.4	-31.2	-21.4	17.9	
		Dead Wood	Gg-CO ₂	58.9	12.8	4.2	4.5	4.0	4.6	11.8	
		Litter	Gg-CO ₂	25.8	5.6	1.8	2.0	1.7	2.2	5.7	
		Soil	Gg-CO ₂	-576.7	-482.4	-385.3	-313.7	-277.1	-261.7	-251.3	

7.6.1. Grassland remaining Grassland (5.C.1)

a) Source/Sink Category Description

In this category carbon stock changes in “Grassland remaining Grassland” during the past 20 years are reported, divided into three subcategories: “pasture land”, “grazed meadow” and “wild land”.

With respect to living biomass, carbon stock changes in pasture land and grazed meadow are assumed to be in a steady state and reported as “NA” in accordance with the Tier 1 estimation method in section 3.4.1.1.1 in the *GPG-LULUCF*. Carbon stock changes in living biomass in wild land are reported as “NE” because the status of carbon pools in wild land is under survey.

Carbon stock changes in dead organic matter in pasture land and grazed meadow are estimated as zero (0) by applying the Tier 1 method described in section 3.4.1.2.1 in the *GPG-LULUCF*, which assumes that the carbon stocks are not changed. Thus, the carbon stock changes are reported as “NA”. Carbon stock changes in dead organic matter in wild land are reported as “NE” because the status of carbon pools in wild land is under survey.

With respect to soil, carbon stock changes in soil in pasture land are presently not estimated because information on carbon stocks and management state in the pasture land is not collected sufficiently for estimating carbon stock changes. Hence, this carbon pool is reported as “NE”. On the other hand, grazed meadows are non-degraded and sustainably managed grassland, but without significant management improvements. Therefore, the default value of the carbon stock change factor for “Nominally managed (non-degraded)” in table 3.4.5 of the *GPG-LULUCF*, which is “1.0”, is applied to grazed meadows. In this case, soil carbon stocks are not changed over time; therefore, the soil carbon stock changes in grazed meadows are reported as “NA”. The carbon stock changes in soil in wild land are reported as “NE” because the actual condition of the carbon stock changes is not clear. CO₂ emissions from organic soils are reported as “NE” because the estimation procedure of the emissions is under examination.

Table 7-28 Areas of “Grassland remaining Grassland” within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Grassland remaining Grassland	kha	869.2	898.1	924.6	945.9	950.3	951.3	952.8
Pasture land	kha	494.2	537.2	557.7	569.4	573.8	574.9	576.4
Grazed meadow	kha	105.0	100.9	96.8	96.5	96.4	96.4	96.4
Wild land	kha	270.0	260.0	270.0	280.0	280.0	280.0	280.0

b) Source-/Sink-specific Recalculations

● *Change of the area of Grazed meadow*

The area of Grazed meadow was recalculated due to the re-examinations of statistical data interpreted as Grazed meadow and the way to reconstruct time series data due to the elimination of the survey item of relevant data from the statistics.

c) Source-/Sink-specific Planned Improvements

● *Carbon Stock Changes in Mineral Soils in “Grassland remaining Grassland”*

Carbon stock changes in mineral soils in this category are currently not estimated. However, research projects on soil carbon stocks in pasture land are in progress. Therefore, Japan is planning to report the carbon stock changes when data will be available for estimation in the future.

● *CO₂ Emissions from Cultivated Organic Soils in Grassland*

With respect to CO₂ emissions from organic soils in Grassland, CO₂ emissions from organic soils are being examined in a cross-cutting manner through the LULUCF sector, including the emissions from Cropland.

7.6.2. Land converted to Grassland (5.C.2)

a) Source/Sink Category Description

This subcategory deals with the carbon stock changes, which occurred in the lands that were converted from other land-use categories to grassland within the past 20 years. The net removal from this subcategory in FY2010 was 216 Gg-CO₂ (excluding 270 Gg-CO₂ of CO₂ emissions resulting from lime application to agricultural soils); this represents a decrease of 51.4% below the FY1990 value and a decrease of 21.9% below the FY2009 value.

With respect to living biomass, its carbon stock changes as a result of land-use conversion from other land use to Grassland are estimated. The carbon stock changes include both temporary loss of living biomass in the land before and subsequent gain after conversion.

With respect to dead organic matter, Japan used the CENTURY-jfos model to estimate carbon stocks in dead organic matter in Forest land, and then estimated carbon stock changes in “Forest land converted to Grassland”. Carbon stock changes in Grassland converted from land-uses other than Forest land were reported as “NE” or “NA” because suitable knowledge for estimating carbon stocks for the land-use categories was not available, or because it was assumed that no carbon stock change occurred, respectively.

Carbon stock changes in soils as a result of land-use conversion from other land use to Grassland are estimated. All soils are temporarily regarded as mineral soils because the actual condition of organic soils is presently being assessed.

b) Methodological Issues

1) Carbon stock changes in Living biomass in “Land converted to Grassland”

● Estimation Method

The Tier 2 method was applied to estimate “Forest land and Cropland (rice fields) converted to Grassland (pasture lands)” using country specific and provisional values of the amount of biomass accumulation. The Tier 1 method was used for “Land uses other than Forest land and Cropland (rice fields) converted to Grassland (pasture lands)” using default value. The equations are given in section 7.5.2.b)1). While the annually converted areas were used for estimating the loss of living biomass upon land-use conversion, the biomass growth after land-use conversion was estimated by summing the converted areas for the latest five years, assuming that biomass growth reaches a steady state at a constant rate over the subsequent five years after conversion.

● Parameters

➤ Biomass stock in each Land-Use Category

The values shown in Table 7-6 are used for the estimation of biomass stock changes upon land-use conversion and subsequent changes in biomass stock due to biomass growth in converted land.

➤ Carbon Fraction of Dry Matter

0.5 (t-C/t-d.m.) (GPG-LULUCF, default value)

● Activity Data (Area)

As shown in Table 7-3, Grassland is treated as a part of arable land in statistics of Japan. Therefore, the procedure to obtain the area of the “Grassland converted from other land-use categories” is as

described in 7.5.2.b)1).

It should be noted that the area presented in the CRF “Table 5.C SECTORAL BACKGROUND DATA FOR LAND USE, LAND-USE CHANGE AND FORESTRY – Grassland” is not the annually converted area in FY2010 but the sum of annually converted areas during the past 20 years.

Table 7-29 Area of land converted to Grassland (single year)

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Grassland	kha	4.0	1.9	1.7	2.4	1.4	1.1	1.2
Forest land converted to Grassland	kha	1.0	0.2	0.1	0.1	0.1	0.1	0.2
Cropland converted to Grassland	kha	0.9	0.6	1.0	1.7	0.8	0.7	0.7
Wetlands converted to Grassland	kha	0.12	0.01	0.03	0	0.20	0	0
Settlements converted to Grassland	kha	IE	IE	IE	IE	IE	IE	IE
Other land converted to Grassland	kha	2.0	1.1	0.6	0.6	0.3	0.4	0.3

Table 7-30 Area of land converted to Grassland within the past 5 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Grassland	kha	27.2	12.2	8.4	9.3	9.6	8.5	7.2
Forest land converted to Grassland	kha	4.9	1.8	0.6	0.3	0.3	0.3	0.5
Cropland converted to Grassland	kha	6.5	3.4	4.5	6.2	6.4	5.7	4.6
Wetlands converted to Grassland	kha	0.32	0.07	0.03	0	0.20	0.20	0.20
Settlements converted to Grassland	kha	IE	IE	IE	IE	IE	IE	IE
Other land converted to Grassland	kha	15.5	6.9	3.3	2.8	2.7	2.3	1.9

2) Carbon Stock Change in Dead Organic Matter and Soils in “Land converted to Grassland”

● Estimation Method

➤ Carbon Stock Changes in Dead Organic Matter

In this category, carbon stock changes in dead organic matter in “Forest land converted to Grassland” were estimated. The Tier 2 estimation method was applied to the subcategory using country specific values of the carbon stocks before and after conversion. It should be noted that the carbon stocks of dead organic matter after conversion to Grassland are assumed as zero (Tier 1 method in 2006 IPCC Guideline Vol.4 section 6.3.2), because there are no quantitative data of them, although a subtle but certain amount of carbon stocks does generally exist on the soil surface. As described in section 7.5.2.b)2), “Cropland and Settlements converted to Grassland” were reported as “NA” since the carbon stocks before and after conversion were assumed as zero. As for “Wetlands and Other land converted to Grassland”, they are estimated to be reclamation and restoration. Thus they were reported as “NA” and “NE”, respectively⁶, for similar reasons as described in section 7.5.2.b)2).

➤ Carbon Stock Changes in Soils

Carbon stock changes in soils were estimated as described in section 7.5.2.b)3). In addition, organic soils were reported as “NE”.

● Parameters

➤ Carbon Stocks in Dead Organic Matter

The average carbon stocks in dead wood and litter in Forest land before conversion are shown in Tables 7-6 and 7-7. The average carbon stocks in these categories from FY1990 to FY2004 are not estimated; therefore those in FY2005 are substituted for them. In addition, it is assumed that they become zero immediately after conversion, and are not accumulated after conversion. All carbon

⁶ Cropland in the Japanese statistics includes Pasture land which falls into Grassland.

stocks in dead organic matter in the subcategory are assumed oxidized and emitted as CO₂ within the year of conversion in accordance with the description in section 3.4.2.2.1 in the *GPG-LULUCF*.

➤ **Carbon Stocks in Soils**

Data listed in Table 7-9 are applied as average carbon stocks before and after conversion. For reference, soil carbon stocks of Grassland are described in detail below.

➤ **Soil carbon stocks in Grassland**

Data from the country-specific soil survey are applied for the carbon stocks in Grassland soils. Although it is difficult to obtain area data by soil types for Grassland, it could be viewed that the area by soil types and the number of samples by soil types have a high correlation; therefore, the area data are calculated by averaging and weighting the soil carbon stock data by the number of samples for each soil group.

Table 7-31 Soil carbon stocks in Grassland

Soil Type	Area [ha]	Proportion	Carbon Stock / ha [t-C/ha]	Carbon Stock [t-C]
Lithosols	*	---	*	---
Sand-Dune Regosols	140	0.6%	79.28	11,099
Andisols	11,364	48.8%	152.19	1,729,487
Wet Andosols	459	2.0%	207.40	95,197
Gleyed Andosols	*	---	*	---
Cambisols	4,071	17.5%	101.27	412,270
Gray Upland Soils	2,008	8.6%	126.44	253,892
Gley Upland Soils	228	1.0%	110.51	25,196
Red Soils	*	---	*	---
Yellow Soils	796	3.4%	74.36	59,191
Dark Red Soils	695	3.0%	54.55	37,912
Fluvisols	2,658	11.4%	107.69	286,240
Gleysols	215	0.9%	78.76	16,933
Gleysols	*	---	*	---
Muck Soils	*	---	*	---
Histosols	663	2.8%	325.18	215,594
Total	23,297	100.0%		3,143,012
Average			128.88	
Weighted Average			134.91	Applied Value

*: Data difficult to obtain with high accuracy.

● **Activity Data (Area)**

The sum of annually converted areas from other land-use categories to Grassland for the past 20 years was regarded as the area of “Land converted to Grassland” during the past 20 years. The areas are shown in Table 7-32.

Table 7-32 Areas of land converted to Grassland within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Grassland	kha	144.8	114.3	80.0	57.2	43.8	40.1	37.2
Forest land converted to Grassland	kha	30.6	25.3	16.5	7.7	5.1	4.0	3.2
Cropland converted to Grassland	kha	25.2	21.2	19.8	20.7	19.4	19.0	18.8
Wetlands converted to Grassland	kha	0.8	0.9	0.7	0.4	0.5	0.4	0.3
Settlements converted to Grassland	kha	IE	IE	IE	IE	IE	IE	IE
Other land converted to Grassland	kha	88.1	67.0	43.0	28.4	18.8	16.6	14.9

c) Uncertainties and Time-series Consistency● **Uncertainty Assessment**

Uncertainties of the parameters and the activity data for living biomass, dead organic matter, and soil were individually assessed on the basis of field study results, expert judgment, or the default values described in the *GPG-LULUCF*. The uncertainty was estimated as 47% for the entire removal from the “Land converted to grassland”. More detailed information on the uncertainty assessment is described in Annex 7. In addition, concrete uncertainty estimates for individual parameters in this category will be calculated in future submissions after investigation is completed.

● **Time-series Consistency**

Although the methods to estimate the area of “Forest land converted to other land use” are different between FY1990-2004 and post FY2005, as described in section 7.5.2.b)1), time-series consistency for this subcategory is basically ensured.

d) Source-/Sink-specific QA/QC and Verification

QC is implemented in accordance with the Tier 1 approach described in GPG (2000) and the *GPG-LULUCF*. The Tier 1 approach includes procedures for checking parameters and activity data, and for archiving references. More detailed information on the QA/QC activity procedures is described in section 6.1 of Annex 6.

e) Source-/Sink-specific Recalculations● **Carbon Stock Changes in Forest land converted to Grassland**

Living biomass accumulation in Forest land subjected to conversion has been estimated by extrapolation of the trend of living biomass accumulation in the D area during the period between 2005 and the latest year. In line with this, the carbon stock changes were recalculated in accordance with the updated living biomass accumulation reflecting the value of FY2010.

f) Source-/Sink-specific Planned Improvements● **Method of Obtaining Data of the “Areas converted from Other Land-use Categories to Grassland”**

The method used to obtain data on the area converted to Grassland needs to be improved. For example, currently, the area of lands converted from Forest land to Grassland is estimated by multiplying the summed areas of “Forest land converted to Cropland and Grassland” by the ratio of grazing land to the summed area. However, this estimation method may not represent the actual status of these areas. Therefore, the validity of the estimation method needs to be reviewed, and, if necessary, a new method of obtaining the area data should be considered.

● **Method of Obtaining Data of the “Area converted from Cropland to Grassland”**

With respect to the method of obtaining data of the area converted from Cropland to Grassland, the converted area cannot be obtained from statistics except for the land-use conversion from Cropland (rice field) to Grassland (pasture land). For this reason, the estimates of the carbon stock changes in this land-use category may not fully reflect the actual conditions. Therefore, the methods used to obtain the following area data need to be investigated.

- from upland field to pasture land
- from orchard to pasture land
- from rice field to grazed meadow

- from upland field to grazed meadow
- from orchard to grazed meadow.

● **Estimation Method of Soil Carbon Stock Change upon “Land-Use Conversion from Other Land to Cropland”**

The estimation method will be considered when new data and information are obtained.

● **Method of Obtaining Data and Revising Estimation Methodologies for Living Biomass Stock in “Grassland other than Pasture Land and grazed Meadow Land”**

It was pointed out by experts that the living biomass stock of the “Grassland other than pasture land and grazed meadow land” is not necessarily identical to the one of “pasture land and grazed meadow land”, which were originally classified in Grassland. Therefore, it is necessary to obtain data which reflect the living biomass stock in grassland other than pasture land and grazed meadow land, and to revise the estimation method for that accordingly.

7.7. Wetlands (5.D)

Wetlands are lands that are covered with or soaked in water throughout the year. They do not fall under the categories of Forest land, Cropland, Grassland, or Settlements”. The *GPG-LULUCF* divides Wetlands into two large groups: peat land and flooded land.

In FY2010, Japan’s wetland area was about 1.33 million ha, which is equivalent to about 3.5% of the national land. The emissions from this category in FY2010 were 82.1 Gg-CO₂; this represents a decrease of 4.3% below the FY1990 value and an increase of 263.8% over the FY2009 value.

This section divides Wetlands into two subcategories, “Wetlands remaining Wetlands (5.D.1.)” and “Land converted to Wetlands (5.D.2.)”, and describes them separately in the following subsections.

Table 7-33 Emissions and removals in Wetlands resulting from carbon stock changes

Gas	Category	Carbon pool	Unit	1990	1995	2000	2005	2008	2009	2010	
CO ₂	5.D. Wetlands	Total	Gg-CO ₂	85.8	360.4	451.4	15.6	15.7	22.6	82.1	
		Living Biomass	Gg-CO ₂	58.8	253.2	324.0	11.5	11.7	17.0	62.0	
		Dead Wood	Gg-CO ₂	18.8	74.5	88.6	2.9	2.8	3.8	13.5	
		Litter	Gg-CO ₂	8.3	32.7	38.9	1.3	1.2	1.8	6.6	
		Soil	Gg-CO ₂	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	
	5.D.1. Wetlands remaining Wetlands	Total	Gg-CO ₂	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE
		Living Biomass	Gg-CO ₂	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE
		Dead Wood	Gg-CO ₂	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE
		Litter	Gg-CO ₂	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE
		Soil	Gg-CO ₂	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE
	5.D.2. Land converted to Wetlands	Total	Gg-CO ₂		85.8	360.4	451.4	15.6	15.7	22.6	82.1
		Living Biomass	Gg-CO ₂		58.8	253.2	324.0	11.5	11.7	17.0	62.0
		Dead Wood	Gg-CO ₂		18.8	74.5	88.6	2.9	2.8	3.8	13.5
		Litter	Gg-CO ₂		8.3	32.7	38.9	1.3	1.2	1.8	6.6
		Soil	Gg-CO ₂	NE	NE	NE	NE	NE	NE	NE	NE

7.7.1. Wetlands remaining Wetlands (5.D.1)

a) Source/Sink Category Description

This subcategory deals with carbon stock changes in Wetlands which have remained as Wetlands during the past 20 years.

Carbon stock changes in organic soils that are managed for peat extraction are reported as “NO”, since peat extraction is not carried out in Japan. (Default value for Japan is not provided in the *GPG-LULUCF* p.3.282 Table 3A3.3).

“Flooded land remaining flooded land” is not calculated at the present time as this will be treated in an appendix and reported as “NE”.

Table 7-34 Areas of “Wetlands remaining Wetlands” within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Wetlands remaining Wetlands	kha	1,236.3	1,257.7	1,285.4	1,299.0	1,301.5	1,301.9	1,302.2
Organic soils managed for peat extraction	kha	NO	NO	NO	NO	NO	NO	NO
Flooded land	kha	1,236.3	1,257.7	1,285.4	1,299.0	1,301.5	1,301.9	1,302.2

7.7.2. Land converted to Wetlands (5.D.2)

a) Source/Sink Category Description

This subcategory deals with the carbon stock changes which occurred in the land that was converted from other land-use categories to Wetlands, particularly to flooded land (i.e., dams) within the past 20 years. The emissions from this subcategory in FY2010 were 82 Gg-CO₂; this represents a decrease of 4.3% below the FY1990 value and an increase of 263.8% over the FY2009 value.

With respect to living biomass, its carbon stock change as a result of land-use conversion from other land use to Wetlands is estimated. This process includes both temporary loss of living biomass in the land before and subsequent gain after conversion.

With respect to dead organic matter, Japan used the CENTURY-jfos model to estimate carbon stocks in dead organic matter in Forest land, and then estimated the carbon stock change in “Wetlands converted from Forest land”. Carbon stock changes in other subcategories were reported as “NA”, supposing that no carbon stock change occur, or “NE” where suitable knowledge for estimating carbon stocks for the land-use categories were not available.

Carbon stock changes in soils in “Land converted to Wetlands” were not estimated due to lack of data. Therefore, the carbon stock changes in the carbon pool were reported as “NE”.

b) Methodological Issues

1) Carbon stock change in Living biomass in “Land converted to Wetlands”

● Estimation Method

The Tier 2 method was applied for the “Land converted to Wetlands (flooded land)”. The equations are given in section 7.5.2.b)1).

● Parameters

➤ Biomass stock in each Land-Use Category

The values shown in Table 7-6 are used for the estimation of biomass stock changes resulting from land-use conversion and subsequent changes in biomass stock due to biomass growth in converted land.

➤ Carbon Fraction of Dry Matter

0.5 (t-C/t-d.m.) (*GPG-LULUCF*, default value)

● **Activity Data (Area)**

The area converted from other land use to Wetlands (dam) was estimated based on the area of dam converted from Forest land and the ratio of Forest land among the area of other land-use categories before conversion. The area for “Forest land converted to Wetlands” was calculated by the method described in section 7.5.2.b)1). As for other land-use categories, the area of converted Cropland is divided proportionately into Cropland and grassland according to the current area ratios of land-use categories. After deducting the areas converted from Forest land, Cropland, grassland, and Settlements from the total dam conversion area, the remainder is considered to be the area converted from other land-use categories.

It should be noted that the area presented in the CRF “Table 5.D SECTORAL BACKGROUND DATA FOR LAND USE, LAND-USE CHANGE AND FORESTRY – Wetlands” is not the annually converted area in FY2010 but the sum of annually converted areas during the past 20 years.

Table 7-35 Area of land annually converted to Wetlands (single year)

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Wetlands	kha	0.43	1.72	2.04	0.07	0.07	0.09	0.34
Forest land converted to Wetlands	kha	0.31	1.24	1.48	0.05	0.05	0.07	0.25
Cropland converted to Wetlands	kha	0.02	0.10	0.13	0.005	0.004	0.01	0.02
Rice field	kha	0.01	0.02	0.09	0.004	0.00	0.00	0.01
Upland field	kha	0.01	0.05	0.03	0.0005	0.001	0.002	0.01
Orchard	kha	0.005	0.018	0.010	0.0001	0.0003	0.0005	0.002
Wetlands converted to Wetlands	kha	0.007	0.029	0.019	0.0003	0.001	0.001	0.003
Settlements converted to Wetlands	kha	0.002	0.006	0.007	0.0002	0.0002	0.0003	0.001
Other land converted to Wetlands	kha	0.09	0.34	0.41	0.01	0.01	0.02	0.07

2) **Carbon Stock Change in Dead Organic Matter in “Land converted to Wetlands”**

● **Estimation Method**

➤ **Carbon stock changes in Dead Organic Matter**

Carbon stock changes in dead organic matter in “Forest land converted to Wetlands” were estimated by applying the Tier 2 estimation method as described in section 7.5.2.b)2). With regard to “Cropland, Grassland, Settlements and Other land converted to Wetlands”, carbon stocks of dead wood and litter carbon pools were assumed to be minor and the stock changes were ignorable as described in section 7.5.2.b)2) and 7.6.2.b)2) for example, and were thus reported as “NA”.

● **Parameters**

➤ **Carbon Stocks in Dead Organic Matter**

The average carbon stocks in dead wood and litter in Forest land before conversion are shown in Tables 7-6 and 7-7. It is assumed that they become zero immediately after conversion, and are not accumulated after conversion.

● **Activity Data (Area)**

The area of land that was converted to Wetlands during the past 20 years is determined by subtracting the estimated area that was not converted during the past 20 years from the total area of Wetlands in those years. The areas are shown in Table 7-36 below.

Table 7-36 Area of land converted to Wetlands within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Wetlands	kha	83.7	62.3	64.6	41.0	28.5	28.1	27.8
Forest land converted to Wetlands	kha	60.6	45.1	46.7	29.6	20.6	20.3	20.1
Cropland converted to Wetlands	kha	5.2	3.7	3.8	2.4	1.7	1.7	1.7
Rice field	kha	1.9	1.3	1.5	1.1	0.8	0.8	0.8
Upland field	kha	2.2	1.7	1.7	1.0	0.7	0.7	0.7
Orchard	kha	1.0	0.7	0.6	0.3	0.2	0.2	0.2
Grassland converted to Wetlands	kha	0.9	0.8	0.9	0.5	0.4	0.4	0.4
Settlements converted to Wetlands	kha	0.3	0.2	0.2	0.1	0.1	0.1	0.1
Other land converted to Wetlands	kha	16.7	12.5	12.9	8.2	5.7	5.6	5.6

c) *Uncertainties and Time-series Consistency*

● *Uncertainty Assessment*

Uncertainties of the parameters and the activity data for living biomass, dead organic matter, and soil were individually assessed on the basis of field study results, expert judgment, or the default values described in the *GPG-LULUCF*. The uncertainty was estimated as 30% of the total emissions from the land converted to Wetlands. More detailed information on the uncertainty assessment is described in Annex 7. In addition, concrete uncertainty estimates for individual parameters in this category will be reported in future submissions after investigation is completed.

● *Time-series Consistency*

Although the methods to estimate the area of “Forest land converted to other land use” are different between FY1990-2004 and post FY2005, as described in section 7.5.2.b)1), time-series consistency for this subcategory is basically ensured.

d) *Source-/Sink-specific QA/QC and Verification*

QC is implemented in accordance with the Tier 1 approach described in the GPG (2000) and the *GPG-LULUCF*. The Tier 1 approach includes procedures for checking parameters and activity data, and for archiving references. More detailed information on the QA/QC activity procedures is described in section 6.1 of Annex 6.

e) *Source-/Sink-specific Recalculations*

● *Converted area and carbon stock changes in FY2005 to 2009*

Due to error correction on converted area, area was recalculated. Carbon stock changes in “Orchard converted to Wetlands” and “Grassland converted to Wetlands” are recalculated accordingly.

● *Carbon Stock Changes in Forest land converted to Wetlands*

Living biomass accumulation in Forest land subjected to conversion has been estimated by extrapolation of the trend of living biomass accumulation in the D area during the period between 2005 and the latest year. In line with this, the carbon stock changes were recalculated in accordance with the updated living biomass accumulation reflecting the value of FY2010.

f) *Source-/Sink-specific Planned Improvements*

● *Validity of the Assumption used in the Method of Estimating the Area of Wetlands*

Under the present estimation method, Wetlands are assumed to consist of “water surfaces”, “rivers” and “canals”, as defined in the national land-use classification, and the whole area is estimated by summing the areas covered by these three land types. However, this estimation method may fail to cover the entire wetland area. The validity of the assumption used in the estimation method is now under revision.

● **Method of Obtaining Data of the Area of Storage Reservoirs**

Storage reservoirs (excluding dams) can be considered as artificial flooded land, but the area they cover are not included in the area of flooded land. Therefore, a method used to obtain data on the area covered by the reservoirs needs to be considered.

● **Estimation Method of Soil Carbon Stock Change upon “Land-Use Conversion from Other Land to Wetlands”**

The estimation method will be considered when new data and information are obtained.

7.8. Settlements (5.E)

Settlements are all developed land, including transportation infrastructure and human habitats, and preclude lands that have been placed in other land-use categories. In Settlements, trees existing in urban green areas such as urban parks and special greenery conservation zones absorb carbon.

In FY2010, Japan’s settlement area was about 3.76 million ha, equivalent to about 9.9% of the national land. The net emissions by this category in FY2010 were 2,518 Gg-CO₂; this represents a decrease of 39.4% below the FY1990 value and an increase of 428.5% over the FY2009 value. The biggest driver for the increase over the previous year is that the single-year converted area from Forest land to Settlements in FY2010 increased by 119% compared to the area in FY2009, and the emissions resulting from carbon stock loss in living biomass in “Forest land converted to Settlements” in FY2010 increased by 127.6% over the 2009 value.

In this section, Settlements are divided into two subcategories, “Settlements remaining Settlements (5.E.1.)” and “Land converted to Settlements (5.E.2.)”, and described separately in the following subsections.

Carbon pools estimated in Settlements are living biomass, dead organic matter and soils. Dead organic matters for several subcategories are included in living biomass stock changes.

With respect to activity data, the Tier 1a and Tier 1b of the *GPG-LULUCF* assume that removals derived from biomass growth are equal to emissions derived from biomass loss where the average tree age in a green area is older than 20 years. Therefore, carbon stock changes in urban green areas older than 20 years after establishment are regarded as zero and not estimated. Moreover, urban green areas included in the activity data are divided into two categories; urban green facilities established as urban parks and others, and special greenery conservation zones for which conservation measures are taken and permanent protection is ensured.

<Urban green areas>

- Urban Green Facilities (urban parks, green areas on roads, green areas at ports, green areas around sewage treatment facilities, green areas by greenery promoting system for private green space, green areas along rivers and erosion control sites, green areas around government buildings and green areas around public rental housing, which are within 20 years after establishment),
- Special Greenery Conservation Zones, which are within 20 years after designation.

Table 7-37 Emissions and removals in Settlements resulting from carbon stock changes

Gas	Category	Carbon pool	Unit	1990	1995	2000	2005	2008	2009	2010
CO ₂	5.E. Settlements	Total	Gg-CO ₂	4,158.4	2,799.6	947.4	125.9	143.8	476.5	2,518.3
		Living Biomass	Gg-CO ₂	2,765.1	1,892.8	606.8	24.5	64.4	321.6	1,870.8
		Dead Wood	Gg-CO ₂	1,155.7	817.7	424.2	257.9	232.8	273.0	596.6
		Litter	Gg-CO ₂	494.0	345.6	172.8	99.8	89.3	119.1	276.9
		Soil	Gg-CO ₂	-256.5	-256.5	-256.5	-256.5	-242.6	-237.2	-226.0
	5.E.1. Settlements remaining Settlements	Total	Gg-CO ₂	-955.5	-1,030.6	-1,071.1	-1,114.2	-1,065.5	-1,058.1	-1,011.4
		Living Biomass	Gg-CO ₂	-758.5	-821.9	-856.2	-891.8	-849.5	-844.8	-806.3
		Dead Wood	Gg-CO ₂	IE,NE	IE,NE	IE,NE	IE,NE	IE,NE	IE,NE	IE,NE
		Litter	Gg-CO ₂	-9.8	-10.2	-10.5	-10.9	-11.0	-10.9	-10.7
		Soil	Gg-CO ₂	-187.3	-198.6	-204.3	-211.5	-205.0	-202.5	-194.4
	5.E.2. Land converted to Settlements	Total	Gg-CO ₂	5,113.9	3,830.2	2,018.5	1,240.1	1,209.4	1,534.6	3,529.7
		Living Biomass	Gg-CO ₂	3,523.6	2,714.7	1,463.0	916.4	913.9	1,166.4	2,677.1
		Dead Wood	Gg-CO ₂	1,155.7	817.7	424.2	257.9	232.8	273.0	596.6
		Litter	Gg-CO ₂	503.8	355.7	183.4	110.7	100.3	130.0	287.6
		Soil	Gg-CO ₂	-69.2	-57.8	-52.1	-45.0	-37.6	-34.7	-31.6

7.8.1. Settlements remaining Settlements (5.E.1)

a) Source/Sink Category Description

This subcategory deals with carbon stock changes in living biomass, litter of dead organic matter and soils in urban green areas in “Settlements remaining Settlements”, which have remained Settlements without conversion during the past 20 years. This subcategory is divided into three subparts: “Special Greenery Conservation Zones”, “Urban Green Facilities” and “Other”. In these subparts, carbon stock changes in the “Special Greenery Conservation Zones” and “Urban Green Facilities” are estimated. In addition, carbon stock changes reported in “Revegetation” activities under Article 3, paragraph 3, of the Kyoto Protocol correspond to those in the “Urban Green Facilities” constructed in and after 1990⁷. However, “Special Greenery Conservation Zones” are not included in the areas of the Revegetation activities. In the CRF tables, “Special Greenery Conservation Zones” are described as “Urban Green Areas not subject to RV”, “Urban Green Facilities” as “Urban Green Areas subject to RV”, and “Other” as “Other than Urban Green Areas”, respectively. Carbon stock changes that are possibly included in the subpart “Other”, such as trees in gardens in personal residences, are reported as “NE” because their activity data are not available. Moreover, with respect to litter and soils, carbon stock changes in urban parks and green areas at ports are reported due to limited availability of parameters. The net removal by this subcategory in FY2010 was 1,011 Gg-CO₂; this represents an increase of 5.9% over the FY1990 value and a decrease of 4.4% below the FY2009 value.

b) Methodological Issues

1) Carbon Stock Changes in Living Biomass in “Settlements remaining Settlements”

● Estimation Method

Due to the differences of characteristics of urban green areas, the Tier 1a method is used for special greenery conservation zones that are communal green areas, and Tier 1b is used for urban green facilities.

⁷ The “Special Greenery Conservation Zones” are not included in Revegetation because they do not meet its definition.

➤ **Tier 1a: Special Greenery Conservation Zones**

$$\Delta C_{SSaLB} = \Delta C_{LBaG} - \Delta C_{LBaL}$$

$$\Delta C_{LBaG} = A \times PW \times BI$$

ΔC_{SSaLB} : changes in carbon stocks in living biomass in special greenery conservation zones (t-C/yr)

ΔC_{LBaG} : gains in carbon stocks due to growth in living biomass in special greenery conservation zones (t-C/yr)

ΔC_{LBaL} : losses in carbon stocks due to losses in living biomass in special greenery conservation zones (t-C/yr).

Note: assumed as “0” (zero) in accordance with the *GPG-LULUCF*

A : area of special greenery conservation zones younger than or equal to 20 years since designation (ha)

PW : rate of forested area (rate of forested area per park area). Note: assumed as 100%

BI : growth per crown cover area (t-C/ha crown cover/yr)

➤ **Tier 1b: Urban Green Facilities**

$$\Delta C_{SSbLB} = \sum (\Delta C_{LBbGi} - \Delta C_{LBbLi})$$

$$\Delta C_{LBbGi} = \Delta B_{LBbG}$$

$$\Delta B_{LBbGi} = \sum NT_{i,j} * C_{Ratei,j}$$

ΔC_{SSbLB} : changes in carbon stocks in living biomass in urban green facilities (t-C/yr)

ΔC_{LBbG} : gains in carbon stocks due to growth in living biomass in urban green facilities (t-C/yr)

ΔC_{LBbL} : losses in carbon stocks due to losses in living biomass in urban green facilities (t-C/yr). Note: assumed as “0” (zero) in accordance with the *GPG-LULUCF*

ΔB_{LBbG} : Annual biomass growth in urban green facilities (t-C/yr)

C_{Rate} : Annual living biomass growth per tree (t-C/tree/yr)

NT : Number of trees

i : Types of urban green facilities (urban parks, green areas on roads, green areas at ports, green areas around sewage treatment facilities, green areas by greenery promoting systems for private green space, green areas along rivers and erosion control sites, green areas around government buildings, or green areas around public rental housing)

j : Tree species

● **Parameters**

➤ **Tier 1a: Annual rate of living biomass growth per crown cover area (special greenery conservation areas)**

The default value, 2.9 t-C/ha crown cover/yr, indicated in the *GPG-LULUCF* (p. 3.297) is taken for the annual rate of living biomass growth of trees per crown cover area in special greenery conservation zones.

➤ **Tier 1b: Annual rate of living biomass growth per tree (urban green facilities)**

The following parameters are taken as the annual living biomass growth rates per tree in urban green facilities.

Table 7-38 Annual biomass growth rate per tree in urban green facilities

Climate category		Annual living biomass growth per tree [t-C/tree/yr]	Remarks
Urban green facilities	Hokkaido	(Other than green areas on roads) 0.0098 (Green areas on roads) 0.0103	Default values 0.0033-0.0142 (t-C/tree/yr) provided in the <i>GPG-LULUCF</i> (p. 3.297, Table 3A.4.1) and the annual growth rates of living biomass for the trees in Japan (0.0204 for Japanese zelkova, 0.0103 for ginkgo, 0.0095, for bamboo-leaf oak and 0.0122 t-C/tree/yr for camphor tree) are combined with the distribution ratio of tree types in sampled urban parks. The annual growth rates of living biomass for these trees are calculated by using the growth curve for each tree species, which were developed based on the results of surveys conducted by the National Institute for Land and Infrastructure Management (NILIM) of the MLITT ⁸ and the average trunk diameter at breast height for each tree species ⁹ , which were determined from the results of surveys in urban parks. For green areas on roads, the distribution ratio of tree species indicated by the surveys in green areas on roads ¹⁰ is taken into account.
	Areas other than Hokkaido	(Other than green areas on roads) 0.0105 (Green areas on roads) 0.0108	

● Activity Data

The areas of “Settlements remaining Settlements” in a certain year reported in the CRF tables are estimated by subtracting the cumulative total area of “Land converted to Settlements” during the past 20 years in a year subject to estimation from the total area of “Settlements” in the year subject to estimation. Moreover, in the CRF tables, the areas of “Settlements remaining Settlements” are reported in three subparts: “Special Greenery Conservation Zones”, “Urban Green Facilities” and “Other”. Within these subparts, carbon stock changes in trees less than or equal to 20-year growth in “Special Greenery Conservation Zones” and “Urban Green Facilities” are estimated.

Japan assumes trees less than or equal to 20-year growth as those growing in urban green areas less than or equal to 20 years since establishment or designation. With respect to Tier 1a, tree crown areas in the “Special Greenery Conservation Zones” are applied as activity data. Tier 1b applies the number of tall trees planted in the “Urban Green Facilities” as activity data.

Table 7-39 Areas of “Settlements remaining Settlements” within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Settlements remaining Settlements	kha	2,350.6	2,655.7	2,863.6	3,049.5	3,171.5	3,210.4	3,232.9
Urban green facilities	kha	64.28	68.18	70.14	72.60	69.23	68.52	65.27
Special greenery conservation zones	kha	1.9	3.6	4.8	5.5	5.6	5.8	5.9
Other	kha	2,284.4	2,583.9	2,788.7	2,971.4	3,096.7	3,136.0	3,161.7

⁸ Reference: Matsue et al., “Estimation equations for the amount of CO₂ fixed by planted trees in cities in Japan”, Journal of the Japanese Society of Revegetation Technology, 35 (2), 318-324, 2009.

⁹ Reference: Parks and Green Spaces Division of the Ministry of Land, Infrastructure and Transport, “FY2004 Survey on evaluation techniques for the effectiveness of greening in urban parks for preventing global warming”, March, 2005.

¹⁰ The distribution ratio of tree types is taken from the Road Tree Planting Status Survey (The Street tree of Japan VI), which covered green areas on roads throughout Japan.

➤ **Tier 1a: Tree crown areas (“Special Greenery Conservation Zones”)**

The tree crown areas of the special greenery conservation zones are calculated by multiplying the area of special greenery conservation zones determined by the Ministry of Land, Infrastructure, Transport and Tourism by the rate of tree crown area, which is assumed to be 100%.

Table 7-40 Areas of special greenery conservation zones younger than or equal to 20 years since notification

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Total	kha	1.9	3.6	4.8	5.5	5.6	5.8	5.9
Green space conservation zones	kha	0.6	0.9	1.4	2.0	2.1	2.3	2.4
Suburban green space conservation zones	kha	1.2	2.7	3.4	3.5	3.5	3.5	3.5

➤ **Tier 1b: Number of tall trees (“Urban Green Facilities”)**

The number of tall trees in urban green facilities is calculated according to the same methods that are used for revegetation activities under Article 3, paragraph 4, of the Kyoto Protocol. Brief descriptions of the calculation methods for each urban green facilities are stated below. In addition, for a detailed description of these calculation methods see section 11.3.2.5.a. in Chapter 11 in this NIR.

- **Urban parks, green areas at ports, green areas around sewage treatment facilities, green areas along rivers and erosion control sites, green areas around government buildings, and green areas around public rental housing**

The number of tall trees is calculated by (1) calculating the areas falling under this category by multiplying each area by the area ratio of land conversion for the whole country, and then (2) calculating the number of tall trees in the calculated areas by multiplying each of the areas by the number of tall trees per area. The number of tall trees per area for each urban green facility is shown in the table below.

Table 7-41 Number of tall trees per area

Item	Unit	Number of tall trees per area	
		Hokkaido	Areas other than Hokkaido
Urban parks	tree/ha	329.5	222.3
Green areas at ports	tree/ha	329.5	222.3
Green areas around sewage treatment facilities	tree/ha	129.8	429.2
Green areas along rivers and erosion control	tree/ha	1470.8	339.0
Green areas around government buildings	tree/ha	108.8	108.8
Green areas around public rental housings	tree/ha	219.9	219.9

- **Green areas on roads**

Activity data (the number of tall trees) in these facilities are calculated by the following procedures.

1. The number of tall trees planted during 20 years after establishing green areas on roads is calculated by using data from the “Road Tree Planting Status Survey” which had been implemented in FY1987, FY1992, FY2007 and each corresponding fiscal year during the commitment period,
2. The number of tall trees calculated in Step 1 is multiplied by the ratio of the number of tall trees planted on the roads whose planted area is more than 500 m²,
3. The number of tall trees calculated in Step 2 is multiplied by the area ratio of “Land remaining Settlements”.

The values of Step 3 become the number of tall trees constituting the activity data on green areas on roads.

- **Green areas by greenery promoting systems for private green space**

Activity data (the number of tall trees) are available for each facility. Therefore, the total number of tall trees is used as activity data.

2) Carbon Stock Changes in litter in “Settlements remaining Settlements”

In this category carbon stock changes in litter in urban parks and green areas at ports are estimated. Carbon stock changes in dead wood result in “IE” because they are included in carbon stock changes in living biomass. Carbon stock changes in litter in the subcategories other than urban parks and green areas at ports are not estimated due to the difficulty of obtaining such activity data.

● **Estimation Method**

A country-specific method is applied for this estimation because no method for carbon stock changes in litter in settlements is provided in the *GPG-LULUCF*. The estimation method is described below.

$$\Delta C_{SSLit} = \sum (A_i \times L_{it,i})$$

ΔC_{SSLit} : Carbon stock changes in litter in “Settlements remaining Settlements” (t-C/yr)

A : Area of urban parks and green areas at ports in “Settlements remaining Settlements” (ha)

L_{it} : Carbon stock change per area in urban parks or green areas at ports (t-C/ha/yr)

i : Type of Urban Green Facilities (urban parks or green areas at ports)

● **Parameters**

For litter, Japan estimates carbon stock changes only in branches and leaves dropped naturally from tall trees. Carbon stock changes in litter per urban park area is calculated by using annual accumulation of litter per tree (Hokkaido and other prefectures: 0.0006 t-C/tree/yr) based on the results of field surveys in urban parks, and the number of tall trees per area and the ratio of litter moved to off-site due to management including cleaning (54.4%). As a result, carbon stock changes in litter per urban park area have been calculated to 0.0882 t-C/ha/yr for Hokkaido and 0.0594 t-C/ha/yr for other prefectures. In addition, the carbon fraction in litter is assumed to be 0.4 t-C/t-d.m. which is a default value provided in the 2006 IPCC Guidelines (p. 8.21).

● **Activity Data**

Activity data on this category are the same as those on living biomass in urban parks and green areas at ports, as described in activity data of “Remaining land: Above-ground biomass, Below-ground biomass” (section 11.4.1.1.d a) of Chapter 11).

3) Carbon Stock Changes in Soils in “Settlements remaining Settlements”

Urban parks, for which the carbon stock changes in soils per area were determined, and Green areas at ports, whose management practices are similar to those for urban parks, are the subject of estimation. In general, soils in RV land are not organic soils (peat soils and muck soils). Therefore, organic soils are reported as “NO”, and only mineral soils are estimated.

● **Estimation Method**

Country specific methodology to estimate carbon stock changes in soils on Settlements is applied

since its default methodology (Tier 1) is not provided by *GPG-LULUCF*.

$$\Delta C_{SSSoils} = \sum_i (\Delta C_{Mineral,i} - L_{Organic,i})$$

$$\Delta C_{Mineral,i} = A_i \times \Delta C_{Soil,i}$$

$\Delta C_{SSSoils}$: Annual carbon stock changes in soils in “Settlements remaining Settlements” (t-C/yr)

$\Delta C_{Mineral}$: Annual carbon stock changes in mineral soils in “Settlements remaining Settlements” (t-C/yr)

$L_{Organic}$: Annual carbon stock changes in organic soils in “Settlements remaining Settlements” (=0) (t-C/yr)

A : Area of “Settlements remaining Settlements” (ha)

C_{Soil} : Annual carbon stock changes in soils per area of “Settlements remaining Settlements” (t-C/ha/yr)

i : Type of Urban Green Facilities (Urban parks and Green areas at ports)

● Parameters

As described in section 11.4.1.1.d d), carbon stock changes in soils per area of Urban parks and Green areas at port are estimated based on the results of surveys conducted in urban parks which have been established within 20 years (1.20 t-C/ha/yr). Thus, this value is applicable to Urban parks and Green areas at port which were established within 20 years.

● Activity Data

Activity data on this category are the same as the area of urban parks and green areas at ports, as described in activity data of “Remaining land: Above-ground biomass, Below-ground biomass” (section 11.4.1.1.d a) of Chapter 11).

c) Uncertainties and Time-series Consistency

● Uncertainty Assessment

The default values shown on page 3.297 in the *GPG-LULUCF* were applied to the annual carbon stock changes for trees in urban parks and special greenery conservation zones. The uncertainty estimates for the emission and removal factors were determined by using the decision tree, to be $\pm 50\%$ through application of the standard value shown in the *GPG-LULUCF* (page 3.298).

Moreover, the uncertainty estimates for living biomass in special greenery conservation zones apply expert judgment according to the decision tree for activity data in the *GPG-LULUCF*. These estimates were determined as 10% for the number of tall trees and existing trees and the areas of existing special greenery conservation zones, 17% for wooded areas, and 20% for forested areas.

Meanwhile, the uncertainty estimates for activity data and the parameters of urban parks, green areas on roads, green areas at ports, green areas around sewage treatment facilities, green areas by greenery promoting systems for private green space, green areas along rivers and erosion control sites, green areas around government buildings and green areas around public rental housing are 67% and 48%, respectively.

As a result, the uncertainty estimate was 76% for the entire removal by “Settlements remaining Settlements”. The methodology of uncertainty assessment was described in Annex 7. In addition, concrete uncertainty estimates for individual parameters in this category will be reported in future submissions after investigation is completed.

- **Time-series Consistency**

Although the methods to estimate the area of “Forest land converted to other land use” are different between FY1990-2004 and post 2005, as described in section 7.5.2.b)1), time-series consistency for this subcategory is basically ensured.

d) Source-/Sink-specific QA/QC and Verification

QC is implemented in accordance with the Tier 1 approach described in the GPG (2000) and the *GPG-LULUCF*. The Tier 1 approach includes procedures for checking parameters and activity data, and for archiving references. The QA/QC activity procedures are described in section 6.1 of Annex 6.

e) Source-/Sink-specific Recalculations

- **Carbon stock changes for living biomass and litter on urban green facilities**

The accuracy of estimation of living biomass was improved by 1) improving the modeled data of tall trees per area due to the re-examination of data and the addition of sample facilities and 2) updating the annual growth rate of living biomass per tree, which reflected the data of the annual living biomass growth rate by tree type obtained for some tree species. By the same token, the accuracy of estimation for litter was improved by updating the annual carbon stock changes in litter in accordance with 1) and 2).

- **Carbon stock changes in soil**

Carbon stock changes in soils, which have been reported as “NE” is estimated for urban parks and green areas at ports, whose management practices are similar to those of urban parks, by calculating soil carbon stock changes per area. The soil carbon stock changes per area were calculated for urban parks, for which new information was available, by applying the difference method, which takes differences in soil carbon stocks between planted areas or lawn areas and bare areas in urban parks into account.

- **Corrections of land area of urban green facilities that have been reported in the NIR and the CRF**

The reported land area of urban green facilities was corrected. This error correction didn't affect the estimates of emissions or removals, since the correct values had originally been used for the estimation. In response to the correction, the area of the subcategory “Other” (“Other than Urban Green Areas” on the CRF) was also changed, since it is calculated by subtracting the areas of “Special Greenery Conservation Zones” and “Urban Green Facilities” from the total of “Settlements remaining Settlements”.

f) Source-/Sink-specific Planned Improvements

- **Growth Rate of Living Biomass per Unit of Greening Area in “Special Greenery Conservation Zones”**

The default values in the *GPG-LULUCF* were applied to the living biomass growth rate per unit of greening area in special greenery conservation zones. However, the growth rate needs to be further examined, and a parameter that can be finally applied as the growth rate should be determined. Therefore, Japan is considering the characteristics of greening activity and will seek a parameter that best suits the actual situation.

- **Validity of the Assumption used in the Method of Estimating the Area of Settlements**

The present estimation method assumes settlement areas as “roads” and “human habitats” in the land-use categorization. However, the validity of the assumption is under re-examination.

7.8.2. Land converted to Settlements (5.E.2)

a) Source/Sink Category Description

Land conversion to Settlements results in carbon stock changes in the living biomass, dead organic matter (dead wood and litter) and soil in the land areas subject to the conversion. This subcategory deals with the carbon stock changes in lands converted to Settlements, which were converted from other land-use categories to Settlements within the past 20 years. With respect to dead organic matter, Japan used the CENTURY-jfos model to estimate carbon stocks in dead organic matter in Forest land, and then estimated carbon stock changes in “Settlements converted from Forest land”. However, the area of “Wetlands converted to Settlements” and “Other land converted to Settlements” cannot be obtained by the current method. Thus, carbon stock changes in these carbon pools were reported as “NO”.

The net emissions by this subcategory in FY2010 were 3,530 Gg-CO₂; this represents a decrease of 31.0% below the FY1990 value and an increase of 130.0% over the FY2009 value.

b) Methodological Issues

1) Carbon stock changes in Living Biomass in “Land converted to Settlements”

● Estimation Method

Carbon stock changes in living biomass in “Land converted to Settlements” are estimated by calculating the carbon stock changes before and after conversion and adding annual carbon stock changes in “Land converted to urban green facilities”. The carbon stock changes in living biomass before and after conversion are estimated by applying the equation in section 3.6.2 in the *GPG-LULUCF* (multiplying the land area converted from each land use to Settlements by the difference between the values of living biomass stock before and after conversion, and by the carbon fraction). Biomass stocks in land converted to urban green areas are increased due to the growth of trees planted after conversion. Hence, carbon stock changes in living biomass in land converted to urban green facilities are estimated by calculating carbon stock changes before and after conversion and adding annual carbon stock changes after conversion that are estimated by applying the Tier 1b method in section 3A.4.1.1.1 in the *GPG-LULUCF*.

$$\Delta C_{LSLB} = \sum (A_i \times (CR_a - CR_{b,i}) \times CF) + \sum (\Delta C_{LS(UG)Gi} - \Delta C_{LS(UG)Li})$$

$$\Delta C_{LS(UG)G} = \Delta B_{LS(UG)G}$$

$$\Delta B_{LS(UG)G} = \sum NT_j \times C_{Ratej}$$

ΔC_{LSLB} : Carbon stock changes in living biomass in land converted to Settlements (t-C/yr)

A_i : Area of land converted annually to Settlements from land-use type i (ha/yr)

CR_a : Carbon reserves immediately following conversion to Settlements (t-d.m./ha), default=0

$CR_{b,i}$: Carbon reserves in land-use type i immediately before conversion to Settlements (t-d.m./ha)

CF : Carbon fraction of dry matter (t-C/t-d.m.)

- I : Type of land before conversion
- $\Delta C_{LS(UG)Gi}$: Annual carbon stock gain in living biomass in land converted to urban green areas due to growth in living biomass (t-C/yr)
- $\Delta C_{LS(UG)Li}$: Annual carbon stock loss in living biomass due to loss of living biomass (t-C/yr) Note: the averaged ages of estimated trees are less than or equal to 20 years old, therefore, the loss is assumed as “0” (zero) in accordance with the *GPG-LULUCF*
- $\Delta B_{LS(UG)G}$: Annual living biomass growth in land converted to urban green areas (t-C/yr)
- C_{Rate} : Annual living biomass growth per tree (t-C/tree/yr)
- NT : Number of trees
- i : Type of urban green area after conversion (urban parks, green areas on roads, green areas at ports, green areas around sewage treatment facilities, green areas by greenery promoting systems for private green space, green areas along rivers and erosion control sites, green areas around government buildings, or green areas around public rental housing)
- j : Tree species

● **Parameters**

➤ **Living biomass stocks for each land-use category**

Table 7-6 shows the living biomass stocks before and after conversion. Carbon stock losses due to loss of living biomass are assumed as “0” (zero) in accordance with the *GPG-LULUCF*, because trees subject to estimation are all younger than or equal to 20 years old. Table 7-38 shows the annual living biomass growth of trees in land converted to urban green areas.

➤ **Carbon fraction of dry matter**

0.5 (t-C/t-d.m.) (default value, *GPG-LULUCF*)

● **Activity Data**

➤ **Land Areas converted to Settlements**

With respect to the area of “Land converted to Settlements”, only the areas converted to Settlements from Forest land, Cropland and Grassland are determined. Since no data is available on the area converted to Settlements from Wetlands or other-land use categories, no figures are reported in those land-use categories. Instead, they are reported as “NO”. It should be noted that the area presented in the CRF “Table 5.E SECTORAL BACKGROUND DATA FOR LAND USE, LAND-USE CHANGE AND FORESTRY—Settlements” is not the annually converted area in FY2010 but the sum of annually converted areas during the past 20 years.

- **Conversion from Forest land**

Areas of “Forest land converted to Settlements” were estimated as described in section 7.5.2.b).1).

- **Conversion from Cropland**

For former rice fields, upland fields, and orchards (according to “Area Statistics for Cultivated and Commercially Planted Land”), the areas of land converted to factories, roads, housing, and forest roads are used.

- **Conversion from Grassland**

For former pasture land and grazed meadow land constituting moved or converted Cropland which is converted to Settlements (according to “Area Statistics for Cultivated and Commercially Planted Land”), the areas of land converted to factories, roads, housing, and forest roads are used.

Table 7-42 Area of land converted to Settlements (single year)

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Settlements	kha	43.8	36.3	23.8	14.8	16.5	14.5	19.3
Forest land converted to Settlements	kha	19.3	13.6	7.1	4.3	4.0	4.9	10.8
Cropland converted to Settlements	kha	21.4	19.5	14.5	9.2	10.9	8.2	7.2
Rice field converted to Settlements	kha	13.0	12.1	9.5	6.0	7.1	5.0	4.1
Upland field converted to Settlements	kha	6.1	5.6	3.8	2.5	3.0	2.5	2.4
Orchard converted to Settlements	kha	2.3	1.8	1.1	0.7	0.8	0.7	0.6
Grassland converted to Settlements	kha	3.2	3.1	2.2	1.4	1.6	1.4	1.3
Wetlands converted to Settlements	kha	IE	IE	IE	IE	IE	IE	IE
Other land converted to settlements	kha	IE	IE	IE	IE	IE	IE	IE

➤ **Area and number of trees in “Land converted to urban green areas”**

The areas of “Land converted to urban green areas” are calculated by multiplying the whole area of each urban green area by the area ratio of land conversion for the whole country. The number of trees is calculated by multiplying each urban green area converted from other land-use categories by the number of trees per area. For detailed information regarding these activity data see section 11.3.2.5.a. in Chapter 11 in this NIR.

2) Carbon Stock Change in Dead Organic Matter in “Land converted to Settlements”

In this category carbon stock changes in dead wood and litter in “Settlements converted from Forest land”, and those in litter in land converted to urban parks and green areas at ports are estimated.

With respect to dead wood, only the carbon stock change in “Forest land converted to Settlements” was estimated. The Tier 2 method was applied to the estimation in accordance with the method for “Conversion from other land use to Cropland” in the *GPG-LULUCF*. Carbon stock changes in dead wood in “Land converted to urban green facilities” are reported as “IE” because they are included in living biomass.

In regards to litter, the carbon stock changes in “Settlements converted from Forest land” and “Land converted to urban parks and green areas at ports” are estimated. The Tier 2 method is applied to the estimation of the carbon stock changes in “Settlements converted from Forest land” in accordance with the method for “Conversion from other land use to Cropland” in the *GPG-LULUCF*. Carbon stock changes in litter in “Land converted to urban parks and green areas at ports” are estimated by applying Japan’s country-specific estimation method due to the lack of an estimation method in the *GPG-LULUCF*. Carbon stock changes in litter in land converted to urban green areas other than urban parks and green areas at ports are not estimated due to the difficulty of obtaining their activity data.

The area of “Wetlands converted to Settlements” and “Other land converted to Settlements” cannot be obtained by the current method. Thus, carbon stock changes in these carbon pools were reported as “NO”.

● **Estimation Method**

$$\Delta C_{LS} = \Delta C_{FS} + \Delta C_{LSLit}$$

ΔC_{FS} : Carbon stock changes in dead organic matter in Settlements converted from Forest land (t-C/yr)

ΔC_{LSLit} : Carbon stock changes in litter in urban parks and green areas at ports converted from land use categories other than Forest land (t-C/yr)

➤ **Carbon stock changes in dead organic matter in “Settlements converted from Forest land”**

Carbon stock changes in dead organic matter in “Forest land converted to Settlements” are estimated by applying the Tier 1 estimation method described in section 2.3.2.2 in Volume 4 of the 2006 IPCC Guidelines. In addition, all carbon stocks in dead organic matter in the subcategory are assumed oxidized and emitted as CO₂ within the year of conversion.

$$\Delta C_{FS} = \sum ((C_{after,i} - C_{before,i}) \times A)$$

- ΔC_{FS} : Carbon stock changes in dead organic matter in “Forest land converted to Settlements” (t-C/yr)
 $C_{after,i}$: Carbon stock in dead wood or litter after conversion (t-C/ha) Note: carbon stocks after conversion are assumed as “0” (zero).
 $C_{before,i}$: Carbon stock in dead wood or litter before conversion (t-C/ha)
 A : Area of “Forest land converted to Settlements” in a year subject to estimation (ha)
 i : type of dead organic matter (dead wood or litter)

➤ **Carbon stock changes in dead organic matter in “Settlements converted from Forest land”**

Carbon stock changes in dead organic matter in “Forest land converted to Settlements” are estimated by applying the Tier 1 estimation method described in section 2.3.2.2 in Volume 4 of the 2006 IPCC Guidelines. In addition, all carbon stocks in dead organic matter in the subcategory are assumed oxidized and emitted as CO₂ within the year of conversion.

$$\Delta C_{LSLit} = \sum (A_i \times (C_{AfterLit,i} - C_{BeforeLit,I}) + A_i \times Lit_i)$$

- ΔC_{LSLit} : Carbon stock changes in litter in urban parks and green areas at ports converted from land-use categories other than Forest land (t-C/yr)
 A : Area of urban parks or green areas at ports converted from land-use categories other than Forest land for the past year (ha)
 $C_{AfterLit}$: Carbon stock in litter after conversion (t-C/ha)
 $C_{BeforeLit}$: Carbon stock in litter before conversion (t-C/ha)
 Lit : Annual carbon stock changes per area in litter in urban parks or green areas at ports converted from land-use categories other than Forest land (t-C/ha/yr)
 I : Land-use type before conversion
 i : Type of urban green facility after conversion (urban parks or green areas at ports)

● **Parameters**

➤ **Carbon stocks in dead organic matter in “Forest land converted to Settlements”**

Average carbon stocks in dead wood and litter in Forest land before conversion are shown in Tables 7-6 and 7-7. The average carbon stocks in these categories from FY1990 to FY2004 are not estimated; therefore the carbon stocks in FY2005 are substituted for them. In addition, it is assumed that they become zero immediately after conversion, and are not accumulated after conversion.

➤ **Carbon stocks in litter in “Urban parks and green areas at ports converted from land-use categories other than Forest land”**

When urban parks and green areas at ports are converted from land-use categories other than forest land, litter stocked before conversion is not moved to off-site because the ground before conversion, including litter, is continuously used after conversion, or covered with additional soils brought externally. Hence, litter stocked before conversion does not decrease after conversion. In addition,

litter stocks scarcely increase immediately after conversion because newly planted trees do not immediately produce litter. Due to these facts, carbon stock changes before and after conversion are regarded as “0” (zero). Litter stocks accumulated in a year after conversion are calculated by the same method as the one used for urban parks and green areas at ports in “Settlements remaining Settlements” due to the research finding that litter stocks are accumulated in the same way as those in “Settlements remaining Settlements”, namely by natural drop of fallen leaves and branches from trees in “Land converted to urban parks and green areas”.

● Activity Data

➤ Carbon stocks in dead organic matter in “Forest land converted to Settlements”

The area of land that was converted from Forest land to Settlements during the past 20 years is determined by aggregating the areas converted from Forest land to Settlements during the past 20 years. For the areas, see Table 7-43 below.

Table 7-43 Area of land converted to Settlements within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Settlements	kha	868.4	773.3	730.4	648.5	581.5	550.6	526.1
Forest land converted to Settlements	kha	288.5	307.0	298.3	259.3	215.1	196.7	188.2
Cropland converted to Settlements	kha	520.6	409.1	376.8	338.8	318.8	307.8	293.7
Rice field converted to Settlements	kha	320.9	252.1	236.6	215.2	204.6	197.6	188.7
Upland field converted to Settlements	kha	137.2	110.5	101.8	91.9	86.1	83.4	79.8
Orchard converted to Settlements	kha	62.4	46.5	38.5	31.6	28.1	26.8	25.2
Grassland converted to Settlements	kha	59.3	57.2	55.4	50.5	47.6	46.1	44.2
Wetlands converted to Settlements	kha	IE	IE	IE	IE	IE	IE	IE
Other land converted to settlements	kha	IE	IE	IE	IE	IE	IE	IE

➤ Carbon stock changes in litter in “Land converted to urban parks and green areas at ports”

Areas of “Land converted to urban green areas” are calculated in the same manner as the carbon stock changes in living biomass in “Land converted to urban green areas”. They are calculated by multiplying the areas of urban parks and green areas at ports by the area ratio of land conversion for the whole country, respectively. For detailed information regarding these areas see section 11.4.1.1.d f) in Chapter 11 in this NIR.

3) Carbon Stock Change in Soils in “Land converted to Settlements”

Likewise the “Settlements remaining Settlements”, urban parks and green areas at ports, whose management practices are similar to those in urban parks, are the only subject of estimation.

● Estimation Method

Country specific methodology to estimate carbon stock changes in soils on Settlements is applied since its default methodology (Tier 1) is not provided by *GPG-LULUCF*.

$$\Delta C_{LSSoils} = \sum_i (\Delta C_{LSMinerali} - L_{LSOrganici})$$

$$\Delta C_{LSMinerali} = \Delta A_i \times (C_{AfterSoil} - C_{BeforeSoil}) + A_i \times \Delta C_{soil,i}$$

$\Delta C_{LSSoils}$: Annual carbon stock changes in soils in urban parks and green areas at port following land-use conversion other than from Forest land (t-C/yr)

- $\Delta C_{LSMineral}$: Annual carbon stock changes in mineral soils in urban parks and green areas at port following land-use conversion other than from Forest land (t-C/yr)
- $L_{LSOrganic}$: Annual carbon stock changes in organic soils in urban parks and green areas at port following land-use conversion other than from Forest land(=0) (t-C/yr)
- ΔA : Area of land (excluding Forest land) converted to Settlements within a year (ha)
- A : Area of land (excluding Forest land) converted to urban parks and green areas at port (ha)
- $C_{AfterSoil}$: Soil carbon stocks immediately after land-use conversion(t-C/ha)
- $C_{BeforeSoil}$: Soil carbon stocks before land-use conversion(t-C/ha)
- ΔC_{Soil} : Annual carbon stock changes in soils per land area of urban parks and green areas at port following land-use conversion other than from Forest land (t-C/ha/yr)
- i : Types of urban green facilities (Urban parks and Green areas at ports)

● **Parameters**

As mentioned in the section for litter, when urban parks are converted from cropland, grassland or wetlands, soils before conversion are almost never moved to off-site. In general, these soils are used after conversion continuously or covered by additional soils. Therefore, soil carbon stocks do not change due to land-use conversion.

Carbon stock changes in soils within a year after conversion is estimated in the same manner as for the remaining Urban parks and Green areas at ports.

● **Activity Data**

The area is as obtained for estimating living biomass. For detail, refer to section 11.4.1.1.d f) in Chapter 11 in this NIR.

c) **Uncertainties and Time-series Consistency**

● **Uncertainty Assessment**

The uncertainties of the parameters and activity data for living biomass and dead organic matter were individually assessed on the basis of field study results, expert judgment, or the default values described in the *GPG-LULUCF*. The uncertainty estimate was 30% for the entire emission from “Land converted to Settlements”. The methodology used in the uncertainty assessment is described in Annex 7. In addition, concrete uncertainty estimates for individual parameters in this category will be presented in future submissions after investigation is completed.

● **Time-series consistency**

Although the methods to estimate the area of “Forest land converted to other land use” are different between FY1990-2004 and post FY2005, as described in section 7.5.2.b)1), time-series consistency for this subcategory is basically ensured.

d) **Source-/Sink-specific QA/QC and Verification**

QC is implemented in accordance with the Tier 1 approach described in the GPG (2000) and the *GPG-LULUCF*. The Tier 1 approach includes procedures for checking parameters and activity data, and for archiving references. The QA/QC activity procedures are described in section 6.1 of Annex 6.

e) **Source-/Sink-specific Recalculations**

● **Carbon stock changes for living biomass and litter on urban green facilities**

The accuracy of estimation of living biomass was improved by 1) improving the modeled data of tall

trees per area due to the re-examination of data and the addition of sample facilities and 2) updating the annual growth rate of living biomass per tree, which reflected the data of the annual living biomass growth rate by tree type obtained for some tree species. By the same token, the accuracy of estimation for litter was improved by updating the annual carbon stock changes in litter in accordance with 1) and 2).

● **Carbon stock changes in soil**

Carbon stock changes in soils, which have been reported as “NE” is estimated for land converted to urban parks and green areas at ports, whose management practices are similar to those of urban parks, by calculating soil carbon stock changes per area. The soil carbon stock changes per area were calculated for urban parks, for which new information was available, by applying the difference method, which takes differences in soil carbon stocks between planted areas or lawn areas and bare areas in urban parks into account.

● **Carbon Stock Changes in Forest land converted to Settlements**

Living biomass accumulation in Forest land subjected to conversion has been estimated by extrapolation of the trend of living biomass accumulation in the D area during the period between 2005 and the latest year. In line with this, the carbon stock changes were recalculated in accordance with the updated living biomass accumulation reflecting the value of FY2010.

f) **Source-/Sink-specific Planned Improvements**

● **Validity of the Assumption used in the Method of Estimating the Area of Settlements**

The areas of Forest land converted to Settlements are presently assumed as “roads”, “human habitats”, “school reservations”, “parks and green areas”, “road sites”, “environmental facility sites”, “golf courses”, “ski courses” and “other recreation sites” in the national land-use categorization; however, this assumption may fail to cover all the areas. Therefore, the validity of the assumption needs to be re-examined.

7.9. Other land (5.F)

Other land consists of land areas that are not included in the other five land-use categories. As concrete examples of other land, the *GPG-LULUCF* indicates bare land, rock, ice, and unmanaged land areas. In FY2010, Japan’s “Other land” area was about 2.77 million ha, which is equivalent to about 7.3% of the national land. The classification of “Other land” is shown in Table 7-44 below¹¹.

Table 7-44 Land included in the Other land category

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Other land	kha	2,381.0	2,511.0	2,556.0	2,592.0	2,647.0	2,669.0	2,671.0
Defense Facility Site	kha	139.0	140.0	140.0	140.0	140.0	140.0	140.0
Cultivation Abandonment Area	kha	217.0	244.0	343.0	386.0	392.0	394.0	396.0
Coast	kha	46.0	46.0	46.0	46.0	46.0	46.0	46.0
Northern Territories	kha	503.6	503.6	503.6	503.6	503.6	503.6	503.6
Other	kha	1,475.4	1,577.4	1,523.4	1,516.4	1,565.4	1,585.4	1,585.4

The emissions from this category in FY2010 were 382 Gg-CO₂; this represents a decrease of 75.4%

¹¹ These land areas are based on the following statistics: “*Defense of Japan*” by the Ministry of Defense for “Defense Facility Site”, “*World Census of Agriculture and Forestry*” for “Cultivation Abandonment Area”, “Digital national land information” by MLIT for “Coast” and “Land Survey of Prefectures, Shi, Ku, Machi and Mura” by the Geographical Survey Institute for “Northern Territories”.

below the FY1990 value and a decrease of 63.6% over the FY2009 value.

This section divides Other land into two subcategories, “Other land remaining Other land (5.F.1.)” and “Land converted to Other land (5.F.2.)”, and describes them separately in the following subsections.

Table 7-45 Emissions and removals resulting from carbon stock changes in Other land

Gas	Category	Carbon pool	Unit	1990	1995	2000	2005	2008	2009	2010	
CO ₂	5.F. Other land	Total	Gg-CO ₂	1,553.9	1,479.2	1,227.9	954.9	834.3	1,049.0	382.2	
		Living Biomass	Gg-CO ₂	1,142.1	1,141.3	983.7	751.9	659.4	834.5	319.8	
		Dead Wood	Gg-CO ₂	286.2	234.8	169.7	141.0	121.4	144.6	42.0	
		Litter	Gg-CO ₂	125.6	103.1	74.5	61.9	53.5	70.0	20.4	
		Soil	Gg-CO ₂	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	
	5.F.1. Other land remaining Other land	Total	Gg-CO ₂								
		Living Biomass	Gg-CO ₂								
		Dead Wood	Gg-CO ₂								
		Litter	Gg-CO ₂								
		Soil	Gg-CO ₂								
	5.F.2. Land converted to Other land	Total	Gg-CO ₂	1,553.9	1,479.2	1,227.9	954.9	834.3	1,049.0	382.2	
		Living Biomass	Gg-CO ₂	1,142.1	1,141.3	983.7	751.9	659.4	834.5	319.8	
		Dead Wood	Gg-CO ₂	286.2	234.8	169.7	141.0	121.4	144.6	42.0	
		Litter	Gg-CO ₂	125.6	103.1	74.5	61.9	53.5	70.0	20.4	
		Soil	Gg-CO ₂	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	NO,NE	

7.9.1. Other land remaining Other land (5.F.1)

a) Source/Sink Category Description

This subcategory deals with carbon stock changes in “Other land remaining Other land” during the past 20 years. The land area of this subcategory is determined by subtracting the summed areas of the other five land-use categories from the total national land area shown in *the Land Use Status Survey* compiled by the Ministry of Land, Infrastructure, Transport, and Tourism. In concrete terms, the land area of this category includes defense facility sites, cultivation abandonment areas, coasts, and northern territories. However, carbon stock changes in this subcategory are not considered in accordance with the *GPG-LULUCF*.

Table 7-46 Areas of “Other land remaining Other land” within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Other land remaining Other land	kha	2,118.4	2,269.0	2,299.9	2,191.5	2,251.0	2,296.4	2,305.5

b) Source-/Sink-specific Recalculations

Due to the change in methodology for estimating area of Grazed meadow, the area of Other land remaining Other land, which is determined by subtracting the summed areas of the other five land-use categories from the total national land area, are recalculated.

c) Source-/Sink-specific Planned Improvements

● Method of Defining Land Areas

7.3% of the nation’s land is categorized as “Other land remaining Other land”, but the validity of the categorization is presently under examination in a cross-cutting manner through the LULUCF sector.

● Carbon Stock Changes in Living Biomass of “Other land remaining Other land”

The carbon stock changes in the living biomass of “Other land remaining Other land” are assumed to be zero, but this assumption may differ from the actual situation. Therefore, the land-use types in the

“Other land” category will be investigated, and the validity of the assumption will be re-examined. If there are some land-use types that contain living biomass, reclassification of land-use categories will be examined.

7.9.2. Land converted to Other land (5.F.2)

a) Source/Sink Category Description

This subcategory deals with carbon stock changes in the land converted to Other land within the past 20 years. The land area of this subcategory includes land converted for soil and stone mining, land damaged by natural disasters, and land in which cultivation is abandoned. The emissions from this subcategory in FY2010 were 382 Gg-CO₂; this represents a decrease of 75.4% below the FY1990 value and a decrease of 63.6% over the FY2009 value.

With respect to living biomass, its carbon stock change as a result of land use conversion from other land use to Other land was estimated.

With respect to dead organic matter, Japan used the CENTURY-jfos model to estimate carbon stocks in dead organic matter in Forest land, and then estimated carbon stock changes in “Forest land converted to Other land”. Carbon stock changes in dead organic matter in other subcategories (conversion from Cropland and Grassland) were reported as “NA”, since dead organic matter pools before and after conversion were assumed to be zero, as described in section 7.5.2.b)2) and 7.6.2.b)2).

Carbon stock changes in soils in “Land converted to Other land” are not estimated due to lack of data. Therefore, the carbon stock changes in the carbon pool were reported as “NE”.

In addition, the area of “Wetlands converted to Other land” and “Settlements converted to Other land” cannot be obtained by the current method. Thus, carbon stock changes in these carbon pools were reported as “NO”.

b) Methodological Issues

1) Carbon stock change in Living Biomass in “Land converted to Other land”

● Estimation Method

The Tier 2 method was applied as described in section 7.5.2.b)1). Carbon stock changes due to biomass growth in Other land was assumed as zero.

● Parameters

➤ Biomass stock in each Land-Use Category

The values shown in Table 7-6 are used for the estimation of biomass stock changes upon land-use conversion and subsequent changes in biomass stock due to biomass growth in converted land.

➤ Carbon Fraction of dry matter

0.5 (t-C/t-d.m.) (GPG-LULUCF, default value)

● Activity Data (Area)

Only the areas converted from Forest land and Cropland to Other land are determined. Since no data were available on the area converted from Wetlands and Settlements to Other land, estimations for those land-use categories could not be made. Therefore, they were reported as “NO.” It should be

noted that the area presented in the CRF “Table 5.F SECTORAL BACKGROUND DATA FOR LAND USE, LAND-USE CHANGE AND FORESTRY – Other land” is not the annually converted area in FY2010 but the sum of annually converted areas during the past 20 years.

➤ **Conversion from Forest Land**

See section 7.5.2.b)1).

➤ **Conversion from Cropland**

For former rice fields, upland fields, and orchards, the area classified as “other, natural disaster damage” is used according to *the Area Statistics for Cultivated and Commercially Planted Land*.

➤ **Conversion from Grassland**

For former pasture land and grazed meadow land, the area of former pasture land classified as “other, natural disaster damage” (according to *the Area Statistics for Cultivated and Commercially Planted Land*) and the area of former grazed meadow land which is classified as “other, classification unknown” (*the Moving and Conversion of Cropland*) are used.

Table 7-47 Area of land converted to Other land (single year)

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Other land	kha	24.0	30.1	28.9	20.4	14.7	16.0	11.2
Forest land converted to Other land	kha	4.8	3.9	2.8	2.4	2.1	2.6	0.8
Cropland converted to Other land	kha	15.4	20.3	17.1	13.2	8.8	8.8	7.2
Rice field	kha	5.0	5.8	6.1	7.2	4.0	2.9	3.1
Upland field	kha	7.6	10.9	8.4	4.7	3.8	4.6	3.2
Orchard	kha	2.8	3.6	2.5	1.3	1.0	1.2	0.8
Grassland converted to Other land	kha	3.9	5.9	9.0	4.9	3.8	4.7	3.2
Wetlands converted to Other land	kha	IE	IE	IE	IE	IE	IE	IE
Settlements converted to Other land	kha	IE	IE	IE	IE	IE	IE	IE

2) Carbon Stock Changes in Dead Organic Matter in “Land converted to Other land”

● Estimation Method

Carbon stock changes in dead organic matter in “Forest land converted to Other land” were estimated by applying the Tier 2 estimation method as described in section 7.5.2.b)2). With regard to “Cropland and Grassland converted to Other land” where Cropland conversion occurred mainly due to abandonment and natural disaster, carbon stock changes were reported as “NE” because suitable knowledge for estimating carbon stock changes from land-use conversion are not available. The area of “Wetlands converted to Other land” and “Settlements converted to Other land” cannot be obtained by the current method. Thus, carbon stock changes in these carbon pools were reported as “NO”.

● Parameters

➤ Carbon Stocks in Dead Organic Matter in “Other Land converted from Forest Land”

The average carbon stocks in dead wood and litter in Forest land before conversion are shown in Tables 7-6 and 7-7. It is assumed that carbon stocks become zero immediately after conversion, and are not accumulated after conversion.

● Activity Data (Area)

The values of annually converted area from each land-use category to Other land during the past 20 years are summed up to obtain the total area that is converted to Other land during the same time period.

Table 7-48 Area of land converted to Other land within the past 20 years

Category	Unit	1990	1995	2000	2005	2008	2009	2010
Land converted to Other land	kha	591.4	516.5	512.9	526.7	506.3	490.4	477.6
Forest land converted to Other land	kha	103.5	103.6	97.3	81.0	70.6	67.4	63.4
Cropland converted to Other land	kha	419.6	338.1	316.5	324.8	311.2	299.7	291.5
Rice field	kha	181.2	120.3	105.0	108.4	106.3	105.1	103.2
Upland field	kha	164.2	153.7	154.8	161.8	154.9	147.9	143.5
Orchard	kha	74.2	64.1	56.6	54.6	50.0	46.8	44.8
Grassland converted to Other land	kha	68.3	74.7	99.1	120.9	124.5	123.3	122.7
Wetlands converted to Other land	kha	IE	IE	IE	IE	IE	IE	IE
Settlements converted to Other land	kha	IE	IE	IE	IE	IE	IE	IE

c) *Uncertainties and Time-series Consistency*

● *Uncertainty Assessment*

The uncertainties of the parameters and the activity data for living biomass and dead organic matter were individually assessed on the basis of field study results, expert judgment, or the default values described in the *GPG-LULUCF*. The uncertainty was estimated as 28% for the entire emission from the “Land converted to Other land”. More detailed information on the uncertainty assessment is described in Annex 7. In addition, concrete uncertainty estimates for individual parameters in this category will be given in future submissions after investigation is completed.

● *Time-series Consistency*

Although the methods to estimate the area of “Forest land converted to other land use” are different between FY1990-2004 and post FY2005, as described in section 7.5.2.b)1), the time-series consistency for this subcategory is basically ensured.

d) *Source-/Sink-specific QA/QC and Verification*

QC is implemented in accordance with the Tier 1 approach described in the GPG (2000) and the *GPG-LULUCF*. The Tier 1 approach includes procedures for checking parameters and activity data, and for archiving references. More detailed information on the QA/QC activity procedures is described in section 6.1 of Annex 6.

e) *Source-/Sink-specific Recalculations*

● *Areas and carbon stock changes in “Grassland converted to Other land” from FY2005 to 2009*

Due to recalculation in the area of “Grassland converted to Wetlands” as described in section 7.7.2. e), the area and carbon stock changes in “Grassland converted to Other land” are recalculated.

● *Carbon Stock Changes in Forest land converted to Other land*

Living biomass accumulation in Forest land subjected to conversion has been estimated by extrapolation of the trend of living biomass accumulation in the D area during the period between 2005 and the latest year. In line with this, the carbon stock changes were recalculated in accordance with the updated living biomass accumulation reflecting the value of FY2010.

f) *Source-/Sink-specific Planned Improvements*

● *Breakdown Analysis of Other Land and Reclassification into Other Land-Use Categories*

A further breakdown analysis of the Other land is required, since it may still include some areas that are supposed to be classified into other land-use categories even after the reallocation carried out in

this year.

● **Carbon Stock Changes in Living Biomass in “Land converted to Other Land”**

The carbon stock changes in living biomass in “Land converted to Other land” were assumed to be zero because of a lack of reference information for “Other land”. However, this assumption may differ from the actual situation. Therefore, the methods used to quantifying the carbon stock are being examined.

● **Estimation Method of Soil Carbon Stock Changes in “Forest land, Cropland and Grassland converted to Other Land”**

The estimation method will be considered when new data and information are obtained.

7.10. Direct N₂O emissions from N fertilization (5. (I))

a) Source/Sink Category Description

It is assumed that the volume of nitrogen-based fertilizer applied to forest soils is included in the amount of applied nitrogen-based fertilizers in the Agriculture sector, although fertilization application in Forest land may rarely be conducted in Japan. Therefore, these sources have been reported as “IE”.

7.11. N₂O emissions from drainage of soils (5.(II))

a) Source/Sink Category Description

Regarding the N₂O emissions from soil drainage activities in Forest land and Wetlands, experts noted that the N₂O emissions were extremely low, because soil drainage activities were very rarely carried out in Japan. Based on this comment, this category is reported as “NO”.

7.12. N₂O emissions from disturbance associated with land-use conversion to Cropland (5.(III))

a) Source/Sink Category Description

This category deals with N₂O emissions from disturbance associated with land-use conversion to Cropland. The emissions by this subcategory in FY2010 were 6.2 Gg-CO₂; this represents a decrease of 93.1% below the FY1990 value and a decrease of 18.7% below the FY2009 value.

Table 7-49 N₂O emissions from disturbance associated with land-use conversion to Cropland

Gas	Category	Unit	1990	1995	2000	2005	2008	2009	2010	
N ₂ O	Total	Gg-N ₂ O	0.29	0.20	0.10	0.05	0.03	0.02	0.02	
		Gg-CO ₂ , eq.	90.02	60.71	31.91	14.74	8.37	7.60	6.18	
	Cropland	Gg-N ₂ O	0.29	0.20	0.10	0.05	0.03	0.02	0.02	
		Forest land converted to Cropland	Gg-N ₂ O	0.23	0.17	0.10	0.04	0.02	0.02	0.02
		Grassland converted to Cropland	Gg-N ₂ O	0.048	0.027	0.004	0.003	0.003	0.003	0.003
		Wetlands converted to Cropland	Gg-N ₂ O	0.0143	0.0027	0.0006	0.0003	0.0003	0.0003	0.0002
		Other land converted to Cropland	Gg-N ₂ O	IE,NE	IE,NE	IE,NE	IE,NE	IE,NE	IE,NE	IE,NE
	Other	Gg-N ₂ O	NA	NA	NA	NA	NA	NA	NA	

b) Methodological Issues**● Estimation Method**

According to the *GPG-LULUCF*, the Tier 1 method is used.

$$N_2O - N_{conv} = N_2O_{net-min} - N$$

$$N_2O_{net-min} - N = EF \times N_{net-min}$$

$$N_{net-min} = C_{released} \times 1 / CN_{ratio}$$

$N_2O - N_{conv}$: N_2O emissions due to land-use conversion to Cropland (kg N_2O -N)

$N_2O_{net-min} - N$: N_2O emissions due to land-use conversion to Cropland (kg N_2O -N/ha/yr)

$N_{net-min}$: annual N emissions from soil disturbance associated with mineralization of soil organic matter (kgN/ha/yr)

EF : emission factor

CN_{ratio} : Carbon Nitrogen ratio of the biomass

$C_{released}$: soil carbon stock that has been mineralized within the past 20 years

● Parameters**➤ *CN ratio for soils***

11.3 (Country specific data (Ministry of the Environment, 2006))

➤ *N-N₂O emission factor for soils*

0.0125 kg-N₂O-N/kg-N (default value stated in the *GPG-LULUCF*, Page 3.94)

● Activity Data

Areas of land converted to Cropland and carbon emissions from soils due to this conversion are used. The areas are the same as those shown in Table 7-25.

c) Uncertainties and Time-series Consistency**● Uncertainty Assessment**

The uncertainties of parameters were individually assessed on the basis of field studies, expert judgment, or default values described in the *GPG-LULUCF*, and the uncertainty estimates for the carbon emissions from soil in “Land converted to Cropland” were applied to the activity data of this category. As a result, the uncertainty estimates of N_2O emissions from disturbance associated with land-use conversion to Cropland were 75%. The methodology of uncertainty assessment was described in Annex 7.

● Time-series Consistency

Time-series consistency for this category is ensured.

d) Source-/Sink-specific QA/QC and Verification

QC is implemented in accordance with the Tier 1 approach described in the GPG (2000) and the *GPG-LULUCF*. The Tier 1 approach includes procedures for checking parameters and activity data, and for archiving references. The QA/QC activity procedures are described in section 6.1 of Annex 6.

e) Source-/Sink-specific Recalculations

None.

f) Source-/Sink-specific Planned Improvements

● **Estimation Method of the “Area converted from Forest Land to Cropland” and “from Grassland to Cropland”**

The methods used to obtain data on the area of “Forest land converted to Cropland” and “Grassland converted to Cropland” need to be improved as mentioned in section 7.5.2.f). Therefore, the validity of the estimates is being reviewed, and the estimation method is being reexamined.

● **Method of Obtaining Data of the “Area converted from Grassland to Cropland”**

Data on the area of land converted from grassland to Cropland cannot be obtained from current statistics, so the carbon stock changes in the areas have not been estimated. Therefore, the methods used to obtain data need to be investigated for the following areas:

- from pasture land to upland field
- from pasture land to orchard
- from grazed meadow to rice field
- from grazed meadow to upland field
- from grazed meadow to orchard.

7.13. CO₂ emissions from agricultural lime application (5.(IV))

a) Source/Sink Category Description

This category deals with CO₂ emissions from agricultural lime application. The emissions from this category in FY2010 were 270 Gg-CO₂; this represents a decrease of 50.9% below the FY1990 value. One of the reasons for the decline compared to FY1990 is that the amount of calcium carbonate fertilizer applied in Japan has decreased because the chemical nature of soils was progressively improved by soil amendment.

Table 7-50 CO₂ emissions from agricultural lime application

Gas	Category	Unit	1990	1995	2000	2005	2008	2009	2010
CO ₂	Total	Gg-CO ₂	550.2	303.5	332.9	231.3	305.6	270.1	270.1
	Cropland	Gg-CO ₂	IE	IE	IE	IE	IE	IE	IE
	Grassland	Gg-CO ₂	IE	IE	IE	IE	IE	IE	IE
	Other	Gg-CO ₂	550.2	303.5	332.9	231.3	305.6	270.1	270.1
	Limestone	Gg-CO ₂	549.9	303.0	332.4	230.7	304.1	269.6	269.6
	Dolomite	Gg-CO ₂	0.3	0.5	0.5	0.6	1.6	0.6	0.6

b) Methodological Issues

● **Estimation Method**

The Tier 1 method is used in accordance with the *GPG-LULUCF* (page 3.80).

$$\Delta C_{CCLime} = (M_{Limestone} \times EF_{Limestone} + M_{Dolomite} \times EF_{Dolomite}) \times 44/12$$

ΔC_{CCLime} : annual CO₂ emissions from agricultural lime application (t-CO₂/yr)

$M_{Limestone}$: annual amount of calcic limestone (t/yr)

$M_{Dolomite}$: annual amount of dolomite (t/yr)

$EF_{Limestone}$: emission factor of calcic limestone (t-C/t)

$EF_{Dolomite}$: emission factor of dolomite (t-C/t)

● **Parameters**

- **Emission factor of calcic limestone (CaCO_3)**
0.120 [t-C/t] (default value, *GPG-LULUCF*).
- **Emission factor of dolomite ($\text{CaMg}(\text{CO}_3)_2$)**
0.122 [t-C/t] (default value, *GPG-LULUCF*).

● **Activity Data**

➤ **Annual amount of lime applied to Cropland**

These data were calculated by adding up lime production and import quantities as listed in *the Yearbook of Fertilizer Statistics (Pocket Edition)* published by the Ministry of Agriculture, Forestry and Fisheries of Japan. Based on expert judgment, all of the “Calcium carbonate fertilizer” and 70% of each of “Fossil seashell fertilizer”, “Crushed limestone” and “Seashell fertilizer” listed in the Yearbook was classified as calcic limestone (CaCO_3), and all of the “Magnesium carbonate fertilizer” and 74% of “Mixed magnesium fertilizer” as dolomite ($\text{CaMg}(\text{CO}_3)_2$).

c) **Uncertainties and Time-series Consistency**

● **Uncertainty Assessment**

The uncertainty for this category was assessed based on the uncertainty of the emission factor (see 2006GL, p.11.27) and that of the statistics that provided the activity data. Consequently, the uncertainty of CO_2 emissions from this category was assessed and estimated as 51%. More detailed information on the uncertainty assessment is described in Annex 7. In addition, concrete uncertainty estimates for each parameter in this category will be given in future submissions after investigation is completed.

● **Time-series Consistency**

Time-series consistency for this category is ensured.

d) **Source-/Sink-specific QA/QC and Verification**

QC is implemented in accordance with the Tier 1 approach described in the *GPG (2000)* and the *GPG-LULUCF*. The Tier 1 approach includes procedures for checking parameters and activity data, and for archiving references. More detailed information on the QA/QC activity procedures is described in section 6.1 of Annex 6.

e) **Source-/Sink-specific Recalculations**

Statistical values of FY2009 were revised and the emissions were recalculated..

f) **Source-/Sink-specific Planned Improvements**

None

7.14. Biomass burning (5.(V))

a) **Source/Sink Category Description**

This category deals with emissions of CH_4 , CO , N_2O and NO_x from biomass burning resulting from forest fires. The emissions resulting from wildfires in “Forest land remaining Forest land” and “Land

converted to Forest land” are reported in a lump in the cell for wildfires in “Forest land remaining Forest land” in the CRF tables, because the data in the statistics for forest fires include the wildfires occurred in both of the categories. Moreover, controlled burning activities in forests are quite rarely implemented in Japan because the activities are stringently restricted by the “Waste Management and Public Cleansing Act” and “Fire Service Act”. Hence, the emissions resulting from controlled burning in Forest land are reported as “NO”.

Controlled burning resulting from land conversion from land-use categories other than forest land to forest land is also very rarely carried out in Japan because of heavy restrictions imposed under the “Waste Management and Public Cleansing Law” and the “Fire Defense Law”. Hence, CH₄, CO, N₂O, and NO_x emissions derived from controlled burning other than in forest land are reported as “NO”.

CH₄, CO, N₂O and NO_x emissions from controlled burning in Cropland are reported as “NE” because they are not estimated due to lack of data. CH₄, CO, N₂O and NO_x emissions from wildfires in Cropland are reported as “NO”. One of the characteristics of Japan’s cropland is intensive management. Under this management style, the occurrence of wildfire is regarded as negligibly small. CH₄, CO, N₂O and NO_x emissions from wildfires in “Land other than Forest land and Cropland” are reported as “NE” because information on wildfires is not sufficiently collected.

The emissions by this subcategory in FY2010 were 2.33 Gg-CO₂; this represents a decrease of 75.1% below the FY1990 value and a decrease of 75.4% below the FY2009 value. These variations originate mainly from variations in the volume of timber damaged in wildfires in private forests (Table 7-53).

Table 7-51 Non-CO₂ emissions from biomass burning

Gas	Category	Unit	1990	1995	2000	2005	2008	2009	2010	
All	Total	Gg-CO ₂ eq.	9.4	9.6	8.6	10.1	23.9	9.5	2.3	
	Total	Gg-CH ₄	0.4	0.4	0.4	0.4	1.0	0.4	0.1	
CH ₄		Gg-CO ₂ eq.	8.5	8.7	7.8	9.2	21.7	8.6	2.1	
	Forest land	Gg-CH ₄	0.4	0.4	0.4	0.4	1.0	0.4	0.1	
	Cropland	Gg-CH ₄	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	
	Grassland	Gg-CH ₄	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	
	Wetlands	Gg-CH ₄	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	
	Settlements	Gg-CH ₄	NO	NO	NO	NO	NO	NO	NO	
	Other land	Gg-CH ₄	NO	NO	NO	NO	NO	NO	NO	
	Other	Gg-CH ₄	NA	NA	NA	NA	NA	NA	NA	
	N ₂ O	Total	Gg-N ₂ O	0.003	0.003	0.003	0.003	0.007	0.003	0.0
			Gg-CO ₂ eq.	0.864	0.886	0.790	0.931	2.205	0.874	0.215
Forest land		Gg-N ₂ O	0.003	0.003	0.003	0.003	0.007	0.003	0.0	
Cropland		Gg-N ₂ O	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	
Grassland		Gg-N ₂ O	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	
Wetlands		Gg-N ₂ O	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	NE,NO	
Settlements		Gg-N ₂ O	NO	NO	NO	NO	NO	NO	NO	
Other land		Gg-N ₂ O	NO	NO	NO	NO	NO	NO	NO	
Other		Gg-N ₂ O	NA	NA	NA	NA	NA	NA	NA	

b) Methodological Issues

● Estimation Method

For CH₄, CO, N₂O and NO_x emissions due to biomass burning, the Tier 1 method is used.

➤ Forest land

(CH₄, CO)

$$bbGHG_f = L_{forestfires} \times ER$$

(N₂O, NO_x)

$$bbGHG_f = L_{forestfires} \times ER \times NC_{ratio}$$

$bbGHG_f$: GHG emissions due to forest biomass burning

$L_{forestfires}$: Carbon released due to forest fires(tC/yr)

ER : Emission ratio (CO : 0.06, CH₄ : 0.012, N₂O : 0.007, NO_x : 0.121)

NC_{ratio} : Nitrogen Carbon ratio of the biomass

● Parameters

➤ Emission ratio

The following values are applied to emission ratios for non-CO₂ gases due to biomass burning.

CO: 0.06, CH₄: 0.012, N₂O: 0.007, NO_x: 0.121

(default value stated in the *GPG-LULUCF*, Table 3A.1.15)

➤ NC ratio

The following values are applied to NC ratio.

NC ratio: 0.01 (default value stated in the *GPG-LULUCF* p.3.50)

● Activity Data

➤ Forest land

As activity data in “Forest land”, carbon released due to forest fire is used. Carbon released due to forest fire is estimated by the Tier 3 method in the *GPG-LULUCF*. For each of the national forest land and private forest land, carbon emissions are calculated from the fire-damaged timber volume multiplied by wood density, the biomass expansion factor and the carbon fraction of dry matter.

$$\Delta C = \Delta C_i + \Delta C_c$$

$L_{forestfires}$: carbon emissions due to fire (t-C/yr)

ΔC_{fn} : carbon emissions due to fire in national forests (t-C/yr)

ΔC_{fp} : carbon emissions due to fire in private forests (t-C/yr)

- National forest

$$\Delta C_{fn} = Vf_n \times D_n \times BEF_n \times CF$$

ΔC_{fn} : carbon emissions due to fire in national forests (t-C/yr)

Vf_n : damaged timber volume due to fire in national forests (m³/yr)

D_n : wood density in national forests (t-d.m./m³)

BEF_n : biomass expansion factor for national forests

CF : carbon fraction of dry matter (t-C/t-d.m.)

- **Private forest**

$$\Delta C_{fp} = Vf_p \times D_p \times BEF_p \times CF$$

- ΔC_{fp} : carbon emissions due to fire in private forests (t-C/yr)
 Vf_p : damaged timber volume due to fire in private forests (m³/yr)
 D_p : wood density in private forests (t-d.m./m³)
 BEF_p : biomass expansion factor for private forests
 CF : carbon fraction of dry matter (t-C/t-d.m.)

The values for wood density and biomass expansion factors for national and private forest land are determined as weighted averages using the ratios of intensively managed forests and semi-natural forests.

Table 7-52 Wood density and biomass expansion factors for national and private forests

Type	Wood density [t-d.m./m ³]	Biomass expansion factor
National forest	0.49	1.61
Private forest	0.46	1.61

Source: Based on Forestry Agency data

Biomass stock change due to fires is separately estimated for national forests and private forests. With regard to national forests, the timber volume of standing trees damaged due to fires in national forests in the *Handbook of Forestry Statistics* is used. With regard to private forests, the damaged timber volume due to fires is estimated by using the actual damaged area and damaged timber volume by age class (inquiry survey by Forestry Agency). Damaged timber volume for age class equal to or under 4 is calculated by multiplying the stand volume per unit area of age class equal to or under 4 estimated by the Forestry Status Survey and the NFRDB by loss ratio (ratio of damaged timber volume to stand volume) of age class equal to or over 5 in private forests. The loss ratio is assumed to be constant regardless of age class.

Table 7-53 Damaged timber volume due to wild fire

Category		Unit	1990	1995	2000	2005	2008	2009	2010
Damaged timber volume due to disturbance in national forest		m ³	3,688.0	1,014.0	1,599.0	359.0	1,901.0	976.0	976.0
Damaged timber volume due to disturbance in private forest		m ³	63,601.8	68,360.7	60,227.9	72,575.5	170,730.3	67,417.1	15,809.9
≧5	Actual damaged area	kha	0.29	0.94	0.48	0.35	0.57	0.37	0.07
	Damaged timber volume	m ³	47,390.0	58,129.0	54,487.0	59,235.0	119,900.0	55,628.0	12,780.0
≦4	Actual damaged area	kha	0.27	0.51	0.16	0.27	0.85	0.28	0.06
	Damaged timber volume	m ³	16,211.76	10,231.74	5,740.89	13,340.50	50,830.31	11,789.06	3,029.92

Source: Based on "Handbook of Forestry Statistics" for national forest, and Forestry Agency data for private forest

Note

In Japan, emissions due to biomass burning are estimated separately for national forests and for private forests, because of different reporting procedures in regards to forest fire information. However, forest fires in Japan are covered by a set of data for both national forests and private forests, and the emissions are thus appropriately estimated.

c) **Uncertainties and Time-series Consistency**

● **Uncertainly Assessment**

The uncertainties for parameters and activity data related to biomass burning were individually assessed on the basis of field studies, expert judgment, or default values described in the *GPG-LULUCF*. As a result, the uncertainty estimates for the emissions resulting from biomass burning were 40% for CH₄ and 42% for N₂O, respectively. The methodology of uncertainty assessment is described in Annex 7. In addition, concrete uncertainty estimates for individual parameters in this category will be reported in future submissions after investigation is completed.

● ***Time-series Consistency***

Time-series consistency for biomass burning in “Forest Land remaining Forest Land” is ensured by using the same data sources (*Handbook of Forestry Statistics* compiled by the Forestry Agency, and the data provided by the Agency) and the same methodology from 1990 to 2010.

d) Source-/Sink-specific QA/QC and Verification

QC is implemented in accordance with the Tier 1 approach described in the *GPG (2000)* and the *GPG-LULUCF*. The Tier 1 approach includes procedures for checking parameters and activity data, and for archiving references. The QA/QC activity procedures are described in section 6.1 of Annex 6.

e) Source-/Sink-specific Recalculations

● ***Revision of damaged timber volume in national forests in FY2009***

Due to change in statistical value of the damaged timber volume in national forests, emissions were revised.

● ***Corrections of the estimation of damaged timber volume in private forests***

The loss ratio (= damaged timber volume per area affected by fire / stand volume per area) is applied to obtain the damaged timber volume in private forests. For the calculation of this ratio, the stand volume per area of overall forest land had been applied instead of the one of private forests. Now the stand volume per area of private forests is appropriately applied to the estimation of the loss ratio, and therefore, emissions were recalculated.

f) Source-/Sink-specific Planned Improvements

● ***Burning of pruned branches from orchard trees***

Although woody biomass such as pruned branches from orchard trees may partially be burnt, non-CO₂ emissions from this burning are not estimated. When data for the treatment of orchard tree residues become available, the emissions will be estimated and reported in the inventories.

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Chapter 8. Waste (CRF Sector 6)

8.1. Overview of Sector

In the waste sector, greenhouse gas emissions from treatment and disposal of waste are estimated for solid waste disposal on land (6.A.), wastewater handling (6.B.), waste incineration (6.C.), and other (6.D.)¹ in accordance with treatment processes.

Waste to be covered in this sector is the waste as defined in the Revised 1996 IPCC Guidelines. In the case of Japan, the waste does not only include municipal waste and industrial waste as defined by the Waste Disposal and Public Cleansing Law, but also recyclables and valuables that are re-used within a company. Since waste statistics are compiled separately for municipal waste and industrial waste in Japan, estimation methodologies for many of emission sources in the waste sector are discussed respectively for municipal waste and industrial waste.

In FY 2010, emissions from the waste sector amounted to 20,874 Gg-CO₂ eq. and represented 1.7% of Japan's total greenhouse gas emissions (excluding LULUCF). These emissions had decreased by 19.1% compared to those of FY 1990.

In Japan, annual waste generation is amounted to around 600 Mt and it has hardly changed since FY 1990. As the latest results (FY 2008 data) from *the Annual Report on the Environment in Japan* (the Ministry of the Environment) shows, waste of biogenic-origin, waste of fossil-origin, and metal and nonmetallic mineral wastes accounted respectively for 55%, 3% and 42% of total amount of waste. With regard to the recycle flow for the waste in FY 2008, for overall waste activities generated, natural decomposition, recycling, volume reduction and final disposal accounted for 27%, 17%, 53% and 2%, respectively, for waste of biogenic-origin; while for waste of fossil-origin, recycling, volume reduction and final disposal accounted for 39%, 48% and 13%, respectively. The final disposal amount in Japan has been decreasing year by year.

8.2. Solid Waste Disposal on Land (6.A.)

This category covers CH₄ emissions from solid waste disposal on land. For this emission source category, estimation methodologies were discussed separately for municipal waste and industrial waste in accordance with Japan's waste classification system, and emissions were estimated for the sources presented in Table 8-1.

¹ Data for some emission source categories in the waste sector are complemented by estimation, when statistical data or related data are not available. The methodologies for this estimation are not described in this chapter. For details, see the *Report of the Waste Panel on Greenhouse Gas Emission Estimate (2006)* (hereinafter referred to as Reference #7).

Table 8 - 1 Categories whose emissions are estimated for solid waste disposal on land (6.A.)

Category	Waste types estimated		Treatment type	
6.A.1. (8.2.1)	Municipal solid waste	Kitchen garbage	Anaerobic landfill and Semi-aerobic landfill	
		Waste paper		
		Waste wood		
		Waste textiles (natural fiber) ^{a)}		
		Human waste treatment, Septic tank sludge		
	Industrial waste	Kitchen garbage	Anaerobic landfill and Semi-aerobic landfill	
		Waste paper		
		Waste wood		
		Waste textiles (natural fiber) ^{a)}		
		Sewage sludge		Digested sewage sludge ^{b)}
				Other sewage sludge
		Waterworks sludge		
		Organic sludge from manufacturing industries		
	Livestock waste ^{c)}			
6.A.3. (8.2.3)	Inappropriate disposal ^{d)}		Anaerobic landfill	

- a) Only natural fiber waste textiles are included in the estimation under the assumption that synthetic fiber waste is not biologically decomposed in landfills.
- b) “Digested sewage sludge” includes sewage sludge landfilled after digested and dehydrated. Because digestion treatment reduces the amount of carbon content biodegraded in sludge decreases, CH₄ emissions were estimated separately by landfilled sewage sludge with and without digestion treatment.
- c) Although livestock waste is not classified as “sludge” under Japanese law, emissions from it were estimated within the category of sludge because of the similarities in their properties.
- d) Waste inappropriately disposed of and containing biodegradable carbon is considered to include waste wood, waste paper, and sludge. However, only the emissions from waste wood were calculated, because only its state of dumping is known at present.

Table 8 - 2 GHG emissions from solid waste disposal on land (6.A.)

Gas	Category	Item	Unit	1990	1995	2000	2005	2008	2009	2010	
CH ₄	6.A.1. Managed Solid Waste Disposal site	Kitchen garbage	Gg CH ₄	62.9	60.2	50.4	32.8	21.1	18.3	15.7	
		Waste paper	Gg CH ₄	147.5	136.4	114.4	92.2	78.8	74.5	69.1	
		Waste textile (natural fiber)	Gg CH ₄	9.6	8.6	7.3	6.7	6.0	5.6	5.1	
		Waste wood	Gg CH ₄	46.4	50.4	49.8	47.7	45.9	45.2	44.5	
		sewage sludge	Digested sewage sludge	Gg CH ₄	5.5	5.0	3.8	2.4	1.6	1.4	1.2
			Other sewage sludge	Gg CH ₄	27.3	24.9	19.2	11.8	7.9	6.8	5.9
		Human waste treatment, Septic tank sludge	Gg CH ₄	12.4	9.0	6.5	4.8	3.7	3.4	3.0	
		Waterworks sludge	Gg CH ₄	3.4	3.2	2.6	1.8	1.4	1.3	1.3	
		Organic sludge from industry	Gg CH ₄	48.2	37.9	22.9	14.0	9.7	8.4	7.4	
		Livestock waste	Gg CH ₄	1.4	1.4	1.3	1.1	1.0	1.0	1.0	
		Recovery	Gg CH ₄	-0.8	-0.7	-0.7	0.0	-0.3	-0.3	-0.4	
			Total	Gg CH ₄	363.7	336.1	277.5	215.2	176.9	165.5	153.8
		6.A.3. Other	Inappropriate disposal	Gg CH ₄	0.3	0.8	2.4	2.4	2.1	2.0	2.0
			Total	Gg CH ₄	364.1	336.9	279.9	217.6	179.0	167.5	155.7
			Gg CO ₂ eq	7,645	7,076	5,878	4,569	3,759	3,517	3,270	

Estimated greenhouse gas emissions from solid waste disposal on land are shown in Table 8-2. In FY

2010, greenhouse gas emissions from this source category were 3,270 Gg-CO₂ eq. and accounted for 0.3% of the national total emissions (excluding LULUCF). Emissions from this category decreased by 57.2% compared to the emissions in FY 1990. This CH₄ emissions decrease is the result of decrease in the amount of biodegradable waste landfilled due to the increase in the practice of waste incineration to reduce waste volume in Japan.

8.2.1. Emissions from Managed Landfill Sites (6.A.1.)

a) Source/Sink Category Description

In Japan, part of kitchen garbage, waste paper, waste textiles, waste wood, and sludge in municipal solid waste (MSW) and industrial waste (ISW) is landfilled without incineration; therefore, CH₄ is generated as a result of biodegradation of organic materials from the landfill sites. Because Japanese landfill sites are appropriately managed pursuant to the Waste Disposal and Public Cleansing Law, the amount of CH₄ emitted from there is reported under this category “Emissions from Managed Landfill Sites (6.A.1.)”. Emissions of CO₂ from waste incineration at the managed landfill sites are reported as NO, because waste incineration is not implemented at that site in Japan.

b) Methodological Issues

● Estimation Method

The revised FOD method given in the *2006 IPCC Guidelines* is applied for its emission estimates since this method assumes a process of delay time from the deposition of waste to the substantial rate of CH₄ production. According to the decision tree indicated in the said guidelines, the revised FOD method with country-specific parameters (Tier 3) is used to estimate emissions from this source.

In Japan, emission factor is defined as “CH₄ emissions from biodegradable waste”, and activity data are defined as “the amount of waste biodegraded within the reporting fiscal year”.

$$E = \left\{ \sum (EF_{ij} \times A_{ij}) - R \right\} \times (1 - OX)$$

Where:

- E : CH₄ emissions from landfill sites (kg CH₄)
- EF_{ij} : Emission factor for a biodegradable waste i (dry basis) that is damped into a landfill site j without incineration (kg CH₄/t)
- A_{ij} : Amount of a biodegradable waste i (dry basis) that is damped into a landfill site j without incineration and is biodegraded within an inventory year
- R : Recovered CH₄ in an inventory year (kg CH₄)
- OX : Oxidation factor of CH₄ related to soil cover

● Emission Factors

Emission factors were defined as the amount of CH₄ (kg) generated through decomposition of one ton of biodegradable landfill wastes (dry basis) without incineration. They were established by the type of biodegradable waste (i.e., kitchen garbage, waste paper, waste natural fibers, waste wood, sewage sludge, human waste, waterworks sludge, organic sludge from manufacturing industries and livestock waste) and by the type of landfill site (i.e., anaerobic or semi-aerobic landfill). Emission factors were estimated as indicated below.

Emission factor = (Carbon content) × (Gas conversion rate) × (Methane correction factor) × (Percentages of CH ₄ in landfill gas) × 1000 × 16/12

Carbon Content (per dry weight)

Carbon content per dry weight, which is used as uniform value every year because the property of each waste type does not vary significantly over time, was determined based on the “*Ministry of the Environment, Survey Study on Improving the Accuracy of Emission Factors for Greenhouse Gas Emissions from the Waste Sector, 2010*” (hereinafter referred to as Reference #15) and Reference #7 as indicated in Table 8-3.

Table 8 - 3 Carbon content of waste disposed of in managed landfill sites (dry base)

Item	Carbon Content (%)	Data source
Kitchen garbage	43.4	MSW: Calculated by taking the averages of carbon contents of MSW provided by Tokyo, Yokohama, Kawasaki, Kobe, and Fukuoka (FY 1990-2004) ISW: Substituted the carbon content for the MSW for ISW because its properties are similar to those of MSW (Reference #15)
Waste paper	40.9	
Waste wood	45.2	
Waste natural fiber textiles	45.0	Calculated by taking a weighted average of carbon content estimated based on the constituent of each natural fiber type (cotton, wool, silk, linen, and recycled textiles) by the domestic demand of natural fibers (FY1990-2004) (Reference #7)
Digested sewage sludge	30.0	Expert judgment based on Reference #49, 50, 58, 62
Other sewage sludge	40.0	GPG2000
Human waste treatment, Septic tank sludge	40.0	Substituted the value for “Other sewage sludge” from GPG2000
Waterworks sludge	6.0	Average values of survey results conducted at 23 water purification plants (Reference #15)
Organic sludge from manufacturing	45.0	Value for papermaking industry was substituted because it generates the largest amount of organic sludge finally disposed of. Estimated based on the carbon content of cellulose because the main constituent of organic sludge generated is paper sludge (Reference #7)
Livestock waste	40.0	Substituted the value for “Other sewage sludge” from GPG2000

Gas conversion rate

Gas conversion rate for the biodegradable waste was set at 50% based on Ito (1992).

Methane correction factor

Default values given in the 2006 IPCC Guidelines were used: 1.0 for anaerobic landfill sites and 0.5 for semi-aerobic landfill sites.

Proportions of CH₄ in generated gas

Default value (50%) given in the Revised 1996 IPCC Guidelines was used.

Table 8 - 4 Emission factors by type of biodegradable waste and by treatment

Item	Anaerobic landfill (kg CH ₄ /t)	Semi-aerobic landfill (kg CH ₄ /t)
Kitchen garbage	145	72
Waste paper	136	68
Waste textiles	150	75
Waste wood	151	75
Digested sewage sludge	100	50
Other sewage sludge	133	67
Human waste treatment, Septic tank sludge	133	67
Waterworks sludge	20	10
Organic sludge from manufacturing	150	75
Livestock waste	133	67

● Activity Data

Out of the amount of waste landfilled without incineration (dry basis), the amount of waste degraded within the reporting year was calculated by multiplying the amount of waste remaining in landfills at the end of the previous reporting year by the decomposition rate for waste landfilled. The amount of biodegradable MSW and ISW were determined by type of waste and landfill site.

The amount of waste landfilled in each fiscal year was calculated by multiplying the amount of biodegradable waste landfilled (wet basis) by the percentage of landfill site by the type of site (wet basis), and subtracting the water content by each type of waste. Activity data were estimated going back as far as FY1954, when the Public Cleansing Law (now the Waste Disposal and Public Cleansing Law) was enforced.

$$W_i(T) = W_i(T-1) \times e^{-k} + w_i(T)$$

$$A_i(T) = W_i(T-1) \times (1 - e^{-k})$$

$$k = \ln(2) / H$$

Where:

$A_i(T)$: Amount of waste i degraded in the calculated year (year T) (activity data: dry basis)

$W_i(T)$: Amount of waste i remaining in a landfill in year T

$w_i(T)$: Amount of waste i landfilled in year T

k : Decomposition rate constant (1/year), and

H : Decomposition half-life of waste i (the time taken by landfilled waste i to reduce in amount by half)

The amount of waste i landfilled in year T

= (Amount of biodegraded waste i landfilled in year T)

× (percentages of landfill sites of each site type) × (1 - percentage of water content in waste i)

Amount of biodegradable waste disposed of in landfills

Table 8-5 shows the annual amount of biodegradable waste landfilled (dry basis) in Japan.

Table 8 - 5 Annual amount of biodegradable waste disposed of in landfills

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Kitchen garbage	kt / year (dry)	505	493	311	135	79	65	66
Waste paper	kt / year (dry)	1,213	949	766	534	387	209	325
Waste textiles (natural fiber)	kt / year (dry)	62	56	44	73	13	7	5
Waste wood	kt / year (dry)	672	526	298	195	85	65	75
Digested sewage sludge	kt / year (dry)	59	50	31	11	4	3	3
Other sewage sludge	kt / year (dry)	219	185	114	42	17	17	17
Human waste treatment, Septic tank sludge	kt / year (dry)	78	51	46	47	17	15	15
Waterworks sludge	kt / year (dry)	199	166	146	66	67	67	67
Organic sludge from manufacturing industries	kt / year (dry)	350	156	69	48	23	22	31
Livestock waste	kt / year (dry)	12	12	11	11	11	11	13
Total	kt / year (dry)	3,369	2,644	1,837	1,162	702	481	618

As indicated in Table 8-6, for the data sources for the amount of biodegradable waste landfilled, the *Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes (Volume on Cyclical Use)*, *Waste Management and Recycling Department of the Ministry of the Environment* (hereinafter referred to as the *Cyclical Use of Waste Report*), and the *annual editions of Sewage Statistics (Admin. Ed.)*, Japan Sewage Works Association (hereinafter referred to as the *Sewage Statistics*) were used.

Table 8 - 6 Data overview for the amount of biodegradable waste disposed of in landfill

Item		Data source	MSW	ISW	Remarks
Kitchen garbage		<i>Research on the State of Wide-range Movement and Cyclical Use of Wastes</i> (MoEJ)	Calculated by multiplying total amount of corresponding landfilled waste by the composition ratio of each waste type	- Amount of animal and plant residues directly landfilled and after intermediate processing. - Amount of livestock carcasses directly landfilled	- Estimated by interpolation for some fiscal years, - Substituted FY 1980 value for the years prior to FY 1980
Waste paper				Amount of waste paper directly landfilled	
Waste wood				Amount of waste wood directly landfilled	
Waste natural fiber textiles			Calculated by multiplying by the ratio of natural fiber textiles in textile products each year from “ <i>Annual Textile Statistics Report</i> ”	Amount of waste natural fiber textiles directly landfilled (considering all the amount as waste natural fiber textiles due to the Waste Disposal and Public Cleansing Law)	
Digested sewage sludge		<i>Data provided by METI</i>		Compiled and provided by MLITT	- For some fiscal years, estimated by interpolation - Substituted FY 1985 value for the years prior to FY 1985
Other sewage sludge		<i>Annual editions of Sewage Statistics (Admin. Ed.)</i>		Total amount of sewage sludge excluding the amount of digested sewage sludge	
Human waste treatment, Septic tank sludge		<i>Research on the State of Wide-range Movement and Cyclical Use of Wastes</i> (MoEJ)	“Direct final disposal & final disposal after treatment” of “Human waste treatment and septic tank sludge” (estimated by subtracting the amount of final disposal from those incinerated within the incineration facilities or sewage sludge treatment facilities)		For the years prior to FY 1998, multiplying the amount of human waste sludge in landfill (volume basis) by the weight-conversion factor (1.0 kg/L)
Waterworks sludge		<i>Waterworks Statistics</i> (Japan Water Works Association)		Estimated by “Total amount of soil disposed” and “landfilled percentage” of each purification plant	Substituted FY 1980 value for the years prior to FY 1980
Organic sludge from manufacturing	Papermaking industry	Data provided by Japan Paper Association, Japan Technical Association of the Pulp and Paper Industry		Total amount of organic sludge landfilled for papermaking industry	Substituted FY 1989 value for the years prior to FY 1989
	Chemicals industry	<i>Report on Results of Trend and Industry-Specific Studies on Industrial Wastes (Mining Industry Waste) and Recyclable Waste</i>		Total amount of organic sludge landfilled for chemicals industry and food manufacturing industry	- For some fiscal years, estimated by interpolation - For the years prior to FY 1998, estimated with the data from the <i>Voluntary Action Plan on Environment</i> , <i>Follow-up Action Result</i> (Japan Federation of Economic Organizations) - Substituted FY 1990 value for the years prior to FY 1990
	Food manufacturing industry				
Livestock waste		Survey conducted by MoEJ			Substituted FY 1980 value for the years prior to FY 1980

Percentage of water content in waste

In Japan, activity data are estimated on a dry basis which can identify the carbon content of waste more precisely. The percentages of water content by each type of waste to estimate activity data on a dry basis and its sources are given in Table 8-7. In order to estimate the CO₂ emissions for the category “8.4. Waste Incineration (6C)” as well as this source category, dry basis activity were used for the same reason.

Table 8 - 7 Percentage of water content in waste disposed of in controlled landfill sites

Category		Water content (%)	Source
Kitchen garbage, animal and plant residues		75 (direct final disposal)	Water percentage of kitchen garbage in <i>Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes</i>
		70 (final disposal after treatment)	
Waste paper		20 (MSW) 15 (ISW)	Expert judgment
Waste wood		45 ¹	Expert judgment
Waste natural fiber textiles		20 (MSW) 15 (ISW)	Expert judgment
Sewage sludge	Digested sewage sludge	Specific to each disposal site	Average moisture content of “delivered or final disposal sludge” in <i>Sewage Statistics (Admin. Ed.)</i>
	Other sewage sludge		
Sludge from human waste treatment and septic tanks		85 (direct final disposal)	Moisture content standard of landfill standard (sludge) specified by enforcement ordinance of Wastes Disposal and Public Cleansing Law
		70 (final disposal after treatment)	Determined by specialists
Waterworks Sludge		- *	—
Organic sludge from manufacturing industries		23 (food manufacturing) 43 (chemical industries) - (paper industries)*	Reference of Clean Japan Center Survey
Livestock waste		83.1 (direct final disposal)	Organic percentage in “ <i>Controlling the Generation of Greenhouse Gases in the Livestock Industry</i> ”
		70 (final disposal after treatment)	Expert judgment

*: The water content of waterworks sludge and organic sludge from paper industries are not included in this table because activity data on a dry basis were provided by the data sources.

Percentages of landfill sites by site structure type

• Percentages of MSW landfill sites by site structure type

The percentages of landfill sites by site structure type for MSW were determined by referring to annual editions of Results of Study on Municipal Solid Waste Disposal Waste Management and Recycling Department, Ministry of the Environment (hereinafter referred to as *Results of Study on MSW Disposal*), which lists Japan’s MSW disposal sites in the section “Facility by Type (Final Disposal Sites)”, regarding as semi-aerobic those sites which have leachate treatment facilities and subsurface containment structures, and regarding the percentage of semi-aerobic landfill disposal volume to be the percentage of their total landfill capacity (m³).

Since the percentages of semi-aerobic landfill sites for the period FY1996 and before are not available, they are determined as indicated below:

- For the period FY1977 and before, all the landfill sites including all the sea area landfills are considered to be anaerobic landfill sites since semi-aerobic landfill technology started in

FY1977.

- For the period FY1997 and after, they are determined based on actual data.
- For the period FY1977-1996, they are estimated by linear interpolation using actual data of FY1997 based on expert judgment.

Table 8 - 8 Percentages of MSW landfill sites by site structure

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Anaerobic landfill percentage	%	74.2	64.2	54.4	43.5	41.5	36.5	36.5
Semi-aerobic landfill percentage	%	25.8	35.8	45.6	56.5	58.5	63.5	63.5

• Percentages of ISW landfill sites by site structure type

The percentages of landfill sites by site structure type for ISW are determined as indicated below:

- For the period FY 2008 and after, they are determined based on the ISW landfill site survey results conducted by the Ministry of the Environment.
- For the period FY1977 and before, all the landfill sites including all the sea area landfills are considered to be anaerobic landfill sites since semi-aerobic landfill technology started in FY1977.
- For the period FY1990 -2007, they are estimated by using the total amount of waste landfilled and the actual data of waste deposited of in semi-aerobic landfill sites in FY2008.
- For the period FY1977-1989, they are estimated by linear interpolation using the data of FY1990 based on expert judgment.

Table 8 - 9 Percentages of ISW landfill sites by site structure

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Anaerobic landfill site	%	90.2	81.1	66.4	48.3	45.8	45.8	45.8
Semi-aerobic landfill site	%	9.8	18.9	33.6	51.7	54.2	54.2	54.2

Decomposition half-life

Decomposition half-life is the time taken for 50% of waste landfilled in a certain year to be degraded from its initial mass. According to Ito (1992) (Reference #52), the half-lives for kitchen waste, waste paper, waste natural fiber textiles, and waste wood are respectively 3, 7, 7, and 36 years. Because no relevant research have been obtained to identify a country specific half life for the sludge, the default value of 3.7 years provided in the spreadsheets attached to the 2006 IPCC Guidelines was applied.

Delay time

Delay time is the time lag since the waste is landfilled until the decomposition actually occurs. As no research is found for making it possible to set a delay time specific to Japan, the default value (6 months) given in the 2006 IPCC Guidelines was used.

Table 8 - 10 Amount of biodegraded waste decomposed in each year (Activity data)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Kitchen garbage	kt / year (dry)	520	518	454	315	208	181	157
Waste paper	kt / year (dry)	1,264	1,212	1,056	892	780	743	693
Waste textiles (natural fiber)	kt / year (dry)	75	69	61	59	56	52	47
Waste wood	kt / year (dry)	349	385	385	373	362	356	351
Digested sewage sludge	kt / year (dry)	63	58	47	31	21	18	16
Other sewage sludge	kt / year (dry)	234	219	176	114	78	68	59
Human waste treatment, Septic tank sludge	kt / year (dry)	111	84	64	51	41	37	33
Waterworks sludge	kt / year (dry)	192	185	157	120	97	92	88
Organic sludge from manufacturing industries	kt / year (dry)	368	295	184	119	85	75	66
Livestock waste	kt / year (dry)	12	12	12	11	11	11	11
Total	kt / year (dry)	13	23	33	32	25	22	1,520

The declining trend in the amount of biodegraded waste is affected by the improvement of waste reduction that causes the decrease of landfilled waste.

Amount of CH₄ recovered from landfills

In order to reduce the amount of organic matter content and CH₄ emissions at landfill sites, certain intermediate treatments and landfill methods have been conducted; CH₄ recovery from landfills is not very common practice in Japan. CH₄ recovery from landfilled MSW for the purpose of electric power generation implemented at the Tokyo Metropolitan Inner Landfill Site for the Central Breakwater "Uchigawa-Shobunjo" is the sole practice example in Japan. For ISW, there is no practice of CH₄ recovery from landfills implemented in Japan. Because CO₂ emitted from the combustion of recovered CH₄ is of biogenic-origin, it is not included in the total emissions.

$$R = r \times f \times 16 / 24.4 / 1,000$$

R : Amount of CH₄ recoved in landfill (g)

r : Amount of recovered landfill gas used for electric power generation (m³N)

f : Ratio of CH₄ to recovered gas (-)

- The amount of recovered landfill gas used for electric power generation in "Uchigawa-Shobunjo" landfill

The amount of recovered gas used for electric power generation was provided by the Waste Disposal Management Office of Tokyo.

- Fraction of CH₄ to the recovered gas

The fraction of CH₄ to recovered landfill gas in the *Uchigawa-Shobunjo* has been annually provided since FY 2005 by the Waste Disposal Management Office of Tokyo. The fraction for the years prior to FY 2005 were determined based on the hearing conducted with the Waste Disposal Management Office of Tokyo: 60% for FY 1987, when the recovery of landfill gas was started; 40% for FY 1996; interpolated for FY 1988 through FY 1995; The FY 1996 value was used for FY 1997 through FY 2004.

Table 8 - 11 Amount of CH₄ recovered at landfill sites in Japan

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Amount of gaseous use	km ³ N	1,985	2,375	2,372	140	1,161	1,154	1,266
CH ₄ ratio	%	53.3	42.2	40.0	48.5	37.1	40.0	43.8
Amount of CH ₄ use	km ³ N	1,059	1,003	949	68	431	462	555
CH ₄ unit conversion	Gg CH ₄	0.76	0.72	0.68	0.05	0.31	0.33	0.40

The consumption of gas used for electric power generation during 1991-1994 had decreased compared to the preceding year and the following year because recovered gas was used for the purposes other than electric power generation. The consumption of recovered gas used for electric power generation had decreased compared to 1996 because no electric power generation using recovered gas was conducted between late 1994 and early 1995 due to the relocation of electric power generation facilities. Amount of gas used in 2005 has dropped to less than 10 percent over the previous year because the electric power generating equipment had been halted from April, 2005 to Mid-February, 2006. After resumption, methane concentration was high through to the end of the fiscal year.

CH₄ oxidation rate related by landfill cover soil

Based on law enforcement ordinances and local government ordinances, daily, intermediate and final soil coverings are practiced in the managed final disposal sites for MSW and ISW in Japan. Therefore, the default oxidation factor for managed landfill sites (0.1) was used in accordance with the 2006 IPCC Guidelines.

c) Uncertainties and Time-series Consistency

● *Uncertainties*

The uncertainty in emission factors was evaluated by integrating the uncertainties for carbon content, gas conversion rate, CH₄ correction factor, and percentage of CH₄ in generated gas, and estimated to be in the range of 42.4-108.6%. The uncertainty in activity data was evaluated by integrating the uncertainties for the residual amount of biodegradable waste (landfilled amount and percentage of water content in waste) at the end of the year before the reporting year and the decomposition rate for the reporting year, and estimated to be in the range of 31.7-56.6%. As a result, the uncertainty in the emissions from solid waste disposal sites was estimated to be in the range of 53-113%.

The methods for evaluation of the uncertainty levels for each component are:

- Use of 95% confidence interval of actual measurement data: carbon content (kitchen garbage, waste paper and waste wood)
- Use of the statistical uncertainties: domestic demand for textile and landfilled amount of biodegradable waste
- Based on expert judgment: carbon content (sewage sludge, human waste treatment sludge and organic sludge from manufacturing industries), gas conversion rate, percentage of CH₄ in landfill gas and percentage of water content in biodegradable waste
- Use of the default values in the IPCC Guidelines: carbon content (livestock waste) and CH₄ correction factor
- Use of the values set by the Committee for GHGs Emissions Estimation Methods: carbon content (waterworks sludge)

- Use of the differences between the adopted values and default ones: residual amount of biodegradable waste.

For more details about basic methods for uncertainty assessment in Japan, see the Annex 7.

● *Time-series consistency*

Although some activity data in FY 1990 and thereafter are not available, they are estimated by using the methods described in “Activity data” to develop consistent time-series data. The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Recalculations

- Determining the amount of ISW disposed of in semi-aerobic landfill sites, emission estimates for the period FY1990-2009 were recalculated.
- Determining the amount of waste landfilled after intermediate treatments other than incineration, emission estimates for the period FY1990-2009 were recalculated.
- Updating the data on MSW and ISW landfill amounts, emission estimates for the period FY2008-2009 were recalculated.
- The emission estimates for the period FY1990-2010 were recalculated due to the correction of error in CH₄ recovery estimates.

f) Source-specific Planned Improvements

Further improvements are planned owing to a lack of sufficient current information. Major issues are:

- Determining the value of methane correction factor taking into account the conditions of the management of landfill sites
- Gas conversion rate for each type of biodegradable waste
- Country-specific half-life for sludge at final disposal sites

8.2.2. Emissions from Unmanaged Waste Disposal Sites (6.A.2.)

Because landfill sites in Japan are appropriately managed pursuant to the Waste Disposal and Public Cleansing Law, there are no unmanaged waste disposal sites in Japan. Therefore, the emissions from this source category are reported as NA.

8.2.3. Emissions from Other Managed Landfill Sites (6.A.3.)

8.2.3.1. Emissions from Inappropriate Disposal (6.A.3.a)

a) Source/Sink Category Description

In Japan, the definition of “inappropriate disposal” is waste disposal violating the Waste Disposal and Public Cleansing Act (illegal dumping and other forms of improper disposal on lands or areas other than landfill sites). The ratio of the amount of inappropriate waste disposal is quite small comparing

to the one of appropriate waste disposal. Although these inappropriate disposal lands or areas generally satisfy the conditions of managed disposal sites defined in the *Revised 1996 IPCC Guidelines*, CH₄ emissions from inappropriate disposal are reported under “Other (6.A.3.)”.

Fires are occasionally observed in inappropriate landfill sites, and they may be emitting fossil-fuel derived CO₂. However, since actual data are not available, the emissions from the fires at inappropriate landfill sites are reported as NE.

b) Methodological Issues

● *Estimation Method*

Waste wood and waste paper are the wastes containing biodegradable carbon and being inappropriately disposed without incineration; however, only waste wood is the subject for the estimation, because the residual amount of waste paper should be very small.

In a similar manner for the “Emissions from Controlled Disposal Sites (6.A.1.)”, a FOD method with Japan’s country-specific parameters is used for the estimation. Emissions are estimated by multiplying the amount of waste wood (dry basis) degraded in a reporting year by an emission factor.

● *Emission Factor*

Since inappropriately disposed wastes are generally covered with soil in Japan, the mechanism for CH₄ emissions from inappropriate disposal is regarded as almost same as for the anaerobic landfill. Therefore the same emission factor is used for the anaerobic disposal sites for “waste wood emissions from managed disposal sites”.

● *Activity Data*

Activity data (dry basis) was obtained by subtracting the water content from the residual amount of inappropriately disposed waste wood (wet basis) and multiplied by decomposition rate. The amount of inappropriately disposed waste wood is provided by “Waste Wood (Construction and Demolition)” in *Study on Residual Amounts of Industrial Waste from Illegal Dumping and other Sources* (Waste Management and Recycling Department, Ministry of the Environment). The percentage of water content and the decomposition rate used for estimating emissions from waste wood in managed disposal sites were also used for this source.

Table 8 - 12 Activity data of inappropriately disposed waste wood (dry basis)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Activity data	kt (dry)	2.3	5.5	16.0	15.7	14.0	13.0	13.0

c) Uncertainties and Time-series Consistency

● *Uncertainties*

The uncertainties in emission factor and activity data were evaluated by using the same methods that were used for “Emissions from Controlled Landfill Sites” (6.A.1). The uncertainty in the CH₄ emissions from inappropriate disposal was estimated to be 79%. For more details, see the Annex 7.

● *Time series consistency*

Because data on inappropriate disposal are available only since FY 2002, activity data prior to FY

2002 are estimated. The emissions are calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Recalculations

Due to the changes in the amount of inappropriate disposal, emission estimates for the period FY1990-2009 were recalculated.

f) Source-specific Planned Improvements

For future inventories, long-term efforts on further scientific investigations will be made to identify country-specific parameters.

8.3. Wastewater Handling (6.B.)

The CH₄ and N₂O emissions from wastewater handling are estimated in the “Wastewater Handling (6.B.)”. The target categories are shown in Table 8-13. Since an emission factor that takes into account emissions from wastewater and sludge treatment processes is used in Japan, emissions from these processes are reported altogether. Therefore, total emission amount is reported in the subcategory “Wastewater” in CRF, 6.B.; while IE is reported in the subcategory “Sludge”.

Table 8 - 13 Categories overview for wastewater handling (6.B.)

Category	Type Estimated	Forms of Treatment	CH ₄	N ₂ O	
6.B.1. (8.3.1)	Industrial wastewater (8.3.1.1)	(Sewage treatment plants)	○	○	
	Landfill leachate treatment (8.3.1.1)	Landfill leachate treatment	○	○	
6.B.2. (8.3.2)	Domestic/commercial wastewater	Sewage treatment plants (8.3.2.1)	○	○	
		Domestic wastewater treatment facilities (mainly septic tanks) (8.3.2.2)	Community plant	○	○
			<i>Gappei-shori johkasou</i>	○	○
			<i>Tandoku-shori johkasou</i>	○	○
			Vault toilet	○	○
		Human waste treatment facilities (8.3.2.3)	High-load denitrification treatment	○	○
			Membrane separation	○	○
			Anaerobic treatment	○	○
			Aerobic treatment	○	
			Standard denitrification treatment	○	
	Other	○			
	Degradation of domestic wastewater in nature (8.3.2.4)	Discharge of untreated domestic wastewater	<i>Tandoku-shori johkasou</i>	○	○
Vault toilet			○	○	
On-site treatment			○	○	
Sludge disposal at sea*		Human waste sludge	○	○	
	Sewage sludge	○	○		

*Due to legal regulations on sludge disposal at sea, there has been no activity since FY2009.

Estimated greenhouse gas emissions from wastewater handling are shown in Table 8-14. In FY 2010, emissions from this source category were 2,401 Gg-CO₂ eq. and accounted for 0.2% of the national total emissions (excluding LULUCF). The emissions from this source category decreased by 30.2% compared to those in FY 1990. This emission decrease is the result of decrease in the amount of CH₄ emissions from “Degradation of Domestic Wastewater in Nature” because the practice of wastewater treatment at wastewater treatment plants increased in Japan. Due to the same reason, the N₂O emissions from the subcategory of “Sewage Treatment Plants (6.B.2.a)” for FY1995 through FY1998 increased.

Table 8 - 14 GHG emissions from wastewater handling (6.B.)

Gas	Category	Item	Unit	1990	1995	2000	2005	2008	2009	2010	
CH ₄	6.B.1. Industrial waste water	(Sewage treatment plants)	Gg CH ₄	6.5	6.3	6.1	5.6	5.5	5.3	4.7	
		Landfill leachate treatment	Gg CH ₄	1.2	1.2	1.1	0.8	0.6	0.6	0.4	
	6.B.2. Domestic/commercial wastewater	Sewage treatment plants	Gg CH ₄	8.6	9.1	11.0	11.8	12.2	12.1	12.1	
		Domestic waste water treatment facilities (mainly septic tank)	Gg CH ₄	21.5	20.4	20.6	20.5	20.6	20.3	20.3	
		Humanwaste treatment facilities	Gg CH ₄	5.2	3.2	1.8	1.0	0.7	0.7	0.7	
		Degradation of domestic wastewater in nature	Gg CH ₄	60.2	50.8	39.5	28.7	23.9	22.4	22.4	
	Total			Gg CH ₄	102.1	89.7	78.9	67.6	63.0	60.6	60.5
			Gg CO ₂ eq	2,144	1,884	1,657	1,419	1,322	1,273	1,270	
N ₂ O	6.B.1. Industrial waste water	(Sewage treatment plants)	Gg N ₂ O	0.39	0.38	0.33	0.39	0.41	0.39	0.39	
		Landfill leachate treatment	Gg N ₂ O	0.03	0.03	0.02	0.02	0.01	0.01	0.01	
	6.B.2. Domestic/commercial wastewater	Sewage treatment plants	Gg N ₂ O	1.59	1.67	2.01	2.16	2.25	2.21	2.21	
		Domestic waste water treatment facilities (mainly septic tank)	Gg N ₂ O	1.51	1.35	1.17	0.99	0.90	0.87	0.87	
		Humanwaste treatment facilities	Gg N ₂ O	0.22	0.26	0.12	0.02	0.02	0.02	0.02	
		Degradation of domestic wastewater in nature	Gg N ₂ O	0.44	0.35	0.27	0.19	0.16	0.15	0.15	
	Total			Gg N ₂ O	4.18	4.04	3.92	3.76	3.74	3.65	3.65
			Gg CO ₂ eq	1,295	1,252	1,216	1,166	1,161	1,133	1,132	
Total of all gases				Gg CO ₂ eq	3,439	3,136	2,874	2,585	2,483	2,405	2,401

8.3.1. Industrial Wastewater (6.B.1.)

CH₄ and N₂O emissions from industrial effluent, which is treated by factories and other facilities in accordance with the regulations based on the Water Pollution Prevention Law and the Sewerage Law, are allocated to “Industrial wastewater treatment (6.B.1.a)” and CH₄ and N₂O emissions from landfill leachate treatment are allocated to “Landfill leachate treatment (6.B.1.b)” under the sub-category of “Industrial Wastewater (6.B.1)”.

8.3.1.1. Industrial Wastewater (6.B.1.a)

a) Source/Sink Category Description

CH₄ and N₂O emissions from industrial effluent, which is treated by factories and other facilities in accordance with the regulations based on the Water Pollution Prevention Law and the Sewerage Law, are allocated to “Industrial wastewater (6.B.1.a)”.

b) Methodological Issues

● Estimation Method

In accordance with the *GPG (2000)* decision tree, CH₄ and N₂O emissions were estimated for the industries that release organic-rich wastewater. Since default values given in the *Revised 1996 IPCC Guidelines* are considered to be unsuited to Japan's circumstances, CH₄ emissions were estimated based on Japan's country-specific methodology, namely, by multiplying the annual amount of organic matter in industrial wastewater subject to report (BOD basis) by the CH₄ emission factor per unit BOD that is based on Japan's country-specific wastewater handling. Because CH₄ is emitted in wastewater biological treatment processes, BOD-based activity data (amount of organic matter in wastewater degraded through biological treatment) is thought to be preferable to COD-based data. For this reason, CH₄ emissions are calculated using BOD in Japan. With regard to N₂O emissions, no estimation methodologies are given in the IPCC guidelines. Therefore, in the same manner for estimating CH₄ emissions, N₂O emissions were estimated by multiplying the amount of nitrogen in industrial wastewater by Japan's country-specific N₂O emission factor.

$$E = EF \times A$$

E : Amount of CH₄ or N₂O emissions generated when treating industrial wastewater (kg CH₄, kg N₂O)

EF : Emission factor (kg CH₄/kg BOD, kg N₂O/kg N)

A : Annual amount of industrial wastewater treated at wastewater treatment facilities (m³)

● *Emission Factor*

No research applicable to the circumstances in Japan has been found for the amounts of CH₄ and N₂O generated from the industrial wastewater treatments; therefore, emission factors were established by using with the ones used for the "Emissions from Treatment of Domestic and Commercial Wastewater (at sewage treatment plants) (6.B.2.a)", which were believed to be relatively similar to the CH₄ and N₂O generation processes in wastewater treatment.

Since the ones used in "6.B.2.a" are expressed in units of volume of wastewater treated (m³), these emission factors were converted to units per amount of organic matter (BOD basis) and nitrogen by dividing the emission factor by the following concentrations of organic matter (BOD basis) and nitrogen in the wastewater intake at sewage treatment plants.

For the BOD concentration of runoff water, the "Planned Runoff Water Quality of Municipal Solid Domestic Wastewater" (180 mgBOD/l) given in *Guidelines and Explanation of Sewerage Facility Design* (Japan Sewerage Works Association, 2001) was used.

For the nitrogen concentration of runoff water, 37.2 mg N/L was used, which was the simple average of total nitrogen concentrations of runoff water of sewage treatment plants obtained from the *Sewerage Statistics 2003 (Admin. Ed.)*.

CH₄ emission factor

= (CH₄ emission factor for emissions from domestic and commercial wastewater treatment & Sewerage treatment plant) / (BOD concentration in influent water)

= 8.8×10^{-4} (kg CH₄/m³) / 180 (mg BOD/L) × 1000

= 0.00489 ≈ 0.0049 (kg CH₄/kg BOD)

N₂O emission factor

$$\begin{aligned}
 &= (\text{N}_2\text{O emission factor for emissions from domestic and commercial wastewater treatment \& Sewage treatment plant}) / (\text{N concentration in influent water}) \\
 &= 1.6 \times 10^{-4} \text{ (kg N}_2\text{O/m}^3) / 37.2 \text{ (mg N/L)} \times 1000 \\
 &= 0.0043 \text{ (kg N}_2\text{O/kg N)}
 \end{aligned}$$

In Japan, CH₄ emissions generated by anaerobic wastewater treatment are entirely recovered. For a small amount of CH₄ emissions generated under partially anaerobic conditions created during aerobic treatment, a country-specific emission factor was applied for emission estimates because the condition for this particular CH₄ emissions differs from that for the use of default value for the CH₄ emissions generated from anaerobic treatment defined in *the 2006 IPCC Guidelines*.

● **Activity Data**

The activity data for CH₄ emission were estimated based on the amount of organic matter contained in wastewater using BOD concentrations. The emission estimates were conducted for the industries which generate large amount of CH₄ emissions with high BOD concentrations from the treatment of wastewater referring to the industry types provided in the *Revised 1996 IPCC Guidelines* (Table 8-15). The amount of organic matter was obtained by sorting and aggregating by industry type according to the middle industrial classification provided by the *Guidelines and Explanation of Sewage Facility Design* (Japan Sewage Works Association, 2001).

The use of COD concentrations is required to report activity data on CRF; however, activity data are reported as “NE” because country-specific methodology was used for this source.

CH₄ emission activity

$$\begin{aligned}
 &= \sum [(\text{Amount of industrial wastewater flowing into wastewater treatment facilities}) \times \\
 &(\text{Percentage of industrial wastewater treated at treatment facilities emitting CH}_4) \times (\text{Percentage of industrial wastewater treated on-site}) \times (\text{BOD concentration of runoff water})]
 \end{aligned}$$

The activity data for N₂O emissions were obtained based on the amount of nitrogen contained in industrial wastewater and aggregated by the same industrial sub-category as that applied to the estimation of CH₄ emissions.

N₂O emission activity

$$\begin{aligned}
 &= \sum [(\text{Amount of industrial wastewater flowing into wastewater treatment facilities}) \times \\
 &(\text{Percentage of industrial wastewater treated at treatment facilities emitting N}_2\text{O}) \times (\text{Percentage of industrial wastewater treated on-site}) \times (\text{Nitrogen concentration of runoff water})]
 \end{aligned}$$

Amount of industrial wastewater inflowed into wastewater treatment facilities

The amount of water used for the treatment of products by industrial sub-category and the volume of water used for washing given in the *Table of Industrial Statistics - Land and Water* (Ministry of Economy, Trade and Industry) were used for the amount of industrial wastewater treated at wastewater treatment facilities.

Percentage of industrial wastewater treated at facilities generating CH₄

Emissions of CH₄ from industrial wastewater treatment are believed to be generated from the treatment of wastewater with the activated sludge method and from the anaerobic treatment. Industrial wastewater treatment percentages for each industry code were set from the percentages of

reported wastewater amounts in total wastewater, as given under “active sludge”, “other biological treatment”, “membrane treatment”, “nitrification and denitrification” and “other advanced treatment” in the *Study on the Control of Burdens Generated* (Water and Air Environment Bureau, Ministry of the Environment).

Percentage of industrial wastewater treated at facilities generating N₂O

Emissions of N₂O from industrial wastewater treatment are believed to be generated mainly from biological treatment processes such as denitrification. Data on the fraction of industrial wastewater treated at facilities generating CH₄ was also used for N₂O emission estimates.

Percentage of industrial wastewater treated on-site

Percentage of industrial wastewater treated on-site is set at 1.0 in all industrial sub-categories because there is no statistical information available making it possible to ascertain this percentage.

BOD and nitrogen concentrations in runoff wastewater

For the BOD concentrations for industrial sub-categories, the BOD raw water quality for industrial sub-categories given in the *Guidelines and Analysis of Comprehensive Planning Surveys for the Provision of Water Mains, by Catchment Area 1999 Edition* (Japan Sewage Works Association) was used. For the nitrogen concentrations for industrial sub-categories, emission intensities (TN: Total Nitrogen) provided by the same survey for industrial sub-categories were used.

Table 8 - 15 BOD and nitrogen concentrations by industry type used for emission estimates

Industry code	Category of Manufacturing	mg BOD/l	mgN/l
9	Food manufacturing	1,467	62
10	Beverage, tobacco and feeding stuff manufacturing	1,138	77
11	Textile manufacturing	386	36
14	Pulp, paper and other paper manufacturing	556	37
16	Chemical industries	1,093	191
17	Petroleum products and coal product manufacturing	975	289
18	Plastic products manufacturing	268	11
19	Rubber products manufacturing	112	32
20	Chamois, chamois products and fur skin manufacturing	1,810	60

Table 8 - 16 BOD load (kt BOD) and TN load (kt N) of industrial wastewater

Item	Unit	1990	1995	2000	2005	2008	2009	2010
BOD load	kt BOD	1,075	1,046	1,032	1,000	1,004	970	970
TN load	kt N	89	87	76	89	94	90	90

c) Uncertainties and Time-series Consistency

● Uncertainties

The level of uncertainty in the CH₄ emission factor was evaluated on the basis of expert judgment. The uncertainty in activity data was estimated to be 37.4% on the basis of the uncertainties in the amount of wastewater used, percentage of industrial wastewater treated at CH₄-generating facilities, percentage of wastewater treated on-site, and BOD concentration in runoff water provided by each middle classification industry. The uncertainties in the amount of wastewater used, percentage of

industrial wastewater treated at facilities generating CH₄, and BOD concentration in runoff water were estimated by using statistical uncertainty. The uncertainty in the percentage of wastewater treated on-site was determined by expert judgment. The uncertainty level for N₂O is evaluated by the same method as was used for the CH₄ and estimated to be 300% and 51.1% for emission factor and activity data, respectively. The uncertainties in CH₄ and N₂O emissions from industrial wastewater handling were estimated to be 71% and 304%, respectively. For details, see the Annex 7.

● ***Time-series consistency***

Data on the percentage of industrial wastewater treated at CH₄- and N₂O-generating facilities since FY 2001 are available only for FY 2004. Therefore, data were interpolated and extrapolated for the remaining years. The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the GPG (2000). The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Recalculations

Updating BOD and TN load, emission estimates for FY2009 were recalculated.

f) Source-specific Planned Improvements

For future inventories, long-term efforts on further scientific investigations will be made to the following items:

- Improving the emission factors for emissions from industrial wastewater treatment for which currently the emission factors used for sewage treatment plants are substituted.
- Determining the amount of CH₄ recovery from industrial wastewater treatment

8.3.1.2. Landfill Leachate Treatment (6.B.1.b)

a) Source/Sink Category Description

CH₄ and N₂O emissions from landfill leachate treatment in MSW and ISW landfill sites are estimated and allocated to “Landfill leachate treatment (6.B.1.b)”.

b) Methodological Issues

● ***Estimation Method***

Potential BOD load (kgBOD/year) and TN load (kgN/year) to be remained in leachate percolated thorough organic waste disposed of in MSW and ISW landfill sites are applied for its activity data, and the methodology for the natural decomposition of domestic wastewater given in *the 2006 IPCC Guidelines* is applied to estimate CH₄ and N₂O emissions from this source as described below:

$$E = EF \times Li$$

E : CH₄ and N₂O emissions

EF : CH₄ and N₂O emission factor

L_i : Potential BOD load (kgBOD/year) and TN load (kgN/year) to be remained in leachate percolated thorough organic waste disposed of in MSW and ISW landfill sites

● Emission factors

Emission factors for CH₄ and N₂O are determined in accordance with the methodology for the natural decomposition of domestic wastewater given in *the 2006 IPCC Guidelines* as described below.

CH₄ Emission factor

According to *the 2006 IPCC Guidelines*, the emission factor for CH₄ is established by multiplying the maximum CH₄ generation potential (B₀) by a CH₄ conversion factor (MCF). The maximum CH₄ generation potential (B₀) is determined to be 0.6 kg CH₄/kg BOD which is a default value for “Domestic waste water” given in the *2006 IPCC Guidelines*, and MCF is determined to be 0.8 which is also a default value for “Anaerobic reactor” of “Treated systems” given in the *2006 IPCC Guidelines*.

$$EF_{CH_4} = B_0 \times MCF$$

$$= 0.6 \text{ (kg CH}_4\text{/kg BOD)} \times 0.8$$

$$= 0.48 \text{ (kg CH}_4\text{/kg BOD)}$$

B_0 : Maximum CH₄ generation potential (kgCH₄/kgBOD), IPCC default value:0.6

MCF : CH₄ conversion factor (IPCC default value: 0.8)

N₂O Emission Factor

The emission factor for N₂O is determined from a default value of 0.005 (kg N₂O-N/kg N) given in the *2006 IPCC Guidelines* after unit conversion.

$$EF_N = 0.005 \text{ (kg N}_2\text{O-N/kg N)} \times 44/28$$

$$= 0.0079 \text{ (kg N}_2\text{O/kg N)}$$

● Activity Data

Based on *the Survey Study on Improving the Accuracy of Emission Factors for Greenhouse Gas Emissions from the Waste Sector*, 2010, the Ministry of the Environment (Reference #15), the activity data for CH₄ and N₂O emission estimates are determined by establishing the ratio of organic and nitrogen contents to be remained in leachate for the amount of organic waste disposed of in MSW and ISW landfill sites to obtain potential BOD load (kgBOD/year) and TN load (kgN/year).

CH₄ activity data

$$L_{BODi} = F_{BOD} \times W \times T_i$$

L_{BODi} : Potential BOD load to be remained in leachate percolated thorough organic waste disposed of in MSW and ISW landfill sites (kgBOD/year)

F_{BOD} : Ratio of organic contents for the amount of organic waste landfilled (kgBOD/t) determined to be 0.188 (kgBOD/t) based on reference #15

W : Amount of organic waste landfilled with or without intermediate treatments including incineration ash (t/ year) obtained by *the Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes*

T_i : Ratio of leachate to be biologically treated in landfill site (%) determined to be 87.6% based on reference #15

N₂O activity data

$$L_{TNi} = F_{TN} \times W \times T_i$$

L_{TNi} : Potential TN load to be remained in leachate percolated thorough organic waste disposed of in MSW and ISW landfill sites (kgN/year)

F_{TN} : Ratio of nitrogen contents for the amount of organic waste landfilled (kgN/t) determined to be 0.254 (kgN/t) based on reference #15

W : Amount of organic waste landfilled with or without intermediate treatments including incineration ash (t/year) obtained by *the Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes*

T_i : Ratio of leachate to be biologically treated in landfill site (%) determined to be 87.6% based on reference #15

Table 8 - 17 BOD load (kt BOD) and NT load (ktN) for landfill leachate treatment

Item	Unit	1990	1995	2000	2005	2008	2009	2010
BOD load	kt BOD	2.6	2.5	2.2	1.6	1.3	1.2	0.9
NT load	kt N	3.5	3.3	3.0	2.2	1.7	1.6	1.2

c) Uncertainties and Time-series Consistency● **Uncertainties**

The level of uncertainty in the CH₄ emission factor was estimated by using the uncertainties in the maximum CH₄ generation potential and the CH₄ correction factor. The default value in the 2006 IPCC Guidelines was used for uncertainty in the N₂O emission factor. The uncertainties in activity data were evaluated for *tandoku-shori*, vault toilets, on-site disposal (determined from the wastewater treatment population and unit BOD or nitrogen in domestic wastewater) and ocean dumping (amount of human waste and septic tank sludge dumped into ocean, and concentration of organic matter or nitrogen in human waste and septic tank sludge). The methods of evaluation of the uncertainty levels for each component are:

- Use of the default values in the 2006 IPCC Guidelines: maximum CH₄ generation potential and CH₄ correction factor
- Based on expert judgment: unit BOD and nitrogen in domestic wastewater
- Use of 95% confidence interval of actual measurement data: concentrations of organic matter and nitrogen in human waste and septic tank sludge

- Use of the statistical uncertainties: wastewater treatment population, amount of human waste and septic tank sludge dumped into ocean

The uncertainties in CH₄ and N₂O emissions from natural decomposition of domestic wastewater were estimated to be 76%. For more details, see the Annex 7.

● *Time series consistency*

As described in detail in the preceding sections, emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials. For more details of QA/QC activities, see the Annex 6.

e) Source-specific Recalculations

Since emissions from landfill leachate treatment for the period FY1990-2010 are estimated based on new scientific findings, emission estimates for “Industrial Wastewater” (6.B.1) for the period FY1990-2009 were recalculated.

f) Source-specific Planned Improvements

No improvements are planned.

8.3.2. Domestic and Commercial Wastewater (6.B.2.)

Domestic and commercial wastewater generated in Japan is treated at various wastewater treatment facilities (e.g., sewage treatment plants, septic tanks, human-waste treatment plants) and greenhouse gas emissions from these sources are reported under “Domestic and Commercial Wastewater (6.B.2.)”. Because the CH₄ and N₂O emission characteristics differ from one wastewater treatment facility to another, a different emission estimation method is established for each facility.

The characteristics, effectiveness, and economic efficiency of wastewater treatment systems were thoroughly reviewed, and the most suitable systems were selected for each area in Japan with care also being taken to avoid excessive expenditure. As indicated in “*Waste Treatment in Japan*” (the Ministry of the Environment) public sewerage system is spreading from large cities to smaller municipalities and used by 68.9% of the population at the end of FY 2009.

Domestic wastewater treatment systems (e.g. *gappei shori jokasou*) are being promoted as an effective means of supplementing sewerage systems in smaller municipalities with low population densities and little flat land. In FY 2009, septic tanks (*jokasou*) were used by 22.4% of the population, with the remainder being treated after collection or on-site.

In CRF (6.B.2.), N₂O emissions from human waste treatment plants are reported in the subcategory “Human sewage (6.B.2.2)”, and other emissions are reported in “Domestic and Commercial (w/o human sludge) (6.B.2.1)”.

“NE” is reported on the CRF table for activity data instead of reporting the amount of organic carbon based on BOD values because the activity data for this source are estimated using a country-specific method by each gas and each wastewater treatment facility.

8.3.2.1. Sewage Treatment Plant (6.B.2.a)

a) Source/Sink Category Description

This category covers CH₄ and N₂O emissions from treatment of wastewater at sewage treatment plants.

b) Methodological Issues

● Estimation Method

Emissions of CH₄ and N₂O from this source were calculated using Japan's country-specific method in accordance with the decision tree of the *GPG (2000)* (Page 5.14, Fig. 5.2). Emissions were calculated by multiplying the volume of sewage treated at sewage treatment plants by the emission factor.

$$E = EF \times A$$

E : Amount of CH₄ or N₂O emitted from sewage treatment plants in conjunction with domestic/commercial wastewater treatment (kg CH₄, kg N₂O)

EF : Emission factor (kg CH₄/m³, kg N₂O/m³)

A : Yearly amount of sewage treated at a sewage treatment plant (m³)

● Emission Factors

Emission factors were established by adding the simple averages for each treatment process, having taken the actual volume of CH₄ and N₂O released from sludge treatment and water treatment processes measured at sewage treatment plants from research studies conducted in Japan (Refer to Reference #7).

CH₄ emission factor

= Average of emission factor for water treatment processes + Average of emission factor for sludge treatment processes = 528.7 [mg CH₄/m³] + 348.0 [mg CH₄/m³] = 8.764 × 10⁻⁴ [kg CH₄/m³]

N₂O emission factor

= Average of emission factor for water treatment processes + Average of emission factor for sludge treatment processes = 160.3 [mg N₂O/m³] + 0.6 [mg N₂O/m³] = 1.609 × 10⁻⁴ [kg N₂O/m³]

● Activity Data

Activity data for CH₄ and N₂O emissions associated with water treatment at sewage treatment plants was derived by subtracting the volumes subject to primary processing from the annual volume of water treated, as given in the *Sewage Statistics (Admin. Ed.)* (Japan Sewage Works Association).

In order to avoid overestimates of activity data, volumes subject to primary processing was subtracted from the annual volume of water treated because CH₄ and N₂O emitted from this source are primarily emitted from biological reaction tanks although the annual volume of water treated as given in the *Sewage Statistics (Admin. Ed.)* (Japan Sewage Works Association) includes primary treatment volumes that are only subject to settling.

Activity data

= (Annual volume of water treated at sewage treatment plants) – (Annual input volume for primary processing at sewage treatment plants)

Table 8 - 18 Activity data for wastewater treated at sewage treatment plant

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Annual amount of wastewater treated	10 ⁶ m ³	9,857	10,392	12,519	13,407	13,963	13,746	13,746

c) Uncertainties and Time-series Consistency

● *Uncertainties*

The uncertainties in CH₄ and N₂O emission factors were estimated by using the 95% confidence interval of actual measurement data. The uncertainty in activity data was evaluated based on the annual throughput and annual primary treatment amount and estimated by using the statistical uncertainties. The uncertainties in CH₄ and N₂O emissions from sewage treatment plants were estimated to be 33% and 146%, respectively. For details, see the Annex 7.

● *Time series consistency*

The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Recalculations

Updating the annual amount of treated wastewater, emission estimates for FY1990 were recalculated.

f) Source-specific Planned Improvements

A revision of the emission factor for sewage treatment plants is planned owing to the high uncertainty.

8.3.2.2. Domestic Sewage Treatment Plant (mainly septic tanks) (6.B.2.b)

a) Source/Sink Category Description

A part of domestic and commercial wastewater not processed in the public sewerage in Japan is processed in *community plants*, *gappei-shori johkasou*, the *tandoku-shori johkasou*, and vaults. The *gappei-shori* and *tandoku-shori* are decentralized wastewater treatment facilities installed at an individual home. The *gappei-shori* processes feces and urine and miscellaneous wastewater, whereas *tandoku-shori* processes only feces and urine. A community plant is small-scale sewage facility, where urine and the miscellaneous wastewater of each region are processed.

Table 8 - 19 Type of sewage and sewage treatment

Sewage treatment type		Sewage type
<i>Community plants</i>	Small-scale wastewater treatment facility regionally established	Human waste and miscellaneous wastewater
<i>Gappei-shori johkasou</i>	Wastewater treatment unit installed at an individual household	Human waste and miscellaneous wastewater
<i>Tandoku-shori johkasou</i>	Wastewater treatment unit installed at an individual household	Human waste
Vaults	Installed at an individual household	Human waste

This category covers CH₄ and N₂O emissions from domestic sewage treatment plants. Emissions from human waste within its residence time in vault toilets were accounted for under this category, whereas the emissions that occur after the waste is collected from vault toilets were accounted for under “Human waste treatment facilities (6.B.2.c)”.

b) Methodological Issues

● *Estimation Method*

Emissions of CH₄ and N₂O from this source were calculated using Japan’s country-specific method, in accordance with decision tree the *GPG (2000)* (Page 5.14, Fig. 5.2). Emissions were calculated by multiplying the annual population of treatment for each type of domestic sewage treatment plant by the emission factor.

$$E = \sum (EF_i \times A_i)$$

E : Emissions of methane and nitrous oxide from the processing of domestic and commercial wastewater at domestic sewage treatment plants (i.e. household septic tanks) (kg CH₄, kg N₂O)

EF_i : Emission factor for domestic sewage treatment plant i (kg CH₄/person, kg N₂O/person)

A : Population (persons) requiring waste processing at domestic sewage treatment plant i per year

● *Emission Factors*

The CH₄ and N₂O emission factors for this source were determined as described below:

- For the CH₄ emission factor for community plants by FY1995, the values indicated in Tanaka, (1998) were used. For the values from FY2005 onwards, the values indicated in Souda (2010) were used taking into account the performance improvement in the plants. The values for FY1996 through FY2004 were interpolated.

- For the N₂O emission factor for community plants by FY1995, the mean values of the upper limit and the lower limit of actual measured values indicated in Tanaka (1997) were used. For the values from FY2005 onwards, the values indicated in Ike and Souda (2010) were used taking into account the performance improvement of the plants. The values for FY1996 through FY2004 were interpolated.

- For the CH₄ and N₂O emission factors for gappei-shori johkasou, the mean values of the upper limit and the lower limit of actual measured values indicated in Tanaka et al. (1998) were used.

- For the CH₄ and N₂O emission factors for tandoku-shori johkasou, the mean value of the upper limit and the lower limit of actual measured values indicated in Takeishi et al., (1993), and Takeishi et al.,

(1994) were used.

- For the CH₄ and N₂O emission factors for vault toilets, the same values as that used for tandoku-shori johkasou were applied because the detention period of human waste is very similar.

Table 8 - 20 CH₄ Emission factors for domestic sewage treatment plants

Item	CH ₄ Emission factor [kg CH ₄ /person-year]		
	FY 1990-1995	FY 1996-2004	FY2005-
Community plants	0.195	Calculated by interpolation using the values of FY1995 and FY 2005	0.062
<i>Gappei-shori johkasou</i>	1.106		
<i>Tandoku-shori johkasou</i>	0.197		
Vault toilets	0.197		

Table 8 - 21 N₂O emission factor for domestic sewage treatment plants

Item	N ₂ O Emission factor [kg N ₂ O-N// person-year]		
	FY 1990-1995	FY 1996-2004	FY2005-
Community plants	0.0394	Calculated by interpolation using the values of FY1995 and FY 2005	0.0048
<i>Gappei-shori johkasou</i>	0.0264		
<i>Tandoku-shori johkasou</i>	0.0200		
Vault toilets	0.0200		

● Activity Data

Annual treatment population by type of domestic sewage treatment plant for community plants, *gappei-shori johkasou*, *tandoku-shori johkasou*, and vault toilets given in the *Waste Treatment in Japan* was used as the activity data for CH₄ and N₂O emitted in association with domestic wastewater treatment facilities.

Table 8 - 22 Annual treatment population by type of domestic sewage treatment plant (1,000 persons)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Community plant	1000 person	493	398	414	554	416	297	297
<i>Gappei-shori johkasou</i>	1000 person	7,983	8,515	10,806	12,770	13,854	13,792	13,792
<i>Tandoku-shori johkasou</i>	1000 person	25,119	26,105	23,289	18,303	15,413	14,712	14,712
Vault toilet	1000 person	38,920	29,409	20,358	13,920	11,301	10,671	10,671
Total	1000 person	72,515	64,427	54,867	45,547	40,984	39,472	39,472

c) Uncertainties and Time-series Consistency

● Uncertainties

The level of uncertainty in the emission factor was evaluated for each treatment facility taking into account the actual measurement data and setting methods. The following data were used:

- The 95% confidence interval of actual measurement data: *gappei-shori* (N₂O) and *tandoku-shori* (CH₄ and N₂O)
- The upper and lower limits of actual measurement data: community plants (CH₄) and *gappei-shori*

(CH₄)

- The values set by the Committee for GHGs Emissions Estimation Methods: community plants (N₂O) and vault toilets (CH₄ and N₂O)

The uncertainty in activity data was evaluated based on the uncertainties in treatment population for each type of treatment facilities by using the statistical uncertainty (10%). The uncertainties in CH₄ and N₂O emissions from domestic wastewater treatment (mainly septic tanks) were estimated to be 87% and 72%, respectively. For details, see the Annex 7.

● *Time series consistency*

The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Recalculations

No recalculations were conducted.

f) Source-specific Planned Improvements

No improvements are planned.

8.3.2.3. Human-Waste Treatment Plant (6.B.2.c)

a) Source/Sink Category Description

This category covers emissions of CH₄ and N₂O emissions from treatment of vault toilet human waste and septic tank sludge collected at human waste treatment plants.

b) Methodological Issues

1) CH₄

● *Estimation Method*

Emissions of CH₄ from this source were calculated using Japan's country-specific methodology in accordance with decision tree of the *GPG (2000)* (Page 5.14, Fig. 5.2). Emissions were calculated by multiplying the volume of domestic wastewater treated at human waste treatment plants by the emission factor.

$$E = \sum (EF_i \times A_i)$$

E : Emission of methane from the processing of domestic and commercial wastewater at human waste treatment plants (kg CH₄)

EF_i : Emission factor for human waste treatment plants (for treatment process i) (kg CH₄/m³)

A_i : Input volume of human waste and septic tank sludge at human waste treatment plants (for treatment process i) (m³)

● Emission factors

Emission factors for CH₄ were determined by treatment processes type, including anaerobic, aerobic, standard denitrification and high-load denitrification treatments as well as membrane separation systems, for each of the human waste treatment plants (Refer to Reference #7).

Table 8 - 23 CH₄ emission factors by each treatment process

Treatment method	CH ₄ emission factor [kg CH ₄ /m ³]	Data source
Anaerobic treatment	0.543	Estimated by multiplying the actual methane emissions given in Reference #36 by the value of 1 – CH ₄ recovery rate (90%).
Aerobic treatment	0.00545	Simple average value of standard de-nitrification and high-load de-nitrification since actual data on emissions is not available.
Standard de-nitrification treatment	0.0059	Reference #63
High load de-nitrification treatment	0.005	Reference #63
Membrane separation	0.00545	Because the current status of its emissions is not identified, substituted the emission factor for aerobic treatment.
Other	0.00545	Because the current status of its emissions is not identified, substituted the emission factor for aerobic treatment.

● Activity Data

Activity data for CH₄ emissions associated with the processing of wastewater at human waste treatment plants was determined from the calculated throughput volume for each of the treatment processes (Table 8-24), by multiplying the total volume of human waste and septic tank sludge processed at human waste treatment plants that were indicated in *Waste Treatment in Japan* (Table 8-25) by the capacity of each treatment process (Table 8-26).

$$\text{Activity data for human waste treatment method } i \\ = (\text{Total amount of human waste and septic tank sludge by treatment method } i) \times \\ (\text{Capacity of waste treatment method } i) / (\text{Total capacity of all waste treatment methods})$$

Table 8 - 24 Volume of human waste and septic tank sludge treated at their treatment plants

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Vault toilet	1000 kl/year	20,406	18,049	14,673	10,400	8,894	8,353	8,353
ST sludge	1000 kl/year	9,224	11,545	13,234	13,790	14,064	13,989	13,989
Total	1000 kl/year	29,630	29,594	27,907	24,190	22,958	22,342	22,342

Source: Waste Treatment in Japan

Table 8 - 25 Trends in treatment capacity by treatment process

Unit	Unit	1990	1995	2000	2005	2008	2009	2010
Anaerobic treatment	kl/day	34,580	19,869	10,996	6,476	4,444	4,144	4,144
Aerobic treatment	kl/day	26,654	19,716	12,166	8,465	7,535	6,961	6,961
Standard denitrification	kl/day	25,196	30,157	31,908	29,655	27,737	27,748	27,748
High-intensity denitrification	kl/day	8,158	13,817	16,498	17,493	14,938	16,285	16,285
Membrane separation	kl/day	0	1,616	2,375	3,055	3,650	3,573	3,573
Other	kl/day	13,777	20,028	25,917	30,277	35,441	34,654	34,654

Table 8 - 26 Activity Data for human waste by treatment type

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Anaerobic treatment	1000 kl/year	9,455	5,589	3,073	1,642	1,088	992	992
Aerobic treatment	1000 kl/year	7,288	5,546	3,400	2,146	1,845	1,666	1,666
Standard denitrification	1000 kl/year	6,889	8,483	8,917	7,518	6,793	6,640	6,640
High-intensity denitrification	1000 kl/year	2,231	3,887	4,611	4,435	3,658	3,897	3,897
Membrane separation	1000 kl/year	0	455	664	774	894	855	855
Other	1000 kl/year	3,767	5,634	7,243	7,676	8,679	8,293	8,293
Total	1000 kl/year	29,630	29,594	27,907	24,190	22,958	22,342	22,342

2) N_2O ● *Estimation Method*

Emissions of N_2O from this source were calculated using Japan's country-specific methodology, in accordance with decision tree of the *GPG (2000)* (Page 5.14, Fig. 5.2). Emissions were calculated by multiplying the volume of nitrogen treated at human waste treatment plants, by the emission factor.

$$E = \sum (EF_i \times A_i)$$

E : Emission of nitrous oxide from the processing of domestic and commercial wastewater at human waste treatment plants (kg N_2O)

EF_i : Emission factor for human waste treatment plants (by treatment process i) (kg N_2O /kg N)

A_i : Amount of nitrous oxide in human waste and septic tank sludge input at human waste treatment plants (by treatment process i) (kg N)

● *Emission factors*

The emission factors for N_2O were determined for each treatment process including high-load denitrification treatment and membrane separation systems using the results of actual case studies in Japan (Refer to Reference #7).

According to the survey study on the emission factors for human waste treatment facilities conducted in FY1994 (Tanaka et al., 1997) and FY2003 (Ohmura et al., 2004) in Japan, because of the advancement of the structure of human waste treatment facilities and the technology of operation and maintenance, actual measurement results show the improvement in the emission factors for high load de-nitrification treatment and membrane separation; therefore, different emission factors were used for FY1994 or before and from FY2003 onwards.

Table 8 - 27 Nitrous oxide emission factors by each treatment process

Treatment method	N_2O emission factors [kg N_2O -N/kg-N]		
	FY1990-1994	FY1995-2002	FY2003 -
High load de-nitrification treatment	0.033 ^a	Calculated by interpolation using the values of FY1994 and FY 2003	0.0029 ^b
Membrane separation	0.033 ^a	Calculated by interpolation using the values of FY1994 and FY 2003	0.0024 ^b
Other (including anaerobic treatment, aerobic treatment, standard de-nitrification treatment)	0.0000045 ^c		

a) Use median value of actual measurements at 13 plants given in Reference #64

b) Use median value of actual measurements at 13 plants given in Reference #57

c) Referred to Reference #63

Note: Calculated by dividing upper limit value for standard de-nitrification treatment (0.00001kg N_2O /m³) by treated

nitrogen concentration in FY1994 (2,211mg/L).

● Activity Data

The volume of nitrogen treated at human waste treatment plants was calculated by multiplying treated nitrogen concentration by the volume of human waste treated at these facilities (the sum of collected human waste and sewage in sewerage tank), given in the *Waste Treatment in Japan*. The treated nitrogen concentration is based on weighted average of the volume of nitrogen contained in collected human waste and sewage in sewerage tank derived using the volume of collected human waste and sewage in sewerage tank treated at human waste treatment plants.

Activity data

$$= [(\text{Input volume of human waste at human waste treatment plants}) \times (\text{Nitrogen concentration in human waste}) + (\text{Input volume of septic tank sludge at human waste treatment plants}) \times (\text{Nitrogen concentration in septic tank sludge})] \times (\text{percentage throughput of treatment process } i)$$

Input volume of human waste and septic tank sludge at human waste treatment plants:

See the data used for the calculation of CH₄ emissions from human waste treatment plants (Table 8-24).

Percentage throughput of the human waste treatment processes:

See the data used for the calculation of CH₄ emission from human waste treatment plants (Table8-25).

Nitrogen concentration in human waste and septic tank sludge input at treatment plants:

For the nitrogen concentration in human waste and septic tank sludge input at treatment plants, the values analyzed for the period FY 1989 - FY 1991, FY 1992 - FY 1994, FY1995 – FY1997, and FY 1998 - FY 2000, respectively, were used based on the research conducted by Okazaki (2001). The value of FY 2000 was substituted for the values from FY 2001 onward. (See Table 8-28).

Table 8 - 28 Concentration of nitrogen contained in collected human waste and sewage in sewerage tank

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Vault toilet	mg N/l	3,940	3,100	2,700	2,700	2,700	2,700	2,700
ST sludge	mg N/l	1,060	300	580	580	580	580	580
Weighted average	mg N/l	3,043	2,008	1,695	1,491	1,401	1,373	1,373

Table 8 - 29 Activity data:

Amount of nitrogen in human waste processed at human waste treatment plants and septic tank sludge

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Anaerobic treatment	kt N	28.8	11.2	5.2	2.4	1.5	1.4	1.4
Aerobic treatment	kt N	22.2	11.1	5.8	3.2	2.6	2.3	2.3
Standard denitrification	kt N	21.0	17.0	15.1	11.2	9.5	9.1	9.1
High-intensity	kt N	6.8	7.8	7.8	6.6	5.1	5.3	5.3
Membrane separation	kt N	0	0.9	1.1	1.2	1.3	1.2	1.2
Other	kt N	11.5	11.3	12.3	11.4	12.2	11.4	11.4
Total	kt N	90.2	59.4	47.3	36.1	32.2	30.7	30.7

c) Uncertainties and Time-series Consistency

● *Uncertainties*

The level of uncertainty in the CH₄ emission factor was evaluated by using the default values set by the Committee for GHGs Emissions Estimation Methods for each type of human waste treatment method (anaerobic treatment, aerobic treatment, standard denitrification, high-intensity denitrification, membrane separation, and other). The uncertainty in the activity data for CH₄ is associated with uncertainties in the amount of human waste and septic tank sludge that entered human waste treatment facilities and the throughput capacity rate by type of human waste treatment. The uncertainties for each component were estimated by using the statistical uncertainties. The uncertainty level in N₂O emission factors was also evaluated by treatment type. For high-intensity denitrification and membrane separation, the 95% confidence interval of actual measurement data on emission factors was used. For other treatments, the default values set by the Committee for GHGs Emissions Estimation Methods were used. The uncertainty in activity data for N₂O was estimated by using the uncertainties in nitrogen concentration in human waste and septic tank sludge that determined from the standard deviations in actual measurement data, in addition to the components of uncertainty for CH₄. The uncertainties in CH₄ and N₂O emissions from human waste treatment were estimated to be 101% and 106%, respectively. For details, see the Annex 7.

● *Time series consistency*

For N₂O emission factor, consistent data over the time series were constructed based on the actual measurement data by using the methods described in Table 8-27. For other parameters, data were constructed consistently for the entire time series. The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the GPG (2000). The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Planned Improvements

No recalculations were conducted.

f) Source-specific Planned Improvements

No improvements are planned.

8.3.2.4. Emission from the Natural Decomposition of Domestic Wastewater (6.B.2.d)

a) Source/Sink Category Description

Although most of the domestic wastewater generated by Japanese households is processed at wastewater treatment plants, some is discharged untreated into public waters. The amounts of CH₄ and N₂O decomposes and emitted from this source are reported under this category.

b) Methodological Issues

● Estimation Method

Estimation method was established in accordance with the method described in the 2006 IPCC Guidelines. In the natural decomposition of wastewater, both the volume of organic matter extracted as sludge and recovered CH₄ were zero. Accordingly, CH₄ emissions were calculated by multiplying the volume of organic matter contained in the untreated domestic wastewater that was discharged into public waters by the emission factor. The N₂O emission was calculated by multiplying the volume of nitrogen contained in the wastewater by the emission factor.

$$E = EF \times A$$

E : Emission of methane or nitrous oxide from the natural decomposition of domestic wastewater (kg CH₄; kg N₂O)

EF : Emission factor (kg CH₄/kg BOD; kg N₂O/kg N)

A : Volume of organic matter (kg BOD) or nitrogen (kg N) in domestic wastewater

● Emission factors

Emission factors were determined in accordance with the 2006 IPCC Guidelines. The emission factor for CH₄ was established by multiplying the maximum CH₄ generation potential (*B*₀) by a CH₄ conversion factor (MCF). The maximum CH₄ generation potential was set to 0.6 kg CH₄/kg BOD, given in the 2006 IPCC Guidelines, and the MCF was set to 0.1, a default value for “Sea, river and lake discharge” of “Untreated systems”.

$$\begin{aligned} EF_{\text{CH}_4} &= B_0 \times \text{MCF} \\ &= 0.6 \text{ (kg CH}_4\text{/kg BOD)} \times 0.1 \\ &= 0.06 \text{ (kg CH}_4\text{/kg BOD)} \end{aligned}$$

The emission factor for N₂O was calculated from the value of 0.005 kg N₂O-N/kg N after conversion of the units.

$$\begin{aligned} EF_{\text{N}_2\text{O}} &= 0.005 \text{ (kg N}_2\text{O-N/kg N)} \times 44/28 \\ &= 0.0079 \text{ (kg N}_2\text{O/kg N)} \end{aligned}$$

● Activity Data

Activity data to be calculated are the following sources:

- Domestic wastewater from households using tandoku-shori johkasou
- Domestic wastewater from households using Vault toilets
- Domestic wastewater from households using on-site disposal systems
- Human waste and septic tank sludge dumped into the ocean
- Sewage sludge dumped into the ocean

Definition for each activity data is provided as in Table 8-30. Estimated activity data are shown in Table 8-31.

Table 8 - 30 Calculation method for activity data used for the calculation of GHG emissions from the natural decomposition of domestic wastewater

Item	Methane emission activity data	Nitrous oxide emission activity data
<i>Tandoku-shori johkasou</i>	User population (persons) × Unit BOD from domestic wastewater (g BOD/person-day)	User population (persons) × Unit nitrogen from domestic wastewater (g N/person-day)
Vault toilet		
On-site disposal *	Population using on-site disposal system (person) × Unit BOD from domestic wastewater (g BOD/person-day)	Population using on-site disposal system (person) × Unit nitrogen from domestic wastewater (g N/person-day)
Ocean dumping (Human waste)	Human waste dumped in ocean (kL) × BOD concentration in human waste (mg BOD/L) + septic tank sludge dumped in ocean (kL) × BOD concentration in septic tank sludge (mg BOD/L)	Human waste dumped in ocean (kL) × nitrogen concentration in septic tank sludge (mg N/L) + septic tank sludge dumped in ocean (kL) × nitrogen concentration in septic tank sludge (mg N/L)
Ocean dumping (Sewage sludge)	Sewage sludge dumped in ocean (kL) × BOD concentration in sewage sludge (mg BOD/L)	Sewage sludge dumped in ocean (kL) × nitrogen concentration in sewage sludge (mg N/L)

Source:

- Volumes for *tandoku-shori johkasou*, vault toilets, on-site disposal systems and ocean dumping: Reference #8
- Unit BOD and unit nitrogen from domestic wastewater: Reference #43
- BOD concentration and nitrogen concentration in human waste and septic tank sludge: Reference #56

* A portion of the human waste in on-site disposal systems is utilized as fertilizer on farmlands in Japan. The nitrous oxide emission from this portion of human waste is already included in the “Direct emission from soil (4.D.)” category in the Agriculture section, and therefore, not included in the calculation for this source.

Table 8 - 31 Activity data: Amount of organic material and nitrogen in domestic wastewater untreated and discharged into public water body

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Tandoku-shori	kt BOD	366.7	381.1	341.0	267.2	225.6	214.8	214.8
Vault toilet	kt BOD	568.2	429.4	298.0	203.2	165.4	155.8	155.8
On-site disposal	kt BOD	46.2	21.0	9.4	3.9	7.6	2.0	2.0
Ocean dumping (Human waste)	kt BOD	21.7	13.5	9.3	3.5	0	0	0
Ocean dumping (sewage sludge)	kt BOD	0.8	0.9	0.0	0	0	0	0
Total	kt BOD	1,002.9	845.1	657.7	477.8	398.7	372.6	372.6

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Tandoku-shori	kt N	18.3	19.1	17.0	13.4	11.3	10.7	10.7
Vault toilet	kt N	28.4	21.5	14.9	10.2	8.3	7.8	7.8
On-site disposal	kt N	2.3	1.1	0.5	0.2	0.4	0.1	0.1
Ocean dumping (Human waste)	kt N	7.2	3.2	2.2	0.8	0	0	0
Ocean dumping (sewage sludge)	kt N	0.1	0.1	0.0	0	0	0	0
Total	kt N	56.3	44.7	34.6	24.5	19.9	18.6	18.6

c) Uncertainties and Time-series Consistency

● Uncertainties

The level of uncertainty in the CH₄ emission factor was estimated by using the uncertainties in the

maximum CH₄ generation potential and the CH₄ correction factor. The default value in the 2006 IPCC Guidelines was used for uncertainty in the N₂O emission factor. The uncertainties in activity data were evaluated for *tandoku-shori*, vault toilets, on-site disposal (determined from the wastewater treatment population and unit BOD or nitrogen in domestic wastewater) and ocean dumping (amount of human waste and septic tank sludge dumped into ocean, and concentration of organic matter or nitrogen in human waste and septic tank sludge). The methods of evaluation of the uncertainty levels for each component are:

- Use of the default values in the 2006 IPCC Guidelines: maximum CH₄ generation potential and CH₄ correction factor
- Based on expert judgment: unit BOD and nitrogen in domestic wastewater
- Use of 95% confidence interval of actual measurement data: concentrations of organic matter and nitrogen in human waste and septic tank sludge
- Use of the statistical uncertainties: wastewater treatment population, amount of human waste and septic tank sludge dumped into ocean

The uncertainties in CH₄ and N₂O emissions from natural decomposition of domestic wastewater were estimated to be 76%. For more details, see the Annex 7.

● *Time series consistency*

The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Recalculations

No recalculations were conducted.

f) Source-specific Planned Improvements

No improvements are planned.

8.3.2.5. Recovery of CH₄ emitted from treating domestic and commercial wastewater (6.B.2.-)

a) Source/Sink Category Description

In Japan, CH₄ emissions generated from sludge digestion at sewage treatment plants and human waste treatment facilities are recovered.

CH₄ emissions generated by anaerobic wastewater treatment are entirely recovered. A small amount of CH₄ emission generated under aerobic conditions is estimated with a country-specific emission factor. These recovered CH₄ emissions treating domestic and commercial wastewater explained in this section are not estimated by the methodology indicated in the *GPG (2000)* and not included in emission estimates.

Therefore, for reference purpose only, the amount of CH₄ recovered treating domestic and commercial wastewater at sewage treatment plants and human waste treatment facilities are reported in this section.

b) Methodological Issues

1) Methane Recovery at Sewage Treatment Plants

● Estimation Method

The amount of CH₄ recovered from sludge digesters at sewage treatment plants is calculated by multiplying the amount of digester gas (volumetric basis) recovered from digesters by an emission factor that takes into account the concentration of CH₄ in digester gas.

$$R = A \times EF$$

R : Amount of recovered CH₄ at final disposal site (Gg CH₄)

A : Amount of generated digester gas (m³)

EF : Emission factor (Gg CH₄/m³)

● Emission factors

Emission factor is set by finding the weight equivalent of the average CH₄ concentration in digester gas.

$$EF = F_{CH_4} \times 16 / 22.4$$

EF : Emission factor (Gg CH₄/m³)

F_{CH_4} : Concentration of methane in digester gas (volumetric basis)

The CH₄ concentration in digester gas (volumetric basis) was set at 60% with reference to the *Manual for Developing Plans for Biosolids Utilization (Draft)* (Ministry of Land, Infrastructure, Transport and Tourism).

● Activity Data

The amount of digester gas recovered from sludge digesters at sewage treatment plants is provided by “amount of digester gas generated by sludge treatment facilities” in the *Sewage Statistics (Admin. Ed.)* (Japan Sewage Works Association). Because entire digester gas generated at sewage treatment plants in Japan is recovered, the total amount of generated digester gas is treated as the amount of digester gas recovered. The amount of digester gas used for energy to be included in the energy category is determined from the amount of digester gas listed in “amount of digester gas used in sludge digester facilities” of the *Sewerage Statistics*.

Table 8 - 32 Amount of CH₄ recovered from sewage treatment plant sludge digesters (Gg- CH₄)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Recovered CH ₄ amount	Gg CH ₄	88.7	110.5	113.3	122.0	130.3	130.2	130.2
Portion used as energy	Gg CH ₄	65.3	73.9	75.3	85.0	93.2	92.4	92.4

2) Methane Recovery from Human Waste Treatment Facilities

● Estimation Method

The amount of CH₄ recovery at human waste treatment facilities was obtained by multiplying the amount of recycled biogas at human waste treatment facilities on a volumetric basis by the emission factor taking into account CH₄ concentration in biogas.

$$R = A \times EF$$

- R : Amount of CH₄ recovered at human waste treatment facilities (Gg CH₄)
 A : Amount of Recycled Biogas (m³)
 EF : Emission Factor (Gg CH₄/m³)

● Emission Factors

Emission factor was determined by taking into account CH₄ concentration in biogas and molecular weight conversion. CH₄ concentration in biogas was determined to be 60% referring to the *JARUS Reference System for Information of Biomass Recycling Technology* (The Japan Association of Rural Resource Recycling Solutions). Because statistical data are aggregated on a volumetric basis, they are converted into molecular weight given the average temperature at the facilities is 18°C.

$$EF = F_{CH_4} \times 16 / 22.4 \times 273 / (273 + 18)$$

- EF : Emission factor (Gg CH₄/m³)
 F_{CH_4} : CH₄ concentration in biogas (volumetric basis)

● Activity Data

For the activity data on CH₄ recovery at human waste treatment facilities, the aggregated amount of recycled biogas at human waste treatment facilities (volumetric basis) provided by *the State of Municipal Waste Treatment Survey*, Ministry of the Environment, Waste Management and Recycling Department was used. The statistical data before FY2005 are not obtained. Therefore, the emissions for FY2004 and before were estimated by applying the amount of CH₄ actually recovered in FY 2005 and in the year that facilities started their operation provided by this survey and in FY 2005, and also using the amount of human waste (vault toilet) and septic tank sludge treated at the facilities for FY 2004 and before.

Table 8 - 33 Amount of CH₄ recovered at human waste treatment facilities

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Recovered CH ₄ amount	Gg CH ₄	0.3	0.5	0.8	0.9	1.6	1.7	1.7

c) Uncertainties and Time-series Consistency

● Uncertainties

The assessment was not conducted, as the amount of CH₄ recovered is reported as a reference value.

● Time series consistency

The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Recalculations

No recalculations were made.

f) Source-specific Planned Improvements

No improvements are planned.

8.4. Waste Incineration (6.C.)

In Japan, waste disposed of has been reduced in volume primarily by incineration. Emissions from waste incineration are categorized as shown in Table 8-34. CO₂, CH₄, N₂O emissions without energy recovery are allocated to this category. Also, waste incineration includes the following practices of waste used as raw material or fuel:

- Energy recovery from waste incineration
- Waste material is used directly as fuel
- Waste material is converted into fuel

Estimated emissions from the sources listed above are allocated to the “Fuel Combustion (Category 1.A.)” in accordance with the Revised 1996 IPCC Guidelines and the GPG (2000).

In order to avoid double-counting or any other confusion, emissions from the categories indicated in Table 8-34 with or without energy use were estimated collectively under the waste sector, thus the estimation methodology for these categories are provided in this section.

Table 8 - 34 Categories for the calculation of emissions from waste incineration (6.C.)

Incineration	Waste category	Estimation classification	Category to be allocated to	CO ₂	CH ₄	N ₂ O
Waste incineration (without energy recovery)	Municipal solid waste	Plastic	6.C.1	○	○ Estimated in bulk	○ Estimated in bulk
		Synthetic textile	6.C.1	○		
		Other (biogenic)	6.C.1	/		
	Industrial solid waste	Waste oil	6.C.2	○	○	○
		Waste plastic	6.C.2	○	○	○
		Other (biogenic)	6.C.2	/	○	○
	Specially controlled industrial waste	Waste oil	6.C.3	○	○	○
		Infectious waste (plastic)	6.C.3	○	○	○
		Infectious waste except plastic (biogenic)	6.C.3	/	○	○
Waste incineration with energy recovery	Municipal solid waste	Plastic	1.A.1	○	○ Estimated in bulk	○ Estimated in bulk
		Synthetic textile	1.A.1	○		
		Other (biogenic)	1.A.1	/		
	Industrial solid waste	Waste oil	1.A.1	○	○	○
		Waste plastic	1.A.1	○	○	○
		Other (biogenic)	1.A.1	/	○	○
Direct use of waste as fuel	Municipal solid waste	Plastic	1.A.1/2	○	○	○
	Industrial solid waste	Waste oil	1.A.2	○	○	○
		Waste plastic	1.A.2	○	○	○
		Waste wood	1.A.2	/	○	○
	Waste tire	Fossil origin	1.A.1/2	○	○	○
Biogenic origin		1.A.1/2	/			
Use of waste processed as fuel	Refuse derived fuel (RDF·RPF)	Fossil origin	1.A.1/2	○	○	○
		Biogenic origin	1.A.1/2	/		

a. CO₂ emissions from the incineration of biomass-derived waste (including biomass-based plastics and waste animal and

vegetable oil) is not included in the total emissions in accordance with the Revised 1996 IPCC Guidelines; instead it is estimated as a reference value and reported under "Biogenic" in Table 6.A,C of the CRF.

Estimated greenhouse gas emissions from waste incineration (category 6.C.) are shown in Table 8-35. In FY 2010, emissions from waste incineration were 14,356 Gg-CO₂ eq. and accounted for 1.1% of the national total emissions (excluding LULUCF). The emissions from this source category increased by 4.1% compared to those in FY 1990. For the period FY1990-FY1997, CO₂ emissions increased as the practice of intermediate treatment by waste incineration increased in order to decrease the total volume of waste landfilled. From FY2001 onwards, as the use of waste as raw material or fuel has been replacing the incineration of fossil-origin waste for intermediate treatments, and these CO₂ emissions which used to be allocated to the waste sector is now allocated to the Energy sector, CO₂ emission estimates from the waste sector decreased.

On the other hand, N₂O emissions increased compared to FY1990 level due to the increase in sewage sludge incineration practice for the period FY 1990 - FY1997. From FY2005 onward, N₂O emissions from this source decreased because the practice of high temperature incineration of sewage sludge increased.

Table 8 - 35 GHG emissions from waste incineration (6.C.)

Gas	Waste category	Estimation Category	Unit	1990	1995	2000	2005	2008	2009	2010	
CO ₂	Municipal solid waste	Plastics	Gg CO ₂	5,041	5,031	5,222	3,060	2,305	2,709	2,617	
		Synthetic textiles	Gg CO ₂	503	539	421	428	548	571	600	
		Other (biogenic)	Gg CO ₂								
	Industrial solid waste	Waste oil	Gg CO ₂	3,652	4,344	4,775	4,249	4,616	3,739	4,013	
		Waste plastics	Gg CO ₂	2,120	4,516	4,358	4,311	4,874	3,539	3,630	
		Other (biogenic)	Gg CO ₂								
	Specially controlled waste	Waste oil	Gg CO ₂	748	1,110	1,636	1,504	1,647	1,334	1,431	
		Infectious plastics	Gg CO ₂	198	327	426	433	492	357	366	
		Infectious waste (except plastics; biogenic)	Gg CO ₂								
	Total			Gg CO ₂	12,263	15,867	16,838	13,984	14,481	12,249	12,658
CH ₄	Municipal solid waste		Gg CH ₄	0.464	0.431	0.381	0.064	0.059	0.060	0.060	
	Industrial solid waste	Waste oil	Gg CH ₄	0.006	0.007	0.008	0.006	0.007	0.006	0.006	
		Waste plastics	Gg CH ₄	0.025	0.053	0.051	0.014	0.015	0.011	0.011	
		Other (biogenic)	Gg CH ₄	0.140	0.207	0.181	0.541	0.427	0.384	0.374	
	Specially controlled waste	Waste oil	Gg CH ₄	0.001	0.002	0.003	0.002	0.002	0.002	0.002	
		Infectious plastics	Gg CH ₄	0.002	0.004	0.005	0.001	0.002	0.001	0.001	
		Infectious waste (except plastics; biogenic)	Gg CH ₄	0.002	0.004	0.005	0.051	0.058	0.042	0.043	
	Total			Gg CH ₄	0.642	0.708	0.635	0.679	0.571	0.507	0.498
	Total			Gg CO ₂ eq	13.481	14.868	13.333	14.267	11.981	10.639	10.457
	N ₂ O	Municipal solid waste		Gg N ₂ O	1.025	1.049	0.979	0.525	0.473	0.485	0.485
Industrial solid waste		Waste oil	Gg N ₂ O	0.015	0.018	0.021	0.098	0.107	0.087	0.093	
		Waste plastics	Gg N ₂ O	0.149	0.318	0.307	0.025	0.028	0.021	0.021	
		Other (biogenic)	Gg N ₂ O	3.692	5.074	5.943	6.062	4.943	4.794	4.798	
Specially controlled waste		Waste oil	Gg N ₂ O	0.003	0.005	0.007	0.032	0.035	0.028	0.031	
		Infectious plastics	Gg N ₂ O	0.014	0.023	0.030	0.003	0.003	0.002	0.002	
		Infectious waste (except plastics; biogenic)	Gg N ₂ O	0.002	0.004	0.005	0.018	0.020	0.014	0.015	
Total			Gg N ₂ O	4.901	6.491	7.290	6.762	5.609	5.431	5.444	
Total			Gg CO ₂ eq	1.519	2.012	2.260	2.096	1.739	1.684	1.688	
Total of all gases			Gg CO ₂ eq	13,796	17,894	19,111	16,095	16,232	13,943	14,356	

* CO₂ emissions from the incineration of biomass-derived waste (including biomass-based plastics and waste animal and

vegetable oil) is not included in the total emissions in accordance with the Revised 1996 IPCC Guidelines; instead it is estimated as a reference value and reported under “Biogenic” in Table 6.A,C of the CRF.

For reference, the greenhouse gas emissions from waste incineration for energy purpose and with energy recovery are shown in Table 8-36. In FY 2010, the emissions from waste incineration including these sources were 28,962 Gg-CO₂, and it accounts for 2.3% of Japan's total greenhouse gas emissions (excluding LULUCF). The emissions from this sources category had increased by 24.1% compared to those in FY 1990.

Table 8 - 36 Total GHG emissions from incineration of waste (reference value)
including emissions from waste incineration for energy use and energy recovery

Gas	Incineration type	Waste category	Estimation Category	Unit	1990	1995	2000	2005	2008	2009	2010	
CO ₂	Waste incineration without energy recovery (simple incineration)			Gg CO ₂	12,263	15,867	16,838	13,984	14,481	12,249	12,658	
	Waste incineration with energy recovery	Municipal solid waste	Plastics	Gg CO ₂	5,857	6,309	8,188	6,611	4,986	5,293	5,113	
			Synthetic textiles	Gg CO ₂	585	676	660	925	1,186	1,115	1,173	
			Other (biogenic)	Gg CO ₂								
		Industrial solid waste	Waste oil	Gg CO ₂	21	30	28	108	117	95	102	
			Waste plastics	Gg CO ₂	31	65	187	306	378	274	281	
			Other (biogenic)	Gg CO ₂								
	Direct use of waste as fuel	Municipal solid waste	Plastics	Gg CO ₂	0	0	91	507	367	410	452	
			Waste oil	Gg CO ₂	2,019	2,504	2,345	3,602	3,232	2,955	3,235	
		Industrial solid waste	Waste plastics	Gg CO ₂	54	36	446	1,203	1,325	1,418	1,453	
			Waste wood	Gg CO ₂								
		Waste tire	Fossil origin	Gg CO ₂	524	841	1,039	865	1,023	946	1,003	
	Use of processed waste as fuel	Refuse derived fuel (RDF, RPF)	Biogenic origin	Gg CO ₂								
			Fossil origin	Gg CO ₂	26	41	159	996	1,361	1,392	1,368	
	Total				Gg CO ₂	21,379	26,369	29,980	29,107	28,458	26,148	26,837
CH ₄	Waste incineration without energy recovery (simple incineration)			Gg CH ₄	0.64196	0.70800	0.63491	0.67938	0.57050	0.50662	0.49797	
	Waste incineration with energy recovery	Municipal solid waste		Gg CH ₄	0.53965	0.54072	0.59747	0.13829	0.12812	0.11812	0.11812	
		Industrial solid waste	Waste oil	Gg CH ₄	0.00004	0.00005	0.00005	0.00016	0.00017	0.00014	0.00015	
			Waste plastics	Gg CH ₄	0.00036	0.00077	0.00221	0.00096	0.00118	0.00086	0.00088	
			Other (biogenic)	Gg CH ₄	0.00039	0.00118	0.00130	0.00828	0.00779	0.00699	0.00680	
	Direct use of waste as fuel	Municipal solid waste	Plastics	Gg CH ₄	0	0	0.00003	0.00005	0.00002	0.00005	0.00001	
			Waste oil	Gg CH ₄	0.01183	0.01626	0.01895	0.02657	0.02410	0.02262	0.02555	
		Industrial solid waste	Waste plastics	Gg CH ₄	0.00025	0.00016	0.03922	0.11568	0.16363	0.16861	0.15828	
			Waste wood	Gg CH ₄	1.75918	1.75918	2.21808	2.88749	4.00783	4.00891	3.72238	
		Waste tire	Fossil origin	Gg CH ₄	0.03095	0.07576	0.09914	0.08004	0.06352	0.05091	0.04372	
	Use of processed waste	Refuse derived fuel (RDF, RPF)	Fossil origin	Gg CH ₄	0.00008	0.00012	0.00056	0.00595	0.01004	0.01193	0.00972	
			Total			Gg CH ₄	2.98468	3.10221	3.61191	3.94285	4.97690	4.89577
					Gg CO ₂ eq	63	65	76	83	105	103	96
	N ₂ O	Waste incineration without energy recovery (simple incineration)			Gg N ₂ O	4.90142	6.49081	7.29001	6.76181	5.60901	5.43120	5.44384
		Waste incineration with energy recovery	Municipal solid waste		Gg N ₂ O	1.19113	1.31566	1.53434	1.13358	1.02358	0.94718	0.94718
Industrial solid waste			Waste oil	Gg N ₂ O	0.00009	0.00013	0.00012	0.00248	0.00271	0.00220	0.00236	
			Waste plastics	Gg N ₂ O	0.00217	0.00460	0.01318	0.00177	0.00219	0.00159	0.00163	
			Other (biogenic)	Gg N ₂ O	0.00838	0.00853	0.01017	0.00540	0.00603	0.00580	0.00584	
Direct use of waste as fuel		Municipal solid waste	Plastics	Gg N ₂ O	0	0	0.00002	0.00004	0.00001	0.00004	0.00001	
			Waste oil	Gg N ₂ O	0.01581	0.02376	0.03221	0.04225	0.03845	0.03677	0.04240	
		Industrial solid waste	Waste plastics	Gg N ₂ O	0.00018	0.00012	0.00356	0.01023	0.01451	0.01495	0.01405	
			Waste wood	Gg N ₂ O	0.01993	0.01993	0.02513	0.03271	0.04540	0.04541	0.04217	
		Waste tire	Fossil origin	Gg N ₂ O	0.00501	0.00999	0.01166	0.01484	0.01773	0.01643	0.01696	
Use of processed waste as fuel		Refuse derived fuel (RDF, RPF)	Fossil origin	Gg N ₂ O	0.00051	0.00079	0.00309	0.01865	0.02524	0.02582	0.02536	
			Total			Gg N ₂ O	6.14462	7.87432	8.92348	8.02375	6.78487	6.52738
				Gg CO ₂ eq	1,905	2,441	2,766	2,487	2,103	2,023	2,028	
Total of all gases				Gg CO ₂ eq	23,346	28,875	32,822	31,678	30,666	28,274	28,962	

* CO₂ emissions from the incineration of biomass-derived waste (including biomass-based plastics and waste animal and vegetable oil) is not included in the total emissions in accordance with the Revised 1996 IPCC Guidelines

8.4.1. Waste Incineration without Energy Recovery (6.C.)

8.4.1.1. Municipal Solid Waste Incineration (6.C.1)

a) Source/Sink Category Description

This category covers the emissions from incineration of MSW without energy recovery. Emissions of CO₂ are reported under either “biogenic” or “plastics and other non-biogenic waste” in accordance with the waste type. Emissions of CH₄ and N₂O are estimated for each type of furnace. The data used for MSW incineration can not distinguish wastes that are either biogenic-origin or non-biogenic origin. Therefore, total emissions including biogenic-origin ones are reported altogether under “plastics and other non-biogenic waste”.

b) Methodological Issues

1) CO₂

● Estimation Method

Emissions of CO₂ from this emission source was calculated based on Japan’s country-specific emission factors, the volume of waste incinerated (dry basis) and the percentage of municipal waste incinerated at the municipal incineration facilities that is accompanied by energy recovery, in accordance with the decision tree in the GPG (2000) (Page 5.26, Fig. 5.5). In order to estimate CO₂ emissions from the incineration of fossil-fuel derived waste² emissions from plastics and synthetic textile wastes in municipal waste were calculated.

$$E = EF \times A \times (1-R)$$

- E* : Emission of carbon dioxide from the incineration of various types of waste (kg CO₂)
EF : Emission factor for the incineration of various types of waste (dry basis) (kg CO₂/t)
A : Volume of each type of waste incinerated (dry basis) (t)
R : Percentage of municipal solid waste incinerated at facilities with energy recovery

● Emission factor

In accordance with the *Revised 1996 IPCC Guidelines*, the emission factor was calculated by multiplying the carbon content of each type of waste by the incineration rate at each incinerator.

$\begin{aligned} & \text{CO}_2 \text{ emission factor (dry basis)} \\ & = 1000 \text{ [kg]} \times \text{Carbon content} \times \text{efficiency of combustion} \times 44/12 \end{aligned}$

Carbon content

The carbon content of waste plastics (fossil-fuel derived and biomass-derived waste) in MSW was estimated based on the averaged value of actual measured data for the period FY1990 - FY2008 provided by four municipalities (Akita city, Kawasaki city, Kobe city and Osaka pref.) and applying it for the entire time-series, according to the *Survey Study on Improving the Accuracy of Emission Factors for Greenhouse Gas Emissions from the Waste Sector, 2010*, Ministry of the Environment (Reference #15).

² Emissions from the incineration of kitchen garbage, waste paper, waste natural fiber textiles and waste wood, and biomass-based plastics were accounted for as the reference figures of biogenic waste. Estimation methods for their emissions are the same as those for emissions from the incineration of fossil-fuel derived plastics and synthetic textile scraps.

For the carbon content of synthetic textile wastes in MSW, the carbon content of the synthetic fibers in the textile products was used. It was set by taking a weighted average of carbon contents determined by the molecular formula of polymer for each type of synthetic textile based on the volume of synthetic textile consumption.

Table 8 - 37 Carbon content of plastics and synthetic textile scrap in MSW

Item	Carbon content	Remarks
Plastics	75.1 %	Averaged value of the data provided by four municipalities
Synthetic textile	63.0 %	Weighted average of carbon content by each type of synthetic textile

Efficiency of Combustion

Taking into account Japan's circumstances, the default value of 99% indicated in the *GPG (2000)* was used.

● ***Activity data***

The activity data for CO₂ emissions from the incineration of fossil-fuel derived waste plastics in MSW on a dry basis were calculated by subtracting water content from the amount of plastics incinerated (wet basis) and also subtracting the amount of biomass-based plastics incinerated (dry basis) in MSW which were estimated separately.

Activity data for plastics (MSW) incinerated (dry basis)

= Volume of plastics incinerated (wet basis) × (1 - Percentage of water content in waste plastics) – Amount of biomass-based plastics incinerated (dry basis)

The amount of biomass-based plastics incinerated (dry basis) is estimated as indicated below:

Amount of biomass-based plastics incinerated (dry basis)

= Amount of biomass-based plastics products consumed (dry basis) × Fraction of biogenic component of biomass-based plastics products × Fraction of biomass-based plastics disposed of as MSW × Fraction of biomass-based plastics incinerated

The activity data of waste synthetic textile in MSW was estimated by multiplying the amount of waste textile in MSW incinerated (wet basis) by the fraction of waste synthetic textile content in waste textile, and subtracting the water content in waste textile.

Activity data for incineration of synthetic textile scraps (MSW) (dry basis)

= Volume of textile scraps incinerated (wet basis) × (1 - Percentage of water content in waste textile) × Percentage of synthetic fiber content in textile scraps

Table 8 - 38 Amount of incineration of plastics and synthetic textile scraps in MSW (dry basis)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Plastics	kt/year (dry)	3,998	4,160	4,919	3,548	2,677	2,937	2,838
Synthetic textile	kt/year (dry)	476	531	473	592	759	737	775

Incineration volume by type of municipal solid waste

The amount of waste plastics including biomass-based plastics and waste textile incinerated were obtained from the *Cyclical Use of Wastes Report*.

Amount of biomass-based plastics products consumed

Because of the fact that most of biomass-based plastics products consumed in Japan are produced abroad, the amount of import of biomass-based plastics products are substituted for the amount of biomass-based plastics products consumed, which was compiled by the Japan Society of Biomass Industries. Currently, available values, the amount of biomass-based plastics products consumed by type and use, are limited to the products from polylactic acid (PLA), chemically modified, and partially biomass-based (Table 8-39).

Table 8 - 39 Amount of biomass-based plastics products consumed

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Containers and packaging (Polylactic acid (PLA))	kt/year (dry)	0	0	0	0	2.81	1.84	2.28
Plastic bags. (Biomass-based)	kt/year (dry)	0	0	0	0	0.50	0.80	1.10
Molded consumer goods (Chemically modified & partially biomass-based)	kt/year (dry)	0	0	0	0	0.23	0.53	0.29

Fraction of biogenic component of biomass-based plastics products

Because of the fact that some types of biomass-based plastics products include substances other than biogenic component, the net amount of biomass-based plastics incinerated by product type was estimated with the fraction of biogenic component (Table 8-40) obtained by the inquiry results from the Japan Society of Biomass Industries.

Table 8 - 40 Fraction of biogenic component of biomass-based plastics products

Product type	Product use	Unit	Fraction of biogenic property
Polylactic acid (PLA)	Containers and packaging	%	100
Chemically modified & partially biomass-based	Plastic bags	%	25
	Molded consumer goods	%	55

Fraction of biomass-based plastics disposed of as municipal waste

The fraction of biomass-based plastics disposed of as municipal waste was considered to be 100% for the each product use of containers and packaging, plastic bags, and molded household consumer goods based on the inquiry results from the Japan Society of Biomass Industries including the fact that all of these products were disposed of as MSW within a relatively short period of time after the their production.

Fraction of incinerated municipal waste, used as raw materials or fuels, and used as RDF

Biomass-based plastics are disposed of as municipal waste and utilized in a few different ways. The fraction of biomass-based plastics incinerated, biomass-based plastics used as raw materials or fuels, and used as RDF were estimated by dividing the amount of biomass-based plastics incinerated, used as raw materials or fuels, and plastic-derived component of RDF by the amount of plastics disposed of as MSW (Table 8-41).

Table 8 - 41 Fraction of incinerated municipal waste used as raw materials and fuels, and used as RDF

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Incineration rate including energy recovery	%	74.9	79.5	86.7	79.4	77.6	87.2	74.5
Fraction of biomass-based plastics used as raw material of fuel (excluding the use of recyclable materials)	%	0	0	0.6	5.3	4.5	5.0	6.4
Fraction of biomass-based plastics used as RDF	%	0.2	0.2	0.8	2.8	2.9	2.9	3.3

The rest of municipal waste other than the above are recycled or landfilled.

Percentage of water content

The percentage of water content in plastics in MSW was determined to be 20% provided by the *Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes*. The percentage of water content in the waste textile contained in MSW was determined to be 20% based on expert judgment and their review of case studies in Japan.

Percentage of synthetic textile in waste textile

Percentage of synthetic textile content in waste textiles contained in the MSW was calculated using the percentage of synthetic textile products in textile products, which was determined by taking the ratio of the annual domestic demand for synthetic textile to the one for all textiles indicated in the *Textile Handbook* and the *Yearbook of Textiles and Consumer Goods Statistics*.

Table 8 - 42 Percentage of synthetic textile in waste textile

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Fraction of Synthetic fabric	%	49.1	50.7	53.5	52.8	55.9	56.6	59.6

- Percentage of municipal waste incinerated at municipal incineration facilities for energy recovery
 Percentage of municipal waste that is incinerated at municipal incineration facilities with energy recovery stands for the one being incinerated at the facilities actually supply electricity or heat outside of them. These values were obtained from the *State of Municipal Waste Treatment Survey* (Ministry of the Environment).

Table 8 - 43 Percentage of municipal solid waste incinerated at incineration facilities with energy recovery

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Without off-field power generation or heat utilization	%	46.3	44.4	38.9	31.6	31.6	33.9	33.9
With off-field power generation or heat utilization	%	53.7	55.6	61.1	68.4	68.4	66.1	66.1

2) CH₄

● Estimation Method

CH₄ emissions from incinerator were estimated by multiplying the amount of MSW (wet basis) by incinerator method by each emission factor. CH₄ emissions from gasification melting furnace were estimated by multiplying the amount of MSW (wet basis) incinerated in gasification melting furnace by emission factors. Emissions from MSW with energy recovery were subtracted from the total emissions from this source and allocated to the waste sector.

$$E = \sum (EF_i \times A_i) \times (1 - R)$$

- E : CH₄ emission from the incineration of MSW (kg CH₄)
 EF_i : Emission factor for incineration method i (or furnace type i) (wet basis) (kg CH₄/t)
 A_i : Amount of incinerated MSW by incineration method i (or furnace type i) (wet basis) (t)
 R : Percentage of MSW incinerated at facilities with energy recovery

● Emission factor

Incinerator

In order to implement countermeasures against dioxins, the renovations, repairs, or rebuilding of incineration facilities took place in the latter half of 1990 through the first half of 2000 in Japan. There have been some improvements made in CH₄ emission factors from the facilities renovated or rebuilt in FY 2000 and later, compared to the values obtained before then (Reference # 15). Therefore, based on the survey (Reference #15) and expert judgment, for the CH₄ emission factors for incinerator by incinerator type (stoker furnace and fluidized bed incinerator) and incineration method (continuous incinerator, semi-continuous incinerator, and batch type incinerator) for the period FY 2001 and before (Reference #7), and from FY 2002 onward (Reference #15), respectively, different values were used. All the emission factors were established based on actual measurement survey.

In order to apply activity data based on the amount of incineration by incineration method, emission factors were established by incineration method (continuous incinerator, semi-continuous incinerator, and batch type incinerator) using the weighted average of fraction of the amount of incineration by incinerator type for each fiscal year. The Correction taking into account CH₄ concentrations in the atmosphere was not made to these emission factors.

Table 8 - 44 CH₄ emission factors by incineration method of incinerator (MSW)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Continuous incinerator	g CH ₄ /t	8.2	8.2	8.3	2.6	2.6	2.6	2.6
Semi-continuous incinerator	g CH ₄ /t	69.6	69.6	75.1	19.9	21.0	20.6	20.6
Batch type incinerator	g CH ₄ /t	80.5	80.5	84.1	13.2	13.2	13.4	13.4

Source: Reference # 6, 8, 15, 23, 47, 52

Gasification Melting Furnace

Different emission factor was used for each furnace type (shaft furnace, fluidized bed, and rotary kiln) (Reference #15). Also, in order to apply activity data based on the total amount of incineration, emission factors were determined by taking the weighted average of the amount of incineration by gasification melting furnace type for each year.

Table 8 - 45 CH₄ emission factors by type of gasification melting furnace (MSW)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Gasification melting furnace	g CH ₄ /t	-	-	5.6	6.9	7.1	7.0	7.0

● Activity Data

The activity data for CH₄ emissions for incinerator and gasification melting furnace were estimated by

multiplying the amount of MSW incinerated (wet basis) provided in the *Report of the Research on the state of wide-range Movement and Cyclical Use of Wastes*(publicized reports and the most current data from the reports prior to publication) by the fraction of incineration by incineration method of incinerator or gasification melting furnace provided by the *Waste Treatment in Japan*.

Table 8 - 46 Amount of incineration of MSW by type of incinerator

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Continuous incinerator	kt /year (wet)	26,215	29,716	32,749	32,246	29,426	28,444	28,444
Semi-Continuous Incinerator	kt /year (wet)	4,810	5,455	5,882	4,047	3,339	3,155	3,155
Batch type Incinerator	kt /year (wet)	5,643	4,328	3,131	1,562	1,346	1,144	1,144

Table 8 - 47 Amount of incineration of MSW from gasification melting furnace

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Gasification melting furnace	kt /year (wet)	0	0	370	2,397	3,122	3,245	3,245

3) N₂O

● Estimation Method

N₂O emissions from incinerator were estimated by multiplying the amount of MSW (wet basis) by incinerator method by each emission factor. N₂O emissions from gasification melting furnace were estimated by multiplying the amount of MSW (wet basis) incinerated in gasification melting furnace by emission factors. Emissions from MSW with energy recovery were subtracted from the total emissions from this source and allocated to the waste sector.

$$E = \sum (EF_i \times A_i) \times (1 - R)$$

E : N₂O emission from the incineration of MSW (kg N₂O)

EF_i : Emission factor for incineration method i (or furnace type i) (wet basis) (kg N₂O /t)

A_i : Amount of incinerated MSW by incineration method i (or furnace type i) (wet basis) (t)

R : Percentage of MSW incinerated at facilities with energy recovery

● Emission factor

Incinerator

Same as for CH₄ emissions estimation, for the N₂O emission factors for incinerator by type and by incineration method, different values were used for the period FY 2001 and before (Reference #7), and from FY 2002 onward (Reference #15), respectively. In order to apply activity data based on the amount of incineration by incineration method, emission factors were established by incineration method (continuous incinerator, semi-continuous incinerator, and batch type incinerator) using the weighted average of fraction of the amount of incineration by incinerator type for each fiscal year calculated based on the *Waste Treatment in Japan*.

Table 8 - 48 N₂O emission factors for incinerator by incineration method (MSW)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Continuous incinerator	g N ₂ O/t	58.8	58.8	59.1	37.9	37.9	37.9	37.9
Semi-continuous incinerator	g N ₂ O/t	56.8	56.8	57.3	71.5	73.3	72.7	72.7
Batch type Incinerator	g N ₂ O/t	71.4	71.4	74.8	76.0	76.0	76.0	76.0

Source: Reference # 7, 8, 15 24, 48, 53

Gasification Melting Furnace

Different emission factor was used for each furnace type (shaft furnace, fluidized bed, and rotary kiln) (Reference #15). In order to apply the activity data based on the total amount of incineration, emission factors were established by taking the weighted average of the amount of incineration by gasification melting furnace type for each year calculated based on the *Waste Treatment in Japan*.

Table 8 - 49 N₂O emission factors for gasification meting furnace (MSW)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Gasification melting furnace	g N ₂ O/t	-	-	16.9	12.0	11.1	11.2	11.2

● **Activity Data**

The activity data for estimating CH₄ emissions from incinerator and gasification melting furnace were also applied for the activity data for N₂O emission estimates from incinerator and gasification melting furnace.

c) Uncertainties and Time-series Consistency

● **Uncertainties**

The level of uncertainty in the CO₂ emission factor was estimated by using the uncertainties in the carbon content of MSW (plastic and synthetic textile) and the incineration rate of MSW incineration facilities. The uncertainty in activity data for CO₂ emissions was estimated from the uncertainties in the amount of MSW incinerated, the percentage of water content and the percentage of synthetic textile (for synthetic textile in MSW).

The uncertainties in the CH₄ and N₂O emission factors were evaluated by type of incineration facilities and determined from the uncertainties in the emission factors for each type of incineration facilities and the ratio of the incinerated amount by type of incineration facilities. The uncertainties in the activity data were estimated based on the uncertainties in the amount of waste incinerated and the ratio of incinerated amount by type of incineration facilities. The methods of evaluation of the uncertainty levels for each component are:

- Use of 95% confidence interval: carbon content, fraction of synthetic textile, emission factors for CH₄ and N₂O by type of incineration facility
- Use of the default value in the 2006 IPCC Guidelines: combustion rate
- Based on expert judgment: percentage of water content
- Use of the statistical uncertainties: incinerated amount of waste and incineration rate by incinerator type

The uncertainties in the CO₂ emissions from incineration of plastics and synthetic textiles of MSW

were estimated to be 17% and 23%, respectively. The uncertainties in the CH₄ and N₂O emissions from incineration of MSW were estimated to be 101% and 42%, respectively. For more details, see the Annex 7.

● ***Time-series consistency***

Because data on the amount of waste incinerated by type of waste were not available for years prior to FY 1997, the data were estimated by using the total incinerated amount of MSW for each year and the ratio of amount of waste incinerated by waste type for FY 1998. The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Recalculations

- Updating the amount of incinerated municipal waste, emission estimates for FY2005 and 2007-2009 were recalculated.

f) Source-specific Planned Improvements

No improvements are planned.

8.4.1.2. Industrial Waste Incineration (6.C.2)

a) Source/Sink Category Description

This category covers CO₂, CH₄ and N₂O emissions from incineration of ISW without energy recovery by each waste type and the emissions are reported in the corresponding category either “biogenic” or “plastics and other non-biogenic waste”.

b) Methodological Issues

1) CO₂

● ***Estimation Method***

Emissions of CO₂ from this source were calculated by using the volume of waste mineral oil and waste plastics incinerated, Japan’s country-specific emission factors, and the percentage of incinerated industrial solid waste with energy recovery at industrial waste incineration facilities in accordance with the decision tree of the *GPG (2000)* (Page 5.26, Fig. 5.5). Since industrial waste textile does not include synthetic textile under the regulation of the Waste Disposal and Public Cleansing Law, the industrial waste textile is regarded as waste natural fiber. Thus the CO₂ emissions from incineration of industrial waste textile were not included in national total because these emissions are biogenic-origin.

$$E = EF \times A \times (1-R)$$

E : Emission of carbon dioxide from incineration of waste (kg CO₂)

EF : Emission factor for waste incineration (wet basis) (kg CO₂/t)

A : Amount of waste incinerated (wet basis) (t)

R : Percentage of industrial solid waste incinerated at facilities with energy recovery (by type of waste)

● Emission factor

In accordance with the approach taken by the *Revised 1996 IPCC Guidelines*, emission factor was calculated by multiplying the carbon content of each type of waste by the incineration rate for incineration facilities.

<p><u>Carbon dioxide emission factor (wet basis)</u> = 1000 [kg] × Carbon content × Efficiency of combustion × 44/12</p>

Carbon content

Carbon content in waste oil was deemed to be 80% based on the factor of 0.8 (t C/t) given in the *Environmental Agency's Report on a Survey of Carbon Dioxide Emissions (1992)*.

Carbon content in waste plastics was deemed to be 70% based on the factor of 0.7 (t C/t) given in the said report .

Efficiency of combustion

Considering Japan's circumstances, the default value for hazardous wastes of 99.5% given in the *GPG (2000)* was used.

● Activity Data

For the activity data for CO₂ emissions from the incineration of waste oil and waste plastics in industrial waste, the amount of incineration provided by *the Report of the Research on the State of Wide-range Movement and Cyclic Use of Wastes* was used. However, the amount of incineration provided in this report includes the amount of incineration of specially controlled industrial waste which is separately reported under "Incineration of Specially Controlled Industrial Waste (6.C.3)", thus it was subtracted from the activity data from this source. The activity data for waste mineral oil was obtained by using the fraction of animal and vegetable waste oil (biogenic-origin waste oil) provided by the survey study conducted by the Ministry of the Environment from the total amount of waste oil (see the methodological equation indicated below). All of the plastics in ISW was considered to be fossil-fuel derived.

<p><u>Activity data for the incineration of waste mineral oil (wet basis)</u> = Amount of waste oil incinerated in industrial waste × (1 – Fraction of waste oil from animal and vegetable origin) – Amount of waste oil incinerated in specially controlled industrial waste*</p>

*All the waste oil in specially controlled industrial waste to be estimated for emissions are waste mineral oil.

Activity data for the incineration of waste oil and plastics (ISW) (wet basis)

= Amount of waste plastics incinerated in industrial waste – Amount of waste plastics incinerated in specially controlled industrial waste

Table 8 - 50 Incinerated ISW (waste oil and waste plastics)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Waste mineral oil	kt / year (wet)	1,258	1,498	1,646	1,493	1,622	1,314	1,410
Waste plastics	kt / year (wet)	842	1,794	1,780	1,808	2,056	1,493	1,532

Table 8 - 51 Fraction of waste animal and vegetable oil

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Fraction of waste animal and vegetable oil	%	2.6	3.5	4.5	5.4	6.0	6.0	6.0

- Percentage of industrial waste incinerated at industrial incineration facilities for energy recovery (by type)

Percentage of industrial waste that is incinerated at industrial incineration facilities with energy recovery stands for the one being incinerated at the facilities actually supply electricity or heat outside of them. The values were obtained from the *FY 2007 Survey of Industrial Waste Treatment Facilities* (Ministry of the Environment).

In Japan, industrial incineration facilities are installed mainly by private sector waste disposal enterprises. In comparison with the municipal waste incinerators installed primarily by municipal governments, energy recovery (for use in power generation and as a heat source) has not yet been so popular. The percentage for the industrial waste category is therefore smaller.

Table 8 - 52 Percentage of ISW incinerated at incineration facilities with energy recovery

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Waste oil ^{a)}	%	0.6	0.7	0.6	2.5	2.5	2.5	2.5
Waste plastics	%	1.4	1.4	4.1	6.6	7.2	7.2	7.2
Waste wood ^{b)}	%	0.2	0.8	1.1	1.5	1.8	1.8	1.8
Sludge	%	0.9	0.8	1.0	1.1	1.6	1.6	1.6
Other ^{c)}	%	0.2	0.8	1.1	1.5	1.8	1.8	1.8

a): "Waste oil" includes waste mineral/animal and vegetable oil.

b): "Waste wood" includes waste paper or waste wood.

c): "Other" includes waste textile, animal and vegetable residues, and animal carcasses.

2) CH₄

● Estimation Method

Emissions of methane from this source have been calculated by multiplying the volume of industrial waste incinerated by Japan's country specific emission factor and by percentage of industrial solid waste incinerated at incineration facilities with energy recovery.

$$E = \sum \{EF_j \times A_j \times (1 - R_j)\}$$

E : Emission of methane from the incineration of industrial waste (kg CH₄)

EF_j : Emission factor for waste type j (wet basis) (kg CH₄/t)

A_j : Incinerated amount of waste type j (wet basis) (t)

R_j : Percentage of industrial solid waste j incinerated at facilities with energy recovery

● Emission factor

Based on expert judgement which takes into account the countermeasures against dioxin emissions from incinerators, for the emission factors by waste type for the period FY1990 - FY2001 (Reference #7) and from FY 2002 onward (Reference #15), respectively, different values were used. These emission factors were established based on actual measurement survey. The correction taking into account CH₄ concentrations in the atmosphere was not made to these emission factors. The emission factor for waste paper or waste wood was substituted for the emission factor for waste textile, animal and vegetable residues, and animal carcasses.

Table 8 - 53 CH₄ emission factors for industrial waste by type

Item	Unit	FY 1990-2001	FY 2002 onward
Waste oil (mineral/animal and vegetable)	g CH ₄ /t	4.8	4.0
Waste plastics	g CH ₄ /t	30	8.0
Waste paper or Waste wood	g CH ₄ /t	22	225
Waste textile	g CH ₄ /t	22	225
Animal and vegetable residues/animal carcasses	g CH ₄ /t	22	225
Sludge	g CH ₄ /t	14	1.5

Reference # 6, 24, 47

● Activity Data

The volume of waste incinerated (wet basis) by waste type was used as the activity data for CH₄ emissions from the incineration of industrial waste.

Paper and wood scraps, waste oil, textile scraps, animal and plant residues or animal carcasses:

The volume of waste incinerated for each type was obtained from the Report of the Research on the State of Wide-range Movement and Cyclical Use of Waste.

Sludge

Activity data was taken as the aggregate of the values obtained from the “Volume of Other Incinerated Organic Sludge” section in the *Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes*, and the “Volume of Incinerated Sewage Sludge” reported in a survey by the Ministry of Lands, Infrastructure, Transport and Tourism.

Waste oil (mineral/animal and vegetable) and waste plastics

The activity data for waste oil and waste plastics were provided by the *Report of the Research on the State of Wide-range Movement and Cyclical Use of Waste*. Because the values provided by this report include the amount of specially controlled industrial waste which is allocated to the category of Specially Controlled Industrial Waste (6.C.3), it was subtracted from the total amount to avoid double counting. Unlike the activity data for CO₂ emissions, waste mineral oil and also waste animal and vegetable oil are included for the estimation of activity data from this source.

Table 8 - 54 Incinerated ISW by waste types

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Waste animal and vegetable oil	kt / year (wet)	40	69	103	115	139	113	121
Waste paper, waste wood	kt / year (wet)	3,014	5,455	3,832	2,188	1,638	1,491	1,444
Waste textile	kt / year (wet)	31	49	50	43	33	26	24
Animal and vegetable remnants,	kt / year (wet)	77	125	272	167	220	181	181
Sludge	kt / year (wet)	5,032	5,850	6,371	7,275	6,820	6,766	6,837

For the amount of waste oil and waste tires incinerated, see Table 8-50.

3) N_2O

● *Estimation Method*

Emissions of N_2O from this source were calculated separately for the major emission source, sewage sludge, and the waste other than sewage sludge. With respect to sewage sludge, emission factors were set by type of flocculants and furnaces; and the ones for “high-molecular-weight, flocculant fluidized bed incinerator” were further determined by the incineration temperatures. Emissions from the industrial waste other than sewage sludge were estimated by multiplying the volume of waste incinerated by Japan’s country-specific emission factor. Among those emissions, the ones to be reported in the waste sector were calculated by multiplying the percentage of industrial waste incinerated at the industrial waste incineration facilities with energy recovery.

$$E = \sum \{EF_j \times A_j \times (1 - R_j)\}$$

- E : Emission of nitrous oxide from the incineration of industrial waste (kg N_2O)
 EF_j : Emission factor for waste type j (wet basis) (kg N_2O /t)
 A_j : Incinerated amount of waste type j (wet basis) (t)
 R_j : Percentage of industrial solid waste j incinerated at facilities with energy recovery

● *Emission factor*

Sewage sludge

Emission factor for N_2O emissions from sewage sludge incineration were determined by taking a weighted average of actually measured emission factors of N_2O at each incineration facility based on the volume of sewage sludge incinerated at the facilities. Since emission factors are different depending on the types of flocculants, incinerators, and furnace temperatures, they were established for each category as given in Table 8-55 (Reference #7).

Table 8 - 55 N₂O emission factors for sewage sludge incineration (wet basis)

Type of flocculant	Type of incinerator	Combustion Temperature	Emission Factor (g N ₂ O/t)
High-molecular weight flocculant	Fluidized Bed Incinerator	Normal temperature combustion (around 800°C)	1,508
	Fluidized Bed Incinerator	High temperature combustion (around 850°C)	645
	Multiple Hearth	—	882
Other	—	—	
Lime Sludge	—	—	294

- Source: Reference #25, 26, 27, 28, 29, 30, 47, 53, 54

- Assume that emission factors for FY1990-2002 are constant.

Waste other than sewage sludge

Based on expert judgement which takes into account the countermeasures against dioxin emissions from incinerators, for the emission factors by waste type for the period FY1990-FY 2001 (Reference #7) and from FY2002 onward (Reference #15), respectively, different values were used. These emission factors were established based on actual measurement survey. The correction taking into account CH₄ concentrations in the atmosphere was not made to these emission factors. The emission factor applied for waste paper or waste wood was also used for waste textile, animal and vegetable residues, and animal carcasses.

Table 8 - 56 N₂O Emission factors for industrial waste by type (wet basis)

Item	Unit	FY 1990-2001	From FY 2002 onward
Waste oil (mineral/animal and vegetable)	g N ₂ O / t	12	62
Waste plastics	g N ₂ O / t	180	15
Waste paper or Waste wood	g N ₂ O / t	21	77
Waste textile	g N ₂ O / t	21	77
Animal and vegetable residues/animal carcasses	g N ₂ O / t	21	77
Sludge (excluding sewage sludge)	g N ₂ O / t	457	99

Source: Reference # 6, 15, 25, 48, 54, 55, 59, 60, 61, 65, 66

● Activity Data

Sewage sludge

Data in the “volume of incinerated sewage sludge, by flocculants and by incinerator types” reported in a survey by the Ministry of Lands, Infrastructure, Transport and Tourism were used as activity data (wet basis).

Table 8 - 57 Amount of sewage sludge incinerated

Item	Unit	1990	1995	2000	2005	2008	2009	2010
High-molecular-weight flocculant Fluidized bed incinerator (normal temp.)	kt / year (wet)	1,112	1,869	2,397	2,839	1,785	1,664	1,664
High-molecular-weight flocculant Fluidized bed incinerator (high temp.)	kt / year (wet)	128	219	723	1,469	2,470	2,561	2,561
High-molecular-weight flocculant multiple hearth	kt / year (wet)	560	656	572	102	56	64	64
Lime sludge	kt / year (wet)	1,070	767	341	289	193	142	142
Other	kt / year (wet)	190	316	267	289	233	229	229

Industrial waste other than sewage sludge

Activity data (wet basis) was determined in the same manner as for the CH₄ emissions from industrial waste, with the exception that the “volume of other incinerated organic sludge” was used as activity data for the sludge (excluding sewage sludge).

c) Uncertainties and Time-series Consistency**● *Uncertainties***

The uncertainties in the CO₂ emission factor and activity data for waste oil and waste plastics were evaluated by the same method as was used for incineration of MSW. The uncertainties in CH₄ and N₂O emission factors were estimated by using the 95% confidence interval of actual measurement data of the emission factors by type of ISW and by type of incineration facility. The uncertainties in the CH₄ and N₂O activity data were estimated by using the statistical uncertainties for incinerated amount of industrial waste by type of waste.

The uncertainties in the CH₄ and N₂O emissions from incineration of industrial waste were estimated to be 150% and 116%, respectively. The uncertainties in the CO₂ emissions from incineration of waste oil and waste plastics were 105% and 100%, respectively. For more details, see the Annex 7.

● *Time series consistency*

Emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Recalculations

Updating the amount of incinerated industrial waste, emission estimates for FY2009 were recalculated.

f) Source-specific Planned Improvements

No improvements are planned.

8.4.1.3. Incineration of Specially controlled Industrial Waste (6.C.3)***a) Source/Sink Category Description***

The specially controlled industrial waste includes wastes with properties that may be harmful to human health and living environment such as explosiveness, toxicity and infectivity. This category covers CO₂, CH₄, and N₂O emissions from incineration of specially controlled industrial waste were estimated by each waste type and reported in the corresponding category either “biogenic” or “plastics and other non-biogenic waste”.

Because the actual state of energy recovery from the incineration of specially controlled industrial

waste is not sufficiently understood, the emissions from specially controlled industrial waste are reported entirely in “Waste Incineration (Category 6.C.)”.

b) Methodological Issues

1) CO₂

● *Estimation Method*

Emissions of CO₂ from the incineration of waste oil and infectious plastic waste contained in specially controlled industrial waste were calculated in accordance with the decision tree given in the *GPG (2000)* (Page 5.26, Fig 5.5) by using Japan’s country-specific emission factors and the volume of waste incinerated.

● *Emission factor*

Emission factors for waste oil and waste plastics in industrial waste were used as the ones for waste oil and waste plastics in specially controlled industrial waste, since their differences in terms of carbon contents and rates of combustion were considered to be small.

● *Activity Data*

On the assumption that the entire volume of waste oil and infectious plastic waste contained in specially controlled industrial waste was incinerated, output volume of waste oil indicated in the *Report on Survey of Organizations in Industrial Waste Administration* (Water Supply Division, Health Service Bureau, the Ministry of Health and Welfare) was used as activity data for the waste mineral oil; while for the plastics in infectious waste, the activity data was calculated by multiplying the output volume of infectious waste reported by the same survey by the percentage of plastic content in infectious waste indicated in the *Waste Handbook* as the result of a composition analysis of infectious waste. All the waste oil in specially controlled industrial waste to be estimated for emissions is waste mineral oil. All of plastics in infectious waste were considered to be fossil-fuel derived.

Activity data for incineration of waste mineral oil (specially controlled ISW) (wet basis)
= Output volume of waste oil

Activity data for incineration of plastics in infectious waste (specially controlled ISW)(wet basis)
= Output volume of infectious waste × percentage of plastic content in infectious waste

2) CH₄

● *Estimation Method*

Emissions of CH₄ from the incineration of waste oil and infectious waste included in the specially controlled industrial waste were calculated by multiplying the volume of incinerated waste by type (wet basis) by Japan’s country-specific emission factor.

● *Emission factor*

Because actual measurement data were not available, the emission factors for the incineration of industrial waste were used as substitutes for the emission factor for the specially controlled industrial waste by type. Specifically, the substitute emission factors used were: the waste mineral oil in industrial waste for the waste mineral oil; the waste plastics in industrial waste for the infectious waste plastics; and the waste paper and waste wood in industrial waste for the waste other than infectious plastics.

● *Activity Data*

Activity data for the waste oil and infectious waste plastics were the same as those used for CO₂ emission. The volume of non-infectious waste plastics incinerated was deemed to be the same as the output volume, and calculated by multiplying the output volume of infectious waste by the percentage of non-plastic content in infectious waste.

3) N₂O

● *Estimation Method*

Emissions of N₂O from the incineration of waste oil and infectious waste in specially controlled industrial waste were calculated by multiplying the incinerated volume of each type of waste (wet basis) by Japan's country-specific emission factor.

● *Emission factor*

Because actual measurement data were not available, the N₂O emission factors for the incineration of industrial waste were used as substitutes for determining the emission factor for each type of specially controlled industrial waste. Specifically, the substitute emission factors used were: the waste oil in industrial waste for the waste oil; the waste plastics in industrial waste for the infectious waste plastics; and the waste paper and waste wood in industrial waste for the waste other than infectious plastics.

● *Activity Data*

The same activity data used for CH₄ emissions was used.

Table 8 - 58 Amount of incineration of specially controlled industrial waste

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Waste mineral oil	kt (wet)	256	380	560	515	564	457	490
Infections Waste (plastic)	kt (wet)	78	128	167	169	193	140	144
Infections Waste (non-plastic)	kt (wet)	105	172	225	228	260	189	193

c) *Uncertainties and Time-series Consistency*

● *Uncertainties*

Since the same CO₂, CH₄, and N₂O emission factors used for the industrial waste were used; their uncertainties were also applied. The uncertainties in activity data were set out separately for waste oil and waste plastics. To the incinerated amount of waste oil and infectious waste, twice the statistical uncertainties were applied by taking into account the fact that the data were recently obtained based on the estimation. For waste plastics, the uncertainties in the percentage of plastics in infectious waste

were determined based on the expert judgment, and then their uncertainties were combined with the ones in the amount of waste incinerated. The uncertainties in the CO₂, CH₄, and N₂O emissions from incineration of specially controlled industrial waste were estimated to be 167%, 142% and 159%, respectively. For details, see the Annex 7.

● *Time series consistency*

Since some basic data used for calculating activity data were available only for part of time series, consistent data over the time series were developed based on the estimation. The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Recalculations

Updating the amount of incinerated specially controlled industrial waste, emission estimates for FY2009 were recalculated.

f) Source-specific Planned Improvements

No improvements are planned.

8.4.2. Emissions from waste incineration with energy recovery (1.A.)

a) Source Category Description

In this category, CO₂, CH₄, and N₂O emissions from the incineration of municipal and industrial waste with energy recovery are estimated and reported. The reporting category for the emissions is “Power Generation/Heat Supply (Category 1.A.1.a)” and the fuel type is classified as “Other fuels”.

b) Methodological Issues

Methodologies similar to that used in “8.4.1.1 Municipal Solid Waste Incineration (6.C.1)” and “8.4.1.2. Industrial Waste Incineration (6.C.2)” were used. Emissions are calculated using the following equation:

1) CO₂

● *Estimation Method*

Municipal Solid Waste

$$E = EF \times A \times R$$

E : Emission of CO₂ from waste incineration (kg CO₂)

EF : Emission factor for incineration (dry basis) (kg CO₂/t)

A : Amount of waste incinerated (dry basis) (t)

R : Percentage of municipal solid waste incinerated at incineration facilities with energy

recovery

Industrial Solid Waste

$$E = EF \times A \times R$$

- E : Emission of CO₂ from waste incineration (kg CO₂)
 EF : Emission factor for waste incineration (wet basis) (kg CO₂/t)
 A : Amount of waste incinerated (wet basis) (t)
 R : Fraction of industrial solid waste incinerated at ISW incineration facilities with energy recovery (by waste type)

2) CH₄, N₂O

● Estimation Method

Municipal Solid Waste

$$E = \sum (EF_i \times A_i) \times R$$

- E : Emissions of CH₄ or N₂O from incineration of municipal solid waste (kgCH₄) (kg N₂O)
 EF_i : Emission factor for municipal solid waste incinerator type i (wet basis) (kgCH₄/t) (kg N₂O/t)
 A_i : Amount of municipal solid waste incinerated for incinerator type i (wet basis) (t)
 R : Percentage of municipal solid waste incinerated at facilities with energy recovery

Industrial Solid Waste

$$E = \sum (EF_j \times A_j \times R_j)$$

- E : Emissions of CH₄ or N₂O from incineration of industrial solid waste (kgCH₄) (kg N₂O)
 EF_j : Emission factor for industrial solid waste type j (wet basis) (kgCH₄/t) (kg N₂O/t)
 A_j : Amount of industrial solid waste type j incinerated (wet basis) (t)
 R : Fraction of industrial solid waste type j incinerated at ISW incineration facilities with energy recovery

● Activity Data converted into energy units (reference value)

Activity data converted into energy units to be reported in CRF was estimated as indicated below.

Municipal Solid Waste

$$A_E = A \times GCV \times R / 10^6$$

- A_E : Calorific value of activity data of MSW (TJ)
 A : Total amount of MSW incinerated (kg[wet])
 GCV : Gross calorific value of MSW (MJ/kg)
 R : Fraction of MSW incinerated at MSW incineration facility with energy recovery

Based on the actual measurement results obtained at municipality, the calorific value of MSW is 9.9 (MJ/kg).

Industrial Solid Waste

$$A_E = \sum A_j \times GCV_j \times R / 10^6$$

A_E : Calorific value of activity data of ISW (TJ)

A_j : Amount of ISW type j incinerated (kg[wet])

GCV_j : Gross calorific value of ISW type j (MJ/kg)

R : Fraction of ISW type j incinerated at ISW incineration facility with energy recovery

Calorific value of ISW is indicated in Table 8-65 (as referred to hereinafter).

c) Uncertainties and Time-series Consistency

Methodologies similar to that used in “8.4.1.1 Municipal Solid Waste Incineration (6.C.1)” and “8.4.1.2. Industrial Waste Incineration of (6.C.2)” were used.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials. For more details of QA/QC activities, see the Annex 6.

e) Source-specific Recalculations

Updating the amount of incinerated municipal and industrial waste, the emission estimates for the period FY2005 and FY2007-2009 were recalculated.

f) Source-specific Planned Improvements

No improvements are planned.

8.4.3. Emissions from direct use of waste as fuel (1.A.)**a) Source/Sink Category Description**

In this category, CO₂, CH₄, and N₂O emissions from waste directly used as fuel are estimated and reported. The reporting category for the emissions for each type of waste is, according to its use as fuel or raw material, either “Energy Industry (Category 1.A.1.)” or “Manufacturing and Construction (1.A.2)”. The fuel type is classified as “Other fuels”.

Greenhouse gas emissions during the direct use of waste as a raw material, such as plastics used as reducing agents in blast furnaces or as a chemical material in coking furnaces, or use of intermediate products manufactured using the waste as a raw material, are estimated in this category. The waste used as raw material and that used as fuel are combined and expressed as “Raw Material/Fuel Use” in this section.

Table 8 - 59 Estimation category for emissions from the direct use of waste as fuel

Emission source	Application breakdown	Major application	Reporting category of Energy sector	
Use of municipal solid waste (plastics) as alternative fuel or rawmaterial	Petrochemical	Fuel	1A2f	Other
	Blast furnace reducing agent	Reducing agent in blast furnace	1A2a	Iron & Steel
	Coke oven chemical feedstock	Alternative fuel or raw material in coke oven	1A1c	Manufacture of solid fuels
	Gasification	Fuel	1A2f	Other
Use of waste oil as alternative fuel or raw material	Cement burning	Cement burning	1A2f	Cement & Ceramics
	Other	Fuel	1A2f	Other
Use of industrial solid waste (waste plastics) as alternative fuel or raw material	Blast furnace reducing agent	Blast furnace reducing agent	1A2a	Iron & Steel
	Boiler	Fuel	1A2b	Chemicals
	Boiler	Fuel	1A2d	Pulp, paper and print
	Cement burning	Cement burning	1A2f:	Cement & Ceramics
	Boiler	Fuel	1A2f	Machinery
Use of industrial solid waste (waste wood) as alternative fuel or material	-	Fuel	1A2f	Other
Use of waste tire as alternative fuel or raw material	Cement burning	Cement burning	1A2f	Cement & Ceramics
	Boiler	Fuel	1A2f	Other
	Iron manufacture	Alternative fuel or raw materials in iron manufacturing	1A2a	Iron & Steel
	Gasification	Fuel in iron manufacturing	1A2a	Iron & Steel
	Metal refining	Fuel in metal refining	1A2b	Non-ferrous metals
	Tire manufacture	Fuel in tire manufacturing	1A2c	Chemicals
	Papermanufacture	Fuel in paper manufacturing	1A2d	Pulp, paper and print
	Power generation	Power generation	1A1a	Public electricity and heat production [*]

*Since the industry category for the use of it is not identified, "1A1a" is applied.

b) Methodological Issues

1) CO₂

● Estimation Method

Emissions were estimated by multiplying the incinerated volume of each type of waste used as raw material or fuel by Japan's country-specific emission factor. The wastes included in the estimation are the portions used as raw material or fuel of: plastics in MSW; waste plastics and waste mineral oil in industrial waste; and waste tires.

● Emission factor

Emission factors were established for the plastics from MSW that were used as chemical raw material in coke ovens and waste tires. The remaining emission sources used the emission factors for "Waste Incineration without Energy Recovery (Chapter 8.4.1.)".

Emission factors for this category	Plastics from municipal solid waste (as chemical raw material in coke ovens) and waste tires
Emission factors for incineration without energy recovery	Plastics from municipal solid waste (other than those used as chemical material in coke ovens) and industrial waste

Table 8 - 60 Country-specific CO₂ emission factors for this category

Item	Unit	1990	1995	2000	2005	2008	2009	2010
MSW-coke oven	kg CO ₂ /t(dry)	1,420	1,420	1,420	1,420	1,420	1,420	1,420
Waste tire	kg CO ₂ /t(dry)	1,858	1,785	1,790	1,737	1,725	1,729	1,750

● Activity Data

Details of the amount of waste used as raw material or alternative fuels, see the 8.4.3.1. - 8.4.3.3.

Table 8 - 61 Use of waste as raw materials or fuels for CO₂ emissions

Item	Unit	1990	1995	2000	2005	2008	2009	2010
MSW-plastics-oilification	kt (dry)	0	0	3	7	3	6	1
MSW-plastics-reducer in blast furnace	kt (dry)	0	0	24	35	17	26	25
MSW-plastics-chemical material in coke-oven	kt (dry)	0	0	10	168	136	144	171
MSW-plastics-gasification	kt (dry)	0	0	1	56	45	43	51
ISW-waste plastics (iron and steel)	kt (wet)	0	0	57	160	74	97	137
ISW-waste plastics (cement)	kt (wet)	0	0	102	302	427	440	413
ISW-waste plastics (boiler)	kt (wet)	21	14	16	9	18	19	19
ISW-waste mineral oil (cement baking furnace)	kt (wet)	137	225	343	423	384	372	436
ISW-waste mineral oil (boiler)	kt (wet)	554	633	460	811	724	640	672
Waste tire	kt (dry)	282	471	580	498	593	547	573

* The amount of biomass-based plastics and waste animal and vegetable oil are not included in any of the items in the table.

2) CH₄, N₂O

● Estimation Method

Emissions were estimated by multiplying the amount of each type of waste used as raw material or fuel by the country-specific emission factor. It should be noted that emissions from some of the emission sources are not estimated. They are summarized below.

Table 8 - 62 CH₄ and N₂O emissions sources not included in emission estimates or allocations

Emission source	Emission source (not calculated)
Use of municipal solid waste as alternative fuel or raw materials	Blast furnace reducing agent (NO), Coke-oven chemical feedstock (IE), Gasification (NE)
Use of industrial solid waste as alternative fuel or raw materials	Blast furnace reducing agent (NO), Petrochemical (NE), Gasification (NE)
Use of waste tire as alternative fuel or raw material	Iron manufacturing (NO)

● Emission factor

Emission factors for waste used as raw material and fuel were determined by multiplying the emission factor for applicable types of furnaces by the calorific value of each waste type, and converting the result to the weight-based values. Table 8-63 shows the data used in the estimation.

Calculation of emission factor (wet basis)

$$= (\text{Emission factor for each type of furnace (kg-CH}_4\text{/TJ, kg-N}_2\text{O/TJ)}) \times (\text{Calorific value of each waste type (MJ/kg)}) / 1000$$

Table 8 - 63 Data used for the calculation of CH₄ and N₂O emission factors for wastes used as raw material and fuel

Item		Emission factor for furnaces and ovens (Energy sector)	Calorific value
Plastics from municipal solid waste	Plastic oil	Boilers (Heavy fuel oil A, gas oil, kerosene, naphtha, other liquid fuels)	Calorific value of waste plastics
Industrial waste	Waste plastics	Cement kilns	Other industrial furnaces (solid fuel)
		Boilers	CH ₄ : Boilers (wood, charcoal, and other solid fuel) N ₂ O: Boilers (other than fluidized-bed) (solid fuel)
	Waste oil (mineral/animal and vegetable)	Cement kilns, boilers	Other industrial furnaces (solid fuel)
		Boilers	Boilers (Heavy fuel oil A, gas oil, kerosene, naphtha, other liquid fuels)
Wood scraps	Boilers	CH ₄ : Boilers (wood, charcoal) N ₂ O: Boilers (other than fluidized-bed) (solid fuel)	
Waste tires	Cement kilns	Other industrial furnaces (solid fuel)	Calorific value of waste tires
	Boilers	CH ₄ : Boilers (Steam coal, coke, other solid fuels) N ₂ O: Boilers (other than fluidized-bed) (solid fuel)	
	Carbonization	Boilers (gas fuels)	
	Gasification	Other industrial furnaces (gas fuels) and other industrial furnaces (liquid fuels) ^{c)}	

a) Calorific value per unit volume was determined by dividing by the specific gravity of waste oil (0.9 kg/L) obtained from the Waste Handbook (1997).

b) Source: 1997 General Survey of Emissions of Air Pollutants

c) The percentage of substances recovered during the gasification of waste tires. A weighted average was calculated using the proportions of gas and oil (22% and 43%) reported in the Hyogo Eco-town documents.

Table 8 - 64 CH₄ and N₂O emission factors for the use of waste as raw material or fuel used in the Energy sector

Furnace type/Fuel type	CH ₄ Emission factor (kg-CH ₄ /TJ)	N ₂ O Emission factor (kg-N ₂ O/TJ)
Boilers (Heavy fuel oil A, gas oil, kerosene, naphtha, other liquid fuels)	0.26	0.19
Boilers (gas fuels)	0.23	0.17
Boilers (steam coal, coke, other solid fuels)	0.13	
Boilers (wood, charcoal)	74.9	
Boilers (other than fluidized-bed) (solid fuels)		0.85
Other industrial furnaces (liquid fuel)	0.83	1.8
Other industrial furnaces (solid fuel)	13.1	1.1
Other industrial furnaces (gas fuel)	2.3	1.2

Emission factors are from Chapter 3, Energy.

Table 8 - 65 Calorific Value of waste incinerated and used as raw material or fuel

Item	Unit	GCV	Source of calorific value	
Waste oil (including reclaimed oil)	TJ/l	40.2	Reference # 22; estimated with 0.9(kg/l) from Reference # 46	
Waste plastics	MJ/kg	29.3	Reference # 22	
Waste paper	MJ/kg	15.1	Reference # 47 (dry basis); value was obtained by subtracting water content	
Waste wood	MJ/kg	14.4	Reference # 22	
Waste textile	MJ/kg	17.9	Reference # 47 (dry basis) ; value was obtained by subtracting water content	
Animal & vegetable residues/animal carcasses	MJ/kg	4.4	Reference # 47 (dry basis) ; value was obtained by subtracting water content	
Sludge (including sewage sludge)	MJ/kg	4.7	Reference # 22 (dry basis) ; value was obtained by subtracting water content	
Waste tires	2004 and before	MJ/kg	20.9	Reference # 22
	2005 and later	MJ/kg	33.2	Reference # 22
RDF	MJ/kg	18	Reference # 22	
RPF	MJ/kg	29.3	Reference # 22	

● Activity Data

Waste used as raw material and fuel

Activity data were determined for each category using the wet-basis values (Table 8-66). For more details, see each section.

Table 8 - 66 Amount of waste used as raw material or fuel for CH₄ and N₂O emissions

Item	Unit	1990	1995	2000	2005	2008	2009	2010
MSW-oilification	kt (wet)	0	0	3	7	3	7	1
ISW-waste wood	kt (wet)	1,635	1,635	2,061	2,683	3,724	3,725	3,459
ISW-waste mineral/animal & vegetable oil (cement baking furnace)	kt (wet)	141	233	359	447	408	396	464
ISW-waste mineral/animal & vegetable oil (boiler)	kt (wet)	569	657	482	858	770	681	715
Waste tire-cement baking furnace	kt (wet)	111	275	361	181	141	112	95
Waste tire-boiler	kt (wet)	119	184	163	255	394	387	428
Waste tire-pyrolysis furnace	kt (wet)	67	37	30	10	2	1	1
Waste tire-gasification	kt (wet)	0	0	0	27	48	48	49

See Table 8-61 for the activity data for ISW-waste plastics (cement manufacturer) and ISW-waste plastics (boiler).

Activity Data converted into energy units (reference value)

Activity data converted into energy units to be reported in CRF are calculated as indicated below.

Activity data converted into energy units

$$= (\text{Amount of waste used as raw material or fuel (kg [wet])}) \times \text{Corresponding calorific value of waste (MJ/kg)} / 10^6$$

c) Uncertainties and Time-series Consistency

See the respective section.

d) Source-specific QA/QC and Verification

See the respective section.

e) Source-specific Recalculations

See the respective section.

f) Source-specific Planned Improvements

See the respective section.

8.4.3.1. Emissions from municipal waste (waste plastics) used as alternative fuel (1.A.1 and 1.A.2)

a) Source/Sink Category Description

This category covers the emissions from municipal waste (waste plastics) used as raw materials or alternative fuels. Plastics in MSW collected under the Containers and Packaging Recycling Law are processed into petrochemical, blast furnace reducing agent, chemical raw material in coke-oven, and gasification to be used as alternative fuel or raw material.

*b) Methodological Issues**1) CO₂*

- *Estimation Method*

Emission estimates were calculated by multiplying the amount of fossil-fuel derived plastics in MSW by each usage (petrochemical, blast furnace reducing agent, chemical raw material in coke-oven, and gasification) by Japan's country-specific emission factor.

- *Emission factor*

For the emission factors for plastics in MSW in the usage of petrochemical, blast furnace reducing agent, and gasification, the same values applied in "8.4.1.1. Municipal Solid Waste Incineration (6.C.1)" were applied. The emission factor for plastics used as chemical raw material in coke ovens was set as the volume of hydrocarbon that is used as chemical raw material and from which no CO₂ is emitted into the air by subtracting the percentage of carbon in the plastics that migrates to hydrocarbon oil in the coke oven (47.9%) from emission factor for plastics (MSW).

Calculation of the emission factor for plastics used as raw material in coke ovens (dry basis)

= (Emission factor for the incineration of plastics in municipal solid waste)
 × [1 – (Fraction of carbon in plastics used as chemical raw material for coke ovens that migrates to hydrocarbon)]

- *Activity Data*

The amount of plastics in MSW used as raw material or fuel by usage (wet basis) was estimated by the total amount collected by designated legal bodies and municipalities to be processed as raw material or fuel by usage (wet basis). The amount of waste plastics (dry basis) was estimated by subtracting water content from the amount of waste plastics (wet basis). The amount of fossil-fuel derived

plastics in MSW used as raw material or fuel (dry basis) was obtained by subtracting the amount of biomass-based plastics in MSW used as raw material or fuel assuming that all of the biomass-based plastics content in each usage is the same.

Amount of fossil-fuel derived waste plastics used as raw material or fuel by usage (dry basis)

= Amount of waste plastics used as raw material or fuel by usage (wet basis)

× (1 – Water content in waste plastics)

– Amount of biomass-based plastics in MSW used as raw material or fuel (dry basis)

× Amount of MSW used as raw material or fuel by usage / amount of waste plastics in MSW used as raw material or fuel

As in the CO₂ emission estimates in “Municipal Solid Waste Incineration (6.C.1)”, the amount of biomass-based plastics in MSW used as raw material or fuel was estimated as indicated below.

Amount of biomass-based plastics in MSW used as raw material or fuel (dry basis)

= Amount of biomass-based plastic products consumed (dry basis)

× Fraction of biogenic component × Fraction of biomass-based plastics disposed of as MSW

× Fraction of biomass-based plastics used as raw material or fuel

■ ***The amount of plastics in MSW used as raw material or fuel by usage (wet basis)***

Processing of plastics collected by designated legal bodies

The amount of the plastics in MSW collected by designated legal bodies into raw material and fuel was determined from the amount reported (pyrolytic oil: petrochemical, blast furnace reducing agent, chemical raw material in coke-oven, syngas, and gasification) in the “Plastic Containers and Packaging (Other Plastics, Food Trays)” section of the *Statistics of Commercial Recycling of Plastics (Recycling)* compiled by the Japan Containers and Packaging Recycling Association. Usage in products that do not emit CO₂ was deducted.

Processing of plastics collected by municipalities

The amount of plastics in MSW collected by municipalities and processed into raw material or fuel was calculated by first subtracting the amount of plastics (wet basis) that was commercially recycled through designated legal bodies¹ from the amount of all plastics that were commercially recycled under the Plastic Containers and Packaging Recycling Law (wet basis)². The amount of waste plastics used as raw material or fuel by usage was estimated by multiplying the amount of plastics in MSW collected by municipalities to be processed into raw material and fuel by the fraction of plastic content by usage³ and the fraction of amount of commercially recycled products⁴.

¹: Amount of plastics commercially recycled through designated legal body channels (wet basis)

The amount was determined from the “Actual Collection of Plastic Containers and Packages” section of the *Statistics of Commercial Recycling of Plastics (Recycling)*.

²: Amount of plastics commercially recycled under the Plastic Containers and Packaging Recycling Law (wet basis)

The results of the selective collections by municipalities and commercial recycling under the Plastic Containers and Packaging Recycling Law were determined from Annual Recycling Statistics by the

Waste Management and Recycling Department of the Ministry of the Environment.

³: Percentage of commercially recycled plastics by recycling method

The rates were obtained from the percentages for various methods of commercial recycling of the plastics collected through municipal channels in the Results of the 2001 Questionnaire to Municipalities on Waste Plastics Processing compiled by the Plastic Waste Management Institute.

⁴: Percentage of commercially recycled plastic products by recycling method

The values for the commercial recycling of the plastics collected through the municipal channels were substituted for the percentage of commercially recycled plastic products collected through designated legal body channels. The percentages were calculated by dividing the amounts of commercially recycled plastic products by various recycling methods, which were established in the activity data for recycling through designated legal body channels, by the amount of commercially recycled plastics. The amount of commercially recycled plastics by each of the recycling methods was calculated by multiplying the amount of plastics commercially recycled through designated legal body channels, by the percentage of commercially recycled plastics by recycling method obtained from the Assessment and Deliberation of the Plastic Containers and Packaging Recycling Law, the Japan Containers and Packaging Recycling Association.

■ **Water content ratio**

Water content ratio of 4% was determined based on the data provided by the *Japan Containers and Packaging Recycling Association*.

■ **Amount of biomass plastics products consumed**

See the section “Municipal Solid Waste Incineration (6.C.1).”

■ **Fraction of biogenic component in biomass-based plastics disposed of as MSW**

See the section “Municipal Solid Waste Incineration (6.C.1).”

■ **Fraction of waste plastics used as raw material or fuel**

See the section “Municipal Solid Waste Incineration (6.C.1).”

2) **CH₄, N₂O**

For estimation method and emission factors, see the section “Emissions from Direct Use of Waste as Fuel (8.4.3)”. The amount of waste plastics used as raw material or fuel by usage (wet basis) was determined by the total amount collected by designated legal bodies and municipalities to be processed as raw material and fuel by usage (wet basis); this value includes the amount of biomass-based plastics consumed.

c) **Uncertainties and Time-series Consistency**

● **Uncertainties**

The same value of uncertainty in “CO₂ emissions from incineration of MSW (6.C.1.a)” was used for the uncertainty in the CO₂ emission factor. The uncertainty in activity data for CO₂ emissions was estimated by using the uncertainties in the amount of plastics used as raw materials or alternative fuels

(statistical uncertainty) and the percentage of water content (same value that was used for the MSW incineration).

The uncertainty in the CH₄ emission factor was estimated by using the uncertainties in emission factors and the calorific value of plastics. For uncertainty in CH₄ and N₂O activity data, the uncertainties in the amount of MSW plastics used as raw materials or alternative fuels were used. The uncertainties in the CO₂, CH₄ and N₂O emissions from MSW plastics used as raw materials or alternative fuels were estimated to be 17%, 180% and 112%, respectively. For details, see the Annex 7.

● *Time series consistency*

Time series consistency in emission estimates has been ensured. However, the statistical data for activity data have been available since FY 2000 because the use of waste as alternative fuel or raw material was not a common practice prior to FY 2000 in Japan.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials. For more details of QA/QC activities, see the Annex 6.

e) Source-specific Recalculations

Updating the amount of biomass-based plastic products consumed, emission estimates for FY2007-2009 were recalculated.

f) Source-specific Planned Improvements

No improvements are planned.

8.4.3.2. Emissions from industrial waste (waste plastics, waste oil, and waste wood) used as raw material or alternative fuels (1.A.2.)

a) Source/Sink Category Description

This category covers greenhouse gas emissions from industrial waste (waste plastics, waste oil, and waste wood) used as raw material or alternative fuels.

b) Methodological Issues

1) CO₂

● *Estimation Method and Emission factor*

Emissions were estimated by multiplying the incinerated amount of waste plastics and waste mineral oil used as raw material or alternative fuels by emission factor used for incineration of ISW.

● *Activity Data*

Industrial waste plastics

Estimated activity data were the amounts of waste plastics (wet basis) in industrial waste used as raw material or fuel in steel industry, chemical industry, paper industry, cement Manufacturer, and

automobile manufacturer. The amount of waste plastics in industrial waste used as raw material or fuel in each industry was provided by the following data sources: for steel industry, the *Current State of Plastic Waste Recycling and Future Tasks* published by the Japan Iron and Steel Federation; for cement manufacturing industry, from the *Cement Handbook* published by the Japan Cement Association; for chemical industry, paper industry, and automobile manufacturer, the amount of waste plastics used for fluid bed boiler provided by the Japan Chemical Industry Association, the Japan Paper Association, the Japan Automobile Manufacturers Association. All of the waste plastics in ISW was considered to be fossil-fuel derived.

Waste mineral oil

Activity data were estimated by subtracting the amount of biogenic-origin waste oil indicated as “Fraction of Animal and Vegetable Origin Waste Oil” provided by the survey conducted by the Ministry of the Environment from the amount of waste oil indicated as “Fuel Usage” of “Direct Recycle Usage” and “Recycle Usage after Treatment” of ISW provided by the *Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes*. The activity data for FY1997 and before were estimated by using the trend of the amount of incinerated industrial waste oil.

2) CH₄, N₂O

● Estimation Method and Emission factor

See the section “Emissions from Direct Use of Waste as Fuel (8.4.3)”

● Activity Data

Waste plastics

Estimated activity data were the amounts of waste plastics used for cement kilns and boilers. Out of the activity data used for CO₂ emission estimates from this source, the amount used as raw materials and fuels in chemical industry, paper industry, cement manufacturer, and automobile manufacturer were used for CH₄ and N₂O emission estimates. Because blast furnace gas generated from steel industry is entirely recovered and not included in the activity data.

Waste oil (Mineral / Animal and Vegetable)

The amount of waste oil used as raw material or fuel is calculated separately for cement kilns and boilers. The amount of waste oil and reclaimed oil, which was produced from the waste oil contained in industrial waste and other waste oil, used as fuel for cement kilns was determined from the annual data in the *Cement Handbook*. The amount used as fuel for boilers was determined by subtracting the amount used as fuel for cement kilns from the amount of waste oil indicated as “Fuel Usage” of “Direct Recycle Usage” and “Recycle Usage after Treatment” of ISW provided by the *Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes*.

Unlike the activity data for CO₂ emissions, waste mineral oil and also waste animal and vegetable oil are included for the estimation of activity data from this source.

Waste wood

The amount of usage of waste wood as raw material or fuel was obtained from the “fuel usage” in the “direct recycle usage” and the “fuel usage” in the “recycle usage after treatment” in the *Report of the*

Research on the State of Wide-range Movement (the volume on Cyclical Use). The values before FY 1997 are estimated by using the average value in the period of FY 1998-2002.

c) Uncertainties and Time-series Consistency

● *Uncertainties*

The same value of uncertainty as was used for “CO₂ emissions from incineration of industrial waste (6.C.1.b)” was applied to uncertainty in CO₂ emission factor. The uncertainties in emission factors for CH₄ and N₂O were evaluated by the same method that was used for municipal waste used as raw materials or alternative fuels. The uncertainty in activity data were evaluated separately for waste plastics, waste oil, and waste wood. For waste plastics, the uncertainty was calculated by combining of the uncertainties in the amount of waste plastics used as raw materials or alternative fuels in the iron and steel industry and in the cement industry. The uncertainty levels for each component were evaluated by using the statistical uncertainties. For waste oil, the values for cement kilns (statistical uncertainty) and boilers (a value for CO₂) were combined. For waste wood, statistical uncertainties for the amount of waste wood used as raw materials or alternative fuels were used.

The uncertainties in CO₂, CH₄ and N₂O emissions from the incineration of industrial waste used as raw material or alternative fuels were estimated to be in the range of 13-105%, 74-128% and 31-110%, respectively. For details, see the Annex 7.

● *Time series consistency*

Data on the amount of waste oil and waste wood used as alternative fuels have been available since FY 1998. For waste oil, consistent data over the time series were developed by using the total amount of waste oil incinerated without the use of waste oil as alternative fuel. For waste wood, the average of FY 1998–2002 data was used to estimate the amount of waste wood for the past years. The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials. For more details of QA/QC activities, see the Annex 6.

e) Source-specific Recalculations

Updating the amount of industrial waste used as raw material or fuel, emission estimates for FY2008 were recalculated.

f) Source-specific Planned Improvements

No improvements are planned.

8.4.3.3. Emissions from waste tires used as raw materials and alternative fuels (1.A.1 and 1.A.2)

a) Source/Sink Category Description

This category includes the emissions from the use of waste tires as raw materials or alternative fuels.

b) Methodological Issues

1) CO₂

● Estimation Method

The emissions were calculated by multiplying the incinerated amount of waste tires used as raw materials or fuels by Japan's country-specific emission factor.

● Emission factor

The emission factor for waste tires was calculated by multiplying the fossil fuel-derived carbon content of the waste tires by the efficiency of combustion of the waste tires at the facilities that use waste tires as fuel. The volume of the fossil fuel-derived carbon in the waste tires was calculated by the material contents of new tires. The efficiency of combustion for waste tires was set to 99.5% based on the maximum default value for hazardous waste in the *GPG (2000)*.

$$\begin{aligned} & \text{Calculation of emission factor for the incineration of waste tires (dry basis)} \\ & = (\text{Fossil fuel-derived carbon content in waste tires}) \times (\text{Efficiency of combustion of waste tires}) \\ & \times 1000 \times 44 / 12 \end{aligned}$$

● Activity Data

Activity data (dry basis) was calculated by subtracting the water content in the waste tires determined from analyses of three constituents of divided tires reported in *the Basic Waste Data Fact Book (2000)* published by Japan Environmental Sanitation Center from the amount of waste tires used as raw material or fuel (wet basis) in the *Tire Industry of Japan (32)*, published by the Japan Automobile Tire Manufacturers Association, Inc.

2) CH₄, N₂O

● Estimation Method and Emission factor

See the section 8.4.3.

● Activity Data

The volume of waste tires used as raw material or fuel by usage that was determined during the calculation of the CO₂ emissions from this source was used. For the activity data, the volume of waste tires recorded in the following categories were used: "Cement kilns" for use in cement kilns; "Medium to small boilers", "Use by tire factories", "Use by paper manufacturers", and "Power generation" for use in boilers; "metal refining" for use in carbonization; and "Gasification" for use in gasification processes.

c) Uncertainties and Time-series Consistency

● Uncertainties

The level of uncertainty in CO₂ emission was estimated by using the carbon content of waste tires and the combustion rate of the furnace using waste tires as alternative fuels. For activity data, the uncertainty was estimated by using the uncertainties in the amount of waste tires used as raw materials or alternative fuels and the percentage of water contents in waste tires. The uncertainties in the

emission factors for CH₄ and N₂O were evaluated by the same method that was applied to MSW used as raw materials or alternative fuels and were estimated by combining the uncertainties in emission factors (CH₄, N₂O of the Energy sector) using waste tires as raw materials or alternative fuels and in the calorific value of waste tires. For activity data, the uncertainties in the amount of waste tires used as raw materials or alternative fuels were used. The methods of evaluation of the uncertainty levels for each component are:

- Use of the values for industrial waste (waste plastics) incineration: carbon content and combustion rate
- Based on expert judgment: percentage of water contents
- Use of the uncertainties set by each statistics: amount of waste tires used as raw materials or alternative fuels

The uncertainties in CO₂, CH₄ and N₂O emissions from the use of waste tires as raw materials or alternative fuels were estimated to be 15%, 91% and 26%, respectively. For details, see the Annex 7.

● *Time series consistency*

The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials. For more details of QA/QC activities, see the Annex 6.

e) Source-specific Recalculations

No recalculations were conducted.

f) Source-specific Planned Improvements

No improvements are planned.

8.4.4. Emissions from incineration of waste processed as fuel (1.A.)

8.4.4.1. Incineration of refuse-based solid fuels (RDF and RPF) (1.A.1 and 1.A.2)

a) Source/Sink Category Description

In this category, CO₂, CH₄, and N₂O emissions from waste that is processed and used as fuel are estimated and reported. Refuse-derived solid fuels (RDF as Refuse Derived Fuel and RPF as Refuse Paper and Plastic Fuel) are used for the estimation of emissions from fuels produced from waste. The reporting categories for the above emissions are included in “Energy Industry (1.A.1)” and “Manufacturing/Construction (1.A.2)” according to the use of waste as fuels. The fuel type is classified as “Other fuels”.

Table 8 - 67 Estimation category for emissions from the use of waste processed as fuel

Emission source	Application breakdown	Major application	Reporting category of Energy sector
Use of refuse-derived fuel (RDF · RPF)	RDF	Fuel use (including power generation)	1A2f Other [*]
	RPF (petroleum products)	boiler fuel	1A1b Petroleum refining
	RPF (chemical industry)	boiler fuel	1A2c Chemicals
	RPF (paper manufacture)	Fuel use in paper manufacturing	1A2d Pulp, paper and print
	RPF (cement burning)	Cement burning	1A2f Cement & ceramics

* : Emissions from power generation and heat supply excluding in-house use should be included in the category 1A1a. However, they are reported in the category 1A2f, because the actual circumstances are not understood at the moment.

b) Methodological Issues

1) CO₂

● Estimation Method

Emissions were estimated by multiplying the incinerated amount of RDF and RPF by Japan's country-specific emission factor.

● Emission factor

Emission factor associated with the use of the refuse-derived solid fuels (RDF and RPF) was calculated separately for RDF and RPF by the equation shown below. For the RPF (refuse paper and plastic fuel), the emission factors were calculated separately for the coal-equivalent and coke-equivalent fuels, and also calculated their average weighted by the percentage used as fuel.

<p><i>Calculation of emission factor for the use of RDF and RPF as fuel (dry basis)</i> $= 1000 \times (1 - \text{Average percentage of water content}) \times (\text{Percentage of plastic-derived constituents, dry basis}) \times (\text{Carbon content of plastics, dry basis}) \times (\text{Efficiency of combustion}) \times 44 / 12$</p>
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Average percentage of water content

Percentage of water contents in the RDF was set to 5.5%, based on the simple average of water content in the RDF manufactured by the facilities listed in the *Proper Management of Refuse-derived Fuels* compiled by the Study Group for Proper Management of RDF.

Percentage of water contents in the RPF was set to 2.6%, based on the water contents of coal-equivalent and coke-equivalent products indicated by the RPF quality standards set by the Japan RPF Industry Association with their average weighted by the manufacturing ratio of these products.

Percentage of plastic-derived content

Calculation of the percentage of the plastics-derived constituents (dry basis) used the wet-based moisture content of the constituents of MSW determined in the "Emission from Controlled Disposal Sites (6.A.1.)" section, which was converted to a dry-based value. The results of the content analysis of the wet-based refuse were obtained from the *Results of Content Analysis of Refuse* for each facility listed in the "Proper Management of Refuse-derived Fuels". The percentage of plastics-derived constituents in the RPF (dry basis) was set at 50% for the coal-equivalent product and 90% for the coke-equivalent product based on the results of a fact-finding survey by the Japan RPF Industry Association.

Carbon content in plastics

Average carbon content used in the “Incineration of Municipal Solid Waste (Plastics)” (Table 8 - 37)” was applied to the carbon content in plastics contained in the RDF (dry basis). The carbon content (73.7%) of plastics contained in the RPF (dry basis) was determined from the carbon content value (70%) used in the “Incineration of Industrial Waste (Waste Plastics)” (95%), which was converted to a dry basis using the moisture content in waste plastics in industrial waste.

Efficiency of combustion

Rate of combustion of the RDF was set to 99%, applying the default value in the *GPG (2000)* in the same manner as for MSW (plastics). The rate for the RPF was set to 99.5%, using the default value in the *GPG (2000)* in the same manner as for industrial waste (waste plastics).

Table 8 - 68 CO₂ emission factors for the emissions from the use of refused-derived fuel (RDF) or refuse paper & plastic fuel (RPF)

Item	Emission Factor [kg CO ₂ /t (dry)]
RDF	808
RPF (coal-equivalent products)	1,419
RPF (coke-equivalent products)	2,445
RPF (weighted average values)	1,627

● Activity Data

RDF

The amount of RDF production was used as the substitute for the amount of use of RDF. Activity data (dry basis) was calculated by subtracting the water content of RDF from the amount of RDF production at RDF production facilities (wet basis) provided by the *Report on Survey of State of Treatment of Municipal Solid Waste* and also subtracting the amount of biomass-based plastics used as RDF. For the fiscal years that the data were unavailable, emission estimates were conducted substituting the values of the refuse processing capacity.

$$\begin{aligned} & \text{Activity data for the use of RDF (dry basis)} \\ & = \text{Amount of use of RDF (wet basis)} \times (1 - \text{water content of RDF}) - \text{Amount of biomass-based} \\ & \text{plastics used as RDF (dry basis)} \end{aligned}$$

As in the activity data for CO₂ emission estimates in “Municipal Solid Waste Incineration (6.C.1)”, the amount of biomass-based plastics used as RDF was estimated as indicated below.

$$\begin{aligned} & \text{Amount of biomass-based plastics used as RDF (dry basis)} \\ & = \text{Amount of import of biomass plastics (dry basis)} \times \text{Fraction of biogenic component} \\ & \times \text{Fraction of biomass-based plastics disposed of as MSW} \times \text{Fraction of use of RDF} \end{aligned}$$

Amount of biomass-based plastics products consumed

See the section “Municipal Solid Waste Incineration (6.C.1).”

Fraction of biogenic component and biomass-based plastics disposed of as MSW

See the section “Municipal Solid Waste Incineration (6.C.1).”

Fraction of use of RDF

See the section “Municipal Solid Waste Incineration (6.C.1).”

RPF

The amounts of RPF used in chemical industry, paper industry, cement manufacturer, and petroleum product manufacturer were estimated. The amount of RPF (dry basis) for paper industry was obtained from the survey results conducted by the Japan Paper Association. The amounts of RPF (dry basis) for chemical industry, cement manufacturer, and petroleum product manufacturer were obtained by using the average water content of RPF and also the survey results (wet basis) conducted by the Japan Cement Association and the Japan Automobile Manufacturers Association. All of the plastics included in RPF was considered to be fossil-fuel derived.

Table 8 - 69 Amount of use of refuse-derived fuel (RDF) or refuse paper & plastic fuel (RPF) (wet basis)

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Refuse-derived fuel (RDF)	kt (dry)	32	37	140	392	365	355	355
Refuse paper & plastic fuel (RPF)	kt (dry)	0	8	32	478	749	776	760

2) CH₄, N₂O**● Estimation Method and Emission factor**

For the estimation method and the emission factors used, see “Emissions from Direct Use of Waste as Fuel (8.4.3)”.

Table 8 - 70 Data used for the calculation of the methane and nitrous oxide emission factors for wastes used as raw material and fuel

Item		Emission factor for furnaces and ovens (Energy sector)	Calorific value
RDF	Boilers	CH ₄ : Boilers (Steam coal, coke, other solid fuels) N ₂ O: Boilers (other than fluidized-bed) (solid fuel)	Calorific value of RDF
RPF	Cement kilns, boilers	Other industrial furnaces (solid fuel)	Calorific value of RPF *
	Boilers	CH ₄ : Boilers (Steam coal, coke, other solid fuels) N ₂ O: Boilers (other than fluidized-bed) (solid fuel)	

*Weighted average of calorific values calculated based on the manufacturing ratio of Coal substitution RPF and Coke substitution RPF given by the Japan RPF Industry Association

● Activity Data**RDF**

The entire amount of RDF production (wet basis) used for CO₂ emission estimates was also used for the amount of use of RDF for boiler. The said amount includes the amount of biomass-based plastics.

RPF

Out of the amount of RPF used for CO₂ emission estimates, the amounts of RPF used in chemical industry, paper industry, and petroleum products manufacturer were applied to the amount of RPF used for boiler (wet basis). The amount of RPF used in cement industry was applied to the amount of RPF used for cement kiln (wet basis). Because the amount of RPF used in paper industry is on a dry basis, the average water content of RPF was added to obtain the value on a wet basis.

Activity data converted into energy units (reference value)

Activity data converted into energy units to be reported in CRF is calculated as indicated below.

$$\begin{aligned} & \text{Activity data converted into energy units} \\ & = (\text{Amount of RDF \& RPF consumed (kg [wet basis])}) \times \text{calorific value of corresponding fuel} \\ & \quad (\text{MJ/kg}) / 10^6 \end{aligned}$$

c) Uncertainties and Time-series Consistency**● Uncertainties**

The level of uncertainty in the CO₂ emission factor for RDF used as fuels was estimated by using the uncertainties in the percentage of plastic-derived constituents in RDF, carbon content in the plastics, and combustion rate of the facilities using RDF as fuels. For RPF, the uncertainty in emission factor for coal-equivalent RPF was used. The uncertainty in activity data was estimated by combining the uncertainty for each element because the activity data were estimated by subtracting water content from the amount of RDF and RPF used as fuels (wet basis) to obtain the values on a dry basis.

The uncertainties in the CH₄ and N₂O emission factors were estimated by using the uncertainties in emission factors by usage of RDF and RPF and the calorific values of the RDF and RPF. For activity data, the uncertainties in the amount of RDF and RPF were used.

The methods of evaluation of the uncertainty levels for each component are:

- Use of 95% confidence interval of data: percentage of plastic-derived constituents of RDF, percentage of water content in RDF
- Use of the values for MSW (plastics) incineration: carbon content of RDF and combustion rate for RDF
- Use of the values for ISW (waste plastics) incineration: carbon content of RPF and combustion rate for RPF
- Expert judgment: percentage of plastic-derived constituents of RPF
- Use of the uncertainties set by each statistics: amount of RDF and RPF used as alternative fuels

The uncertainties in CO₂, CH₄, and N₂O emissions from the use of RDF and RPF as raw materials or alternative fuels were estimated to be 44%, 49%, and 33%, respectively. For details, see the Annex 7.

● Time-series consistency

Because data on the amount of RDF produced were not available for the years prior to FY 1997, these data were estimated by using the trend on capacity of refuse-based fuel-producing facilities. The emissions were calculated in a consistent manner.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials. For more details of QA/QC activities, see the Annex 6.

e) Source-specific Recalculations

Updating the amount of biomass-based plastic products consumed, CO₂ emission estimates for FY2007-2009 were recalculated.

f) Source-specific Planned Improvements

No improvements are planned.

8.5. Other (6.D.)

In this category, CO₂ emissions as a result of the decomposition of petroleum-derived surfactants and CH₄ and N₂O emissions from the composting of organic waste are calculated. Estimated greenhouse gas emissions from category 'Other' are shown in Table 8-71. In FY 2010, emissions from this source category were 847 Gg-CO₂ eq. and accounted for 0.07% of the national total emissions (excluding LULUCF). The emissions from this source category had decreased by 7.3% compared to those in FY 1990. This emission decrease is primarily due to the decrease in CO₂ emissions for FY2001 through FY2004 from the use of alkylbenzenes by introduction of the Pollutant Release and Transfer Register (PRTR).

Table 8 - 71 GHG emissions from category 'Other' (6.D.)

Gas	Category	Unit	1990	1995	2000	2005	2008	2009	2010
CO ₂	6.D.2. Decomposition of petroleum-derived surfactants	Gg CO ₂	703	668	656	507	530	514	528
CH ₄	6.D.1. Composting of organic waste	Gg CH ₄	5.3	5.1	4.6	6.0	7.5	5.6	8.0
		Gg CO ₂ eq	112	106	96	126	157	118	169
N ₂ O		Gg N ₂ O	0.32	0.30	0.27	0.36	0.45	0.34	0.48
		Gg CO ₂ eq	99	94	85	112	139	105	150
Total of all gases		Gg CO ₂ eq	914	868	837	744	826	737	847

8.5.1. Emissions from Composting of Organic Waste (6.D.1)**a) Source/Sink Category Description**

Part of the MSW and industrial waste generated in Japan is composted, and CH₄ and N₂O generated in that process are emitted from composting facilities. Emissions from composting of livestock waste are accounted for under "Emissions from manure treatment (4.B)" in the agriculture sector.

b) Methodological Issues**● Estimation Method**

Emissions were calculated by taking the amount of organic waste composted, which was obtained

from the statistical information available in Japan, and multiplying it by the default emission factor provided in the *IPCC 2006 Guidelines*. The calculation method is the same for both CH₄ and N₂O emissions.

$$E = EF \times A$$

E : Amount of CH₄ (N₂O) emissions generated by composting organic waste (kg CH₄ or kgN₂O)

EF : Emission factor for (dry basis) (kg CH₄/t, (kg N₂O/t)

A_{dry} : Amount of composted organic waste (dry basis)

● *Emission factor*

In accordance with the *2006 IPCC Guidelines*, emission factors (dry basis) are set as 10.0 (kg CH₄/t) for CH₄ and 0.6 (kg N₂O/t) for N₂O, respectively, for all fiscal years.

● *Activity data*

Activity data (amount composted on a dry basis) was obtained by subtracting the water content appropriate to the properties of composted waste from the amount of composted waste (wet basis) listed below:

Municipal Solid Waste

- Amount of composted waste by waste types calculated by multiplying the amount of MSW treated at waste composting facilities indicated in the *Waste Treatment in Japan* by the fraction of waste types in MSW treated at high-rate composting facilities provided in the *Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes*.
- Amount of human waste composted at waste composting facilities indicated in the Ministry of the Environment, Waste Management and Recycling Department, *The state of municipal waste treatment survey*.

Industrial Solid Waste

- Amount of sludge treated at composting facilities provided by the *Sewage Statistics*
- Amount of composted animal and plant residues generated by food and beverage manufacturing is obtained in the *Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes*.
- Amount of composted food waste* other than the above is obtained in the *Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes*.*

*Although under the *Waste Disposal and Public Cleansing Act*, they fall under the category of municipal waste, it is included in industrial waste because of its source and properties.

Percentage of water content in composted waste, as indicated in the “Emissions from Controlled Disposal Sites (6.A.1)” section, are; 20% in waste paper, 75% in kitchen waste, 20% in textile waste, 45% in waste wood, and 70% in sewage sludge.

Table 8 - 72 Amounts of composted waste

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Municipal solid waste	kt (dry)	38	22	29	36	54	68	68
Industrial solid waste	kt (dry)	494	485	429	564	693	495	736

c) Uncertainties and Time-series Consistency

● *Uncertainties*

The uncertainty in emission factor was estimated by using the upper and lower limits for the uncertainty range provided in the *2006 IPCC Guidelines*. For activity data, uncertainty was evaluated on the basis of the statistical uncertainties. The uncertainties in CH₄ and N₂O emissions from composting of organic wastes were estimated to be 74% and 86.3%, respectively. For more details, see the Annex 7.

● *Time series consistency*

The amount of composted animal and plant residues generated by food manufacturing and food waste other than those for the period FY1990-2006 are estimated based on the results of the “the *Committee for Improving Survey on Cyclical Use of Wastes, FY2009* (MoE)”. Since the results for FY2008 are unavailable, the data for FY2007 are also applied for FY2008; thus, time series consistency in emission estimates has been ensured.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials.

e) Source-specific Recalculations

- Estimating the amount of composted animal and plant residues and other food waste, emission estimates for FY1990-2009 were recalculated.
- Updating the ratio of waste disposed of into high speed composting facility, emission estimates for the period FY2008-2009 were recalculated.

f) Source-specific Planned Improvements

- For future inventories, detailing of emission estimates will be conducted upon new scientific findings because the necessity of establishing country-specific emission factor from this source has been well recognized.
- The implementation of emission estimates from domestic and commercial composting machine will be further considered because this kind of research could not be completed in a short period of time, and a long-term efforts on scientific investigations will be necessary.

8.5.2. Emissions from the Decomposition of Petroleum-Derived Surfactants (6.D.2)

a) Source/Sink Category Description

Surfactants are used for various cleaning activities at home and factories in Japan. Petroleum-derived surfactants discharged into wastewater treatment facilities and into the environment, and emit CO₂. As this emission source did not correspond to any of the existing waste categories (6.A. to 6.C.), it was included in the “Other (6.D.)” section. Because “CH₄ and N₂O emissions from wastewater treatment” and “CO₂ emissions from the decomposition of petroleum-derived surfactants” concern different types of gas, they are unrelated to each other and pose no duplicate inventory issues.

b) Methodological Issues

● Estimation Method

As neither the *Revised 1996 IPCC Guidelines* nor the *GPG (2000)* specified a method for determining CO₂ emissions, a method specifically established in Japan was applied to the calculation. Because carbon contained in surfactants that emitted into wastewater treatment facilities and into the environment is eventually oxidized to CO₂ and emitted into the atmosphere as a result of surfactants decomposition, CO₂ emissions were estimated based on the amount of carbon contained in surfactants that emitted into wastewater treatment facilities and into the environment.

Based on the facts stated above, the CO₂ emissions were calculated by multiplying the volume of the petroleum-derived surfactant for each type of raw material by the carbon content of each of the materials. The calculation covered synthetic alcohols, alkylbenzenes, alkylphenols, and ethylene oxide. Some of the carbon contained in surfactants discharged into wastewater treatment facilities are adsorbed and assimilated by sludge. However, this portion of carbon is not decomposed biologically. It is released into the atmosphere as CO₂ through incineration and landfilling of sludge. Therefore, the emission is included in CO₂ emission estimates.

● Emission factor

Emission factor was determined for each type of material by calculating the amount of CO₂, expressed in kg that was emitted from the decomposition of 1 t of a surfactant using the average carbon content in the molecules.

$$EF_i = C_i \times 1,000 \times 44/12$$

EF_i : Emission factor of petroleum-derived raw material i used in a surfactant

C_i : Average carbon content of petroleum-derived raw material i used in a surfactant

Table 8 - 73 Average carbon content of surfactants, by petroleum-derived raw material

Raw material	Carbon number	Molecular weight	Carbon content	Basis for determination
<i>Synthetic alcohol</i>	12	186	77.4%	C12-alcohol as the main constituent.
<i>Alkylbenzene</i>	18	250	86.4%	C12-alkylbenzene as the main constituent.
Alkylphenol	15	210	85.7%	C9-alkylphenol as the main constituent.
Ethylene oxide	2	44	54.5%	Based on ethylene oxide molecules (C ₂ H ₄ O)

● Activity Data

Activity data is the amount of raw materials consumed for petroleum-derived surfactants. As some of the surfactants produced in Japan are exported, the activity data were determined by multiplying the volume of raw materials used in the surfactants obtained from the statistical data for surfactant use by an import/export adjustment factor.

Volume of surfactants used

The volumes of the use of surfactant by material were obtained from the consumption of raw materials for surfactants indicated in the *Chemical Industry Statistical Yearbook*. As there was no compilation of usage since FY 2002, the volume of use was estimated using the simple averages (k value) of ratio of

consumption and production in the period from FY 1990 to FY 2001.

Export/import correction factor

Correction factor was calculated from the export/import statistics in *International Trade Statistics* by the Customs Bureau of the Ministry of Finance for categories of anionic surfactants, cationic surfactants, non-ionic surfactants, and other organic surfactants and the volume of surfactants used. As some of the materials for surfactants were used in several types of surfactants, an average of the export/import correction factor was weighted by surfactant production volume as necessary to calculate the correction factor for each classification of surfactant.

Export/Import correction factor

$$= (\text{Surfactant production} + \text{Surfactants imported} - \text{surfactants exported}) / \text{surfactant production}$$

Table 8 - 74 Activity data associated with decomposition of petroleum-based surfactants

Item	Unit	1990	1995	2000	2005	2008	2009	2010
Synthetic alcohol	t	29,239	16,253	28,285	31,609	32,988	32,872	33,750
Alkyl benzene	t	105,432	102,794	80,832	47,349	55,442	50,206	51,005
Alkyl phenol	t	10,141	8,798	7,454	3,448	2,338	2,044	2,054
Ethylene oxide	t	124,984	132,175	146,509	127,150	125,628	126,301	131,158

c) Uncertainties and Time-series Consistency

● ***Uncertainty***

The level of uncertainty associated with emission factor was evaluated by using the differences in carbon content in the major constituents of raw materials for surfactants and was found to be 19% (calculated by using standard deviation). With respect to uncertainties in activity data, twice of the statistical uncertainties set out for the statistics (Survey of total population (rounding) and Other statistics) was used and evaluated to be 40%.

● ***Time-series consistency***

Consistent methodology was used in the estimation. However, data on the amount of raw materials consumed for surfactants have become not available since FY 2002 and activity data were estimated from the production amount of the surfactants.

d) Source-specific QA/QC and Verification

Tier 1 QC activities are implemented in accordance with the *GPG (2000)*. The Tier 1 QC activities include the verification of parameters such as activity data and emission factors, and the archive of reference materials. For more details of QA/QC activities, see the Annex 6.

e) Source-specific Recalculations

Updating activity data of surfactants, emission estimates for FY2009 were recalculated.

f) Source-specific Planned Improvements

No improvements are planned.

References

1. IPCC, *Revised 1996 IPCC Guidelines for National Greenhouse Gas Inventories*, 1997
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Chapter 9. Other (CRF sector 7)

9.1. Overview of Sector

UNFCCC Reporting Guidelines (FCCC/SBSTA/2006/9) para.29 indicates that Annex I Parties should report and explicitly describe the details of emissions from each country-specific source of gases which are not included in the IPCC Guidelines. According to this requirement, emissions from the Other category (CRF sector7) are indicated below.

9.2. CO₂, CH₄, N₂O, HFCs, PFCs, and SF₆

The national inventory submitted this year does not include the emissions and removals of gases targeted under the Kyoto Protocol (CO₂, CH₄, N₂O, HFCs, PFCs, SF₆) from sources and sinks which are not included in the IPCC Guidelines.

9.3. NO_x, CO, NMVOC, and SO₂

The inventory submitted this year includes CO emissions from smoking as emissions of indirect greenhouse gases (NO_x, CO, NMVOC) and SO₂ from sources which are not included in the IPCC Guidelines.

Chapter 10. Recalculation and Improvements

10.1. Explanation and Justification for Recalculations

This section explains improvements on estimation of emissions and removals in the inventory submitted in 2012.

In accordance with the *Good Practice and Uncertainty Management in National Greenhouse Gas Inventories (2000)* and the *IPCC Good Practice Guidance for Land Use, Land-Use Change and Forestry*, recalculations of previously reported emissions and removals are recommended in the cases of 1) application of new estimation methods, 2) addition of new categories for emissions and removals and 3) data refinement. Major changes in the inventory submitted last year are indicated below.

10.1.1. General Issues

In general, activity data for the latest year available at the time when the inventory is compiled are often revised in the year following the submission year because of such as the publication of data in the fiscal year basis. In the national inventory submitted this year, activity data in many sources for 2009 have been changed and as a result, the emissions from those sources for the inventory year have been recalculated.

10.1.2. Recalculations in Each Sector

The information of recalculation for sectors (energy; industrial processes; solvent and other product use; agriculture; land use, land-use change and forestry; and waste) is described separately at sections named as “Source/Sink-specific Recalculations” in Chapters 3 to 8. Information on recalculations for KP-LULUCF activities are described in section 11.4.1.4 of Chapter 11.

10.2. Implications for Emission Levels

Table 10-1 shows the changes made to the overall emission estimates due to the recalculations indicated in “Section 10.1. Explanation and Justification for Recalculations”.

10.2.1. GHG Inventory

Compared to the values reported in the previous year’s inventory, total emissions excluding LULUCF sector in the base year (1990) under the UNFCCC increased by 0.01%, and the total emissions in year 2009 decreased by 0.15% compared to the data reported in last year (Table 10-1).

Table 10-1 Comparison of emissions and removals in the inventories submitted in 2011 and 2012

		[Mt-CO ₂ eq.]																			
		1990	1991	1992	1993	1994	1995	1996	1997	1998	1999	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009
CO ₂ with LULUCF	JNGI2011	1071.5	1073.2	1082.0	1071.4	1129.5	1142.1	1150.3	1144.9	1109.5	1144.2	1164.2	1149.0	1184.9	1180.9	1180.7	1192.0	1178.5	1212.5	1134.9	1073.0
	JNGI2012	1071.0	1072.7	1081.4	1070.9	1129.0	1141.6	1149.9	1144.5	1109.1	1143.8	1163.8	1148.5	1184.4	1180.4	1180.3	1191.5	1178.1	1212.2	1134.5	1070.4
	<i>difference</i>	<i>-0.05%</i>	<i>-0.05%</i>	<i>-0.05%</i>	<i>-0.05%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.04%</i>	<i>-0.03%</i>	<i>-0.04%</i>	<i>-0.25%</i>
CO ₂ without LULUCF	JNGI2011	1141.2	1150.1	1158.6	1150.9	1210.7	1223.7	1236.6	1231.5	1195.9	1230.9	1251.6	1236.4	1273.5	1278.6	1278.0	1282.3	1263.1	1296.3	1213.3	1144.6
	JNGI2012	1141.2	1150.1	1158.6	1150.9	1210.7	1223.7	1236.6	1231.5	1195.9	1230.9	1251.6	1236.4	1273.5	1278.6	1278.0	1282.3	1263.1	1296.3	1213.2	1142.3
	<i>difference</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>-0.20%</i>
CH ₄ with LULUCF	JNGI2011	31.9	31.7	31.4	31.2	30.5	29.6	28.9	27.8	27.0	26.4	25.8	25.0	24.1	23.5	23.1	22.7	22.3	21.8	21.2	20.7
	JNGI2012	32.0	31.8	31.5	31.3	30.6	29.7	29.0	27.9	27.1	26.5	25.9	25.1	24.2	23.7	23.2	22.9	22.5	22.1	21.5	20.9
	<i>difference</i>	<i>0.40%</i>	<i>0.39%</i>	<i>0.39%</i>	<i>0.39%</i>	<i>0.43%</i>	<i>0.41%</i>	<i>0.39%</i>	<i>0.37%</i>	<i>0.38%</i>	<i>0.39%</i>	<i>0.40%</i>	<i>0.49%</i>	<i>0.59%</i>	<i>0.66%</i>	<i>0.72%</i>	<i>0.79%</i>	<i>1.00%</i>	<i>1.35%</i>	<i>1.45%</i>	<i>0.83%</i>
CH ₄ without LULUCF	JNGI2011	31.9	31.7	31.4	31.1	30.5	29.6	28.9	27.8	27.0	26.4	25.8	25.0	24.0	23.5	23.1	22.7	22.3	21.8	21.2	20.7
	JNGI2012	32.0	31.8	31.5	31.2	30.6	29.7	29.0	27.9	27.1	26.5	25.9	25.1	24.2	23.7	23.2	22.9	22.5	22.1	21.5	20.9
	<i>difference</i>	<i>0.40%</i>	<i>0.39%</i>	<i>0.39%</i>	<i>0.39%</i>	<i>0.43%</i>	<i>0.41%</i>	<i>0.39%</i>	<i>0.37%</i>	<i>0.38%</i>	<i>0.39%</i>	<i>0.40%</i>	<i>0.49%</i>	<i>0.59%</i>	<i>0.66%</i>	<i>0.72%</i>	<i>0.79%</i>	<i>1.00%</i>	<i>1.35%</i>	<i>1.45%</i>	<i>0.83%</i>
N ₂ O with LULUCF	JNGI2011	31.7	31.2	31.1	31.1	32.3	32.7	33.7	34.4	32.8	26.4	29.0	25.5	24.8	24.5	24.5	24.0	24.0	22.7	22.5	22.1
	JNGI2012	31.7	31.2	31.4	31.1	32.3	32.7	33.7	34.4	32.8	26.4	29.0	25.6	24.8	24.5	24.6	24.1	24.1	22.8	22.8	22.6
	<i>difference</i>	<i>0.11%</i>	<i>0.08%</i>	<i>0.08%</i>	<i>0.07%</i>	<i>0.08%</i>	<i>0.00%</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.05%</i>	<i>0.06%</i>	<i>0.06%</i>	<i>0.08%</i>	<i>0.12%</i>	<i>0.13%</i>	<i>0.18%</i>	<i>0.23%</i>	<i>0.49%</i>	<i>1.67%</i>	<i>1.95%</i>
N ₂ O without LULUCF	JNGI2011	31.6	31.1	31.3	31.0	32.2	32.7	33.7	34.3	32.8	26.4	28.9	25.5	24.8	24.5	24.5	24.0	24.0	22.7	22.4	22.1
	JNGI2012	31.6	31.1	31.3	31.0	32.2	32.7	33.6	34.3	32.8	26.4	29.0	25.5	24.8	24.5	24.5	24.1	24.1	22.8	22.8	22.6
	<i>difference</i>	<i>0.11%</i>	<i>0.08%</i>	<i>0.08%</i>	<i>0.07%</i>	<i>0.08%</i>	<i>0.00%</i>	<i>-0.03%</i>	<i>-0.05%</i>	<i>-0.03%</i>	<i>-0.05%</i>	<i>0.06%</i>	<i>0.06%</i>	<i>0.08%</i>	<i>0.12%</i>	<i>0.13%</i>	<i>0.18%</i>	<i>0.23%</i>	<i>0.49%</i>	<i>1.67%</i>	<i>1.95%</i>
HFCs	JNGI2011	NE	NE	NE	NE	NE	20.3	19.9	19.9	19.4	19.9	18.8	16.2	13.7	13.8	10.6	10.6	11.7	13.3	15.3	16.7
	JNGI2012	NE	NE	NE	NE	NE	20.3	19.9	19.9	19.4	19.9	18.8	16.2	13.7	13.8	10.6	10.5	11.7	13.3	15.3	16.6
	<i>difference</i>	<i>NA</i>	<i>NA</i>	<i>NA</i>	<i>NA</i>	<i>NA</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>-0.46%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>-0.71%</i>
PFCs	JNGI2011	NE	NE	NE	NE	NE	14.2	14.8	16.2	13.4	10.4	9.5	7.9	7.4	7.2	7.5	7.0	7.3	6.4	4.6	3.3
	JNGI2012	NE	NE	NE	NE	NE	14.2	14.8	16.2	13.4	10.4	9.5	7.9	7.4	7.2	7.5	7.0	7.3	6.4	4.6	3.3
	<i>difference</i>	<i>NA</i>	<i>NA</i>	<i>NA</i>	<i>NA</i>	<i>NA</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.04%</i>	<i>-0.11%</i>
SF ₆	JNGI2011	NE	NE	NE	NE	NE	17.0	17.5	15.0	13.6	9.3	7.2	6.0	5.6	5.3	5.1	4.8	4.9	4.4	3.8	1.9
	JNGI2012	NE	NE	NE	NE	NE	17.0	17.5	15.0	13.6	9.3	7.2	6.0	5.6	5.3	5.1	4.8	4.9	4.4	3.8	1.9
	<i>difference</i>	<i>NA</i>	<i>NA</i>	<i>NA</i>	<i>NA</i>	<i>NA</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>	<i>0.00%</i>
Total with LULUCF	JNGI2011	1135.1	1136.1	1144.7	1133.7	1192.2	1255.9	1265.1	1288.2	1215.8	1236.7	1254.5	1229.5	1260.4	1255.1	1251.5	1261.1	1248.8	1281.1	1202.3	1137.7
	JNGI2012	1134.8	1135.7	1144.3	1133.3	1191.9	1255.6	1264.8	1287.8	1215.5	1236.3	1254.2	1229.2	1260.1	1254.8	1251.2	1260.8	1248.6	1281.1	1202.6	1135.5
	<i>difference</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.03%</i>	<i>-0.02%</i>	<i>-0.02%</i>	<i>-0.02%</i>	<i>-0.02%</i>	<i>-0.01%</i>	<i>-0.01%</i>	<i>0.00%</i>	<i>0.02%</i>	<i>-0.19%</i>
Total without LULUCF	JNGI2011	1204.7	1212.9	1221.2	1213.1	1273.3	1337.4	1351.3	1344.7	1302.2	1323.3	1341.8	1317.0	1349.0	1352.8	1348.7	1351.3	1333.3	1364.9	1280.6	1209.2
	JNGI2012	1204.9	1213.0	1221.4	1213.2	1273.5	1337.5	1351.4	1344.8	1302.3	1323.4	1341.9	1317.1	1349.1	1353.0	1348.9	1351.5	1333.6	1365.3	1281.3	1207.4
	<i>difference</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.01%</i>	<i>0.02%</i>	<i>0.03%</i>	<i>0.05%</i>	<i>-0.15%</i>

10.2.2. KP-LULUCF Inventory

Compared to the values reported in the previous year's inventory, total emissions/removals arising from KP-LULUCF activities in 2008 and 2009 increased by 0.74% and 0.70%, respectively (Table 10-2).

Table 10-2 Comparison of emissions and removals in the inventories submitted in 2011 and 2012 for KP-LULUCF activities

		[Gg-CO ₂ eq.]			
Activity	Gas		1990	2008	2009
Afforestation and Reforestation	CO ₂	JNGI2011	-	-389.6	-415.1
		JNGI2012	-	-389.6	-415.0
		<i>difference</i>	-	<i>-0.01%</i>	<i>-0.01%</i>
CH ₄	JNGI2011	-	0.0	0.0	
	JNGI2012	-	0.0	0.0	
	<i>difference</i>	-	<i>0.38%</i>	<i>-1.29%</i>	
N ₂ O	JNGI2011	-	0.0	0.0	
	JNGI2012	-	0.0	0.0	
	<i>difference</i>	-	<i>0.38%</i>	<i>-1.29%</i>	
Deforestation	CO ₂	JNGI2011	-	2,427.0	3,083.4
		JNGI2012	-	2,454.3	3,112.0
		<i>difference</i>	-	<i>1.13%</i>	<i>0.93%</i>
N ₂ O	JNGI2011	-	2.37	3.12	
	JNGI2012	-	2.37	3.12	
	<i>difference</i>	-	<i>0.00%</i>	<i>0.00%</i>	
Forest Management	CO ₂	JNGI2011	-	-45,402.8	-49,011.4
		JNGI2012	-	-45,402.8	-49,011.4
		<i>difference</i>	-	<i>0.00%</i>	<i>0.00%</i>
CH ₄	JNGI2011	-	12.7	5.1	
	JNGI2012	-	12.8	5.3	
	<i>difference</i>	-	<i>0.38%</i>	<i>3.36%</i>	
N ₂ O	JNGI2011	-	1.3	0.5	
	JNGI2012	-	1.3	0.5	
	<i>difference</i>	-	<i>0.38%</i>	<i>3.36%</i>	
Revegetation	CO ₂	JNGI2011	-47.1	-729.7	-754.8
		JNGI2012	-77.8	-1,081.8	-1,112.3
		<i>difference</i>	<i>65.27%</i>	<i>48.25%</i>	<i>47.36%</i>
Total	JNGI2011	-47.1	-44,078.8	-47,089.2	
	JNGI2012	-77.8	-44,403.3	-47,417.8	
	<i>difference</i>	<i>65.27%</i>	<i>0.74%</i>	<i>0.70%</i>	

10.3. Implication for Emission Trends, including Time Series Consistency

Table 10-3 shows the changes made to the emission trends due to the recalculations indicated in “Section 10.1. Explanation and Justification for Recalculations”. The comparison between the 2011 submission (JNGI 2011) and the 2012 submission (JNGI 2012) is made through the comparison of values between the base year and FY2009.

The actual emissions of HFCs, PFCs, and SF₆ prior to CY1995 are not reported; hence, the comparison between 2010 and 2011 of these emissions applies the comparison values between CY1995 and CY2009.

10.3.1. GHG Inventory

Total emissions excluding the LULUCF sector in the 2012 submission decreased by approximately 2.0 million tons (in CO₂ equivalents) and increased by 0.04 percentage points, compared to the data reported in the previous submission.

Table 10-3 Comparison of increase and decrease from the base year, between the inventories submitted in 2011 and 2012 excluding LULUCF sector

		Trend [Mt-CO ₂ eq.]			Trend (%)		
		JNGI2011	JNGI2012	Difference	JNGI2011	JNGI2012	Difference
CO ₂	1)	3.4	1.1	-2.3	0.3%	0.1%	-0.2%
CH ₄	1)	-11.2	-11.1	0.0	-35.1%	-34.8%	0.3%
N ₂ O	1)	-9.5	-9.1	0.4	-30.0%	-28.7%	1.3%
HFCs	2)	-3.6	-3.7	-0.1	-17.7%	-18.3%	-0.6%
PFCs	2)	-11.0	-11.0	0.0	-77.0%	-77.1%	0.0%
SF ₆	2)	-15.1	-15.1	0.0	-89.1%	-89.1%	0.0%
Total	3)	-47.0	-49.0	-2.0	1.9%	2.0%	0.04%

1) Comparison of emissions between FY1990 and FY2009

2) Comparison of emissions between CY1995 and CY2009

3) Comparison of emissions between the base year of the Kyoto Protocol (CO₂, CH₄, N₂O: FY1990, HFCs, PFCs, SF₆: CY1995) and 2009

10.4. Recalculations and improvement plan, including response to the review process

10.4.1. Improvements after submission of inventory

The major improvements carried out since submission of the 2011 inventory are listed below.

10.4.1.1. Methodology for estimating emissions and removals of GHGs

Changed calculation methods are as follows. See each category for details.

10.4.1.1.a. GHG Inventory

1. For “1.A.3.b. Road Transportation”, the recalculation were implemented in response to the updates of the emission factors used for the estimation of the CH₄ and N₂O emissions from road transportation.

2. For “2.A.3. Limestone and Dolomite Use,” and “2.A.4. Soda Ash Production and Use, the activity data were re-examined and the emissions have been recalculated.
3. For “2.F.1. Refrigeration and Air Conditioning Equipment,” the activity data were updated etc and the emissions have been recalculated.
4. For “4.B. Manure Management”, new ‘proportion of separated and mixed treatment of manure, by type of livestock’ and ‘percentage of manure management by type of animal’ were reported. Therefore, emissions for this category were revised.
5. For “4.D.1. Crop Residue”, the revision of nitrogen content in crop residue for some crops, the amount of nitrogen put into soils as crop residues were revised. Therefore, the emissions were revised.
6. For “5.A. Forest land”, the implied emission/removal factor for the afforestation, reforestation and deforestation, which was used to separate the carbon stock changes in living biomass in Forest land into the “5.A.1 Forest land remaining Forest land” and the “5.A.2 Land converted to Forest land”, was re-investigated. As a result, the implied factor was found not to concern the losses of 5.A.2. Therefore, the losses of 5.A.2, which used to be reported as “IE” in the previous submission, were newly accounted.
7. For “5.E.1 Settlements remaining Settlements”, the recalculations were implemented in response to the updates of the parameters used for the estimation of the carbon stock changes in living biomass and litter in the urban green facilities.
8. For “5.E.1 Settlements remaining Settlements”, the carbon stock changes in soils, which used to be reported as “NE” for the urban green facilities until the previous submission, were newly estimated, since the new knowledge became available.
9. For “5(V) Biomass burning”, the emissions were recalculated in response to the corrections of parameters used for estimating the damaged timber volume.
10. For “6.A.1. Solid Waste Disposal on Land”, determining the amount of ISW disposed of in semi-aerobic landfill sites, and CH₄ emissions from ISW landfill sites were recalculated.
11. For “6.D.1. Emissions from Composting of Organic Waste”, CH₄ and N₂O emission estimates from composted animal and plant residues generated by food and beverage manufacturing and some other food waste were newly accounted

10.4.1.1.b. KP-LULUCF Inventory

1. For the RV activity, the carbon stock changes in living biomass and litter were recalculated, as the parameters were updated.
2. For the RV activity, the carbon stock changes in soils, which used to be reported as “NE”, were recalculated, since new knowledge became available.
3. For the RV activity, the area was recalculated, since the way to obtain the subject areas were re-examined.

10.4.1.2. National Greenhouse Gas Inventory Report

The report has been modified to follow the structure in the Annotated outline of the National Inventory Report including reporting elements under the Kyoto Protocol, the application of which is recommended by the UNFCCC secretariat. This results in reporting supplementary information under Article 7.1 of the Kyoto Protocol in Chapters 11 (Annex 11 in the 2011 submission), 12 through 15 (Annex 10 in the 2011 submission).

10.4.1.3. Improvements by following UNFCCC-ERT recommendations

Actions taken in response to UNFCCC-ERT recommendations are summarized below. See relative sections for details.

Table 10-2 Improvements to the NIR and the CRF in response to UNFCCC review

Sector/Category	recommendations by ERT	Taken Actions	NIR/CRF
General	The provision in future NIRs of an overview of the main drivers of emission trends in order to enhance the transparency of the emission estimates (paragraph 29(a) in ARR 2009 and so on)	Main drivers of emission trends are provided in Chapter 2 of the NIR by sector and by gas.	Chapter 2 of the NIR
Energy/ Fuel combustion (1.A)	Japan has provided a detailed discussion and analysis of discrepancies between the figures reported in the CRF tables and the IEA statistics in annex 2 of the NIR, but the analysis examines data for the year 2005. The ERT encourages Japan to update this initiative for the latest inventory year in future annual submissions. (paragraph 36 in ARR 2010)	The detailed information regarding the discrepancies of the reported value between the CRF and IEA statistics is updated with the FY2009 actual data in the NIR Annex 2..	NIR Annex 2 p. Annex2-1~2-9
Agriculture/ Enteric Fermentation (4.A)	Japan revises and improves the explanations about animal characterization, in particular tables 6.2 and 6.9 of its NIR. (paragraph 70 in ARR draft 2011)	Information in table 6-2 (Categorization and assumption underlying calculation for cattle) and 6-9 (Livestock population for cattle) of the NIR were added.	Table 6-2 and 6-9 of the NIR
Agriculture/ Manure Management (4.B)	According to Table 6-27, the amount of livestock manure for grazing Buffalo, Sheep, Goats and Horses will be presented in the Pasture, Range and Paddock column of CRF Table 4B(b). (Assignment by the Review in 2011)	The amount of livestock manure for grazing Buffalo, Sheep, Goats and Horses were presented in the Pasture, Range and Paddock column of CRF Table 4B(b).	CRF Table4.B(b)
Agriculture/ Prescribed Burning of Savanna (4.E)	Emissions from this activity are reported, in some years (1990 - 2003), as 'NE' but the use of 'NE' is not in line with the UNFCCC reporting guidelines, because Japan does not have this emission source; instead, these emissions should always be reported as 'NO'. (paragraph 62 in ARR draft 2011)	Emissions from prescribed burning of savannas in all years are reported as 'NO'.	CRF Table4. E
KP-LULUCF	The gains for below-ground living biomass for deforestation are reported as "NO" while both gains and losses are reported for above-ground living biomass.	The gains for below-ground living biomass were included in the gains for above-ground biomass; therefore, the appropriate notation key "IE" should have been used. In this submission, the gains for above- and below-ground biomass were reported separately.	KP-CRF, 5(KP-I)A.2

10.4.2. Planned Improvements

The main planned improvements are as follows.

1. Review of estimation methods, activity data, emission factors and other elements

Japan will hold meetings of a Committee for Greenhouse Gas Emission Estimation Methods and will consider improvements of estimation methods, activity data, emission factors and other elements used in the current inventory. In case of implementation, Japan will prioritize highly important issues such as those relevant to key-categories and those pointed out in the past review reports.

2. Improvement of transparency

Japan will further improve transparency of the inventory by examining descriptions of methodologies, assumptions, data, and other elements in NIR, and by adding necessary information to NIR.

Chapter 11. Supplementary Information on LULUCF activities under Article 3, Paragraphs 3 and 4 of the Kyoto Protocol

11.1. Summary of removal related trends, and emissions and removals from KP LULUCF activities

Japan reports supplementary information on Afforestation/Reforestation (AR), Deforestation (D), Forest management (FM) and Revegetation (RV) as LULUCF activities under Article 3, Paragraphs 3 and 4 of the Kyoto Protocol. Table 11-1 shows the activity coverage and other information relating to activities under Article 3.3 and elected activities under Article 3.4. The net removals in FY2010 by those activities were 49,985 Gg-CO₂ eq. (Table 11-2).

Table 11-1 Activity coverage and other information relating to activities under Article 3.3 and elected activities under Article 3.4 (CRF-Table NIR 1)

Activity	Change in carbon pool reported ⁽¹⁾					Greenhouse gas sources reported ⁽²⁾							
	Above-ground biomass	Below-ground biomass	Litter	Dead wood	Soil	Fertilization ⁽³⁾	Drainage of soils under forest management	Disturbance associated with land-use conversion to croplands	Liming	Biomass burning ⁽⁴⁾			
						N ₂ O	N ₂ O	N ₂ O	CO ₂	CO ₂	CH ₄	N ₂ O	
Article 3.3 activities													
Afforestation and Reforestation	R	R	R	R	R	IE			NO	IE	R	R	
Deforestation	R	R	R	R	R			R	R	NO	NO	NO	NO
Article 3.4 activities													
Forest Management	R	R	R	R	R	IE	NO		NO	IE	R	R	
Cropland Management	NA	NA	NA	NA	NA			NA	NA	NA	NA	NA	NA
Grazing Land Management	NA	NA	NA	NA	NA				NA	NA	NA	NA	NA
Revegetation	R	R	R	IE	R				R	NO	NO	NO	NO

*R: Reported. See Annex 5 for the definitions of the other notation leys.

Table 11-2 Accounting summary for activities under Articles 3.3 and 3.4 of the Kyoto Protocol (CRF Information Table)

GREENHOUSE GAS SOURCE AND SINK ACTIVITIES	BY	Net emissions/removals				Accounting Parameters	Accounting Quantity
		2008	2009	2010	Total		
		(Gg CO ₂ equivalent)					
A. Article 3.3 activities							
A.1. Afforestation and Reforestation						-1,230.68	
A.1.1. Units of land not harvested since the beginning of the commitment period		-389.54	-415.03	-426.11	-1,230.68	-1,230.68	
A.1.2. Units of land harvested since the beginning of the commitment period							
A.2. Deforestation		2,456.72	3,115.09	4,822.89	10,394.70	10,394.70	
B. Article 3.4 activities							
B.1. Forest Management (if elected)		-45,388.77	-49,005.55	-53,251.78	-147,646.10	-147,646.10	
3.3 offset						9,164.02	
FM cap						238,333.33	
						-138,482.08	
B.2. Cropland Management (if elected)		NA	NA	NA	NA	NA	
B.3. Grazing Land Management (if elected)		NA	NA	NA	NA	NA	
B.4. Revegetation (if elected)		-77.78	-1,081.76	-1,112.34	-1,130.14	-3,324.24	
						-233.34	
						-3,090.90	

- ※ The net removals by FM after application of 3.3 offset are lower than the upper limit (13 Mt-C times 5 (238,333 Gg-CO₂)) given in the Appendix to decision 16/CMP.1.
- ※ Since the total anthropogenic GHG emissions by sources and removals by sinks in managed forests since 1990 are larger than the net source of emissions incurred under Article 3.3, the offset rule according to paragraph 10 of the Annex to decision 16/CMP.1 is applied to Japan.
- ※ Methodologies for estimation and accounting of Article 3.3 and 3.4 activities are continuously reviewed. The values in Table 11-2 are estimated by using the current methodologies, and are only reported but not accounted for in the 2012 submission since Japan elected accounting for the entire commitment period. The issuance of removal units from LULUCF activities under the Kyoto Protocol is to be performed at the end of the first commitment period.
- ※ The total values and results of summing up each figure are not always the same because of the difference in display digit.

11.2. General information

11.2.1. Definition of forest and any other criteria

Japan's definitions of forest are identified as the following, in accordance with decision 16/CMP.1 and the requirement from GPG-LULUCF.

- Minimum value for forest area: 0.3 [ha]
- Minimum value for tree crown cover: 30 [%]
- Minimum value for tree height: 5 [m]
- Minimum value for forest width: 20 [m]

Forests with minimum values for forest area, tree crown cover and forest width (mentioned above) are consistent with forests under the existing forest planning system in Japan. Although any minimum value for tree height is not defined under the existing system, forests with usual composition of tree species and under usual climate conditions in Japan usually reach a tree height of 5 m at maturity *in situ*. Each prefecture has surveyed and compiled information on forest resources under the forest planning system into Forest Registers, which are primarily intended to be prepared for establishing forest plans. Therefore, forests under the forest planning system are considered as forests under the Kyoto Protocol, and Forest Registers are suitable as basic data source for reporting. This is the same concept as the one used for reporting the LULUCF forest sector under the Convention.

The definitions of forest mentioned above are consistent with those in the Global Forest Resources Assessment 2005 (FRA2005) by the Food and Agriculture Organization of the United Nation (FAO) (Table 11-3).

Table 11-3 Japan's forest categories and definition used in reporting to FAO

Category	Definition
Forest	Land on which trees and/or bamboo grow collectively, together with those trees and bamboo, or any other land that is provided for collective growth of trees and/or bamboo which are 0.3 ha or more. Lands that are utilized mainly for agriculture, residential use or other similar purposes, and trees and bamboo on these lands, are not included.
Forest with standing trees	Forest that has a tree crown cover of 30 percent or higher (including young stands).
Forest with less standing trees (Cut-over forest, lesser stocked forest)	Forest that does not fall under "forest with standing trees" or "bamboo forest".
Bamboo forest	Forest that does not fall under "forest with standing trees" and is mainly dominated by bamboo (excluding sasa).

※ See section 7.2.2. for a more detailed definition of each category

Before 1995, Japan classified forests with standing trees into two sub-categories, "Intensively managed forests" and "Semi-natural forests" in the Forestry Status Survey. Since 2002, Japan has introduced new sub-categories which are "Ikusei-rin forest" and "Tennensei-rin forest". In these new sub-categories, the degree of human-induced activities and stratification of forest have been taken into account. In ikusei-rin forests, intensively managed forests regenerated mainly by planting after felling and semi-natural forests regenerated by supplementary works such as site preparation are included. The definitions of intensively managed forest, semi-natural forest, ikusei-rin forest and tennensei-rin forest are shown below.

Table 11-4 Definitions of intensively managed forest, semi-natural forest, ikusei-rin forest and tennensei-rin forest

Sub-categories by regeneration method		Sub-categories by management types	
Intensively managed forest	Forest regenerated by planting and so on.	Ikusei-rin forest	Forest where practices for establishment and maintenance of single-storied forests (“Ikusei-tansou-rin” practices) have been carried out after clear-cutting, or where forest practices for establishment and maintenance of multi-storied forests (“Ikusei-fukusou-rin” practices) have been carried out after selective cutting (including temporally single-storied forest in practice).
Semi-natural forest	Forest which is not classified as intensively managed forest.	Tennensei-rin forest	Forest where practices for establishment and maintenance of forest mainly depending on natural power are carried out. These practices include logging prohibition for land, natural environment conservation and preservation of the species.

11.2.2. Elected activities under Article 3, paragraph 4 of the Kyoto Protocol

Japan elected FM and RV defined by decision 16/CMP.1 in paragraph 6 of the Annex, as “additional human-induced activities related to changes in GHG emissions by sources and removals by sinks in the agricultural soils and the land-use change and forestry categories” defined by Article 3, paragraph 4 of the Kyoto Protocol.

11.2.2.1. Forest Management

FM is defined by decision 16/CMP.1 in paragraph 1 (f) of the Annex as “a system of practices for stewardship and use of forest land aimed at fulfilling relevant ecological (including biological diversity), economic and social functions of the forest in a sustainable manner”. Japan interprets the definition of FM as the following by recalling GPG-LULUCF which the party is requested to use in accordance with decision 16/CMP.1, paragraph 2.

- Activities for FM in “Ikusei-rin forests” are appropriate forest practices including regeneration (land preparation, soil scarification, planting, etc.), tending (weeding, pre-commercial cutting, etc.), thinning and harvesting which have been carried out since 1990.
- Activities for FM in “Tennensei-rin forests” are practices for protection or conservation of forests including controlling logging activities and land-use change which have been carried out by law.

11.2.2.2. Revegetation

RV is defined by decision 16/CMP.1 ANNEX paragraph 1 (e) as “a direct human-induced activity to increase carbon stocks on sites through the establishment of vegetation that covers a minimum area of 0.05 ha and does not meet the definitions of AR”. Japan interprets the definition of RV as the following by recalling GPG-LULUCF.

- Practices for the creation of “parks and green space”, “public green space”, and “private green space guaranteed by administration” which have been carried out in settlements since 1990¹. Activities which cover less than an area of 0.05 ha or meet the definitions of AR are not included.

¹ In Japan, the urban green facilities subject to RV activities are: “Urban parks”, “Green areas on roads”, “Green areas at ports”, “Green areas around sewage treatment facilities”, “Green areas by greenery promoting systems for private green space”, “Green areas along rivers and erosion control sites”, “Green areas around government buildings”, and “Green areas around public rental housing”.

11.2.3. Description of how the definitions of each activity under Article 3.3 and each elected activity under Article 3.4 have been implemented and applied consistently over time

The forest definition explained in section 11.2.1. has not changed over time. The same forest definition is used for AR and D under Article 3.3 as well as FM under Article 3.4. The definitions of FM and RV explained in section 11.2.2. above have been implemented and applied consistently over time.

11.2.4. Description of precedence conditions and/or hierarchy among elected Article 3.4 activities, and how they have been consistently applied in determining how land was classified

Japan interprets that FM activities occur only in forest land and RV activities only in settlements. Therefore, there is no overlapping between FM and RV.

11.3. Land-related information

11.3.1. Spatial assessment unit used for determining the area of the units of land under Article 3.3

In accordance with the definition of forest explained in section 11.2.2. , Japan determines the spatial assessment unit used for determining the area of the units of land under Article 3.3 as 0.3 ha.

11.3.2. Methodology used to develop the land transition matrix

11.3.2.1. Description of land transition matrix (CRF-NIR Table 2)

Table 11-5 shows the land transition matrix related to the activities under Articles 3.3 and 3.4. The FM area in Japan is estimated by using the narrow approach concept described in section 4.2.7.1, Chapter 4 of the GPG-LULUCF. Therefore, new FM areas are identified every year due to the progress of FM practices in managed forests which previously had not been categorized as FM area. These areas appear as land transition from “Other” to FM in Table 11-5. In a similar fashion, sites where RV practices have been newly performed become new RV areas and appear as land transition from “Other” to RV in Table 11-5.

While there are some cases where the activity categories of land before transition cannot be separated at the moment (e.g. D in FM land and D in non-FM land), transition from “Other” to certain activities is temporarily used for such cases in this table.

Table 11-5 Land Transition Matrix of Kyoto Protocol Activities (CRF-Table NIR 2)

TO 2010		Article 3.3 activities		Article 3.4 activities			Other	Total
		Afforestation and reforestation	Deforestation	Forest Management (if elected)	Cropland Management (if elected)	Grazing Land Management (if elected)		
FROM 2009		(kha)						
Article 3.3 activities	Afforestation and Reforestation	28.26	0.00					28.26
	Deforestation		309.44					309.44
Article 3.4 activities	Forest Management (if elected)		IE	14,314.85				14,314.85
	Cropland Management ⁽⁴⁾ (if elected)	-	-		-	-	-	0.00
	Grazing Land Management ⁽⁴⁾ (if elected)	-	-		-	-	-	0.00
	Revegetation ⁽⁴⁾ (if elected)	0.00			-	-	76.23	76.23
Other		0.49	13.29	795.72	-	-	1.21	22,250.51
Total area		28.75	322.73	15,110.57	0.00	0.00	77.44	22,250.51
								37,790.00

11.3.2.2. Overview of the procedures to estimate emissions and removals

This section gives an overview of the procedures to estimate emissions and removals for AR, D and FM activities in Japan. For AR and D activities, emissions and removals are estimated in AR and D areas which are detected for each prefecture based on sample survey data. For FM activity, emissions and removals are estimated by firstly subtracting emissions and removals in AR land from those in all managed forests for each prefecture, and then applying the FM ratio determined by the sample survey to the remaining emissions and removals.

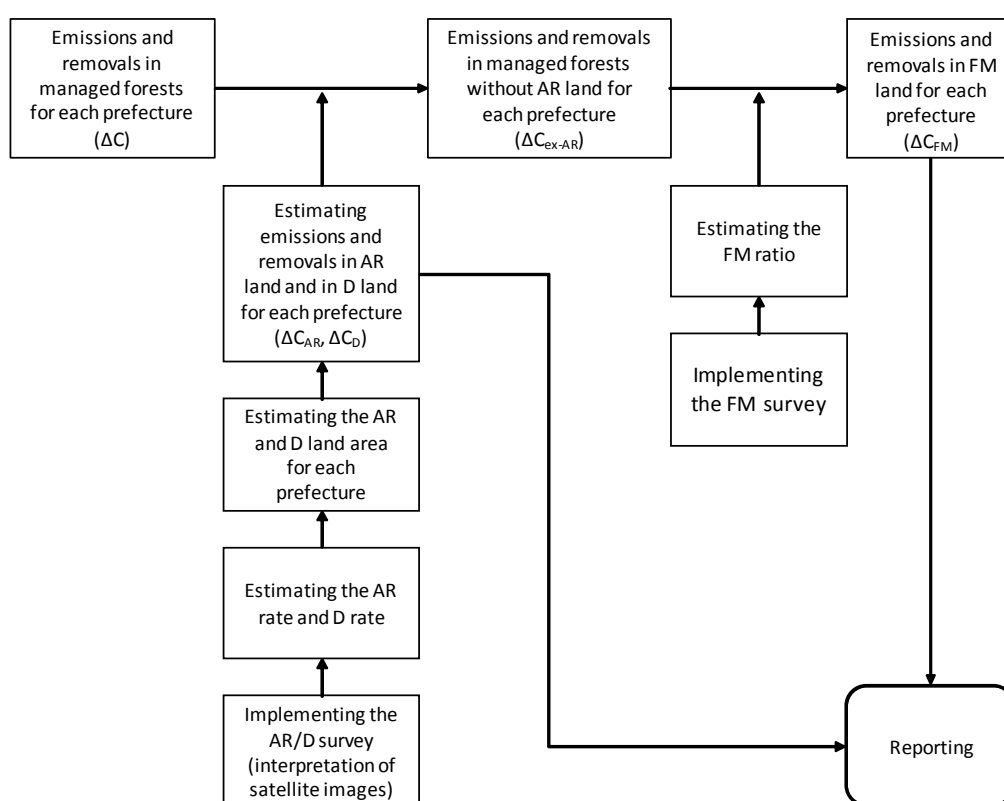


Figure 11-1 Procedures to estimate emissions and removals for AR, D and FM activities

11.3.2.3. Afforestation/Reforestation and Deforestation

11.3.2.3.a. Procedure

Japan identifies the change of forest cover in each sample plot by using orthophotos taken at the end of 1989 and recent satellite images, taking into account the spatial assessment unit (0.3 ha). Plots identified as non-forest land converted to forest land due to human-induced forestation practice are categorized as AR plot, and plots identified as forest land converted to non-forest land are categorized as D plot (Hayashi *et al.*, 2008). Satellite images of the country are updated and interpreted in the subsequent two years (e.g. satellite images taken in 2007 were interpreted in FY2008 and FY2009), and AR and D land areas are calculated based on the results of the interpretation. The detailed procedures are as follows:

1. The plot points on the whole country are set in a grid with an interval of 500 m (approximately 1,500 thousand plots).
2. Land conversion between forest and non-forest is detected at each plot point. Plots which are

difficult to interpret are excluded from “available sample plots” which are used for the following estimation.

3. The AR rate for FY1990-FY2010 is estimated as follows: The number of AR plots for FY1990-FY2007 is calculated by using orthophotos taken at the end of 1989 and satellite images taken in 2005 and 2007. The increase in AR plots for FY2008 is estimated as the average of two-year increase of AR plots obtained from the interpretation of satellite images of 2005 and 2007. The increase in AR plots for FY2009 is estimated as the average of two-year increase of AR plots obtained from the interpretation of satellite images of 2007 and 2009 (the interpretation work for the 2009 estimation covers half of the national land). In a similar way, the increase in AR plots for FY2010 is estimated from the interpretation of satellite images of 2007 and 2009 (the interpretation work for the 2010 estimation covers the other half of the national land). The AR rate for FY1990-FY2010 is estimated through dividing the increase of AR plots in each year by the number of “available sample plots” of each interpretation and then summing them.
4. The D rate for FY1990-FY2010 is estimated as follows: The number of D plots for each fiscal year during FY1990-FY2007 is estimated by multiplying the total number of D plots during FY1990-FY2007, obtained by using orthophotos taken at the end of 1989 and satellite images in 2005 and 2007, by the land-conversion ratio in each fiscal year provided by statistics. The increase in D plots for FY2008 is estimated as the average of two-year increase of D plots obtained from the interpretation of satellite images of 2005 and 2007. The increase in D plots for FY2009 is estimated as the average of two-year increase of D plots obtained from the interpretation of satellite images of 2007 and 2009 (the interpretation work for the 2009 estimation covers half of the national land). In a similar way, the increase in D plots for FY2010 is estimated from the interpretation of satellite images of 2007 and 2009 (the interpretation work for the 2010 estimation covers the other half of the national land). The D rate for FY1990-FY2010 is estimated through dividing the increase of D plots in each year by the number of “available sample plots” of each interpretation and then summing them. The land-use status after D is analyzed at each plot point and these data are used for the estimation of new land-use status in D land.
5. The AR land area during FY1990- FY2010 is calculated by multiplying the land area for each prefecture by the AR rate. In the same way, the D land area for each prefecture during FY1990-FY2010 is calculated by multiplying the land area for each prefecture by the D rate.

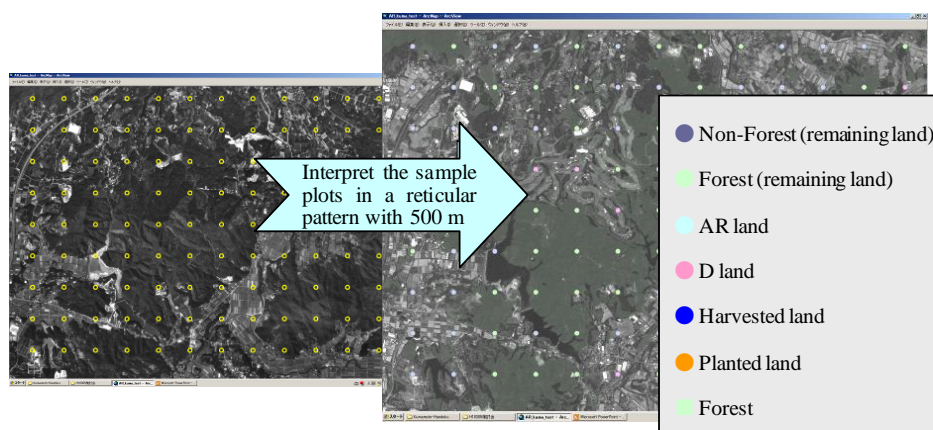


Figure 11-2 ARD land identification by interpreting remote sensing images

Although Forest Registers are used as basic data source for reporting since forests under the forest planning system are considered as forests under the Kyoto Protocol in Japan, orthophotos and satellite images are used for AR and D detection. This is because it is difficult to reconstruct the forest status during FY1990-FY2005 from the data in the Forest Registers, and to distinguish human-induced AR from forest expansion due to other causes.

11.3.2.3.b. Data

Japan determines the ARD land area by using the following data.

Table 11-6 Data used in ARD land detection

	Resolution [m]	Data format
Ortho air-photo (at the end of 1989)	1	Raster
SPOT-5/HRV-P(after 2005, 2007 and 2009)	2.5	Raster

11.3.2.3.c. Land-use change in deforested land

Japan determines the area of D land in accordance with the procedures mentioned in section 11.3.2.3.a. However, these procedures do not cover the continuous tracking of land-use change in D land. Therefore, the land-use change status in the D land has been assessed separately.

Japan has compiled land-use mesh data in the so called “Digital National Land Information” continuously over time. Although this mesh data cannot be used directly to monitor land-use change in the plots identified as D land because this mesh data is not absolutely consistent with the system mentioned in section 11.3.2.3.a (e.g. definition, resolution and land identification method), it can detect the overall tendency of land-use transition in the D plot. The results of the analysis of this mesh data show that D land is seldom converted to other land use again. Therefore, Japan assumes that the status of land use after D will continue to be the same and secondary land-use change will not occur.

11.3.2.4. Forest Management

11.3.2.4.a. Procedure

Japan estimates the FM land area for Ikusei-rin forests and Tennensei-rin forests according to the following procedures.

a) Ikusei-rin forests

1. A field survey in private forests and national forests is implemented each year to identify lands which have been subject to FM activities (the number of sample plots are systematically distributed by tree species and regions; then, sample plots are selected randomly from the National Forest Resource Database (NFRDB)).

Survey items: current status of forests (tree species, stand age, number of trees, etc), status and contents of practices since 1990, etc.

2. The ratio of these FM land areas (FM ratio) is estimated according to the survey findings.
3. After the AR land area for each prefecture is subtracted from the total forest area, the remaining forest area for each prefecture is multiplied by the FM ratio for each tree species, regions and age class.

Table 11-7 FM ratio for Ikusei-rin forests (private forests / national forests)

Sub-category / Tree species		Region	Private forest	National forest
Intensively managed forest	Japanese cedar	Tohoku, Kita-kanto, Hokuriku, Tosan	0.82	0.84
		Minami-kanto, Tokai	0.62	0.80
		Kinki, Chugoku, Shikoku, Kyusyu	0.67	0.81
	Hinoki cypress	Tohoku, Kanto, Chubu	0.78	0.82
		Kinki, Chugoku, Shikoku, Kyusyu	0.75	0.85
	Japanese larch	All	0.76	0.71
Other	All	0.58	0.73	
Semi-natural forest / All		All	0.26	0.59

* Data at the end of FY2010. About 20,000 sample plots are located all across the country.

* These regions generally used broad boundaries which aggregated several prefectures.

* FM ratios shown in this table are area-weighted average values of FM ratio for each age class.

b) *Tennensei-rin forests*

For Tennensei-rin forests, forest lands subject to practices for protection or conservation of forests such as controlling logging activities and land-use change which have been carried out by law are identified by using the NFRDB.

Table 11-8 Area of protected/conserved Tennensei-rin forests

Protected / Conserved forest type	[Unit: kha]		
	Private forest	National forest	Total
Protection Forest	2,726	4,761	7,487
Area for Conservation facility installation project	1	0	1
Protected Forest	0	903	903
Special Protected Zones in National Parks	41	214	255
Class I Special Zones in National Parks	36	165	201
Class II Special Zones in National Parks	120	212	331
Special Protected Zones in Quasi-National Parks	9	56	65
Class I Special Zones in Quasi-National Parks	31	120	152
Class II Special Zones in Quasi-National Parks	98	85	183
Special Zone in National Environment Conservation Area	0	10	10
Special Seed Forest	1	1	1
Total	3,063 (2,628)	6,526 (4,601)	9,590 (7,228)

* NFRDB (1st April 2011)

* This table includes forests with less standing trees.

* () means total land area excluding overlaps.

11.3.2.4.b. Data

a) *Basic data for estimation*

The basic data sources for FM estimation are Forest Registers and yield tables developed by prefectures or Regional Forest Offices. Some of the yield tables are developed by the Forestry and Forest Products Research Institute. These Forest Registers and yield tables are also used for reporting under the convention. Detailed information on Forest Registers and yield tables is provided in section 7.4.1.b).1, Chapter 7 of this report.

b) Development of the National Forest Resources Database

To estimate emissions from or removals by forests, the Forestry Agency has developed the National Forest Resources Database (NFRDB). In the NFRDB, Forest Registers which are the basic data source for estimating and reporting, administrative information including Forest Planning Maps and geographical location information such as orthophotos and satellite images like Landsat-TM and SPOT are archived.

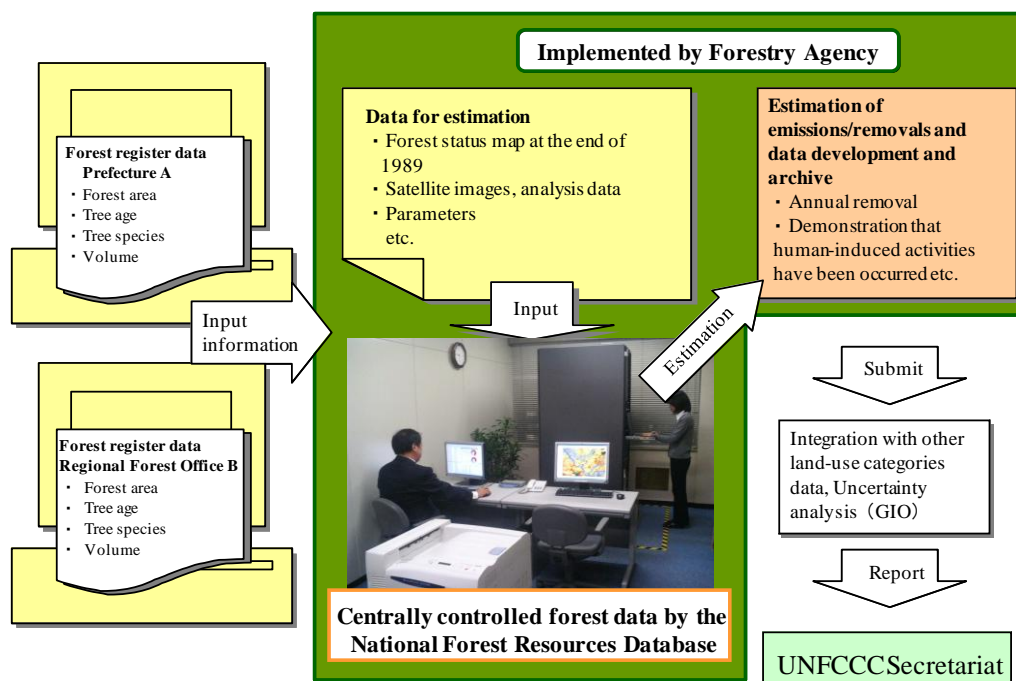


Figure 11-3 Summary of the National Forest Resources Database

11.3.2.5. Revegetation

11.3.2.5.a. Procedure

Japan estimates the RV land area by types of urban green facilities according to the following procedures.

a) Urban parks

1. The information on the notification date and the establishment areas are rearranged as of the end of each corresponding fiscal year during the commitment period for all urban parks which are installed in our country.
2. The urban parks which have been notified since 1st January 1990 and whose establishment area is 500 m² or more are extracted.
3. The urban parks extracted in Step 2 are rearranged after address and the establishment areas determined by geographical boundary (prefecture) are counted.
4. The area of land which was classified as forest land on 31st December 1989 is calculated by multiplying the establishment area estimated in Step 3 by the sum of the area ratios of "Land that has been converted from Forest land to Settlements per annum" since 1990 until each

corresponding fiscal year during the commitment period. This area is excluded from the establishment areas because it is classified as D. The remaining area is considered as RV land area.

5. The areas of “Remaining land (Settlements remaining Settlements)” and “Land converted from other land-use categories (Cropland / Grassland / Wetlands / Other land converted to Settlements)” are calculated by multiplying the land area estimated in Step 4 by the area ratio of “Land converted to Settlements” in the single year².

b) Green areas on roads

1. The number of tall trees at the end of each corresponding fiscal year during the commitment period is calculated for each geographical boundary (prefecture) based on the results of “Road Tree Planting Status Survey”.
2. The number of tall trees on 31st March 1990 is calculated by using linear regression of two surveyed data sets (1986 and 1991) from the “Road Tree Planting Status Survey”. Next, the number of tall trees for each prefecture on 31st March 1990 is calculated by multiplying these values by the ratio of the number of tall trees for each prefecture on 31st March 2007. The ratio of the number of tall trees on 31st March 1990 is fixed to the value on 31st March 2007.
3. The number of tall trees which have been planted since 1st April 1990 is calculated by subtracting the value estimated in Step 1 from the value in Step 2 (RV is considered to be an activity which takes place after 1st January 1990. However, Japan considers RV as an activity after 1st April 1990 because the “Road Tree Planting Status Survey” has been implemented on a fiscal year basis).
4. The ratio of the number of tall trees planted on roads with a planted area less than 500 m² is estimated by using data from the sampling survey implemented in 2006 (general road: 1.00%, expressway: 0.00%, significance level: 95%).
5. The land area per tall tree is estimated by using modeled data from the sampling survey implemented in 2006 (general road: 0.0062 ha/tree, expressway: 0.0008 ha/tree, significance level: 95%). These modeled data are calculated by dividing randomly sampled RV land areas by the number of tall trees planted on the land).
6. The area of land planted with tall trees, which is 500 m² or more, is calculated by multiplying the values estimated in Steps 4 and 5 by the number of tall trees for each geographical boundary (prefecture) estimated in Step 3.

Area of land where tall trees have been planted since 1st April 1990 and whose size is 500 m² or more (ha)
 = 3. Number of tall trees planted since 1st April 1990 (tree)
 * 4. Ratio of the number of tall trees planted on land which is 500 m² or more (%)
 * 5. Land area per tall tree (ha/tree)

7. The area of land which was classified as forest land on 31st December 1989 is calculated by multiplying the area estimated in Step 6 by the sum of the area ratios of “Land that has been

² Land-use change from the previous year to each corresponding year is applied when the area ratio of “single year” is used.

converted from Forest land to Settlements per annum” since 1990 until each corresponding fiscal year during the commitment period.. This area is excluded because it is classified as D area. The remaining area is considered as RV land area.

8. The areas of “Remaining land (Settlements remaining Settlements)” and “Land converted from other land-use categories (Cropland / Grassland / Wetlands / Other land converted to Settlements)” are calculated by multiplying the land area estimated in Step 7 by the area ratio of “Land converted to Settlements” in the single year.

c) Green areas at ports

1. The green areas at ports which have been established since 1st January 1990 and which have a service area of 500 m² or more are extracted. Then, their areas are rearranged according to geographical boundaries (All green areas at ports can be reported because they are considered not to be classified as forest land on 31st December 1989).
2. The areas of “Remaining land (Settlements remaining Settlements)” and “Land converted from other land-use categories (Cropland / Grassland / Wetlands / Other land converted to Settlements)” are calculated by multiplying the land area estimated in Step 1 by the area ratio of “Land converted to Settlements” in the single year.

d) Green areas around sewage treatment facilities

1. The green areas around sewage treatment facilities which have been established since 1st January 1990 and which have a greening area of 500 m² or more are extracted. Then, their areas are rearranged according to geographical boundaries.
2. The area of land which was classified as forest land on 31st December 1989 is calculated by multiplying the greening areas estimated in Step 1 by the sum of the area ratios of “Land that has been converted from Forest land to Settlements per annum” since 1990 until each corresponding fiscal year during the commitment period. This area is excluded because it is classified as D area. The remaining area is considered as RV land area.
3. The areas of “Remaining land (Settlements remaining Settlements)” and “Land converted from other land-use categories (Cropland / Grassland / Wetlands / Other land converted to Settlements)” are calculated by multiplying the land area estimated in Step 2 by the area ratio of “Land converted to Settlements” in the single year.

e) Green areas by greenery promoting systems for private green space

1. The green areas by greenery promoting systems for private green space which have a greening area (excluding wall green areas) of 500 m² or more are extracted and their areas are rearranged according to geographical boundaries. All of them are activities which took place after 1st January 1990 because greenery promoting systems have been implemented since May 2001.
2. All green areas by greenery promoting systems for private green space to be reported are “Remaining land (Settlements remaining Settlements)” because they were not classified as Forest land on 31st December 1989, and land-use conversion, if any in recent years, occurred only within Settlements.

f) Green areas along rivers and erosion control sites

1. The greening works and erosion and sediment control works including hillside works in river zones which have been established since 1st January 1990 and which have a greening area of 500 m² or more are extracted (greening works: (1) – (8), erosion and sediment control works: (9) – (11) in the following table).

Table 11-9 RV projects in green areas along rivers and erosion control sites
and definition of planted land area

RV works in green areas along rivers and erosion control sites	Definition of planted land area
(1) Planting in inspection passage of excavated channel	Area of land from levee wall shoulder to private land
(2) Planting in face of river bank of excavated channel	Area of land from levee wall shoulder to private land
(3) Planting in backslope banquette	Area of embanked land
(4) Planting in levee marginal strip (second-class and third-class)	Area of marginal strip which is subject to greening works
(5) Planting in high water channel	Area of land from low-flow channel shoulder to foot of levee slope
(6) Planting in retarding basin	Area of retarding basin
(7) Planting in lake foreshore	Area of land from low-flow channel shoulder to foot of levee slope
(8) Planting in super levee	(Same as planting in excavated channel)
(9) Greening under erosion and sediment control works	Area of land which is subject to hillside works
(10) Greening under landslide control works	Area of land which is subject to hillside works
(11) Greening under steep slope failure prevention works	Area of land which is subject to hillside works

2. The planted land area in green areas along rivers and erosion control sites for each geographical boundary (prefecture) extracted in Step 1 is calculated. Double-counting between RV land and D land is prevented because forested land (on 1st January 1990) is not included in Step 1.
3. The land areas of “Remaining land (Settlements remaining Settlements)” and “Land converted from other land-use categories (Cropland / Grassland / Wetlands / Other land converted to Settlements)” are calculated by multiplying the land area estimated in Step 2 by the area ratio of “Land converted to Settlements (excluding Forest land converted to Settlements)” in the single year.

g) Green areas around government buildings

1. The green areas around government buildings which have been established since 1st January 1990 and whose RV land area (= total land area - building area) is 500 m² or more are extracted.
2. The RV land area for each geographical boundary (prefecture) extracted in Step 1 is calculated.
3. The area of land which was classified as Forest land on 31st December 1989 is calculated by multiplying the land area estimated in Step 2 by the sum of the area ratios of “Land that has been converted from Forest land to Settlements per annum” since 1990 until each corresponding fiscal year during the commitment period. This area is excluded because it is classified as D area. The remaining area is considered as RV land area.
4. The areas of “Remaining land (Settlements remaining Settlements)” and “Land converted from other land-use categories (Cropland / Grassland / Wetlands / Other land converted to Settlements)” are calculated by multiplying the land area estimated in Step 3 by the area ratio of “Land converted to Settlements” in the single year.

h) Green areas around public rental housing

1. The green areas around public rental housing which have been established since 1st January 1990 and which have a RV land area (= total land area - building area) of 500 m² or more are extracted.
2. The RV land area for each geographical boundary (prefecture) extracted in Step 1 are calculated.
3. The area of land which was classified as Forest land on 31st December 1989 is calculated by multiplying the land area estimated in Step 2 by the sum of the area ratios of “Land that has been converted from Forest land to Settlements per annum” since 1990 until each corresponding fiscal year during the commitment period. This area is excluded because it is classified as D area. The remaining area is considered as RV land area.
4. The areas of “Remaining land (Settlements remaining Settlements)” and “Land converted from other land-use categories (Cropland / Grassland / Wetlands / Other land converted to Settlements)” are calculated by multiplying the land area estimated in Step 3 by the area ratio of “Land converted to Settlements” in the single year.

11.3.2.5.b. Data

The data applied in estimating RV land area are shown below.

Table 11-10 Data applied in estimating RV land area

Sub-division	Data type	Method for data collection
Urban parks	• Area for each urban park	• Urban Parks Status Survey (FY2008, 2009, 2010)
Green area on roads	• Number of tall trees	• Road Tree Planting Status Survey (FY1987, 1992, 1997, 2002, 2007, 2008, 2009, 2010, 2011)
	• Land area per tall tree	• Basic Data Collection Survey on Tall Tree Planting on Roads (February, 2007)
Green areas at ports	• Service area	• Complete census for FY2008, 2009, 2010
Green areas around sewage treatment facilities	• Green area	• Sewage Treatment Facility Status Survey (FY2008, 2009, 2010)
Green areas by greenery promoting systems for private green space	• Greening area • Wall greening area • Number of tall trees	• Application form for greenery promoting systems for private green space • Urban Greening Status Survey (FY2008, 2009, 2010)
Green areas along river and erosion control sites	• Planted land area	• Survey on carbon dioxide absorption at source in river works (FY2008, 2009, 2010)
Green areas around government buildings	• Total land area and building area	• Complete census for FY2008, 2009, 2010
Green areas around public rental housing	• Total land area and building area	• Progress survey on tree planting for public rental housing (FY2008, 2009, 2010)

11.3.3. Maps and/or database to identify the geographical locations, and the system of identification codes for the geographical locations

Section 4.2.2.2 of the GPG-LULUCF shows two methods for identifying and reporting the units of land subject to Article 3.3 activities and lands subject to Article 3.4 activities. Reporting Method 1 entails delineating areas that include multiple land units subject to Article 3.3 and 3.4 activities by using legal, administrative, or ecosystem boundaries. Reporting Method 2 is based on the spatially explicit and complete geographical identification of all units of land subject to Article 3.3 activities and all lands subject to Article 3.4 activities.

Japan elects Reporting Method 1 in accordance with the decision tree indicated in Figure 4.2.4 in

Chapter 4 of the GPG-LULUCF, which means that the entire national land is stratified by using the geographic boundary of prefectures, and the total area of each “unit of land” subject to each Article 3.3 activity and each “land” subject to each Article 3.4 activity is reported within each boundary. The identification code is determined for each prefecture as shown in the following map. Each activity under Articles 3.3 and 3.4 is detected as described in sections 11.3.2.3-11.3.2.5, and the units of land or lands subject to it are identified within prefectural boundaries in accordance with Reporting Method 1.

This geographical boundary is applied for all units of land: units of land subject to activities under Article 3.3, units of land subject to activities under 3.3 which would otherwise be included in land subject to elected activities under Article 3.4, under the provisions of paragraph 8 of the Annex to the decision 16/CMP.1, and lands subject to elected activities under Article 3.4.

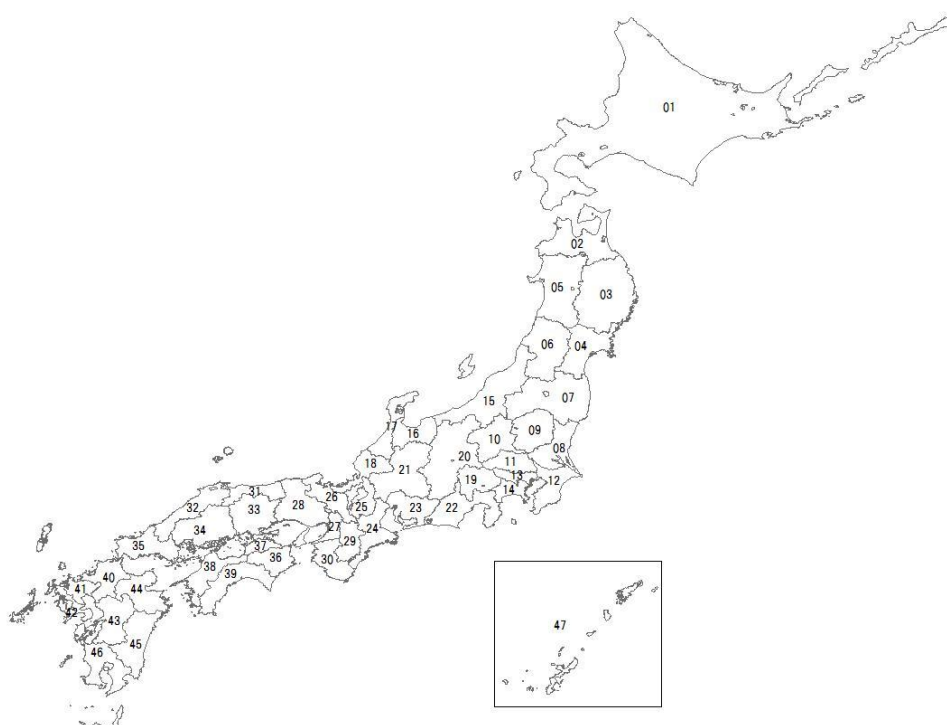


Figure 11-4 Japan’s determination of identification codes

Table 11-11 Identification codes and prefectures

ID	Prefecture	ID	Prefecture	ID	Prefecture	ID	Prefecture	ID	Prefecture
01	Hokkaido	11	Saitama	21	Gifu	31	Tottori	41	Saga
02	Aomori	12	Chiba	22	Shizuoka	32	Shimane	42	Nagasaki
03	Iwate	13	Tokyo	23	Aichi	33	Okayama	43	Kumamoto
04	Miyagi	14	Kanagawa	24	Mie	34	Hiroshima	44	Oita
05	Akita	15	Niigata	25	Shiga	35	Yamaguchi	45	Miyazaki
06	Yamagata	16	Toyama	26	Kyoto	36	Tokushima	46	Kagoshima
07	Fukushima	17	Ishikawa	27	Osaka	37	Kagawa	47	Okinawa
08	Ibaraki	18	Fukui	28	Hyogo	38	Ehime		
09	Tochigi	19	Yamanashi	29	Nara	39	Kochi		
10	Gunma	20	Nagano	30	Wakayama	40	Fukuoka		

11.4. Activity-specific information

11.4.1. Methods for carbon stock change and GHG emission and removal estimates

11.4.1.1. Description of the methodologies and the underlying assumptions used

11.4.1.1.a. Afforestation/Reforestation

a) Above-ground biomass, Below-ground biomass

● Methodology

The carbon stock change in living biomass in AR land is calculated using the Tier 2 stock change method in accordance with the GPG-LULUCF. In this method, the biomass stock change is estimated by subtracting the biomass stock change due to land conversion from the difference between the total amount of biomass at two times.

$$\Delta C_{LB} = \Delta C_{SC} - \Delta C_L$$

- ΔC_{LB} : Annual carbon stock change in living biomass [t-C/yr]
 ΔC_{SC} : Annual carbon stock change due to biomass growth, felling, fuelwood gathering, disturbance after land conversion [t-C/yr]
 ΔC_L : Annual carbon stock change due to land conversion [t-C/yr]

Carbon stock change due to biomass growth, felling, fuelwood gathering and disturbance after land conversion

$$\Delta C_{SC} = \sum_k \{(C_{t_2} - C_{t_1}) / (t_2 - t_1)\}_k$$

- ΔC_{SC} : Annual carbon stock change in living biomass [t-C/yr]
 t_1, t_2 : Time point of carbon stock measurement
 C_{t_1} : Total carbon in biomass calculated at time t_1 [t-C]
 C_{t_2} : Total carbon in biomass calculated at time t_2 [t-C]
 k : Type of forest management

The carbon stocks in living biomass are calculated from the volume for each tree species multiplied by wood density, biomass expansion factor, root-to-shoot ratio and carbon fraction.

$$C = \sum_j \{ [V_j \times D_j \times BEF_j] \times (1 + R_j) \times CF \}$$

- C : Carbon stock in living biomass [t-C]
 V : Volume [m³]
 D : Wood density [t-d.m./m³]
 BEF : Biomass expansion factor [dimensionless]
 R : Root-to-shoot ratio [dimensionless]
 CF : Carbon fraction (= 0.5[t-C/t-d.m.]
 j : Tree species

Carbon stock change due to land conversion

The carbon stock change due to land conversion is calculated as below, in accordance with the GPG-LULUCF.

$$\Delta C_L = \sum_i \{A_i \times (B_a - B_{b,i}) \times CF\}$$

- ΔC_L : Annual biomass carbon stock change in land that has been converted from other land-use type to forest [t-C/yr]
 A_i : Annual increase of land area that has been converted from land-use type i to forest [ha/yr]
 B_a : Dry matter weight per unit area immediately after conversion to forest [t-d.m./ha]
 $B_{b,i}$: Dry matter weight per unit area before conversion from land-use type i to forest [t-d.m./ha]
 CF : Carbon fraction of dry matter [t-C/t-d.m.]
 i : Type of land use

● Parameters

Data such as volume, biomass expansion factor, root-to-shoot ratio, wood density and carbon fraction are the same as those for reporting of LULUCF under the Convention. Detailed information is provided in section 7.4.1, Chapter 7 of this report.

The biomass stock data for each land use category which is used for estimation of biomass stock change due to land conversion are also the same as those for reporting of LULUCF under the Convention. Detailed information is provided in Table 7-5, Chapter 7 of this report.

● Activity data

The activity data is AR land area which was calculated by using the procedure described in section 11.3.2.3. of this report.

b) Dead wood, Litter and Soils

● Methodology

The carbon stock change in dead wood and litter in AR land is calculated in accordance with the basic stock change method provided by the GPG-LULUCF under the assumption that carbon stocks would change linearly over 20 years from those in non-forest land to those in forest land at the age of 20. The calculation is conducted by using average carbon stocks derived from the CENTURY-jfos model, and carbon stocks in dead wood and litter before land conversion are assumed to be zero.

$$\Delta C_{DW} = \sum_i \{A_i \times (C_{DW20} - C_{DW,i}) / 20\}$$

$$\Delta C_{LT} = \sum_i \{A_i \times (C_{LT20} - C_{LT,i}) / 20\}$$

- ΔC_{DW} : Annual carbon stock change in dead wood [t-C/yr]
 ΔC_{LT} : Annual carbon stock change in litter [t-C/yr]
 A_i : Afforested or reforested land area converted from land use i [ha]
 C_{DW20} : Average carbon stocks in dead wood per unit area of 20-year-old forests [t-C/ha]
 C_{LT20} : Average carbon stocks in litter per unit area of 20-year-old forests [t-C/ha]
 $C_{DW,i}$: Average carbon stocks in dead wood per unit area of land-use i [t-C/ha] (assumed to be zero)
 $C_{LT,i}$: Average carbon stocks in litter per unit area of land-use i [t-C/ha] (assumed to be zero)
 i : Type of land-use (cropland, grassland, wetlands, settlements and other land)

The carbon stock change in soils in AR land is calculated in accordance with the basic stock change method provided by the GPG-LULUCF under the assumption that carbon stocks would change linearly over 20 years from those in non-forest land to those in forest land at the age of 20. This calculation is conducted by using average carbon stocks derived from the CENTURY-jfos model.

$$\Delta C_{Soil} = \sum_i \{A_i \times (C_{Soil20} - C_{Soil,i}) / 20\}$$

ΔC_{Soil} : Annual carbon stock change in soils [t-C/yr]

A_i : Afforested or reforested land area converted from land-use i [ha]

C_{Soil20} : Average carbon stocks in soils per unit area of 20-year-old forests [t-C/ha]

$C_{Soil,i}$: Average carbon stocks in soils per unit area in land-use i [t-C/ha]

i : Type of land use (cropland, grassland, wetlands, settlements and other land)

● Parameters

The parameters are determined based on the CENTURY-jfos model and relevant literature.

● Activity data

The AR land area is calculated by using the procedure described in section 11.3.2.3. of this report.

c) Other gases

1) Direct N₂O emissions from N fertilization

It is assumed that the amount of nitrogen-based fertilizer applied in Forest land is counted in the Agriculture sector. Therefore, this category has been reported as “IE”.

2) CO₂ emissions from agricultural lime application

According to a survey in 2009 for private forests, all prefectures answered that no lime was applied to forest management practices like regeneration and tending in private forests. No lime was applied to such forest management practices in national forests either. Therefore, lime application in Forest land is considered as “not occurred” in Japan. This category has been reported as “NO” for all time series.

3) Biomass burning

GHG emissions from wild fire exist in Japan as explained in section 7.14.a), Chapter 7 of this report. Since there is no data which directly express biomass burning status in AR land, GHG emissions in AR land are estimated by multiplying GHG emissions due to fire for all forest land by the ratio of AR land area to all forest land area. Carbon released due to fire for all forest land (national forests and private forests) is estimated by multiplying the damaged timber volume due to fire by wood density, biomass expansion factor and carbon fraction of dry matter. Calculations only for non CO₂ emissions are performed since CO₂ emissions are already included in the calculation of carbon stock change.

d) Results

Table 11-12 Net emissions and removals from AR activity

	2008	2009	2010
	[Gg-CO ₂]	[Gg-CO ₂]	[Gg-CO ₂]
AR	-389.54	-415.03	-426.11
Above-ground biomass	-223.58	-242.50	-245.60
Below-ground biomass	-56.89	-60.60	-64.08
Dead wood	-65.69	-67.40	-74.03
Litter	-28.49	-29.24	-29.66
Soils	-14.91	-15.30	-12.74
Other gases	0.03	0.01	0.003

* CO₂ +: Emissions, -: Removals

11.4.1.1.b. Deforestation

a) Above-ground biomass, Below-ground biomass

● Methodology

The carbon stock change of living biomass (above-ground biomass and below-ground biomass) in D land is estimated by adding the living biomass loss in forests due to land conversion and carbon stock change due to growth of living biomass in D land after land conversion, in accordance with the GPG-LULUCF.

The forest living biomass loss due to land conversion is estimated from data in the NFRDB taking into account the status of D land such as tree species and forest, and all loss is allocated as emissions for the year of land conversion.

The carbon stock change due to growth of living biomass is estimated according to land use after conversion in D land. The land-use categories, except forest land where living biomass growth after conversion is calculated, are “Land converted to Grassland” and “Land converted to Settlements” as explained in Table 7-6 in Chapter 7 of this report. D land which is converted to settlement with living biomass growth is the land subject to RV practices. This is the land subject to both Article 3.3 and 3.4 activities, and the carbon stock change in this land is reported under D activity. The calculation is performed according to land-use status immediately after conversion in D land taking into account that D land is assumed to be seldom converted to other land uses again as explained in section 11.3.2.3.c.

$$\Delta C_{D-LB} = \Delta C_{DG-LB} + \Delta C_{DS-LB}$$

$$\Delta C_{DG-LB} = A_{5,DG} \times C_{G-LB}$$

$$\Delta C_{DS-LB} = \Delta C_{RV-LB} \times RA_{DS-RV}$$

ΔC_{D-LB} : Annual carbon stock change due to living biomass growth after D activity [t-C/yr]

ΔC_{DG-LB} : Carbon stock change due to living biomass growth in grassland subject to D activity [t-C/yr]

ΔC_{DS-LB} : Carbon stock change due to living biomass growth in settlements subject to D activity [t-C/yr]

ΔC_{RV-LB} : Carbon stock change in living biomass due to all RV practices [t-C/yr] (see section 11.4.1.1.d)

$A_{5,DG}$: Area of grassland subject to D activity within the past 5 years [ha]

C_{G-LB} : Carbon stock change per area in grassland [t-C/ha/yr]

RA_{DS-RV} : Ratio of the area subject to both D and RV activities within all areas subject to RV activities

● Parameters

Information relating to loss of forest biomass is obtained from the NFRDB. The parameter in Table 7-6 in Chapter 7 of this report is used for estimating carbon stock change due to living biomass growth after D activity in grassland. The parameters for estimating carbon stock change due to RV practices are the same as those used for RV activity.

● Activity data

The D land area is calculated by the method described in section 11.3.2.3. . The D land area where RV practices have been taken place is calculated by the method described in section 11.4.1.1.d.

b) Dead wood, Litter and Soils

The carbon stock change in dead wood, litter and soils associated with D is calculated in accordance with the Tier 2 method in the GPG-LULUCF. Japan assumes that all carbon stocks in dead wood and

litter are emitted at the time when D activities occur. The carbon stock change in soils is calculated under the assumption that soil carbon stocks change linearly over 20 years to those in non-forest land. Carbon stocks before and after conversion are established based on the data in Tables 7-7, 7-8 and 7-9 in Chapter 7 of this report, and data obtained from the CENTURY-jfos model.

c) Other gases

1) N₂O emissions from disturbance associated with land-use conversion to cropland

The GPG-LULUCF Tier 1 method, which utilizes mineralized soil carbon stocks due to disturbances associated with land-use conversion to cropland as an activity data, is applied to estimate N₂O emissions. The same methodology and parameters that were explained in section 7.12 b), Chapter 7 of this report are used. The carbon loss due to mineralization as a result of conversion to cropland in D area is calculated by all carbon loss due to deforestation multiplied by the ratio of land-use change to cropland in D area.

2) CO₂ emissions from agricultural lime application

CO₂ emissions from lime application in D land are estimated by the total CO₂ emissions from lime application in cropland in accordance with the GPG-LULUCF Tier 1 method (NIR Chapter 7 Section 7.13 b)) multiplied by the ratio of D area to the total area of cropland. Japan did not elect Cropland Management (CM) under Article 3.4 of the Kyoto Protocol; therefore CO₂ emissions from agricultural lime application to be reported under the Kyoto Protocol are only those in “Cropland converted from Forest land” since 1990 (identified as D land). However, it is difficult to directly determine the amount of lime and dolomite applied in such lands. Therefore it is assumed that lime application is conducted uniformly in all cropland.

3) Biomass burning

Prescribed fire associated with D activity is very rarely performed in Japan because of severe restrictions imposed by the “Waste Management and Public Cleansing Law” and the “Fire Defense Law”. Therefore, CH₄, CO, N₂O, and NO_x emissions are reported as “NO”.

d) Results

Table 11-13 Net emissions and removals from D activity

	2008	2009	2010
	[Gg-CO ₂]	[Gg-CO ₂]	[Gg-CO ₂]
D	2,456.72	3,115.09	4,822.89
Above-ground biomass	1,296.59	1,645.55	2,661.16
Below-ground biomass	331.96	421.09	681.43
Dead wood	434.84	543.13	864.04
Litter	174.12	217.92	347.64
Soils	215.11	282.71	264.00
Other gases	4.10	4.69	4.62

* CO₂ +: Emissions, -: Removals

11.4.1.1.c. Forest Management

a) Above-ground biomass, Below-ground biomass

● Methodology

1. Emissions/removals in all forest land are estimated by using biomass stock data stored in the NFRDB (based on the stock change method).
2. Emissions/removals relating to ARD activities are subtracted from emissions/removals in all forest land. For Ikusei-rin forest, emissions/removals in FM land are estimated by applying the FM ratio for each tree species, region and age class. For Tennensei-rin forest, the area of forest land with standing trees subject to practices for protection or conservation of forests such as controlling logging activities and land-use change which have been implemented under laws are identified by using the NFRDB, and emissions/removals are estimated.

● Parameters

The parameters are the same as those used for AR.

b) Dead wood, Litter and Soils

● Methodology

The carbon stock change in each pool is estimated by the Tier 3 method. It is estimated by multiplying carbon emissions/removals per area in each pool, which are calculated by the CENTURY-jfos model for each type of forest management, by the land area of each type of forest management and then summing them.

$$\Delta C_{dls} = \sum_{k,m,j} (A_{k,m,j} \times (d_{k,m,j} + l_{k,m,j} + s_{k,m,j}))$$

ΔC_{dls} : Carbon stock change in dead wood, litter and soil [t-C/yr]

A : Area [ha]

d : Average carbon stock change in dead wood per unit area [t-C/ha/yr]

l : Average carbon stock change in litter per unit area [t-C/ha/yr]

s : Average carbon stock change in soils per unit area [t-C/ha/yr]

k : Type of forest management

m : Age class or forest age

j : Tree species

● Parameters

The average carbon stock changes per unit area for dead wood, litter and soils are calculated by the CENTURY-jfos model, which was the modified version of the CENTURY model (Colorado State University) to accommodate for Japanese climate, soil, and vegetation conditions. Detailed explanation of the CENTURY-jfos model is provided in section 7.4.1.b).2), Chapter 7 of this report.

c) Other gases

1) Direct N₂O emissions from N fertilization

It is assumed that the amount of nitrogen-based fertilizer applied in forest land is included in the amount of nitrogen-based fertilizer counted in the Agriculture sector. Therefore, this category is reported as “IE”.

2) N₂O emissions from drainage of soils

Based on expert judgment, N₂O emissions are extremely low, because soil drainage activities are very rarely conducted in Japan. Therefore, this category is reported as “NO”.

3) CO₂ emissions from agricultural lime application

According to a survey in 2009 for private forests, all prefectures answered that no lime was applied to forest management practices like regeneration and tending in private forests. No lime was applied to such forest management practices in national forests either. Therefore, lime application in forest land is considered as “not occurred” in Japan. This category has been reported as “NO” for all time series.

4) Biomass burning

Emissions due to biomass burning are estimated in the same way as in the case of AR by multiplying GHG emissions due to fire for all forest land by the ratio of FM land area to all forest land area.

d) Results

Table 11-14 Net emissions and removals from FM activity

	2008	2009	2010
	[Gg-CO ₂]	[Gg-CO ₂]	[Gg-CO ₂]
FM	-45,388.77	-49,005.55	-53,251.78
Above-ground biomass	-34,747.68	-37,955.21	-41,795.32
Below-ground biomass	-8,758.73	-9,581.09	-10,549.16
Dead wood	134.69	540.79	1,050.80
Litter	-472.06	-394.24	-356.19
Soils	-1,559.02	-1,621.64	-1,603.43
Other gases	14.05	5.83	1.52

* CO₂ +: Emissions, -: Removals

11.4.1.1.d. Revegetation

Methodologies for estimating GHG emissions and removals from RV activity are described in two cases: when RV activity is performed on the land where no land conversion has occurred (remaining land) and on the land where land conversion has occurred (Conversion Land).

a) Remaining land: Above-ground biomass, Below-ground biomass

Japan estimates the carbon stock change in above-ground biomass and below-ground biomass of tall trees planted in RV lands. Tall trees are consistent with the definition in “Standards for the quality and size of planted trees for the public (draft)”³.

● Methodology

$$\Delta C_{RVLB} = \sum_i (\Delta C_{LBG,i} - \Delta C_{LBL,i})$$

$$\Delta C_{LBG,i} = \Delta B_{LBG,i}$$

$$\Delta B_{LBG,i} = \sum_j (NT_{i,j} \times C_{Ratei,j})$$

ΔC_{RVLB} : Annual carbon stock changes in living biomass in remaining RV land [t-C/yr]

ΔC_{LBG} : Annual carbon stock changes due to living biomass growth in remaining RV land [t-C/yr]

ΔC_{LBL} : Annual carbon stock changes due to living biomass loss in remaining RV land [t-C/yr]

ΔB_{LBG} : Annual living biomass growth in RV land [t-C/yr]

³ “Standards for the quality and size of planted trees for the public (draft)” was decided by the Ministry of Land, Infrastructure, Transport and Tourism in order to promote proper enforcement of projects such as greening in public spaces. Tall tree is defined in the standards as tree which reaches 3 ~ 5 m in height.

C_{Rate}	: Annual living biomass growth rate per tree [t-C/tree/yr]
NT	: Number of trees
i	: Type of urban green facilities (Urban parks, Green areas on roads, Green areas at ports, Green areas around sewage treatment facilities, Green areas by greenery promoting systems for private green space, Green areas along rivers and erosion control sites, Green areas around public rental housing and Green areas around government buildings)
j	: Tree species

● Parameters⁴

➤ **Urban parks**

Carbon stock changes due to the loss of living biomass in urban parks are assumed to be zero based on Tier 1 method in GPG-LULUCF (p. 3.297), because the average age of trees is found to be less than or equal to 20 years in the tree survey for sample urban parks⁵.

The annual living biomass growth of trees in urban parks is calculated by using the country-specific value for annual growth rate of living biomass per tree, which was developed by combining the default values (0.0033-0.0142 t-C/tree/yr) provided in the GPG-LULUCF (p. 3.297, Table 3A.4.1) and the country specific annual growth rates of living biomass for the trees in Japan (0.0204 for Japanese zelkova, 0.0103 for ginkgo, 0.0095, for bamboo-leaf oak and 0.0122 t-C/tree/yr for camphor tree) by taking into account the distribution ratio of tree species in sample urban parks⁶. The annual growth rates of living biomass for Japanese zelkova, ginkgo, bamboo-leaf oak and camphor tree are calculated by using the growth curve for each tree species⁷, which were developed based on the results of surveys conducted by the National Institute for Land and Infrastructure Management (NILIM) of the Ministry of Land, Infrastructure, Transport and Tourism (MLIT) and the average trunk diameter at breast height for each tree species⁸, which were determined from the results of surveys in urban parks. For the ratio of above-ground biomass/below-ground biomass, the default value (root-to-shoot ratio: 0.26) provided in the 2006 IPCC Guidelines (p. 8.9) is applied.

➤ **Green areas on roads**

Carbon stock changes due to the loss of living biomass in green areas on roads are assumed to be zero, because the average age of trees is found to be less than or equal to 20 years for those trees planted in randomly extracted green areas on roads.

The annual living biomass growth in green areas on roads is calculated by using the country-specific value for annual growth rate of living biomass per tree, which was developed by combining the default values and the annual growth rates of living biomass for the trees in Japan (4 species), which were also used for the urban parks, taking into account the distribution ratio of tree species indicated by the surveys in green areas on roads⁹.

⁴ The Tier 1b method described in the GPG-LULUCF and the Tier 2 method with country-specific annual biomass growth rates are applied for the estimation of the annual growth rate of living biomass per tree. Japan will further improve the accuracy of this estimation.

⁵ 129 samples were randomly extracted from the urban parks notified after 1st of January 1990 and located in Kanagawa prefecture, which is located in Japan's typical climate zone and has various types of urban parks. In addition, the same survey was implemented in 3 urban parks in Chiba prefecture, which is located next to Kanagawa prefecture, in order to cover the park types that did not exist in Kanagawa prefecture.

⁶ The distribution ratio of tree types was calculated by using tree registers and plantation maps for all urban parks in Kushiro city and Yubari city in Hokkaido and for 321 randomly extracted urban parks in the other prefectures.

⁷ Reference: Matsue et al., "Estimation equations for the amount of CO₂ fixed by planted trees in cities in Japan", Journal of the Japanese Society of Revegetation Technology, 35 (2), 318-324, 2009.

⁸ Reference: Parks and Green Spaces Division of the Ministry of Land, Infrastructure and Transport, "FY2004 Survey on evaluation techniques for the effectiveness of greening in urban parks for preventing global warming", March, 2005.

⁹ The distribution ratio of tree types is taken from the *Road Tree Planting Status Survey (The Street tree of Japan VI)*, which covered green areas on roads throughout Japan.

For the ratio of above-ground biomass/below-ground biomass, the same value used for urban parks is applied.

➤ **Urban green areas other than Urban parks, Green areas on roads and Green areas by greenery promoting systems for private green space**

Carbon stock changes due to the loss of living biomass in these green areas are assumed to be zero, because the standard of planted trees, tree types and their distribution are applied in the same manner as in urban parks.

The annual living biomass growth and the ratio of above-ground biomass/below-ground biomass are the same parameters as for urban parks.

➤ **Green areas by greenery promoting systems for private green space**

Carbon stock changes due to the loss of living biomass in these green areas are assumed to be zero, because the standard of planted trees is selected in the same manner as in urban parks and all facilities have been certified since 2002.

The annual living biomass growth and the ratio of above-ground biomass/below-ground biomass are the same parameters as for urban parks.

● **Activity data**

➤ **Urban parks**

The area of land remaining urban parks is calculated by multiplying the area of urban parks by the area ratio of land conversion for the whole country. The activity data for carbon stock changes in living biomass in urban parks is the number of tall trees planted in urban parks which is calculated by multiplying the area of urban parks obtained from the “Urban Parks Status Survey” by the number of tall trees per area (Hokkaido: 329.5 tree/ha, the other prefectures: 222.3 tree/ha). The number of tall trees per area is calculated based on the number of tall trees and the land areas of sample urban parks, whose sample number was intended to satisfy the significance level of 95%.¹⁰

Table 11-15 Area of urban parks which were not classified as forest land on 31st December 1989

		At the end of 2010		
	Land-use category	Area ratio of land which has been converted from Forest land to Settlements since 1990 until the corresponding fiscal year during the commitment period	Area [ha]	Classified as RV land
Urban parks which have been notified since 1st January 1990 and whose establishment area is 500 m ² or more	Forest	5.52%	2,966.28	No
	Non-forest	94.48%	50,768.14	Yes
	Total	100.00%	53,734.42	-

¹⁰ The number of tall trees per area in urban parks was calculated by using data from tree registers and planting maps for randomly extracted 176 sample urban parks in Hokkaido and 321 sample urban parks in the other prefectures. For Hokkaido, the number of samples was not sufficient to satisfy the significant level of 95% because the tree register has not been developed completely.

Table 11-16 Area of urban parks (remaining land / converted land)

At the end of 2010

	Land-use category	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Urban parks which have been notified since 1st January 1990 and whose establishment area is 500 m ² or more (classified as RV land)	Converted (except land converted from forest land)	0.24%	121.18	28,452
	Remaining	99.76%	50,646.96	11,891,495
	Total	100.00%	50,768.14	11,919,947

➤ **Green areas on roads**

The activity data (the number of tall trees) in “Remaining green areas on roads” is calculated by the following procedures.

1. The number of tall trees in all green areas on roads on 31st of March 1990 and in the end of each corresponding fiscal year during the commitment period is estimated by using data from the “Road Tree Planting Status Survey” which have been implemented in FY1987, FY1992 and each corresponding fiscal year.
2. The number of tall trees planted after 1st of April 1990 was calculated by subtracting the number for 31st of March 1990 from the number for the end of each corresponding fiscal year (RV is an activity which takes place after 1st of January 1990. However, Japan considers it an activity after 1st of April 1990 because it is impossible to estimate the number of tall trees which have been planted between 1st of April 1990 and 31st of March 1990).
3. The number of tall trees calculated in Step 2 is multiplied by the ratio of the number of tall trees planted on roads whose planted area is more than 500 m².
4. The number of tall trees calculated in Step 3 is multiplied by the area ratio of green areas on roads, which were classified as “Forest land” on 31st December 1989.
5. The number of tall trees calculated in Step 4 is multiplied by the area ratio of “Land remaining Settlements”.

Table 11-17 Area of green areas on roads which have been classified as RV

At the end of 2010

	Area of green areas on roads per tall tree [ha/tree]	Number of planted tall trees [tree]			Area ratio of planted lands which are 500 m ² or more [%]	Area ratio of land which was classified as forest land on 31st December 1989 [%]	Area of green areas on roads which was classified as RV land [ha]	Activity data [Number of tall trees]
		31st March 1990	31st March 2011	FY1990 - FY2010				
		a	b	c-b				
General roads (managed by the MLITT, Prefectures, local authorities, public corporations)	0.006237	4,342,070	6,884,950	2,542,880	99.00%	5.52%	14,834	2,378,401
Expressway (managed by now-defunct public corporation)	0.000829903	1,096,380	8,208,260	7,111,880	100.00%	5.52%	5,576	6,719,285
Total	—	5,438,450	15,093,210	9,654,760	—	—	20,410	9,097,686

MLIT: Ministry of Land, Infrastructure, Transport and Tourism

Table 11-18 Area of green areas on roads which have been classified as RV and activity data [number of tall trees] (remaining land / converted land)

At the end of 2010

	Land-use category	Area ratio of land which has been converted in the current year	Activity data [Number of tall trees]	Area [ha]
Green areas on roads which have been notified since 1st January 1990 and whose establishment area is 500 m ² or more (classified as RV land)	Converted	0.24%	21,716	48.72
	Remaining	99.76%	9,075,970	20,361.72
	Total	100.00%	9,097,686	20,410.44
General roads	Converted	0.24%	5,677	35.41
	Remaining	99.76%	2,372,724	14,798.68
	Total	100.00%	2,378,401	14,834.09
Expressway	Converted	0.24%	16,039	13.31
	Remaining	99.76%	6,703,247	5,563.04
	Total	100.00%	6,719,285	5,576.35

➤ **Green areas at ports**

The activity data for carbon stock changes in living biomass in green areas at ports is the number of tall trees planted in green areas at ports. The activity data is calculated by multiplying the service area obtained from complete census by the number of tall trees per unit area of urban parks (329.5 trees/ha for Hokkaido and 222.3 trees/ha for the other prefectures). These values were adopted by taking into account the similarities between the urban parks and the green areas at ports as mentioned above. All green areas at ports are located in “Settlements” and judged not being classified as “Forest land” on 31st of December 1989.

Table 11-19 Area of green areas at ports and activity data (remaining land / converted land)

At the end of 2010

Land-use category	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Converted	0.24%	3.78	859
Remaining	99.76%	1,578.48	359,164
Total	100.00%	1,582.26	360,023

➤ **Green areas around sewage treatment facilities**

The area of land remaining green areas around sewage treatment facilities is calculated in the same manner as for urban parks. The activity data for carbon stock change in living biomass in green areas around sewage treatment facilities are obtained from the “Sewage Treatment Facility Status Survey” for each fiscal year during the commitment period. The number of tall trees planted in green areas around sewage treatment facilities is calculated by multiplying the greening areas by the number of tall trees per greening area (129.8 tree/ha for Hokkaido and 429.2 tree/ha for the other prefectures).¹¹ All green areas around sewage treatment facilities are located in “Settlements”.

¹¹ The number of tall trees per area for green areas around sewage treatment facilities was established by using data on the number of tall trees and greening areas measured in 59 green areas.

Table 11-20 Area of green areas around sewage treatment facilities which were not classified as “Forest land” on 31st December 1989

At the end of 2010

Land-use category	Area ratio of land which has been converted from Forest land to Settlements since 1990 until the corresponding fiscal year during the commitment period	Area (ha) (Green areas)	Classified as RV land
Forest	5.52%	38.22	No
Non-forest	94.48%	654.10	Yes
Total	100.00%	692.32	-

Table 11-21 Area and activity data of “Green areas around sewage treatment facilities” [number of tall trees] (remaining land / converted land)

At the end of 2010

Land-use category	Area ratio of land has been converted for the current year	Area [ha] (Green areas)	Activity data [Number of tall trees]
Converted (except land converted from forest land)	0.24%	1.56	633
Remaining	99.76%	652.54	264,513
Total	100.00%	654.10	265,146

➤ **Green areas by greenery promoting systems for private green space**

Activity data (the number of tall trees) is available for each facility. Therefore, the total number of tall trees is used as activity data.

Table 11-22 Activity data and area of “Green areas by greenery promoting systems for private green space”

Certification year	Location	Area [m ²]	Breakdown of area [m ²]			Area Wall green area by greenery promoting system for private green space [m ²]	Activity data Number of tall trees [tree]
			Ground	Roof	Wall		
2002	Minato-ku, Tokyo	17,244	1,314	2,042	106	3,356	335
2002	Minato-ku, Tokyo	19,708	3,285	736		4,021	147
2002	Minato-ku, Tokyo	52,766	10,679			10,679	672
2002	Minato-ku, Tokyo	84,780	8,846	7,493		16,339	813
2003	Minato-ku, Tokyo	5,519	1,654			1,654	167
2003	Osaka City	22,282	1,527	3,164	110	4,691	500
2005	Kawaguchi City	1,995	586	164	18	750	153
2006	Kyoto City	3,857	1,271			1,271	90
2006	Hiroshima City	4,453	130	783		913	1
2007	Hiroshima City	14,353	4,058			4,058	261
2007	Fukuoka City	5,689	602	799		1,401	19
2008	Ishikawa Prefecture	7,281	682	1,411		2,093	19
2009	Setagaya-ku, Tokyo	5,526	1,116			1,116	51
2009	Setagaya-ku, Tokyo	6,459	1,370			1,370	15
2010	—						
	Total	251,912	37,120	16,592	234	53,712	3,243

* There were no areas certified in FY2010.

➤ **Green areas along rivers and erosion control sites**

The area of land remaining green areas along rivers and erosion control sites is calculated by multiplying the area of this green area by the area ratio of land conversion for the whole country. The activity data for living biomass (the number of tall trees) is calculated by multiplying this area by the number of tall trees per area (Hokkaido: 1470.8 tree/ha, the other prefectures: 339.0 tree/ha).¹²

The green areas along rivers and erosion control sites exclude lands, which were classified as Forest land at the time of survey. Therefore, land conversion from Forest land is not taken into account for the estimation of activity data.

Table 11-23 Activity data and area of "Green areas along rivers and erosion control sites"
(remaining land / converted land)

At the end of 2010

	Land-use category	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Green areas along rivers and erosion control sites which have been established since 1st January 1990 and whose establishment area is 500 m ² or more (classified as RV land)	Converted (except land converted from forest land)	0.01%	3.72	2,133
	Remaining	99.99%	1,553.24	891,440
	Total	100.00%	1,556.96	893,573

➤ **Green areas around government buildings**

The area of land remaining green area around government buildings is calculated by multiplying the area of this green area by the area ratio of land conversion for the whole country. The activity data for living biomass (the number of tall trees) is calculated by multiplying this area by the number of tall trees per area (all prefecture: 108.8 tree/ha).¹³

Table 11-24 Area of "Green areas around government buildings" which were not classified as "Forest land" on 31st of December 1989

At the end of 2010

	Land-use category	Area ratio of land which has been converted from Forest land to Settlements since 1990 until the corresponding fiscal year during the commitment period	Area [ha] (Green areas)	Classified as RV land
Green areas around government buildings which have been established since 1st January 1990 and whose establishment area is 500 m ² or more	Forest	5.52%	16.46	No
	Non-forest	94.48%	281.71	Yes
	Total	100.00%	298.17	-

¹² For green areas along rivers and erosion control sites, the number of tall trees was measured in approximately 95% of this green area. Based on these data, the number of planted trees per area was estimated in order to simplify the estimation of the number of tall trees in all green areas.

¹³ For green areas around government buildings, the number of tall trees per area was estimated by dividing the number of tall trees by the "total land area – building area" (these data were based on 30 facilities where planting maps were available). The common value is used for all prefectures, since the sample data were not sufficient enough to set values for Hokkaido and the other prefectures, respectively.

Table 11-25 Area and activity data of “Green areas around government buildings” (remaining land / converted land)

At the end of 2010

	Land-use category	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Green areas around government buildings which have been established since 1st January 1990 and whose establishment area is 500 m ² or more (classified as RV land)	Converted (except land converted from forest land)	0.24%	0.67	73
	Remaining	99.76%	281.04	30,577
	Total	100.00%	281.71	30,650

➤ **Green areas around public rental housing**

The area of land remaining green areas around public rental housing is calculated by multiplying the area of this green area by the area ratio of land conversion for the whole country. The activity data for living biomass (the number of tall trees) is calculated by multiplying this area by the number of tall trees per area (all prefecture: 219.9 tree/ha).¹⁴

Table 11-26 Area of “Green areas around public rental housing” which were not classified as “Forest land” on 31st of December 1989

At the end of 2010

	Land-use category	Area ratio of land which has been converted from Forest land to Settlements since 1990 until the corresponding fiscal year during the commitment period	Area [ha] (Green areas)	Classified as RV land
Green areas around public rental housing which have been established since 1st January 1990 and whose establishment area is 500 m ² or more	Forest	5.52%	127.51	No
	Non-forest	94.48%	2,182.35	Yes
	Total	100.00%	2,309.86	-

Table 11-27 Area and activity data of “Green areas around public rental housing” (remaining land / converted land)

At the end of 2010

	Land-use category	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Green areas around public rental housing which have been established since 1st January 1990 and whose establishment area is 500 m ² or more (classified as RV land)	Converted (except land converted from forest land)	0.24%	5.21	1,145
	Remaining	99.76%	2,177.14	478,753
	Total	100.00%	2,182.35	479,898

b) Remaining land: Dead wood

➤ **Urban parks**

The number of tall trees per land area used in the estimation of activity data for living biomass

¹⁴ For green areas around public rental housing, the number of tall trees per area was estimated for 33 facilities, where planting maps were available, by dividing the number of tall trees by the area “total land area – building area”. The common value is used for all prefectures, since the sample data were not sufficient enough to set values for Hokkaido and the other prefectures, respectively.

includes trees which have died and have been complementary planted since the establishment of the park. Thus the carbon stock changes in dead wood are thought to be included in the carbon stock changes in living biomass. Therefore, this category is reported as “IE”.

➤ **Green areas on roads**

The number of tall trees used in the estimation of activity data for living biomass is surveyed every 5 years (implemented every year since 2007). These data include the effects of dead wood and complementary planting, thus the carbon stock change in dead wood is included in the carbon stock changes in living biomass. Therefore, this category is reported as “IE”.

➤ **Urban green facilities other than Urban parks and Green areas on roads**

These categories are reported as “IE” based on the same assumption as urban parks.

c) **Remaining land: Litter**

Carbon stock changes in litter are estimated for urban parks and green areas at ports.

● **Methodology**

$$\Delta C_{RVLit} = \sum_i (A_i \times L_{it,i})$$

ΔC_{RVLit} : Annual carbon stock changes in litter in remaining RV land [t-C/yr]

A : Area of remaining RV land [ha]

L_{it} : Annual carbon stock changes in litter per RV land [t-C/ha/yr]

i : Type of urban green facilities (Urban parks and Green areas at ports)

● **Parameters**

➤ **Urban parks and Green areas at ports**

For litter, Japan estimates carbon stock changes only in branches and leaves dropped naturally from tall trees. The carbon stock changes in litter per urban park area are calculated by using the annual accumulation of litter per tall tree (all prefectures: 0.0006 t-C/tree/yr) based on the results of a field survey in urban parks¹⁵, the number of tall trees per area and the ratio of litter moved to off-site due to management including cleaning (54.4%). As a result, carbon stock change in litter per urban park area is calculated to be 0.0882 t-C/ha/yr for Hokkaido and 0.0594 t-C/ha/yr for other prefectures. In addition, the carbon fraction in litter is assumed to be 0.4 t-C/t-d.m., which is a default value provided in the 2006 IPCC Guidelines (p.8.21).

➤ **Urban green facilities other than Urban parks and Green areas at ports**

Litter in these urban green facilities includes branches and leaves dropped naturally and dead roots. A part of the litter remains on-site and leads to an increase in carbon stocks, although other litter is moved to off-site due to management such as cleaning (such litter is dropped from trees planted after the establishment of green areas). Dead roots also lead to an increase in carbon stocks because they are not moved to off-site.

¹⁵ The annual accumulation of litter dropped naturally was measured for some tree types by using litter traps installed in Takino Suzuran Kyuryo National Government Park (Hokkaido) and Showa Kinen National Government Park (Tokyo). Litter is defined as branches and leaves dropped on the surface. In the selection of parks for the survey, large-sized and intensively managed national government parks in which continuous monitoring is available and different types of trees have been planted are considered to meet the measurement requirements. In addition, it is also considered that the distribution of tree types differs between Hokkaido and other prefectures. Therefore, Japan selected two parks for the survey, one in Hokkaido and the other in a typical climate zone excluding Hokkaido.

The carbon stock changes in these urban green facilities cannot be estimated accurately because it is difficult to obtain detailed information on various managements (such as cleaning). However, it is clear that the input of litter and dead roots increases carbon stocks. Therefore, these sub-categories are not sources of GHGs and are not included in the reporting (the exclusion of these sub-categories is assumed to be conservative).

● Activity data

The activity data is the same as for living biomass.

d) Remaining land: Soils

Urban parks, for which the carbon stock changes in soils per area were determined, and Green areas at ports, whose management practices are similar to those for urban parks, are the subject of estimation. In general, soils in RV land are not organic soils (peat soils and muck soils). Therefore, organic soils are reported as “NO”, and only mineral soils are estimated.

● Methodology

$$\Delta C_{RVSoils} = \sum_i (\Delta C_{Mineral,i} - L_{Organic,i})$$

$$\Delta C_{Mineral,i} = A_i \times \Delta C_{Soil,i}$$

$\Delta C_{RVSoils}$: Annual carbon stock changes in soils in remaining RV land¹⁶ [t-C/yr]

$\Delta C_{Mineral}$: Annual carbon stock changes in mineral soils in RV land [t-C/yr]

$L_{Organic}$: Annual carbon stock changes in organic soils in RV land (=0) [t-C/yr]

A : Area of remaining RV land [ha]

ΔC_{Soil} : Annual carbon stock changes in soils per area of remaining RV land [t-C/ha/yr]

i : Type of urban green facilities (Urban parks and Green areas at ports)

● Parameters

➤ Urban parks and Green areas at ports

Carbon stock changes in soils per area of RV land are estimated based on the results of surveys¹⁷ conducted in urban parks which have been established within 20 years (1.20 t-C/ha/yr)¹⁸.

This value is applicable to land, which is the subject of revegetation activity and was established within 20 years, because the value is based on the results of surveys conducted in urban parks which have been established within 20 years.

¹⁶ Soil organic carbon pools are the subject of estimation of carbon stock changes in soils in RV land.

¹⁷ Soil carbon stocks (at 30 cm depth) were measured for areas with different types of vegetation cover in urban parks (planted: 21 areas, lawn: 19 areas, bare: 21 areas), which are located in Tokyo and were established in different years.

¹⁸ Since urban parks are generally established by turning entire sites into urban parks, soil carbon stocks within the site immediately after establishment are assumed to be uniform irrespective of previous types of vegetation cover. The soil carbon stocks of the area, where basically carbon is not supplied by plants (bare area), are assumed to be the same as soil carbon stocks of sites immediately after conversion. Based on the soil carbon stocks in the areas with different types of vegetation cover (planted, lawn and bare) in urban parks, which were established in different years, “carbon accumulation rates in planted areas” and “carbon accumulation rates in lawn areas” are calculated:

- Carbon accumulation rates in planted areas = “Difference in soil carbon stocks between planted and bare areas” / “Average years after establishment of surveyed planted areas”
- Carbon accumulation rates in lawn areas = “Difference in soil carbon stocks between lawn and bare areas” / “Average years after establishment of surveyed lawn areas”

Furthermore, changes in soil carbon stocks per area are determined by taking the weighted average based on the typical area ratio among planted, lawn and bare sites in urban parks. The soil carbon stocks of bare area are about 38 t-C/ha when converted from the sample data.

➤ **Urban green facilities other than Urban parks and Green areas at ports**

It is assumed that the patterns of carbon stock changes in soils in these urban green facilities are similar to those in urban parks, because planting, establishment and management practices are implemented in a similar way. The expressway slopes, where different plantation practices are applied, are assumed to be a sink, because field surveys have revealed that the carbon stocks keep increasing for at least 20 years after establishment.

Nevertheless, in this submission, these urban green facilities are reported as “NR”, because available data are not sufficient to estimate carbon stock changes in soils in these facilities. The estimation for urban green facilities other than urban parks and green areas at ports will be further considered in the future.

● **Activity data**

The area is as obtained for estimating the activity data for living biomass.

e) **Remaining land: Other gases**

1) **Direct N₂O emissions from N fertilization**

It is assumed that the volume of nitrogen-based fertilizer applied to urban parks is included in the demand for nitrogen-based fertilizers in the Agriculture sector, although fertilization in urban parks has been conducted in Japan. Therefore, these sources have been reported as “IE”.

2) **Carbon emissions from lime application**

Japan estimates carbon emissions from lime application in all urban green facilities. For urban parks and green areas on roads (lime application is implemented only in green areas on general roads), the amount of lime applied per area is estimated. For other urban green facilities, the amount of lime applied per area of urban parks is applied.

Carbon emissions are estimated for all RV land together because the estimation method is similar regardless of remaining or converted land.

● **Methodology**

$$C_{RVLm} = C_{RVCaCO_3} + C_{RVCaMg(CO_3)_2}$$

$$C_{RVCaCO_3} = \sum_i (A_i \times \Delta C_{RVCaCO_3,i}) \times 12.01/100.09$$

$$C_{RVMg(CO_3)_2} = \sum_i (A_i \times \Delta C_{RVMg(CO_3)_2,i}) \times 12.01/184.41$$

C_{RVLm}	: Annual carbon emissions in RV lands due to lime application [t-C/yr]
C_{RVCaCO_3}	: Carbon emissions in RV lands due to calcic limestone application [t-C/yr]
$C_{RVCaMg(CO_3)_2}$: Carbon emissions in RV lands due to dolomite application [t-C/yr]
A_i	: Area of RV land (total of remaining and converted lands of urban green facilities, i) [ha]
$\Delta C_{RVCaCO_3,i}$: Amount of calcic limestone application per area of RV land (urban green facilities, i) [t-C/ha]
$\Delta C_{RVCaMg(CO_3)_2,i}$: Amount of dolomite application per area of RV land (urban green facilities i) [t-C/ha]
12.01/100.09	: Ratio of molecular weight of carbon in calcic limestone
12.01/184.41	: Ratio of molecular weight of carbon in dolomite
i	: Types of urban green facilities (Urban parks, Green areas on roads, Green areas at ports, Green areas around sewage treatment facilities, Green areas by greenery promoting systems for private green space, Green areas along rivers and erosion control sites, Green areas around public rental housing and Green areas around government buildings)

● Parameters

➤ Urban parks

The amount of calcic limestone application per area is established as 298.4 g/ha/yr based on the results of a questionnaire survey carried out for 11,274 urban parks. The amount of dolomite application per area is established as 1,088.4 g/ha/yr based on the results of a questionnaire survey carried out for 9,346 urban parks.

In estimating carbon emissions, it is assumed that all carbon included in applied calcic limestone and dolomite is released to the atmosphere within the application year.

➤ Green areas on roads

The amount of calcic limestone application per tall tree is established as 0.3311 g/tree/yr and the amount of dolomite application per tall tree is established as 1.5431 g/tree/yr based on the results of a questionnaire survey implemented for 40 road managers.

In estimating carbon emissions, it is assumed that all carbon included in applied calcic limestone and dolomite is released to the atmosphere within the application year.

➤ Urban green facilities other than Urban parks and Green areas on roads

The parameter values for urban parks are applied because lime application in these green facilities is implemented in the same manner as in urban parks (application pattern and frequency).

● Activity data

The area of all RV lands (regardless of remaining or converted land) is used as activity data.

3) Biomass burning

In settlements subjected to RV activities, burning of residues is essentially prohibited by the Law for Waste Treatment and Cleaning. In addition, wild fires do not usually occur in lands subjected to RV activities because these lands are managed. Therefore, biomass burning activities which lead to carbon emissions do not occur and Japan reports this category as “NO”.

f) Land converted from other land-use categories: Above-ground biomass, Below-ground biomass

● Methodology

For RV activities, land conversion occurs due to the establishment or building of “facilities” and all living biomass is basically replaced in one year (In the case of urban parks converted from cropland, new planting in urban parks is carried out after removal of trees in cropland).

In Japan’s basic estimation principles for land converted to RV land, the facilities established newly by land conversion in the reporting year are defined as “Land converted to RV land”. The estimation methods are shown below.

$$\Delta C_{RVLUC} = \sum_i \left\{ A_i \times (C_{AfterLBi} - C_{BeforeLBi}) + (\Delta C_{RVLUCGi} - \Delta C_{RVLUCLi}) \right\}$$

$$\Delta C_{RVLUCGi} = \Delta B_{RVGi}$$

$$\Delta B_{RVGi} = \sum_j (NT_{i,j} \times C_{Ratei,j})$$

ΔC_{RVLUC}	: Annual carbon stock changes in living biomass in converted RV land [t-C/yr]
A	: Area of converted RV land [ha/yr]
$C_{AfterLB}$: Carbon stocks in living biomass immediately following land conversion [t-C/ha]
$C_{BeforeLB}$: Carbon stocks in living biomass immediately before land conversion [t-C/ha]
ΔC_{RVLUCG}	: Annual carbon stock changes due to living biomass growth in converted RV land [t-C/yr]
ΔC_{RVLUCL}	: Annual carbon stock changes due to living biomass loss in converted RV land [t-C/yr]
ΔB_{RVG}	: Annual biomass growth in RV land [t-C/yr]
C_{Rates}	: Annual living biomass growth per tree [t-C/tree/yr]
NT	: Number of trees
i	: Type of urban green facilities (Urban parks, Green areas on roads, Green areas at ports, Green areas around sewage treatment facilities, Green areas by greenery promoting systems for private green space, Green areas along rivers and erosion control sites, Green areas around public rental housing and Green areas around government buildings)
j	: Tree species

● **Parameters**

➤ **Urban parks**

The carbon stocks in living biomass immediately before conversion [t-C/ha] are the same as those for Grassland, Cropland, Wetlands and Other land. The carbon stocks in living biomass immediately following conversion are assumed to be zero (When urban parks classified as RV land were established, planting activities occurred and living biomass was stocked. Japan assumes that these biomass stocks are zero because they were carried from other fields and they have not grown by the RV activities). In addition, it is assumed that living biomass before conversion is emitted due to the establishment of RV land. The other parameters are assumed to be the same as the ones for “Remaining urban parks”.

➤ **Urban green facilities other than Urban parks**

The carbon stocks in living biomass immediately before and after conversion [t-C/ha] are the same as those for urban parks converted from other land uses.

The other parameters are assumed to be the same as the ones for “Remaining green area on roads”, “Remaining green area at ports”, “Remaining green area around sewage treatment facilities”, “Remaining green area along rivers and erosion control sites”, “Remaining green area around public rental housing” and “Remaining green area around government buildings”.

● **Activity data**

➤ **Urban parks**

The area of land converted to urban parks is calculated by multiplying the area of urban parks by the area ratio of land conversion for the whole country. The activity data for living biomass (the number of tall trees) is estimated in the same manner as for “Remaining urban parks”.

Table 11-28 Area of "Urban parks" and activity data (remaining land / converted land)

At the end of 2010

	Land-use category before conversion	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Urban parks which have been notified since 1st January 1990 and whose establishment area is 500 m ² or more	Remaining land	99.76%	50,646.96	11,891,495
	Cropland	0.20%	102.60	24,089
	Grassland	0.04%	18.58	4,363
	Wetlands	IE	IE	IE
	Other land	IE	IE	IE
	Total	100.00%	50,768.14	11,919,947

➤ **Green areas on roads**

The area of land converted to green area on roads is calculated by multiplying the area of green areas on roads by the area ratio of land conversion for the whole country. The activity data for living biomass (the number of tall trees) is estimated in the same manner as for "Remaining green area on roads".

Table 11-29 Area of "Green areas on roads" and activity data for each land-use category

At the end of 2010

	Land-use category before conversion	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Green areas on roads which have been notified since 1st January 1990 and whose establishment area is 500 m ² or more	Remaining	99.76%	20,361.72	9,075,970
	Cropland	0.20%	41.25	18,386
	Grassland	0.04%	7.47	3,330
	Wetlands	IE	IE	IE
	Other land	IE	IE	IE
	Total	100.00%	20,410.44	9,097,686

➤ **Green areas at ports**

The area of land converted to green areas at ports is calculated by multiplying the service area of green areas at ports by the area ratio of land conversion for the whole country. The activity data for living biomass (the number of tall trees) is estimated in the same manner as for "Remaining green areas at ports".

Table 11-30 Area of "Green areas at ports" and activity data for each land-use category

At the end of 2010

Land-use category before conversion	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Remaining land	99.76%	1,578.48	359,164
Cropland	0.20%	3.20	728
Grassland	0.04%	0.58	132
Wetlands	IE	IE	IE
Other land	IE	IE	IE
Total	100.00%	1,582.26	360,024

➤ **Green areas around sewage treatment facilities**

The area of land converted to green areas around sewage treatment facilities is calculated by multiplying the green areas around sewage treatment facilities by the area ratio of land conversion for the whole country. The activity data for living biomass (the number of tall trees) is estimated in the same manner as for “Remaining green area around sewage treatment facilities”.

Table 11-31 Area of “Green areas around sewage treatment facilities” and activity data for each land-use category

At the end of 2010

Land-use category before conversion	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Remaining land	99.76%	652.54	264,513
Cropland	0.20%	1.32	536
Grassland	0.04%	0.24	97
Wetlands	IE	IE	IE
Other land	IE	IE	IE
Total	100.00%	654.10	265,146

➤ **Green areas along rivers and erosion control sites**

The area of land converted to green areas along rivers and erosion control sites is calculated by multiplying the planted land area by the area ratio of land conversion for the whole country. The activity data for living biomass (the number of tall trees) is estimated in the same manner as for “Remaining Green area along rivers and erosion control sites”.

Table 11-32 Area of “Green areas along rivers and erosion control sites” and activity data for each land-use category

At the end of 2010

Land-use category before conversion	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Remaining land	99.76%	1,553.24	891,440
Cropland	0.20%	3.15	1,806
Grassland	0.04%	0.57	327
Wetlands	IE	IE	IE
Other land	IE	IE	IE
Total	100.00%	1,556.96	893,573

➤ **Green areas around government buildings**

The area of land converted to green areas around government buildings is calculated by multiplying the “total land area – building area” by the area ratio of land conversion for the whole country. The activity data for living biomass (the number of tall trees) is estimated in the same manner as for “Remaining green area around government buildings”.

Table 11-33 Area of “Green areas around government buildings” and activity data for each land-use category

At the end of 2010

Land-use category before conversion	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Remaining land	99.76%	281.04	30,577
Cropland	0.20%	0.57	62
Grassland	0.04%	0.10	11
Wetlands	IE	IE	IE
Other land	IE	IE	IE
Total	100.00%	281.71	30,650

➤ **Green areas around public rental housing**

The area of land converted to green areas around public rental housing is calculated by multiplying the “total land area – building area” by the area ratio of land conversion for the whole country. The activity data for living biomass (the number of tall trees) is estimated in the same manner as for “Remaining green area around public rental housing”.

Table 11-34 Area of “Green areas around public rental housing” and activity data for each land-use category

At the end of 2010

Land-use category before conversion	Area ratio of land which has been converted in the current year	Area [ha]	Activity data [Number of tall trees]
Remaining land	99.76%	2,177.14	478,753
Cropland	0.20%	4.41	970
Grassland	0.04%	0.80	176
Wetlands	IE	IE	IE
Other land	IE	IE	IE
Total	100.00%	2,182.35	479,899

g) Land converted from other land use categories: Dead wood

When a RV activity following land-use conversion is implemented, dead wood is removed off-site and supplemental planting is implemented before conversion because almost all of such lands are managed and trees are assumed to be “property”. Therefore, dead wood is not left on the ground immediately before land-use conversion. The carbon stocks in dead wood immediately after conversion are assumed to be zero, the same as living biomass. Therefore, the carbon stocks in dead wood before and after conversion are assumed to be zero.

The carbon stocks in dead wood accumulated for a year after conversion are reported as “IE”, the same as for “Remaining land”.

h) Land converted from other land-use categories: Litter

Likewise the “Remaining land”, Japan estimates carbon stock changes in litter in urban parks and green areas at ports only. The other urban green facilities (Green areas on roads, Green areas around

sewage treatment facilities, Green areas along rivers and erosion control sites, Green areas around public rental housing and Green areas around government buildings) are not the subject of estimation and are not reported (NR), since these facilities are not net sources.

● Methodology

$$\Delta C_{LUCRVLit} = \sum_i \{A_i \times (C_{AfterLiti} - C_{BeforeLiti}) + A_i \times Lit_i\}$$

$\Delta C_{LUCRVLit}$: Annual carbon stock changes in litter in land converted to RV land [t-C/yr]

$C_{AfterLit}$: Carbon stocks in litter immediately following land conversion [t-C/ha]

$C_{BeforeLit}$: Carbon stocks in litter immediately before land conversion [t-C/ha]

A : Area of converted RV land [ha/yr]

Lit : Annual carbon stock changes in litter per area of RV land [t-C/ha/yr]

i : Types of urban green facilities (Urban parks and Green areas at ports)

● Parameters

➤ Urban parks and Green areas at ports

When urban parks are converted from cropland, grassland or wetlands, soils before conversion are not moved to off-site and in general, these soils are used continuously after conversion or covered by additional soils. Therefore, litters and dead roots accumulated before conversion do not decrease due to the land-use conversion.

In addition, litter in urban parks immediately following conversion is very little, because the parks are newly planted. Therefore, carbon stock changes in litter due to land conversion are assumed to be zero.

The amount of carbon in litter accumulated for a year after conversion is estimated in the same manner as for “Remaining urban parks”.

➤ Urban green facilities other than Urban parks and Green areas at ports

The carbon stock changes in litter due to land-use conversion are assumed to be zero for the same reasons as for urban parks.

The amount of carbon in litter accumulated in the year after conversion is not estimated likewise the “Remaining green area on roads”, “Remaining green area around sewage treatment facilities”, “Remaining green area along rivers and erosion control sites”, “Remaining green area around public rental housings” and “Remaining green area around government buildings”.

For the above mentioned reasons, these urban green facilities are not reported (NR), even though these facilities are sinks.

● Activity data

The activity data is the same as for living biomass.

i) Land converted from other land-use categories: Soils

Likewise the “Remaining land”, urban parks and green areas at ports, whose management practices are similar to those in urban parks, are the only subject of estimation.

● Methodology

$$\Delta C_{LUCRVSoils} = \sum_i (\Delta C_{LUCMinerali} - L_{LUCOrganici})$$

$$\Delta C_{LUCMinerali} = \Delta A_i \times (C_{AfterSoil} - C_{BeforeSoil}) + A_i \times \Delta C_{soil,i}$$

$\Delta C_{LUCRVSoils}$: Annual carbon stock changes in soils in RV land following land-use conversion [t-C/ha]
$\Delta C_{LUCMineral}$: Annual carbon stock changes in mineral soils in RV land following land conversion [t-C/ha]
$L_{LUCOrganic}$: Annual carbon stock changes in organic soils in RV land following land conversion (=0) [t-C/ha]
ΔA	: Area of land converted to RV land within a year [ha/yr]
A	: Area of land converted to RV land [ha]
$C_{AfterSoil}$: Soil carbon stocks immediately after land-use conversion [t-C/ha]
$C_{BeforeSoil}$: Soil carbon stocks before land-use conversion [t-C/ha]
ΔC_{Soil}	: Annual carbon stock changes in soils per RV land area [t-C/ha/yr]
i	: Types of urban green facilities (Urban parks and Green areas at ports)

● Parameters

➤ *Urban parks and Green areas at ports*

As mentioned in the section for litter, when urban parks are converted from cropland, grassland or wetlands, soils before conversion are almost never moved to off-site (even if moved to off-site, carbon in these soils are not emitted due to combustion). In general, these soils are used after conversion continuously or covered by additional soils.

Therefore, soil carbon stocks do not change due to land-use conversion (the carbon stocks may increase due to additional soils. However, Japan assumes that soil carbon stocks do not change because additional soils do not lead to carbon sequestration from the atmosphere).

Carbon stock changes in soils within a year after conversion is estimated in the same manner as for the remaining Urban parks and Green areas at ports.

➤ *Urban green areas other than Urban parks and Green areas at ports*

The urban green facilities other than Urban parks and Green areas at ports are not sources of GHGs and are not reported (NR) in this submission because of the same reasons as for the “Land converted to urban parks”.

● Activity data

The area is as used for living biomass.

j) Land converted from other land-use categories: Other gases

1) Direct N₂O emissions from N fertilization

It is assumed that the volume of nitrogen-based fertilizer applied to urban parks is included in the demand for nitrogen-based fertilizers in the Agriculture sector, although fertilization in urban parks has been conducted in Japan. Therefore, these sources have been reported as “IE”.

2) Carbon emissions from lime application

Carbon emissions from lime application are estimated based on methodologies described in “Remaining land: Other gases” for all RV land together because the estimation method is similar regardless of remaining or converted land.

3) Biomass burning

As in the case of “Remaining RV land”, biomass burning activities which release carbon do not occur. Therefore, this category has been reported as “NO”.

k) Results

Table 11-35 Emissions and removals from RV activity

	1990	2008	2009	2010
	[Gg-CO ₂]	[Gg-CO ₂]	[Gg-CO ₂]	[Gg-CO ₂]
RV	-77.78	-1,081.76	-1,112.34	-1,130.14
Above-ground biomass	-47.30	-675.97	-694.81	-704.59
Below-ground biomass	-12.30	-175.75	-180.65	-183.19
Dead wood	IE	IE	IE	IE
Litter	-0.92	-11.42	-11.77	-12.04
Soils	-17.27	-218.63	-225.14	-230.34
Other gases	0.002	0.02	0.02	0.02

* CO₂ +: Emissions, -: Removals

11.4.1.2. Justification when omitting any carbon pool or GHG emissions/removals from activities under Article 3.3 and elected activities under Article 3.4

Some carbon pools under RV activities (litter and soils: Green areas on roads, Green areas around sewage treatment facilities, Green areas along river and erosion control sites, Green areas around public rental housing and Green areas around government buildings) are not included in the reporting. Some intermediate results of the ongoing research project relating to RV land by the MLIT show a clear tendency that those carbon pools have been increasing although more research and analysis are necessary to quantify carbon stock changes in these carbon pools (Handa et al., 2008). This does not lead to over-estimation of removals because these carbon pools are not sources of GHGs, although further information and data are needed for estimating carbon stock changes in these carbon pools.

11.4.1.3. Information on whether or not indirect and natural GHG emissions and removals have been factored out

Japan does not factor out indirect, natural and pre-1990 effects specified in paragraph 7 in the Annex to decision 15/CMP.1 in estimating emissions/removals from activities under Articles 3.3 and 3.4.

11.4.1.4. Changes in data and methods since the previous submission (recalculations)

■ New accounting of carbon stock changes in soils in RV land

Carbon stock changes in soils in RV land were earlier reported as zero, because this pool was not regarded as an emission source. However, these changes have been newly estimated for urban parks and green areas at ports, whose management practices are similar to those of urban parks, by calculating soil carbon stock changes per area. The soil carbon stock changes per area were calculated for urban parks, for which new information was available, by applying the difference method, which takes differences in soil carbon stocks between planted areas or lawn areas and bare areas in urban parks into account.

■ Improvement of the estimation accuracy of living biomass and litter in the RV accounting

The accuracy of estimation of living biomass in the RV land was improved by 1) refining the data of tall trees per area through re-examination of the background data and by covering a larger number

of sample facilities and 2) updating the annual growth rate of living biomass per tree, incorporating new data of the annual living biomass growth rate by tree type obtained for some tree species. By the same token, the accuracy of estimation for litter was improved by updating the annual carbon stock changes in litter.

■ Revision of the methodology for classifying RV land

A part of RV land, which used to be classified as wetlands under the Convention, was reclassified as settlements by taking into account the actual land-use status. In addition, the estimation methodology with data of land-use conversion from Forest land to Settlements since 1990, which is the beginning of activities under the Kyoto Protocol, was adopted in order to estimate and exclude from RV land the units of land otherwise subject to elected activities under Article 3.4.

■ Removals of living-biomass and litter at the lands subject to D activities

Due to the revision of methodology for estimating the units of land otherwise subject to elected activities under Article 3.4, removals by RV activities at the land were recalculated.

■ Correction of reporting on “Gains” of below-ground biomass of D activities in the CRF table

In the CRF table submitted last year, “Gains” for below-ground living biomass of D activities were reported, though “NO”, as they were reported altogether in the “Gains” column for above-ground biomass. In this submission, the values were reported for above- and below-ground biomass respectively, since it was possible to separate these values.

■ Uncertainty assessment

Due to the reassessment of data and parameters used for the uncertainty assessment, the uncertainty estimates were recalculated.

11.4.1.5. Uncertainty estimates

As a result of uncertainty assessment implemented by the method provided in Annex 7, “7.1 Methodology of Uncertainty Assessment”, the uncertainty of the total emissions/removals from activities under Articles 3.3 and 3.4 in 2010 has been assessed at 12%.

Table 11-36 Uncertainty of emissions and removals from activities under Articles 3.3 and 3.4

Greenhouse gas source and sink activities	GHGs	Emissions/Removals [Gg CO ₂ eq.]		Emissions/Removals Uncertainty [%]	Rank	Emissions/Removals Uncertainty as % of total national emissions [%]	Rank
			%				
Article 3.3 activities Afforestation and Reforestation	CO ₂ , N ₂ O, CH ₄	-426	-1%	36%	1	0%	3
Article 3.3 activities Deforestation	CO ₂ , N ₂ O, CH ₄	4,823	10%	26%	2	-2%	4
Article 3.4 activities Forest management	CO ₂ , N ₂ O, CH ₄	-53,252	-107%	11%	4	12%	1
Article 3.4 activities Revegetation	CO ₂ , N ₂ O, CH ₄	-1,130	-2%	17%	3	0%	2
Total		-49,985	-100%	12%			

11.4.1.5.a. Afforestation/Reforestation

The uncertainty of emissions/removals from AR activities in 2010 has been assessed at 36%.

Table 11-37 Uncertainty of emissions and removals from afforestation and reforestation activities

Greenhouse gas source and sink activities		GHGs	Emissions/ Removals [Gg CO ₂ eq.]	AD Uncertainty [%]	EF/RF Uncertainty [%]	Combined Uncertainty [%]	Rank	Combined Uncertainty as % of total national emissions [%]	Rank	
Article 3.3 activities	Change in carbon pool reported									
	Afforestation and Reforestation	Above-ground biomass	CO ₂	-246	43%	9%	44%	4	32%	1
Below-ground biomass		CO ₂	-64	IE	IE	IE	-	IE	-	
Litter		CO ₂	-30	-	-	25%	5	2%	3	
Dead wood		CO ₂	-74	-	-	97%	1	17%	2	
Soil		CO ₂	-13	-	-	25%	6	1%	4	
Afforestation and Reforestation	Greenhouse gas sources reported									
	Fertilization	N ₂ O	IE	-	-	-	-	-	-	
	Drainage of soils under forest management	N ₂ O	-	-	-	-	-	-	-	
	Disturbance associated with land-use conversion to croplands	N ₂ O	-	-	-	-	-	-	-	
	Liming	CO ₂	NO	NO	NO	NO	-	-	-	
	Biomass burning	CO ₂	IE	IE	IE	IE	-	-	-	
		CH ₄	0.003	-	-	-	59%	3	0%	6
		N ₂ O	0.0003	-	-	-	61%	2	0%	5
	Total			-426			36%			

11.4.1.5.b. Deforestation

The uncertainty of emissions/removals from D activities in 2010 has been assessed at 26%.

Table 11-38 Uncertainty of emissions and removals from deforestation activities

Greenhouse gas source and sink activities		GHGs	Emissions/ Removals [Gg CO ₂ eq.]	AD Uncertainty [%]	EF/RF Uncertainty [%]	Combined Uncertainty [%]	Rank	Combined Uncertainty as % of total national emissions [%]	Rank
Article 3.3 activities	Change in carbon pool reported								
	Dforestation	Above-ground biomass	CO ₂	2,661	26%	9%	27%	4	19%
Below-ground biomass		CO ₂	681	IE	IE	IE	-	IE	-
Litter		CO ₂	348	-	-	25%	5	2%	3
Dead wood		CO ₂	864	-	-	97%	1	17%	2
Soil		CO ₂	264	-	-	25%	6	1%	4
Dforestation	Greenhouse gas sources reported								
	Fertilization	N ₂ O	-	-	-	-	-	-	-
	Drainage of soils under forest management	N ₂ O	-	-	-	-	-	-	-
	Disturbance associated with land-use conversion to croplands	N ₂ O	3	-	-	86%	2	0%	5
	Liming	CO ₂	2	-	-	58%	3	0%	6
	Biomass burning	CO ₂	NO	NO	NO	NO	-	-	-
		CH ₄	NO	NO	NO	NO	-	-	-
		N ₂ O	NO	NO	NO	NO	-	-	-
Total			4,823			26%			

11.4.1.5.c. Forest Management

The uncertainty of emissions/removals from FM activities in 2010 has been assessed at 11%.

Table 11-39 Uncertainty of emissions/removals from forest management activities

Greenhouse gas source and sink activities		GHGs	Emissions/ Removals [Gg CO ₂ eq.]	AD Uncertainty [%]	EF/RF Uncertainty [%]	Combined Uncertainty [%]	Rank	Combined Uncertainty as % of total national emissions [%]	Rank
Article 3.4 activities	Change in carbon pool reported								
	Forest manafement	Above-ground biomass	CO ₂	-41,795	6%	9%	11%	6	11%
Below-ground biomass		CO ₂	-10,549	IE	IE	IE	-	IE	-
Litter		CO ₂	-356	-	-	25%	4	0%	3
Dead wood		CO ₂	1,051	-	-	97%	1	-2%	6
Soil		CO ₂	-1,603	-	-	25%	5	1%	2
Greenhouse gas sources reported									
Forest manafement	Fertilization	N ₂ O	IE	IE	IE	IE	-	-	-
	Drainage of soils under forest management	N ₂ O	NO	NO	NO	NO	-	-	-
	Disturbance associated with land-use conversion to croplands	N ₂ O	-	-	-	-	-	-	-
	Liming	CO ₂	NO	NO	NO	NO	-	-	-
	Biomass burning	CO ₂	IE	IE	IE	IE	-	-	-
		CH ₄	1	-	-	40%	3	0%	5
		N ₂ O	0.1	-	-	43%	2	0%	4
Total			-53,252			11%			

11.4.1.5.d. Revegetation

The uncertainty of emissions/removals from RV activities in 2010 has been assessed at 17%.

Table 11-40 Uncertainty of emissions/removals from revegetation activities

Greenhouse gas source and sink activities		GHGs	Emissions/ Removals [Gg CO ₂ eq.]	AD Uncertainty [%]	EF/RF Uncertainty [%]	Combined Uncertainty [%]	Rank	Combined Uncertainty as % of total national emissions [%]	Rank
Article 3.4 activities	Change in carbon pool reported								
	Revegetation	Above-ground biomass	CO ₂	-705	-	-	20%	4	15%
Below-ground biomass		CO ₂	-183	IE	IE	IE	-	IE	-
Litter		CO ₂	-12	-	-	116%	1	1%	3
Dead wood		CO ₂	IE	IE	IE	IE	-	-	-
Soil		CO ₂	-230	-	-	38%	3	8%	2
Greenhouse gas sources reported									
Revegetation	Fertilization	N ₂ O	IE	IE	IE	IE	-	-	-
	Drainage of soils under forest management	N ₂ O	-	-	-	-	-	-	-
	Disturbance associated with land-use conversion to croplands	N ₂ O	-	-	-	-	-	-	-
	Liming	CO ₂	0.02	-	-	51%	2	0%	4
	Biomass burning	CO ₂	NO	NO	NO	NO	-	-	-
		CH ₄	NO	NO	NO	NO	-	-	-
		N ₂ O	NO	NO	NO	NO	-	-	-
Total			-1,130			17%			

11.4.1.6. Information on other methodological issues (methods dealing with the effects of natural disturbances¹⁹)

11.4.1.6.a. Afforestation/Reforestation and Deforestation

The effects of natural disturbances have been reflected in forest resources data when Forest Registers are updated every 5 years in each planning area.

¹⁹ Including fires, windstorms, insects, droughts, flooding and ice storms, etc.

11.4.1.6.b. Forest Management

The effects of natural disturbances have been reflected in forest resources data when Forest Registers are updated every 5 years in each planning area.

11.4.1.6.c. Revegetation

It is considered that windstorms, floods and insects are natural disturbances which have a considerable impact on carbon stock changes in RV land. However, all land classified as RV is under human-induced management by administration etc. In addition, when tall trees disappear and outflow of soils occur in RV land located in settlements, the business budget is often appropriated and urgent restoration measures are administered from the viewpoint of safety and view.

Consequently, the effects of natural disturbances are not considered in the estimation because it looks that carbon stocks do not change. Furthermore, carbon stock change due to post-disaster restoration practices which are not implemented in the year when the disaster occurred does not lead to double-counting because it is not considered in this reporting.

11.4.1.7. The year of the onset of an activity, if after 2008

In this submission, all units of land and lands which start to be subject to activities under Article 3.3 or elected activities under Article 3.4 until 2010 are reported. The emissions and removals from the units of land and the lands which start to be subject to the activities in 2010 for the first time are included in the calculation of emissions and removals in neither 2008 nor 2009. Likewise, the emissions and removals from the units of land and the lands which start to be subject to the activities in 2009 for the first time are not included in the calculation of emissions and removals in 2008. The areas of such lands are shown below.

Table 11-41 Afforestation/Reforestation, Deforestation and Forest Management

Area of activities	Afforestation/ Reforestation [kha]	Deforestation [kha]	Forest Management [kha]		
			Ikusei-rin forest	Tennensei- rin forest	Total
FY1990~2010	28.7	322.7	7,882	7,228	15,111
(FY2010)	—	13.3	—	—	—

Table 11-42 Revegetation

Categories	Urban parks [ha]	Green areas on roads [ha]	Green areas at ports [ha]	Green areas around sewage treatment facilities [ha]	Green areas by greenery promoting systems for private green space [ha]
FY1990	3,730	1,621	195	48	0
FY1990-FY2010	50,768	20,410	1,582	654	5
(FY2010)	1,144	-127	39	11	0
Categories	Green areas along rivers and erosion control sites [ha]	Green areas around government buildings [ha]	Green areas around public rental housing [ha]	Total [ha]	
FY1990	58	12	199	5,862	
FY1990-FY2010	1,557	282	2,182	77,441	
(FY2010)	123	4	13	1,207	

* Areas for green areas on roads subject to RV decreased in FY2010 compared to the areas in FY1990-FY2009.

11.5. Article 3.3

11.5.1. Information that demonstrates that activities under Article 3.3 began on or after 1 January 1990 and before 31 December 2012 and are direct human-induced

Japan identifies AR and D by detecting a change of the forest cover which has occurred since 1 January 1990 using orthophotos taken at the end of 1989 and recent satellite images. In doing so, AR and forest restoration through natural succession are distinguished through imagery interpretation whether each forest cover change is human-induced or not. Whether land conversion is human-induced or not is judged by the imagery interpretation from the condition that whether any signs of human activity such as uniform tree species and uniform tree height, artificial forestation blocks, or work roads for forestation are observed or not.

11.5.2. Information on how harvesting or forest disturbance that is followed by the re-establishment of forest is distinguished from deforestation

In Japan, land conversion from forest land to other land use means exclusion of the land from forest plans. Therefore, as far as the area of harvested forest would remain included in forest plans, the area would be considered to be subject not to D but to temporary loss of biomass stock, and in Forest Registers would be distinguished from D which means conversion to other land use.

Japan identifies forest cover change as D only in the case when land form transformation or artificial construction are observed or obvious conversion to non-forest land such as cropland are detected through imagery interpretation using aerial photos and satellite images. By this methodology, D is distinguished from temporary loss of biomass stock in forest land such as clearcut under ongoing forestry activities.

Sample field surveys are conducted at plots which are interpreted as D areas in several prefectures every year, and accuracy of D interpretation is approximately 70% on average.

11.5.3. Information on the size and geographical location of forest areas that have lost forest cover but which are not yet classified as deforested

The total area of forest land that has temporarily lost forest cover due to harvesting or disturbance and which are not classified as deforested but as “Forest with less standing trees” (cut-over forests, lesser stocked forests) in Forest Registers was about 1.16 million ha in 2010.

11.5.4. Information on emissions and removals of greenhouse gases from lands harvested during the first commitment period following afforestation and reforestation

Japan assumes that all AR units of land have not been harvested during the first commitment period. Therefore, paragraph 4 of the Annex to decision 16/CMP.1 is not applied to.

11.6. Article 3.4

11.6.1. Information that demonstrates that activities under Article 3.4 have occurred since 1 January 1990 and are human-induced

11.6.1.1. Forest Management

The status of FM activities since 1 January 1990 has been investigated since FY2007 by sample surveys including field surveys, interviews with forest owners' associations and detection of

administrative information on subsidies for forest practices, of Ikusei-rin forests throughout the country. The results of the survey have been used to estimate the FM ratio.

11.6.1.2. Revegetation

Japan demonstrates that RV activities have occurred since 1990 and are human induced based on the following reasons.

Table 11-43 Information that demonstrates that Revegetation activities have occurred since 1st January 1990 and are human induced

Urban green facilities	Information that demonstrates that Revegetation activities have occurred since 1 st January 1990 and are human induced
Urban parks	<p><u>Extraction of activities which have occurred since 1st January 1990</u> MLIT has implemented the “Urban Parks Status Survey” and has collected data on the notification year of urban parks. In the reporting, only urban parks which have been notified since 1st January 1990 are included. Although some urban parks were established before the notification year, Japan considers that RV activities have occurred since the notification year under the “Urban Park Act”.</p> <p><u>Demonstration that activities are human induced</u> Activity data (the number of tall trees) is calculated based on the number of tall trees per land area (tree/ha) which is developed by using data on planted tall trees. Its calculation procedure ensures that Japan extracts human-induced activities.</p>
Green areas on roads	<p><u>Extraction of activities which have occurred since 1st January 1990</u> MLIT has implemented the “Road Tree Planting Status Survey” every 5 years (implemented every year since 2007) and has collected data on the number of planted tall trees. Activity data after 1990 is calculated by extrapolating or interpolating these data.</p> <p><u>Demonstration that activities are human induced</u> In the “Road Tree Planting Status Survey”, only planted tall trees have been measured. Their measurement procedure ensures that Japan extracts human induced activities.</p>
Green areas at ports	<p><u>Extraction of activities which have occurred since 1st January 1990</u> MLIT has implemented complete census since 2006 and has collected relevant data (year of establishment and service area) for green areas at ports which had been established since 1990.</p> <p><u>Demonstration that activities are human induced</u> Activity data (the number of tall trees) is calculated by using parameters of urban parks which are based on human-induced activities data.</p>
Green areas around sewage treatment facilities	<p><u>Extraction of activities which have occurred since 1st January 1990</u> MLIT has implemented the “Sewage treatment Facility Status Survey” since 2006 and has collected relevant data (year of establishment and greening area) for green areas around sewage treatment facilities which had been established since 1990.</p> <p><u>Demonstration that activities are human induced</u> Activity data (the number of tall trees) is calculated based on the number of tall trees per land area (tree/ha) which is developed by using data on planted tall trees. Its calculation procedure ensures that Japan extracts human-induced activities.</p>
Green areas by greenery promoting systems for private green space	<p><u>Extraction of activities which have occurred since 1st January 1990</u> It is clear that all green areas by greenery promoting systems for private green space have been established since 1st January 1990 because greenery promoting systems have been implemented since 2001. Existing tall trees before 1990 in some green areas are reported when it have been notified by the local authority mayor. They are excluded from RV land area.</p> <p><u>Demonstration that activities are human induced</u> All green areas by greenery promoting systems for private green space have been established by human-induced activities.</p>
Green areas along rivers and erosion control sites	<p><u>Extraction of activities which have occurred since 1st January 1990</u> MLIT has implemented the “Survey on carbon dioxide absorption at source in river works” since 2007 and has collected relevant data (name, location, year of establishment, planted land area [projected area] and the number of tall trees) for river works and erosion and sediment control works which had been implemented since 1990.</p> <p><u>Demonstration that activities are human induced</u> Activity data (the number of tall trees) is calculated based on the number of tall trees per land area (tree/ha) which is developed by using data on planted tall trees. Its calculation procedure</p>

Urban green facilities	Information that demonstrates that Revegetation activities have occurred since 1 st January 1990 and are human induced
	ensures that Japan extracts human-induced activities.
Green areas around government buildings	<u>Extraction of activities which have occurred since 1st January 1990</u> MLIT has implemented complete census since 2007 and has collected relevant data (name, location, year of establishment, total land area and building area) for government buildings which had been established since 1990. <u>Demonstration that activities are human induced</u> Activity data (the number of tall trees) is calculated based on the number of tall trees per land area (tree/ha) which is developed by using data on planted tall trees. Its calculation procedure ensures that Japan extracts human-induced activities.
Green areas around public rental housing	<u>Extraction of activities which have occurred since 1st January 1990</u> MLIT has implemented the “Progress survey on tree planting for public rental housing” since 2007 and has collected relevant data (name, location, year of establishment, total land area and building area) for public rental housing which had been established since 1990. <u>Demonstration that activities are human induced</u> Activity data (the number of tall trees) is calculated based on the number of tall trees per land area (tree/ha) which is developed by using data on planted tall trees. Its calculation procedure ensures that Japan extracts human-induced activities.

11.6.2. Information relating to Revegetation for the base year and the commitment period

The anthropogenic greenhouse gas removals in “Revegetation” for the base year are those from RV area in 1990. The area where RV activity took place in 1990 is directly obtained by activity data in each subcategory of RV. The anthropogenic greenhouse gas removals in “Revegetation” for the commitment period are those from RV area in each year. Those removals are reported within the relevant geographical location. The data and the methodologies used are provided in sections 11.3.2.5. and 11.4.1.1.d.

11.6.3. Information that demonstrates the emissions and removals resulting from elected Article 3.4 activities are not accounted for under activities under Article 3.3 activities

11.6.3.1.a. Information on emissions and removals by FM activities are not accounted for under Article 3.3 activities

AR and D are of higher hierarchy than FM in the land classification system of Articles 3.3 and 3.4 in Japan. Emissions and removals by AR and D are estimated in the first step, then emissions and removals by FM are estimated by subtracting emissions and removals by AR from emissions and removals in managed forests as explained in section 11.3.2.2. (see Figure 11-1). Therefore, emissions and removals by FM could not be included in those by AR nor D.

11.6.3.1.b. Information on emissions and removals from RV activities are not accounted under Article 3.3 activities

RV land is defined as the land which is not included in AR land as described in the definition section 11.2.2.2. Therefore, emissions and removals from RV could not be included in those from AR theoretically.

The area of D land which would otherwise be included in RV lands is reported in the CRF Table 5(KP-I) A.2.1. Since this land is classified as D land and is not included in RV land, all emissions and removals from this land are reported under D activity as described in the explanation of methodologies of D in section 11.4.1.1.b and those of RV in section 11.4.1.1.d. Therefore, there is no double count between D and RV and emissions and removals from RV could not be included in those from D.

11.6.4. Information relating to Forest Management

11.6.4.1. The definition of forest for this category conforms with the definition in item 11.2 above

In Japan, the area and carbon stock change in land subject to FM activities are estimated by applying FM ratios to data of all forests which meet our country's forest definition. Therefore, the definition of land subject to FM activities is consistent with our country's forest definition.

On the other hand, not all managed forests reported under the Convention are subject to FM reported as Article 3.4 activity under the Kyoto Protocol in Japan, because FM forests consist of only the area where FM activities have been taken place since 1990 as described in section 11.3.2.4. .

11.6.4.2. The definition of forest management conforms with the definition in paragraph 1 (f) of the Annex to decision 16/CMP.1

Japan considers that FM activities which are reported under the Kyoto Protocol should be of sustainable system and whether this is fulfilled or not is judged from whether appropriate forest practices have been carried out in Ikusei-rin forests or whether practices for the protection or conservation of forests such as controlling logging activities and land-use change have been carried out by laws. Therefore, Japan's definition of FM is consistent with the definition provided in "Decision 16/CMP.1" (a system of practices for stewardship and use of forest land aimed at fulfilling relevant ecological, economic and social functions of the forest in a sustainable manner).

11.6.4.3. Information on the extent to which GHG removals by sinks offset the debit incurred under Article 3.3.

The total amount of FM removals that offset the debit incurred under Article 3.3 was 9,164 Gg-CO₂ eq. from 2008 until 2010. Related information is provided in Table 11-2.

11.7. Other information

11.7.1. Key category analysis for Article 3.3 activities and any elected activities under Article 3.4

In accordance with the GPG-LULUCF, Chapter 5, the activity which meets the following requirements is considered as a key category.

-The associated category under the UNFCCC is identified as key. In addition, emissions/removals from the activity are greater than the smallest category that is identified as key in the UNFCCC inventory (Tier 1 level assessment).

-The estimation method is changed from previous reporting.

● Corresponding key categories under the UNFCCC

Japan's LULUCF key categories under the UNFCCC for 2010 (Annex 1 of this report) are as follows;

5.A.1. Forest land remaining Forest land (CO₂)

5.A.2. Land converted to Forest land (CO₂)

5.B.2. Land converted to Cropland (CO₂)

5.E.2. Land converted to Settlements (CO₂)

In accordance with GPG-LULUCF, AR, D, FM and RV may be identified as key categories under the Kyoto Protocol.

Table 11-44 Relationship between UNFCCC categories and Kyoto Protocol activities

UNFCCC category under the Convention	Kyoto Protocol category
5.A.1. Forest land remaining Forest land	FM
5.A.2. Land converted to Forest land	AR
5.B.1. Cropland remaining Cropland	
5.B.2. Land converted to Cropland	D
5.C.1. Grassland remaining Grassland	
5.C.2. Land converted to Grassland	D
5.D.1. Wetlands remaining Wetlands	
5.D.2. Land converted to Wetlands	D
5.E.1. Settlements remaining Settlements	RV
5.E.2. Land converted to Settlements	D、RV
5.F.1. Other land remaining Other land	—
5.F.2. Land converted to Other land	D

The relationship between conventional categories and the Kyoto categories in this table is based on the GPG-LULUCF, p. 5.39, Table 5.4.4., and the definitions of Articles 3.3 and 3.4 activities in Japan. Shade indicates the key categories under the UNFCCC.

● *Comparison with the smallest key category under the UNFCCC*

The smallest category for the UNFCCC (Tier 1 level assessment) for 2010 was 2.A.3. Limestone and Dolomite Use (CO₂) [8,073 Gg-CO₂]. As a result of the comparison, only FM activity was greater than this category.

● *Qualitative considerations*

Since the removals by RV lands have been increasing since FY1990 and the estimation methodologies for RV have been improved in this submission, RV was regarded as a key category.

Therefore, AR, D, FM and RV activities (CO₂) are identified as key activities for 2010.

11.7.2. Further improvements

Methodological issues relating to Articles 3.3 and 3.4 are identified under the Committee for Greenhouse Gas Emissions Estimation Methods- Breakout Group on LULUCF. They are updated every year taking into account the progress of the inventory-related work and issues identified by the Expert Review Team. Many of the improvement plans on LULUCF reporting under the Convention described in Chapter 7 of this report are closely linked to activities under Articles 3.3 and 3.4 of the Kyoto Protocol. Therefore, both the reporting under the Convention and the reporting under the Kyoto Protocol are discussed together. Major issues to be improved are as follows:

- Improvement of methodology and data to estimate carbon stock change in soil due to land-use conversion which reflects changes in management practices more properly is under discussion in Japan.
- With regard to the annual growth rate of living biomass per tree in RV land, Japan plans to improve the accuracy when new country-specific data by tree species become available.
- Except for Urban parks and Green areas at ports, carbon stock change in soils is not included in the reporting because soils are not sources of GHGs under RV activities. Japan will continue to collect fundamental information on soil carbon and consider about estimation methods.

11.8. Information relating to Article 6

Japan has not carried out any projects under Article 6 of the Kyoto Protocol. Therefore, a special indication of whether the boundary of the geographical location encompasses land subject to the Article 6 project is not prepared.

11.9. Information on the reporting status of paragraphs 5 to 9 of the Annex to decision 15/CMP.1

The requirements for reporting about Articles 3.3 and 3.4 which are set out in paragraphs 5 to 9 of the annex to decision 15/CMP.1 are provided in sections shown in Table 11-45.

Table 11-45 List of reference sections for the requirements set in paragraphs 5 to 9 of the Annex to decision 15/CMP.1

Checklist for KP reporting (paragraphs 5-9 in the annex to decision 15/CMP.1)		Paragraph	Main sections of Chapter 11 providing relevant information
Information on how inventory methodologies have been applied taking into account GPG-LULUCF and decision 16/CMP.1		6 (a)	Detailed information is provided in each section
Information on the geographical location of the boundaries of areas that encompass:		6 (b)	11.3.3, 11.3.2
	Units of land subject to activities under Article 3.3	6 (b) (i)	11.3.3, 11.3.2
	Units of land subject to activities under Article 3.3, which would otherwise be included in land subject to elected activities under Article 3.4	6 (b) (ii)	11.3.3, 11.3.2 and CRF table 5(KP-D)A.2.1
	Land subject to elected activities under Article 3.4	6 (b) (iii)	11.3.3, 11.3.2
Information on the spatial assessment unit for determining the area of accounting for ARD		6 (c)	11.3.1
GHG emissions by sources and removals by sinks from LULUCF activities under Articles 3.3 and 3.4:			
	Emissions by sources and removals by sinks are clearly distinguished from emissions from Annex A sources.	5	11.4.1: Methodology
	Emissions by sources and removals by sinks are reported for all geographical locations reported in current and previous years	6 (d)	11.3.2.3, 11.3.2.4, 11.3.2.5
	Emissions/removals from Articles 3.3 or (elected) 3.4 activities are reported since the beginning of the commitment period or the onset of the activity	6 (d)	11.4.1.7
	Information on which pools (above-ground / below-ground biomass, litter, dead wood and soil organic carbon) were not accounted for.	6 (e)	11.4.1.2
	Information on whether Articles 3.3 and (elected) 3.4 emissions/removals factor out removals from (i) elevated CO ₂ concentrations above pre-industrial levels; (ii) indirect N deposition; and (iii) dynamic effects of age structure resulting from pre-1 January 1990 activities.	7	11.4.1.3
Specific information to be reported for Article 3.3 activities			
	Information that activities under Article 3.3 began on or after 1 January 1990 and before 31 December of the last year of the commitment period	8 (a)	11.5.1
	Information on how harvesting or forest disturbance that is followed by the re-establishment of forest is distinguished from deforestation	8 (b)	11.5.2, 11.5.3

Checklist for KP reporting (paragraphs 5-9 in the annex to decision 15/CMP.1)		Paragraph	Main sections of Chapter 11 providing relevant information
	Information on emissions/removals from lands harvested during the 1 st commitment period following AR on these units of land since 1990.	8 (c)	11.5.4
Specific information to be reported for Article 3.4 activities			
	Information that activities under Article 3.4 occurred since 1 January 1990 and are human induced.	9 (a)	11.6.2
	CM, GM, RV: emissions/removals reported for each year of the commitment period and for the base year for each of the elected activities on the geographical locations reported.	9 (b)	11.6.1, 11.3.2.5, 11.4.1.1.d
	Information that emissions/removals from Article 3.4 activities are not accounted for under activities under Article 3.3.	9 (c)	11.6.3
	FM: information on the extent to which GHG removals by sinks offsets the debit incurred under Article 3.3.	9 (d)	11.6.4.3

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21. Parks and Green Spaces Division of the Ministry of Land, Infrastructure and Transport, "FY2004 Survey on evaluation techniques for the effectiveness of greening in urban parks for preventing global warming", March, 2005.
22. National Institute for Land and Infrastructure Management, Ministry of Land, Infrastructure, Transport and Tourism, Japan, *Technical note of National Institute for Land and Infrastructure Management No.506 The Street tree of Japan VI*, 2009.

Chapter 12. Information on accounting of Kyoto units

In line with paragraph 10 of decision 15/CMP.1, Japan reports the information on holdings and transactions of Kyoto units (ERUs, CERs, ICERs, tCERs, AAUs and RMUs)¹. For the reporting, in accordance with paragraph 11 of annex of decision 15/CMP.1, Japan uses Standard electronic format (SEF) defined in the annex of decision 14/CMP.1. Apart from this NIR, SEF is submitted to the UNFCCC Secretariat with the file name “SEF_JP_2012_1_14-40-7 21-2-2012”.

12.1. Summary of information reported in the SEF tables

For information on Kyoto units in Japan’s National Registry, see the annex “Standard Electric Format for Reporting of Information on Kyoto Protocol Units” (SEF_JP_2012_1_14-40-7 21-2-2012) submitted on the basis of Decision 14/CMP.1.

12.2. Discrepancies and notifications

Regarding Japan’s national registry, discrepancies and notifications to be reported in accordance with paragraphs 12-17 of annex to decision 15/CMP.1 are as follows.

Table 12-1 Discrepancies and notifications

Reporting item	Description
Para. 12 of the annex to decision 15/CMP.1 Discrepancies	There were discrepancies to be reported. For detailed information on discrepant transaction, see the annex “annex 2-2 disc transaction list JP2011” .
Para. 13 of the annex to decision 15/CMP.1 Notification from Executive Board of CDM	There was no notification regarding ICERs to be replaced due to a reversal of storage.
Para. 14 of the annex to decision 15/CMP.1 Failure of certification	There was no notification regarding ICERs to be replaced due to non-submission of certification report.
Para. 15 of the annex to decision 15/CMP.1 List of non-replacements	There was no record of non-replacement identified by the transaction log.
Para. 16 of the annex to decision 15/CMP.1 Invalid Kyoto units	There were no units that are invalid for use towards compliance with commitments.
Para. 17 of the annex to decision 15/CMP.1 Discrepant transaction that needs actions to correct problem	There was no discrepant transaction that needs actions to correct problem.

12.3. Publicly accessible information

As presented in the section IV of Part 2 of “Report on Japan’s Assigned Amount”, a list of the information publicly accessible by means of the user interface to the national registry is as below:

- Account information as required by paragraph 45, annex, Decision 13/CMP.1
- Article 6 project information as required by paragraph 46, annex, Decision 13/CMP.1

¹ Kyoto units are: emission reduction units (ERU) from joint implementation (JI) projects, certified emission reductions (CERs) from clean development mechanism (CDM) projects, temporary certified emission reductions (tCERs) and long-term certified emission reductions (ICERs) from afforestation/reforestation CDM projects, assigned amount units

- Kyoto units information as required by paragraph 47, annex, Decision 13/CMP.1
- Legal entities as required by paragraph 48, annex, Decision 13/CMP.1

The information is provided in “Publicly Accessible Information” of Japan’s national registry website.

- URL of the Japan’s national registry system: http://www.registry.go.jp/index_e.html
- Publicly Accessible Information: http://www.registry.go.jp/public_info_en.html

The following information is not published due to confidentiality concerns:

- Unit holdings at an individual account level
- Identity of accounts to which Japan’s national registry transferred units and those from which it acquired units.

In addition, for better readability, information on units is not associated with their respective serial numbers.

12.4. Calculation of the commitment period reserve (CPR)

According to the paragraph 6 of Annex to Decision 11/CMP.1, each Party included in Annex I shall maintain, in its national registry, a commitment period reserve which should not drop below 90 per cent of the Party’s assigned amount calculated pursuant to Article 3, paragraphs 7 and 8 of the Kyoto Protocol, or 100 per cent of five times its most recently reviewed inventory, whichever is lowest.

Japan’s commitment period reserve is 5,335,431,899 t-CO₂ eq.², the same as the value reported in the previous submission.

12.5. KP-LULUCF accounting

As stated in the section II.3 of Part 2 of “Report on Japan’s Assigned Amount”, Japan will account for the credits issued by activities under Article 3, paragraphs 3 and 4 of the Kyoto Protocol for the entire commitment period.

(AAUs) and removal units (RMUs) from KP-LULUCF activities within Annex I Parties.

² See section I of part 2 of “Report on Japan’s Assigned Amount Pursuant to Article 3, Paragraphs 7 and 8 of the Kyoto Protocol – Under Article 7, paragraph 4 of the Kyoto Protocol, United Nations Framework convention on Climate Change (The Government of Japan, August 2006 (updated in March 2007))” for the calculation of CPR.

Chapter 13. Information on changes in national system

In line with paragraph 21 of decision 15/CMP.1, Japan reports the changes in its national system from the previous inventory submission.

- Japan's national system has not changed from the previous inventory submission.

Chapter 14. Information on changes in national registry

In line with paragraph 22 of decision 15/CMP.1, Japan reports changes in national registry of Japan from the previous inventory submission.

14.1. Summary of changes made on national registry of Japan in 2011

Table 14-1 Summary of changes made on national registry of Japan in 2011

Reporting Items	Descriptions of Changes
15/CMP.1, annex II, para 32. (a) Change of name or contact	No change
15/CMP.1, annex II, para 32. (b) Change of cooperation arrangement	No change
15/CMP.1, annex II, para 32. (c) Change to database or the capacity of national registry	No change
15/CMP.1, annex II, para 32. (d) Change of conformance to technical standards	No change
15/CMP.1, annex II, para 32. (e) Change of procedures to minimize discrepancies	No change
15/CMP.1, annex II, para 32. (f) Change of security measures	No change
15/CMP.1, annex II, para 32. (g) Change of a list of publicly accessible information	Information on unit holdings and transactions is made publicly available on the basis of Standard Electronic Format (SEF) to meet the requirement specified in decision 14/CMP.1. In April 2011, the information for 2010 was published. The following information is not published due to confidentiality concerns: - Unit holdings at an individual account level - Identity of accounts to which national registry of Japan transferred units and those from which it acquired units. In addition, for better readability, information on units is not associated with their respective serial numbers.
15/CMP.1, annex II, para 32. (h) Change of the internet address	No change
15/CMP.1, annex II, para 32. (i) Change of measures for ensuring data integrity	No change
15/CMP.1, annex II, para 32. (j) Change of test results	No change

14.2. Information relevant to the changes made on national registry of Japan

- In December 2010, some documents of DES were revised, in which a response code was added regarding a list of national registries. The revised documents and their impacts on national registry of Japan are described as follows:
 - The revised DES main text (version 1.1.8) was released. There is no change made on national registry of Japan in relation to the release.
 - The revised DES annex E (List of checks and Response Codes for Message Processing, version 1.1.10) was released. The response code was added in national registry of Japan as

well.

- Public information on the unit holdings and transactions conducted was updated in April 2011, on the basis of the SEF for 2010, for the purpose of meeting the requirement specified in decision 13/CMP.1. The following information, which is requested to be made publicly available in decision 13/CMP.1, has not been made so due mostly to confidentiality concerns (relevant paragraph numbers of the annex to decision 13/CMP.1 are indicated in parentheses):
 - The full name of the representative of the account holder (paragraph 45(e))
 - Serial numbers of ERUs, CERs, AAUs and RMUs those are subject of this public information (paragraph 47)
 - The total quantity of ERUs, CERs, AAUs and RMUs in each account at the beginning of the year (the total quantity is only available by account type) (paragraph 47(a))
 - The identity of the transferring accounts from which ERUs, CERs, AAUs and RMUs were acquired by national registry of Japan during the year (the identity of the transferring registries is available) (paragraph 47(d))
 - The identity of the acquiring accounts to which ERUs, CERs, AAUs and RMUs were transferred from national registry of Japan during the year (the identity of the acquiring registries is available) (paragraph 47(g))
 - Current holdings of ERUs, CERs, AAUs and RMUs in each account (the current holdings are only available by account type) (paragraph 47(l))
- In June 2011, the security patches were checked for the middleware and middleware was updated with the security patches after the investigation. There is no impact on the functions of the ITL and other national registries.
- In September 2011, the security patches were checked for the middleware and middleware was updated with the security patches after the investigation. There is no impact on the functions of the ITL and other national registries.
- In December 2011, the DNS server was updated with security patches after the investigation. There is no impact on the functions of the ITL and other national registries.

Chapter 15. Information on minimization of adverse impacts in accordance with Article 3, paragraph 14

In line with paragraphs 23-26 of decision 15/CMP.1, Japan reports the information on adverse impacts in accordance with Article 3, paragraph 14.

15.1. Overview

Japan takes actions, taking into account the importance to make effort to minimize adverse impacts in accordance with Article 3, paragraph 14. On the other hand, it should be noted that we have difficulty in assessing accurately specific adverse impacts due to the implementation of response measures to address climate change issues. For example, the fluctuation in price of crude oil is caused by balance between supply and demand and numerous other factors (e.g., trend in crude oil futures market or the economy), and it is uncertain whether there exists a causal link or if so what extent is from adverse impacts of climate change policy and measures.

In addition, it is necessary to change the perception of response measures in order to address climate change issues effectively, and sustainable development could be the one of the key options. For instance, the introduction of renewable energy lead to improve the energy access, prepare for a disaster and create employment through a new industry, as well as contributing to reducing GHG emissions, As discussed in COP17 and Rio+20, the transition to green economy and the attainment to low-carbon growth are the key elements of addressing climate change and achieving the sustainable development to make balance between environment and economy. Efforts toward the establishment of low-carbon society should be accelerated throughout the world. Promoting further measures to mitigation as well as to adaptation taking into account the needs of vulnerable countries lead to maximize the positive impacts of response measures.

15.2. Actions to minimize adverse impacts in accordance with Article 3, paragraph 14

Japan has given a priority to the efforts below, taking into consideration that these efforts are important to minimize adverse social, environmental and economic impacts on developing country Parties, particularly those identified in Article 4, paragraphs 8 and 9, of the Convention in implementing the commitments under Article 3, paragraph 1 of the Kyoto Protocol.

At the same time, it should be noted that it is impossible to evaluate these efforts since the method of evaluation is currently under international negotiation.

● *Technical assistance in the energy and environmental sectors*

Based on the Japan's Cooperation Initiative for Clean Energy and Sustainable Growth presented at the 2nd East Asia Summit in January 2007 and the agreement reached at Asian Ministerial Energy Roundtable held in April 2009, we provided the cooperation in human resource development through accepting trainees and dispatching experts in the area of energy conservation and renewable energy to countries in East Asia and Middle East. We assisted these countries in the establishment and implementation of legal systems of energy conservation and renewable energy. In addition, in a joint

policy studies among research institutions from Japan and countries like China and India, we compared country policies related to energy conservation that will benefit the host countries' policy making process and also estimated possibilities of energy use reductions of highly energy-consuming industries.

Additionally, technical assistance in the field of energy and environment by Japan has long been provided throughout the world, contributing to the sustainable economic growth of developing countries. Through Japan International Cooperation Agency (JICA), depending on the needs of developing countries, Japan has been providing assistance in human resource development such as dispatching experts and providing training programs in Japan.

● ***Assistance to oil producing countries in diversifying their economies***

In April 2009, the 3rd Asian Ministerial Energy Roundtable was held in Japan where we requested that regulatory agencies take more coordinated action to strengthen surveillance on commodity futures trading markets and enhance its transparency. Furthermore, parties have agreed to conduct specific projects such as formulation of a demand and supply projection, sharing of leading projects concerning energy conservation and renewable energy, and provision of training opportunities (e.g., Japan will accept 2000 trainees over 3 years).

● ***Development of carbon capture and storage (CCS) technologies***

Recognizing that CCS is an innovative technology that may achieve highly efficient carbon emissions reductions, Japan has been implementing a large-scale demonstration projects toward practical use of CCS by 2020, and researches and developments on cost reductions and safety improvements. Also, Japan actively exchanged information on CCS technologies with other countries such as the United States of America and European countries.

In terms of institutions regarding to the sub-seabed geological storage of CO₂ (offshore CCS), Japan amended the Marine Pollution Prevention Law in 2007 and build up the system of permission by the Minister of the Environment with the point of view of preserving the marine environment. It is examining the methods of the potential environmental impact assessment and monitoring technology.

Annex 1. Key Categories

A1.1. Outline of Key Category Analysis

The *UNFCCC Inventory Reporting Guidelines*¹ require the application of the *Good Practice Guidance (2000)*, and the key category analysis² given in the Guidance. The guidelines for national system under Article 5 of the Kyoto Protocol also require countries, in compiling their inventories, to follow the method given in Chapter 7 of the *GPG* and identify the key categories.

The key category analyses were done for both data of FY 2009 and of FY 1990, the base year for the UNFCCC³. Their results are presented here.

A1.2. Results of Key Category Analysis

A1.2.1. Key Categories

Key categories were assessed in accordance with the *GPG* assessment methods (Tier 1 level assessment, Tier 1 trend assessment, Tier 2 level assessment and Tier 2 trend assessment).

The key category for Land use, land use change and forestry (LULUCF) sector were assessed in accordance with *GPG-LULUCF*. The key categories were identified for the inventory excluding LULUCF first, and then the key category analysis was repeated for the full inventory including the LULUCF categories.

As a result, 39 and 35 sources and sinks were detected as the key source categories for FY 2010 and FY 1990, respectively (Table A1-1 and A1-2).

¹ Guidelines for the preparation of national communications by Parties included in Annex I to the Convention, Part I: UNFCCC reporting guidelines on annual inventories (following incorporation of the provisions of decision 14/CP.11) (FCCC/SBSTA/2006/9)

² The *IPCC Good Practice Guidance for Land Use, Land-Use Change and Forestry* (2003), which was welcomed in COP9, extends the key source analysis to LULUCF categories. In the UNFCCC reporting guidelines (FCCC/SBSTA/2004/8) and afterwards, the term “key source category” was revised to “key category”.

³ With respect to HFCs, PFCs, SF₆, the data used for this analysis were the FY 1995 values.

Table A1-1 Japan's Key Categories (FY2010)

	A IPCC Category		B Direct GHGs	L1	T1	L2	T2
#1	1A Stationary Combustion	Solid Fuels	CO2	#1	#2	#4	#7
#2	1A Stationary Combustion	Liquid Fuels	CO2	#2	#1	#10	#8
#3	1A Stationary Combustion	Gaseous Fuels	CO2	#3	#3		
#4	1A3 Mobile Combustion	b. Road Transportation	CO2	#4	#6	#6	#24
#5	5A Forest Land	1. Forest Land remaining Forest Land	CO2	#5		#2	
#6	2A Mineral Product	1. Cement Production	CO2	#6	#7	#11	#10
#7	2F(a) Consumption of Halocarbons and SF6	1. Refrigeration and Air Conditioning Equipment	HFCs	#7	#5	#3	#1
#8	1A Stationary Combustion	Other Fuels	CO2	#8	#11	#7	#11
#9	6C Waste Incineration		CO2	#9		#5	
#10	1A3 Mobile Combustion	d. Navigation	CO2	#10	#14		
#11	1A3 Mobile Combustion	a. Civil Aviation	CO2	#11			
#12	2A Mineral Product	3. Limestone and Dolomite Use	CO2	#12		#20	
#13	4A Enteric Fermentation		CH4			#25	
#14	2A Mineral Product	2. Lime Production	CO2			#23	
#15	4B Manure Management		N2O			#8	
#16	4C Rice Cultivation		CH4			#17	
#17	1A Stationary Combustion		N2O			#16	#19
#18	5E Settlements	2. Land converted to Settlements	CO2			#22	#22
#19	6A Solid Waste Disposal on Land		CH4		#13	#19	#9
#20	4D Agricultural Soils	1. Direct Soil Emissions	N2O			#9	#13
#21	4D Agricultural Soils	3. Indirect Emissions	N2O			#13	#16
#22	1A3 Mobile Combustion	b. Road Transportation	N2O			#14	#12
#23	4B Manure Management		CH4			#15	#18
#24	2F(a) Consumption of Halocarbons and SF6	7. Semiconductor Manufacture	PFCs			#18	#15
#25	6C Waste Incineration		N2O			#12	
#26	2F(a) Consumption of Halocarbons and SF6	5. Solvents	PFCs		#9		#4
#27	6B Wastewater Handling		CH4				#26
#28	6B Wastewater Handling		N2O			#21	
#29	2F(a) Consumption of Halocarbons and SF6	8. Electrical Equipment	SF6		#8		#2
#30	2B Chemical Industry	3. Adipic Acid	N2O		#10		#17
#31	5B Cropland	2. Land converted to Cropland	CO2				#20
#32	5A Forest Land	2. Land converted to Forest Land	CO2				#25
#33	2E Production of Halocarbons and SF6	2. Fugitive Emissions	PFCs				#21
#34	2E Production of Halocarbons and SF6	2. Fugitive Emissions	SF6		#12		#3
#35	1A3 Mobile Combustion	a. Civil Aviation	N2O			#1	#6
#36	1A3 Mobile Combustion	d. Navigation	N2O			#24	
#37	2E Production of Halocarbons and SF6	2. Fugitive Emissions	HFCs				#23
#38	2E Production of Halocarbons and SF6	1. By-product Emissions (Production of HCFC-22)	HFCs		#4		#14
#39	1B Fugitive Emission	1a i. Coal Mining and Handling (under gr.)	CH4				#5

N.B. Figures recorded in the Level and Trend columns indicate the ranking of individual level and trend assessments.

Table A1-2 Japan's Key Categories (FY 1990)

	A IPCC Category		B Direct GHGs	L1	L2
#1	1A Stationary Combustion	Liquid Fuels	CO2	#1	#7
#2	1A Stationary Combustion	Solid Fuels	CO2	#2	#6
#3	1A3 Mobile Combustion	b. Road Transportation	CO2	#3	#8
#4	1A Stationary Combustion	Gaseous Fuels	CO2	#4	
#5	5A Forest Land	1. Forest Land remaining Forest Land	CO2	#5	#1
#6	2A Mineral Product	1. Cement Production	CO2	#6	#10
#7	2E Production of Halocarbons and SF6	1. By-product Emissions (Production of HCFC-22)	HFCs	#7	#27
#8	1A3 Mobile Combustion	d. Navigation	CO2	#8	
#9	6C Waste Incineration		CO2	#9	#3
#10	2F(a) Consumption of Halocarbons and SF6	8. Electrical Equipment	SF6	#10	#4
#11	2A Mineral Product	3. Limestone and Dolomite Use	CO2	#11	#23
#12	2F(a) Consumption of Halocarbons and SF6	5. Solvents	PFCs	#12	#9
#13	1A Stationary Combustion	Other Fuels	CO2	#13	#17
#14	4A Enteric Fermentation		CH4	#14	#28
#15	6A Solid Waste Disposal on Land		CH4	#15	#15
#16	2B Chemical Industry	3. Adipic Acid	N2O	#16	
#17	1A3 Mobile Combustion	a. Civil Aviation	CO2	#17	
#18	4C Rice Cultivation		CH4	#18	#22
#19	2A Mineral Product	2. Lime Production	CO2		#26
#20	4B Manure Management		N2O		#14
#21	5E Settlements	2. Land converted to Settlements	CO2		#21
#22	2E Production of Halocarbons and SF6	2. Fugitive Emissions	SF6		#5
#23	4D Agricultural Soils	1. Direct Soil Emissions	N2O		#11
#24	1A3 Mobile Combustion	b. Road Transportation	N2O		#13
#25	4D Agricultural Soils	3. Indirect Emissions	N2O		#16
#26	2B Chemical Industry	1. Ammonia Production	CO2		#30
#27	2F(a) Consumption of Halocarbons and SF6	7. Semiconductor Manufacture	PFCs		#19
#28	4B Manure Management		CH4		#18
#29	1B Fugitive Emission	1a i. Coal Mining and Handling (under gr.)	CH4		#12
#30	6B Wastewater Handling		CH4		#29
#31	6C Waste Incineration		N2O		#20
#32	6B Wastewater Handling		N2O		#24
#33	5E Settlements	1. Settlements remaining Settlements	CO2		#32
#34	1A3 Mobile Combustion	d. Navigation	N2O		#25
#35	1A3 Mobile Combustion	a. Civil Aviation	N2O		#2

N.B. Figures recorded in the Level columns indicate the ranking of individual level assessments.

The data of HFCs, PFCs and SF₆ utilized for this analysis are the 1995 values.

A1.2.2. Level Assessment

Level assessment involves an identification of categories as a key by calculating the proportion of emissions and removals in each category to the total emissions and removals. The calculated values of proportion are added from the category that accounts for the largest proportion, until the sum reaches 95% for Tier 1, 90% for Tier 2. Tier 1 level assessment uses emissions and removals from each category directly and Tier 2 level assessment analyzes the emissions and removals of each category, multiplied by the uncertainty of each category.

The key category analysis was first conducted for the inventory excluding LULUCF and the key categories for source sectors were identified (1). Then the key category analysis was repeated again for the full inventory including the LULUCF categories and key categories for LULUCF sector were identified (2). In accordance with the *GPG-LULUCF*, a source category, which was identified as key in (1) but not in (2), was still regarded as key; while a source category, which was not identified as key

in (1) but was done in (2), was not regarded as key (gray rows in tables below).

Tier 1 level assessment of the latest emissions and removals (FY 2010) gives the following 12 sub-categories as the key categories (Table A1-3). Tier 2 level assessment of the latest emissions and removals (FY 2010) gives the following 25 sub-categories as the key categories (Table A1-4).

Table A1-3 Results of Tier 1 Level Assessment (FY 2010)

A	IPCC Category		B	D	E	F	Cumulative
			Direct GHGs	Current Year Estimate [Gg-CO ₂ e _{q.}]	Level Assessment	% Contribution to Level	
#1	1A Stationary Combustion	Solid Fuels	CO2	431,476.35	0.322	32.2%	32.2%
#2	1A Stationary Combustion	Liquid Fuels	CO2	256,176.89	0.191	19.1%	51.3%
#3	1A Stationary Combustion	Gaseous Fuels	CO2	210,774.39	0.157	15.7%	67.0%
#4	1A3 Mobile Combustion	b. Road Transportation	CO2	204,276.92	0.152	15.2%	82.3%
#5	5A Forest Land	1. Forest Land remaining Forest Land	CO2	76,372.11	0.057	5.7%	88.0%
#6	2A Mineral Product	1. Cement Production	CO2	23,784.44	0.018	1.8%	89.7%
#7	2F(a) Consumption of Halocarbons and SF6 (actual emissions - Tier 2)	1. Refrigeration and Air Conditioning Equipment	HFCs	17,088.19	0.013	1.3%	91.0%
#8	1A Stationary Combustion	Other Fuels	CO2	14,179.79	0.011	1.1%	92.1%
#9	6C Waste Incineration		CO2	12,657.57	0.009	0.9%	93.0%
#10	1A3 Mobile Combustion	d. Navigation	CO2	10,885.45	0.008	0.8%	93.8%
#11	1A3 Mobile Combustion	a. Civil Aviation	CO2	9,193.00	0.007	0.7%	94.5%
#12	2A Mineral Product	3. Limestone and Dolomite Use	CO2	8,073.22	0.006	0.6%	95.1%

Table A1-4 Results of Tier 2 Level Assessment (FY 2010)

A	IPCC Category		B	D	I	K	Cumulative
			Direct GHGs	Current Year Estimate [Gg-CO ₂ e _{q.}]	Source/Sink Uncertainty	Contribution to Total L2	
#1	1A3 Mobile Combustion	a. Civil Aviation	N2O	92.85	10000%	11.3%	11.3%
#2	5A Forest Land	1. Forest Land remaining Forest Land	CO2	76,372.11	11%	10.2%	21.5%
#3	2F(a) Consumption of Halocarbons	1. Refrigeration and Air Conditioning	HFCs	17,088.19	45%	9.3%	30.8%
#4	1A Stationary Combustion	Solid Fuels	CO2	431,476.35	1%	7.8%	38.7%
#5	6C Waste Incineration		CO2	12,657.57	50%	7.7%	46.4%
#6	1A3 Mobile Combustion	b. Road Transportation	CO2	204,276.92	2%	5.7%	52.1%
#7	1A Stationary Combustion	Other Fuels	CO2	14,179.79	25%	4.4%	56.5%
#8	4B Manure Management		N2O	5,475.35	48%	3.2%	59.7%
#9	4D Agricultural Soils	1. Direct Soil Emissions	N2O	2,918.17	91%	3.2%	62.9%
#10	1A Stationary Combustion	Liquid Fuels	CO2	256,176.89	1%	3.2%	66.1%
#11	2A Mineral Product	1. Cement Production	CO2	23,784.44	10%	3.0%	69.1%
#12	6C Waste Incineration		N2O	1,687.59	105%	2.2%	71.3%
#13	4D Agricultural Soils	3. Indirect Emissions	N2O	2,688.76	63%	2.1%	73.3%
#14	1A3 Mobile Combustion	b. Road Transportation	N2O	2,267.10	71%	2.0%	75.3%
#15	4B Manure Management		CH4	2,205.06	68%	1.8%	77.1%
#16	1A Stationary Combustion		N2O	3,942.82	33%	1.6%	78.7%
#17	4C Rice Cultivation		CH4	5,451.67	22%	1.5%	80.2%
#18	2F(a) Consumption of Halocarbons	7. Semiconductor Manufacture	PFCs	1,818.65	64%	1.4%	81.6%
#19	6A Solid Waste Disposal on Land		CH4	3,270.04	34%	1.4%	83.0%
#20	2A Mineral Product	3. Limestone and Dolomite Use	CO2	8,073.22	14%	1.4%	84.3%
#21	6B Wastewater Handling		N2O	1,131.61	96%	1.3%	85.7%
#22	5E Settlements	2. Land converted to Settlements	CO2	3,529.72	30%	1.3%	87.0%
#23	2A Mineral Product	2. Lime Production	CO2	6,284.59	16%	1.2%	88.2%
#24	1A3 Mobile Combustion	d. Navigation	N2O	91.32	1000%	1.1%	89.3%
#25	4A Enteric Fermentation		CH4	6,673.27	12%	1.0%	90.2%

Tier 1 level assessment of the latest emissions and removals (FY 1990) gives the following 18 sub-categories as the key categories (Table A1-5). Tier 2 level assessment of the latest emissions and removals (FY 1990) gives the following 31 sub-categories as the key categories (Table A1-6).

Table A1-5 Results of Tier 1 Level Assessment (FY 1990)

	A IPCC Category		B Direct GHGs	C Base Year Estimate [Gg-CO ₂ e q.]	E level Assessment	F % Contribution to Level	Cumulative
#1	1A Stationary Combustion	Liquid Fuels	CO2	435,168.99	0.323	32.3%	32.3%
#2	1A Stationary Combustion	Solid Fuels	CO2	308,620.23	0.229	22.9%	55.2%
#3	1A3 Mobile Combustion	b. Road Transportation	CO2	189,227.88	0.141	14.1%	69.3%
#4	1A Stationary Combustion	Gaseous Fuels	CO2	104,300.83	0.077	7.7%	77.1%
#5	5A Forest Land	1. Forest Land remaining Forest Land	CO2	76,762.09	0.057	5.7%	82.8%
#6	2A Mineral Product	1. Cement Production	CO2	37,904.87	0.028	2.8%	85.6%
#7	2E Production of Halocarbons and SF6	1. By-product Emissions (Production of HCFC-22)	HFCs	16,965.00	0.013	1.3%	86.8%
#8	1A3 Mobile Combustion	d. Navigation	CO2	13,730.95	0.010	1.0%	87.9%
#9	6C Waste Incineration		CO2	12,262.95	0.009	0.9%	88.8%
#10	2F(a) Consumption of Halocarbons and SF6 (actual emissions - Tier 2)	8. Electrical Equipment	SF6	11,004.99	0.008	0.8%	89.6%
#11	2A Mineral Product	3. Limestone and Dolomite Use	CO2	10,522.25	0.008	0.8%	90.4%
#12	2F(a) Consumption of Halocarbons and SF6 (actual emissions - Tier 2)	5. Solvents	PFCs	10,263.55	0.008	0.8%	91.1%
#13	1A Stationary Combustion	Other Fuels	CO2	9,115.90	0.007	0.7%	91.8%
#14	4A Enteric Fermentation		CH4	7,676.61	0.006	0.6%	92.4%
#15	6A Solid Waste Disposal on Land		CH4	7,645.06	0.006	0.6%	92.9%
#16	2B Chemical Industry	3. Adipic Acid	N2O	7,501.25	0.006	0.6%	93.5%
#17	1A3 Mobile Combustion	a. Civil Aviation	CO2	7,162.41	0.005	0.5%	94.0%
#18	4C Rice Cultivation		CH4	6,959.68	0.005	0.5%	94.5%
#19	2A Mineral Product	2. Lime Production	CO2	6,674.45	0.005	0.5%	95.0%

Table A1-6 Results of Tier 2 Level Assessment (FY 1990)

	A IPCC Category		B Direct GHGs	C Base Year Estimate [Gg-CO ₂ e q.]	I Source/Sink Uncertainty	K Contribution to Total L2	Cumulative
#1	5A Forest Land	1. Forest Land remaining Forest Land	CO2	76,762.09	11%	8.3%	8.3%
#2	1A3 Mobile Combustion	a. Civil Aviation	N2O	69.75	10000%	6.9%	15.2%
#3	6C Waste Incineration		CO2	12,262.95	50%	6.1%	21.3%
#4	2F(a) Consumption of Halocarbons	8. Electrical Equipment	SF6	11,004.99	49%	5.4%	26.7%
#5	2E Production of Halocarbons	2. Fugitive Emissions	SF6	4,708.30	100%	4.7%	31.4%
#6	1A Stationary Combustion	Solid Fuels	CO2	308,620.23	1%	4.5%	35.9%
#7	1A Stationary Combustion	Liquid Fuels	CO2	435,168.99	1%	4.4%	40.3%
#8	1A3 Mobile Combustion	b. Road Transportation	CO2	189,227.88	2%	4.3%	44.6%
#9	2F(a) Consumption of Halocarbons	5. Solvents	PFCs	10,263.55	40%	4.1%	48.7%
#10	2A Mineral Product	1. Cement Production	CO2	37,904.87	10%	3.9%	52.6%
#11	4D Agricultural Soils	1. Direct Soil Emissions	N2O	4,121.85	91%	3.7%	56.3%
#12	1B Fugitive Emission	1a i. Coal Mining and Handling (under gr.)	CH4	2,785.23	117%	3.2%	59.5%
#13	1A3 Mobile Combustion	b. Road Transportation	N2O	3,901.71	71%	2.7%	62.2%
#14	4B Manure Management		N2O	5,533.01	48%	2.7%	64.9%
#15	6A Solid Waste Disposal on Land		CH4	7,645.06	34%	2.6%	67.5%
#16	4D Agricultural Soils	3. Indirect Emissions	N2O	3,730.52	63%	2.3%	69.8%
#17	1A Stationary Combustion	Other Fuels	CO2	9,115.90	25%	2.3%	72.1%
#18	4B Manure Management		CH4	3,094.12	68%	2.1%	74.2%
#19	2F(a) Consumption of Halocarbons	7. Semiconductor Manufacture	PFCs	3,144.23	64%	2.0%	76.1%
#20	6C Waste Incineration		N2O	1,519.44	105%	1.6%	77.7%
#21	5E Settlements	2. Land converted to Settlements	CO2	5,113.88	30%	1.5%	79.3%
#22	4C Rice Cultivation		CH4	6,959.68	22%	1.5%	80.8%
#23	2A Mineral Product	3. Limestone and Dolomite Use	CO2	10,522.25	14%	1.4%	82.2%
#24	6B Wastewater Handling		N2O	1,295.25	96%	1.2%	83.5%
#25	1A3 Mobile Combustion	d. Navigation	N2O	112.87	1000%	1.1%	84.6%
#26	2A Mineral Product	2. Lime Production	CO2	6,674.45	16%	1.0%	85.6%
#27	2E Production of Halocarbons	1. By-product Emissions	HFCs	16,965.00	5%	0.9%	86.5%
#28	4A Enteric Fermentation		CH4	7,676.61	12%	0.9%	87.4%
#29	6B Wastewater Handling		CH4	2,143.81	42%	0.9%	88.3%
#30	2B Chemical Industry	1. Ammonia Production	CO2	3,384.68	23%	0.8%	89.1%
#31	2E Production of Halocarbons	2. Fugitive Emissions	PFCs	762.85	100%	0.8%	89.8%
#32	5E Settlements	1. Settlements remaining Settlements	CO2	955.53	76%	0.7%	90.6%

A1.2.3. Trend Assessment

The difference between the rate of change in emissions and removals in a category and the rate of change in total emissions and removals is calculated. The trend assessment is calculated by multiplying this value by the ratio of contribution of the relevant category to total emissions and removals. The calculated results, regarded as trend assessment values, are added from the category of which the proportion to the total of trend assessment values is the largest, until the total reaches 95% for Tier 1, 90% for Tier 2. At this point, these categories are defined as the key categories. Tier 1 level assessment uses emissions and removals from each category directly and Tier 2 level assessment

analyzes the emissions and removals of each category, multiplied by the uncertainty of each category.

The key category analysis was first conducted for the inventory excluding LULUCF and the key categories for source sectors were identified (1). Then the key category analysis was repeated again for the full inventory including the LULUCF categories and key categories for LULUCF sector were identified (2). In accordance with the *GPG-LULUCF*, a source category, which was identified as key in (1) but not in (2), was still regarded as key; while a source category, which was not identified as key in (1) but was done in (2), was not regarded as key (gray rows in tables below).

Tier 1 trend assessment of the latest emissions and removals (FY 2010) gives the following 14 sub-categories as the key categories (Table A1-7). Tier 2 trend assessment of the latest emissions and removals (FY 2010) gives the following 26 sub-categories as the key categories (Table A1-8).

Table A1-7 Results of Tier 1 Trend Assessment (FY 2010)

A IPCC Category		B Direct GHGs	C Base Year Estimate [Gg-CO ₂ eq.]	D Current Year Estimate [Gg-CO ₂ eq.]	H % Contribution to Trend	Cumulative
#1 1A Stationary Combustion	Liquid Fuels	CO ₂	435169	256177	32.4%	32.4%
#2 1A Stationary Combustion	Solid Fuels	CO ₂	308620	431476	22.7%	55.1%
#3 1A Stationary Combustion	Gaseous Fuels	CO ₂	104301	210774	19.6%	74.7%
#4 2E Production of Halocarbons and SF ₆	1. By-product Emissions (Production of HCFC-22)	HFCs	16965	42	3.1%	77.8%
#5 2F(a) Consumption of Halocarbons and SF ₆ (actual emissions - Tier 2)	1. Refrigeration and Air Conditioning Equipment	HFCs	840	17088	3.0%	80.7%
#6 1A3 Mobile Combustion	b. Road Transportation	CO ₂	189228	204277	2.9%	83.6%
#7 2A Mineral Product	1. Cement Production	CO ₂	37905	23784	2.6%	86.2%
#8 2F(a) Consumption of Halocarbons and SF ₆ (actual emissions - Tier 2)	8. Electrical Equipment	SF ₆	11005	652	1.9%	88.1%
#9 2F(a) Consumption of Halocarbons and SF ₆ (actual emissions - Tier 2)	5. Solvents	PFCs	10264	1376	1.6%	89.7%
#10 2B Chemical Industry	3. Adipic Acid	N ₂ O	7501	516	1.3%	91.0%
#11 1A Stationary Combustion	Other Fuels	CO ₂	9116	14180	0.9%	91.9%
#12 2E Production of Halocarbons and SF ₆	2. Fugitive Emissions	SF ₆	4708	198	0.8%	92.7%
#13 6A Solid Waste Disposal on Land		CH ₄	7645	3270	0.8%	93.5%
#14 1A3 Mobile Combustion	d. Navigation	CO ₂	13731	10885	0.5%	94.0%
#15 1B Fugitive Emission	1a i. Coal Mining and Handling (under gr.)	CH ₄	2785	35	0.5%	94.5%
#16 2A Mineral Product	3. Limestone and Dolomite Use	CO ₂	10522	8073	0.4%	95.0%
#17 1A3 Mobile Combustion	a. Civil Aviation	CO ₂	7162	9193	0.4%	95.3%

Table A1-8 Results of Tier 2 Trend Assessment (FY 2010)

A IPCC Category		B Direct GHGs	C Base Year Estimate [Gg-CO ₂ eq.]	D Current Year Estimate [Gg-CO ₂ eq.]	I Source/Sink Uncertainty	M Contribution to Total T2	Cumulative
#1 2F(a) Consumption of Halocarbons	1. Refrigeration and Air Conditioning	HFCs	840.40	17,088.19	45%	15.2%	15.2%
#2 2F(a) Consumption of Halocarbons	8. Electrical Equipment	SF ₆	11,004.99	652.25	49%	10.6%	25.8%
#3 2E Production of Halocarbons	2. Fugitive Emissions	SF ₆	4,708.30	198.37	100%	9.4%	35.2%
#4 2F(a) Consumption of Halocarbons	5. Solvents	PFCs	10,263.55	1,375.99	40%	7.4%	42.6%
#5 1B Fugitive Emission	1a i. Coal Mining and Handling (under gr.)	CH ₄	2,785.23	34.69	117%	6.7%	49.3%
#6 1A3 Mobile Combustion	a. Civil Aviation	N ₂ O	69.75	92.85	10000%	4.9%	54.1%
#7 1A Stationary Combustion	Solid Fuels	CO ₂	308,620.23	431,476.35	1%	3.8%	58.0%
#8 1A Stationary Combustion	Liquid Fuels	CO ₂	435,168.99	256,176.89	1%	3.7%	61.7%
#9 6A Solid Waste Disposal on Land		CH ₄	7,645.06	3,270.04	34%	3.1%	64.8%
#10 2A Mineral Product	1. Cement Production	CO ₂	37,904.87	23,784.44	10%	3.0%	67.8%
#11 1A Stationary Combustion	Other Fuels	CO ₂	9,115.90	14,179.79	25%	2.7%	70.5%
#12 1A3 Mobile Combustion	b. Road Transportation	N ₂ O	3,901.71	2,267.10	71%	2.4%	72.9%
#13 4D Agricultural Soils	1. Direct Soil Emissions	N ₂ O	4,121.85	2,918.17	91%	2.2%	75.1%
#14 2E Production of Halocarbons	1. By-product Emissions	HFCs	16,965.00	42.12	5%	1.9%	77.0%
#15 2F(a) Consumption of Halocarbons	7. Semiconductor Manufacture	PFCs	3,144.23	1,818.65	64%	1.8%	78.8%
#16 4D Agricultural Soils	3. Indirect Emissions	N ₂ O	3,730.52	2,688.76	63%	1.4%	80.1%
#17 2B Chemical Industry	3. Adipic Acid	N ₂ O	7,501.25	516.10	9%	1.3%	81.5%
#18 4B Manure Management		CH ₄	3,094.12	2,205.06	68%	1.2%	82.7%
#19 1A Stationary Combustion		N ₂ O	2,160.43	3,942.82	33%	1.2%	83.9%
#20 5B Cropland	2. Land converted to Cropland	CO ₂	2,513.21	452.41	28%	1.2%	85.1%
#21 2E Production of Halocarbons	2. Fugitive Emissions	PFCs	762.85	200.24	100%	1.2%	86.3%
#22 5E Settlements	2. Land converted to Settlements	CO ₂	5,113.88	3,529.72	30%	1.0%	87.3%
#23 2E Production of Halocarbons	2. Fugitive Emissions	HFCs	480.12	86.22	100%	0.8%	88.1%
#24 1A3 Mobile Combustion	b. Road Transportation	CO ₂	189,227.88	204,276.92	2%	0.8%	88.9%
#25 5A Forest Land	2. Land converted to Forest Land	CO ₂	1,830.25	304.80	24%	0.8%	89.6%
#26 6B Wastewater Handling		CH ₄	2,143.81	1,269.65	42%	0.7%	90.4%

Data utilized for the key category analysis are shown in Table A1-9 and A1-10 as references.

Table A1-9 Data used for the key category analysis (FY 2010)

A	IPCC Category	B	C	D	E	F	G	H	I	J	K	L	M	
		Direct GHGs	Base Year Estimate (Gg-CO ₂ e.g.)	Current Year Estimate (Gg-CO ₂ e.g.)	Level Assessment	% Contribution to Level	Trend Assessment	% Contribution to Trend	Source/Sink Uncertainty	Level Uncertainty (x 1000)	Contribution to Total L2	Trend Uncertainty (x 1000)	Contribution to Total T2	
#1	1A Stationary Combustion	Liquid Fuels	CO2	435,168.99	256,176.89	0.191	19.1%	0.1327	32.4%	1%	1.94	0.03	1.35	0.04
#2	1A Stationary Combustion	Solid Fuels	CO2	308,620.23	431,476.35	0.322	32.2%	0.0930	22.7%	1%	4.79	0.08	1.38	0.04
#3	1A Stationary Combustion	Gaseous Fuels	CO2	104,300.83	210,774.39	0.157	15.7%	0.0801	19.6%	0%	4.47	0.01	0.24	0.01
#4	1A Stationary Combustion	Other Fuels	CO2	9,115.90	14,179.79	0.011	1.1%	0.0038	0.9%	25%	2.66	0.04	0.96	0.03
#5	1A Stationary Combustion		CH4	543.43	588.64	0.000	0.0%	0.0000	0.0%	47%	0.21	0.00	0.02	0.00
#6	1A Stationary Combustion		N2O	2,160.43	3,942.82	0.003	0.3%	0.0013	0.3%	33%	0.97	0.02	0.44	0.01
#7	1A Stationary Combustion		CH4	49.20	85.80	0.000	0.0%	0.0000	0.0%	117%	0.07	0.00	0.03	0.00
#8	1A Stationary Combustion		N2O	385.39	340.36	0.000	0.0%	0.0000	0.0%	36%	0.99	0.00	0.01	0.00
#9	1A3 Mobile Combustion	a. Civil Aviation	CO2	7,162.41	9,193.00	0.007	0.7%	0.0015	0.4%	3%	0.17	0.00	0.04	0.00
#10	1A3 Mobile Combustion	b. Road Transportation	CO2	189,227.88	204,276.92	0.152	15.2%	0.0119	2.9%	2%	3.50	0.06	0.27	0.01
#11	1A3 Mobile Combustion	c. Railways	CO2	932.45	588.10	0.000	0.0%	0.0003	0.1%	2%	0.01	0.00	0.01	0.00
#12	1A3 Mobile Combustion	d. Navigation	CO2	13,730.95	10,885.45	0.008	0.8%	0.0021	0.5%	2%	0.19	0.00	0.05	0.00
#13	1A3 Mobile Combustion	a. Civil Aviation	CH4	2.94	4.58	0.000	0.0%	0.0000	0.0%	200%	0.01	0.00	0.00	0.00
#14	1A3 Mobile Combustion	b. Road Transportation	CH4	266.66	140.78	0.000	0.0%	0.0001	0.0%	64%	0.07	0.00	0.06	0.00
#15	1A3 Mobile Combustion	c. Railways	CH4	1.18	0.72	0.000	0.0%	0.0000	0.0%	14%	0.00	0.00	0.00	0.00
#16	1A3 Mobile Combustion	d. Navigation	CH4	26.76	21.65	0.000	0.0%	0.0000	0.0%	200%	0.03	0.00	0.01	0.00
#17	1A3 Mobile Combustion	a. Civil Aviation	N2O	69.75	92.85	0.000	0.0%	0.0000	0.0%	1000%	0.11	0.00	1.75	0.05
#18	1A3 Mobile Combustion	b. Road Transportation	N2O	3,901.71	2,367.10	0.002	0.2%	0.0012	0.3%	71%	1.20	0.02	0.86	0.02
#19	1A3 Mobile Combustion	c. Railways	N2O	121.39	75.01	0.000	0.0%	0.0000	0.0%	11%	0.01	0.00	0.00	0.00
#20	1A3 Mobile Combustion	d. Navigation	N2O	112.87	91.32	0.000	0.0%	0.0000	0.0%	100%	0.68	0.01	0.16	0.00
#21	1B Fugitive Emission	1a. Coal Mining and Handling (under gr.)	CH4	2,785.23	34.69	0.000	0.0%	0.0021	0.5%	117%	0.03	0.00	2.40	0.07
#22	1B Fugitive Emission	1a. Coal Mining and Handling (surface)	CH4	21.20	9.80	0.000	0.0%	0.0000	0.0%	185%	0.01	0.00	0.02	0.00
#23	1B Fugitive Emission	2a. Oil	CO2	0.14	0.10	0.000	0.0%	0.0000	0.0%	21%	0.00	0.00	0.00	0.00
#24	1B Fugitive Emission	2a. Oil	CH4	28.32	24.66	0.000	0.0%	0.0000	0.0%	17%	0.00	0.00	0.00	0.00
#25	1B Fugitive Emission	2a. Oil	N2O	0.00	0.00	0.000	0.0%	0.0000	0.0%	27%	0.00	0.00	0.00	0.00
#26	1B Fugitive Emission	2b. Natural Gas	CO2	0.25	0.41	0.000	0.0%	0.0000	0.0%	25%	0.01	0.00	0.00	0.00
#27	1B Fugitive Emission	2b. Natural Gas	CH4	187.94	295.56	0.000	0.0%	0.0001	0.0%	22%	0.05	0.00	0.02	0.00
#28	1B Fugitive Emission	2c. Venting & Flaring	CO2	36.23	32.64	0.000	0.0%	0.0000	0.0%	18%	0.00	0.00	0.00	0.00
#29	1B Fugitive Emission	2c. Venting & Flaring	CH4	14.45	11.02	0.000	0.0%	0.0000	0.0%	20%	0.00	0.00	0.00	0.00
#30	1B Fugitive Emission	2c. Venting & Flaring	N2O	0.11	0.11	0.000	0.0%	0.0000	0.0%	18%	0.00	0.00	0.00	0.00
#31	2A Mineral Product	1. Cement Production	CO2	37,904.87	23,784.44	0.018	1.8%	0.0105	2.6%	10%	1.85	0.03	1.09	0.03
#32	2A Mineral Product	2. Lime Production	CO2	6,674.45	6,284.59	0.005	0.5%	0.0003	0.1%	16%	0.74	0.01	0.04	0.01
#33	2A Mineral Product	3. Limestone and Dolomite Use	CO2	10,522.25	8,073.22	0.006	0.6%	0.0018	0.4%	14%	0.83	0.01	0.25	0.01
#34	2A Mineral Product	4. Soda Ash Production and Use	CO2	267.28	157.94	0.000	0.0%	0.0001	0.0%	16%	0.02	0.00	0.00	0.00
#35	2B Chemical Industry	1. Ammonia Production	CO2	3,384.68	2,104.42	0.002	0.2%	0.0009	0.2%	33%	0.36	0.01	0.22	0.01
#36	2B Chemical Industry	other products except Ammonia	CO2	824.39	630.81	0.000	0.0%	0.0001	0.0%	77%	0.36	0.01	0.11	0.00
#37	2B Chemical Industry	2. Nitric Acid	N2O	765.70	561.64	0.000	0.0%	0.0002	0.0%	46%	0.19	0.00	0.07	0.00
#38	2B Chemical Industry	3. Adipic Acid	N2O	7,501.25	516.10	0.000	0.0%	0.0052	1.3%	9%	0.04	0.00	0.48	0.01
#39	2B Chemical Industry	4. Carbide Production	CH4	0.42	0.66	0.000	0.0%	0.0000	0.0%	100%	0.00	0.00	0.00	0.00
#40	2B Chemical Industry	5. Carbon Black, Ethylene, Ethylene	CH4	337.80	103.32	0.000	0.0%	0.0002	0.0%	89%	0.07	0.00	0.16	0.00
#41	2C Metal Production	1. Iron and Steel Production	CO2	356.09	159.86	0.000	0.0%	0.0001	0.0%	5%	0.01	0.00	0.01	0.00
#42	2C Metal Production	1. Iron and Steel Production	CH4	15.47	12.31	0.000	0.0%	0.0000	0.0%	163%	0.01	0.00	0.00	0.00
#43	2C Metal Production	2. Ferrous Production	CH4	3.89	2.56	0.000	0.0%	0.0000	0.0%	163%	0.00	0.00	0.00	0.00
#44	2C Metal Production	3. Aluminum Production	PFCs	69.74	19.38	0.000	0.0%	0.0000	0.0%	23%	0.00	0.00	0.01	0.00
#45	2C Metal Production	4. SF6 Used in Aluminum and Magnesium	SF6	119.50	307.90	0.000	0.0%	0.0001	0.0%	5%	0.01	0.00	0.01	0.00
#46	2E Production of Halocarbons	1. By-product Emissions	HFCs	16,965.00	42.12	0.000	0.0%	0.0126	3.1%	5%	0.00	0.00	0.68	0.02
#47	2E Production of Halocarbons	2. Fugitive Emissions	HFCs	480.12	86.22	0.000	0.0%	0.0003	0.1%	100%	0.06	0.00	0.30	0.01
#48	2E Production of Halocarbons	2. Fugitive Emissions	PFCs	762.85	200.24	0.000	0.0%	0.0004	0.1%	100%	0.15	0.00	0.42	0.01
#49	2E Production of Halocarbons	2. Fugitive Emissions	SF6	4,708.30	198.37	0.000	0.0%	0.0054	0.8%	100%	0.15	0.00	3.38	0.09
#50	2F(a) Consumption of Halocarbons	1. Refrigeration and Air Conditioning	HFCs	840.40	17,088.19	0.013	1.3%	0.0122	3.5%	45%	5.72	0.09	5.46	0.15
#51	2F(a) Consumption of Halocarbons	2. Foam Blowing	HFCs	451.76	290.97	0.000	0.0%	0.0001	0.0%	50%	0.11	0.00	0.06	0.00
#52	2F(a) Consumption of Halocarbons	3. Fire Extinguishers	HFCs	0.00	6.72	0.000	0.0%	0.0000	0.0%	64%	0.00	0.00	0.00	0.00
#53	2F(a) Consumption of Halocarbons	4. Aerosols/ Metered Dose Inhalers	HFCs	1,365.00	640.00	0.000	0.0%	0.0005	0.1%	20%	0.13	0.00	0.14	0.00
#54	2F(a) Consumption of Halocarbons	5. Solvents	PFCs	10,263.57	1,735.99	0.001	0.1%	0.0066	1.6%	40%	0.41	0.01	2.63	0.07
#55	2F(a) Consumption of Halocarbons	7. Semiconductor Manufacture	HFCs	157.89	102.19	0.000	0.0%	0.0000	0.0%	64%	0.05	0.00	0.03	0.00
#56	2F(a) Consumption of Halocarbons	7. Semiconductor Manufacture	PFCs	3,144.23	1,818.65	0.001	0.1%	0.0010	0.2%	64%	0.87	0.01	0.63	0.02
#57	2F(a) Consumption of Halocarbons	7. Semiconductor Manufacture	SF6	1,128.66	703.91	0.001	0.1%	0.0003	0.1%	64%	0.34	0.01	0.20	0.01
#58	2F(a) Consumption of Halocarbons	8. Electrical Equipment	SF6	11,004.99	652.25	0.000	0.0%	0.0077	1.9%	49%	0.24	0.00	3.82	0.11
#59	2F(a) Consumption of Halocarbons	9. Other - Railway Silicon Rectifiers	PFCs	0.00	0.00	0.000	0.0%	0.0000	0.0%	49%	0.00	0.00	0.00	0.00
#60	3 Solvent & Other Product Use	Using Laughing Gas in Hospital	N2O	287.07	98.95	0.000	0.0%	0.0001	0.0%	5%	0.00	0.00	0.01	0.00
#61	4A Enteric Fermentation		CH4	7,676.61	6,673.27	0.005	0.5%	0.0007	0.2%	12%	0.58	0.01	0.09	0.00
#62	4B Manure Management		CH4	3,094.12	2,205.06	0.002	0.2%	0.0007	0.2%	68%	1.12	0.02	0.45	0.01
#63	4B Manure Management		N2O	5,533.01	5,475.35	0.004	0.4%	0.0000	0.0%	48%	1.98	0.03	0.01	0.00
#64	4C Rice Cultivation		CH4	6,959.68	5,451.67	0.004	0.4%	0.0011	0.3%	22%	0.91	0.01	0.25	0.01
#65	4D Agricultural Soils	1. Direct Soil Emissions	N2O	4,121.85	2,918.17	0.002	0.2%	0.0009	0.2%	91%	1.97	0.03	0.81	0.02
#66	4D Agricultural Soils	2. Pasture, Range and Paddock Manure	N2O	11.91	11.80	0.000	0.0%	0.0000	0.0%	133%	0.01	0.00	0.43	0.01
#67	4D Agricultural Soils	3. Indirect Emissions	N2O	3,730.52	2,688.76	0.002	0.2%	0.0008	0.2%	63%	1.27	0.02	0.49	0.01
#68	4F Field Burning of Agricultural Residues		CH4	100.68	57.39	0.000	0.0%	0.0000	0.0%	217%	0.09	0.00	0.07	0.00
#69	4F Field Burning of Agricultural Residues		N2O	32.65	18.12	0.000	0.0%	0.0000	0.0%	161%	0.02	0.00	0.02	0.00
#70	5A Forest Land	1. Forest Land remaining Forest Land	CO2	76,762.09	76,372.11	0.057	5.7%	0.0001	0.0%	11%	6.23	0.10	0.01	0.00
#71	5A Forest Land	2. Land converted to Forest Land	CO2	1,830.25	304.80	0.000	0.0%	0.0011	0.3%	24%	0.05	0.00	0.27	0.01
#72	5A Forest Land		CH4	8.51	2.12	0.000	0.0%	0.0000	0.0%	40%	0.00	0.00	0.00	0.00
#73	5A Forest Land		N2O	0.86	0.22	0.000	0.0%	0.0000	0.0%	42%	0.00	0.00	0.00	0.00
#74	5B Cropland	1. Cropland remaining Cropland	CO2	0.00	0.00	0.000	0.0%	0.0000	0.0%	0%	0.00	0.00	0.00	0.00
#75	5B Cropland	2. Land converted to Cropland	CO2	2,513.21	452.41	0.000	0.0%	0.0015	0.4%	28%	0.10	0.00	0.43	0.01
#76	5B Cropland		CH4	0.00	0.00	0.000	0.0%	0.0000	0.0%	0%	0.00	0.00	0.00	0.00
#77	5B Cropland		N2O	90.02	6.18	0.000	0.0%	0.0001	0.0%	75%	0.00	0.00	0.05	0.00
#78	5C Grassland	1. Grassland remaining Grassland	CO2	0.00	0.00	0.000	0.0%	0.0000	0.0%	0%	0.00	0.00	0.00	0.00
#79	5C Grassland	2. Land converted to Grassland	CO2	444.03	215.86	0.000	0.0%	0.0002	0.0%	47%	0.08	0.00	0.08	0.00
#80	5C Grassland		CH4	0.00	0.00	0.000	0.0%	0.0000	0.0%	0%	0.00	0.00	0.00	0.00
#81	5C Grassland		N2O	0.00	0.00	0.000	0.0%	0.0000	0.0%	0%	0.00	0.00	0.00	0.00
#82	5D Wetlands	1. Wetlands remaining Wetlands	CO2	0.00	0.00	0.000	0.0%	0.0000	0.0%	0%	0.00	0.00	0.00	0.00
#83	5D Wetlands	2. Land converted to Wetlands	CO2	85.84	82.13	0.000	0.0%	0.0000	0.0%	30%	0.02	0.00	0.00	0.00
#84	5D Wetlands		CH4	0.00	0.00	0.000	0.0%	0.0000	0.0%	0%	0.00	0.00	0.00	0.00
#85														

Table A1-10 Data used for the key category analysis (FY 1990)

A	IPCC Category	B	C	E	F	I	J	K	
		Direct GHGs	Base Year Estimate [Gg-CO ₂ eq.]	Level Assesment	% Contribution to Level	Source/Sink Uncertainty	Level Uncertainty (x 1000)	Contribution to Total L2	
#1	1A Stationary Combustion	Liquid Fuels	CO2	435,168.99	0.323	32.3%	1%	3.28	0.04
#2	1A Stationary Combustion	Solid Fuels	CO2	308,620.23	0.229	22.9%	1%	3.41	0.05
#3	1A Stationary Combustion	Gaseous Fuels	CO2	104,300.83	0.077	7.7%	0%	0.23	0.00
#4	1A Stationary Combustion	Other Fuels	CO2	9,115.90	0.007	0.7%	25%	1.70	0.02
#5	1A Stationary Combustion		CH4	543.43	0.000	0.0%	47%	0.19	0.00
#6	1A Stationary Combustion		N2O	2,160.43	0.002	0.2%	33%	0.53	0.01
#7	1A Stationary Combustion		CH4	49.20	0.000	0.0%	117%	0.04	0.00
#8	1A Stationary Combustion		N2O	385.39	0.000	0.0%	36%	0.10	0.00
#9	1A3 Mobile Combustion	a. Civil Aviation	CO2	7,162.41	0.005	0.5%	3%	0.13	0.00
#10	1A3 Mobile Combustion	b. Road Transportation	CO2	189,227.88	0.141	14.1%	2%	3.23	0.04
#11	1A3 Mobile Combustion	c. Railways	CO2	932.45	0.001	0.1%	2%	0.02	0.00
#12	1A3 Mobile Combustion	d. Navigation	CO2	13,730.95	0.010	1.0%	2%	0.24	0.00
#13	1A3 Mobile Combustion	a. Civil Aviation	CH4	2.94	0.000	0.0%	200%	0.00	0.00
#14	1A3 Mobile Combustion	b. Road Transportation	CH4	266.66	0.000	0.0%	64%	0.13	0.00
#15	1A3 Mobile Combustion	c. Railways	CH4	1.18	0.000	0.0%	14%	0.00	0.00
#16	1A3 Mobile Combustion	d. Navigation	CH4	26.76	0.000	0.0%	200%	0.04	0.00
#17	1A3 Mobile Combustion	a. Civil Aviation	N2O	69.75	0.000	0.0%	10000%	5.18	0.07
#18	1A3 Mobile Combustion	b. Road Transportation	N2O	3,901.71	0.003	0.3%	71%	2.05	0.03
#19	1A3 Mobile Combustion	c. Railways	N2O	121.39	0.000	0.0%	11%	0.01	0.00
#20	1A3 Mobile Combustion	d. Navigation	N2O	112.87	0.000	0.0%	1000%	0.84	0.01
#21	1B Fugitive Emission	1a i. Coal Mining and Handling (under gr.)	CH4	2,785.23	0.002	0.2%	117%	2.42	0.03
#22	1B Fugitive Emission	1a i. Coal Mining and Handling (surface)	CH4	21.20	0.000	0.0%	185%	0.03	0.00
#23	1B Fugitive Emission	2a. Oil	CO2	0.14	0.000	0.0%	21%	0.00	0.00
#24	1B Fugitive Emission	2a. Oil	CH4	28.32	0.000	0.0%	17%	0.00	0.00
#25	1B Fugitive Emission	2a. Oil	N2O	0.00	0.000	0.0%	27%	0.00	0.00
#26	1B Fugitive Emission	2b. Natural Gas	CO2	0.25	0.000	0.0%	25%	0.00	0.00
#27	1B Fugitive Emission	2b. Natural Gas	CH4	187.94	0.000	0.0%	22%	0.03	0.00
#28	1B Fugitive Emission	2c. Venting & Flaring	CO2	36.23	0.000	0.0%	18%	0.00	0.00
#29	1B Fugitive Emission	2c. Venting & Flaring	CH4	14.45	0.000	0.0%	20%	0.00	0.00
#30	1B Fugitive Emission	2c. Venting & Flaring	N2O	0.11	0.000	0.0%	18%	0.00	0.00
#31	2A Mineral Product	1. Cement Production	CO2	37,904.87	0.028	2.8%	10%	2.94	0.04
#32	2A Mineral Product	2. Lime Production	CO2	6,674.45	0.005	0.5%	16%	0.78	0.01
#33	2A Mineral Product	3. Limestone and Dolomite Use	CO2	10,522.25	0.008	0.8%	14%	1.08	0.01
#34	2A Mineral Product	4. Soda Ash Production and Use	CO2	267.28	0.000	0.0%	16%	0.03	0.00
#35	2B Chemical Industry	1. Ammonia Production	CO2	3,384.68	0.003	0.3%	23%	0.58	0.01
#36	2B Chemical Industry	other products except Ammonia	CO2	824.39	0.001	0.1%	77%	0.47	0.01
#37	2B Chemical Industry	2. Nitric Acid	N2O	765.70	0.001	0.1%	46%	0.26	0.00
#38	2B Chemical Industry	3. Adipic Acid	N2O	7,501.25	0.006	0.6%	9%	0.51	0.01
#39	2B Chemical Industry	4. Carbide Production	CH4	0.42	0.000	0.0%	100%	0.00	0.00
#40	2B Chemical Industry	5. Carbon Black, Ethylene, Ethylene	CH4	337.80	0.000	0.0%	89%	0.22	0.00
#41	2C Metal Production	1. Iron and Steel Production	CO2	356.09	0.000	0.0%	5%	0.01	0.00
#42	2C Metal Production	1. Iron and Steel Production	CH4	15.47	0.000	0.0%	163%	0.02	0.00
#43	2C Metal Production	2. Ferroalloys Production	CH4	3.89	0.000	0.0%	163%	0.00	0.00
#44	2C Metal Production	3. Aluminium Production	PFCs	69.74	0.000	0.0%	33%	0.02	0.00
#45	2C Metal Production	4. SF6 Used in Aluminium and Magnesium	SF6	119.50	0.000	0.0%	5%	0.00	0.00
#46	2E Production of Halocarbons	1. By-product Emissions	HFCs	16,965.00	0.013	1.3%	5%	0.68	0.01
#47	2E Production of Halocarbons	2. Fugitive Emissions	HFCs	480.12	0.000	0.0%	100%	0.36	0.00
#48	2E Production of Halocarbons	2. Fugitive Emissions	PFCS	762.85	0.001	0.1%	100%	0.57	0.01
#49	2E Production of Halocarbons	2. Fugitive Emissions	SF6	4,708.30	0.003	0.3%	100%	3.51	0.05
#50	2F(a) Consumption of Halocarbons	1. Refrigeration and Air Conditioning	HFCs	840.40	0.001	0.1%	45%	0.28	0.00
#51	2F(a) Consumption of Halocarbons	2. Foam Blowing	HFCs	451.76	0.000	0.0%	50%	0.17	0.00
#52	2F(a) Consumption of Halocarbons	3. Fire Extinguishers	HFCs	0.00	0.000	0.0%	64%	0.00	0.00
#53	2F(a) Consumption of Halocarbons	4. Aerosols/ Metered Dose Inhalers	HFCs	1,365.00	0.001	0.1%	26%	0.27	0.00
#54	2F(a) Consumption of Halocarbons	5. Solvents	PFCs	10,263.55	0.008	0.8%	40%	3.05	0.04
#55	2F(a) Consumption of Halocarbons	7. Semiconductor Manufacture	HFCs	157.89	0.000	0.0%	68%	0.08	0.00
#56	2F(a) Consumption of Halocarbons	7. Semiconductor Manufacture	PFCs	3,144.23	0.002	0.2%	64%	1.50	0.02
#57	2F(a) Consumption of Halocarbons	7. Semiconductor Manufacture	SF6	1,128.66	0.001	0.1%	64%	0.54	0.01
#58	2F(a) Consumption of Halocarbons	8. Electrical Equipment	SF6	11,004.99	0.008	0.8%	49%	4.05	0.05
#59	2F(a) Consumption of Halocarbons	9. Other - Railway Silicon Rectifiers	PFCs	0.00	0.000	0.0%	49%	0.00	0.00
#60	3 Solvent & Other Product Use	Using Laughing Gas in Hospital	N2O	287.07	0.000	0.0%	5%	0.01	0.00
#61	4A Enteric Fermentation		CH4	7,676.61	0.006	0.6%	12%	0.67	0.01
#62	4B Manure Management		CH4	3,094.12	0.002	0.2%	68%	1.56	0.02
#63	4B Manure Management		N2O	5,533.01	0.004	0.4%	48%	1.99	0.03
#64	4C Rice Cultivation		CH4	6,959.68	0.005	0.5%	22%	1.16	0.02
#65	4D Agricultural Soils	1. Direct Soil Emissions	N2O	4,121.85	0.003	0.3%	91%	2.77	0.04
#66	4D Agricultural Soils	2. Pasture, Range and Paddock Manure	N2O	11.91	0.000	0.0%	133%	0.01	0.00
#67	4D Agricultural Soils	3. Indirect Emissions	N2O	3,730.52	0.003	0.3%	63%	1.75	0.02
#68	4F Field Burning of Agricultural Residues		CH4	100.68	0.000	0.0%	217%	0.16	0.00
#69	4F Field Burning of Agricultural Residues		N2O	32.65	0.000	0.0%	161%	0.04	0.00
#70	5A Forest Land	1. Forest Land remaining Forest Land	CO2	76,762.09	0.057	5.7%	11%	6.24	0.08
#71	5A Forest Land	2. Land converted to Forest Land	CO2	1,830.25	0.001	0.1%	24%	0.32	0.00
#72	5A Forest Land		CH4	8.51	0.000	0.0%	40%	0.00	0.00
#73	5A Forest Land		N2O	0.86	0.000	0.0%	42%	0.00	0.00
#74	5B Cropland	1. Cropland remaining Cropland	CO2	0.00	0.000	0.0%	0%	0.00	0.00
#75	5B Cropland	2. Land converted to Cropland	CO2	2,513.21	0.002	0.2%	28%	0.53	0.01
#76	5B Cropland		CH4	0.00	0.000	0.0%	0%	0.00	0.00
#77	5B Cropland		N2O	90.02	0.000	0.0%	75%	0.05	0.00
#78	5C Grassland	1. Grassland remaining Grassland	CO2	0.00	0.000	0.0%	0%	0.00	0.00
#79	5C Grassland	2. Land converted to Grassland	CO2	444.03	0.000	0.0%	47%	0.16	0.00
#80	5C Grassland		CH4	0.00	0.000	0.0%	0%	0.00	0.00
#81	5C Grassland		N2O	0.00	0.000	0.0%	0%	0.00	0.00
#82	5D Wetlands	1. Wetlands remaining Wetlands	CO2	0.00	0.000	0.0%	0%	0.00	0.00
#83	5D Wetlands	2. Land converted to Wetlands	CO2	85.84	0.000	0.0%	30%	0.02	0.00
#84	5D Wetlands		CH4	0.00	0.000	0.0%	0%	0.00	0.00
#85	5D Wetlands		N2O	0.00	0.000	0.0%	0%	0.00	0.00
#86	5E Settlements	1. Settlements remaining Settlements	CO2	955.53	0.001	0.1%	76%	0.54	0.01
#87	5E Settlements	2. Land converted to Settlements	CO2	5,113.88	0.004	0.4%	30%	1.16	0.02
#88	5E Settlements		CH4	0.00	0.000	0.0%	0%	0.00	0.00
#89	5E Settlements		N2O	0.00	0.000	0.0%	0%	0.00	0.00
#90	5F Other Land	1. Other Land remaining Other Land	CO2	0.00	0.000	0.0%	0%	0.00	0.00
#91	5F Other Land	2. Land converted to Other Land	CO2	1,553.92	0.001	0.1%	28%	0.52	0.00
#92	5F Other Land		CH4	0.00	0.000	0.0%	0%	0.00	0.00
#93	5F Other Land		N2O	0.00	0.000	0.0%	0%	0.00	0.00
#94	5G Other	CO2 emissions from agricultural lime	CO2	550.22	0.000	0.0%	51%	0.21	0.00
#95	6A Solid Waste Disposal on Land		CH4	7,645.06	0.006	0.6%	34%	1.94	0.03
#96	6B Wastewater Handling		CH4	2,143.81	0.002	0.2%	42%	0.66	0.01
#97	6B Wastewater Handling		N2O	1,295.25	0.001	0.1%	96%	0.93	0.01
#98	6C Waste Incineration		CO2	12,262.95	0.009	0.9%	50%	4.55	0.06
#99	6C Waste Incineration		CH4	13.48	0.000	0.0%	119%	0.01	0.00
#100	6C Waste Incineration		N2O	1,519.44	0.001	0.1%	105%	1.18	0.02
#101	6D Other		CO2	702.83	0.001	0.1%	25%	0.13	0.00
#102	6D Other		CH4	111.85	0.000	0.0%	74%	0.06	0.00
#103	6D Other		N2O	99.06	0.000	0.0%	86%	0.06	0.00
	TOTAL			1,346,245.93	1.00	100.0%		75.02	1.00

A1.2.4. Qualitative Analysis

Key categories identified in the qualitative analysis include the categories in which: mitigation techniques have been employed, significant variance of emissions and removals has been confirmed, a high uncertainty exists due to the solo implementation of the Tier 1 analysis of key categories, unexpectedly high or low estimates are identified, and major changes in the estimation methodology or data have occurred.

In Japan, the categories in which mitigation techniques have been employed, emissions and removals have been newly estimated, and estimation methods have been changed, were identified as key in terms of the qualitative analysis. In this year, the key categories were identified only based on the quantitative results of the level and trend assessments, including both Tier 1 and Tier 2.

Annex 2. Detailed Discussion on Methodology and Data for Estimating CO₂ Emissions from Fossil Fuel Combustion

A2.1. Discrepancies between the figures reported in the CRF tables and the IEA statistics

In the report of the individual review of the greenhouse gas inventory of Japan submitted in 2006 (FCCC/ARR/2006/JPN), which was conducted from January to February 2007, the ERT (Expert Review Team) recommended that in the next NIR submission Japan provide a clear explanation for the discrepancies found between the data in the CRF tables and the IEA statistics.

In response to this recommendation, Japan has provided the detailed information on the Annex 2 regarding the discrepancies of the FY 2005 data between the CRF tables and the IEA statistics. Also in the individual review report of the GHG inventory of Japan submitted in 2010 (FCCC/ARR/2010/JPN), the updating of this information with the latest available inventory year data was recommended by the ERT. In response to this recommendation, the detailed information regarding the discrepancies of the reported value between the CRF and the IEA statistics is hereby updated with the FY 2009 actual data. The IEA statistical data used in the explanation were extracted from the Energy Statistics of OECD Countries 2008–2009, 2011 Edition, OECD/IEA (CD-ROM version).

In summary, these discrepancies occurred because (a) Japan and the IEA treat international aviation and marine bunker fuels differently in their respective energy balances and (b) because of the different classifications of fuel oil A. The IEA energy balances include fuel consumption by international flights and international marine; whereas the energy balances of Japan do not include them as these are not regarded as domestic consumption. Consequently, the data for the bonded exports and imports of jet kerosene and fuel oil C are differently accounted for. With respect to fuel oil A, Japan includes it under Residual Fuel Oil in its energy balances but reports it to the IEA under Gas/Diesel Oil according to the classifications used in Europe and the United States.

According to Japanese definition, fuel oil A has a flash point of more than 60 °C, kinematic viscosity of 20 mm²/s below, carbon residue content of 4% below and sulfur content of 2.0 % below. Fuel oil B has a flash point of more than 60 °C, kinematic viscosity of 50 mm²/s below, carbon residue content of 8% below and sulfur content of 3.0 % below. Fuel oil B is rarely used nowadays in Japan, for this reason, fuel oil B is treated as “fuel oil B/C” together with fuel oil C in Japanese statistics. Fuel oil C has a flash point of more than 70 °C, kinematic viscosity of less than 1,000 mm²/s and sulfur content of less than 3.5%.

Further explanations are provided below for each of the discrepancies noted by the ERT.

a) Differences in exports of jet kerosene and residual fuel oil

<ERT findings on FCCC/ARR/2006/JPN>

Exports of liquid fuels are between 40 and 70 per cent lower in the IEA data; the differences are due in particular to differences in the figures for jet kerosene and residual fuel oil, with the largest errors occurring in recent years.

<Explanation 1: Exports of jet kerosene>

The figures for jet kerosene exports reported in the CRF tables are different from those in the IEA statistics because the CRF figures include bonded exports whereas the export figures in the IEA statistics do not. The IEA statistics accounted the final consumption of jet kerosene by international aviation as an aggregate of the bonded exports and imports. (See Chapter 3, for bonded exports and imports.)

<Reference: Exports of jet kerosene in 2009>

CRF Table 1.A(b)	IEA Statistics
Exports: $8,333.93 \times 10^3$ kl	Exports: $1,967 \times 10^3$ t [$2,512.55 \times 10^3$ kl (exports excluding bonded exports) $\times 0.7834$ (specific gravity) = $1,968 \times 10^3$ t]
<Breakdown> Exports excluding bonded exports: $2,512.55 \times 10^3$ kl Bonded exports: $5,821.38 \times 10^3$ kl	<Remarks 1> Because each exported amount per destination country is rounded off before aggregation in the IEA statistics, the aggregation sometimes differ slightly from the multiplication product of the total exported amount and the specific gravity. <Remarks 2> International aviation: $4,898 \times 10^3$ t [$5,821.38 \times 10^3$ kl (bonded exports) + 430.268×10^3 kl (bonded imports)* = $6,251.648 \times 10^3$ kl; $6,251.648 \times 10^3$ kl $\times 0.7834$ (specific gravity) = $4,898 \times 10^3$ t]

<Explanation 2: Exports of residual fuel oil>

The figures for exports of residual fuel oil reported in the CRF tables are different from those in the IEA statistics because the CRF figures for residual fuel oil include the bonded exports, whereas the export figures for heavy fuel oil in the IEA statistics do not. The bonded exports portion of the heavy fuel oil was reported in the IEA statistics as an aggregate of the bonded exports and imports of heavy fuel oil under International Marine Bunkers. (See Chapter 3, for bonded exports and imports.)

Further, the figures for exports of residual fuel oil reported in the CRF include fuel oil A, whereas the figures reported under Heavy Fuel Oil in the IEA statistics do not. The IEA reports fuel oil A together with gas oil under Gas/Diesel Oil in its statistics. Because fuel oil A, which is treated as a fuel oil that is distinguished from diesel oil in Japan, is grouped together with diesel oil in Europe and the United States, the fuel oil A data have been included in the diesel oil data in Japan's report to the IEA.

<Reference: Exports of residual fuel oil in 2009>

CRF Table 1.A(b)	IEA Statistics/ Heavy Fuel oil
Exports: $8,406.47 \times 10^3$ kl [608.07 $\times 10^3$ kl (fuel oil A) + 7,798.40 $\times 10^3$ kl (fuel oils B and C) = 8,406.47 $\times 10^3$ kl]	Exports: $2,623 \times 10^3$ t [2,914.59 $\times 10^3$ kl (exports of fuel oils B and C excluding bonded exports) $\times 0.9$ (specific gravity) = 2,623 $\times 10^3$ t]
<Breakdown> Exports of fuel oil A: 608.07×10^3 kl Exports excluding bonded exports: 485.84 $\times 10^3$ kl Bonded exports: 122.23×10^3 kl Exports of fuel oils B and C: 7,798.40 $\times 10^3$ kl Exports excluding bonded exports: 2,914.59 $\times 10^3$ kl Bonded exports: $4,883.81 \times 10^3$ kl	<Remarks> International marine bunkers: $4,509 \times 10^3$ t [4,883.81 $\times 10^3$ kl (bonded exports of fuel oils B and C) + 126.42 $\times 10^3$ kl (bonded imports of fuel oils B and C) = 5,010.23 $\times 10^3$ kl; 5,010.23 $\times 10^3$ kl $\times 0.9$ (specific gravity) = 4,509 $\times 10^3$ t]

b) Differences in imports of jet kerosene and gas/diesel oil

<ERT findings on FCCC/ARR/2006/JPN>

Imports of jet kerosene have been reported to the IEA, but are shown as zero in the CRFs for the years 1990–1997, while imports of gas/diesel oil are systematically about 80 per cent lower in the CRF tables than in the IEA figures.

<Explanation 1: Imports of jet kerosene>

The figures for jet kerosene imports reported in the CRF tables are different from those in the IEA statistics because the CRF figures are the sums of imports including bonded imports and bonded exports while the IEA statistics figures are only the imports including bonded imports. (See Chapter 3, for bonded exports and imports.)

<Reference: Jet kerosene imports in 2009>

CRF Table 1.A(b)	IEA Statistics
Imports: $6,251.65 \times 10^3$ kl <Jet kerosene imports> Imports: $6,251.65 \times 10^3$ kl Imports excluding bonded imports: 0 Bonded imports: 430.27×10^3 kl Bonded exports: $5,821.38 \times 10^3$ kl	Imports: 337×10^3 t [430.27 $\times 10^3$ kl (imports including bonded imports) $\times 0.7834$ (specific gravity) = 337 $\times 10^3$ t]

<Explanation 2: Imports of gas/diesel oil>

The figures for imports of gas/diesel oil reported in the CRF tables are different from those in the IEA statistics, because the CRF figures are the sums of imports (including bonded imports) and bonded

exports of diesel oil, which excludes fuel oil A, while the figures for imports of gas/diesel oil in the IEA statistics are the aggregate of imports of diesel oil and fuel oil A, both of which included the bonded imports.

<Reference: Imports of gas/diesel oil in 2009>

CRF Table 1.A(b)	IEA Statistics
Imports: 317.33×10^3 kl <Imports of gas/diesel oil> Imports excluding bonded imports: 310.11×10^3 kl Bonded imports: 0 Bonded exports: 7.22×10^3 kl	Imports: 325×10^3 t [310.11×10^3 kl (imports of gas/diesel oil including bonded imports) + 75.73×10^3 kl (imports of fuel oil A including bonded imports) = 385.84×10^3 kl; 385.84×10^3 kl \times 0.843 (specific gravity) = 325×10^3 t]

c) Differences in imports of coking coal

<ERT findings on FCCC/ARR/2006/JPN>

Furthermore, the figures for imports of coking coal are systematically lower in the CRF tables than those in the IEA statistics, with the largest discrepancy occurring in 1999.

<Explanation: Imports of coking coal>

The coking coal imports in the CRF is slightly different from that in the IEA statistics, because the imported amount of coking coal for coke is reported in the CRF, while the estimated figure for the imported coking coal by the weighted average for calorific value of coking coal and PCI coal is reported in the IEA statistics. The reason of such treatment is that the coking coal and PCI coal are separately calculated in the estimation of CO₂ emission reported in the CRF, while they are not separated in the IEA statistics.

<Reference: Imports of coking coal in 2009>

CRF Table 1.A(b)	IEA Statistics
Imports: $52,334.58 \times 10^3$ t <Remarks> Imports of coking coal: $52,334.58 \times 10^3$ t Imports of PCI coal: 0	Imports: $52,514 \times 10^3$ t [Imports of coking coal: $52,334 \times 10^3$ t \times 29.10 GJ/t ⁽¹⁾ / 29.00 GJ/t ⁽²⁾ = $52,514 \times 10^3$ t] (1) Calorific value of coking coal (2) Weighted average calorific value of coking coal and PCI coal

d) Differences in stock changes in liquid and gaseous fuels

<ERT findings on FCCC/ARR/2006/JPN>

In addition, the data on stock changes are not consistent for liquid and gaseous fuels.

<Explanation 1: Changes in crude oil stock>

The difference between the CRF table and the IEA statistics with respect to changes in crude oil stock occurred because the figures reported in the CRF were calculated using the stock of crude oil after customs clearance (or more precisely, after inspection in the presence of customs officers). The stock changes reported in the IEA statistics were calculated based on stock that included crude oil carried by oil tankers in Japanese territorial waters but which was yet to clear customs as well as the crude oil in the national stockpile. This discrepancy arose because the UNFCCC and the IEA had different objectives.

Also, the plus-minus signs of stock changes in the CRF differ from those of the IEA, because the changes in the CRF are defined as plus for stock increase and as minus for stock release, while the changes in the IEA are defined as minus for stock increase and as plus for stock release.

<Reference: Changes of crude oil stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: - 1,605.05× 10 ³ kl	Stock changes: 1,566× 10 ³ t

<Explanation 2: Changes in NGL stock>

Stock changes concerning NGL were reported in the CRF. The NGL stock changes reported in the IEA statistics were zero because the NGL stock figure in the Monthly Oil Statistics (MOS) of the IEA was zero. This discrepancy resulted from the direction given by the IEA that the figures in the IEA statistics must be consistent with the MOS figures.

Furthermore, the figures for “stock changes” required by the CRF tables are not included in the MOS. On the other hand, the MOS requires figures for Opening Stock and Closing Stock, but Japan does not collect such statistical data for NGL. As a result, Japan reported zero values to the IEA for both Opening Stock and Closing Stock data for the MOS. In light of the fact that no statistical data exists for stock changes in NGL, even though the stock actually existed, with respect to the CRF tables the estimated value is reported.

<Reference: Changes in NGL stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: 114.33× 10 ³ kl	Stock changes: 0

<Explanation 3: Changes in gasoline stock>

The figures for changes in gasoline stock reported in the CRF tables are the same as the figures in the IEA statistics. The changes in gasoline stock in the CRF correspond to the stock changes in motor gasoline and in white spirit of the IEA statistics.

<Reference: Changes in gasoline stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: -353.33×10^3 kl	Stock changes in motor gasoline: 255×10^3 t $[353.33 \times 10^3 \text{ kl} \times 0.737 \text{ (specific gravity)}]$ $= 260 \times 10^3$ t 260×10^3 t - (Stock changes in white spirit) $= 255 \times 10^3$ t] Stock changes in white spirit: 5×10^3 t $[6.42 \times 10^3 \text{ kl} \times 0.737 \text{ (specific gravity)}]$ $= 5 \times 10^3$ t]

<Explanation 4: Changes in jet kerosene stock>

The figures for changes in jet kerosene stock reported in the CRF tables are the same as the figures in the IEA statistics.

<Reference: Changes in jet kerosene stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: 23.92×10^3 kl	Stock changes: -19×10^3 t $[-23.92 \times 10^3 \text{ kl} \times 0.7834 \text{ (specific gravity)}]$ $= -19 \times 10^3$ t]

<Explanation 5: Changes in kerosene stock>

The figures for changes in kerosene stock reported in the CRF tables are the same as the figures in the IEA statistics.

<Reference: Changes in kerosene stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: -168.94×10^3 kl	Stock changes: 137×10^3 t $[168.938 \times 10^3 \text{ kl} \times 0.814 \text{ (specific gravity)}]$ $= 137 \times 10^3$ t]

<Explanation 6: Changes in gas/diesel oil stock>

The figures for gas/diesel stock reported in the CRF tables were different from those in the IEA statistics because the CRF figures did not include stock changes in fuel oil A while the IEA statistics did.

<Reference: Changes in gas/diesel oil stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: -237.87×10^3 kl	Stock changes: 334×10^3 t $[237.87 \times 10^3 \text{ kl} \times 0.843 \text{ (specific gravity)}$ $= 201 \times 10^3 \text{ t (stock changes in gas/diesel oil);}$ $158.10 \times 10^3 \text{ kl} \times 0.843 \text{ (specific gravity)}$ $= 133 \times 10^3 \text{ t (stock changes in fuel oil A);}$ $201 \times 10^3 \text{ t} + 133 \times 10^3 \text{ t} = 334 \times 10^3 \text{ t}]$

<Explanation 7: Changes in residual fuel oil stock>

The figures for residual fuel oil stock reported in the CRF tables were different from those in the IEA statistics because the CRF figures included changes in fuel oil A stock, whereas stock change data under Heavy Fuel Oil in the IEA statistics did not include fuel oil A. (See the explanation for the gas/diesel oil data above.)

<Reference: Changes in residual fuel oil stock in 2009>

CRF Table 1.A(b)	IEA Statistics/Heavy Fuel oil
Stock changes: -69.69×10^3 kl <Breakdown> Stock changes in fuel oil A: -158.10×10^3 kl Stock changes in fuel oil C: 88.41×10^3 kl	Stock changes: -80×10^3 t $[-88.41 \times 10^3 \text{ kl (stock changes in fuel oil C)}$ $\times 0.900 \text{ (specific gravity)} = -80 \times 10^3 \text{ t}]$

<Explanation 8: Changes in LPG stock>

The figures for changes in LPG stock reported in the CRF tables are the same as the figures in the IEA statistics.

<Reference: Changes in LPG stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: -440.02×10^3 t	Stock changes: 440×10^3 t

<Explanation 9: Changes in naphtha stock>

The figures for changes in naphtha stock reported in the CRF tables are the same as the figures in the IEA statistics.

<Reference: Changes in naphtha stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: - 421.88× 10 ³ kl	Stock changes: 311× 10 ³ t [421.88× 10 ³ kl × 0.737 (specific gravity) = 311× 10 ³ t]
	<Remarks> The amount of stock changes was 294× 10 ³ t in “Energy Statistics of OECD Countries 2008-2009, 2011 Edition, OECD/IEA”. After this edition, this amount was revised to 421.88× 10 ³ kl (311× 10 ³ t) due to the revision of statistic value in naphtha stock, and this revision was reported on the annual report to the IEA.

<Explanation 10: Changes in bitumen stock>

The figures for changes in bitumen stock reported in the CRF tables were slightly different from the figures reported under Bitumen in the IEA statistics because the Bitumen data in the CRF tables included asphalt and other heavy oil and paraffin products. The IEA statistics reported figures for only asphalt under Bitumen, and the figures for other heavy oil and paraffin products reported in the CRF tables under Bitumen were included in the figures reported under Paraffin Waxes in the IEA statistics.

<Reference: Changes in bitumen stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: -20.64× 10 ³ t <Breakdown> Asphalt: - 19.87× 10 ³ t Other fuel oils and paraffin products: - 0.77× 10 ³ t	Stock changes in bitumen: 20× 10 ³ t
	<Remarks> In the IEA statistics, the figures for other <i>heavy oil</i> and paraffin products (which were reported under Bitumen in the CRF tables) are reported under Paraffin Waxes.

<Explanation 11: Changes in lubricants stock>

The figures for changes in lubricants stock reported in the CRF tables are the same as the figures in the IEA statistics.

<Reference: Changes in lubricating oil stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: -50.99× 10 ³ kl	Stock changes: 45× 10 ³ t [50.989× 10 ³ kl × 0.891 (specific gravity) = 45× 10 ³ t]

<Explanation 12: Changes in oil coke stock>

The figures for changes in oil coke stock reported in the CRF tables are the same as the figures in the IEA statistics.

<Reference: Changes in oil coke stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: -1.36×10^3 t	Stock changes: 2×10^3 t [18.561 $\times 10^3$ t (stocks at the end of March 2009) - 17.201 $\times 10^3$ t (stocks at the end of March 2010) = 1.36 $\times 10^3$ t] <Remarks> The above calculated value differs from the IEA published value, because when inputting on the IEA reporting file the stock changes (or the differences) are calculated after the stock amount at the end of each month is rounded off due to file formatting by the IEA,.

<Explanation 13: Changes in refinery feedstock stock>

The figures for changes in refinery feedstock stock reported in the CRF were different from those in the IEA statistics because the IEA statistics included the figures for stock changes in slack wax and slack coke in addition to the semi-refined products reported in the CRF tables.

The changes in slack wax and coke stocks were not reported in the CRF tables because the both items were solids used as raw materials for the production of paraffin and oil coke, and unlikely to be returned to oil refining processes. In addition, shipments of paraffin and oil coke produced using slack wax and slack coke were separately accounted for.

<Reference: Changes in refinery feedstock stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Stock changes: $- 902.90 \times 10^3$ kl <Breakdown> Slack gasoline: $- 450.89 \times 10^3$ kl Slack kerosene: 19.31×10^3 kl Slack diesel oil or gas oil: $- 254.93 \times 10^3$ kl Slack fuel oil: $- 216.38 \times 10^3$ kl (Slack fuel oil is the aggregate of $- 101.33 \times 10^3$ kl for slack fuel oil and $- 115.06 \times 10^3$ kl for slack lubricant)	Stock changes: 737×10^3 t <Breakdown> Slack gasoline: 450.89×10^3 kl Slack kerosene: $- 19.31 \times 10^3$ kl Slack diesel oil or gas oil: 254.93×10^3 kl Slack fuel oil: 101.33×10^3 kl Slack lubricant: 115.06×10^3 kl Slack wax: 6.36×10^3 kl Slack coke: 5.88×10^3 kl Each of the above figures is multiplied by its specific gravity for conversion to weight for reporting purposes.
<Remarks> The stock changes are different in some years between the CRF tables and the IEA statistics because of the differences between monthly statistics and yearly statistics. The figures for the supply and stock of oil in the IEA statistics use the figures in the Monthly Oil Statistics compiled by the IEA. The report to the IEA for the MOS is submitted on a monthly basis. The monthly data may be adjusted for the yearly statistics. The CRF tables report annual data.	

<Explanation 14: Changes in natural gas stock>

The figures for changes in natural gas stock (imported LNG and domestic natural gas) reported in the CRF tables were different from those in the IEA statistics because of the differences in the methods used for estimation of changes in the imported LNG stock. Although the same figure for the domestic natural gas stock was reported in the CRF and the IEA statistics because the statistical data existed in Japan, however the data were estimated for imported LNG because the statistics do not catch whole stocks.

The figures for changes in LNG stock reported in the CRF tables were estimated as the difference between the LNG imports and the consumption. The figures for stock changes reported to the IEA were the difference between the stock of imported LNG at the end of the previous year and the stock at the end of the current year, with the former calculated as one-half of the LNG import in March of the previous year, and the latter as one-half of the LNG import in March of the current year.

<Reference: Changes in natural gas stock in 2009>

CRF Table 1.A(b)	IEA Statistics
Changes in LNG stock: $- 3,612 \times 10^3$ t Changes in domestic natural gas stock: 15.63×10^6 m ³	Stock changes: $- 24,574$ TJ (GCV) <Remarks> The figures for LNG and natural gas were combined under Natural Gas as the IEA statistics do not separate them.

A2.2. General Energy Statistics

A2.2.1. General Energy Statistics Overview

The data given in the *General Energy Statistics* compiled by the Agency for Natural Resources and Energy were used for the activity data of fuel combustion in energy sector.

The *General Energy Statistics* (Energy Balance Table) provides a comprehensive overview of domestic energy supply and demand to grasp what are converted from energy sources, such as coal, oil, natural gas and others, provided in Japan and what are consumed in what sectors. The supply/conversion and consumption data in *General Energy Statistics* use official statistics and are structured with the minimum of estimation and adjustment.

General Energy Statistics (Energy Balance Table) indicates an overview of domestic energy supply and demand, shows the main energy sources used in Japan as “Columns” and the supply, conversion and consumption sectors as “Rows”, in a matrix. Specifically, columns comprise 11 major categories (coal [code \$100], coal products [code \$150], oil [code \$200], oil products [code \$250], natural gas [code \$400], town gas [code \$450], new and renewable energy [code \$500], large-scale hydropower [code \$550], nuclear power [code \$600], electricity [code \$700], and heat [code \$800]) and the necessary sub-categories and a more detailed breakdown of the sub-categories. The *General Energy Statistics* supply and demand sectors (rows) comprise 3 major sectors — primary energy supply (primary supply) [code #1000], energy conversion (conversion) [code #2000], and final energy consumption (final consumption) [code #5000] — plus the necessary sub-categories and a more detailed breakdown of the sub-categories. (Refer to the following General Energy Statistics simplified table.)

The *General Energy Statistics* (complete Energy Balance Tables) for the years since FY 1990 are available on the following internet site:

<http://www.enecho.meti.go.jp/info/statistics/jukyu/result-2.htm>

The following is the energy balance simplified table (Table A 2-1 - Table A 2-5).

Table A 2-1 Energy balance simplified table (General Energy Statistics, FY1990)

1990FY	Code	100	150	200	250	400	450	500	550	600	700	800	900	910	920	
<Energy balance simplified table>		Coal	Coal Product	Oil	Oil Products	Natural Gas	Town Gas	Renewable E	Hydraulic	Nuclear Ener	Electricity	Heat	Total	Energy Total	Non-Energy	
<<Energy units>>		TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	
Code																
1000	Primary Energy Supply	3345244	15352	9164033	2354044	2059168	0	524099	833304	1887390	0	0	20182635	18632722	1549913	
1100	Indigenous Production	187036	0	24484	0	89203	0	524099	833304	1887390	0	0	3545517	0	0	
1200	Import	3158208	15352	9139549	2354044	1969965	0	0	0	0	0	0	16637118	0	0	
1500	TPES Total Primary Energy Supply	3345244	15352	9164033	2354044	2059168	0	524099	833304	1887390	0	0	20182635	18632722	1549913	
1600	Export	-53	-56644	0	-302130	0	0	0	0	0	0	0	-358828	0	0	
1700	Stockpile Change	1689	1951	-190171	-22710	42651	0	0	0	0	0	0	-166610	0	0	
1900	DPEs Domestic Primary Energy Supply	3346859	-39341	8973862	2029203	2101819	0	524099	833304	1887390	0	0	19657197	18107284	1549913	
													consumption side	19785779	18235866	1549913
2000	Energy Transformation & Own use	-3039243	1595040	-9032036	5785908	-2039503	629852	-470769	-833304	-1887390	2698536	696058	-589853	-5865031	-31822	
2100	Power Generation	-673045	-204274	-874209	-1055765	-1531630	0	-19259	-767173	-1879280	2691329	0	-4313307	-4313307	0	
2200	Auto Power Generation	-116820	-96004	0	-399646	-5054	-12280	-170874	-66131	-8110	304022	0	-570897	-570897	0	
2300	Industrial Steam Generation	-123177	-69991	0	-444065	-2693	-15028	-278052	0	0	784558	0	-148448	-148448	0	
2350	District Heat Supply	-824	0	0	-2633	0	-6169	-2028	0	0	8464	0	-4419	-4419	0	
2400	Town Gas Production	0	-19178	0	-142210	-503865	664661	-546	0	0	0	0	-1139	-1139	0	
2500	Coal Products	-2142396	2081208	0	-38206	0	0	0	0	0	0	0	-99394	-99394	0	
2600	Oil Products	0	0	-8143167	8175884	5121	0	0	0	0	0	-94149	-56212	0	-56212	
2700	Other Conversions & Blending	30171	2880	0	-18897	0	18897	0	0	0	0	0	33051	0	33051	
2800	TC Total Conversion	-3026090	1694639	-9017376	6074562	-2038122	650081	-470758	-833304	-1887390	2994122	698872	-5160764	-5137603	-23161	
2900	Own Use & Loss	-3015	-101777	-1017	-301251	-1738	-20230	0	0	0	-295586	-2814	-727428	-727428	0	
3000	OI Other Input/Output	0	0	0	12924	0	0	0	0	0	0	0	12924	0	12924	
3500	FS Stock Change	-10138	2177	-13642	-327	357	0	-10	0	0	0	0	-21584	0	-21584	
4000	DC Stastical Discrepancy	-75007	0	-58202	3856	769	0	0	0	0	2	0	-128582	-128582	0	
5000	Final Energy Consumption	382623	1555699	28	7811256	61547	629852	53330	0	0	2698534	696058	13888926	12370836	1518091	
6000	Industry	365162	1532019	28	3019423	57690	110593	0	0	0	1220265	687697	6892876	5516717	1476159	
6100	NMFC Non-Manufacturing	263	1141	28	759211	3757	20677	0	0	0	21251	0	806329	553571	252758	
6500	MFC Manufacturing	364899	1530877	0	2260212	53933	88916	0	0	0	1199013	687697	6186547	4963145	1223401	
6520	Pulp & Paper	126	0	0	27726	2	1272	0	0	0	121360	249523	400009	400009	0	
6550	Chemical	5443	46803	0	1356286	26599	1028	0	0	0	186050	185545	1807754	670574	1137180	
6570	Cement & Ceramics	235223	40381	0	104386	20	743	0	0	0	79708	6706	467168	456544	10624	
6580	Iron & Steel	143931	1103634	0	119268	25030	8746	0	0	0	265486	92916	1759011	1758326	685	
6600	Machinery	15	16700	0	85879	2132	22135	0	0	0	212915	0	339776	339776	0	
6700	Duplication Adjustment	-36513	-8421	0	-56803	-3000	-2137	0	0	0	-49573	-22295	-178742	-169225	-9517	
6900	Other Industries & SMEs	1164	320931	0	354525	2014	31396	0	0	0	235503	121650	1067184	982755	84429	
7000	ResCom	17461	23680	0	1634972	3857	519258	53330	0	0	1417755	8361	3678676	3677496	1180	
7100	RES Residential	0	2880	0	594332	0	342157	51488	0	0	662933	1284	1655075	1655075	0	
7150	HokkaidoTohoku,Hokuriku	0	0	0	214484	0	14146	0	0	0	104048	0	359948	359948	0	
7180	Kantou,Tokai,Kansai	0	0	0	285397	0	337114	0	0	0	416516	0	1039026	1039026	0	
7170	Chuugoku,Shikoku,Kyushu,Okinawa	0	0	0	119753	0	48044	0	0	0	143216	0	311012	311012	0	
7500	COM Commercial & Others	17461	20801	0	1040640	3857	177101	1842	0	0	754822	7077	2023801	2022421	1180	
7510	Water supply, Sewage & Waste Dis	262	0	0	73615	0	3295	0	0	0	67696	4	144872	144872	0	
7540	Telecommunication & Broadcasting	0	0	0	9009	0	2257	0	0	0	19005	395	30666	30666	0	
7800	Trade & Finance Service	0	0	0	259263	0	25973	0	0	0	188251	2656	476143	476143	0	
7700	Public Service	12038	0	0	274167	0	49255	0	0	0	214702	1346	551508	551508	0	
7810	Commercial Service	235	261	0	97285	0	4358	0	0	0	55712	413	158265	158265	0	
7850	Retail Service	2406	1906	0	219818	0	67360	0	0	0	135481	1576	428547	428547	0	
8000	Transportation	0	0	0	3156861	0	0	0	0	0	60514	0	3217375	3176823	40752	
8100	PAS Passenger	0	0	0	1614051	0	0	0	0	0	56610	0	1670661	1638859	31802	
8110	Car	0	0	0	1375786	0	0	0	0	0	0	0	1375786	1344140	31646	
8120	Rail	0	0	0	11264	0	0	0	0	0	56610	0	67874	67718	156	
8130	Ship	0	0	0	67628	0	0	0	0	0	0	0	67628	67628	0	
8140	Air	0	0	0	88429	0	0	0	0	0	0	0	88429	88429	0	
8500	FRT Freight	0	0	0	1542810	0	0	0	0	0	3905	0	1546714	1537764	8950	
8510	Truck & Lorry	0	0	0	1391105	0	0	0	0	0	0	0	1391105	1386473	4632	
8520	Rail	0	0	0	2638	0	0	0	0	0	3905	0	6543	6374	169	
8530	Ship	0	0	0	130812	0	0	0	0	0	0	0	130812	126662	4149	
8540	Air	0	0	0	18256	0	0	0	0	0	0	0	18256	18256	0	
9000	FEEC Final Energy Consumption	382112	1538556	28	6324859	47544	629814	53330	0	0	2698534	696058	12370836	12370836	0	
9500	Non-Energy	511	17143	0	1486397	14003	38	0	0	0	0	0	1518091	0	1518091	
9600	Industry	511	17143	0	1444465	14003	38	0	0	0	0	0	1476159	0	1476159	
9800	ResCom & others	0	0	0	1180	0	0	0	0	0	0	0	1180	0	1180	
9850	Transport	0	0	0	40752	0	0	0	0	0	0	0	40752	0	40752	

Table A 2-2 Energy balance simplified table (General Energy Statistics, FY1995)

1995FY	Code	100	150	200	250	400	450	500	550	600	700	800	900	910	920	
(Energy balance simplified table)		Coal	Coal	Product Oil	Oil Products	Natural Gas	Town Gas	Renewable E/Hydraulic	Nuclear	Electricity	Heat		Total	Energy Total	Non-Energy	
<<(Energy units)>>		TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	
Code																
1000	Primary Energy Supply	3732254	18016	10204290	2225292	2479453	0	564207	761329	2700257	0	0	22665097	20955245	1729852	
1100	Indigenous Production	149495	0	32455	0	95250	0	564207	761329	2700257	0	0	4302993	0	0	
1200	Import	3582759	18016	10171835	2225292	2384203	0	0	0	0	0	0	18382105	0	0	
1500	TPES Total Primary Energy Supply	3732254	18016	10204290	2225292	2479453	0	564207	761329	2700257	0	0	22665097	20955245	1729852	
1600	Export	-75	-103811	0	-733696	0	0	0	0	0	0	0	-837582	0	0	
1700	Stockpile Change	-2710	-6113	-30486	134344	58576	0	0	0	0	0	0	153611	0	0	
1900	DPES Domestic Primary Energy Supply	3729468	-81908	10173804	1625939	2538029	0	564207	761329	2700257	0	0	22001126	20271274	1729852	
													consumption side	21947773	20217921	1729852
2000	Energy Transformation & Own use	-3286798	1395073	-10108952	7217819	-2474669	823061	-518878	-761329	-2700257	3090955	694292	-6829583	-6626513	-3070	
2100	Power Genertion	-1072304	-210723	-669401	-838649	-1750818	0	-36870	-700065	-2687729	3071160	0	-4895399	-4895399	0	
2200	Auto Power Generation	-150687	-115758	-880	-459430	-5691	-32050	-199357	-61264	-12528	364710	0	-672935	-672935	0	
2300	Industrial Steam Generation	-133278	-60234	-328	-446810	-2879	-30180	-278056	0	0	784719	0	-167044	-167044	0	
2350	District Heat Supply	-638	0	0	-1638	0	-11101	-4577	0	0	-2548	16423	-4079	-4079	0	
2400	Town Gas Production	0	-12205	0	-157821	-723643	892307	-37	0	0	0	0	-1400	-1400	0	
2500	Coal Products	-1963775	1893360	0	-30083	0	0	0	0	0	0	0	-100498	-100498	0	
2600	Oil Products	0	0	-9421404	9490043	5773	0	0	0	0	0	-103260	-28847	-0	-28847	
2700	Other Conversions & Blending	36411	1637	0	-22539	0	22539	0	0	0	0	0	38047	0	38047	
2800	TC Total Conversion	-3284272	1496077	-10082012	7533073	-2477258	841515	-518897	-761329	-2700257	3433322	697882	-5832154	-5841355	9200	
2900	Own Use & Loss	-2978	-93780	-1058	-321669	-1261	-18454	0	0	0	-342367	-3590	-785158	-785158	0	
3000	OI Other Input/Output	0	0	0	9078	0	0	0	0	0	0	0	9078	0	9078	
3500	FS Stock Change	452	-7224	-15882	-2563	3650	0	19	0	0	0	0	-21348	0	-21348	
4000	DC Statistical Discrepancy	-7652	0	64852	-8469	4822	0	-0	0	0	0	0	53353	53353	0	
5000	Final Energy Consumption	450322	1303165	0	8852328	58738	823061	45329	0	0	3090955	694292	15318190	13591408	1726782	
6000	Industry	428876	1299570	0	3267149	56329	163883	36	0	0	1269782	678469	7164096	5471642	1692454	
6100	NMFC Non-Manufacturing	191	528	0	735650	1776	26151	0	0	0	20464	0	784760	557911	226848	
6500	MFG Manufacturing	428685	1299042	0	2531499	54553	137733	36	0	0	1249318	678469	6378336	4913731	1465605	
6520	Pulp & Paper	0	0	0	30072	5	5747	36	0	0	126598	246261	408718	408718	0	
6550	Chemical	6176	34647	0	1705864	21627	6650	0	0	0	189040	193896	2167901	799104	1368797	
6570	Cement & Ceramics	235274	37704	0	118517	341	628	0	0	0	84884	8266	485615	475539	10076	
6580	Iron & Steel	201778	958301	0	114033	26245	20866	0	0	0	255475	94083	1670781	1670574	208	
6600	Machinery	4	14083	0	89461	3476	32517	0	0	0	236790	0	376331	376331	0	
6700	Duplication Adjustment	-26421	-5593	0	-81902	-1529	-3384	0	0	0	-49200	-20224	-188251	-182224	-6028	
6900	Other Industries & SMEs	1841	250502	0	261747	2808	40947	0	0	0	244492	104443	906581	814028	92553	
7000	ResCom	21446	3594	0	1846240	2409	659177	45293	0	0	1753655	15823	4347637	4345809	1828	
7100	RES Residential	0	1637	0	700079	0	398516	43786	0	0	827334	1368	1972720	1972720	0	
7150	Hokkaido	0	0	0	245943	0	46561	0	0	0	135963	0	428467	428467	0	
7160	Tohoku	0	0	0	330756	0	367163	0	0	0	541005	0	1238924	1238924	0	
7170	Kantou, Toukai, Kansai	0	0	0	145241	0	50323	0	0	0	187224	0	382789	382789	0	
7170	Chuugoku,Shikoku,Kyushu,Okinaawa	0	0	0	145241	0	50323	0	0	0	187224	0	382789	382789	0	
7500	COM Commercial & Others	21446	1958	0	1146160	2409	260662	1507	0	0	926321	14455	2374918	2373090	1828	
7510	Water supply, Sogawa & Waste Dis	426	0	0	113365	0	4750	0	0	0	63661	8	182210	182210	0	
7540	Telecommunication & Broadcasting	0	0	0	10732	0	2438	0	0	0	22209	384	35764	35764	0	
7600	Trade & Finance Service	0	0	0	269831	0	40343	0	0	0	201589	6828	518593	518593	0	
7700	Public Service	16599	0	0	364499	0	75063	0	0	0	277487	1960	735608	735608	0	
7810	Commercial Service	330	254	0	98680	0	5580	0	0	0	66005	587	171435	171435	0	
7850	Retail Service	3820	1682	0	261577	0	154955	0	0	0	160825	3268	586127	586127	0	
8000	Transportation	0	0	0	3738939	0	0	0	0	0	67518	0	3806457	3773957	32500	
8100	PAS Passenger	0	0	0	2044897	0	0	0	0	0	63676	0	2108573	2083686	24887	
8110	Car	0	0	0	1787886	0	0	0	0	0	0	0	1787886	1762915	24771	
8120	Rail	0	0	0	9759	0	0	0	0	0	63676	0	73435	73319	116	
8130	Ship	0	0	0	79258	0	0	0	0	0	0	0	79258	79258	0	
8140	Air	0	0	0	128698	0	0	0	0	0	0	0	128698	128698	0	
8500	FRT Freight	0	0	0	1694042	0	0	0	0	0	3842	0	1697884	1690271	7613	
8510	Truck & Lorry	0	0	0	1566432	0	0	0	0	0	0	0	1566432	1562452	3980	
8520	Rail	0	0	0	2400	0	0	0	0	0	3842	0	6242	6130	112	
8530	Ship	0	0	0	131840	0	0	0	0	0	0	0	131840	128319	3521	
8540	Air	0	0	0	24397	0	0	0	0	0	0	0	24397	24397	0	
9000	FEEC Final Energy Consumption	449885	1291322	0	7149862	46702	823061	45329	0	0	3090955	694292	13591408	13591408	0	
9500	Non-Energy	437	11843	0	1702466	12036	0	0	0	0	0	0	1726782	0	1726782	
9800	Industry	437	11843	0	1668138	12036	0	0	0	0	0	0	1692454	0	1692454	
9800	ResCom & others	0	0	0	1828	0	0	0	0	0	0	0	1828	0	1828	
9850	Transport	0	0	0	32500	0	0	0	0	0	0	0	32500	0	32500	

Table A 2-3 Energy balance simplified table (General Energy Statistics, FY2000)

2000FY	Code	100	150	200	250	400	450	500	550	800	700	800	900	910	920	
<Energy balance simplified table>		Coal	Coal Product	Oil	Natural Gas	Town Gas	Renewable E	Hydraulic	Nuclear Ener	Electricity	Heat		Total	Energy Total	Non-Energy	
<<Energy units>>		TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	
Code																
1000	Primary Energy Supply	4210040	76219	9761365	2246246	3060666	0	616335	778417	2873130	0	0	23622418	21719570	1902848	
1100	Indigenous Production	66013	0	28034	0	106340	0	616335	778417	2873130	0	0	4468269	0	0	
1200	Import	4144027	76219	9733330	2246246	2954327	0	0	0	0	0	0	19154149	0	0	
1500	TPES Total Primary Energy Supply	4210040	76219	9761365	2246246	3060666	0	616335	778417	2873130	0	0	23622418	21719570	1902848	
1600	Export	-112	-78077	0	-627862	0	0	0	0	0	0	0	-706051	0	0	
1700	Stockpile Change	-2958	-1963	-116285	-106335	72387	0	0	0	0	0	0	-155155	0	0	
1900	DPEs Domestic Primary Energy Supply	4206970	-3821	9645079	1512049	3133054	0	616335	778417	2873130	0	0	22761213	20858365	1902848	
													consumption side	22790985	20888136	1902848
2000	Energy Transformation & Own use	-3738666	1287540	-9721175	7518258	-3072804	986782	-562115	-778417	-2873130	3396151	739685	-6815890	-6685083	-130807	
2100	Power Generation	-1515218	-212244	-301245	-548677	-2131672	-1447	-46226	-711603	-2866777	3332394	0	-5001815	-5001815	0	
2200	Auto Power Generation	-199734	-148205	-99	-425144	-9644	-38900	-211258	-66814	-6353	423092	0	-683058	-683058	0	
2300	Industrial Steam Generation	-191460	-34306	-119	-428955	-6884	-30434	-298304	0	0	0	857666	-132897	-132897	0	
2350	District Heat Supply	-708	0	0	-1725	0	-14515	-6275	0	0	-3940	23428	-3735	-3735	0	
2400	Town Gas Production	0	-9573	0	-126581	-925315	1061122	-31	0	0	0	0	-377	-377	0	
2500	Coal Products	-1816696	1790538	0	-39481	0	0	0	0	0	0	0	-65640	-65640	0	
2600	Oil Products	0	0	-9431042	9467009	6972	0	0	0	0	0	0	-137327	-94389	-94389	
2700	Other Conversions & Blending	17846	0	0	-23232	0	23232	0	0	0	0	0	17846	0	17846	
2800	TC Total Conversion	-3705970	1386210	-9732505	7873214	-3066843	999058	-562094	-778417	-2873130	3752445	743767	-5964065	-5887523	-76543	
2900	Own Use & Loss	-4240	-93659	-518	-325749	-743	-12276	0	0	0	-356294	-4082	-797561	-797561	0	
3000	OI Other Input/Output	0	0	0	-32610	0	0	0	0	0	0	0	-32610	0	-32610	
3500	FS Stock Change	-26456	-5012	11849	3404	-5418	0	-21	0	0	0	0	-21654	0	-21654	
4000	DC Statal Discrepancy	43208	0	-76095	-6521	9637	0	0	0	0	0	0	-29772	-29772	0	
5000	Final Energy Consumption	427096	1283719	0	9036828	50613	986782	54220	0	0	3396151	739685	15975094	14203053	1772041	
6000	Industry	402587	1281740	0	3284658	49960	159109	18388	0	0	1307620	717036	7221098	5490897	1730201	
6100	NMFC Non-Manufacturing	178	603	0	608480	1930	25527	0	0	0	17223	0	653942	474431	179511	
6500	MFC Manufacturing	402409	1281136	0	2676177	48030	133583	18388	0	0	1290397	717036	6567156	5016466	1550890	
6520	Pulp & Paper	0	0	0	20792	70	563	12142	0	0	132838	253277	419682	419682	0	
6550	Chemical	19	37438	0	1809648	23095	3181	0	0	0	179582	256781	2309744	843806	1465939	
6570	Cement & Ceramics	184710	23143	0	85120	175	489	6235	0	0	79974	10800	390646	390154	492	
6580	Iron & Steel	223836	977757	0	100256	22175	31828	0	0	0	253494	105469	1714614	1714463	152	
6600	Machinery	0	6359	0	37273	945	18302	2	0	0	262650	0	325731	325731	0	
6700	Duplication Adjustment	-12253	-1231	0	-27736	-176	-676	-10	0	0	-40768	-89866	-171817	-171817	-0	
6800	Other Industries & SMEs	1927	227946	0	423747	0	46382	0	0	0	266689	124234	1090926	1006819	84107	
7000	ResCom	24509	1979	0	1891287	653	827673	35833	0	0	2021667	22648	4826249	4818571	7678	
7100	RES Residential	0	0	0	731171	0	418454	34912	0	0	928274	1306	2114117	2114117	0	
7150	Hokkaido,Tohoku,Hokuriku	0	0	0	258987	0	52403	0	0	0	166607	0	477997	477997	0	
7180	Kantou, Toukai, Kansai	0	0	0	339898	0	417463	0	0	0	624718	0	1382078	1382078	0	
7170	Chuugoku,Shikoku,Kyushu,Okinawa	0	0	0	147430	0	50550	0	0	0	210400	0	408379	408379	0	
7500	COM Commercial & Others	24509	1979	0	1160116	653	409219	921	0	0	1093394	21342	2712132	2704454	7678	
7510	Water supply, Sewage & Waste Dis	521	0	0	100189	0	7316	0	0	0	76046	12	184085	184085	0	
7540	Telecommunication & Broadcasting	0	0	0	18618	0	5698	0	0	0	36920	605	61841	61841	0	
7600	Trade & Finance Service	0	0	0	258854	0	54325	0	0	0	230281	9039	552499	552499	0	
7700	Public Service	17507	0	0	419901	0	124201	0	0	0	363562	3024	928195	928195	0	
7810	Commercial Service	464	334	0	106094	0	8911	0	0	0	88762	1066	205631	205631	0	
7850	Retail Service	4658	1567	0	280109	0	205098	0	0	0	196235	4745	692411	692411	0	
8000	Transportation	0	0	0	3860884	0	0	0	0	0	66864	0	3927748	3893585	34162	
8100	PAS Passenger	0	0	0	2283876	0	0	0	0	0	63385	0	2347261	2321514	25746	
8110	Car	0	0	0	2086803	0	0	0	0	0	0	0	2086803	2061151	25652	
8120	Rail	0	0	0	8598	0	0	0	0	0	63385	0	71983	71889	94	
8130	Ship	0	0	0	78498	0	0	0	0	0	0	0	78498	78498	0	
8140	Air	0	0	0	134790	0	0	0	0	0	0	0	134790	134790	0	
8500	FRT Freight	0	0	0	1577008	0	0	0	0	0	3479	0	1580487	1572071	8416	
8510	Truck & Lorry	0	0	0	1558126	0	0	0	0	0	0	0	1558126	1555516	2610	
8520	Rail	0	0	0	1878	0	0	0	0	0	3479	0	5357	5274	83	
8530	Ship	0	0	0	137346	0	0	0	0	0	0	0	137346	131623	5722	
8540	Air	0	0	0	24246	0	0	0	0	0	0	0	24246	24246	0	
9000	FEEC Final Energy Consumption	427096	1288259	0	7288772	42088	986782	54220	0	0	3396151	739685	14203053	14203053	0	
9500	Non-Energy	0	15460	0	1748057	8525	0	0	0	0	0	0	1772041	0	1772041	
9600	Industry	0	15460	0	1706216	8525	0	0	0	0	0	0	1730201	0	1730201	
9800	ResCom & others	0	0	0	7678	0	0	0	0	0	0	0	7678	0	7678	
9850	Transport	0	0	0	34162	0	0	0	0	0	0	0	34162	0	34162	

Table A 2-4 Energy balance simplified table (General Energy Statistics, FY2005)

2005FY	Code	100	150	200	250	400	450	500	550	600	700	800	900	910	920	
<Energy balance simplified table>		Coal	Coal Product	Oil	Oil Products	Natural Gas	Town Gas	Renewable E	Hydraulic	Nuclear Ener	Electricity	Heat	Total	Energy Total	Non-Energy	
<<Energy units>>		TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	
Code																
1000	Primary Energy Supply	4747650	81314	9506203	2135196	3288496	0	676443	671713	2678958	0	0	23783974	21767429	2016545	
1100	Indigenous Production	0	0	33051	0	134612	0	676443	671713	2678958	0	0	4192776	0	0	
1200	Import	4747650	81314	9473152	2135196	3153885	0	0	0	0	0	0	19591198	0	0	
1500	TPES Total Primary Energy Supply	4747650	81314	9506203	2135196	3288496	0	676443	671713	2678958	0	0	23783974	21767429	2016545	
1600	Export	-85	-49279	0	-897381	0	0	0	0	0	0	0	-946745	0	0	
1700	Stockpile Change	0	-16228	-96075	-73435	105352	0	0	0	0	0	0	-80386	0	0	
1900	DPES Domestic Primary Energy Supply	4747565	15807	9410128	1164381	3393848	0	676443	671713	2678958	0	0	supply side consumption side	22756843	20740297	2016545
2000	Energy Transformation & Own use	-4380236	1328905	-9637342	7534806	-3318058	1206465	-645344	-671713	-2678958	3515694	714918	-7028662	-6832682	-188862	
2100	Power Generation	-2146038	-186507	-301537	-546923	-1912210	-58869	-76110	-613992	-2678958	3440416	0	-5071412	-5071412	0	
2200	Auto Power Generation	-225239	-138544	-24	-396248	-18506	-67598	-247349	-57720	0	464983	0	-686246	-686246	0	
2300	Industrial Steam Generation	-201817	-33452	-33	-364073	-10580	-53178	-314989	0	0	0	832833	-145289	-145289	0	
2350	District Heat Supply	-633	0	0	-1058	0	-18102	-6739	0	0	-4129	25984	-4677	-4677	0	
2400	Town Gas Production	0	-1994	0	-76818	-1315225	1391962	-46	0	0	0	0	-2121	-2121	0	
2500	Coal Products	-1852761	1802622	0	-19827	0	0	0	0	0	0	0	-69966	-69966	0	
2600	Oil Products	0	0	-9331018	9324866	8203	0	0	0	0	0	-139784	-137714	0	-137714	
2700	Other Conversions & Blending	18933	0	0	-22505	0	22505	0	0	0	0	0	18933	0	18933	
2800	TC Total Conversion	-4407555	1442124	-9632613	7897434	-3248318	1216719	-645232	-671713	-2678958	3901270	719033	-6105809	-5979711	-118781	
2900	Own Use & Loss	-6994	-94941	-85	-309370	-41736	-10254	0	0	0	-385576	-4115	-852972	-852972	0	
3000	OI Other Input/Output	0	0	0	-53184	0	0	0	0	0	0	0	-53184	0	-53184	
3500	FS Stock Change	34314	-18378	-4644	-73	-28004	0	-112	0	0	0	0	-16897	0	-16897	
4000	DC Statistical Discrepancy	-48131	0	-227214	-2538	9378	0	-0	0	0	0	0	-268505	-261187	0	
5000	Final Energy Consumption	415460	1344712	0	8701725	66413	1206465	31099	0	0	3515694	714918	15996485	14168802	1827683	
6000	Industry	394168	1342658	0	3142673	65661	191539	6329	0	0	1231595	689846	7064470	5273144	1791326	
6100	NMFC Non-Manufacturing	100	191	0	503751	2758	30491	0	0	0	10887	0	548178	423510	124667	
6500	MFG Manufacturing	394067	1342467	0	2638922	62903	161049	6329	0	0	1220708	689846	6516292	4849634	1666658	
6520	Pulp & Paper	0	0	0	18699	119	762	25	0	0	127812	242031	389447	389447	0	
6550	Chemical	4351	37042	0	1880133	31475	5702	0	0	0	171601	242225	2372528	789974	1582554	
6570	Cement & Ceramics	161134	20463	0	75555	185	842	6300	0	0	78074	9075	351627	348954	2673	
6580	Iron & Steel	248848	971128	0	84755	25945	47754	0	0	0	253662	97734	1729825	1729695	130	
6600	Machinery	1	5255	0	38649	3007	25317	5	0	0	285518	0	355752	355752	0	
6700	Duplication Adjustment	-24479	0	0	-20151	-500	-754	0	0	0	-33744	-77425	-157052	-154564	-2488	
6900	Other Industries & SMEs	1409	299506	0	386675	0	28603	0	0	0	191277	129120	1036590	952801	83789	
7000	ResCom	21292	2054	0	1872067	751	1014925	24769	0	0	2215492	25072	5176423	5174228	2195	
7100	RES Residential	0	0	0	701600	0	435817	24033	0	0	1019088	1326	2181864	2181864	0	
7150	HokkaidoTohoku,Hokuriku	0	0	0	252024	0	57970	0	0	0	182318	0	492311	492311	0	
7160	Kantou,Toukai,Kansai	0	0	0	329849	0	472168	0	0	0	705199	0	1507215	1507215	0	
7170	Chugoku,Shikoku,Kyushu,Okinawa	0	0	0	151797	0	55495	0	0	0	243104	0	450396	450396	0	
7500	COM Commercial & Others	21292	2054	0	1170467	751	579108	736	0	0	1196404	23746	2945559	2992364	2195	
7510	Water supply, Sewage & Waste Dis	707	0	0	97018	0	10275	0	0	0	77690	10	185689	185689	0	
7540	Telecommunication & Broadcasting	0	0	0	16400	0	7767	0	0	0	33240	687	58094	58094	0	
7600	Trade & Finance Service	0	0	0	228066	0	184556	0	0	0	369264	7442	789329	789329	0	
7700	Public Service	15580	0	0	396400	0	165795	0	0	0	345691	2515	925981	925981	0	
7810	Commercial Service	785	220	0	83668	0	8214	0	0	0	87825	947	181659	181659	0	
7850	Retail Service	2159	1798	0	264254	0	238811	0	0	0	193168	2954	703145	703145	0	
8000	Transportation	0	0	0	3688985	0	0	0	0	0	68607	0	3755892	3721430	34162	
8100	PAS Passenger	0	0	0	2242955	0	0	0	0	0	65029	0	2307984	2282238	25746	
8110	Car	0	0	0	1968839	0	0	0	0	0	0	0	1968839	1943187	25652	
8120	Rail	0	0	0	7833	0	0	0	0	0	65029	0	72862	72768	94	
8130	Ship	0	0	0	70204	0	0	0	0	0	0	0	70204	70204	0	
8140	Air	0	0	0	137208	0	0	0	0	0	0	0	137208	137208	0	
8500	FRT Freight	0	0	0	1444030	0	0	0	0	0	3578	0	1447608	1439192	8417	
8510	Truck & Lorry	0	0	0	1333297	0	0	0	0	0	0	0	1333297	1330687	2610	
8520	Rail	0	0	0	1718	0	0	0	0	0	3578	0	5296	5212	84	
8530	Ship	0	0	0	117819	0	0	0	0	0	0	0	117819	112097	5722	
8540	Air	0	0	0	23641	0	0	0	0	0	0	0	23641	23641	0	
9000	FEEC Final Energy Consumption	415460	1329123	0	6905897	50146	1206465	31099	0	0	3515694	714918	14168802	14168802	0	
9500	Non-Energy	0	15589	0	1795828	16286	0	0	0	0	0	0	1827683	0	1827683	
9800	Industry	0	15589	0	1795970	16286	0	0	0	0	0	0	1791326	0	1791326	
9800	ResCom & others	0	0	0	2195	0	0	0	0	0	0	0	2195	0	2195	
9850	Transport	0	0	0	34162	0	0	0	0	0	0	0	34162	0	34162	

Table A 2-5 Energy balance simplified table (General Energy Statistics, FY2010)

2010FY (Energy balance simplified table) Code	Code Code (Energy units)	100	150	200	250	400	450	500	550	600	700	800	900	910	920
		Coal	Coal	Oil	Natural Gas	Town Gas	Renewable E	Hydraulic	Nuclear Ener	Electricity	Heat	Total	Energy Total	Non-Energy	
		TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	TJ	
1000	Primary Energy Supply	4967481	29909	8170567	1930339	4001721	0	816048	711607	2494679	0	0	23122551	21344376	1778175
1100	Indigenous Production	0	0	30638	0	149324	0	816048	711607	2494679	0	0	4202496	0	0
1200	Import	4967481	29909	8139929	1930339	3852397	0	0	0	0	0	0	18920055	0	0
1500	TPES Total Primary Energy Supply	4967481	29909	8170567	1930339	4001721	0	816048	711607	2494679	0	0	23122551	21344376	1778175
1600	Export	-87	-19695	0	-1208528	0	0	0	0	0	0	0	-1228309	0	0
1700	Stockpile Change	0	3901	-27191	-12021	232175	0	0	0	0	0	0	196864	0	0
1900	DPES Domestic Primary Energy Supply	4967395	14115	8143377	709790	4233896	0	816048	711607	2494679	0	0	22091107	20312932	1778175
													supply side		
													consumption side		
2000	Energy Transformation & Own use	-4378708	1324833	-8099635	6748180	-4183366	1450226	-790497	-711607	-2494679	3591136	646732	-6897586	-6856549	-37004
2100	Power Generation	-2084699	-200982	-189297	-374299	-2347369	-58859	-116517	-568728	-2494679	3478244	0	-4953354	-4953354	0
2200	Auto Power Generation	-229402	-131208	-59	-274946	-31593	-72109	-321610	-142879	0	499018	0	-704788	-704788	0
2300	Industrial Steam Generation	-209478	-37764	-77	-255544	-22812	-66356	-346236	0	0	0	774086	-164200	-164200	0
2350	District Heat Supply	0	0	0	-841	0	-17003	-5562	0	0	-4126	25462	-2070	-2070	0
2400	Town Gas Production	0	0	0	-48495	-1670107	1697063	0	0	0	0	0	-21539	-21539	0
2500	Coal Products	-1825984	1804431	0	-14315	0	0	-87	0	0	0	0	-35955	-35955	0
2600	Oil Products	0	0	-7914831	8074786	4667	0	0	0	0	0	-147048	17573	0	17573
2700	Other Conversions & Blending	21802	0	0	-19358	0	19358	0	0	0	0	0	21802	0	21802
2800	TC Total Conversion	-4327761	1434477	-8104264	7086988	-4067214	1501094	-790031	-711607	-2494679	3974136	652499	-5846562	-5881905	39376
2900	Own Use & Loss	-21642	-112202	-62	-283374	-117728	-50869	0	0	0	-383000	-5767	-974644	-974644	0
3000	OI Other Input/Output	0	0	0	-45542	0	0	0	0	0	0	0	-45542	0	-45542
3500	FS Stock Change	-29305	2557	4692	-9892	1576	0	-466	0	0	0	0	-30839	0	-30839
4000	DC Statistical Discrepancy	201144	0	43742	-9521	-16285	0	-0	0	0	67	0	219145	223177	0
5000	Final Energy Consumption	387543	1338948	0	7467492	66815	1450226	25550	0	0	3591070	646732	14974376	13233205	1741171
6000	Industry	366977	1336815	0	2757841	66238	225185	7376	0	0	1189032	622314	6571777	4864769	1707008
6100	NMFC Non-Manufacturing	76	191	0	378385	5064	33406	0	0	0	9253	0	426374	341572	84802
6500	MFC Manufacturing	366901	1336624	0	2379456	61174	191779	7376	0	0	1179780	622314	6145403	4523197	1622206
6520	Pulp & Paper	0	0	0	14871	334	1852	433	0	0	113560	207044	330895	330895	0
6550	Chemical	16	40777	0	1803571	33223	6099	251	0	0	158172	226698	2268808	732073	1536735
6570	Cement & Ceramics	125879	17973	0	65852	431	972	6410	0	0	77576	12752	307844	305477	2366
6580	Iron & Steel	253852	975580	0	70803	20328	69826	387	0	0	252475	99338	1742570	1742385	185
6600	Machinery	0	4259	0	30432	2596	27763	0	0	0	282866	0	347916	347916	0
6700	Duplication Adjustment	-15892	0	0	-9724	0	-4013	-251	0	0	-36304	-74945	-141030	-138718	-2311
6900	Other Industries & SMEs	1012	290080	0	296364	0	11792	0	0	0	163320	109814	872383	787151	85231
7000	ResCom	20566	2133	0	1346973	578	1225041	18175	0	0	2334488	24418	4972372	4972372	0
7100	RES Residential	0	0	0	609480	0	428875	17516	0	0	1098953	1282	2154107	2154107	0
7150	Hokkaido	0	0	0	214344	0	48592	0	0	0	181877	0	444812	444812	0
7180	Tohoku	0	0	0	274505	0	476774	0	0	0	634920	0	1386199	1386199	0
7170	Kansai	0	0	0	110838	0	62080	0	0	0	214842	0	387760	387760	0
7170	Chugoku,Shikoku,Kyushu,Okinawa	0	0	0	737493	578	798166	658	0	0	1235535	23136	2818265	2818265	0
7500	COM Commercial & Others	20566	2133	0	68644	0	12568	0	0	0	78386	7	160517	160517	0
7510	Water supply, Sewage & Waste Dis	912	0	0	10881	0	9727	0	0	0	31519	806	52933	52933	0
7540	Telecommunication & Broadcasting	0	0	0	144426	0	266894	0	0	0	442630	4783	858733	858733	0
7800	Trade & Finance Service	13621	0	0	290937	0	205023	0	0	0	318038	1803	829422	829422	0
7700	Public Service	1081	96	0	47483	0	6129	0	0	0	81086	762	136637	136637	0
7810	Commercial Service	2198	2001	0	200688	0	246249	0	0	0	184404	1147	636886	636886	0
7850	Retail Service	0	0	0	3362677	0	0	0	0	0	67549	0	3430226	3396064	34162
8000	Transportation	0	0	0	2068627	0	0	0	0	0	64280	0	2132908	2107157	25751
8100	PAS Passenger	0	0	0	1904164	0	0	0	0	0	0	0	1904164	1878511	25652
8110	Car	0	0	0	7248	0	0	0	0	0	64280	0	71528	71429	99
8120	Rail	0	0	0	53094	0	0	0	0	0	0	0	53094	53094	0
8130	Ship	0	0	0	115381	0	0	0	0	0	0	0	115381	115381	0
8140	Air	0	0	0	1294050	0	0	0	0	0	3269	0	1297319	1288908	8411
8500	FRT Freight	0	0	0	1208308	0	0	0	0	0	0	0	1208308	1205698	2610
8510	Truck & Lorry	0	0	0	1493	0	0	0	0	0	3269	0	4762	4683	79
8520	Rail	0	0	0	106166	0	0	0	0	0	0	0	106166	100443	5722
8530	Ship	0	0	0	21549	0	0	0	0	0	0	0	21549	21549	0
8540	Air	0	0	0	0	0	0	0	0	0	0	0	0	0	0
9000	FEEC Final Energy Consumption	387543	1323777	0	5788091	50216	1450226	25550	0	0	3591070	646732	13233205	13233205	0
9500	Non-Energy	0	15171	0	1709401	16599	0	0	0	0	0	0	1741171	0	1741171
9600	Industry	0	15171	0	1675239	16599	0	0	0	0	0	0	1707008	0	1707008
9800	ResCom & others	0	0	0	0	0	0	0	0	0	0	0	0	0	0
9850	Transport	0	0	0	34162	0	0	0	0	0	0	0	34162	0	34162

A2.2.2. General Energy Statistics and CRF

In order to report CO₂ emissions in CRF, emissions reported under the sectors in *General Energy Statistics* (Energy Balance Table) were reported under each sector in CRF as indicated in Table A 2-6 and Table A 2-7.

Values subtracting energy consumption reported under ‘Non-energy’ [#9500] from energy consumption reported under ‘Energy Conversion & Own use’ [#2000], ‘Industry’ [#6000], ‘Residential’ [#7100], ‘Commercial & Others’ [#7500], and ‘Transportation’ [#8000] in *General Energy Statistics* (Energy Balance Table) are used for activity data. Because energy consumption reported under ‘Non-energy’ [#9500] was used for the purposes other than combustion and was considered not emitting CO₂, these values were deducted. However, out of this amount deducted as feedstocks and non-energy use, the emissions from what is used or collected as energy during waste incineration are separately estimated and reported.

The *Revised 1996 IPCC Guidelines* requires carbon dioxide emitted from auto power generation, etc., to be counted in the corresponding sector. In Japan’s Energy Balance Table (*General Energy Statistics*), fuel consumption used for auto power generation and industrial steam generation are presented under ‘Auto Power Generation’ [#2200], ‘Industrial Steam Generation’ [#2300] in the Energy Conversion Sector. However, auto power generation and industrial steam generation actually belong to industrial sector. Hence, carbon dioxide emissions from “Auto Power Generation” and “Industrial Steam Generation” are allocated to each section of ‘1.A.2 Manufacturing Industries and Construction’.

Table A 2-6 Correspondence between sectors of General Energy Statistics (Miner Sector) and of the CRF

CRF		Japan's Energy Balance Table	
1A1	Energy Industries		
1A1a	Public Electricity and Heat Production	Power Generation, General Electric Utilities	#2110
		Own use, General Electric Utilities	#2911
		Power Generation, Independent Power Producing	#2150
		Own use, Independent Power Producing	#2912
		District Heat Supply	#2350
1A1b	Petroleum Refining	Own use, District Heat Supply	#2913
		Own use, Oil Refinery	#2916
1A1c	Manufacture of Solid Fuels and Other Energy Industries	Coal Products	#2500
		Own use, Town Gas	#2914
		Own use, Steel Coke	#2915
		Own use, Other Conversion	#2917
1A2	Manufacturing Industries and Construction		
1A2a	Iron and Steel	Auto: Iron & Steel	#2217
		Steam Generation: Iron & Steel	#2307
		Final Energy Consumption, Iron & Steel	#6580
		Non-Energy, Iron & Steel	#9680
1A2b	Non-Ferrous Metals	Auto: Non-Ferrous Metal	#2218
		Steam Generation: Non-Ferrous Metal	#2308
		Final Energy Consumption, Non-Ferrous Metal	#6590
		Non-Energy, Non-Ferrous Metal	#9690
1A2c	Chemicals	Auto: Chemical Textiles	#2212
		Steam Generation: Chemical Textiles	#2302
		Final Energy Consumption, Chemical Textiles	#6530
		Non-Energy, Chemical Textiles	#9630
		Auto: Chemical	#2214
		Steam Generation: Chemical	#2304
		Final Energy Consumption, Chemical	#6550
		Non-Energy, Chemical	#9650
1A2d	Pulp, Paper and Print	Auto: Pulp & Paper	#2211
		Steam Generation: Pulp & Paper	#2301
		Final Energy Consumption, Pulp & Paper	#6520
		Non-Energy, Pulp & Paper	#9620
1A2e	Food Processing, Beverages and Tobacco	Final Energy Consumption, Food	#6510
		Non-Energy, Non-Manufacturing Industry (Food)	#9610
1A2f	Other		
	Mining	Final Energy Consumption, Mining	#6120
		Non-Energy, Non-Manufacturing Industry (Mining)	#9610
	Construction	Final Energy Consumption, Construction	#6150
		Non-Energy, Non-Manufacturing Industry (Construction)	#9610
	Oil Products	Auto: Oil products	#2213
		Steam Generation: Oil products	#2303
		Final Energy Consumption, Oil products	#6540
		Non-Energy, Oil products	#9640
	Glass Wares	Auto: Glass Wares	#2215
		Steam Generation: Glass Wares	#2305
		Final Energy Consumption, Glass Wares	#6560
		Non-Energy, Glass Wares	#9660
	Cement & Ceramics	Auto: Cement & Ceramics	#2216
		Steam Generation: Cement & Ceramics	#2306
		Final Energy Consumption, Cement & Ceramics	#6570
		Non-Energy, Cement & Ceramics	#9670
	Machinery	Auto: Machinery & Others	#2219
		Steam Generation: Machinery & Others	#2309
		Final Energy Consumption, Machinery	#6600
		Non-Energy, Machinery	#9700
	Duplication Adjustment	Auto: Duplication Adjustment	#2220
		Steam Generation: Duplication Adjustment	#2310
		Final Energy Consumption, Duplication Adjustment	#6700
Non-Energy, Duplication Adjustment		#9710	
Other Industries & Small and Medium Enterprises	Auto: Others	#2250	
	Final Energy Consumption, Other Industries & Small and Medium Enterprises	#6900	
	Non-Energy, Other Industries & Small and Medium Enterprises	#9720	

Table A 2-7 Correspondence between sectors of General Energy Statistics (Miner Sector) and of the CRF (cont.)

CRF		Japan's Energy Balance Table	
1A3	Transport		
1A3a	Civil Aviation	Final Energy Consumption, Passenger Air	#8140
		Final Energy Consumption, Freight Air	#8540
		Non-Energy, Transportation (Air)	#9850
1A3b	Road Transportation	Final Energy Consumption, Passenger Car	#8110
		Final Energy Consumption, Freight, Freight Truck & Lorry	#8510
		Final Energy Consumption, Passenger Bus	#8115
		Final Energy Consumption, Passenger, Transportation fraction estimation error	#8190
		Final Energy Consumption, Freight, Transportation fraction estimation error.	#8590
		Non-Energy, Transportation (Car, Truck & Lorry, Bus)	#9850
		Final Energy Consumption, Passenger Rail	#8120
1A3c	Railways	Final Energy Consumption, Freight Rail	#8520
		Non-Energy, Transportation (Rail)	#9850
		Final Energy Consumption, Passenger Ship	#8130
1A3d	Navigation	Final Energy Consumption, Freight Ship	#8530
		Non-Energy, Transportation (Ship)	#9850
		1A3e	Other Transportation
1A4	Other Sectors		
1A4a	Commercial/Institutional	Final Energy Consumption, Commercial & Others	#7500
		Non-Energy, ResCom & others (Commercial & Others)	#9800
1A4b	Residential	Final Energy Consumption, Residential	#7100
		Non-Energy, ResCom & others (Residential)	#9800
1A4c	Agriculture/Forestry/Fisheries	Final Energy Consumption, Agriculture, Forestry & Fishery	#6110
		Non-Energy, Non-Manufacturing Industry (Agriculture, Forestry & Fishery)	#9610
1A5	Other		
1A5a	Stationary	-	-
		-	-
1A5b	Mobile	-	-

- Auto: Non-utility power generation
- #9xxx items are not energy use activity.

In 'Energy Conversion & Own use', 'Power Generation' [#2100], 'Auto Power Generation' [#2200], 'Industrial Steam Generation' [#2300], 'District Heat Supply' [#2350], 'Coal Products' [#2500], and 'Own Use & Loss' [#2900] are calculated, and other sectors ('Town Gas Production', 'Oil Products', 'Other Conversions & Blending', 'Other Input/Output' and 'Stock Change') are excluded from calculations.

Energy consumptions reported under 'Town Gas Production' are feedstocks of town gas production, and was not used to purposes combustion. Therefore, they are excluded from calculations. Meanwhile, CO₂ emissions from carbon contained in these feedstocks are calculated with town gas consumption in final energy consumption sector (industry, residential, commercial & others, and transportation).

The energy consumption recorded under coal products corresponds to the difference between the coke-making carbon input and carbon output. This is the portion that is oxidized in the atmosphere (burned) from the time that red-hot coke is extruded from a coke oven until it enters the coke dry quenching facility. It was considered appropriate to count this as CO₂ emissions, and it was calculated as carbon emissions from this sector.

Energy consumptions reported under 'Oil Products' are feedstocks for oil products, and was not used for the purpose of combustion. Meanwhile, CO₂ emissions from carbon contained in these feedstocks

are calculated with each kind of energy consumption in energy conversion sector and final energy consumption sector (industry, residential, commercial & others, and transportation).

A2.2.3. Duplication adjustment for Energy Balance Table

The data set of the manufacturing sector indicated in Japan's Energy Balance Table (*General Energy Statistics*) and used as the reference of activity data are based on the Ministry of Economy, Trade and Industry's *Yearbook of the Current Survey of Energy Consumption*. *The Yearbook of the Current Survey of Energy Consumption* is a statistical survey on factories and business institutions of key manufacturing. Factories and business institutions which produce items indicated in Table A 2-8 are surveyed.

In Japan, it is rare that single factory or business institution produces single item. Most factories and business institutions produce various items extending across categories of industry utilizing by-products and surplus business resources. For example, most integrated steelworks produce not only steel products falling into iron & steel industry but also coke and slag cement falling into cement & ceramics industry and chemical products delivered from coal tar and industrial gas falling into chemical industry; i.e. one factory can conduct three different categories of industries and produces many kinds of items at the same time.

Because single factory may report duplicated energy consumption data which can not be classified to certain sector or item, total energy consumption summed up by sector or by item can be larger than actual total energy consumption when totalizing by sector or by item is conducted under the *Yearbook of the Current Survey of Energy Consumption*.

Hence, to avoid duplication adjustment and to adjust the data in the *Yearbook of the Current Survey of Energy Consumption*, the following steps were taken: (1) to calculate total energy consumption by factory and business institution, (2) to calculate total energy consumption by sector and by item including duplication among sectors and items, (3) to express the difference between total energy consumption by sector and item and total energy consumption by factory and business as negative values as "duplication adjustment".

In the *Yearbook of the Current Survey of Energy Consumption*, the adjustment stated above is applied indicating values for "duplication adjustment" when total energy consumption is calculated by sector or by item for Auto Power Generation, Industrial Steam Generation, and Manufacturing.

Calculation method for duplication adjustment

$$\text{Values of duplication adjustment} = E_p - E_t$$

E_p : Total energy consumption of designated sectors and items by factories and business institutions

E_t : Total energy consumption by factories and business institutions

Subjects to be surveyed to obtain the data for the *Yearbook of the Current Survey of Energy Consumption* were changed in December, 1997. As shown in Table A 2-8, the survey for the industries

of Dyeing, Rubber Product, and Non-ferrous Metals has been discontinued since 1998. Also, since 1998, business institutions or designated items to be surveyed for the industries of Chemical Ceramics, Clay and Stone Products, Glass Products, Iron and Steel, Non-ferrous Metals, and Machinery has been changed. Therefore, energy consumption for the said industries during 1990-1997 is chronologically inconsistent comparing to that from 1998 and onward. Also, the classification of industries was revised during this period. Because of these changes, energy consumption for duplication adjustment, other industries, and small-to-medium-sized manufacturing significantly fluctuates.

Table A 2-8 Surveyed industries and products in *Yearbook of the Current Survey of Energy Consumption*

Surveyed industry	from 1990 to 1997		after 1997	
	Products	Scope of survey	Products	Scope of survey
Pulp and paper industry	* Pulp * Paper * Sheet paper	All Establishments with 50 or more employees Establishments with 50 or more employees	* Pulp * Paper * Sheet paper	All Establishments with 50 or more employees Establishments with 50 or more employees
Chemical industry (except chemical fiber industry)	* Petrochemical products * Ammonia and ammonia-derived products * Soda industries chemicals * High pressure gas (O ₂ , N ₂ , Ar) * Inorganic chemicals and colorant (titanic oxide, active char, chinese white, iron oxide) * Oil and fat products and surfactant	All All All All (except high pressure gas products by air fraction method(gas container)) All Establishments with 30 or more employees	* Petrochemical products * Ammonia and ammonia-derived products * Soda industries chemicals	All
Chemical fiber industry	* Chemical fibers	Establishments with 30 or more employees	* Chemical fibers	Establishments with 30 or more employees
Petroleum products industry	* Petroleum products (except grease)	All	* Petroleum products (except grease)	All
Ceramics, clay and stone products industry (except glass product industry, with the exception of sheet glass industry)	* Cement * Sheet glass * Lime * Fire brick * Carbon products	All All Establishments with 30 or more employees Establishments with 30 or more employees All	* Cement * Sheet glass * Lime	All All Establishments with 30 or more employees
Glass product industry (except sheet glass industry)	* Glass products	Establishments with 10 or more employees	* Glass products	Establishments with 100 or more employees
Iron and steel industry	Manufacturers of pig iron, ferroalloys, crude steel, semi-finished steel products, forged steel products, cast steel products, general steel and hot-rolled steel materials, cold-rolled wide steel strips, cold-rolled electrical steel strips, plated steel materials, special steel hot-rolled steel materials, steel pipes (except cold working steel pipes), or cast iron tubes. Iron and steel.	All	Manufacturers of pig iron, ferroalloys, crude steel, semi-finished steel products, forged steel products, cast steel products, general steel and hot-rolled steel materials, cold-rolled wide steel strips, cold-rolled electrical steel strips, plated steel materials, special steel hot-rolled steel materials, steel pipes (except cold working steel pipes), or cast iron tubes. Iron and steel.	All
Non-ferrous metal industry	* Non-ferrous metals	All	* Copper * Lead * Zinc * Aluminum * Aluminum secondary ground metal	All All All All Establishments with 30 or more employees
Machinery industry	* Machinery and appliances * cast and forged products	Establishments with 500 or more employees Establishments with 100 or more employees	* Civil engineering machinery, tractors, metal working and metal processing machinery, parts and accessories for communication and electronics equipment, electron tubes, semiconductors, ICs, electronics applied equipment, automobiles and parts (including motorcycles)	Establishments with 500 or more employees which are designated by the Minister of International Trade and Industry
Dyeing	* Dyeing wool * Dyeing fabric	Establishments with 20 or more employees	demise	
Rubber product	* Tires and tube	Establishments with 30 or more employees	demise	
Non-ferrous metal product	* Copper and brass * Flat-rolled aluminum * Electric cable * Aluminum secondary bare metal	All All Establishments with 30 or more employees Establishments with 30 or more employees	demise	

References

1. Environmental Agency , *The Estimation of CO₂ Emissions in Japan*, 1992
2. Research Institute of Economy, Trade & Industry, Kazunari Kaino, *Interpretation of General Energy Statistics*, 2009

Annex 3. Other Detailed Methodological Descriptions for Individual Source or Sink Categories

A3.1. Methodology for Estimating Emissions of Precursors

In addition to the greenhouse gases (e.g., CO₂, CH₄, N₂O, HFCs, PFCs, SF₆) reported under the Kyoto Protocol, Japan reports on the emissions of precursors (NO_x, CO, NMVOC, SO₂) calculated by established methods. This section explains the source categories for which methodologies for estimating emissions have been provided.

Emissions from the source categories for which estimation methods have not been established are considered to be minimal, and accordingly reported as either “NO” or “NE” (or as “IE” as the case may be) based on the results of historical investigations.

A3.1.1. Energy Sector

A3.1.1.1. Stationary Combustion (1.A.1., 1.A.2., 1.A.4.: NO_x, CO, NMVOC, SO₂)

A3.1.1.1.a. Facilities emitting soot and smokes

1) NO_x and SO₂

● *Methodology for Estimating Emissions*

General Survey of the Emissions of Air Pollutants by the Ministry of the Environment was used as the basis for estimation of NO_x and SO₂ emitted from fixed sources (see Page 3.12 for details of the survey). So as to ensure consistency with the *Revised 1996 IPCC Guidelines* and the *IPCC Good Practice Guidance (2000)*, the following operation isolated the emissions from the energy sector from the emissions listed in the *General Survey of the Emissions of Air Pollutants*:

1. All emissions from the following facilities and operations are reported under Energy:
 - Facility: [0101–0103: Boilers]; [0601–0618: Metal rolling furnaces, metal furnaces, and metal forge furnaces]; [1101–1106: Drying ovens]; [2901–3202: Gas turbines, diesel engines, gas engines, and gasoline engines]
 - Operation: [A–D: Accommodation/eating establishments, health care/educational and academic institutions, public bathhouses, laundry services]; [F–L: Agriculture/fisheries, mining, construction, electricity, gas, heat distribution, building heating/other operations]
2. Emissions from the facilities and operations other than the above and [1301–1304: Waste incinerators], are reported under the Industrial Processes sector. Accordingly, the emissions from the specified sources, calculated by the following methods, are subtracted from the emissions listed in the *General Survey of the Emissions of Air Pollutants* to determine the emissions from the Energy sector.

➤ NO_x

If raw material falls under either [44: Metallurgical coal] or [45: Metallurgical coke], the following equation is used:

Calculation of NO_x emissions from metallurgical coal or coke (to be included in the Industrial Processes sector)

$$\begin{aligned} & \text{NO}_x \text{ emissions from metallurgical coal or coke [t-NO}_x\text{]} \\ & = \text{NO}_x \text{ emission factor per material [t-NO}_x\text{/kcal]} \times \text{energy consumed per material [kcal]} \\ & \quad \times (1 - \text{denitrification rate [\%]}) \end{aligned}$$

If raw material falls under either [41: Iron/ironstone] or [46: Other], the following equation is used:

Calculation of NO_x emissions from iron/ironstone or other material (to be included in the Industrial Processes sector)

$$\begin{aligned} & \text{NO}_x \text{ emissions from iron/ironstone or other material [t-NO}_x\text{]} \\ & = \text{Nitrogen content per material [t-NO}_x\text{]} \times (1 - \text{denitrification rate [\%]}) \end{aligned}$$

If, however, the emissions from the Industrial Processes sector calculated by the above equations exceed the emission volume listed in the *General Survey of the Emissions of Air Pollutants*, the total emissions listed in the Survey are considered to be the emissions from the Industrial Processes sector. Materials listed in the categories [42: Sulfide minerals] and [43: Non-ferrous metal ores] are excluded from the calculation due to the lack of data.

➤ **SO₂**

Emissions from the Industrial Processes sector is calculated from the consumption and sulfur contents of the materials in categories from [41: Iron/ironstone] to [46: Other materials], and subtracted from the emissions listed in the *General Survey of the Emissions of Air Pollutants* to determine SO₂ emissions in the energy sector.

Calculation of SO₂ emissions (in the Industrial Processes sector)

$$\text{SO}_2 \text{ emissions [t-SO}_2\text{]} = \text{Sulfur content per material [t-SO}_2\text{]} \times (1 - \text{desulphurization rate [\%]})$$

● **Emission factors**

➤ **NO_x emission factors for metallurgical coal and coke**

NO_x emission factors for the materials used in the calculation of NO_x emissions from metallurgical coal and coke (in the Industrial Processes sector) were established for each facility and material type based on the *General Survey of the Emissions of Air Pollutants*.

➤ **Denitrification rate**

The denitrification rate was calculated by the following equation:

Calculation of denitrification rate

$$\begin{aligned} & \text{Denitrification rate [\%]} \\ & = \text{Denitrification efficiency [\%]} \times (\text{Hours of operation of denitrification unit [h/yr]} / \\ & \quad \text{Hours of operation of furnace [h/yr]} \times (\text{Processing capacity of denitrification unit [m}^3\text{/yr]} / \\ & \quad \text{max exhaust gas emission [m}^3\text{/yr]}) \end{aligned}$$

The General Survey of the Emissions of Air Pollutants data were used for all items.

Denitrification efficiency: (NO_x volume before treatment – NO_x volume after treatment) / volume of smoke and soot

➤ **Desulphurization rate**

Desulphurization rate was calculated by the following equation:

Calculation of desulphurization rate

$$\begin{aligned} & \text{Desulphurization rate [\%]} \\ &= \text{Desulphurization efficiency [\%]} \times (\text{Hours operation of desulphurization unit [h/yr]} / \\ & \text{Hours operation of furnace [h/yr]} \times (\text{Processing capacity of desulphurization unit [m}^3\text{/yr]} / \\ & \text{max exhaust gas emission [m}^3\text{/yr]}) \end{aligned}$$

The General Survey of the Emissions of Air Pollutants data were used for all items.

Desulphurization efficiency: (SO₂ volume before treatment – SO₂ volume after treatment) / volume of smoke and soot

- **Activity data**

- **Energy consumption of metallurgical coal or coke**

The activity data was calculated by multiplying the consumption of materials (under [44: Metallurgical coal] and [45: Metallurgical coke]) provided in the *General Survey of the Emissions of Air Pollutants* by gross calorific value.

- **Nitrogen content of iron/ironstone and other materials**

The activity data was calculated by multiplying the weighted average of nitrogen content, calculated from the nitrogen content and consumption of the materials (under [41: Iron/ironstone] and [46: Other materials]) provided in the *General Survey of the Emissions of Air Pollutants*, by the consumption volume of the material.

- **Sulfur content of various materials**

The activity data was calculated by multiplying the weighted average of sulfur content, calculated on the basis of sulfur content and consumption of the material (under [44: Metallurgical coal] through [46: Other materials]) provided in the *General Survey of the Emissions of Air Pollutants*, by the consumption volume of the material.

2) CO

- **Methodology for Estimating Emissions**

Emissions of CO from the specified sources were calculated by multiplying the energy consumption per facility type by Japan's own emission factor.

- **Emission factors**

CO emission factors were established based on the summary data in the *Reports on Greenhouse gas emissions estimation methodology* (Japan Sociality Atmospheric Environment, 1996).

- **Activity data**

Energy consumption according to facility type determined from General Energy Statistics was used for activity data.

3) NMVOC

- **Methodology for Estimating Emissions**

Emissions of NMVOC from the specified sources were calculated by multiplying the energy consumption per facility type by Japan's own emission factor.

- **Emission factors**

NMVOC emission factors were established by multiplying the CH₄ emission factor for each facility per fuel type by the ratio of NMVOC emission to CH₄ emission factor per fuel type. The CH₄ emission factors were established from the summary data provided in the *Reports on Greenhouse gas emissions estimation methodology* (Japan Sociality Atmospheric Environment, 1996), while the NMVOC/CH₄ emission factor ratios were determined from the *report on Screening Survey Regarding Measures to Counter Global Warming* (Japan Environmental Sanitation Center) and *Study of Establishment of Methodology for Estimation of Hydrocarbon Emissions* (Institute of Behavioral Science).

- **Activity data**

Energy consumption according to facility type determined from General Energy Statistics (Agency for Natural Resources and Energy) was used for activity data.

A3.1.1.1.b. Small facilities (commercial and other sector, manufacturing sector)

- **Methodology for Estimating Emissions**

NO_x, CO, NMVOC, and SO₂ emitted by the specified sources were calculated by multiplying energy consumption per facility type by Japan's own emission factor.

- **Emission factors**

- **NO_x and SO₂**

Emission factors for NO_x and SO_x were established for each fuel type for [0102: Heating system boilers] for facilities listed in [L: Heating systems for buildings/other places of business] in the *General Survey of the Emissions of Air Pollutants* by aggregating emission and energy consumption per fuel type.

- **CO**

The emission factors established for [0102: Heating system boilers] based on the *Reports on Greenhouse gas emissions estimation methodology* (Japan Sociality Atmospheric Environment, 1996) were adopted as the CO emission factors.

- **NMVOC**

NMVOC emission factors were established by multiplying the CH₄ emission factors for [0102: Heating system boilers] by the ratio of NMVOC emission to CH₄ emission factor per fuel type. The CH₄ emission factors were established from the *Reports on Greenhouse gas emissions estimation methodology* (Japan Sociality Atmospheric Environment, 1996), while the NMVOC/CH₄ emission factor ratios were determined from the *report on Screening Survey Regarding Measures to Counter Global Warming* (Japan Environmental Sanitation Center) and *Study of Establishment of Methodology for Estimation of Hydrocarbon Emissions* (Institute of Behavioral Science).

- **Activity data**

To determine NO_x and SO₂, energy consumption by small facilities per fuel type was calculated by subtracting energy consumption per fuel type, identified by the *General Survey of the Emissions of Air Pollutants*, from energy consumption per fuel type provided in the *General Energy Statistics* (Agency for Natural Resources and Energy). If the activity data shown in the *General Survey of the Emissions of Air Pollutants* exceeded the activity data provided in the *General Energy Statistics*, the activity data

for the specified sources was deemed to be zero. The fuels covered were town gas, LPG, kerosene, and heating oil A. Energy consumption from General Energy Statistics (Agency for Natural Resources and Energy) was used for CO and NMVOCs.

A3.1.1.1.c. Residential sector

- **Methodology for Estimating Emissions**

NO_x, CO, NMVOC, and SO₂ emissions from the target source were calculated by multiplying energy consumed per facility type by Japan's own emission factor or the IPCC default emission factor.

- **Emission factors**

- **NO_x**

For solid fuels (steaming coal and coal briquettes), emission factors were established by converting the default values provided in the *Revised 1996 IPCC Guidelines* to gross calorific values.

For liquid (kerosene) and gaseous (LPG, town gas) fuels, the emission factors per usage per fuel type provided in the reports by Air Quality Management Bureau, Ministry of the Environment were used. This report calculated the emission factors by weighting the average concentration of NO_x emissions per source unit, obtained through questionnaires and interviews in the household gas appliances industry.

- **CO**

For solid fuels (steaming coal and coal briquettes), emission factors were established by converting the default values provided in the *Revised 1996 IPCC Guidelines* to gross calorific values.

For liquid (kerosene) and gaseous (LPG, town gas) fuels, the emission factors per usage per fuel type provided in the reports by Institute of Behavioral Science were used. This report tabulated the emission factors by usage and fuel using the actual values measured in Tokyo, Yokohama city and Chiba Prefecture.

- **NMVOC**

For all of the solid (steaming coal and coal briquettes), liquid (kerosene), and gaseous (LPG and town gas) fuels, emission factors were established by converting the default values provided in the *Revised 1996 IPCC Guidelines* to gross calorific values.

- **SO₂**

For solid fuels (steaming coal and coal briquettes), emission factors were established by converting the default values provided in the *Revised 1996 IPCC Guidelines* to gross calorific values.

For liquid fuel (kerosene), emission factors were calculated from energy consumption, specific gravity and sulfur content based on the fuel characteristics of kerosene described in information material compiled by the Petroleum Association of Japan.

- **Activity data**

Consumption by type of fuel for residential use in *General Energy Statistics* has been taken for the activity data. The fuels covered were steaming coal, coal briquettes, kerosene, LPG, and town gas. For the amount of residential fuel consumption by type of use, the ratio of consumption by energy source

and by type of use per household, in the Handbook of Energy & Economic Statistics in Japan (The Energy Data and Modeling Center) is used.

A3.1.1.1.d. Incineration of waste for energy purposes and with energy recovery

Emissions of NO_x, CO, NMVOC and SO₂ from the incineration of waste for energy purposes and from the incineration of waste with energy recovery are reported in the data input cells for “Other Fuels” under the relevant subcategories of 1.A.1 and 1.A.2. Explanations for methodology for estimating emissions, emission factors, and activity data are all given in the section “3.1.5. Wastes”.

A3.1.1.2. Mobile Combustion (1.A.3: NO_x, CO, NMVOC, and SO₂)

A3.1.1.2.a. Road Transportation (1.A.3.b.)

1) NO_x, CO, and NMVOC

● Methodology for Estimating Emissions

NO_x, CO, and NMVOC emissions from the specified mobile sources were calculated by multiplying the distance traveled per year for each vehicle type per fuel by Japan’s own emission factor.

● Emission factors

Emission factors were established from the measured values for each vehicle class per fuel type (Ministry of the Environment). The NMVOC emission factors, however, were calculated by multiplying the emission factor of total hydrocarbon (THC) (per Ministry of the Environment) by the percentage of NMVOC in the THC emission (per Ministry of the Environment).

Table A 3-1 NO_x emission factors for automobiles

Fuel	Vehicle Type	Unit	1990	1995	2000	2005	2008	2009	2010
Gasoline	Light Vehicle	gNO _x /km	0.230	0.159	0.157	0.079	0.045	0.035	0.027
	Passenger Vehicle (including LPG)	gNO _x /km	0.237	0.203	0.199	0.080	0.047	0.037	0.028
	Light Cargo Truck	gNO _x /km	0.873	0.658	0.375	0.200	0.128	0.106	0.089
	Small Cargo Truck	gNO _x /km	1.115	0.897	0.478	0.087	0.042	0.032	0.025
	Regular Cargo Truck	gNO _x /km	1.833	1.093	0.560	0.162	0.061	0.043	0.032
	Bus	gNO _x /km	4.449	3.652	2.438	0.090	0.052	0.040	0.034
Diesel	Special Vehicle	gNO _x /km	1.471	0.873	0.429	0.121	0.052	0.037	0.029
	Passenger Vehicle	gNO _x /km	0.636	0.526	0.437	0.448	0.384	0.361	0.339
	Small Cargo Truck	gNO _x /km	1.326	1.104	1.005	1.009	0.829	0.744	0.658
	Regular Cargo Truck	gNO _x /km	5.352	4.586	4.334	4.497	4.028	3.759	3.422
	Bus	gNO _x /km	4.226	3.830	3.597	4.070	3.502	3.212	2.880
	Special Vehicle	gNO _x /km	3.377	2.761	2.152	3.626	3.164	2.923	2.633

Source: Ministry of the Environment

Table A 3-2 CO emission factors for automobiles

Fuel	Vehicle Type	Unit	1990	1995	2000	2005	2008	2009	2010
Gasoline	Light Vehicle	gCO/km	1.749	1.549	1.543	0.971	0.692	0.607	0.537
	Passenger Vehicle	gCO/km	2.325	2.062	2.034	0.936	0.667	0.582	0.509
	Light Cargo Truck	gCO/km	10.420	8.540	5.508	2.773	2.032	1.887	1.787
	Small Cargo Truck	gCO/km	9.656	10.079	8.309	2.075	1.013	0.785	0.607
	Regular Cargo Truck	gCO/km	12.624	10.601	8.950	3.616	1.601	1.208	0.941
	Bus	gCO/km	26.209	25.079	21.938	2.072	1.320	1.140	1.066
Diesel	Special Vehicle	gCO/km	12.466	10.666	8.924	2.298	1.138	0.886	0.746
	Passenger Vehicle	gCO/km	0.480	0.432	0.429	0.374	0.317	0.288	0.258
	Small Cargo Truck	gCO/km	0.975	0.896	0.808	0.601	0.413	0.343	0.284
	Regular Cargo Truck	gCO/km	3.221	2.988	2.440	2.042	1.437	1.205	0.995
	Bus	gCO/km	2.579	2.534	2.200	2.035	1.386	1.131	0.913
	Special Vehicle	gCO/km	2.109	1.893	1.297	1.601	1.075	0.881	0.713

Source: Ministry of the Environment

Table A 3-3 NMVOC emission factors for automobiles

Fuel	Vehicle Type	Unit	1990	1995	2000	2005	2008	2009	2010
Gasoline	Light Vehicle	gHC/km	0.128	0.050	0.048	0.043	0.027	0.023	0.019
		%	60%	60%	60%	60%	60%	60%	60%
		gNMVOC/km	0.077	0.030	0.029	0.026	0.016	0.014	0.012
	Passenger Vehicle (including LPG)	gHC/km	0.189	0.112	0.104	0.030	0.020	0.017	0.015
		%	60%	60%	60%	60%	60%	60%	60%
		gNMVOC/km	0.113	0.067	0.062	0.018	0.012	0.010	0.009
	Light Cargo Truck	gHC/km	1.058	0.610	0.274	0.151	0.096	0.079	0.066
		%	60%	60%	60%	60%	60%	60%	60%
		gNMVOC/km	0.635	0.366	0.165	0.091	0.058	0.048	0.040
	Small Cargo Truck	gHC/km	1.188	0.882	0.346	0.068	0.030	0.022	0.017
		%	60%	60%	60%	60%	60%	60%	60%
		gNMVOC/km	0.713	0.529	0.208	0.041	0.018	0.013	0.010
	Regular Cargo Truck	gHC/km	1.658	0.959	0.471	0.103	0.043	0.029	0.020
		%	60%	60%	60%	60%	60%	60%	60%
		gNMVOC/km	0.995	0.575	0.283	0.062	0.026	0.018	0.012
	Bus	gHC/km	3.604	3.164	2.193	0.065	0.029	0.023	0.020
		%	60%	60%	60%	60%	60%	60%	60%
		gNMVOC/km	2.162	1.899	1.316	0.039	0.017	0.014	0.012
Special Vehicle	gHC/km	1.619	0.786	0.317	0.081	0.035	0.025	0.020	
	%	60%	60%	60%	60%	60%	60%	60%	
	gNMVOC/km	0.972	0.472	0.190	0.048	0.021	0.015	0.012	
Diesel	Passenger Vehicle	gHC/km	0.109	0.098	0.097	0.089	0.078	0.072	0.066
		%	60%	60%	60%	60%	60%	60%	60%
		gNMVOC/km	0.065	0.059	0.058	0.053	0.047	0.043	0.040
	Small Cargo Truck	gHC/km	0.389	0.343	0.258	0.206	0.119	0.090	0.067
		%	60%	60%	60%	60%	60%	60%	60%
		gNMVOC/km	0.233	0.206	0.155	0.124	0.071	0.054	0.040
	Regular Cargo Truck	gHC/km	1.634	1.488	1.040	0.753	0.488	0.394	0.315
		%	60%	60%	60%	60%	60%	60%	60%
		gNMVOC/km	0.980	0.893	0.624	0.452	0.293	0.237	0.189
	Bus	gHC/km	1.273	1.255	0.995	0.807	0.495	0.381	0.291
		%	60%	60%	60%	60%	60%	60%	60%
		gNMVOC/km	0.764	0.753	0.597	0.484	0.297	0.229	0.175
Special Vehicle	gHC/km	1.101	0.965	0.526	0.575	0.350	0.276	0.216	
	%	60%	60%	60%	60%	60%	60%	60%	
	gNMVOC/km	0.661	0.579	0.316	0.345	0.210	0.165	0.129	

Top row: THC emission factors;

Middle row: Percentage of NMVOC in the THC emission;

Source: Ministry of the Environment

- **Activity data**

The activity data used the travel distance per year for each vehicle class per fuel type, which were calculated by multiplying distances traveled in a year for each vehicle class per fuel type, provided in the *Statistical Yearbook of Motor Vehicle Transport* (Ministry of Land, Infrastructure, Transport and

Tourism), by the percentage of the distances per fuel types calculated from fuel consumption and cost data.

2) SO₂

● *Methodology for Estimating Emissions*

The emissions of SO₂ from these sources were calculated by multiplying fuel consumption by vehicle class and fuel types by Japan's own emission factor.

● *Emission factor*

Sulfur content (by weight) of each fuel type was used to establish emission factors.

Table A 3-4 Sulfur content (by weight) by fuel type

		1990	1995	2000	2005	2008	2009	2010
Gasoline	%	0.008%	0.008%	0.008%	0.008%	0.008%	0.008%	0.008%
Diesel	%	0.350%	0.136%	0.136%	0.136%	0.136%	0.136%	0.136%
LPG	%	0.002%	0.002%	0.002%	0.002%	0.002%	0.002%	0.002%

Source: Gasoline/LPG – The Institute of Behavioral Science, Diesel oil – Petroleum Association of Japan

● *Activity data*

Activity data was calculated by multiplying fuel consumption for each vehicle class per fuel type by specific gravity of each fuel type, and converting the resultant values to weight. The fuel consumption data was reported in the *Statistical Yearbook of Motor Vehicle Transport* (Ministry of Land, Infrastructure, Transport and Tourism).

● *Completeness*

Emissions of NO_x, CO, NMVOCs, and SO₂ from natural gas vehicles and motorcycles are reported as “NE”.

A3.1.1.2.b. Civil Aviation (1.A.3.a: NO_x, CO, NMVOC)

● *Methodology for Estimating Emissions*

NO_x, CO, and NMVOC emissions from the specified sources were calculated by multiplying the fuel consumption converted to net calorific value by the default emission factors provided in the *Revised 1996 IPCC Guidelines*.

● *Emission factors*

The default emission factors provided for the “Jet and Turboprop Aircraft” category in the *Revised 1996 IPCC Guidelines* were used.

Table A 3-5 IPCC default emission factors for civil aviation

Gas	EF[g/MJ]
NO _x	0.29
CO	0.12
NMVOC	0.018

Source: Revised 1996 IPCC Guidelines, Vol. 3; Page 1.90, Table 1-47

● *Activity data*

Figures for jet fuel consumption (for domestic scheduled flights and others [commuter, sightseeing and charter flights]) in the *Statistical Yearbook of Air Transport* (Ministry of Land, Infrastructure,

Transport and Tourism) were converted to net calorific value for the calculation of activity data.

- **Completeness**

Emissions of NO_x, CO, and NMVOCs from aviation fuel consumption are reported as “NE”.

A3.1.1.2.c. Navigation (1.A.3.d.: NO_x, CO, NMVOC)

1) NO_x, CO, and NMVOC

- **Methodology for Estimating Emissions**

NO_x, CO, and NMVOC emissions from the specified sources were calculated by multiplying the fuel consumption converted to net calorific value by the default emission factors provided in the *Revised 1996 IPCC Guidelines*.

- **Emission factors**

The default emission factors provided for the “Ocean-Going Ships” category in the *Revised 1996 IPCC Guidelines* were used.

Table A 3-6 IPCC default emission factors for ocean-going ships

Gas	Emission factor [g/MJ]
NO _x	1.8
CO	0.18
NMVOC	0.052

Source: Revised 1996 IPCC Guidelines, Vol. 3; Page 1.90, Table 1-48

- **Activity data**

The marine fuel consumption data per fuel type (diesel, heating oil A, heating oil B, and heating oil C) provided in the *General Energy Statistics* (Agency for Natural Resources and Energy) were converted to net calorific value for the calculation of activity data. The consumption data were based on the statistical data on marine transport (coastal services [passenger and freight]) in the *The Survey on Transport Energy* (Ministry of Land, Infrastructure, Transport and Tourism).

2) SO₂

- **Methodology for Estimating Emissions**

Emissions from the specified sources were calculated by multiplying the fuel consumption by the emission factors.

- **Emission factors**

The multiplied product of the specific gravity of each marine fuel, the sulfur ratio of each fuel, and the molecular weight ratio of sulfur dioxide versus sulfur is used for the emission factor. The sulfur ratio of each fuel is restricted by law or Japanese Industrial Standard, therefore the regulation value is used for the sulfur ratio in the estimation.

Table A 3-7 Specific Gravity and Sulfur Ratio of Fuel for Ocean-Going Ships

Fuel	Specific Gravity[kg/l]	Sulfur Ratio[% in weight]
Diesel Oil	0.83	0.001
Fuel Oil A	0.84	2.0
Fuel Oil B	0.91	3.0
Fuel Oil C	0.93	3.5

Source: Sulfur ratio of diesel oil based on Act on the Quality Control of Gasoline and Other Fuels

Sulfur ratio of each fuel oil based on Japanese Industrial Standard K2205

Specific gravity based on Regulation of Total NO_x Emission Manual

● Activity data

The marine fuel consumption data per fuel type (diesel, heating oil A, heating oil B, and heating oil C) provided in the *General Energy Statistics* (Agency for Natural Resources and Energy) were used for the activity data. The consumption data were based on the statistical data on marine transport (coastal services [passenger and freight]) in the *The Survey on Transport Energy* (Ministry of Land, Infrastructure, Transport and Tourism).

A3.1.1.2.d. Railways (1.A.3.c.: NO_x, CO, and NMVOC)

● Methodology for Estimating Emissions

NO_x, CO, and NMVOC emissions from the specified sources were calculated by multiplying fuel consumption converted to net calorific value by the default emission factors provided in the *Revised 1996 IPCC Guidelines*.

● Emission factors

The default emission factors provided for the “Locomotives” category in the *Revised 1996 IPCC Guidelines* were used.

Table A 3-8 IPCC default emission factors for locomotives

Gas	Emission factor [g/MJ]
NO _x	1.8
CO	0.61
NMVOC	0.13

Source: Revised 1996 IPCC Guidelines, Vol. 3; Page 1.89, Table 1-47

● Activity data

The diesel oil consumption by railways in the *General Energy Statistics* (Agency for Natural Resources and Energy) was used for the calculation of activity data.

A3.1.1.3. Fugitive emissions from fuels (1.B.: NMVOC)

A3.1.1.3.a. NMVOCs fugitive emissions at oil refinery

● Methodology for Estimating Emissions

NMVOC emissions from the specified sources were calculated by multiplying the capacity of oil refineries (BPSD: Barrels Per Served Day) by Japan’s own emission factors and annual days of operation.

- **Emission factor**

Based on the *Study on the total system for prevention of HC-Vapor in petroleum industries* (Agency of Natural Resources and Energy, 1975), the emission factor was established as 0.05767 (g-NMVOC/BPSD). The number of days of operation for atmospheric distillation was established as 350 days.

- **Activity data**

Figures for the BPSD based on the results of surveys conducted by the Ministry of Economy, Trade and Industry, were used for the calculation of activity data.

A3.1.1.3.b. NMVOCs emissions from lubricant oil production

- **Methodology for Estimating Emissions**

NMVOC emissions from the specified sources were calculated by multiplying gross sales amount to consumers by Japan's own emission factors for toluene and methyl ethyl ketone.

- **Emission factors**

Based on internal documents of Yokohama city, emission factors were established for toluene and methyl ethyl ketone.

Table A 3-9 Toluene and methyl ethyl ketone emission factors in lubricant oil production

Gas	Emission factor (g/kl)
Toluene	333.2
Methyl ethyl ketone	415.5

Source: Yokohama city

- **Activity data**

Figures for gross sales amount to consumers, provided in the *Yearbook of Mineral Resources and Petroleum Production Statistics* (Ministry of Economy, Trade and Industry), were used for the calculation of activity data.

A3.1.1.3.c. NMVOCs fugitive emissions at storage facilities

- **Methodology for Estimating Emissions**

NMVOC emissions from the specified sources were calculated on the assumption that yearly emissions were the same as the 1983 volume of losses from breathing and acceptance for cone-roof type storage tanks and shipping losses from floating-roof type storage tanks at refineries and storage tanks (Petroleum Association of Japan).

- **Emission factor**

No emission factors were established.

- **Activity data**

No activity data were used.

A3.1.1.3.d. NMVOCs fugitive emissions at shipping facilities

● Methodology for Estimating Emissions

NMVOC emissions from specified sources were calculated by multiplying the 1983 figures for NMVOC emissions from ships and tank lorries/freight cars by the 1983 ratio of amount of shipment or that of sales to consumers.

● Emission factor

No emission factors were established.

● Activity data

Figures for shipment of crude oil not to be refined, gross sales amount of gasoline to consumers, export of gasoline, gross sales amount of naphtha to consumers, export of naphtha, gross sales amount of jet fuel to consumers and export of jet fuel provided in the *Yearbook of Mineral Resources and Petroleum Products Statistics* (Ministry of Economy, Trade and Industry) were used for the calculation of activity data. Table A3-9 shows the relationship between the NMVOC emission sources and activity data.

Table A 3-10 Relationship between the NMVOC emission sources and activity data

NMVOC emission source		Activity data used in calculation
Ships	Crude oil	shipment of crude oil not to be refined
	Gasoline	gross sales amount of gasoline to consumers
		export of gasoline
	Naphtha	gross sales amount of naphtha to consumers
export of naphtha		
Jet fuel	gross sales amount of jet fuel to consumers	
	export of jet fuel	
Tank lorries /Freight cars	Gasoline	gross sales amount of gasoline to consumers
	Naphtha	gross sales amount of naphtha to consumers
	Jet fuel	gross sales amount of jet fuel to consumers

A3.1.1.3.e. NMVOCs fugitive emissions from gas stations

● Methodology for Estimating Emissions

NMVOC emissions from specified sources were calculated by multiplying amount of sales to consumers by Japan's own emission factors for oil accepting and providing, and subtracting the portion of fuels prevented from fugitive emissions by a vapor return facility.

● Emission factor

Emission factors were established for oil accepting and for oil providing, based on the *Study on the total system for prevention of HC-Vapor in petroleum industries* (Agency of Natural Resources and Energy, 1975).

Table A 3-11 Emission factors at gas stations during oil accepting and providing

	Emission factor (kg/kl)
Oil accepting	1.08
Oil providing	1.44

Source: Study on the total system for prevention of HC-Vapor in petroleum industries (Agency of Natural Resources and Energy, 1975)

● **Activity data**

Figures for sales amount of gasoline (for automobiles) in the *Yearbook of Mineral Resources and Petroleum Products Statistics* (Ministry of Economy, Trade and Industry) were used for the calculation of activity data.

Fugitive emissions prevented by a vapor return facility during oil accepting at gas stations were calculated by the following equation:

<p><i>Calculation of fugitive emissions prevented by vapor return facility during oil accepting</i></p> <p>Fugitive emissions prevented by vapor return facility during oil accepting [t]</p> $= \sum \text{Prefecture} \{ (\text{gasoline sales per prefecture [ML]} \times \text{emission factor for oil accepting [kg/kl]}) \times (\text{No. of service stations with vapor return facility per prefecture} / \text{No. of service stations per prefecture}) \}$
--

Based on the data provided in the *Yearbook of Mineral Resources and Petroleum Products Statistics* (Ministry of Economy, Trade and Industry). For the number of service stations after FY 2001, the number of service stations registered under law was used.

A3.1.2. Industrial Processes

A3.1.2.1. Mineral Products, Chemical Industry, Metal Production, and Other Production (2.A., 2.B., 2.C., 2.D.; NO_x, SO₂)

● **Methodology for Estimating Emissions**

NO_x and SO₂ emissions from the specified sources were calculated for sources not included in the following facilities or operations by isolating the emissions from the Industrial Processes sector.

Facility: [0101– 0103: Boilers]; [0601– 0618: Metal rolling furnaces, metal furnaces, and metal forge furnaces]; [1101–1106: Drying ovens]; [1301–1304: Waste incinerators]; [2901–3202: Gas turbines, diesel engines, gas engines, and gasoline engines]

Operation: [A–D: Accommodation/eating establishments, health care/educational and academic institutions, public bathhouses, laundry services]; [F–L: Agriculture/fisheries, mining, construction, electricity, gas, heat distribution, building heating/other operations]

➤ **NO_x**

If raw material falls under either [44: Metallurgical coal] or [45: Metallurgical coke], the following equation is used:

<p><i>Calculation of NO_x emissions from metallurgical coal or coke (for Industrial Processes sector)</i></p> <p>NO_x emissions from metallurgical coal or coke [t-NO_x]</p> $= \text{NO}_x \text{ emission factor per origin [t-NO}_x\text{/kcal]} \times \text{energy consumed per material [kcal]} \times (1 - \text{denitrification rate [\%]})$
--

If raw material falls under either [41: Iron/ironstone] or [46: Other], the following equation is used:

<p><i>Calculation of NO_x emissions from iron/ironstone or other material (for Industrial Processes sector)</i></p> <p>NO_x emissions from iron/iron ore or other material [t-NO_x]</p> $= \text{Nitrogen content per material [t-NO}_x\text{]} \times (1 - \text{denitrification rate [\%]})$
--

If, however, the emissions from the Industrial Processes sector calculated by the above equations exceed the emission volume listed in the *General Survey of the Emissions of Air Pollutants*, the total emissions listed in the Survey are considered to be the emissions from the Industrial Processes sector.

Materials listed in the categories [42: Sulfide minerals] and [43: Non-ferrous metal ores] are excluded from the calculation due to the lack of data.

➤ **SO₂**

Based on the consumption and sulfur contents of the materials in the categories from [41: Iron/ironstone] to [46: Other materials], SO₂ emissions from the Industrial Processes sector are calculated as follows:

Calculation of SO₂ emissions (in the Industrial Processes sector)

SO₂ emissions [t-SO₂]

= Sulfur content per material [t-SO₂] × (1 – desulphurization rate [%])

● **Emission factor**

➤ **NO_x emission factors for metallurgical coal and coke**

NO_x emission factors for the materials used in calculation of NO_x emissions from metallurgical coal and coke (in the Industrial Processes sector) were established for each facility and material type based on the *General Survey of the Emissions of Air Pollutants*.

➤ **Denitrification rate**

The denitrification rate was calculated by the following equation:

Calculation of denitrification rate

Denitrification rate [%]

= Denitrification efficiency [%] × (Hours of operation of denitrification unit [h/yr]

/ Hours of operation of furnace [h/yr]) × (Processing capacity of denitrification unit [m³/yr]

/ max. exhaust gas emission [m³/yr])

The General Survey of the Emissions of Air Pollutants data were used for all items.

Denitrification efficiency: (NO_x volume before treatment – NO_x volume after treatment) / volume of smoke and soot

➤ **Desulphurization rate**

The desulphurization rate was calculated by the following equation:

Calculation of desulphurization rate

Desulphurization rate [%]

= Desulphurization efficiency [%] × (Hours operation of desulphurization unit [h/yr]

/ Hours operation of furnace [h/yr]) × (Processing capacity of desulphurization unit [m³/yr]

/ max. exhaust gas emission [m³/yr])

The General Survey of the Emissions of Air Pollutants data were used for all items.

Desulphurization efficiency: (SO₂ volume before treatment – SO₂ volume after treatment) / volume of smoke and soot

● **Activity data**

➤ **Energy consumption of metallurgical coal or coke**

The activity data was calculated by multiplying the consumption of materials (under [44: Metallurgical coal] and [45: Metallurgical coke]) provided in the *General Survey of the Emissions of Air Pollutants* by gross calorific value.

➤ **Nitrogen content of iron/ironstone and other materials**

The activity data was calculated by multiplying the weighted average of nitrogen content, calculated from the nitrogen content and consumption of the materials (under [41: Iron/ironstone] and [46: Other raw materials]) provided in the *General Survey of the Emissions of Air Pollutants*, by the consumption

volume of the material.

➤ **Sulfur content of various materials**

The activity data was calculated by multiplying the weighted average of sulfur content, calculated on the basis of sulfur content and consumption of the material (under [41: Iron/ironstone] through [46: Other materials]) provided in the *General Survey of the Emissions of Air Pollutants*, by the consumption volume of the material.

A3.1.2.2. Other (2.G.: NMVOC)

A3.1.2.2.a. NMVOCs emissions from petrochemical manufacturing

● **Methodology for Estimating Emissions**

NMVOCs emissions from petrochemical manufacturing were calculated by multiplying the production volume per type of petrochemical product by Japan's own emission factors.

● **Emission factors**

Emission factors were established based on the *Basic Study on HC Sources* (Institute of Behavioral Science, 1987).

Table A 3-12 NMVOC emission factors by petrochemical product

Petrochemical product	Emission factor (kg/t)
Propylene oxide	0.828
Vinyl chloride monomer	3.288
Styrene monomer	0.529
Vinyl acetate	1.299
B.T.X.	0.080
Ethylene oxide	0.421
Acrylonitrile	1.035
Butadiene	0.210
Polyethylene (produced under middle-low pressure)	1.851
Polyethylene (produced under high pressure)	1.088
ABS, AS resins	1.472
Synthetic rubber	0.248
Acetaldehyde	0.016
Terephthalic acid	0.534
Polypropylene	2.423
Ethylene and Propylene	0.016

Source: Basic Study on HC Sources (Institute of Behavioral Science, 1987).

● **Activity data**

Figures in the petrochemical production volume by type in the *Yearbook of Mineral Resources and Petroleum Products Statistics* (Ministry of Economy, Trade and Industry) were used for the calculation of activity data.

A3.1.2.2.b. NMVOCs emissions from storage facilities for chemical products

● **Methodology for Estimating Emissions**

NMVOCs emissions from storage facilities for chemical products were calculated on the assumption that the emission volumes were same as the 1983 combined yearly emissions of "Petrochemicals" and

“Others”, given in the *Basic Study on HC Sources* (Institute of Behavioral Science, 1987). “Petrochemicals” covered base chemicals (for the chemical industry); “Other” covered solvents (shipped primarily for non-feedstock use).

● **Emission factors**

No emission factors were established.

● **Activity data**

No activity data were calculated.

A3.1.2.2.c. NMVOCs emissions from shipping facilities for chemical products

● **Methodology for Estimating Emissions**

NMVOCs emissions from shipping facilities for chemical products were calculated on the assumption that the emission volumes were same as the 1983 combined yearly emissions of “Petrochemicals” and “Others”, shown in the *Basic Study on HC Sources* (Institute of Behavioral Science, 1987). “Petrochemicals” covered base chemicals (for the chemical industry); “Other” covered solvents (shipped primarily for non-feedstock use).

● **Emission factors**

No emission factor has been established.

● **Activity data**

No activity data has been established.

A3.1.3. Sectors that use solvents and other products

A3.1.3.1. NMVOCs emissions from paint solvent use (3.A.: NMVOC)

● **Methodology for Estimating Emissions**

Emissions of NMVOC were calculated by multiplying the consumption of solvent by the NMVOC emission rate (the percentage of NMVOC not removed but released into atmosphere).

● **Emission factors**

The NMVOC emission rate ($92.54[\%] = 100[\%] - 7.46[\%]$) calculated from the NMVOC removal rate (7.46[%]) estimated by the Ministry of the Environment (1983) was used as the emission factor.

● **Activity data**

Consumption of solvent was calculated by multiplying the 1990 data for solvent consumption per solvent type by the 1990 ratio of solvent consumption in paint production. The consumption data were extracted from the *Present condition and prospect about VOCs in Paint Industry* (Japan Paint Manufacturers Association). The solvent consumption ratio was provided in the *Yearbook of Chemical Industries Statistics* (Ministry of Economy, Trade and Industry). As the statistical records on solvent consumption in paint production were discontinued, the data for 2001 were substituted for values for years 2002 and beyond.

Calculation of annual consumption of paint solvent A in Year X

$$\begin{aligned} & \text{Annual consumption of paint solvent A in Year X [t]} \\ & = \text{Annual consumption of paint solvent A in 1990 [t]} \\ & \times (\text{Annual consumption of paint production solvent B in Year X [t]} \\ & / \text{Annual consumption of paint production solvent B in 1990 [t]}) \end{aligned}$$

Table A 3-13 Relationship of types of paint solvents and solvents for paint production used in calculation

Types of Paint Solvent (A)	Types of Paint Production Solvents Used in Calculation (B)
Aliphatic compound hydrocarbon	Mineral spirit
Alicyclic compound hydrocarbon	Toluene, xylene, and other aromatic hydrocarbon
Aromatic compound hydrocarbon	Toluene, xylene, and other aromatic hydrocarbon
Petroleum mixed solvent	Mineral spirit
Alcohol solvent	Alcohol solvent
Ether, Ether Alcohol solvent	Alcohol solvent
Ester solvent	Ester solvent
Ketone solvent	Ketone solvent
Chloric solvent	Solvent with a high boiling point
Other non-chloric solvent	Solvent with a high boiling point

A3.1.3.2. Degreasing, dry cleaning (3.B.: NMVOC)

A3.1.3.2.a. NMVOCs emissions from metal cleansing

- **Methodology for Estimating Emissions**

NMVOCs emissions from metal cleansing were calculated by multiplying the shipping amount of solvents (trichloro ethylene and tetrachloro ethylene) in degreasing by Japan's own emission factor.

- **Emission factors**

Emission factors were established as the ratio of emission to shipment ($0.66 \text{ [Mg/t]} = 88,014 / 133,000$), based on data for 1983 in the *Report on the Survey of Measures for Stationary Sources of Hydrocarbons* (Institute of Behavioral Science, 1991).

- **Activity data**

Shipping amount of solvents was calculated by multiplying the sales volume of trichloro ethylene and tetrachloro ethylene, provided in the *Yearbook of Chemical Industries Statistics* (Ministry of Economy, Trade and Industry), by the ratio of consumption for metal cleansing use to total consumption of organic chloric solvent (3 type) ($0.2 = 11,266 / 56,350$), shown in documents from the Perchlo Association.

A3.1.3.2.b. NMVOCs emissions from dry cleaning

- **Methodology for Estimating Emissions**

NMVOCs emissions from dry cleaning were calculated on the assumption that the volume of NMVOC emissions was the same as the volume of solvents used in dry cleaning (petroleum solvents and tetrachloro ethylene).

- **Emission factors**

No emission factors were established, as all the solvents used in dry cleaning were assumed to be

discharged into the atmosphere.

- **Activity data**

Estimates by the Institute of Cleaning Research were used for the calculation of the annual consumption of petroleum solvents and tetrachloro ethylene in 1990 and 1991.

Annual consumption in 1992 and in subsequent years was calculated by the following equation on the assumption that solvent consumption was proportional to the number of machines in operation:

<p><u>Calculation of annual consumption of solvents in Year X</u></p> <p>Annual consumption of solvents in Year X [t]</p> <p>= Σpetroleum-based solvent/tetrachloroethylene {annual consumption of petroleum solvents or tetrachloroethylene in 1991 [t] \times (the number of machines in operation in Year X / the number of machines in operation in 1991)}</p>

A3.1.3.3. Chemical products, manufacture and processing (3.C.: NMVOC)

A3.1.3.3.a. NMVOCs emissions from paint production

- **Methodology for Estimating Emissions**

NMVOCs emissions from paint production were calculated by multiplying the amount of solvent treated in paint production by Japan's own emission factors.

- **Emission factors**

Emission factors were established based on the *Manual to control HC emissions* (Air Quality Management Bureau, Ministry of the Environment, 1982).

Table A 3-14 Emission factors for solvents used as raw material for paints

Solvent	Emission factor (%)
Toluene	0.3
Xylene	0.2
Other aromatic hydrocarbon	0.2
Mineral spirit	0.2
Alcohol solvent	0.3
Ester solvent	0.3
Methyl isobutyl ketone	0.3
Other ketones	0.2
Solvent with a high boiling point	0.1

Source: Manual to control HC emissions (Air Quality Management Bureau, Ministry of the Environment, 1982)

- **Activity data**

Amount of solvent treated in paint production in the *Yearbook of Chemical Industries Statistics* (Ministry of Economy, Trade and Industry) was used for the calculation of activity data. The usage of ketone solvents was allocated to "Methyl isobutyl ketone" and "Other ketones" (with approx. 63% allocated to methyl isobutyl ketones), based on the interview survey results included in *Manual to control HC emissions* (Air Quality Management Bureau, Ministry of the Environment, 1982). For 2002 and subsequent years, the 2001 values were used because the statistics were discontinued.

A3.1.3.3.b. NMVOCs emissions from printing ink production

- **Methodology for Estimating Emissions**

NMVOCs emissions from printing ink production were calculated by multiplying amount of solvent treated in paint production, by Japan's own emission factors.

- **Emission factors**

Emission factors were established based on the results of surveys conducted by the Ministry of the Environment, as well as *Basic study on HC sources* (Institute of Behavioral Science, 1987).

Table A 3-15 Emission factors for solvents used as materials in printing ink

Solvent	Emission factor
Petroleum solvent ^{a)}	0.00033
Aromatics hydrocarbon ^{a)}	0.00108
Alcohol solvent ^{a)}	0.00105
Ester, ether solvent ^{b)}	0.00117

Source: a: Surveys by the Ministry of the Environment

b: Basic Study on HC sources (Institute of Behavioral Science, 1987) Activity data

- **Activity data**

Amount of solvent treated in paint production in the Yearbook of Chemical Industries Statistics (Ministry of Economy, Trade and Industry) were used for the calculation of activity data. For 2002 and subsequent years, the 2001 values were used because the statistics were discontinued.

A3.1.3.3.c. NMVOCs emissions from printing ink solvent use

- **Methodology for Estimating Emissions**

NMVOCs emissions from printing ink solvent use were calculated by multiplying the 1983 figures for NMVOC emissions from printing ink solvent use by the ratio of 1983 and each year about shipment amount of solvent.

- **Emission factor**

Emission factors were established as "0.3".

- **Activity data**

Shipment amount of solvent in the *Yearbook of Chemical Industries Statistics* (Ministry of Economy, Trade and Industry) were used for the calculation of activity data.

A3.1.3.3.d. NMVOCs emissions from polyethylene laminate

- **Methodology for Estimating Emissions**

NMVOCs emissions from polyethylene laminate were calculated on the assumption that the yearly emissions equaled the 1983 emissions data provided in the *Basic study on HC sources* (Institute of Behavioral Science, 1987)

- **Emission factor**

No emission factors were established.

- **Activity data**

No activity data were calculated.

A3.1.3.3.e. NMVOCs emissions from solvent-type adhesive use

- **Methodology for Estimating Emissions**

NMVOCs emissions from solvent-type adhesive use were assumed to equal the amount of solvents (xylene, toluene) used in adhesives.

- **Emission factors**

No emission factors were established as all the solvents used in adhesives were assumed to be discharged into the atmosphere.

- **Activity data**

Shipment amount of adhesive were calculated by multiplying amount of adhesives shipment by type (on calendar year basis), shown in the *Current survey report on adhesive* (Japan Adhesive Industry Association), by solvent content rate for each type shown in the *Current survey report on adhesive* (Japan Adhesive Industry Association).

Table A 3-16 Solvent content in adhesives by type

Adhesive	Solvent content (%)
Vinyl acetate resin solvent type	65
Other resin solvent type	50
CR solvent type	71
Other synthetic rubber solvent type	76
Natural rubber solvent type	67

Source: Current survey report on adhesive (Japan Adhesive Industry Association)

A3.1.3.3.f. NMVOCs emissions from gum solvent use

- **Methodology for Estimating Emissions**

NMVOCs emissions from gum solvent use were calculated by multiplying the consumption of solvents in rubber by NMVOC emission rate (the percentage of NMVOC not removed but released into atmosphere).

- **Emission factors**

The NMVOC emission rate ($92.7[\%] = 100[\%] - 7.3[\%]$) was used. This was calculated from the 1983 estimate of the NMVOC removal rate (7.3%), provided in the *Basic study on HC sources* (Institute of Behavioral Science, 1987).

- **Activity data**

The annual consumption of solvents in rubber was calculated by multiplying the consumption of petrol for solvent use by the ratio of the amount of rubber petrol use to total amount of gum solvent use ($0.42 = 21,139 / 50,641$). The consumption data were obtained either from the *Statistics of rubber products* (Ministry of Economy, Trade and Industry) or the results of surveys by the Japan Rubber Manufacturers Association; the usage rate was provided by the *Basic study on HC sources* (Institute of Behavioral Science, 1987).

A3.1.3.4. Other (3.D.: NMVOC)**A3.1.3.4.a. NMVOCs emissions from other solvent use for production**● **Methodology for Estimating Emissions**

NMVOCs emissions from other solvent use for production were calculated on the assumption that the yearly emissions equaled the 1983 emissions shown in the *Basic study on HC sources* (Institute of Behavioral Science, 1987).

● **Emission factor**

No emission factors were established.

● **Activity data**

No activity data were calculated.

A3.1.4. Agriculture**A3.1.4.1. Field burning of agricultural residues (4.F.)****A3.1.4.1.a. Rice Straw, Rice Chaff & Straw of Wheat, Barley, Oats and Rye (4.F.1.: CO)**● **Methodology for Estimating Emissions**

CO emissions from the specified sources were calculated by using Japan's own Methodology for Estimating Emissions shown below (Rye and oats were excluded from the estimate because there are no Japan-specific emission factors for them):

Calculation of CO emission from burning of rice straw, chaff, and wheat straw

CO emission from burning of rice and wheat straw and chaff [t-CH₄]
 = $\sum_{\text{rice straw, wheat straw, chaff}}$ (amount of rice or wheat straw or chaff burnt [t])
 × carbon content (dry weight) × percentage of carbon released as CO
 × mol ratio of CO to CO₂ in emitted gases

● **Emission factors**

Emission factors were established for each parameter based on the measured data available in Japan.

Table A 3-17 Carbon content of rice/wheat straw and chaff

	Carbon content	Note
Rice straw	0.356	Adopted the mean value between 0.369 ^a and 0.342 ^b .
Chaff	0.344	Value measured by Bando et al. ^a
Wheat straw	0.356	Assumed to be the same as for rice straw

Source: a: Bando, Sakamaki, Moritomi, and Suzuki, "Study of analysis of emissions from biomass burning" (from the 1991 Report on Studies on Comprehensive Promotion Cost of Environmental Studies (National Institute of Environmental Studies, 1992))

b: Y Miura and T Kan'no, "Emissions of trace gases (CO₂, CO, CH₄, and N₂O) resulting from rice straw burning", *Soil Sci. Plant Nutr.*, 43(4),849–854, 1997

Table A 3-18 Percentage of carbon emitted as CO from rice and wheat straw and chaff

	Percentage of carbon emitted as CO	Note
Rice straw	0.684	Adopted the median value between 0.8 ^a and 0.567 ^b .
Chaff	0.8	Value measured by Bando et al. ^a
Wheat straw	0.684	Assumed to be the same as for rice straw

Source: a: Bando, Sakamaki, Moritomi, and Suzuki, "Study of analysis of emissions from biomass burning" (from the 1991 Report on Studies on Comprehensive Promotion Cost of Environmental Studies (National Institute of Environmental Studies, 1992))

b: Y Miura and T Kan'no, "Emissions of trace gases (CO₂, CO, CH₄, and N₂O) resulting from rice straw burning", Soil Sci. Plant Nutr., 43(4),849–854, 1997

Table A 3-19 Mol ratio of CO to CO₂ in gases emitted from burning rice and wheat straw and chaff

	Mol ratio of CO to CO ₂ in emitted gas	Note
Rice straw	0.219	Adopted the mean value between values by a and b.
Chaff	0.255	Value measured by Bando et al. ^a
Wheat straw	0.219	Assumed to be the same as for rice straw

Source: a: Bando, Sakamaki, Moritomi, and Suzuki, "Study of analysis of emissions from biomass burning" (from the 1991 Report on Studies on Comprehensive Promotion Cost of Environmental Studies (National Institute of Environmental Studies, 1992))

b: Y Miura and T Kan'no, "Emissions of trace gases (CO₂, CO, CH₄, and N₂O) resulting from rice straw burning", Soil Sci. Plant Nutr., 43(4),849–854, 1997

● Activity data

Amounts of rice straw, chaff, and wheat straw burned were drawn from amounts used in 4.F.1. to calculate CH₄ and N₂O emissions from the burning of agricultural residue. Amounts of wheat straw burned were obtained by using the following equation.

$$\text{Amount of wheat/barley straw burned} = (\text{amounts of wheat and barley burned}) \times 0.5$$

Note: Based on expert judgment, the ratio of straw to chaff was set at 1:1.

A3.1.5. Land Use, Land-Use Change and Forestry

A3.1.5.1. Biomass burning (5(V))

● Methodology for Estimating Emissions

For CO and NO_x emissions due to biomass burning, Tier 1 method is used.

➤ Forest land

CO

$$bbGHG_f = L_{forestfires} \times ER$$

NO_x

$$bbGHG_f = L_{forestfires} \times ER \times NC_{ratio}$$

$bbGHG_f$: GHG emissions due to forest biomass burning

$L_{forestfires}$: Carbon released due to forest fires(tC/yr)

ER : Emission ratio (CO : 0.06, NO_x : 0.121)

NC_{ratio} : NC ratio

- **Emission Factor**

- **Emission ratio**

The following values are applied to emission ratios for CO and NO_x due to biomass burning.

CO: 0.06, NO_x: 0.121

(default value stated in the GPG-LULUCF, Table 3A.1.15)

- **NC ratio**

The following values are applied to NC ratio of NO_x.

NC ratio: 0.01 (default value stated in the GPG-LULUCF p.3.50)

- **Activity data**

For activity in Forest land, carbon released by forest fire is used. For detailed information, see the description on the activity data in section 7.14 in Chapter 7.

A3.1.6. Wastes

A3.1.6.1. Waste incineration (6.C.)

A3.1.6.1.a. Municipal Solid Waste Incineration (6.C.–)

- **Methodology for Estimating Emissions**

The NO_x, CO, NMVOC, and SO₂ emissions from the specified sources were calculated by multiplying the incineration amount of MSW in each incinerator type (Continuous Incinerators, Semi-continuous Incinerators, Batch type Incinerators, Gasification melting furnaces) by Japan's own emission factors. These emissions are categorized following the methods given in chapter 8 based on incinerations either with or without energy recovery. The former emissions are reported in the Energy sector, while the latter are reported in the Waste sector.

- **Emission factors**

- **NO_x, SO₂**

For incinerators, emission factors were established for each incinerator type by using the emission volume and volume of treated waste identified in the *General Survey of the Emissions of Air Pollutants*. (The categories of incinerator types included: [1301: Waste incinerator (municipal solid waste; continuous system)] and [1302: Waste incinerator (municipal solid waste; batch system)]). The incineration material was [53: Municipal solid waste].) It should be noted that while the *General Survey of the Emissions of Air Pollutants* classified the incinerators into two classes (Continuous and Batch), this report classifies incinerators into three classes (“Continuous”, “Semi-continuous”, and “Batch type”) by dividing the Continuous system and assigning those which operated for less than 3,000 hours to the “Semi-continuous” class.

For gasification melting furnaces, the value for Continuous Incinerators with a similar incineration method was used.

Table A 3-20 NO_x and SO₂ emission factors for municipal waste incineration by facility type

	Item	Unit	単位	1990	2000	2005	2008	2009	2010
NO _x	Continuous Incinerator	kg-NO _x /t	kg-NO _x /t	1.238	1.127	1.127	1.127	1.127	1.127
	Semi-Continuous Incinerator	kg-NO _x /t	kg-NO _x /t	1.055	1.226	1.226	1.226	1.226	1.226
	Batch type Incinerator	kg-NO _x /t	kg-NO _x /t	1.137	1.850	1.850	1.850	1.850	1.850
	Gasification melting furnace	kg-NO _x /t	kg-NO _x /t	1.238	1.127	1.127	1.127	1.127	1.127
SO ₂	Continuous Incinerator	kg-SO ₂ /t	kg-SO ₂ /t	0.555	0.361	0.361	0.361	0.361	0.361
	Semi-Continuous Incinerator	kg-SO ₂ /t	kg-SO ₂ /t	0.627	0.712	0.712	0.712	0.712	0.712
	Batch type Incinerator	kg-SO ₂ /t	kg-SO ₂ /t	1.073	1.714	1.714	1.714	1.714	1.714
	Gasification melting furnace	kg-SO ₂ /t	kg-SO ₂ /t	0.555	0.361	0.361	0.361	0.361	0.361

The data after 2000 were used for 2001 and subsequent years.

Source: Research of Air Pollutant Emissions from Stationary Sources (Ministry of the Environment)

➤ CO

For incinerators, based on the emission factors for individual facilities summarized in the *Reports on Greenhouse gas emissions estimation methodology* (Japan Sociality Atmospheric Environment, 1996) as well as other reports, the emission factors were established for each incinerator class. It should be noted that while the Atmospheric Environment Society report subdivided the facilities by furnace type (e.g., stoker, fluidized bed, etc.), this report determined the emission factors for three classes of “Continuous”, “Semi-continuous” and “Batch type” by weighting the average of incinerated volume for each furnace.

For gasification melting furnaces, the value for continuous stoker furnaces with a similar incineration method was used.

Table A 3-21 CO emission factors for municipal waste incineration by facility type

	Furnace Type	Unit	単位	1990	2000	2005	2008	2009	2010
CO	Continuous Incinerator	gCO/t	gCO/t	557	555	554	554	554	554
	Semi-Continuous Incinerator	gCO/t	gCO/t	548	567	591	613	605	605
	Batch type Incinerator	gCO/t	gCO/t	8,237	8,298	8,341	8,343	8,351	8,351
	Gasification melting furnace	gCO/t	gCO/t	567	567	567	567	567	567

Source: Reports on Greenhouse gas emissions estimation methodology (Japan Sociality Atmospheric Environment, 1996), and others.

➤ NMVOC

For both incinerators and gasification melting furnaces, NMVOC emission factors were established by multiplying the CH₄ emission factors for each furnace type per fuel type by “NMVOC/CH₄”, the emission ratio for fuel type. The ratio was determined by using the reference material by Japan Environmental Sanitation Center and Institute of Behavioral Science, which estimated CH₄ and NMVOC emissions per unit calorific value.

Table A 3-22 NMVOC emission factors for municipal waste incineration by facility type

	Furnace Type	Unit	単位	1990	2000	2005	2008	2009	2010
NMVOC	Continuous Incinerator	gNMVOC/t	gNMVOC/t	0.9	0.9	0.3	0.3	0.3	0.3
	Semi-Continuous Incinerator	gNMVOC/t	gNMVOC/t	7.8	8.5	2.2	2.4	2.3	2.3
	Batch type Incinerator	gNMVOC/t	gNMVOC/t	9.1	9.5	1.5	1.5	1.5	1.5
	Gasification melting furnace	gNMVOC/t	gNMVOC/t	-	0.6	0.8	0.8	0.8	0.8

Source: Report on Screening Survey Regarding Measures to Counter Global Warming (Japan Environmental Sanitation Center, 1989), Study of Establishment of Methodology for Estimation of Hydrocarbon Emissions (Institute of Behavioral Science, 1984)

● Activity data

For incinerators, the activity data used was the incineration volume for each facility type as calculated by multiplying the incineration volume of municipal waste by the incineration rate for each facility type. The incineration volume data were extracted from the *Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes (the Volume on Cyclical Use)* by the Ministry of the Environment. The incineration rate was calculated in the *Waste Treatment in Japan* published by the Ministry of the Environment.

For gasification melting furnaces, the activity data used was the volume incinerated in gasification melting furnaces, calculated from data in the Ministry of the Environment's "*Waste Treatment in Japan*."

A3.1.6.1.b. Industrial Wastes Incineration (6.C.–)

● Methodology for Estimating Emissions

NO_x, CO, NMVOC, and SO₂ emissions from the specified sources were calculated by multiplying the incineration amount of industrial waste for each waste type by Japan's own emission factors. These emissions are categorized following the methods given in chapter 8 based on incinerations either with or without energy recovery. The former emissions are reported in the Energy sector, while the latter are reported in the Waste sector.

● Emission factors

➤ NO_x, SO₂

An emission factor was established for each type of industrial solid waste using the emission volume and volume of treated industrial solid waste identified by the *General Survey of the Emissions of Air Pollutants*. The categories of incinerator types included: [1303: Waste incinerator (industrial solid waste; continuous system)] and [1304: Waste incinerator (industrial solid waste; batch system)]. The incinerator fuel covered the categories [23: Fuel Wood] and [54: Industrial solid waste]). The six types of industrial waste were "Waste paper or waste wood", "Sludge", "Waste oil", "Waste plastics", "Waste textiles", and "Animal/plant residue, livestock carcasses". Category [23: Sawn Timber] was used for "Waste paper or waste wood", "Waste textiles", and "Animal/plant residues, livestock carcasses", while category [54: Industrial waste] was used for "Sludge", "Waste oil", and "Waste plastics". However, no emission factor was set for the mixed burning of multiple waste types.

Table A 3-23 NO_x and SO₂ emission factors for industrial waste by facility type

	Item	Unit	単位	1990	2000	2005	2008	2009	2010
NO _x	"Fuel Wood 23"	kg-NO _x /t	kg-NO _x /t	1.545	5.828	5.828	5.828	5.828	5.828
	"Industrial Waste 54"	kg-NO _x /t	kg-NO _x /t	0.999	1.415	1.415	1.415	1.415	1.415
SO ₂	"Fuel Wood 23"	kg-SO ₂ /t	kg-SO ₂ /t	1.528	2.118	2.118	2.118	2.118	2.118
	"Industrial Waste 54"	kg-SO ₂ /t	kg-SO ₂ /t	1.179	1.352	1.352	1.352	1.352	1.352

The data after 2000 were used for 2001 and subsequent years.

Source: Research of Air Pollutant Emissions from Stationary Sources (Ministry of the Environment)

➤ CO

Based on the emission factors for individual facilities summarized in the *Reports on Greenhouse gas emissions estimation methodology* (Japan Sociality Atmospheric Environment, 1996) as well as other reports, an emission factor was established for each type of industrial solid waste. The six types of industrial waste were "Waste paper or waste wood", "Sludge", "Waste oil", "Waste plastics", "Waste textiles", and "Animal/plant residues, livestock carcasses". The emission factor for "wood waste" was used for "Waste textiles" and "Animal/plant residues, livestock carcasses", for which there are no measurements. No emission factor was set for the mixed burning of multiple waste types.

Table A 3-24 CO emission factors for industrial waste incinerators by operation type

Item	Unit	単位	1990	2000	2005	2008	2009	2010
Waste Paper, Waste Wood	gCO/t	gCO/t	1,334	1,334	1,334	1,334	1,334	1,334
Waste Oil	gCO/t	gCO/t	127	127	127	127	127	127
Waste Plastics	gCO/t	gCO/t	1,790	1,790	1,790	1,790	1,790	1,790
Sludge	gCO/t	gCO/t	2,285	2,285	2,285	2,285	2,285	2,285
Waste textile	gCO/t	gCO/t	1,334	1,334	1,334	1,334	1,334	1,334
Animal and Plant residues	gCO/t	gCO/t	1,334	1,334	1,334	1,334	1,334	1,334

Source: Reports on Greenhouse gas emissions estimation methodology (Japan Sociality Atmospheric Environment, 1996) and others

➤ NMVOC

NMVOC emission factors were established by multiplying the CH₄ emission factors for each furnace type per fuel type by "NMVOC/CH₄", the emission ratio for fuel type. The ratio was determined by using the reference materials by Japan Environmental Sanitation Center and Institute of Behavioral Science, which estimated CH₄ and NMVOC emissions per unit calorific value.

Table A 3-25 NMVOC emission factors for industrial waste incineration by facility type

Item	Unit	単位	1990	2000	2005	2008	2009	2010
Waste Paper, Waste Wood	gNMVOC/t	gNMVOC/t	2.48	2.48	25.28	25.28	25.28	25.28
Waste Oil	gNMVOC/t	gNMVOC/t	0.54	0.54	0.45	0.45	0.45	0.45
Waste Plastics	gNMVOC/t	gNMVOC/t	3.40	3.40	0.90	0.90	0.90	0.90
Sludge	gNMVOC/t	gNMVOC/t	1.61	1.61	0.17	0.17	0.17	0.17
Waste textile	gNMVOC/t	gNMVOC/t	2.48	2.48	25.28	25.28	25.28	25.28
Animal and Plant residues	gNMVOC/t	gNMVOC/t	2.48	2.48	25.28	25.28	25.28	25.28

Source: Report on Screening Survey Regarding Measures to Counter Global Warming (Japan Environmental Sanitation Center, 1989)

Study of Establishment of Methodology for Estimation of Hydrocarbon Emissions (Institute of Behavioral Science, 1984)

Activity Data The activity data used the incineration volume data for each type of waste extracted from

the Report of the Research on the State of Wide-range Movement and Cyclical Use of Wastes (the Volume on Cyclical Use) and the Waste Treatment in Japan published by the Ministry of the Environment.

A3.1.6.1.c. Incineration in Conjunction with Use of Waste as Fuel and Raw Material (1.A.-)

● *Methodology for Estimating Emissions*

CO and NMVOC emissions from this source were estimated by multiplying the amounts of fuel/raw material burned for each waste type by a Japan-specific emission factor. These emissions are reported in Energy sector (1.A.) following the methodologies given in chapter 8 (Waste).

● *Emission Factors*

➤ *CO*

The CO emission factors (fixed unit basis) for furnace types, which are used for counting emissions from 1A Stationary Sources, were determined by using the calorific values in General Energy Statistics to convert to weight-based emission factors. For the calorific values of waste tires from FY2005 and on, values from the Agency for Natural Resources and Energy's "The Reexamination of Standard Calorific Values and Their Revised Values to Be Applied from FY2005 and on" (2007) were used.

Table A 3-26 CO emission factors from incineration in conjunction with use of waste as fuel and raw material

Application	Units	Waste oil	RDF	RPF	Waste tires (FY2004 and before)	Waste tires (FY2005 and after)	Waste plastics	Waste wood
Simple incineration	kgCO/t	0.13	1.79	1.79	1.79	1.79	-	-
Boilers	kgCO/t	0.052	0.24	0.39	0.28	0.44	0.034	3.64
Cement kilns	kgCO/t	49.1	19.8	32.2	23.0	36.5	32.2	-
Other furnaces	kgCO/t	0.052	0.24	0.39	0.28	0.44	-	-
Pyrolysis furnaces	kgCO/t	-	-	-	0.021	0.033	-	-
Gasification	kgCO/t	-	-	-	0.015	0.024	-	-

➤ *NMVOC*

Just as for the incineration of municipal solid waste and industrial waste, emission factors were determined from documents with estimates of emissions of CH₄ and NMVOCs per unit calorific values.

Table A 3-27 NMVOC emissions factors from incineration in conjunction with use of waste as fuel and raw material

Application	Units	Waste oil	RDF	RPF	Waste tires (FY2004 and before)	Waste tires (FY2005 and after)	Waste plastics	Waste wood
Boilers	kgNMVOC/t	0.015	0.00027	0.00043	0.00031	0.00049	0.010	0.12
Cement kilns	kgNMVOC/t	0.048	-	0.043	0.031	0.049	0.043	-
Pyrolysis furnaces	kgNMVOC/t	-	-	-	0.0051	0.0080	-	-
Gasification	kgNMVOC/t	-	-	-	0.0089	0.0141	-	-

● *Activity data*

We used the same activity data that were used when estimating CH₄ emissions from the use of waste

as fuel and raw material.

A3.1.7. Other sectors

A3.1.7.1. Smoking (7.-: CO)

- ***Methodology for Estimating Emissions***

CO emissions were calculated by multiplying the volume of cigarette sales by Japan's own emission factor.

- ***Emission factor***

The emission factor (0.055 [g-CO/cigarette]) was provided by Japan Tobacco Inc.

- ***Activity data***

The volume of cigarette sales published on Tobacco Institute of Japan website (<http://www.tioj.or.jp/>) was used for activity data.

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Annex 4. CO₂ Reference Approach and Comparison with Sectoral Approach, and Relevant Information on the National Energy Balance

This chapter explains a comparison between reference approach and sectoral approach in accordance with the UNFCCC Reporting Guidelines on Annual Inventories (FCCC/SBSTA/2006/9, paragraph 31).

A4.1. Difference in Energy Consumption

As shown in Table A 4-1, fluctuations of difference of energy consumption between the reference approach and the sectoral approach during 1990-2010 range between -3.11% and -0.28%. It is relatively low compared to the inventories from other countries.

Energy consumption from wastes used for energy and from the incineration of wastes with energy recovery, which had been reported as NE (Not Estimated) in previous submissions, are calculated in the sectoral approach from the 2011 inventory submission in accordance with the *1996 Revised IPCC Guidelines* and the *GPG (2000)*. Therefore, the energy consumption from sectoral approach and the difference in energy consumption between the reference approach and the sectoral approach are changed from the 2011 inventory submission.

Difference of solid fuels in 2008 was quite a large value (5.91%), because of coal (Imported Steam Coal [130^1]) stock change increasing.

Table A 4-1 Comparison of Energy Consumption

[10 ¹⁵ J]	1990	1995	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010
Reference Approach													
Liquid fuels	9,689	10,191	9,503	9,200	9,211	9,167	8,926	8,913	8,468	8,528	7,850	7,174	7,267
Solid fuels	3,270	3,603	4,175	4,267	4,409	4,534	4,967	4,736	4,796	5,010	4,894	4,354	4,940
Gaseous fuels	2,097	2,534	3,130	3,126	3,215	3,365	3,354	3,388	3,746	4,082	4,013	3,975	4,228
Other fuels	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
Total RA	15,056	16,328	16,809	16,593	16,835	17,066	17,246	17,037	17,010	17,620	16,757	15,503	16,436
Sectoral Approach													
Liquid fuels	9,550	10,051	9,450	9,133	9,275	9,094	8,934	8,903	8,390	8,402	7,726	7,103	7,185
Solid fuels	3,354	3,635	4,118	4,220	4,484	4,605	4,721	4,808	4,787	4,955	4,621	4,402	4,737
Gaseous fuels	2,106	2,548	3,136	3,137	3,238	3,371	3,371	3,368	3,756	4,106	4,021	4,011	4,238
Other fuels	259	294	348	359	379	408	416	436	438	444	437	415	418
Total	15,268	16,529	17,052	16,848	17,375	17,478	17,443	17,515	17,371	17,907	16,805	15,931	16,577
Difference (%)													
Liquid fuels	1.46%	1.39%	0.56%	0.74%	-0.69%	0.80%	-0.10%	0.10%	0.93%	1.50%	1.60%	1.00%	1.14%
Solid fuels	-2.50%	-0.88%	1.39%	1.10%	-1.65%	-1.54%	5.20%	-1.51%	0.19%	1.11%	5.91%	-1.08%	4.30%
Gaseous fuels	-0.44%	-0.55%	-0.20%	-0.32%	-0.72%	-0.19%	-0.50%	0.62%	-0.28%	-0.57%	-0.18%	-0.91%	-0.22%
Other fuels	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
Total	-1.39%	-1.21%	-1.43%	-1.51%	-3.11%	-2.36%	-1.13%	-2.73%	-2.08%	-1.60%	-0.28%	-2.68%	-0.85%

¹ Code number of the *General Energy Statistics* (Energy Balance Table)

A4.2. Difference in CO₂ Emissions

As shown in Table A 4-2, fluctuations of a difference of CO₂ emissions between the reference approach and the sectoral approach during 1990-2010 range between -1.92% and 2.01%. Emissions from wastes used for energy and from the incineration of wastes with energy recovery, which had been reported in waste sector (6.C.) in previous submissions, are reported in the energy sector (1.A.) from the 2009 inventory submission in accordance with the *1996 Revised IPCC Guidelines* and the *GPG (2000)*.

Difference of solid fuels in 2008 was quite a large value (5.26%), because of coal (Imported Steam Coal [S130]) stock change increasing.

Table A 4-2 Comparison of CO₂ Emissions

[Tg CO ₂]	1990	1995	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010
Reference Approach													
Liquid fuels	659.1	692.4	647.0	626.3	626.7	623.9	607.8	606.4	575.7	580.5	534.5	488.8	495.3
Solid fuels	294.6	324.2	377.6	385.5	399.0	410.3	450.0	428.7	434.2	453.7	442.6	394.1	447.5
Gaseous fuels	103.7	125.3	154.8	154.6	159.0	166.4	165.8	167.6	185.2	201.9	198.5	196.6	209.1
Other fuels	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
Total RA	1,057	1,142	1,179	1,166	1,185	1,201	1,224	1,203	1,195	1,236	1,176	1,080	1,152
Sectoral Approach													
Liquid fuels	646.2	677.3	635.1	613.1	622.9	611.4	600.4	597.8	562.0	563.7	518.4	475.0	481.1
Solid fuels	308.6	331.7	376.5	384.9	409.6	419.7	431.1	437.9	436.7	451.5	420.5	401.6	431.5
Gaseous fuels	104.3	126.2	155.3	155.3	160.4	167.0	166.9	166.8	186.4	203.3	199.5	198.7	210.8
Other fuels	9.1	10.5	13.1	14.2	15.0	15.8	15.6	15.1	14.2	14.4	14.0	13.9	14.2
Total	1,068	1,146	1,180	1,167	1,208	1,214	1,214	1,218	1,199	1,233	1,152	1,089	1,138
Difference (%)													
Liquid fuels	1.99%	2.23%	1.87%	2.17%	0.62%	2.05%	1.22%	1.43%	2.44%	2.98%	3.11%	2.91%	2.94%
Solid fuels	-4.54%	-2.26%	0.29%	0.17%	-2.60%	-2.24%	4.38%	-2.11%	-0.57%	0.49%	5.26%	-1.85%	3.72%
Gaseous fuels	-0.57%	-0.71%	-0.32%	-0.45%	-0.88%	-0.40%	-0.65%	0.45%	-0.61%	-0.69%	-0.53%	-1.07%	-0.81%
Other fuels	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
Total	-1.01%	-0.33%	-0.06%	-0.08%	-1.92%	-1.10%	0.79%	-1.24%	-0.34%	0.26%	2.01%	-0.88%	1.26%

A4.3. Comparison between Differences in Energy Consumption and that of CO₂ Emissions

The difference in energy consumption and the difference in CO₂ emissions generally show a similar tendency for their trends.

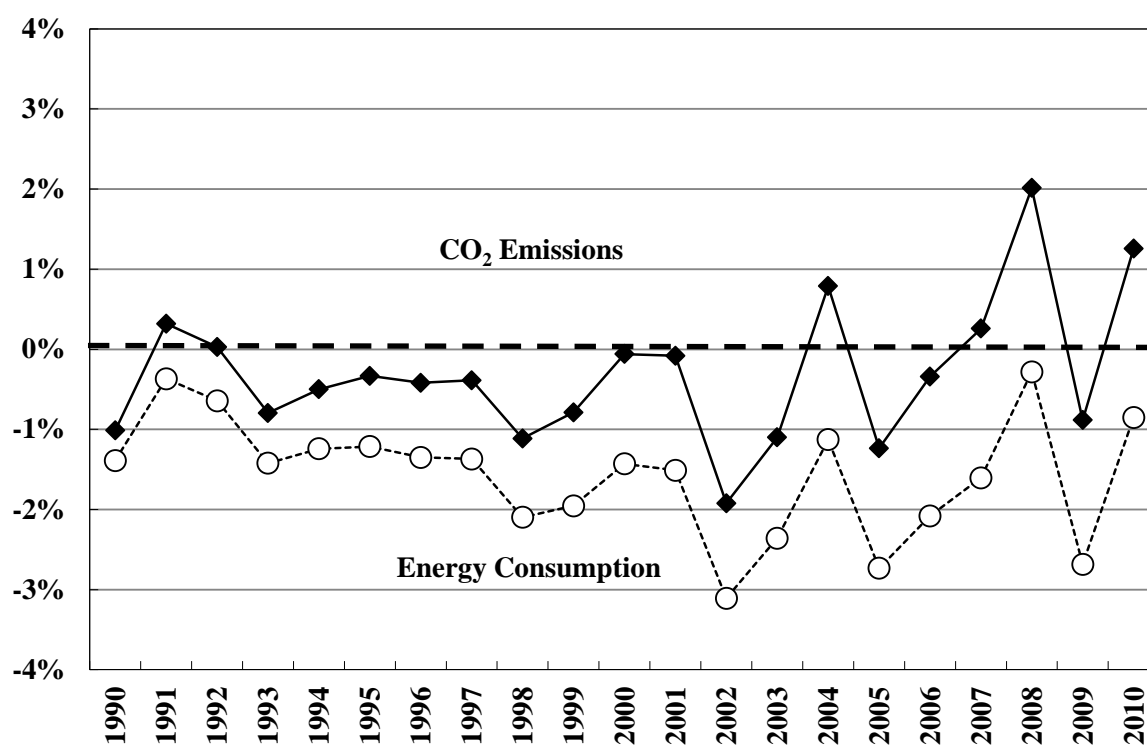


Figure A 4-1 Trends in Difference of Energy Consumption and CO₂ Emissions

A4.4. Causes of the difference between Reference Approach and Sectoral Approach

The difference in energy consumption and in CO₂ emissions can be explained by the difference of the amount of carbon which were deducted as feedstock and non-energy use in each approach, and ‘Other Conversions & Blending’ [#2700], ‘Other Input/Output’ [#3000], ‘Stock Change’ [#3500], ‘Statistical Discrepancy’ [#4000], and “energy loss” and “carbon imbalance” of ‘Oil Products’ [#2600] of the Energy Balance Table (*General Energy Statistics*).

The default values given in the *Revised 1996 IPCC Guidelines* were used for the fractions of carbon stored for feedstock and non-energy in reference approach.

1) Matters not sufficiently considered in the calculation process of Reference Approach

In the current estimation of reference approach, it was assumed that the amount of energy subtracted the energy amount for non-energy use from the national energy amount supplied was completely combusted. However, in real situations, some of the energy amount combusted is left without being combusted. The increase or decrease of the remaining energy amount were not considered in the current estimation of reference approach.

【Other Input/Output [#3000]】

In oil refining and other parts of the energy conversion sector, energy source shipment/drawdown amounts do not necessarily match production/receipt amounts. Other than energy received through one's own imports or that produced by refining, factors involved include returns from consumption/sales sectors of products once shipped, transactions of small amounts of byproduct energy from other companies, stock buildups and drawdowns due to product storage tank installation or decommissioning at factories and business sites, and losses due to accidents or fires.

When energy source inconsistencies due to such causes in the energy conversion sector are determined, the other input/output sector accounts for the amount. However, this input/output are not reflected under reference approach emission calculation.

【Stock Change [#3500]】

The increase or decrease of stock were not reflected under reference approach emission calculation.

CO₂ emissions from wastes used for energy and from the incineration of wastes with energy recovery originate from carbon in waste oil, waste plastics, waste tire, synthetic textile scrap and other non-biogenic waste which were incinerated. These amounts of carbons may not be reflecting the actual conditions in the deduction of carbon for feedstock and non-energy use in the calculation of the reference approach. The methodology for calculating the amount of stored carbon as feedstock and non-energy use in the reference approach should be examined and revised in the future.

2) Matters which cannot be avoided for the characteristics of survey data

【Statistical Discrepancy [#4000]】

Statistical discrepancy is originally the intrinsic error arising at the sampling stage in statistical studies (source error), and mutual discrepancies among the statistics for supply, conversion, and consumption. It is sometimes difficult to guess where discrepancies come from (relative error).

These errors induce the discrepancies among domestic supply, conversion, and final energy consumption, calculated as difference between both approaches.

3) Matters related to the difference of energy and carbon balance between energy input and output

【Other Conversions & Blending [#2700]】

This sector represents energy conversion that does not belong to any of the sectors from #2100 Commercial Power Generation to #2600 Oil Products, and actions considered to be energy conversion in which coal or oil product brands are changed by only simple operations such as blending or moisture adjustment.

Carbon weight is considered to be consistent before and after blending or conversions. However, given that carbon content per calorific value is changed following such as blending, in statistics, carbon weight could be varied before and after blending or conversions. This difference can generate the variation between two approaches.

【Oil Products [#2600]】

Energy loss and carbon imbalance during the process of oil production produce the difference between

input and output of energy or carbon.

Table A 4-3 Comparison of CO₂ emissions (detail)

	[Gg-CO ₂]												
	1990	1995	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010
RA	1,057,427	1,141,966	1,179,346	1,166,441	1,184,667	1,200,526	1,223,561	1,202,642	1,195,192	1,236,089	1,175,623	1,079,518	1,151,863
Liquid fuels	659,104	692,444	646,974	626,340	626,747	623,890	607,770	606,374	575,734	580,471	534,521	488,831	495,250
Solid fuels	294,611	324,221	377,604	385,525	398,965	410,252	449,953	428,702	434,223	453,747	442,626	394,132	447,538
Gaseous fuels	103,711	125,302	154,767	154,575	158,955	166,384	165,837	167,566	185,235	201,872	198,476	196,554	209,075
Other fuels	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
SA	1,068,260	1,145,769	1,180,044	1,167,384	1,207,886	1,213,888	1,213,986	1,217,696	1,199,277	1,232,913	1,152,418	1,089,142	1,137,551
Liquid fuels	646,223	677,349	635,121	613,057	622,889	611,372	600,423	597,813	562,037	563,675	518,395	474,999	481,120
Solid fuels	308,620	331,720	376,521	384,881	409,624	419,659	431,080	437,937	436,698	451,548	420,521	401,560	431,476
Gaseous fuels	104,301	126,198	155,261	155,279	160,359	167,045	166,918	166,823	186,374	203,273	199,525	198,684	210,774
Other fuels	9,116	10,503	13,142	14,167	15,014	15,812	15,565	15,123	14,168	14,417	13,976	13,899	14,180
RA-SA	-10,833	-3,803	-698	-943	-23,219	-13,362	9,575	-15,055	-4,085	3,176	23,205	-9,625	14,312
Liquid fuels	12,881	15,095	11,854	13,284	3,858	12,519	7,348	8,560	13,697	16,795	16,126	13,832	14,130
Solid fuels	-14,009	-7,499	1,084	644	-10,659	-9,407	18,873	-9,235	-2,475	2,199	22,105	-7,428	16,062
Gaseous fuels	-589	-896	-494	-704	-1,404	-662	-1,081	743	-1,139	-1,402	-1,050	-2,130	-1,700
Other fuels	-9,116	-10,503	-13,142	-14,167	-15,014	-15,812	-15,565	-15,123	-14,168	-14,417	-13,976	-13,899	-14,180
Statistical Discrepancy	-10,465	3,381	-1,258	-1,504	-12,510	-9,485	-3,088	-19,607	8,471	8,797	12,460	4,853	19,717
Liquid fuels	-3,708	3,839	-5,664	-5,292	-12,641	-10,667	-15,985	-15,724	2,881	2,443	1,239	814	2,322
Solid fuels	-6,796	-693	3,915	3,343	-320	836	12,409	-4,361	6,111	6,428	11,586	4,663	18,224
Gaseous fuels	39	236	491	446	450	346	488	478	-521	-73	-366	-624	-830
Other Conversions & Blending	-2,828	-3,076	-1,189	-1,277	-782	-775	-601	-1,110	-1,233	-1,475	-1,134	-979	-1,404
Liquid fuels	803	1,058	1,119	1,091	1,136	1,171	1,161	1,193	1,151	1,093	1,082	1,055	986
Solid fuels	-2,807	-3,078	-1,121	-1,168	-709	-709	-546	-1,059	-1,131	-1,361	-1,044	-901	-1,284
Gaseous fuels	-825	-1,056	-1,186	-1,201	-1,210	-1,237	-1,216	-1,244	-1,253	-1,206	-1,172	-1,134	-1,106
Stock Change	1,452	1,878	2,225	4,268	-8,722	-6,234	9,121	556	-2,851	-2,625	15,694	-9,876	2,749
Liquid fuels	788	1,311	-976	1,209	-3,753	-1,853	-2,369	270	2,234	-1,292	1,740	-689	457
Solid fuels	681	757	2,934	2,912	-4,286	-4,504	12,005	-1,097	-5,567	-990	13,635	-8,585	2,369
Gaseous fuels	-18	-190	268	148	-683	123	-515	1,383	482	-344	318	-602	-78
Other Input/Output	-895	-642	2,106	623	1,878	2,010	1,625	2,577	-1,385	1,174	1,374	1,429	2,504
Liquid fuels	-895	-642	2,106	623	1,878	2,010	1,625	2,577	-1,385	1,174	1,374	1,429	2,504
Solid fuels	0	0	0	0	0	0	0	0	0	0	0	0	0
Gaseous fuels	0	0	0	0	0	0	0	0	0	0	0	0	0
Oil Products	1,257	1,057	6,121	8,664	9,025	10,777	8,166	10,182	875	4,019	3,016	4,255	-870
Liquid fuels	1,518	1,351	6,476	9,032	9,399	11,162	8,548	10,600	1,278	4,393	3,387	4,498	-631
Solid fuels	0	0	0	0	0	0	0	0	0	0	0	0	0
Gaseous fuels	-261	-294	-355	-368	-374	-385	-382	-418	-403	-374	-371	-244	-239
Total	-11,478	2,598	8,004	10,775	-11,111	-3,707	15,222	-7,401	3,877	9,890	31,410	-318	22,695
Liquid fuels	-1,493	6,917	3,060	6,663	-3,981	1,822	-7,021	-1,083	6,160	7,811	8,824	7,107	5,638
Solid fuels	-8,921	-3,015	5,727	5,086	-5,314	-4,377	23,868	-6,517	-587	4,077	24,177	-4,822	19,310
Gaseous fuels	-1,064	-1,304	-783	-975	-1,816	-1,152	-1,626	199	-1,695	-1,997	-1,591	-2,604	-2,253
(RA-SA)-(Total)	645	-6,401	-8,703	-11,718	-12,107	-9,655	-5,647	-7,653	-7,963	-6,715	-8,205	-9,306	-8,383
Liquid fuels	14,375	8,178	8,794	6,620	7,839	10,696	14,368	9,643	7,537	8,985	7,303	6,724	8,491
Solid fuels	-5,088	-4,484	-4,643	-4,443	-5,345	-5,030	-4,995	-2,718	-1,888	-1,878	-2,072	-2,606	-3,248
Gaseous fuels	475	408	289	271	412	490	545	544	556	595	542	474	554
Other fuels	-9,116	-10,503	-13,142	-14,167	-15,014	-15,812	-15,565	-15,123	-14,168	-14,417	-13,976	-13,899	-14,180

Annex 5. Assessment of Completeness and (Potential) Sources and Sinks of Greenhouse Gas Emissions and Removals Excluded

A5.1. Assessment of Completeness

Current inventory is submitted in accordance with the common reporting format (CRF), which requires entering emission data or a notation key¹ such as “NO”, “NE”, or “NA” for all sources. This chapter presents the definition of notation keys and decision trees for the application of them, both of which are based on the UNFCCC reporting Guidelines (FCCC/CP/1999/7, FCCC/CP/2002/8, FCCC/SBSTA/2004/8 or FCCC/SBSTA/2006/9) and the results of Committee for Greenhouse Gases Emissions Estimation Methods in 2002.

This chapter also reports source categories which have not been estimated because i) applicability of IPCC default values is not assured, ii) default methodologies and default values are not provided, iii) activity data is not available, iv) actual condition of GHG emissions or removals is not understood clearly.

A5.2. Definition of Notation Keys

When reviewing the appropriateness of applying notation keys shown in the UNFCCC reporting guideline, it is necessary to establish a common concept for an application of these keys for each sector, but unclear points described in Table A5-1 are found as below regarding the use of the notation key.

- The explanation of “NO” in the UNFCCC reporting guidelines can be taken that “NO” may be applied to both situations when there are no emissions or removals because the activities do not exist in Japan, and when emissions or removals do not occur in principle although the activities do exist.
- The first sentence of the “NA” explanation in the UNFCCC reporting guidelines seems to imply that “NA” may be applied to both situations as for “NO”. However, because the second sentence states that “If categories... are shaded, they do not need to be filled in”, it also seems to mean that “NA” is applied only when the activities exist but there are no emissions or removals in principle.

In the Committee for Greenhouse Gases Emissions Estimation Methods in 2002, the meanings of the notation keys are defined based on the following policy (as shown in Table A5-2).

- It was decided that “NA” is applied when the activity does exist in Japan, but in principle there are no GHG emissions or removals, while “NO” will apply when the activity itself does not exist and there are no emissions or removals.

¹ These were called "standard indicators" in FCCC/CP/1999/7, but were changed to "notation keys" in FCCC/CP/2002/8.

If the UNFCCC reporting guidelines are revised in future, the review of the definitions of notation keys and the way to fill them in CRF will be conducted.

Table A 5-1 Notation keys indicated in UNFCCC reporting guidelines

Notation Key	Explanation
NO (Not Occurring)	“NO” (not occurring) for emissions by sources and removals by sinks of greenhouse gases that do not occur for a particular gas or source/sink category within a country;
NE (Not Estimated)	“NE” (not estimated) for existing emissions by sources and removals by sinks of greenhouse gases which have not been estimated. Where “NE” is used in an inventory for emissions or removals of CO ₂ , CH ₄ , N ₂ O, HFCs, PFCs or SF ₆ , the Party should indicate why emissions could not be estimated, using the completeness table of the common reporting format;
NA (Not Applicable)	“NA” (not applicable) for activities in a given source/sink category that do not result in emissions or removals of a specific gas. If categories in the common reporting format for which “NA” is applicable are shaded, they do not need to be filled in;
IE (Included Elsewhere)	“IE” (included elsewhere) for emissions by sources and removals by sinks of greenhouse gases estimated but included elsewhere in the inventory instead of the expected source/sink category. Where “IE” is used in an inventory, the Party should indicate, using the completeness table of the common reporting format, where in the inventory the emissions or removals from the displaced source/sink category have been included and the Party should give the reasons for this inclusion deviating from the expected category;
C (Confidential)	“C” (confidential) for emissions by sources and removals by sinks of greenhouse gases which could lead to the disclosure of confidential information, given the provisions of paragraph 27 above; (para 27: Emissions and removals should be reported on the most disaggregated level of each source/sink category, taking into account that a minimum level of aggregation may be required to protect confidential business and military information.

Source : UNFCCC reporting guidelines on annual inventories (FCCC/SBSTA/2004/8)

* The notation key “0” was deleted at COP8 from the revised UNFCCC reporting guidelines (FCCC/CP/2002/8).

Table A 5-2 Definition of Notation Keys

Notation Key	Definition
NO (Not Occurring)	Used when there are no activities that are linked to emissions or removals for a certain source.
NE (Not Estimated)	Used when the emissions or removals of a certain source cannot be estimated.
NA (Not Applicable)	Used when an activity associated with a certain source does exist, but in principle it accompanies no occurrence of specific GHG emissions or removals. “NA” is not applied when there are no GHG emissions or removals because the GHGs in raw materials have been removed.
IE (Included Elsewhere)	IE is used when an emissions or removals are already included in other sources. For assuring the completeness of CRF, the sources in which the emissions or removals are included and the reasons for including it elsewhere are to be recorded in the table.
C (Confidential)	Used for confidential information relating to business or the military. However, in consideration of transparency in calculation of emissions or removals, information will be reported to the extent that it does not hinder business or other operations (for example, reporting the aggregated total of several substances).

A5.3. Decision Tree for Application of Notation Keys

Decision tree for the application of notation keys, based on UNFCCC reporting Guidelines (FCCC/CP/1999/7 FCCC/CP/2002/8, FCCC/SBSTA/2004/8 or FCCC/SBSTA/2006/9) and the results of Committee for Greenhouse Gases Emissions Estimation Methods in 2002, is shown in Figure A5-1.

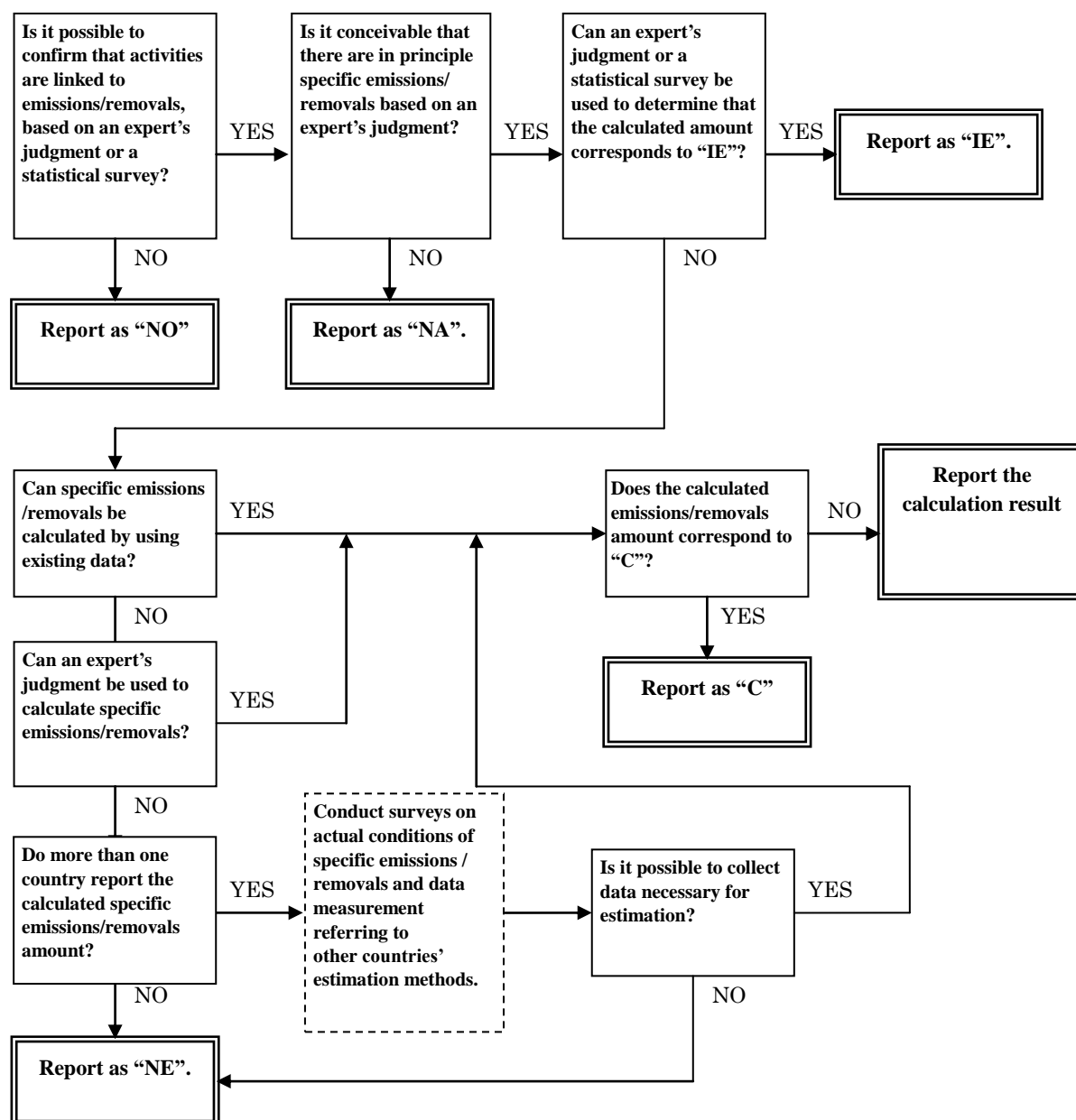


Figure A 5-1 Decision tree for application of notation keys

A5.4. Source categories not estimated in Japan's inventory

Source categories dissolved not estimate status in this year and categories still not estimated in Japan's inventory are listed below. Note that the actual emissions 1990-1994 of HFCs, PFCs and SF₆ are not estimated.

Table A 5-3 Dissolution of “NE” categories for 2010

Code	Sector	Source category				Gas
1	Land - use Change and Forestry	Settlements	Settlements remaining Settlements	Urban Green Areas subject to RV	Soil	Carbon Stock Change
2	Land - use Change and Forestry	Settlements	Land Converted to Settlements	Forest Land Converted to Settlements	Soil	Carbon Stock Change
3	Land - use Change and Forestry	Settlements	Land Converted to Settlements	Cropland Converted to Settlements	Soil	Carbon Stock Change
4	Land - use Change and Forestry	Settlements	Land Converted to Settlements	Grassland Converted to Settlements	Soil	Carbon Stock Change

Table A 5-4 “NE” categories for 2010

Code	Sector	Source category				GHG
1	Energy	Fugitive Emissions from Fuels	Solid Fuels	Coal Mining		CO ₂
2	Energy	Fugitive Emissions from Fuels	Solid Fuels	Coal Mining		N ₂ O
3	Energy	Fugitive Emissions from Fuels	Solid Fuels	Solid Fuel Transformation		CO ₂
4	Energy	Fugitive Emissions from Fuels	Solid Fuels	Solid Fuel Transformation		CH ₄
5	Energy	Fugitive Emissions from Fuels	Solid Fuels	Solid Fuel Transformation		N ₂ O
6	Energy	Fugitive Emissions from Fuels	Oil and Natural Gas	Oil	Refining/Storage	CO ₂
7	Energy	Fugitive Emissions from Fuels	Oil and Natural Gas	Oil	Distribution of Oil Products	CO ₂
8	Energy	Fugitive Emissions from Fuels	Oil and Natural Gas	Oil	Distribution of Oil Products	CH ₄
9	Industrial Processes	Mineral Products	Asphalt roofing			CO ₂
10	Industrial Processes	Mineral Products	Road Paving with Asphalt			CO ₂
11	Industrial Processes	Chemical Industry	Ammonia Production			CH ₄
12	Industrial Processes	Metal Production	Aluminium Production			CH ₄
13	Solvent and Other Product Use	Degreasing and Dry-Cleaning				CO ₂
14	Solvent and Other Product Use	Chemical Product, Manufacture and Processing				CO ₂
15	Solvent and Other Product Use	Other	Other Use of N ₂ O			N ₂ O
16	Agriculture	Enteric Fermentation	Poultry			CH ₄
17	Agriculture	Field Burning of Agricultural Residues	Other			CH ₄
18	Agriculture	Field Burning of Agricultural Residues	Other			N ₂ O
19	Land - use Change and Forestry	Cropland	Cropland remaining Cropland	Soil		Carbon Stock Change
20	Land - use Change and Forestry	Cropland	Cropland remaining Cropland	Biomass Burning	Controlled Burning	CO ₂
21	Land - use Change and Forestry	Cropland	Cropland remaining Cropland	Biomass Burning	Controlled Burning	CH ₄
22	Land - use Change and Forestry	Cropland	Cropland remaining Cropland	Biomass Burning	Controlled Burning	N ₂ O
23	Land - use Change and Forestry	Cropland	Land Converted to Cropland	Forest Land converted to Cropland	Soil (Organic soils)	Carbon Stock Change
24	Land - use Change and Forestry	Cropland	Land Converted to Cropland	Grassland converted to Cropland	Soil (Organic soils)	Carbon Stock Change
25	Land - use Change and Forestry	Cropland	Land Converted to Cropland	Wetland converted to Cropland	Soil (Organic soils)	Carbon Stock Change
26	Land - use Change and Forestry	Cropland	Land Converted to Cropland	Other Land converted to Cropland	Dead Organic Matter	Carbon Stock Change
27	Land - use Change and Forestry	Cropland	Land Converted to Cropland	Other Land converted to Cropland	Soil	Carbon Stock Change
28	Land - use Change and Forestry	Cropland	Land Converted to Cropland	N ₂ O emissions from disturbance	Soil	N ₂ O
29	Land - use Change and Forestry	Grassland	Grassland remaining Grassland	Wild land	Living Biomass	Carbon Stock Change
30	Land - use Change and Forestry	Grassland	Grassland remaining Grassland	Wild land	Dead Organic Matter	Carbon Stock Change
31	Land - use Change and Forestry	Grassland	Grassland remaining Grassland	Wild land	Soil	Carbon Stock Change
32	Land - use Change and Forestry	Grassland	Grassland remaining Grassland	Grazed meadow	Soil	Carbon Stock Change
33	Land - use Change and Forestry	Grassland	Grassland remaining Grassland	Pasture land	Soil	Carbon Stock Change
34	Land - use Change and Forestry	Grassland	Grassland remaining Grassland	Biomass Burning	Wildfires	CO ₂
35	Land - use Change and Forestry	Grassland	Grassland remaining Grassland	Biomass Burning	Wildfires	CH ₄
36	Land - use Change and Forestry	Grassland	Grassland remaining Grassland	Biomass Burning	Wildfires	N ₂ O
37	Land - use Change and Forestry	Grassland	Grassland remaining Grassland	Biomass Burning	Controlled Burning	CO ₂
38	Land - use Change and Forestry	Grassland	Grassland remaining Grassland	Biomass Burning	Controlled Burning	CH ₄
39	Land - use Change and Forestry	Grassland	Grassland remaining Grassland	Biomass Burning	Controlled Burning	N ₂ O
40	Land - use Change and Forestry	Grassland	Land Converted to Grassland	Forest Land converted to Grassland	Soil (Organic soils)	Carbon Stock Change
41	Land - use Change and Forestry	Grassland	Land Converted to Grassland	Cropland converted to Grassland	Soil (Organic soils)	Carbon Stock Change
42	Land - use Change and Forestry	Grassland	Land Converted to Grassland	Wetland converted to Grassland	Soil	Carbon Stock Change
43	Land - use Change and Forestry	Grassland	Land Converted to Grassland	Other Land converted to Grassland	Dead Organic Matter	Carbon Stock Change
44	Land - use Change and Forestry	Grassland	Land Converted to Grassland	Other Land converted to Grassland	Soil	Carbon Stock Change
45	Land - use Change and Forestry	Grassland	Land Converted to Grassland	Biomass Burning	Wildfires	CO ₂
46	Land - use Change and Forestry	Grassland	Land Converted to Grassland	Biomass Burning	Wildfires	CH ₄
47	Land - use Change and Forestry	Grassland	Land Converted to Grassland	Biomass Burning	Wildfires	N ₂ O
48	Land - use Change and Forestry	Wetlands	Wetlands remaining Wetlands	Flooded land	Living Biomass	Carbon Stock Change
49	Land - use Change and Forestry	Wetlands	Wetlands remaining Wetlands	Flooded land	Dead Organic Matter	Carbon Stock Change
50	Land - use Change and Forestry	Wetlands	Wetlands remaining Wetlands	Flooded land	Soil	Carbon Stock Change
51	Land - use Change and Forestry	Wetlands	Wetlands remaining Wetlands	Biomass Burning	Wildfires	CO ₂
52	Land - use Change and Forestry	Wetlands	Wetlands remaining Wetlands	Biomass Burning	Wildfires	CH ₄
53	Land - use Change and Forestry	Wetlands	Wetlands remaining Wetlands	Biomass Burning	Wildfires	N ₂ O
54	Land - use Change and Forestry	Wetlands	Wetlands remaining Wetlands	Biomass Burning	Controlled Burning	CO ₂
55	Land - use Change and Forestry	Wetlands	Wetlands remaining Wetlands	Biomass Burning	Controlled Burning	CH ₄
56	Land - use Change and Forestry	Wetlands	Wetlands remaining Wetlands	Biomass Burning	Controlled Burning	N ₂ O

Table A 5-4 “NE” categories for 2010 (cont.)

Code	Sector	Source category				GHG
57	Land - use Change and Forestry	Wetlands	Land converted to Wetlands	Forest Land converted to Wetlands	Soil	Carbon Stock Change
58	Land - use Change and Forestry	Wetlands	Land converted to Wetlands	Cropland converted to Wetlands	Soil	Carbon Stock Change
59	Land - use Change and Forestry	Wetlands	Land converted to Wetlands	Grassland converted to Wetlands	Soil	Carbon Stock Change
60	Land - use Change and Forestry	Wetlands	Land converted to Wetlands	Settlements converted to Wetlands	Soil	Carbon Stock Change
61	Land - use Change and Forestry	Wetlands	Land converted to Wetlands	Other Land converted to Wetlands	Dead Organic Matter	Carbon Stock Change
62	Land - use Change and Forestry	Wetlands	Land converted to Wetlands	Other Land converted to Wetlands	Soil	Carbon Stock Change
63	Land - use Change and Forestry	Wetlands	Land converted to Wetlands	Biomass Burning	Wildfires	CO ₂
64	Land - use Change and Forestry	Wetlands	Land converted to Wetlands	Biomass Burning	Wildfires	CH ₄
65	Land - use Change and Forestry	Wetlands	Land converted to Wetlands	Biomass Burning	Wildfires	N ₂ O
66	Land - use Change and Forestry	Settlements	Settlements remaining Settlements			CH ₄
67	Land - use Change and Forestry	Settlements	Settlements remaining Settlements			N ₂ O
68	Land - use Change and Forestry	Settlements	Settlements remaining Settlements	Other than Urban Green Areas	Living Biomass	Carbon Stock Change
69	Land - use Change and Forestry	Settlements	Settlements remaining Settlements	Other than Urban Green Areas	Dead Organic Matter	Carbon Stock Change
70	Land - use Change and Forestry	Settlements	Settlements remaining Settlements	Other than Urban Green Areas	Soil	Carbon Stock Change
71	Land - use Change and Forestry	Settlements	Settlements remaining Settlements	Urban Green Areas not subject to RV	Dead Organic Matter	Carbon Stock Change
72	Land - use Change and Forestry	Settlements	Settlements remaining Settlements	Urban Green Areas not subject to RV	Soil	Carbon Stock Change
73	Land - use Change and Forestry	Other land	Land Converted to Other land	Forest Land Converted to Other land	Soil	Carbon Stock Change
74	Land - use Change and Forestry	Other land	Land Converted to Other land	Cropland Converted to Other land	Soil	Carbon Stock Change
75	Land - use Change and Forestry	Other land	Land Converted to Other land	Grassland Converted to Other land	Soil	Carbon Stock Change
76	Land - use Change and Forestry	Harvested Wood Product				CO ₂
77	Land - use Change and Forestry	Harvested Wood Product				CH ₄
78	Land - use Change and Forestry	Harvested Wood Product				N ₂ O

Annex 6. Additional Information to be Considered as Part of the NIR

Submission or Other Useful Reference Information

A6.1. Details on Inventory Compilation System and QA/QC Plan

The main parts of the QA/QC Plan for Japan's greenhouse gas inventory are excerpted.

A6.1.1. Introduction to QA/QC Plan

The QA/QC Plan is an internal document that documents, among other things, the specifics of all QA/QC activities in all processes from the start of National Inventory Report compilation to the final report, the compilation schedule, and the apportionment of all involved entities' roles. It organizes and systematizes the QA/QC activities of inventory compilation and clarifies what each entity involved in compilation is supposed to do. Additionally, it is prepared for the purpose of guaranteeing the implementation of QA/QC activities.

A6.1.2. QA/QC plan's scope

The QA/QC Plan's scope includes the processes of preparing, reporting, and reviewing the inventory under the Framework Convention on Climate Change, and the supplementary information on sinks under Kyoto Protocol Articles 3.3 and 3.4, as stipulated in Article 7.1 of the Protocol.

A6.1.3. Roles and responsibilities of each entity involved in the inventory preparation process

Following are the agencies involved in the inventory compilation process, and the roles of those agencies.

1) Ministry of the Environment (Low-carbon Society Promotion Office, Global Environment Bureau)

- The single national agency responsible for preparing Japan's inventory, which was designated pursuant to the Kyoto Protocol Article 5.1.
- It is responsible for editing and submitting the inventory.

2) Greenhouse Gas Inventory Office of Japan (GIO), Center for Global Environmental Research, National Institute for Environmental Studies

- Performs the actual work of inventory compilation. Responsible for inventory calculations, editing, and the archiving and management of all data.

3) Relevant Ministries/Agencies

The relevant ministries and agencies have the following roles and responsibilities regarding inventory compilation.

- Preparation of activity data, emission factor data, and other data needed for inventory compilation, and submission of the data by the submission deadline.
- Quality control (QC) of the data provided to the Ministry of the Environment and the GIO.
- Confirmation and verification of the inventory (CRF, NIR, spreadsheets, and other information) prepared by the Ministry of the Environment and the GIO.
- (When necessary), responding to questions from expert review teams about the statistics

controlled by relevant ministries and agencies, or about certain data they have prepared, and preparing comments on draft reviews.

- (When necessary), responding to visits by expert review teams.

4) *Relevant Organizations*

Relevant organizations have the following roles and responsibilities regarding inventory compilation.

- Preparation of activity data, emission factor data, and other data needed for inventory compilation, and submission of the data by the submission deadline.
- (When necessary), responding to questions from expert review teams about the statistics controlled by relevant organizations, or about certain data they have prepared, and preparing comments on draft reviews.

5) *Committee for the Greenhouse Gas Emissions Estimation Methods*

The Committee for the Greenhouse Gas Emissions Estimation Methods (Committee) is a committee created and run by the Ministry of the Environment. Its role is to consider the methods for calculating inventory emissions and removals, and consider the selection of parameters such as activity data and emission factors. Under the Committee is the inventory working group (WG) that examines crosscutting issues, and breakout groups that consider sector-specific problems (Breakout group on Energy and Industrial Processes, Breakout group on Transport, Breakout group on F-gas [HFCs, PFCs, and SF₆], Breakout group on Agriculture, Breakout group on Waste, and Breakout group on LULUCF). The inventory WG and breakout groups comprise experts in various fields, and consider suggestions for inventory improvements. Improvement suggestions are considered once more by the Committee before approval.

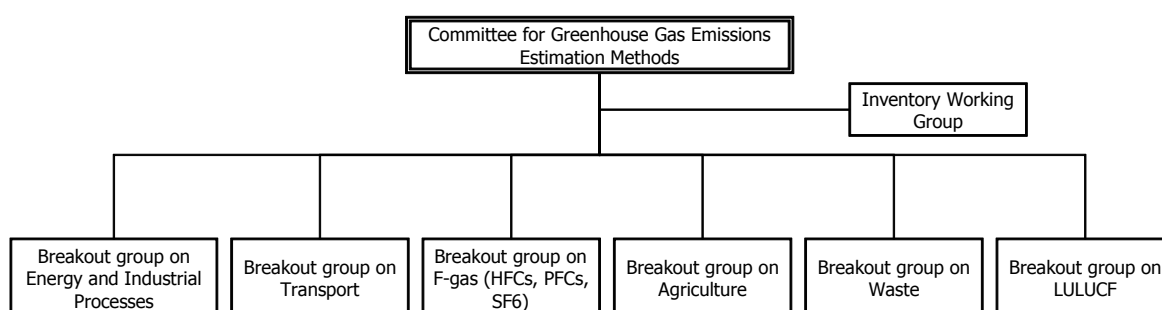


Figure A 6-1 Structure of the Committee for the Greenhouse Gas Emissions Estimation Methods

6) *Private Consulting Companies*

Private consultant companies that are contracted by the Ministry of the Environment to perform tasks related to inventory compilation play the following roles in inventory compilation based on their contracts.

- Quality control (QC) of inventory (CRF, NIR, spreadsheets, and other information) compiled by the Ministry of the Environment and the GIO.
- (When necessary), providing support for responding to questions from expert review teams and for preparing comments on draft reviews.
- (When necessary), providing support for responding to visits by expert review teams.

7) GHG Inventory Quality Assurance Working Group (Expert Peer Review) (QAWG)

The GHG Inventory Quality Assurance Working Group (the QAWG) is an organization that is for QA activities, and comprises experts who are not directly involved in inventory compilation. Its role is to assure inventory quality and to identify places that need improvement by conducting detailed reviews of each emission source and sink in the inventory.

A6.1.4. Collection process of activity data

When the activity data needed for calculations are available from sources such as publications and the internet, the necessary data are gathered from these media. Data that are not released in publications, the internet, or in other media, and unpublished data that are used when compiling the inventory are obtained by the Ministry of the Environment or the GIO by requesting them from the relevant ministries and agencies and the relevant organizations which control those data. Since FY2009, the Network of Official Greenhouse Gas Inventory System for Total Management (NOGISTOM) has been used for data collection. This system manages all the information related to provision of data. The main relevant ministries and agencies and relevant organizations that provide data are as shown in Table A 6-1.

Table A 6-1 List of the main relevant ministries and agencies and the relevant organizations (data providers)

Ministries/Agencies/Organizations		Major data or statistics
Relevant Ministries/ Agencies	Ministry of the Environment	Research of Air Pollutant Emissions from Stationary Sources / volume of waste in landfill / volume of incinerated waste / number of people per <i>johkasou</i> facility / volume of human waste treated at human waste treatment facilities
	Ministry of Economy, Trade and Industry	General Energy Statistics / Yearbook of Production, Supply and Demand of Petroleum, Coal and Coke / Yearbook of Iron and Steel, Non-ferrous Metals, and Fabricated Metals Statistics / Yearbook of Chemical Industry Statistics / Yearbook of Ceramics and Building Materials Statistics / Census of Manufactures / General outlook on electric power supply and demand
	Ministry of Land, Infrastructure, Transport and Tourism	Annual of Land Transport Statistics / Survey on Transport Energy / Statistical Yearbook of Motor Vehicle Transport / Survey on Current State of Land Use, Survey on Current State of Urban Park Development / Sewage Statistics
	Ministry of Agriculture, Forestry and Fisheries	Crop Statistics / Livestock Statistics / Vegetable Production and Shipment Statistics / World Census of Agriculture and Forestry / Statistics of Arable and Planted Land Area / Handbook of Forest and Forestry Statistics / Table of Food Supply and Demand
	Ministry of Health, Labour and Welfare	Statistics of Production by Pharmaceutical Industry
Relevant Organizations	Federation of Electric Power Companies	Amount of Fuel Used by Pressurized Fluidized Bed Boilers
	Japan Coal Energy Center	Coal Production
	Japan Cement Association	Amount of clinker production / Amount of waste input to in raw material processing / Amount of RPF incineration
	Japan Iron and Steel Federation	Emissions from Coke Oven Covers, Desulfurization Towers, and Desulfurization Recycling Towers
	Japan Paper Association	Amount of final disposal of industrial waste / Amount of RPF incineration

A6.1.5. Selection process of emission factors and estimation methods

Calculation methods for Japan's emission and removal amounts are determined by having the Committee explore calculation methods suited to Japan's situation for all the activity categories necessary for calculating Japan's greenhouse gas emission and removal amounts, based on the 1996 Revised IPCC Guidelines, GPG (2000), GPG-LULUCF, and the 2006 IPCC Guidelines.

A6.1.6. Improvement process of estimations for emissions and removals

In Japan, improvements in calculation methods are considered in accordance with necessity whenever an inventory item requiring improvement is identified because of, for example, a UNFCCC review or an observation by the QAWG, progress in international negotiations such as the creation of new guidelines, progress or changes in scientific research or in the compilation of statistics, or the acquisition of new information by the system for calculating, reporting, and publishing GHG emissions. Proposals for improving the estimation of emissions and removals are considered by scientific research or the Committee, and the results are incorporated into the inventory. Figure A 6-2 below is a diagram of the inventory improvement process.

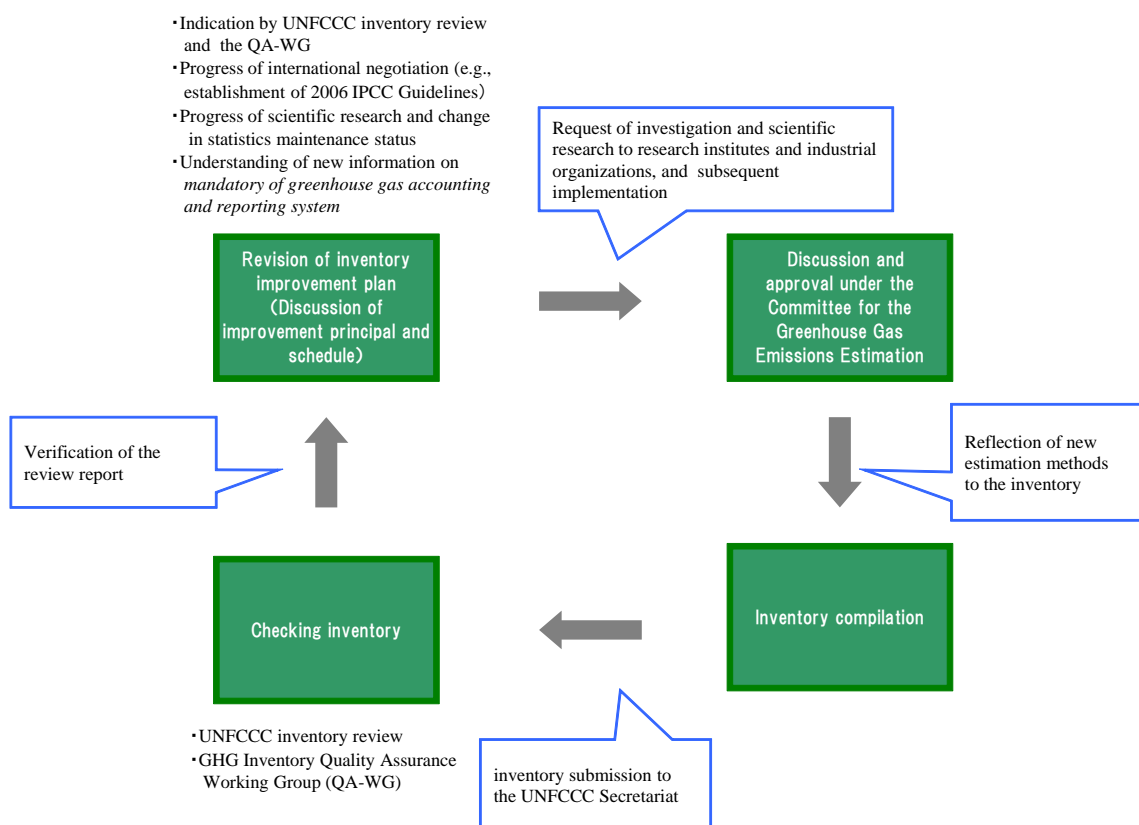


Figure A 6-2 Diagram of the inventory improvement process

A6.1.7. QA/QC activity

When compiling the inventory in Japan, inventory quality is controlled by performing quality control (QC) activities (such as checking the correctness of calculations and archive of documents) at each step in accordance with GPG (2000) and GPG-LULUCF. In Japan, the quality control activities relating to inventory compilation performed by personnel belonging to agencies involved in inventory compilation—that is, the Ministry of the Environment (including the GIO and private consultant companies), relevant ministries and agencies—are considered to be QC. External reviews by experts who are outside the inventory compilation system (QAWG) are considered to be QA (quality assurance). They verify and assess data quality from the perspectives of scientific knowledge and data availability with respect to current calculation methods. Table A 6-2 sketches Japan's QA/QC activities.

Table A 6-2 Summary of Japan's QA/QC activity

	Implementing entity	Main contents of activity
QC (Quality Control)	Ministry of the Environment (Low-carbon Society Promotion Office, Global Environment Bureau)	<ul style="list-style-type: none"> • Progress management of the inventory compilation and overall control • Check of inventory compiled by the GIO (CRF, NIR, spreadsheets, and other information) • Establishment and revision of QA/QC plan • Check of the inventory improvement plan • Holding the meeting of the Committee for the Greenhouse Gas Emissions Estimation Methods
	Greenhouse Gas Inventory Office of Japan, Center for Global Environmental Research, National Institute for Environmental Studies (GIO)	<ul style="list-style-type: none"> • QC check in inventory compilation • Archiving of QA/QC activity records and relevant data and documents • Development of information system • Making of inventory improvement plan • Making of revised QA/QC plan
	Relevant Ministry and Agencies (including the Ministry of the Environment)	<ul style="list-style-type: none"> • Preparation of activity data, emission factor, and other data needed for inventory compilation, and submission of the data by the submission deadline. • Check of various data supplying to the GIO • Check and validation of inventory compiled by the GIO (CRF, NIR, spreadsheets, and other information)
	Committee for the Greenhouse Gas Emissions Estimation Methods	<ul style="list-style-type: none"> • Discussion and Assessment for estimation methods, emission factors, and activity data
	Private Consultant Companies	<ul style="list-style-type: none"> • Check of inventory compiled by the GIO (CRF, NIR, spreadsheets, and other information)
QA (Quality Assurance)	Inventory Quality Assurance Working Group (QAWG)	<ul style="list-style-type: none"> • Expert peer review to validate estimation methods, emission factors, and activity data • Inventory assessment

A6.1.7.1. QC activity

A6.1.7.1.a. General QC procedures (Tier 1)

General QC procedures include the general items to be confirmed which are related to the calculation, data processing, completeness, and documentation applicable to all emission source and sink categories. General QC procedures are implemented by each inventory compiler.

Following are the QC activities conducted by the sectoral experts (SEs), who perform the work of compiling the emissions/removals estimation files for each category, the CRF master files and NIR; the National Inventory compiler (NIC), who integrates the information from the individual SEs and compiles the inventory; and the data providers, who provide the activity data and other data used to calculate emissions and removals.

This section describes the QC activities of the GIO and private consultant companies in parts 1) and 2).

1) Sectoral expert (SE)

SEs perform mainly the following QC activities.

- Checking for transcription errors in data entry and referencing
- Checking to ensure that emissions are accurately estimated
- Checking to see that parameters and emission units are accurately recorded, and that proper conversion factors are used
- Checking the conformity of databases and/or files
- Checking the consistency of data from one category to another

- Checking the accuracy of inventory data behavior from one processing step to the next
- Checking completeness
- Checking time series consistency
- Checking trends
- Conducting comparisons with past estimated values
- Checking that uncertainties in emissions and removals are accurately estimated and calculated
- Carrying out reviews of internal documentation
- Checking that the assumptions and criteria for selecting activity data and emission factors are documented

2) *National inventory compiler (NIC)*

The NIC performs mainly the following QC activities when preparing CRF files.

- Confirming that CRF Reporter data provided by SEs are imported without omission
- Confirming that the information needed for the documentation box is properly entered
- Confirming that the reasons for “NE” and “IE” are correctly entered
- Confirming that the key category analysis results are correctly entered
- Confirming that the reasons for recalculations are provided for all categories
- Confirming that data are corrected after the coordination with the relevant ministries and agencies

A6.1.7.1.b. QC procedure for each category (Tier 2)

As part of the QC activities in Japan, private consultant companies perform external QC on the estimation files prepared by the GIO, and on the CRF and NIR drafts. In addition to confirming the data entered into estimation files for each emission source category and the equations for calculating emissions, private consultant companies use estimation files like those of the GIO to calculate total greenhouse gas emissions, and carry out mutual verification of emission estimation results. They also send to the relevant ministries and agencies the sets of files for estimation files, CRF, NIR, and the drafts of published documents for domestic release showing estimated values for emissions and removals. And they confirm and verify the content of categories relevant to each ministry or agency (coordination with the relevant ministries and agencies).

Since the Committee considers and selects the methodologies, activity data and parameters including emission factors, which are actually applied to the estimation of emissions/removals from each category, it also implements Tier 2 QC activities.

A6.1.7.2. QA activity

Quality assurance (QA) refers to assessment of inventory quality by third units that are not directly involved in inventory compilation. In Japan the following QA is conducted to assure inventory quality.

- GHG Inventory Quality Assurance Working Group (Expert Peer Review)

A6.1.7.2.a. GHG Inventory Quality Assurance Working Group (Expert Peer Review) (QAWG)

1) Summary

The QAWG performs detailed reviews (expert peer reviews) by experts not directly involved in inventory compilation for each emission source and sink in order to assure inventory quality and to

identify places that need improvement.

2) *Scope of review*

The GHG Inventory Quality Assurance Working Group performs reviews mainly in the following areas.

- Confirming the soundness of estimation methods, activity data, emission factors, and other items.
- Confirming the soundness of content reported in the CRF and NIR.

3) *QAWG in FY2011*

The QAWG was newly established in FY2009 as a result of discussions within the Committee held in FY2008 in order to enhance Japan's QA/QC activities. The QAWG fulfils QA activities for inventory preparation, reporting and reviewing as required for the Annex I Parties under the FCCC as well as the Kyoto Protocol by implementing a detailed review by experts, who are not directly involved in or related to the inventory preparation process, for each source and/or sink. The secretariat for the QAWG was established within the GIO. The secretariat and the Ministry of the Environment determined the sectors and categories to be reviewed by the QAWG. The experts for the QAWG were selected by taking the following requirements into account.

<Requirements for QAWG review expert>

- | |
|---|
| <p>a. No direct involvement in the inventory preparation process for estimating emissions/removals from the sectors/categories to be reviewed (i.e., no involvement in the Committee, the data creation and the data provision for those sectors/categories)</p> <p>b. No specific interests related to the inventory and the capability to judge objectively without being affected by any specific organizations and/or stakeholders.</p> <p>c. Sufficient skills, knowledge and experiences to assure the quality of the inventory</p> |
|---|

The Energy sector was reviewed by two experts in FY2011, and the schedule was as follows.

Table A 6-3 Schedule for the QAWG in FY2011

Schedule	Matter
Until March 2011	Selection of experts by the Ministry of Environment of Japan and the secretariat
May	Visit and briefing of the experts
May to July	Review by the experts (The detailed review of the Inventory, the listing of dubious and controversial points, and proposal for improvements)
30 August	Holding of the QAWG meeting
September to January 2012	Bringing up and discussion of suggestions from the QAWG to each breakout group in the Committee

Key data and the methods of estimation used in this sector have been validated by QAWG. The QAWG identified some issues and submitted them to the Committee. Other issues that have not been resolved by the committee are presented in each category of the "f) Source-specific Planned Improvement" section in this report. In addition, the QAWG identified insufficient explanations and incorrect descriptions in the NIR 2011 and addressed them in this report to improve transparency and accuracy.

The MOE and the secretariat will annually determine the sectors/categories to be reviewed by the QAWG, with the aim of reviewing the entire inventory within the next few years.

A6.1.8. Response for UNFCCC inventory review

The convention inventory and Kyoto Protocol supplementary information on sinks that Japan submits each year are to be reviewed by an expert review team (ERT) pursuant to UNFCCC inventory review guidelines¹, Kyoto Protocol Article 8, Decision 22/CMP.1, and other requirements. Specifically, rigorous checks are performed in accordance with Japan's prescribed estimation method guidelines² from perspectives including: Are emissions and removals accurately and completely estimated and reported? Are transparent explanations provided for estimation methods? Are QA/QC activities and uncertainty assessments performed appropriately?

Because the inventory review has great significance for attaining Japan's emission reduction targets under the Kyoto Protocol, it is necessary to address this matter after having made careful preparations. The system shown in Figure A 6-3 is used for responding to reviews.

The Ministry of the Environment, which in Japan is responsible for editing and submitting the inventory, is assigned to be the agency with overall control (responsibility) for review response, while the GIO performs the actual work, such as preparing source materials. Communication with the UNFCCC Secretariat is performed by the Ministry of Foreign Affairs. The relevant ministries and agencies, relevant organizations, and private consultant companies³ that are involved in inventory compilation cooperate with review response through activities including providing relevant information, support for source material preparation, and QC implementation.

¹ FCCC/CP/2002/8

² 1996 Revised IPCC Guidelines, Good Practice Guidance (2000), GPG-LULUCF

³ Private consultant companies cooperate in correspondence of the reviews based on the operating agreement with the Ministry of the Environment.

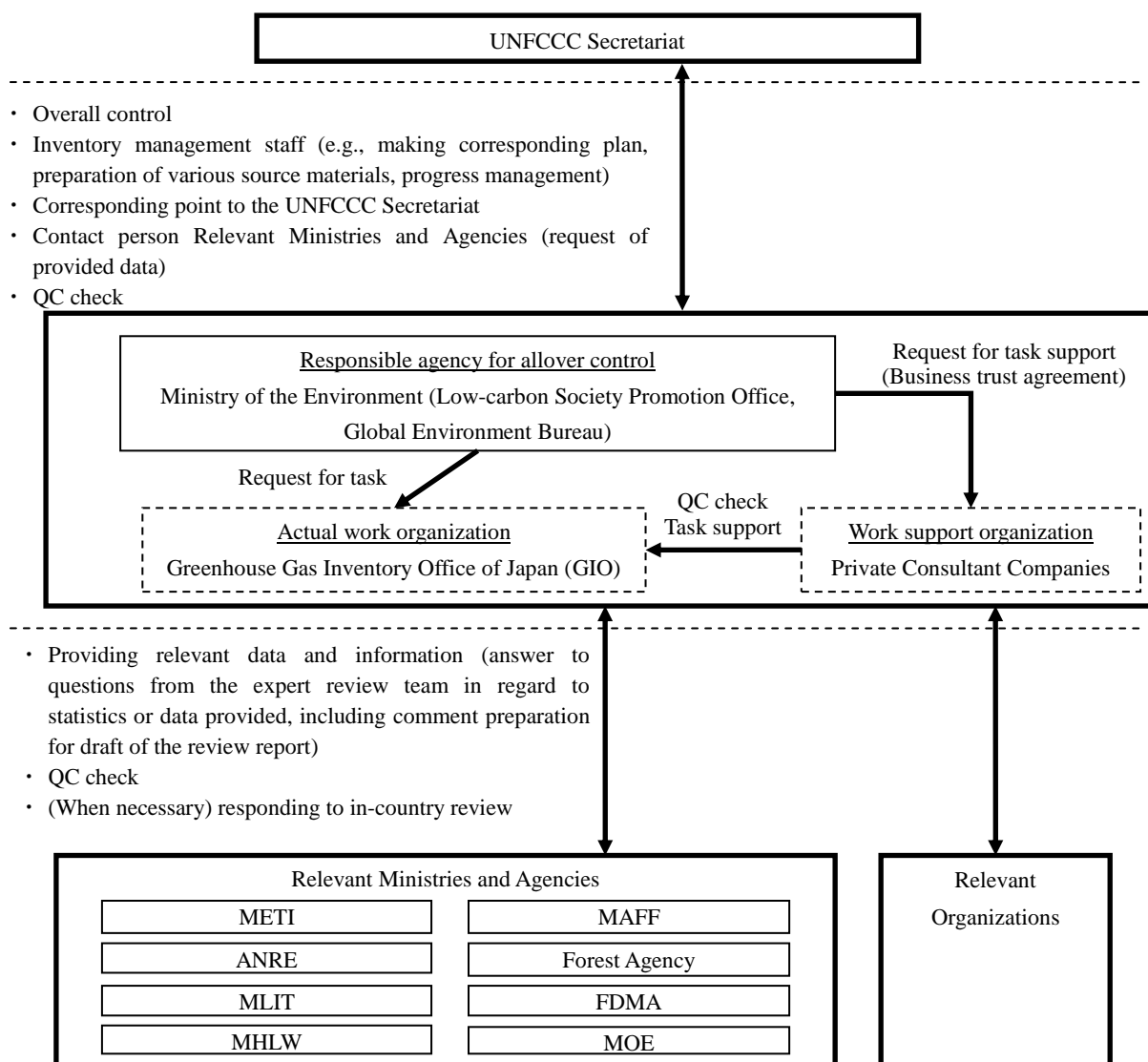


Figure A 6-3 Basic structure of Japan's national system corresponding to inventory review

A6.1.9. Documentation and archiving of inventory information

In Japan, the information needed for inventory compilation is documented and as a rule archived by the agency which compiles the inventory (GIO).

A6.1.9.1. Documentation of information

The GIO documents all the inventory-related information in electronic or printed form and archives it. Examples of information that must be archived follow.

- The inventories submitted every year to the UNFCCC Secretariat, and the related files
- Published materials for preliminary and finalized data
- Statistical data and provided data (including data providers, time period when provided, and other related information) used in compiling the inventory
- Information on the discussion process and discussion results related to the selection of activity data, estimation methods, emission factors, and other items (relevant source materials for the discussion process by the Committee for the Greenhouse Gas Emissions Estimation Methods)
- Records of communications with related entities in the inventory compilation process

- Information on inventory recalculations (such as reasons for recalculations, and when performed)
- Record of QA/QC activities conducted
- Comments by experts on the inventory
- In relation to UNFCCC inventory reviews, review reports and records of questions and answers with expert review teams
- Internal documents on inventory compilation, including the QA/QC Plan

A6.1.9.2. Archiving of information

1) Archiving electronic information

i) Inventory-related electronic information

- Each year's emissions/removals estimation files and CRF- and NIR-related files have file names with the year the estimation is for and the year it was performed, and files are saved in folders prescribed for each year.
- Electronic files of statistical data, provided data, etc. used to prepare the inventory's emissions/removals estimates and other, related data are given file names with the date on which the data were obtained and the data provider, and saved in prescribed folders. Furthermore, since data collection from relevant ministries and agencies and the relevant organizations is done via the NOGISTOM, the received data are also stored within this database.
- Source materials in electronic form (files in Word, PDF, or other format) used when considering emissions/removals estimation methods are labeled with the source material title and the date the file was obtained (and if necessary the file provider), and saved in prescribed folders.
- If the exchange of information on the inventory has been conducted by email, the email files are saved in prescribed folders.

ii) Backup and risk management of electronic information

- The CGER server, where inventory-related information is stored, is automatically backed up to two other locations every day.
- Once a year, after submission of the annual inventory to the UNFCCC Secretariat, all inventory-related electronic information is saved to CD-ROMs and other electronic media and archived.

2) Archiving printed form

- Books of statistics, data and source materials (including faxes) in printed form that have been provided, and other source materials in printed form that have been used in inventory emissions/removals estimates are filed in a prescribed storage location.

A6.1.9.3. QC activity for documentation and archiving of inventory information

Immediately after the inventory is submitted to the UNFCCC Secretariat, the GIO carries out QC activities related to the documentation and archive of inventory information.

Annex 7. Methodology and Results of Uncertainty Assessment

A7.1. Methodology of Uncertainty Assessment

A7.1.1. Background and Purpose

Under the United Nations Framework Convention on Climate Change (UNFCCC), Annex I Parties are required to submit their inventories on greenhouse gases emissions and removals (hereafter, ‘inventory’) to the UNFCCC secretariat. *Good Practice Guidance (2000)*, adopted in May 2000, further requires parties to quantitatively assess and report the uncertainty of their inventories. It should be noted that uncertainty assessment is intended to contribute to continuous improvement in the accuracy of inventories and that a high or low uncertainty assessed will not affect the justice of an inventory nor result in the comparison of accuracy among parties’ inventories.

Japan considered uncertainty of its inventory in the Committee for the Greenhouse Gases Emissions Estimation Methods in FY 2001 and again in FY 2006. Japan has annually conducted uncertainty assessment based on the Committee’s results since then.

This chapter will be used as a guideline for conducting the uncertainty assessment of Japan’s inventories. It may be subjected to be adjusted as appropriate.

A7.1.2. Overview of Uncertainty Assessment Indicated in the Good Practice Guidance

A7.1.2.1. About Uncertainty Assessment

A7.1.2.1.a. What is uncertainty?

- The term “uncertainty” refers to the degree of discrepancy in various data in comparison with a true value, stemming from number of characteristics with lack of sureness including representational reliability of measurements, and it is a concept that is much broader than that of accuracy.
- The uncertainty of emissions from a particular source is obtained by calculating and applying the uncertainty associated with the source’s emission factor, and the uncertainty of activity data.
- The *Good Practice Guidance* requires uncertainty of emissions from a source to be calculated using the method given below.

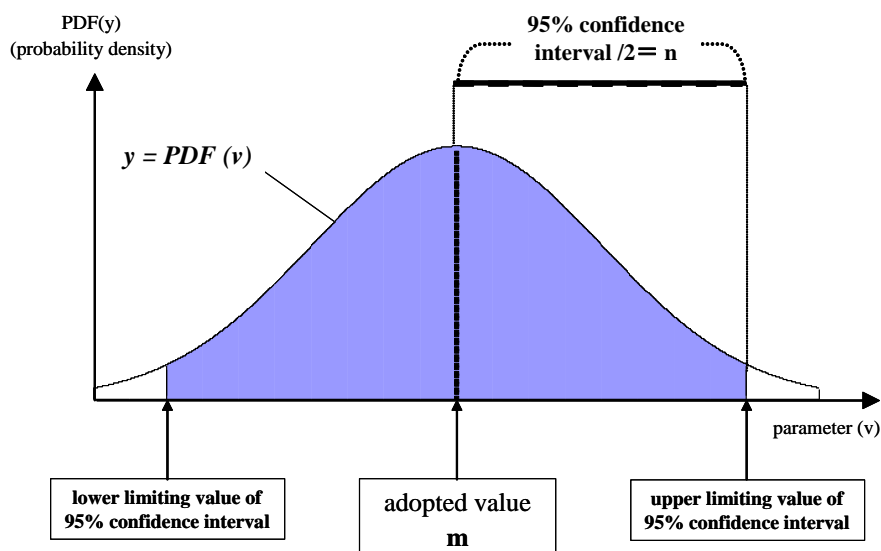
$$U = \sqrt{U_{EF}^2 + U_A^2}$$

U : Uncertainty of the emissions of the source (%)
 U_{EF} : Uncertainty of the emission factor (%)
 U_A : Uncertainty of the activity data (%)

A7.1.2.1.b. Methodology of identifying the uncertainties of emission factors and activity data of each source

- The standard deviations of the observed values of an emission factor are used to set the probability density function, and uncertainty is assessed by seeking a 95 percent confidence interval.

$$\text{Uncertainty of EF or A} = \frac{95\% \text{ confidential interval} / 2 (n)}{|\text{Adopted Value of EF or A} (m)|}$$



A7.1.2.1.c. Method of determining the uncertainty of total national emissions

- By combining the uncertainties of emissions from all sources, it is possible to assess the uncertainty of Japan’s total inventory.
- When there is no correlation between multiple uncertainties, and they are normally distributed, the *Good Practice Guidance* suggests two rules of expedience that relate to combining method (addition and multiplication) of uncertainties. This report adopts Rule A, given in Table 6.1 of the *Good Practice Guidance*, for the calculations.

$$U_{Total} = \frac{\sqrt{(U_1 \times E_1)^2 + (U_2 \times E_2)^2 + \dots + (U_n \times E_n)^2}}{E_1 + E_2 + \dots + E_n}$$

U_{Total} : Uncertainties of National Total Emissions (%)
 U_i : Uncertainties of the Emissions from Source “ i ” (%)
 E_i : the Emissions from Source “ i ” (Gg)

A7.1.2.2. Targets of the Uncertainty Assessment

The *Good Practice Guidance* suggests that all uncertainties be taken into account when estimating emissions. It indicates that the following may be the reasons of uncertainty in emission factors or activity data.

Examples of common reasons of uncertainty in emission factors	
➤	Uncertainties associated with a continuous monitoring of emissions - Refers to uncertainties arising from differences in conditions at the time of measurement, such as measurements that are taken annually.
➤	Uncertainties associated with an establishment of emission factors - Startup and shutdown in operation of machinery, etc., can give different emission rates relative to activity data. In these cases, the data should be partitioned, with separate emission factors and probability density functions derived for steady-state, startup and shutdown conditions. - Emission factors may depend on load of operation. In these cases, the estimation of total

emissions and the uncertainty analysis may need to be stratified to take account of load, which is expressed, for example, as a percentage of full capacity. This could be done by the regression analysis and scatter plots of the emission rate against seemingly influential variables (e.g., emissions versus load) with load becoming a part of the required activity data.

- Adoption of results from measurements taken for other purposes may not be representative. For example, methane measurements made for safety reasons at coalmines and landfills may not reflect total emissions. In such cases, the ratio between the measured data and total emissions should be estimated for the uncertainty analysis.

- Uncertainties associated with an estimation of emission factors from limited measured data
 - The distribution of emission factors may often differ from the normal distribution. When the distribution is already known, it is appropriate to estimate according to expert judgment, by appending a document that provides the theoretical background.

Examples of common reasons of uncertainty in activity data

- Interpretation of statistical differences: Statistical differences in energy balances usually represent a difference between amounts of primary fuels and amounts of fuels identified in the categories under 'final consumption' and 'in transformation'. They can give an indication of sizes of the uncertainties of the data, especially where long time series are considered.
- Interpretation of energy balances: Production, use, and import/export data should be consistent. If not, this may give an indication of the uncertainties.
- Crosschecks: It may be possible to compare two types of activity data that apply to the same source to provide an indication of uncertainty ranges. For example, the sum of vehicle fuel consumption should be commensurate with the total of fuel consumption calculated by multiplying vehicle-km by fuel consumption efficiency for all types of vehicles.
- Vehicle numbers and types: Some countries maintain detailed vehicle registration databases with data on vehicles by type, age, fuel type, and emission control technology, all of which can be important for a detailed bottom-up inventory of methane (CH₄) and nitrous oxide (N₂O) emissions from such vehicles. Others do not have such detailed information and this will tend to increase the uncertainty.
- Smuggling of fuel across borders: Imported fuel and the sum of sectoral fuel consumption may be compared as a crosscheck.
- Biomass fuels: Where formal markets for these fuels do not exist, consumption estimates may be much less accurate than for fuels in general.
- Livestock population data: Accuracy will depend on the extent and reliability of national census and survey methods, and there may be different accounting conventions for animals that do not live for a whole year.

A7.1.2.3. Methodology of Uncertainty Assessment

The *Good Practice Guidance* suggests that uncertainty is assessed through expert judgment and actual data with consideration to the sources of uncertainty indicated in section above.

A7.1.3. Methodology of Uncertainty Assessment in Japan's Inventories

A7.1.3.1. Principle of Uncertainty Assessment

The following method of uncertainty assessment is used, with regard for both convenience of the compilation and suggestions made in the *Good Practice Guidance*, in a manner that as far as possible ensures there is no deviation from assessment standards among categories.

A7.1.3.2. Separation between Emission Factors and Activity Data

The equation for estimating emissions from individual sources is generally represented as follows.

$$E \text{ (Emissions)} = EF \text{ (Emission Factor)} \times A \text{ (Activity Data)}$$

There are sources of emissions, however, where emissions are derived from stochastic equations comprising three or more parameters, and it becomes unclear which combination of parameters should be deemed as the emission factor and the activity data.

In such cases, emission factor and activity data are basically defined in accordance with the concept of emission factor described in the *Enforcement Ordinance for the Act on Promotion of Global Warming Counter Measures* (March 1999).

Example: A stochastic equation comprising three or more parameters

- Emission source: Methane emissions from a waste burial site (food scraps)
- Stochastic equation :
 Volume of emissions from the source
 = Carbon content in food scraps × Gas conversion rate of food scraps
 × Proportion of methane in generated gas × 16/12
 × Food scraps broken down during the basic period of calculation, expressed in tons
 = (*Emission Factor*: Carbon content of food scraps
 × Gas conversion rate of food scraps
 × Proportion of methane in gas generated × 16/12)
 × (*Activity Data*: Food scraps broken down during the basic period of calculation,
 expressed in tons)

A7.1.3.3. Uncertainty Assessment of Emission Factors

The uncertainty of emission factors is assessed using the following decision tree.

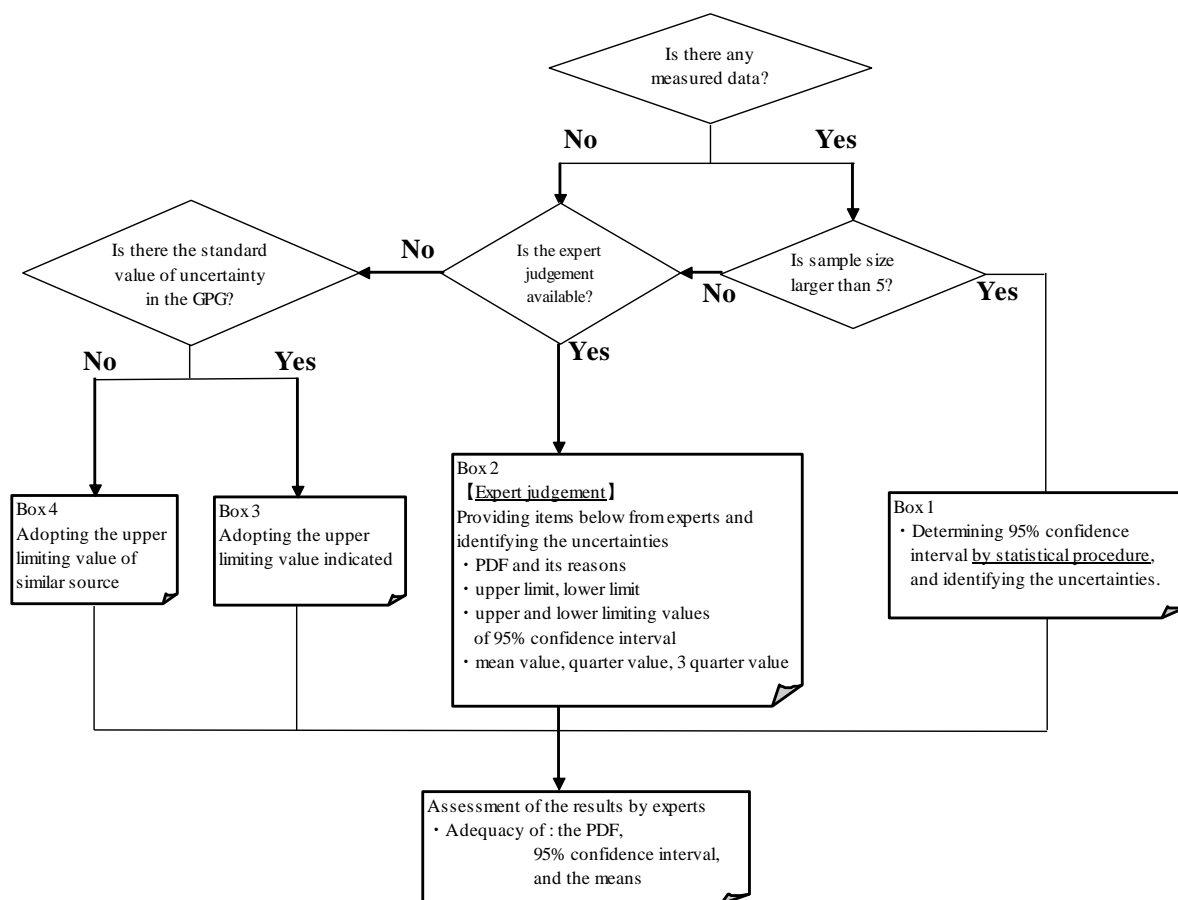


Figure A7-1 Decision tree for assessing uncertainty associated with emission factors established by the *Committee for the GHGs Emissions Estimation Methods*

- If an appropriate assessment cannot be made using the decision tree above, it may be done using a method that has been considered and deemed as appropriate. In such cases, the reason why an appropriate assessment could not be achieved using the decision tree, and the method applied, will both need to be clearly explained.

A7.1.3.3.a. Case where there is measurement data with five or more samples (Box 1)

Where data from actual measurements is available and there are five or more¹ samples, uncertainty is assessed quantitatively in accordance with the guidelines below.

Guidelines for assessment of uncertainty associated with emission factors	
Guideline 1	Where data from actual measurements is available and there are five or more samples, the central limit theorem says that the distribution of averages will follow a normal distribution curve. Assuming that all averages \bar{x} and standard deviations σ / \sqrt{n} follow a normal distribution curve, uncertainty need to be assessed on the basis of the data used to establish the emission factor only.

¹ The *Good Practice Guidance* cites “adequate samples”, but for convenience, the Secretariat of *Committee for the GHGs Estimation Methods* suggests the use of five or more.

Guideline 2

In assessing uncertainty, it is assumed that systematic error inherent to individual items of data is already a factor in the distribution. Therefore, systematic error inherent to individual items of data need not be investigated.

Guideline 3

Items that may contribute to uncertainty, but which may not be readily quantitatively assessable, should be recorded for the future investigation. If, through expert judgment, it is possible to estimate their uncertainty, the uncertainty shall be estimated in accordance with expert judgment.

a) When it is not possible to use statistical methods to derive the distribution of data used in calculating emission factors

1) Emission factor has been established by calculating a simple average of the sample data

Where the emission factor has been calculated using a simple average, it is assumed that the data used in calculating the emission factor follows a normal distribution curve. Therefore, the standard deviation of the sample is divided by the square root of the number of samples to estimate the standard deviation of the emission factor σ_{EF} , and uncertainty is calculated by finding the 95 percent confidence interval in accordance with Equation 1.1.

$$\text{Uncertainty of Emission Factor (\%)} = \frac{1.96 \times \sigma_{EF}}{|EF|} \quad \dots \text{Equation 1.1.}$$

σ_{EF} : Standard Deviation of Average
 EF : Emission Factor

2) Emission factor has been calculated using a weighted average of the sample data

Where the emission factor has been derived using a weighted average of the sample data, it is assumed that the data used in calculating the emission factor follows a normal distribution.

Therefore, the standard deviation σ_{EF} of the sample is derived using the equation below. Uncertainty is calculated by finding the 95 percent confidence interval of the averages in accordance with Equation 1.1. Note that the equation does not account for the uncertainty of weights w_i .

The weight applied in the weighted average, w_i ($\sum w_i = 1$)

Sample averages : $EF = \sum (w_i \times EFi)$

Unbiased variance of sample averages :

$$\sigma_{EF^2} = \sum \{w_i \times (EF_i - \overline{EF})^2\} / (1 - \sum w_i^2) \times \sum w_i^2$$

b) When the distribution of data used in calculating emission factor is derived using statistical methods

When it is possible to derive the distribution of data used in calculating the emission factor by using statistical methods, it is assumed that the data follows a normal distribution, and the uncertainty of each piece of data is estimated on the basis of section “a) When it is not possible to use statistical methods to derive the distribution of data used in calculating emission factors”. The uncertainty of

each piece of data is then determined using Equation 1.2, and the standard deviation of the emission factor σ_{EF} is calculated, to obtain the uncertainty.

When weight averaging is done to obtain at emission factors, the emission factor EF is expressed as follows, where the emission factor of each sub-category is EF_i , the weight variable is A_i , and the total of weight variables is A .

$$EF = \frac{\sum_i EF_i \times A_i}{\sum_i A_i} = \frac{\sum_i EF_i \times A_i}{A}$$

Substituting the distribution of the emission factor EF , σ_{EF}^2 , and the distributions of the individual emission factors EF_i and individual weight variables A_i , $\sigma_{EF_i}^2$ and $\sigma_{A_i}^2$, then σ_{EF}^2 is calculated as follows, using an equation known as the Error Propagation Equation.

$$\sigma_{EF}^2 = \sum_i \left\{ \left(\frac{\partial EF}{\partial EF_i} \right)^2 \sigma_{EF_i}^2 + \left(\frac{\partial EF}{\partial A_i} \right)^2 \sigma_{A_i}^2 \right\} = \sum_i \left\{ \frac{A_i^2}{A^2} \sigma_{EF_i}^2 + \frac{(EF_i - EF)^2}{A^2} \sigma_{A_i}^2 \right\}$$

... Equation 1.2.

Thus, the uncertainty of the emission factor U is obtained using the following equation.

$$U = \frac{1.96 \times \sigma_{EF}}{|EF|}$$

If experts at *Working Group on Inventory of Committee for the GHGs Emissions Estimation Methods* indicate that statistical analysis is inappropriate, even using five or more samples, then uncertainty should be assessed by expert judgment. Conversely, if an expert determines that it is possible to carry out statistical analysis, even with less than five samples, uncertainty shall be assessed statistically.

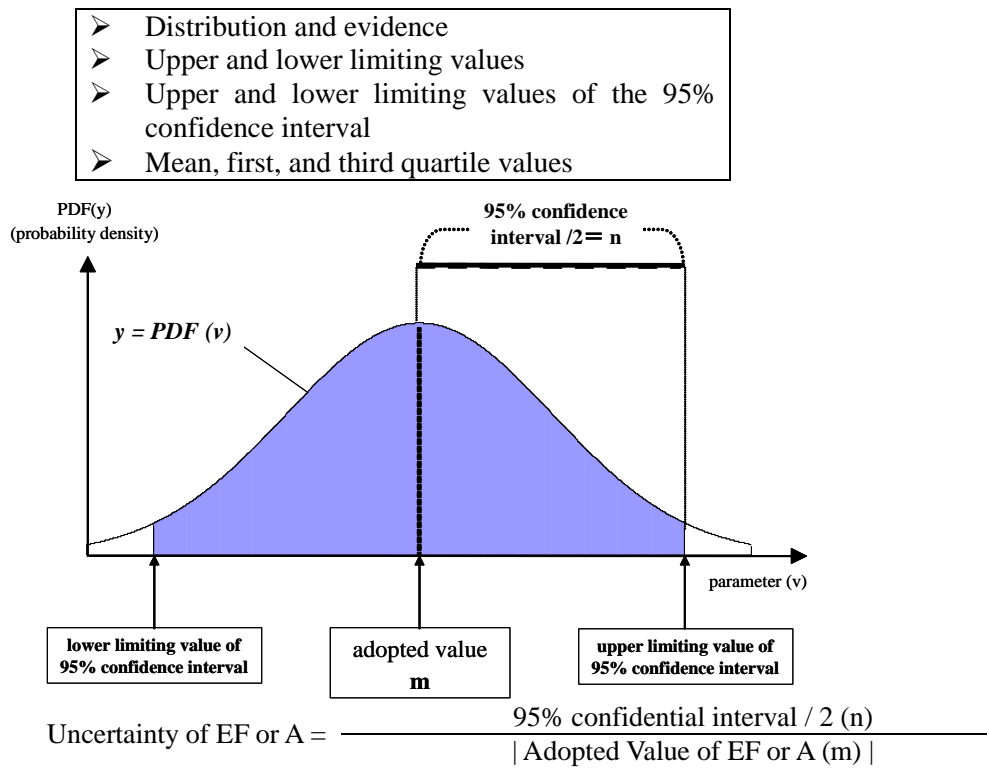
A7.1.3.3.b. Case where there is no actual measurement data, or there are less than five samples

When there is no actual measurement data, or there are less than five samples, uncertainty shall be assessed by expert judgment.

a) When expert judgment is feasible (Box 2)

1) When the distribution of the probability density function of emission factors can be obtained using expert judgment

In this case, uncertainty should be assessed in accordance with expert judgment for the following. The expert providing the expert judgment, the basis for their decision, and factors contributing to uncertainty that are excluded from consideration, should be documented, and the document should be retained.

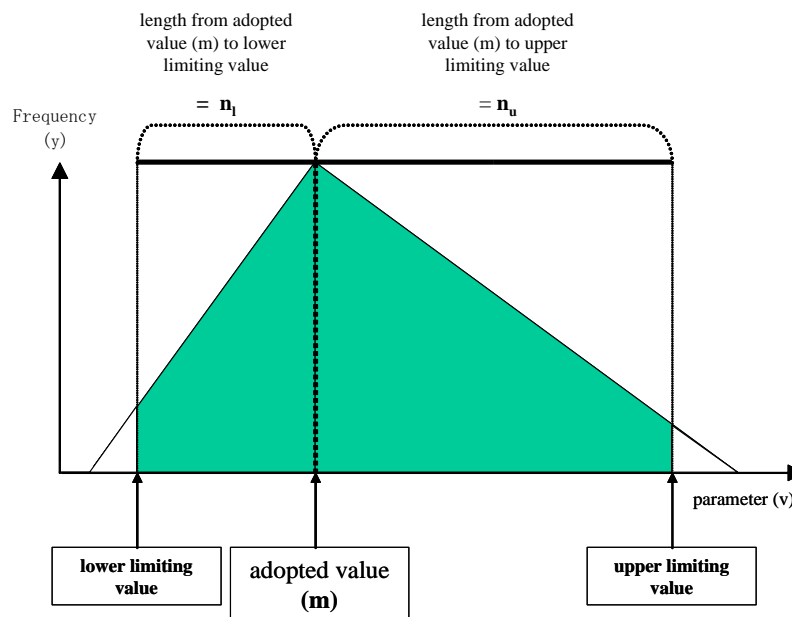


2) When the distribution of the probability density function of emission factors cannot be obtained using expert judgment

Ask an expert for the upper and lower limiting values appropriate to emission factors in Japan, and draw a triangular distribution for the emission factors with the Japanese emission factor as the vertex, and such that the upper and lower limiting values of a 95 percent confidence interval correspond to the upper and lower limiting values appropriate to the Japanese emission factor (see diagram below).

If the emission factor used is larger than the upper limiting value, the emission factor should be used as the upper limiting value. If the emission factor used is smaller than the lower limiting value, the emission factor should be used as the lower limiting value.

The expert providing the expert judgment, the basis for their decision, and factors contributing to uncertainty that are excluded from consideration, should be documented, and the document should be retained.



Uncertainty in this context is calculated using the following equation.

<p>Uncertainty to the lower limiting value U_l (%) $= - \{ \text{distance to lower limiting value } (n_l) / \text{mode } (m) \}$</p> <p>Uncertainty to the upper limiting value U_u (%) $= + \{ \text{distance to upper limiting value } (n_u) / \text{mode } (m) \}$</p> <p>Uncertainty is expressed in the form, $-○\%$ to $+●\%$, but in assessing overall uncertainty for Japan, the largest absolute value should be used.</p>

b) When expert judgment is not possible

1) A standard value for uncertainty is provided in the Good Practice Guidance (Box 3)

When the *Good Practice Guidance* provides a standard value for uncertainty for a particular emission source, an estimate of uncertainty should err on the safe side, and the upper limiting value of the standard uncertainty value given in the *Good Practice Guidance* should be used.

2) No standard value for uncertainty is provided in the Good Practice Guidance (Box 4)

When the *Good Practice Guidance* does not provide a standard uncertainty for a particular emission source, the standard uncertainty given in the *Good Practice Guidance* for a similar emission source should be used for the upper limiting value.

Category	Uncertainty of EF
1. Energy	
1.A. CO ₂	5%
1.A. CH ₄ , N ₂ O	3%~10%
1.A.3. Transport(CH ₄ , N ₂ O)	5%
2. Industrial Processes	
Excluding HFCs, PFCs, SF ₆	1%~100%
HFCs, PFCs, SF ₆	5%~50%
3. Solvent and Other Product Use	-*
4. Agriculture	2%~60%
5. Land Use Change and Forestry	-**
6. Waste	5%~100%

* Category 3: The use of organic solvents and other such products are not dealt within the GPG (2000).

** Category 5: Changes in land use and forestry are not dealt with in the GPG (2000).

A7.1.3.3.c. Methods for Combining Uncertainties of Emission Factors

The basic method for combining uncertainties is Tier 1 in the *Good Practice Guidance*. When a correlation between elements is strong, uncertainties may be combined using the Monte Carlo method (Tier 2 in the *Good Practice Guidance*).

a) Uncertainty of emission factor derived from a combination of multiple parameters

The uncertainty of an emission factor may be obtained at from the uncertainty of multiple parameters using the equation given below, in situations of the type described in the example in Section A7.1.3.2..

$$U_{EF} = \sqrt{U_1^2 + U_2^2 + \dots + U_n^2}$$

U_{EF} : Uncertainties of Emission Factors (%)
 U_i : Uncertainties of Parameter "i" (%)

A7.1.3.4. Uncertainty Assessment of Activity Data

The uncertainty of activity data is assessed in accordance with the decision tree depicted below.

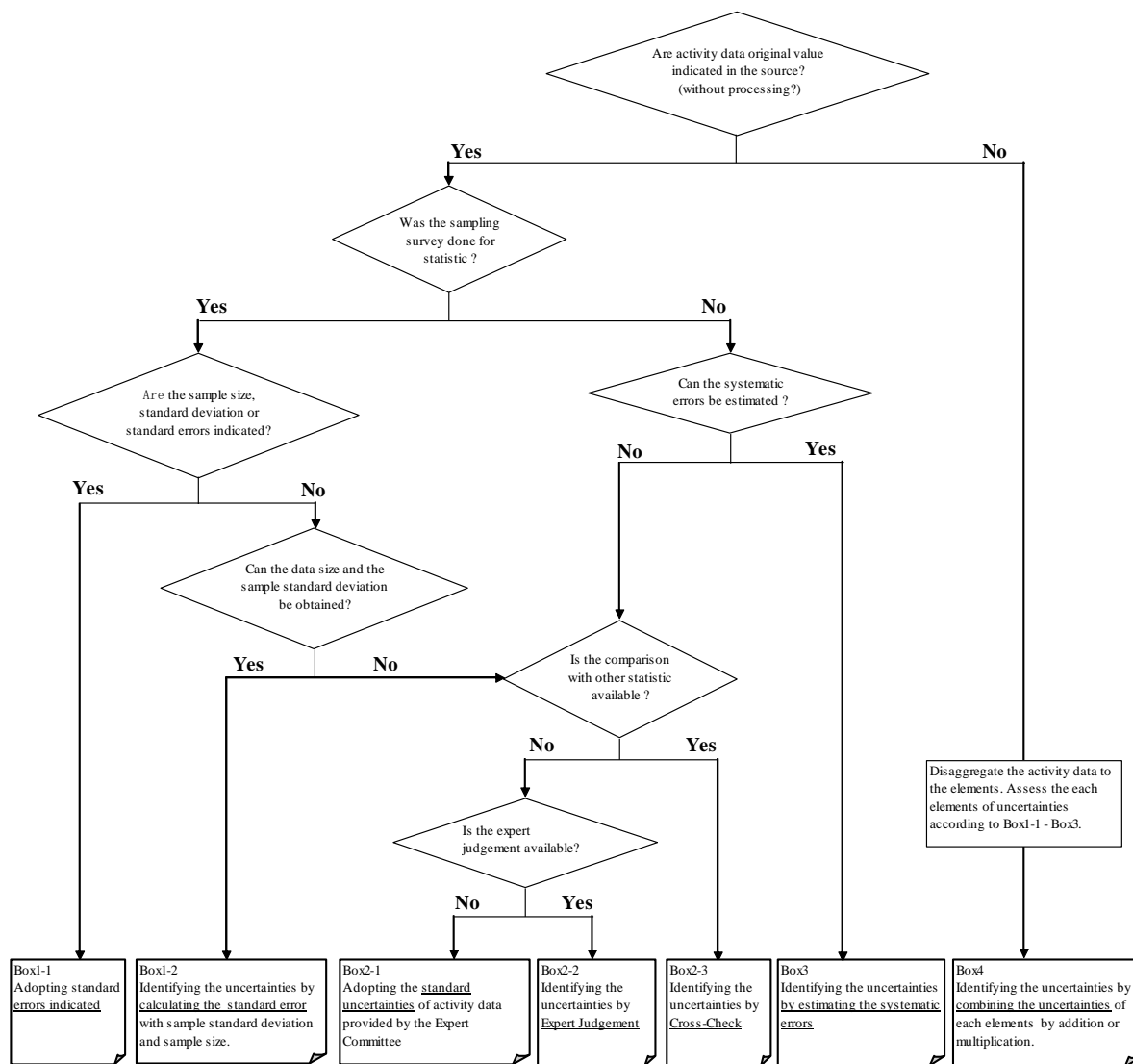


Figure A7-2 Decision tree for assessing uncertainty associated with activity data established by the *Committee for the GHGs Emissions Estimation Methods*

➤ If an appropriate assessment cannot be made using the decision tree above, it may be done using a method that has been considered and deemed as appropriate. The reason why an appropriate assessment could not be achieved using the decision tree, and the method applied, will both need to be clearly explained.

A7.1.3.4.a. Using statistical values for activity data

When using statistical values for activity data, uncertainty should be quantitatively assessed in accordance with the following guidelines.

Guidelines for assessment of uncertainty associated with activity data
<p><u>Guideline 1</u> Only the sample error needs to be considered as part of uncertainty assessment in sample surveys.</p>

Guideline 2

In situations other than sample surveys, if it is possible to estimate a systemic error, it should be considered as part of an uncertainty assessment.

Guideline 3

In situations other than sample surveys, if it is not possible to estimate a systemic error, uncertainty should be assessed through crosschecks, or by expert judgment.

Guideline 4

Where quantitative assessment is difficult, factors that would contribute to uncertainty should be recorded for a future investigation.

a) Statistical values based on a sample survey**1) The publisher has made errors public (Box 1-1)**

When the publisher of a statistical document has made the sampling errors public in the sample survey, it should be used as the uncertainty of the activity data.

2) The publisher has not made errors public (Box 1-2)

Enquire the publisher of the statistical document for the size of the sample, the sample average, and the standard deviation of the sample. Under the assumption that the distribution of the sample reproduces the distribution of the population, assessment of uncertainty from the statistical values should be done.

$$\text{Uncertainty } U = (1.96 \times s / \sqrt{n}) / X_{ad}$$

X_{ad} : Sample average

S : Standard deviation of sample

n : Number of items of data

If, however, distribution is asymmetrical, the uncertainty U is calculated by dividing the difference between the value of the 95 percent confidence limit furthest from X_{ad} and the average value, by X_{ad} .

Confirmation of the estimation method for Japan from values drawn from the sample survey and, as far as possible, estimation of the uncertainty associated with the estimation method should be done also (e.g., multiply the sample average of the number of head of livestock raised per farm by the number of farms).

3) Amount of data and sample standard deviation are not available, and crosschecking is possible (Box 2-3)

In the case of statistics drawn from a sample survey, where the amount of data and the sample standard deviation are not available, but it is possible to compare the relevant statistical value with multiple other statistical values, uncertainty should be assessed using the same means as in the second case described at section A1.2.3 in the page A1.7 of the *Good Practice Guidance*.

$$\text{Uncertainty } U = (1.96 \times s) / X_{ap}$$

X_{ap} : Value used for activity data

s : Standard deviation (data to be cross-checked)

However, if a distribution is asymmetrical, the uncertainty U may be calculated by dividing the difference between the value of the 95 percent confidence limit furthest from X_{ap} and the average value, by X_{ap} .

Also, when there is a single other statistical value only, the assessment should be done using the same method described at 2) “When the distribution of the probability density function of emission factors cannot be obtained using expert judgment” in *Section 7.1.3.3.b.*

4) Amount of data and sample standard deviation are not available, and expert judgment is available (Box 2-2)

In the case of statistics drawn from a sample survey where the amount of data and sample standard deviation are not available, ask an expert for the upper and lower limiting values appropriate to activity data in Japan, and draw a triangular distribution for activity data (see diagram at Section A7.1.3.3.b.) with the Japanese activity data as the vertex, and such that the upper and lower limiting values of a 95 percent confidence interval correspond to the upper and lower limiting values appropriate to the Japanese activity data.

If the activity data used is larger than the upper limiting value, that activity data should be used as the upper limiting value. If the activity data used is smaller than the lower limiting value, that emission factor should be taken as the lower limiting value.

The experts providing the expert judgment, the basis for their decision, and factors contributing to uncertainty that are excluded from consideration, should be documented, and the document should be retained.

5) Amount of data and sample standard deviation are not available, and expert judgment is unavailable (Box 2-1)

The following standard values established by the *Committee for the GHGs Emissions Estimation Methods* will be used.

Table A7-1 Uncertainty of sample statistics established by the Committee for the GHGs Emissions Estimation Methods

	Fundamental statistics	Other statistics
Sample survey	50 [%]	100 [%]

The values for fundamental statistics, approved statistics, and reported statistics have been established by the Committee for the GHGs Emissions Estimation Methods, with reference to the *Good Practice Guidance* and other material. Statistics other than fundamental statistics have been deemed to be twice the fundamental statistics.

b) Statistical values not based on a sample survey

1) Systemic error can be estimated (Box 3)

Where a systemic error can be estimated, it should be estimated and used. The method by which the systemic error is calculated should be documented, and the document should be retained.

2) Systemic error cannot be estimated, and crosschecking is possible (Box 2-3)

Where systemic error cannot be estimated, but it is possible to compare the relevant statistical value with other statistical values, uncertainty should be assessed using the same means as in Case 2 described at A1.2.3 of Section A1.7 of the *Good Practice Guidance*.

3) Systemic error cannot be estimated, crosschecking is not possible, and expert judgment is available (Box 2-2)

Same as for “4) Amount of data and sample standard deviation are not available, and expert judgment is available (Box 2-2)” in Section 7.1.3.4.a.

4) Systemic error cannot be estimated, crosschecking is not possible, and expert judgment is unavailable (Box 2-1)

The following standard values established by the Committee for the GHGs Emissions Estimation Methods should be used.

Table A7-2 Uncertainty of sample statistics established by the Committee for the GHGs Emissions Estimation Methods

	Fundamental statistics	Other statistics
Survey of total population (no rounding)	5 [%]	10 [%]
Survey of total population (rounding)	20 [%]	40 [%]

The values for fundamental statistics, approved statistics, and reported statistics have been established by the Committee for the GHGs Emissions Estimation Methods with reference to the *Good Practice Guidance* and other material. Statistics other than fundamental statistics have been deemed to be twice the fundamental statistics.

A7.1.3.4.b. Using statistical values processed as activity data (Box 3)

a) Breakdown of each element of activity data and assessment

Activity data should be broken down as shown in the following example.

<ul style="list-style-type: none"> ➤ Emission source : CO₂ emission from incineration of naphtha in the chemical industry ➤ Stochastic equation : <p>Activity data for relevant emission source = Naphtha consumption × 20% (remaining 80% is fixed in the product) ² - ammonia raw material</p>
--

After being broken down, each element of the statistical values should be assessed for uncertainty using the method shown at section “7.1.3.4.a. Using statistical values for activity data”.

In the example above, for elements based on survey research, such as the figure of 20%, uncertainty should be assessed on the basis of the method shown at section “7.1.3.3. Uncertainty Assessment of Emission Factors”.

b) Combining elements

Combine each element using the sum and product methods of combination, and assess the uncertainty.

- Sum method (Rule A): Where uncertainty quantities are to be combined by addition.
Activity data is expressed as $A_1 + A_2$

² Environmental Agency, *The Estimation of CO₂ Emission in Japan*, 1992

$$U_{A-total} = \frac{\sqrt{(U_{A1} \times A_1)^2 + (U_{A2} \times A_2)^2}}{A_1 + A_2}$$

U_{An} : Uncertainty of element An (%)

- Product method: Where uncertainty quantities are to be combined by multiplication. Activity data is expressed as $A_1 \times A_2$

$$U_A = \sqrt{U_{A1}^2 \times U_{A2}^2}$$

U_{An} : Uncertainty of element An (%)

A7.1.3.5. Uncertainty Assessment of Emissions

A7.1.3.5.a. Uncertainty assessment of emissions from individual emission sources

1) Emissions estimated from emission factor and activity data

Use the product combination equation given at Tier 1 of the *Good Practice Guidance* on the results of emission factor assessment from the previous section and the activity data, and assess the uncertainty of emissions from each emission source.

$$U_{Ei} = \sqrt{U_{EFi}^2 + U_{Ai}^2}$$

U_{Ei} : Uncertainty of emissions from emission source i (%)
 U_{EFi} : Uncertainty of element An (%)
 U_{Ai} : Uncertainty of element An (%)

2) Actual measurements taken of emissions

When emissions are derived from actual measurement, uncertainty of emissions should be assessed directly, in accordance with “7.1.3.3. *Uncertainty Assessment of Emission Factors*”.

A7.1.3.5.b. Calculating uncertainty of total emissions

Combine the results of assessments of emission uncertainty for multiple emission sources to assess the uncertainty of total Japanese emissions of greenhouse gases. The uncertainty of emissions from multiple sources should be combined using the product combination equation given at Tier 1 in the *Good Practice Guidance*.

$$U_{Total} = \frac{\sqrt{(U_1 \times E_1)^2 + (U_2 \times E_2)^2 + \dots + (U_n \times E_n)^2}}{E_1 + E_2 + \dots + E_n}$$

U_{Total} : Uncertainty of total Japanese emissions (%)
 U_i : Uncertainty of emission source i (%)
 E_i : Emissions from emission source i (Gg)

When the uncertainties of emissions from multiple sources are combined, only the uncertainty of emissions should be indicated. Combination of the uncertainties for both emission factor and activity data should not be done.

A7.2. Results of Uncertainty Assessment

A7.2.1. Assumption of Uncertainty Assessment

Uncertainty Assessment is basically conducted based on the results of uncertainty assessment in

Committee for the Greenhouse Gases Emissions Estimation Methods in FY 2006.

A7.2.2. Uncertainty of Japan's Total Emissions

In FY 2010, total net emissions in Japan were approximately 1,185 million tons (carbon dioxide equivalents). Uncertainty of total net emissions has been assessed at 2% and uncertainty introduced into the trend in total net emissions has been assessed at 1%.

Table A7-3 Uncertainty of Japan's Total Net Emissions

IPCC Category	GHGs	Emissions / Removals [Gg-CO ₂ eq.]		Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions ¹⁾	rank
		A	[%]				
1A. Fuel Combustion (CO ₂)	CO ₂	1,137,550.9	90.4%	1%	10	0.74%	2
1A. Fuel Combustion (Stationary:CH ₄ ,N ₂ O)	CH ₄ , N ₂ O	4,957.6	0.4%	27%	4	0.11%	8
1A. Fuel Combustion (Transport:CH ₄ ,N ₂ O)	CH ₄ , N ₂ O	2,694.0	0.2%	351%	1	0.80%	1
1B. Fugitive Emissions from Fuels	CO ₂ , CH ₄ , N ₂ O	409.0	0.0%	19%	5	0.01%	9
2. Industrial Processes (CO ₂ ,CH ₄ ,N ₂ O)	CO ₂ , CH ₄ , N ₂ O	42,373.9	3.4%	7%	8	0.25%	7
2. Industrial Processes (HFCs,PFCs,SF ₆)	HFCs, PFCs, SF ₆	23,524.2	1.9%	33%	2	0.66%	4
3. Solvent & other Product Use	N ₂ O	99.0	0.0%	5%	9	0.00%	10
4. Agriculture	CH ₄ , N ₂ O	25,499.6	2.0%	18%	6	0.39%	6
5. LULUCF	CO ₂ , CH ₄ , N ₂ O	-73,179.1	-5.8%	12%	7	0.71%	3
6. Waste	CO ₂ , CH ₄ , N ₂ O	20,873.8	1.7%	32%	3	0.57%	5
Total Net Emissions		(D) 1,184,802.8		(E) ²⁾ 2%			

1) $C = A \times B / D$

2) $E = \sqrt{C_1^2 + C_2^2 + \dots}$

Hereafter, the same method for calculating uncertainty assessment has been used in each sector appearing in Table A7-4 and the following tables.

A7.2.3. Energy Sector

A7.2.3.1. Fuel Combustion (CO₂)

Carbon-Hydrogen ratio of hydrocarbons is strongly correlating with calorific value in theory, then, standard deviation of sample data of each fuel's calorific value are used for uncertainty assessment based on assumption that deviation of carbon content and that of calorific value is equal. The uncertainty of energy consumption in TJ given in the *General Energy Statistics* was assessed based on the given statistical error of solid fuels, liquid fuels, and gaseous fuels, since it was difficult to set uncertainty by fuel types and industry.

Table A7-4 Results of uncertainty assessment of fuel combustion (CO₂)

IPCC Category			GHGs	Emissions / Removals [Gg-CO ₂ eq.]	EF/RF Uncertainty [%]	AD Uncertainty [%]	Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions	rank	
				A	a	b	B ³⁾		C		
1A. Fuel Combustion	Solid Fuels	Steel Making Coal	CO ₂	16,498.3	3.5%	1.2%	4%	19	0.05%	13	
		Steam Coal (imported)	CO ₂	249,085.2	2.0%	1.2%	2%	31	0.49%	1	
		Steam Coal (indigenous)	CO ₂	0.0	2.0%	1.2%	2%	31	0.00%	38	
		Hard Coal	CO ₂	0.0	4.5%	1.2%	5%	16	0.00%	38	
		Coke	CO ₂	94,312.3	1.7%	1.2%	2%	39	0.17%	5	
		Coal Tar	CO ₂	3,332.8	5.0%	1.2%	5%	14	0.01%	26	
		Coal Briquette	CO ₂	0.0	5.0%	1.2%	5%	14	0.00%	38	
		Coke Oven Gas	CO ₂	14,654.1	2.0%	1.2%	2%	31	0.03%	18	
		Blast Furnace Gas	CO ₂	43,333.5	3.8%	1.2%	4%	17	0.15%	7	
		Converter Furnace Gas	CO ₂	10,260.1	2.9%	1.2%	3%	20	0.03%	19	
		Liquid Fuels	Crude Oil for Refinery	CO ₂	0.0	0.8%	2.3%	2%	26	0.00%	38
			Crude Oil for Power Generation	CO ₂	12,925.0	0.9%	2.3%	2%	25	0.03%	20
			Vituminous Mixture Fuel	CO ₂	0.0	0.4%	2.3%	2%	30	0.00%	38
	NGL & Condensate		CO ₂	39.8	1.6%	2.3%	3%	21	0.00%	36	
	Naphtha		CO ₂	588.1	0.1%	2.3%	2%	34	0.00%	31	
	Reformed Material Oil		CO ₂	0.0	0.1%	2.3%	2%	34	0.00%	38	
	Gasoline		CO ₂	134,709.0	0.03%	2.3%	2%	38	0.26%	3	
	Jet Fuel		CO ₂	12,715.7	1.0%	2.3%	3%	24	0.03%	21	
	Kerosene		CO ₂	48,423.2	0.05%	2.3%	2%	37	0.09%	10	
	Gas Oil or Diesel Oil		CO ₂	84,665.7	1.2%	2.3%	3%	23	0.19%	4	
	Heating Oil A		CO ₂	43,332.9	1.5%	2.3%	3%	22	0.10%	9	
	Heating Oil B		CO ₂	51.3	5.0%	2.3%	6%	10	0.00%	35	
	Heating Oil C		CO ₂	54,969.8	0.6%	2.3%	2%	27	0.11%	8	
	Lubricating Oil		CO ₂	208.7	5.0%	2.3%	6%	10	0.00%	32	
	Asphalt		CO ₂	10,816.8	0.6%	2.3%	2%	27	0.02%	24	
	Non Asphalt Heavy Oil Products		CO ₂	0.0	0.6%	2.3%	2%	27	0.00%	37	
	Oil Coke		CO ₂	14,250.6	5.0%	2.3%	6%	10	0.07%	12	
	Galvanic Furnace Gas		CO ₂	143.1	2.9%	2.3%	4%	18	0.00%	33	
	Refinery Gas		CO ₂	33,990.7	5.0%	2.3%	6%	10	0.16%	6	
	LPG		CO ₂	26,384.0	0.1%	2.3%	2%	34	0.05%	14	
	Gaseous Fuels	LNG	CO ₂	124,372.8	0.1%	0.3%	0%	42	0.03%	17	
		Indigenous Natural Gas	CO ₂	2,626.9	0.6%	0.3%	1%	40	0.00%	30	
		Town Gas*	CO ₂	85,536.1	0.5%	0.3%	1%	41	0.04%	16	
		Small Scale Town Gas*	CO ₂	1,144.7	0.1%	0.3%	0%	42	0.00%	34	
	Other Fuels	Municipal Solid Waste (Plastics)	CO ₂	5,112.9	4.3%	16.0%	17%	6	0.07%	11	
		Municipal Solid Waste (Waste textile)	CO ₂	1,172.8	4.3%	22.4%	23%	5	0.02%	23	
		Industrial Solid Waste (Waste Mineral Oil)	CO ₂	102.0	4.8%	104.4%	105%	1	0.01%	28	
		Industrial Solid Waste (Plastics)	CO ₂	281.3	4.8%	100.0%	100%	3	0.02%	22	
		Raw material and fuel use of MSW	CO ₂	451.7	4.3%	16.0%	17%	6	0.01%	29	
		Raw material and fuel use of ISW (Waste Mineral Oil)	CO ₂	3,234.6	4.8%	104.4%	105%	1	0.29%	2	
		Raw material and fuel use of ISW (Waste Plastics)	CO ₂	1,453.5	4.8%	12.3%	13%	9	0.02%	25	
		Raw material and fuel use of Waste tire	CO ₂	1,002.7	4.8%	14.5%	15%	8	0.01%	27	
		Fuel use of RDF and RPF	CO ₂	1,368.3	42.6%	10.6%	44%	4	0.05%	17	
		Sub Total			1,137,550.9			1%		0.74%	
	Total Emissions	(D)		1,184,802.8			2%				

* Reported in Gaseous Fuels according to the main material; LNG

3) $B = \sqrt{a^2 + b^2}$ (Hereafter, the same method has been used in each sector appearing in Table A7-5 and following)A7.2.3.2. Stationary Combustion (CH₄ and N₂O)

Table A7-5 Results of uncertainty assessment of fuel combustion

IPCC Category			GHGs	Emissions / Removals [Gg-CO ₂ eq.]	EF/RF Uncertainty [%]	AD Uncertainty [%]	Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions	rank	
				A	a	b	B		C		
1A. Fuel Combustion (Stationary)			CH ₄	588.6	— ⁴⁾	— ⁴⁾	47%	12	0.02%	2	
			N ₂ O	3,942.8	— ⁴⁾	— ⁴⁾	33%	15	0.11%	1	
C. Waste Incineration	Municipal Solid Waste	Waste	CH ₄	2.5	—	—	101%	7	0.00%	9	
		Waste	N ₂ O	293.6	—	—	42%	13	0.01%	3	
	Industrial Solid Waste	Waste	CH ₄	0.2	111.5%	100.0%	150%	2	0.00%	15	
		Waste	N ₂ O	3.0	58.8%	100.0%	116%	4	0.00%	7	
	Raw material and fuel use of MSW			CH ₄	0.0	179.4%	10.0%	180%	1	0.00%	18
				N ₂ O	0.0	111.2%	10.0%	112%	5	0.00%	17
	Raw material and fuel use of ISW	Waste Oil (total)	Waste	CH ₄	0.5	—	—	74%	10	0.00%	10
			Waste	N ₂ O	13.1	—	—	41%	14	0.00%	11
		Waste Plastics	Waste	CH ₄	3.3	91.7%	10.0%	92%	8	0.00%	14
			Waste	N ₂ O	4.4	29.7%	10.0%	31%	17	0.00%	6
		Waste Wood	Waste	CH ₄	78.2	80.2%	100.0%	128%	3	0.01%	4
			Waste	N ₂ O	13.1	45.3%	100.0%	110%	6	0.00%	5
	Raw material and fuel use of Waste tire			CH ₄	0.9	—	—	91%	9	0.00%	13
		N ₂ O	5.3	—	—	26%	18	0.00%	12		
Fuel use of RDF and RPF			CH ₄	0.2	—	—	49%	11	0.00%	16	
			N ₂ O	7.9	—	—	33%	16	0.00%	8	
Sub Total				4,957.6			27%		0.11%		
Total Emissions	(D)			1,184,802.8			2%				

4) Because “—” means aggregation of detailed sub-categories, uncertainties of EF/RF and AD can not be calculated for this level of disaggregation of categories.

A7.2.3.3. Mobile Combustion (CH₄ and N₂O)Table A7-6 Results of uncertainty assessment of mobile combustion (CH₄ and N₂O)

IPCC Category		GHGs	Emissions / Removals [Gg-CO ₂ eq.]	EF/RF Uncertainty [%]	AD Uncertainty [%]	Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions	rank	
			A	a	b	B		C		
1A. Fuel Combustion (Transport)	a. Civil Aviation	CH ₄	4.6	200.0%	10.0%	200%	4	0.00%	6	
		N ₂ O	92.9	10000.0%	10.0%	10000%	1	0.78%	1	
	b. Road Transportation	CH ₄	140.8	40.0%	50.0%	64%	6	0.01%	4	
		N ₂ O	2,267.1	50.0%	50.0%	71%	5	0.14%	2	
	c. Railways	CH ₄	0.7	—	—	14%	7	0.00%	8	
		N ₂ O	75.0	—	—	11%	8	0.00%	7	
	d. Navigation	CH ₄	21.7	200.0%	13.0%	200%	3	0.00%	5	
		N ₂ O	91.3	1000.0%	13.0%	1000%	2	0.08%	3	
	Sub Total			2,694.0			351%		0.80%	
	Total Emissions		(D)	1,184,802.8				2%		

(Note) CO₂ emissions from 1A Fuel Combustion (Transport) have been reported in Table A7-4.

A7.2.3.4. Fugitive Emissions from Fuel

Table A7-7 Results of uncertainty assessment of fugitive emissions from fuel

IPCC Category				GHGs	Emissions / Removals [Gg-CO ₂ eq.]	EF/RF Uncertainty [%]	AD Uncertainty [%]	Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions	rank	
					A	a	b	B		C		
1B. Fugitive Emissions from Fuels	1. Solid Fuels	a. Coal Mining	i. Underground Mines	Mining Activities	CH ₄	14.4	—	—	5%	24	0.00%	12
			Post-Mining Activities	CH ₄	20.3	200.0%	10.0%	200%	1	0.00%	2	
		ii. Surface Mines	Mining Activities	CH ₄	9.0	200.0%	10.0%	200%	1	0.00%	3	
			Post-Mining Activities	CH ₄	0.8	200.0%	10.0%	200%	1	0.00%	11	
	2. Oil and Natural Gas	a. Oil	i. Exploration	CO ₂	0.02	25.0%	10.0%	27%	7	0.00%	20	
				CH ₄	0.02	25.0%	10.0%	27%	6	0.00%	21	
				N ₂ O	0.00006	25.0%	10.0%	27%	4	0.00%	24	
				CH ₄	1.4	25.0%	5.0%	25%	9	0.00%	17	
			ii. Production	CO ₂	0.08	25.0%	5.0%	25%	9	0.00%	9	
				CH ₄	8.9	25.0%	5.0%	25%	9	0.00%	9	
				CO ₂	0.0047	25.0%	5.0%	25%	9	0.00%	22	
				CH ₄	1.4	25.0%	5.0%	25%	9	0.00%	14	
		iv. Refining / Storage	CH ₄	14.3	25.0%	0.9%	25%	23	0.00%	7		
			b. Natural Gas	ii. Production / Processing	CO ₂	0.4	25.0%	5.0%	25%	9	0.00%	16
					CH ₄	256.2	25.0%	5.0%	25%	9	0.01%	1
				iii. Transmission	CH ₄	23.1	25.0%	10.0%	27%	4	0.00%	4
	CH ₄	16.3			25.0%	8.7%	26%	8	0.00%	6		
	c. Venting and Flaring	Venting	i. oil	CO ₂	0.0	25.0%	5.0%	25%	9	0.00%	23	
				CH ₄	8.5	25.0%	5.0%	25%	9	0.00%	10	
			Flaring	i. oil	CO ₂	19.6	25.0%	5.0%	25%	9	0.00%	5
					CH ₄	0.85	25.0%	5.0%	25%	9	0.00%	15
		ii. Gas		N ₂ O	0.058	25.0%	5.0%	25%	9	0.00%	18	
				CO ₂	13.0	25.0%	5.0%	25%	9	0.00%	8	
CH ₄		1.7	25.0%	5.0%	25%	9	0.00%	13				
N ₂ O		0.048	25.0%	5.0%	25%	9	0.00%	19				
Sub Total					409.0			19%		0.01%		
Total Emissions				(D)	1,184,802.8				2%			

A7.2.4. Industrial Processes

A7.2.4.1. CO₂, CH₄ and N₂O

For emissions sources with actual data available for emission factors, the emission factor dataset is deemed to be a sample of the total dataset, and the uncertainty assessment is achieved statistically. It is not a synthesis of the uncertainties of measured error of emissions from each operating site.

Table A7-8 Results of uncertainty assessment of industrial processes (CO₂, CH₄ and N₂O)

IPCC Category			GHGs	Emissions / Removals [Gg-CO ₂ eq.]	EF/RF Uncertainty [%]	AD Uncertainty [%]	Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions	rank	
				A	a	b	B		C		
2. Industrial Processes	A. Mineral Products	1. Cement Production		CO ₂	23,784.4	3.0%	10.0%	10%	15	0.21%	1
		2. Lime Production		CO ₂	6,284.6	15.0%	5.0%	16%	14	0.08%	3
		3. Limestone & Dolomite Use	Limestone	CO ₂	6,483.7	16.4%	4.8%	17%	12	0.09%	2
			Dolomite	CO ₂	1,589.5	3.5%	3.9%	5%	17	0.01%	8
	4. Soda Ash Production and Use		CO ₂	137.9	15.0%	6.3%	16%	13	0.00%	10	
	B. Chemical Industries	1. Ammonia Production		CO ₂	2,106.4	22.5%	5.0%	23%	11	0.04%	5
		Chemical Industries other than Ammonia		CO ₂	630.8	77.2%	5.0%	77%	8	0.04%	4
		2. Nitric Acid		N ₂ O	561.6	46.0%	5.0%	46%	10	0.02%	6
		3. Adipic Acid		N ₂ O	516.1	9.0%	2.0%	9%	16	0.00%	9
		4. Carbide		CH ₄	0.66	100.0%	10.0%	100%	5	0.00%	17
		5. Other	Carbon Black	CH ₄	5.4	54.8%	5.0%	55%	9	0.00%	14
			Ethylene	CH ₄	2.2	77.2%	5.0%	77%	7	0.00%	16
			Dichloroethylene	CH ₄	0.33	100.7%	5.0%	101%	4	0.00%	18
			Styrene	CH ₄	2.0	113.2%	5.0%	113%	3	0.00%	15
			Methanol	CH ₄	0.0	NA	NA	NA	NA	NA	NA
	Coke		CH ₄	93.5	98.5%	5.0%	99%	6	0.01%	7	
	C. Metal Production	1. Iron and steel		CO ₂	159.9	—	—	5%	18	0.00%	12
				CH ₄	12.3	163.0%	5.0%	163%	1	0.00%	11
		2. Ferroalloy		CH ₄	2.6	163.0%	5.0%	163%	1	0.00%	13
	Sub Total				42,373.9			7%		0.25%	
	Total Emissions			(D)	1,184,802.8			2%			

A7.2.4.2. F-gases

Table A7-9 Results of uncertainty assessment of industrial processes (F-gases)

IPCC Category				GHGs	Emissions / Removals [Gg-CO ₂ eq.]	EF/RF Uncertainty [%]	AD Uncertainty [%]	Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions	rank		
					A	a	b	B		C			
2. Industrial Processes (F-gas)	C. Metal Production	3. Aluminium		PFCs	10.4	33.0%	5.0%	33%	30	0.00%	21		
		4. SF ₆ Used in Aluminium and Magnesium Foundries		SF ₆	307.9	—	5.0%	5%	32	0.00%	19		
	E. Production of F-gas	1. By-product Emissions (HCFC-22)		HFCs	42.1	2.0%	5.0%	5%	31	0.00%	22		
		2. Fugitive Emissions		HFCs	86.2	100.0%	10.0%	100%	1	0.01%	14		
	F. Consumption of F-gas	1. Refrigeration and Air Conditioning Equipment	Domestic Refrigerator	manufacturing stock	HFCs	359.1	50.0%	40.0%	64%	6	0.02%	8	
				disposal	HFCs	IE	50.0%	40.0%	64%	6	0.00%	24	
					HFCs	IE	—	40.0%	40%	20	0.00%	24	
			Commercial Refrigerator	manufacturing stock	HFCs	11,336.4	50.0%	40.0%	64%	6	0.61%	1	
				disposal	HFCs	IE	50.0%	40.0%	64%	6	0.00%	24	
					HFCs	IE	—	40.0%	40%	20	0.00%	24	
			Stationary Air-Conditioning	manufacturing stock	HFCs	2,890.2	50.0%	40.0%	64%	6	0.16%	2	
				disposal	HFCs	IE	50.0%	40.0%	64%	6	0.00%	24	
					HFCs	IE	—	40.0%	40%	20	0.00%	24	
			Mobile Air-Conditioning	manufacturing stock	HFCs	2,502.4	50.0%	40.0%	64%	6	0.14%	3	
		disposal		HFCs	IE	50.0%	40.0%	64%	6	0.00%	24		
				HFCs	IE	—	40.0%	40%	20	0.00%	24		
		2. Foam Blowing	manufacturing stock		HFCs	138.6	50.0%	50.0%	71%	4	0.01%	13	
					HFCs	152.4	50.0%	50.0%	71%	4	0.01%	12	
		3. Fire Extinguisher		manufacturing stock		HFCs	6.7	50.0%	40.0%	64%	6	0.00%	20
		4. Aerosols / MDI	Aerosols	manufacturing stock		HFCs	99.8	—	40.0%	40%	20	0.00%	18
				HFCs	372.0	—	40.0%	40%	20	0.01%	11		
	MDI		manufacturing stock		HFCs	3.1	—	40.0%	40%	20	0.00%	23	
					HFCs	165.2	—	40.0%	40%	20	0.01%	16	
5. Solvents				PFCs	1,376.0	—	40.0%	40%	20	0.05%	5		
7. Semiconductor Manufacture				HFCs	102.2	50.0%	40.0%	64%	6	0.01%	17		
				PFCs	1,818.6	50.0%	40.0%	64%	6	0.10%	4		
				SF ₆	703.9	50.0%	40.0%	64%	6	0.04%	6		
8. Electrical Equipment	manufacturing stock		SF ₆	164.9	30.0%	40.0%	50%	19	0.01%	15			
			SF ₆	487.3	50.0%	40.0%	64%	6	0.03%	7			
9. Other - Railway Silicon Rectifiers				PFCs	0.0	—	40.0%	40%	20	0.00%	24		
Sub Total				23,524.2			33%		0.66%				
Total Emissions			(D)	1,184,802.8			2%						

(Note) Uncertainty of SF₆ emissions from 2.C.4 Magnesium Foundries applies the same value as that of 2.C.3 Aluminium

A7.2.5. Solvents and Other Product Use

Table A7-10 Results of uncertainty assessment of solvent and other product use

IPCC Category			GHGs	Emissions / Removals [Gg-CO ₂ eq.]	EF/RF Uncertainty [%]	AD Uncertainty [%]	Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions	rank
				A	a	b	B		C	
3. Solvent and Other Product Use	D_Other	Anaesthesia	N ₂ O	99.0	—	5.0%	5%	1	0.00%	1
	Sub Total			99.0			5%		0.00%	
Total Emissions			(D)	1,184,802.8			2%			

A7.2.6. Agriculture

Table A7-11 Results of uncertainty assessment of Agriculture

IPCC Category			GHGs	Emissions / Removals [Gg-CO ₂ eq.]	EF/RF Uncertainty [%]	AD Uncertainty [%]	Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions	rank	
				A	a	b	B		C		
4. Agriculture	A. Enteric Fermentation	Dairy Cattle	CH ₄	3,149.0	—	5.0%	15%	63	0.04%	14	
		Non-Dairy Cattle	CH ₄	3,265.3	—	5.0%	19%	62	0.05%	13	
		Buffalo	CH ₄	0.09	50.0%	100.0%	112%	44	0.00%	58	
		Sheep	CH ₄	1.23	50.0%	100.0%	112%	44	0.00%	43	
		Goat	CH ₄	1.19	50.0%	100.0%	112%	44	0.00%	44	
		Swine	CH ₄	225.3	50.0%	0.9%	50%	58	0.01%	20	
		Horse	CH ₄	31.1	50.0%	100.0%	112%	44	0.00%	25	
	B. Manure Management	Dairy Cattle		CH ₄	1,925.051	—	—	78%	54	0.13%	4
				N ₂ O	961.3	—	—	91%	52	0.07%	11
		Non-Dairy Cattle		CH ₄	89.406	—	—	73%	56	0.01%	22
				N ₂ O	1,012.8	—	—	125%	42	0.11%	6
		Buffalo		CH ₄	0.003	100.0%	100.0%	141%	31	0.00%	63
				N ₂ O	0.013	100.0%	100.0%	141%	31	0.00%	62
		Swine		CH ₄	119.872	—	0.9%	106%	48	0.01%	18
				N ₂ O	1,952.8	—	0.9%	92%	51	0.15%	1
		Poultry (Hen, Broiler)		CH ₄	66.998	—	7.3%	54%	57	0.00%	24
				N ₂ O	1,511.0	—	7.3%	80%	53	0.10%	7
		Sheep		CH ₄	0.083	100.0%	100.0%	141%	31	0.00%	56
				N ₂ O	1.4	100.0%	100.0%	141%	31	0.00%	40
		Goat		CH ₄	0.051	100.0%	100.0%	141%	31	0.00%	59
				N ₂ O	5.1	100.0%	100.0%	141%	31	0.00%	31
	Horse		CH ₄	3.597	100.0%	100.0%	141%	31	0.00%	32	
			N ₂ O	30.9	100.0%	100.0%	141%	31	0.00%	23	
	C. Rice Cultivation	Continuously Flooded		CH ₄	190.1	116.3%	0.3%	116%	43	0.02%	17
		Intermittently Flooded	Straw amendment	CH ₄	3,480.7	—	0.3%	32%	61	0.09%	9
			Various compost	CH ₄	1,161.4	—	0.3%	32%	60	0.03%	15
		No-amendment		CH ₄	619.5	—	0.3%	46%	59	0.02%	16
	D. Agricultural Soils	1. Direct Soil Emissions	Synthetic Fertilizers	N ₂ O	1,132.9	—	—	139%	39	0.13%	3
			Animal Waste Applied to Soils	N ₂ O	1,040.1	—	—	152%	30	0.13%	2
			N-Fixing Crops	N ₂ O	78.4	—	—	99%	49	0.01%	21
			Crop residues	N ₂ O	550.5	—	—	211%	16	0.10%	8
			Organic soil	N ₂ O	116.3	—	—	712%	1	0.07%	12
			2. Pasture, Range	N ₂ O	11.8	—	—	133%	40	0.00%	28
3. Indirect Emissions		Atmospheric Deposition	N ₂ O	1,205.9	—	—	75%	55	0.08%	10	
	N Leaching & Run-off	N ₂ O	1,482.9	—	—	97%	50	0.12%	5		
F. Field Burning of Agricultural Residue	1. Cereals	Wheat	CH ₄	5.3	—	—	186%	20	0.00%	30	
			N ₂ O	1.1	—	—	185%	24	0.00%	39	
		Barley	CH ₄	1.3	—	—	185%	22	0.00%	37	
			N ₂ O	0.3	—	—	187%	18	0.00%	48	
		Maize	CH ₄	28.4	418.0%	50.0%	421%	7	0.01%	19	
			N ₂ O	6.0	423.0%	50.0%	426%	3	0.00%	27	
		Oats	CH ₄	0.6	—	—	156%	28	0.00%	46	
			N ₂ O	0.5	—	—	170%	27	0.00%	47	
		Rye	CH ₄	0.029	—	—	130%	41	0.00%	60	
			N ₂ O	0.014	—	—	154%	29	0.00%	61	
		Rice	CH ₄	17.5	178.0%	50.0%	185%	23	0.00%	26	
			N ₂ O	7.1	175.0%	50.0%	182%	26	0.00%	29	
	2. Pulse	Peas	CH ₄	0.08	481.0%	20.0%	481%	2	0.00%	50	
			N ₂ O	0.05	423.0%	20.0%	423%	5	0.00%	54	
		Soybeans	CH ₄	1.58	176.0%	50.0%	183%	25	0.00%	36	
			N ₂ O	1.05	182.0%	50.0%	189%	17	0.00%	41	
		Other (Adzuki beans)	CH ₄	0.24	179.0%	50.0%	186%	21	0.00%	49	
			N ₂ O	0.15	180.0%	50.0%	187%	19	0.00%	52	
		Other (kidney beans)	CH ₄	0.06	418.0%	50.0%	421%	7	0.00%	53	
			N ₂ O	0.04	418.0%	50.0%	421%	7	0.00%	55	
	Other (peanuts)	CH ₄	0.08	418.0%	50.0%	421%	7	0.00%	51		
		N ₂ O	0.03	418.0%	50.0%	421%	7	0.00%	57		
	3. Tuber & Roots	Potatoes	CH ₄	0.4	418.0%	20.0%	418%	15	0.00%	42	
			N ₂ O	0.5	419.0%	20.0%	419%	14	0.00%	38	
Other: Sugarbeet		CH ₄	1.1	417.0%	50.0%	420%	13	0.00%	33		
		N ₂ O	1.0	419.0%	50.0%	422%	6	0.00%	34		
4. Sugar Cane	CH ₄	0.8	418.0%	50.0%	421%	7	0.00%	35			
	N ₂ O	0.3	423.0%	50.0%	426%	3	0.00%	45			
Sub Total				25,499.6			18%		0.39%		
Total Emissions			(D)	1,184,802.8			2%				

A7.2.7. LULUCF

Table A7-12 Results of uncertainty assessment of LULUCF

IPCC Category			GHGs	Emissions / Removals [Gg-CO ₂ eq.]	EF/RF Uncertainty [%]	AD Uncertainty [%]	Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions	rank 5)		
				A	a	b	B		C			
5. LULUCF	A. Forest Land	1. Forest Land remaining Forest Land	CO ₂	-76,372.1	—	—	11%	12	0.71%	1		
		2. Land converted to Forest Land	CO ₂	-304.8	—	—	24%	11	0.01%	8		
	B. Cropland	1. Cropland remaining Cropland	2. Land converted to Cropland	CH ₄	2.1	25.0%	31.2%	40%	6	0.00%	11	
				N ₂ O	0.2	28.6%	31.2%	42%	5	0.00%	12	
		2. Land converted to Cropland	CO ₂	IE,NA,NE,NO	452.4	—	—	—	28%	9	0.01%	5
			CH ₄	NE,NO	—	—	—	—	—	—	—	—
	C. Grassland	1. Grassland remaining Grassland	2. Land converted to Grassland	N ₂ O	6.2	—	—	75%	2	0.00%	10	
			CO ₂	IE,NA,NE	—	—	—	—	—	—	—	—
		2. Land converted to Grassland	CO ₂	—	-215.9	—	—	47%	4	0.01%	7	
			CH ₄	NE,NO	—	—	—	—	—	—	—	—
	D. Wetlands	1. Wetlands remaining Wetlands	2. Land converted to Wetlands	N ₂ O	—	—	—	—	—	—	—	
			CO ₂	NE,NO	82.1	—	—	—	30%	8	0.00%	9
		2. Land converted to Wetlands	CH ₄	NE,NO	—	—	—	—	—	—	—	—
			N ₂ O	NE,NO	—	—	—	—	—	—	—	—
	E. Settlements	1. Settlements remaining Settlements	2. Land converted to Settlements	CO ₂	-1,011.4	—	—	76%	1	0.06%	3	
			CO ₂	—	3,529.7	—	—	30%	7	0.09%	2	
		2. Land converted to Settlements	CH ₄	NE,NO	—	—	—	—	—	—	—	
			N ₂ O	NE,NO	—	—	—	—	—	—	—	
	F. Other Land	1. Other Land remaining Other Land	2. Land converted to Other Land	CO ₂	—	—	—	—	—	—	—	
			CO ₂	—	382.2	—	—	28%	10	0.01%	6	
2. Land converted to Other Land		CH ₄	NO	—	—	—	—	—	—	—		
		N ₂ O	NO	—	—	—	—	—	—	—		
G. Other	CO ₂	CO ₂ , emissions from agricultural lime application	270.1	—	—	51%	3	0.01%	4			
Sub Total				-73,179.1			12%		0.71%			
Total Emissions			(D)	1,184,802.8			2%					

5) Numbers of the rank have been assessed based on the absolute values of “Combined uncertainty as % of total national emissions”.

A7.2.8. Waste

Table A7-13 Results of uncertainty assessment of Waste

IPCC Category			GHGs	Emissions / Removals [Gg-CO ₂ eq.]	EF/RF Uncertainty [%]	AD Uncertainty [%]	Combined Uncertainty [%]	rank	Combined uncertainty as % of total national emissions	rank		
				A	a	b	B		C			
6. Waste	A. Solid Waste Disposal on Land	1. Managed Waste Disposal on Land	Kitchen Garbage	CH ₄	330.6	42.4%	32.4%	53%	30	0.01%	13	
			Waste PAQer	CH ₄	1,452.0	42.4%	42.7%	60%	26	0.07%	6	
			Waste Textile	CH ₄	107.7	43.8%	42.9%	61%	25	0.01%	22	
			Waste Wood	CH ₄	933.5	42.5%	56.6%	71%	21	0.06%	7	
			Digested Sewage Sludge	CH ₄	24.5	44.2%	32.0%	55%	28	0.00%	31	
			Other Sewage Sludge	CH ₄	124.0	44.2%	32.0%	55%	28	0.01%	21	
			Human Waste Sludge	CH ₄	62.9	44.2%	32.6%	55%	27	0.00%	25	
			Water Purification Sludge	CH ₄	26.4	108.6%	31.7%	113%	8	0.00%	27	
			Organic Sludge from Manufacture	CH ₄	154.6	54.0%	33.4%	63%	24	0.01%	18	
			Livestock Waste	CH ₄	20.9	46.9%	49.4%	68%	23	0.00%	29	
	3. Other	CH ₄	41.1	42.5%	66.8%	79%	16	0.00%	26			
	B. Wastewater Handling	1. Industrial Wastewater		CH ₄	107.8	60.0%	37.4%	71%	22	0.01%	20	
				N ₂ O	123.7	300.0%	51.1%	304%	1	0.03%	9	
			2. Domestic and Commercial Wastewater	Sewage Treatment Plant	CH ₄	253.1	30.9%	10.4%	33%	32	0.01%	19
				Private Sewerage Tank	N ₂ O	685.8	145.7%	10.4%	146%	5	0.08%	5
				Human-Waste Treatment Plant	CH ₄	425.5	86.8%	10.0%	87%	14	0.03%	10
		Degradation of domestic wastewater in nature	N ₂ O	270.9	71.0%	10.0%	72%	20	0.02%	12		
			CH ₄	13.8	100.0%	12.5%	101%	11	0.00%	30		
			N ₂ O	5.8	100.0%	33.9%	106%	9	0.00%	33		
			CH ₄	469.5	—	—	76%	17	0.03%	11		
		N ₂ O	45.4	—	—	76%	17	0.00%	24			
C. Waste Incineration	Municipal Solid Waste	Plastics	CO ₂	2,617.0	4.3%	16.0%	17%	35	0.04%	8		
		Waste textile	CO ₂	600.3	4.3%	22.4%	23%	34	0.01%	14		
			CH ₄	1.3	—	—	101%	12	0.00%	35		
			N ₂ O	150.3	—	—	42%	31	0.01%	23		
	Industrial Solid Waste	Waste mineral oil	CO ₂	4,012.6	4.8%	104.4%	105%	10	0.35%	1		
		Plastics	CO ₂	3,629.9	4.8%	100.0%	100%	13	0.31%	2		
			CH ₄	8.2	111.5%	100.0%	150%	4	0.00%	32		
			N ₂ O	1,522.6	58.8%	100.0%	116%	7	0.15%	4		
	Specially Controlled Industrial Solid Waste		CO ₂	1,797.9	—	—	167%	2	0.25%	3		
			CH ₄	1.0	—	—	142%	6	0.00%	34		
		N ₂ O	14.7	—	—	159%	3	0.00%	28			
D. Other	Decomposition of petroleum-derived surface-active agent	CO ₂	528.5	—	—	25%	33	0.01%	15			
	Composting of Organic Waste	CH ₄	168.8	—	—	74%	19	0.01%	17			
		N ₂ O	149.5	—	—	86%	15	0.01%	16			
Sub Total				20,873.8			32%		0.57%			
Total Emissions			(D)	1,184,802.8			2%					

- 6) Regarding 6A1, uncertainty of “Anaerobic landfill”, which is the largest source under this sub-category, has been used.
- 7) Regarding 6A2, uncertainty of “Gappei-shori johkasou”, which is the largest source under this sub-category, has been used.
- 8) Regarding CH₄ of 6C MSW, uncertainty of “Semi-Continuous Incinerator” has been used.
- 9) Regarding CH₄ of 6C ISW, uncertainty of “Waste Paper and Waste Wood” has been used.
- 10) Regarding N₂O of 6C ISW, uncertainty of “Waste Plastics” has been used.
- 11) Regarding 6C Fuel use of RDF and RPF, uncertainty of “RDF” has been used.

A7.2.9. Consideration of the results

The result of uncertainty assessment shows that Japan's uncertainty of total net emissions is approximately 2%. This value is relatively smaller compared to 21.3% of UK indicated in the *Good Practice Guidance*. It is attributed to the fact that the ratio of Japan's N₂O emission from "4.D.1. Agricultural Soils (Direct Soil Emissions)" to the national total emissions is small compared to that of UK (the ratios of Japan and UK reported in their inventories submitted in 2003 were 0.28% and 4.1%, respectively).

Below are the results of sensitivity analysis with N₂O emissions from this source, uncertainty of emission factor and national total emissions (calculation used the reported values of inventories submitted in 2003).

Table A7-14 Sensitivity Analysis on N₂O emissions from "4.D. Agricultural Soils 1 Direct Emissions"

	N ₂ O Emissions [Gg-CO ₂ eq.]	Uncertainty of EF	Uncertainty of Total Emissions	Note
Original	3,597.58	129.9%	2.4%	2001 Emissions contained in the GHG inventory submitted in 2003.
Case 1	3,597.58	500%	2.6%	EF uncertainty was assumed to be same as UK's case.
Case 2	71,951.53	129.9%	4.8%	Emissions were assumed to be approximately 5% of national total emissions in 2001.

A7.2.10. Issues in Uncertainty Assessment

- According to the method indicated in the *Revised 1996 IPCC Guidelines*, only emission sources of which emissions had already been calculated were the subject of uncertainty assessment. No assessment has been made for emission sources not estimated (NE), or of those portions unconfirmed in emission sources for which only partial calculation has been done (PART). Therefore, it should be remembered that the uncertainty of total emissions prepared by compiling the uncertainty of emissions from each source, does not depict the uncertainty of inventory in the context of the realities of emissions.
- In the sources recalculated, consideration is needed whether to re-assess the uncertainties or not.
- Where it was not possible to carry out a statistical assessment of the uncertainty of activity data, the values were derived from those established by the Committee for the GHGs Emissions Estimations Methods, which have established the uncertainty values in relation to whether the data were derived from specified statistics, or whether they were obtained from total population surveys. But further consideration needs to be given to improve the appropriateness of this approach.
- In carrying out a statistical assessment of uncertainty, it was assumed that the averages of all samples followed a normal distribution. In some cases, however, it means that the emission factor or activity data could, in fact, be negative. Emissions can only be positive under the present IPCC guidelines, so further consideration would need to be given for the possibility to assume that the emission factor or activity data follows some other distribution.
- Consideration on application of probability density function (PDF) with Monte-Carlo analysis is further issue. Further consideration on analysis with more disaggregated sources or each coefficients are needed.

- The number of decimal places to be used when depicting uncertainty was set as follows for the uncertainty assessments conducted, but as the precision of uncertainty assessment varies between emission sources, further consideration needs to be given to the number of decimal places that are effective in uncertainty assessment.
 - 1) Uncertainty of emission factor is given to one decimal place.
 - 2) Uncertainty of activity data is also given to one decimal place.
 - 3) Uncertainty of emissions is given as an integer. (Proportion of total emissions attributable to the uncertainty of a particular source = two decimal places.)

A7.2.11. Reference Material

Results of the uncertainty assessment for this year in accordance with Table 6.1 of *GPG (2000)* are indicated below.

Table 6.1 Tier 1 Uncertainty Calculation & Reporting																							
A IPCC Source Category	B Gas	C		D		E		F		G		H		I		J		K		L		M	
		Base year emissions / removals		2010 emissions / removals		Activity Data		EForRF Uncertainty		Combined Uncertainty		Combined Uncertainty as % of Total National Emissions in 2010		Type A Sensitivity		Type B Sensitivity		Uncertainty in trend in National Emissions introduced by EForRF Uncertainty		Uncertainty in trend in National Emissions introduced by Activity Data Uncertainty		Uncertainty introduced into the Total National Emissions	
		Input Data		Input Data		Input Data		Input Data		(E/F)²/(G)²/(H)²		O'D/D / H²		Note B		D/E/C		F/F Note C		F/E'/2		(K/L)²/(M)²	
		Gg CO ₂ equivalent		Gg CO ₂ equivalent		%		%		%		%		%		%		%		%		%	
Total			1,186,712.60	1,184,804.18								2%	0.0%									1%	
2. Industrial Processes	A. Mineral Products	1. Cement Production	CO ₂	37,904.87	23,784.44	10.0%	3.0%	10%	0.2%	0.0%	-1.2%	2.0%	0.0%	0.3%	0.0%	0.3%	0.0%	0.3%	0.0%	0.3%	0.0%	0.3%	
		2. Lime Production	CO ₂	6,674.45	6,284.59	5.0%	15.0%	16%	0.1%	0.0%	0.5%	0.0%	0.5%	0.0%	0.0%	0.5%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		3. Limestone & Dolomite Use	CO ₂	9,054.75	6,483.72	4.8%	16.4%	17%	0.1%	0.0%	-0.2%	0.5%	0.0%	0.0%	0.0%	0.5%	0.0%	0.0%	0.0%	0.0%	0.0%	0.1%	
		4. Soda Ash Production and Use	CO ₂	1,467.80	1,589.50	3.9%	3.5%	5%	0.0%	0.0%	0.0%	0.1%	0.0%	0.0%	0.0%	0.1%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
	B. Chemical Industries	1. Ammonia Production	CO ₂	3,384.68	2,106.42	5.0%	22.5%	23%	0.0%	0.0%	-0.1%	0.2%	0.0%	0.0%	0.0%	0.2%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		Chemical Industries other than Nitric Acid	CO ₂	824.39	630.81	5.0%	77.2%	77%	0.0%	0.0%	0.0%	0.1%	0.0%	0.0%	0.0%	0.1%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		2. Nitric Acid	N ₂ O	765.70	561.64	5.0%	46.0%	46%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		3. Adipic Acid	N ₂ O	7,501.25	516.10	2.0%	9.0%	9%	0.0%	0.0%	-0.6%	0.0%	0.0%	0.0%	-0.1%	0.0%	0.0%	-0.1%	0.0%	0.0%	0.0%	0.1%	
		4. Carbide	CH ₄	0.42	0.66	10.0%	100.0%	100%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		5. Other	CH ₄	5.83	5.37	5.0%	54.8%	55%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		Ethylene	CH ₄	1.88	2.20	5.0%	77.2%	77%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		Dichloroethylene	CH ₄	0.28	0.33	5.0%	100.7%	101%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		Styrene	CH ₄	1.45	1.97	5.0%	113.2%	113%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		Methanol	CH ₄	3.52	0.00	5.0%	113.2%	113%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
	Coke	CH ₄	324.84	93.45	5.0%	98.5%	99%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
	C. Metal Production	1. Iron and steel	CO ₂	356.09	159.86	4.5%	0.0%	5%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		2. Ferroalloy	CH ₄	3.89	2.56	5.0%	163.0%	163%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
	C. Metal Production	3. Aluminum	PPCs	69.74	10.38	5.0%	33.0%	33%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		4. SP6 Used in Aluminium and Magnesium Foundries	SP	119.50	307.90	5.0%	0.0%	5%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
	E. Production of F-gases	1. By-product Emissions (HCFC-22)	HCFCs	16,965.30	42,112	5.0%	2.9%	4%	0.0%	0.0%	-1.4%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
2. Fugitive Emissions		HCFCs	480.12	86.22	10.0%	100.0%	100%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
2. Industrial Processes (F-gas)	F. Consumption of F-gases	1. Refrigeration and Air Conditioning Equipment	Domestic Refrigerator	manufacturing stock	HFCs	11.34	359.15	40.0%	50.0%	64%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
			disposal	HFCs	0.00	0.00	40.0%	50.0%	64%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		Commercial Refrigerator	manufacturing stock	HFCs	42.48	11,336.43	40.0%	50.0%	64%	0.0%	0.0%	0.0%	1.0%	0.5%	0.5%	0.7%	0.0%	0.0%	0.0%	0.0%	0.0%		
		disposal	HFCs	0.00	0.00	40.0%	50.0%	64%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		Stationary Air-Conditioning	manufacturing stock	HFCs	0.00	2,890.19	40.0%	50.0%	64%	0.2%	0.0%	0.2%	0.2%	0.1%	0.1%	0.2%	0.0%	0.0%	0.0%	0.0%	0.0%		
		disposal	HFCs	0.00	0.00	40.0%	50.0%	64%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		Mobile Air-Conditioning	manufacturing stock	HFCs	786.58	2,502.43	40.0%	50.0%	64%	0.1%	0.0%	0.1%	0.2%	0.1%	0.1%	0.1%	0.1%	0.1%	0.1%	0.1%	0.1%		
		disposal	HFCs	0.00	0.00	40.0%	50.0%	64%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		2. Foam Blowing	manufacturing stock	HFCs	451.76	138.58	50.0%	50.0%	71%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
		disposal	HFCs	0.00	152.39	50.0%	50.0%	71%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		3. Fire Extinguisher	manufacturing stock	HFCs	0.00	6.72	40.0%	50.0%	64%	0.0%	0.0%	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA		
		disposal	HFCs	0.00	99.82	40.0%	50.0%	64%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		4. Aerosols / MDI	manufacturing stock	HFCs	1,365.00	371.99	40.0%	0.0%	40%	0.0%	0.0%	-0.1%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		disposal	HFCs	0.00	3.07	40.0%	0.0%	40%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		MDI	manufacturing stock	HFCs	0.00	165.21	40.0%	0.0%	40%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		disposal	HFCs	0.00	0.00	40.0%	0.0%	40%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		5. Solvents	PPCs	10,263.55	1,375.99	40.0%	0.0%	40%	0.0%	0.0%	-0.7%	0.1%	0.0%	0.0%	0.1%	0.0%	0.1%	0.1%	0.1%	0.1%			
		7. Semiconductor Manufacture	PPCs	3,144.23	102.19	40.0%	50.0%	64%	0.0%	0.0%	-0.3%	0.0%	0.0%	-0.1%	0.0%	-0.1%	0.0%	0.1%	0.1%				
		disposal	PPCs	157.89	1,838.65	40.0%	50.0%	64%	0.1%	0.2%	0.1%	0.2%	0.1%	0.1%	0.1%	0.1%	0.1%	0.1%	0.1%				
		SP	1,128.66	703.91	40.0%	50.0%	64%	0.0%	0.0%	0.0%	0.1%	0.0%	0.1%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%				
8. Electrical Equipment	manufacturing stock	SP	9,560.00	164.91	40.0%	30.0%	50%	0.0%	0.0%	-0.8%	0.0%	-0.2%	0.0%	-0.2%	0.0%	0.2%	0.2%						
disposal	SP	1,444.99	487.34	40.0%	50.0%	64%	0.0%	0.0%	-0.1%	0.0%	0.0%	-0.1%	0.0%	0.0%	0.0%	0.0%							
9. Other - Railway Silicon Rectifiers	PPCs	0.00	0.00	40.0%	0.0%	40%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%							
3. SOFU	D. Other	Anaesthesia	N ₂ O	287.07	98.95	5.0%	0.0%	5%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%						
4. Agriculture	A. Enteric Fermentation	Dairy Cattle	CH ₄	4,044.60	3,149.00	5.0%	14.2%	15%	0.0%	0.0%	-0.1%	0.3%	0.0%	0.0%	0.3%	0.0%	0.0%						
		Non-Dairy Cattle	CH ₄	3,322.55	3,265.30	5.0%	18.0%	19%	0.1%	0.0%	0.0%	0.3%	0.0%	0.0%	0.3%	0.0%							
		Buffalo	CH ₄	0.25	0.09	100.0%	50.0%	112%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%							
		Sheep	CH ₄	1.88	1.23	100.0%	50.0%	112%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%								
		Goat	CH ₄	2.22	1.19	100.0%	50.0%	112%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%								
		Swine	CH ₄	261.75	225.33	0.9%	50.0%	50%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%								
		Horses	CH ₄	43.37	31.13	100.0%	50.0%	112%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%								
		Dairy Cattle	CH ₄	2,387.79	1,925.05	10.0%	77.0%	78%	0.1%	0.0%	-0.1%	0.2%	0.0%	0.0%	0.0%								
		Non-Dairy Cattle	N ₂ O	840.93	961.30	10.0%	90.1%	91%	0.1%	0.0%	0.0%	0.1%	0.0%	0.0%	0.0%								
		Buffalo	N ₂ O	89.12	1,012.80	10.0%	125.1%	125%	0.1%	0.0%	0.0%	0.1%	0.0%	0.0%	0.0%								
	Swine	CH ₄	0.01	0.00	100.0%	100.0%	141%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%									
	N ₂ O	0.04	0.01	100.0%	100.0%	141%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%										
	Goat	CH ₄	333.44	119.87	0.9%	100%	100%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%										
	Sheep	CH ₄	1,479.89	1,952.78	0.9%	91.6%	92%	0.2%	0.0%	0.0%	0.2%	0.0%	0.0%										
	Poultry (Hen, Broiler)	CH ₄	73.82	67.00	7.3%	53.1%	54%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%										
	N ₂ O	2,288.25	1,511.03	7.3%	79.2%	80%	0.1%	0.0%	-0.1%	0.1%	-0.1%	0.0%	0.1%										
	Goat	CH ₄	0.13	0.08	100.0%	100.0%	141%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%										

Annex 7. Methodology and Results of Uncertainty Assessment

Table 6.1 Tier 1 Uncertainty Calculation & Reporting																							
A IPCC Source Category	B Gas	C		D		E		F		G		H		I		J		K		L		M	
		Base year emissions / removals		2010 emissions / removals		Activity Data Uncertainty		EForRF Uncertainty		Combined Uncertainty		Combined Uncertainty as % of Total National Emissions in 2010		Type A Sensitivity		Type B Sensitivity		Uncertainty in trend in National Emissions introduced by EForRF Uncertainty		Uncertainty in trend in National Emissions introduced by Activity Data Uncertainty		Uncertainty introduced into the Trend in Total National Emissions	
		Input Data		Input Data		Input Data		Input Data		(F2)-(F3)/(F2)		(G)-(H)/(G)		Note B		D-E/C		FF Note C		FF1-2		(M2)-(M3)/(M2)	
Gg CO ₂ equivalent		Gg CO ₂ equivalent		%		%		%		%		%		%		%		%		%		%	
Total		1,186,712.60		1,184,804.18								2%		0.0%								1%	
5. LULUCF	A. Forest Land	1. Forest Land remaining Forest Land	CO ₂	-76,762.09	-76,372.11	-	-	-	-	11%	-0.7%	0.0%	0.0%	-6.4%	-	-	-	-	-	-	-	-	-
		2. Land converted to Forest Land	CO ₂	-1,830.25	-304.80	-	-	-	-	24%	0.0%	0.0%	0.1%	0.0%	0.0%	-	-	-	-	-	-	-	-
	B. Cropland	1. Cropland remaining Cropland	CO ₂	2,513.21	452.41	-	-	-	-	28%	0.0%	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA
			CH ₄	8.51	2.12	31.2%	25.0%	-	-	40%	0.0%	0.0%	0.0%	0.0%	0.0%	-	-	-	-	-	-	-	0.0%
		2. Land converted to Cropland	N ₂ O	0.86	0.22	31.2%	28.6%	-	-	42%	0.0%	0.0%	0.0%	0.0%	0.0%	-	-	-	-	-	-	-	0.0%
	C. Grassland	1. Grassland remaining Grassland	CO ₂	IE,NA,NE	IE,NA,NE	-	-	-	-	-	0.0%	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA
			CO ₂	-444.03	-215.86	-	-	-	-	47%	0.0%	0.0%	0.0%	0.0%	0.0%	-	-	-	-	-	-	-	0.0%
2. Land converted to Grassland		CH ₄	NE,NO	NE,NO	-	-	-	-	-	0.0%	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	
D. Wetlands	1. Wetlands remaining Wetlands	CO ₂	NE,NO	NE,NO	-	-	-	-	-	0.0%	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	
		CO ₂	85.84	82.13	-	-	-	-	30%	0.0%	0.0%	0.0%	0.0%	-	-	-	-	-	-	-	-	0.0%	
	2. Land converted to Wetlands	CH ₄	NE,NO	NE,NO	-	-	-	-	-	0.0%	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	
E. Settlements	1. Settlements remaining Settlements	CO ₂	-955.53	-1,011.43	-	-	-	-	76%	-0.1%	0.0%	0.0%	-0.1%	-	-	-	-	-	-	-	-	-	
		CO ₂	5,113.88	3,529.72	-	-	-	-	30%	0.1%	0.0%	-0.1%	0.3%	-	-	-	-	-	-	-	-	0.0%	
	2. Land converted to Settlements	CH ₄	NE,NO	NE,NO	-	-	-	-	-	0.0%	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	
F. Other Land	1. Other Land remaining Other Land	CO ₂	NE,NO	NE,NO	-	-	-	-	-	0.0%	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	
		CO ₂	1,553.92	382.22	-	-	-	-	28%	0.0%	0.0%	-0.1%	0.0%	-	-	-	-	-	-	-	-	0.0%	
	2. Land converted to Other Land	CH ₄	NO	NO	-	-	-	-	-	0.0%	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	0.0%	NA	NA	
G. Other	CO ₂ emissions from agricultural lime application	CO ₂	550.22	270.12	-	-	-	-	51%	0.0%	0.0%	0.0%	0.0%	-	-	-	-	-	-	-	-	-	
		CH ₄	1,319.91	350.64	32.4%	42.4%	-	-	55%	0.0%	0.0%	-0.1%	0.0%	0.0%	-	-	-	-	-	-	-	0.0%	
	3. Other	CH ₄	7.29	41.05	66.8%	42.5%	79%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
6. Waste	A. Solid Waste Disposal on Land	1. Managed Waste Disposal on Land	Kitchen Garbage	CH ₄	3,096.96	1,452.05	42.7%	42.4%	60%	0.1%	0.0%	-0.1%	0.1%	-0.1%	0.1%	-0.1%	0.1%	0.0%	0.0%	0.0%	0.1%	0.1%	
			Waste Paper	CH ₄	202.64	107.72	42.9%	43.8%	61%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%
			Waste Textile	CH ₄	973.45	933.53	56.6%	42.5%	71%	0.1%	0.0%	0.0%	0.1%	0.0%	0.1%	0.0%	0.1%	0.0%	0.1%	0.0%	0.1%	0.0%	
			Waste Wood	CH ₄	115.01	24.55	32.0%	44.2%	55%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%
			Sewage Sludge	CH ₄	573.33	124.00	32.0%	44.2%	55%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%
			Other Sewage Sludge	CH ₄	260.92	62.87	32.6%	44.2%	55%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%
			Human Waste Sludge	CH ₄	70.40	26.40	31.7%	108.6%	113%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%
			Water Purification Sludge	CH ₄	1,012.28	154.61	33.4%	54.0%	68%	0.0%	0.0%	-0.1%	0.0%	-0.1%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%
			Organic Sludge from Manufacture	CH ₄	28.75	20.93	49.4%	46.9%	68%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%
			Illegal Disposal	CH ₄	7.29	41.05	66.8%	42.5%	79%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%
B. Wastewater Handling	1. Industrial Wastewater	CH ₄	135.76	107.78	37.4%	60.0%	71%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		N ₂ O	127.81	123.74	51.1%	300.0%	304%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
	2. Domestic and Commercial Wastewater	Sewage Treatment Plant	CH ₄	181.48	253.08	10.4%	30.9%	33%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		Private Sewerage Tank	CH ₄	491.78	685.52	10.4%	145.7%	146%	0.1%	0.0%	0.0%	0.1%	0.0%	0.0%	0.1%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		Human Waste Treatment Plant	CH ₄	451.84	425.51	10.0%	86.8%	87%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		Human Waste Treatment Plant	CH ₄	468.72	270.93	10.0%	71.0%	72%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		Degradation of domestic wastewater in nature	CH ₄	110.14	13.77	12.3%	100.0%	101%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		Degradation of domestic wastewater in nature	CH ₄	1,264.60	469.50	10.0%	75.4%	76%	0.0%	0.0%	0.0%	-0.1%	0.0%	-0.1%	0.0%	-0.1%	0.0%	0.0%	0.0%	0.0%	0.0%		
		Degradation of domestic wastewater in nature	N ₂ O	137.38	45.38	10.0%	75.4%	76%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
		Degradation of domestic wastewater in nature	CO ₂	5,040.90	2,616.95	16.0%	4.3%	17%	0.0%	0.0%	-0.2%	0.2%	0.0%	0.2%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.1%		
C. Waste Incineration	Municipal Solid Waste	Plastics	CO ₂	803.19	600.28	22.4%	4.3%	28%	0.0%	0.0%	0.0%	0.0%	0.1%	0.0%	0.1%	0.0%	0.1%	0.0%	0.0%	0.0%			
		Waste textile	CH ₄	9.75	1.27	10.0%	100.2%	101%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%			
	Industrial Solid Waste	Plastics	CO ₂	317.82	150.29	10.0%	40.6%	42%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%			
		Waste Mineral Oil	CO ₂	3,651.84	4,012.55	104.4%	4.8%	105%	0.4%	0.0%	0.0%	0.3%	0.0%	0.3%	0.0%	0.3%	0.0%	0.5%	0.0%	0.5%			
		Plastics	CO ₂	2,120.24	3,629.87	100.0%	4.8%	100%	0.3%	0.0%	0.1%	0.3%	0.0%	0.3%	0.0%	0.4%	0.0%	0.4%	0.0%				
		Plastics	CH ₄	3.60	8.21	100.0%	111.5%	150%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%			
		Plastics	N ₂ O	1,095.67	1,522.58	100.0%	58.8%	116%	0.1%	0.0%	0.0%	0.1%	0.0%	0.1%	0.0%	0.2%	0.0%	0.2%	0.0%				
		Specially Controlled Industrial Solid Waste	CO ₂	946.78	1,797.90	100.0%	133.1%	167%	0.3%	0.0%	0.1%	0.2%	0.1%	0.2%	0.1%	0.2%	0.0%	0.2%	0.0%				
		Specially Controlled Industrial Solid Waste	CH ₄	0.12	0.98	100.0%	100.3%	142%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%				
		Specially Controlled Industrial Solid Waste	N ₂ O	5.95	14.72	100.0%	123.2%	159%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%				
D. Other	Decomposition of petroleum-derived surface-active agent	CO ₂	702.83	528.50	10.0%	22.4%	25%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%					
		CH ₄	111.85	168.82	10.0%	73.3%	74%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%						
		N ₂ O	99.06	149.53	10.0%	85.7%	86%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%						

Annex 8. Hierarchical Structure of Japan's National GHG Inventory File System

Multiple MS Excel files have been used when estimating Japanese inventory. The explanation of each MS Excel file and the hierarchical structure of Japanese National GHGs Inventory (JNGI) file system are shown below.

Table A 8-1 Explanation of each MS Excel file

category	file name	contents
	JPN-2012-1990-v1.1.xls ~ JPN-2012-2010-v1.1.xls, KP-JPN-2012-1990-v1.1.xls, KP-JPN-2012-2008-v1.1.xls ~ KP-JPN-2012-2010-v1.1.xls	Common reporting format provided by UNFCCC secretariat
1. Energy	1A-L3-nonCO2-1990-2012.xls ~ 1A-L3-nonCO2-2010-2012.xls	Non-CO ₂ emissions from stationary facilities
	1A-L3-CO2-1990-2012.xls ~ 1A-L3-CO2-2010-2012.xls	CO ₂ emissions from fuel combustions
	1A-L3-NOxSO2-2012.xls	Emissions of Non-CO ₂ from stationary combustion
	1A-L3-CRF-2012.xls	CRF format data of GHG emissions from fuel combustion (including emissions by energy use of waste)
	1A-L3-timeseries-2012.xls	Time-series data of GHG emissions from fuel combustion
	1A-L2-MAP-IEF-1990-2012.xls ~ 1A-L2-MAP-IEF-2010-2012.xls	Implied Emission Factors of Non-CO ₂ from stationary combustion
	1A-L2-nonCO2-ADEF-2012.xls	Activity Data and Emission Factors of Non-CO ₂ from fuel combustion
	1A-L2-nonCO2-EF-2012.xls	Emission Factors of Non-CO ₂ from fuel combustion
	1A-L2-EBEF-2012.xls	Emission Factors for CO ₂ from fuel combustion
	1A-L1-EB-2012.xls	Data of the General Energy Statistics using in Mobile (CH ₄ , N ₂ O), Fugitive emissions from fuels and IP sector
	1A3-L3-CH4N2O-2012.xls	GHG emissions from Mobile Combustion (transport sector) (except CO ₂)
	1A3-L2-ADEF-2012.xls	Activity Data and Emission Factors for Mobile Combustion (transport sector)
	1B-L3-2012.xls	Fugitive GHG emissions from fuels
	1B-L2-ADEF-2012.xls	Activity Data and Emission Factors for Fugitive Emissions from Fuels
2. Industrial Processes	2-L2-ADEF-2012.xls	Activity Data and Emission Factors of Caotegory2 (except F-gas)
	2-L3-2012.xls	GHG emissions from Category2 (Industrial Processes)
	2-L3-Fgas-2012.xls	F-gas (HFCs, PFCs, SF ₆) emissions
3. Solvent and Other Product Use	3-L3-2012.xls	N ₂ O emissions from anesthesia
4. Agriculture	4A-L3-CH4-2012.xls	CH ₄ emissions from enteric fermentation
	4B-L3-CH4N2O-2012.xls	GHG emissions from manure management
	4C-L3-CH4-2012.xls	CH ₄ emissions from rice cultivation
	4D-L3-N2O-2012.xls	N ₂ O emissions from agricultural soils
	4F-CH4N2OCO-2012.xls	GHG emissions from field burning of agricultural residues
	4-L2-ADEF-2012.xls	Activity Data and Emission Factors of Caotegory4
5. LULUCF	5-L3-nonCSC-2012.xls	GHG emissions excluding carbon stock change
	5A-L3-CO2-2012.xls	CO ₂ emissions and removals from forest land
	5B-L3-CO2-2012.xls	CO ₂ emissions and removals from cropland
	5C-L3-CO2-2012.xls	CO ₂ emissions and removals from grassland
	5D-L3-CO2-2012.xls	CO ₂ emissions and removals from wetlands
	5E-L3-CO2-2012.xls	CO ₂ emissions and removals from settlements
	5F-L3-CO2-2012.xls	CO ₂ emissions and removals from other land
	5-L2-LandArea-2012.xls	Land area for each land use category
	5-L2-Parameter-2012.xls	Parameters for each land use category
	5-L2-nonCSC-2012.xls	Activity data for GHG emissions excluding carbon stock change
6. Waste	6A3-L2-AD-2012.xls	Activity data of solid waste disposal on land (other)
	6A-L3-2012.xls	GHGs emissions from solid waste disposal on land
	6A-L2-AD-2012.xls	Activity data of solid waste disposal on land
	6B-L3-2012.xls	GHGs emissions from wastewater handling
	6B-L2-AD-2012.xls	Activity data of wastewater handling
	6B-L2-EF-2012.xls	Emission Factor of wastewater handling
	6C-L3-nonCO2-2012.xls	GHGs emissions from waste incineration (exclude CO ₂)
	6C-L2-AD-2012.xls	Activity data of waste incineration
	6C-L3-CO2-2012.xls	CO ₂ emissions from waste incineration
	6C-L3-Energy-2012.xls	GHGs (CO ₂ , CH ₄ , N ₂ O, CO, NO _x , SO _x , NMVOC) Emissions from the incineration of waste for energy and use as alternative fuels
	6D-L3-2012.xls	GHGs emissions from other waste
	6D-L2-2012.xls	Activity data of other waste
7. Other	7-L3-2012.xls	CO Emissions from tobaccos
Memo Item	1C-L3-bunker-2012.xls	GHGs emissions from bunker fuels
KP-LULUCF	KP-3-RV-CRPreporter-2012.xls	GHG emissions and removals from Revetation
	KP-3-Summary-2012.xls	GHG emissions and removals from KP3.3 and 3.4 activities
	KP-2-AR-2012.xls	GHG emissions and removals from Afforestation/Reforestation
	KP-2-D-2012.xls	GHG emissions and removals from Deforestation
	KP-2-FM-2012.xls	GHG emissions and removals from Forest Management
	KP-2-RV-2012.xls	GHG emissions and removals from Revegetation

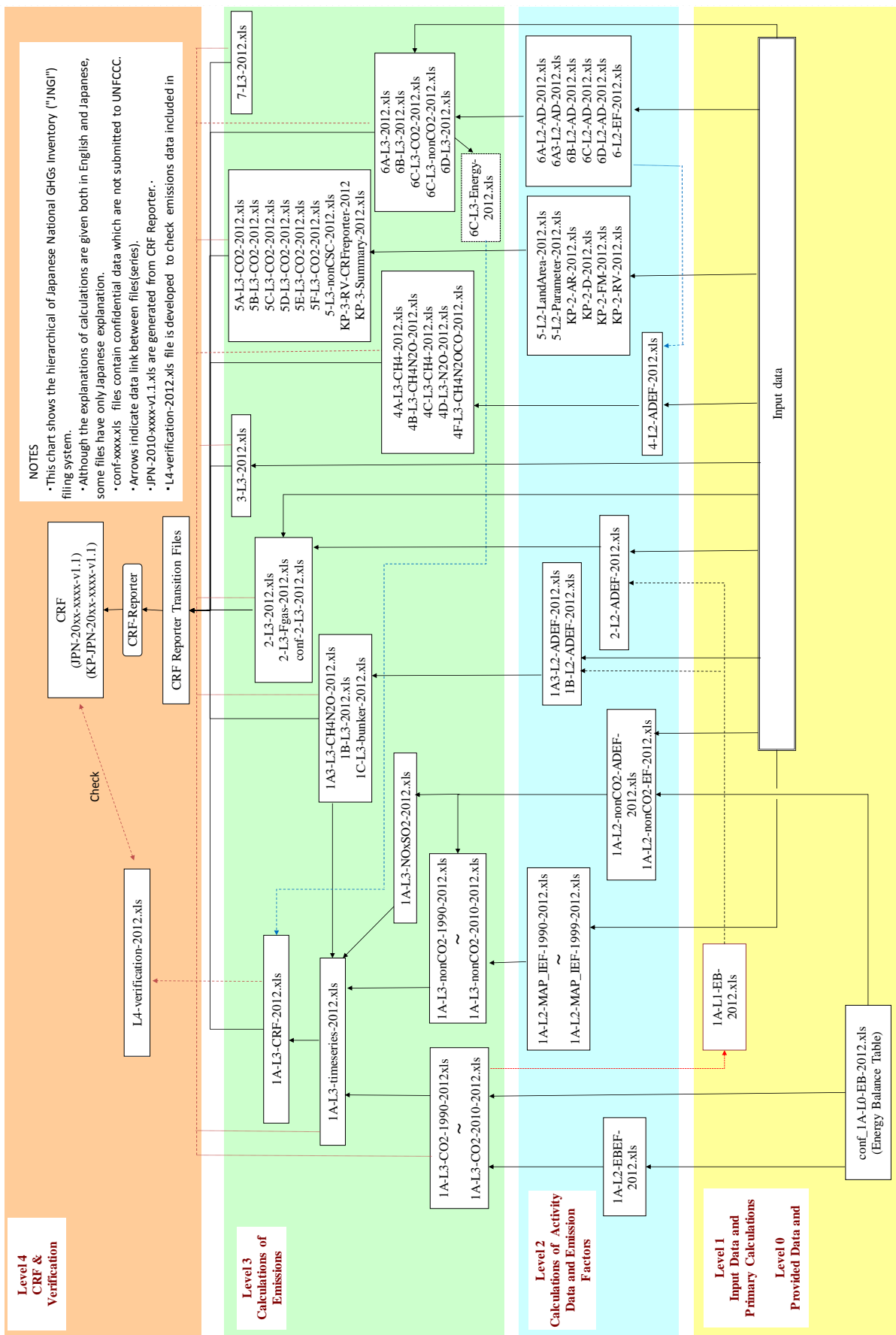


Figure A 8-1 Hierarchical structure of Japan's National GHG Inventory File System

Annex 9. Summary of Common Reporting Format

“Summary.2 Table” of the CRF indicated below shows emissions and removals for every year.

During 1990-1994, Japan had reported only potential emissions of HFCs, PFCs, and SF₆. In Table.10 of the CRF showing the trend each year, between 1990 and 1994, the potential emissions of HFCs, PFCs, and SF₆ are shown, and from 1995 onward, actual emissions of HFCs, PFCs, SF₆ are shown.

A9.1. Emissions¹ and Removals in 1990

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS (Sheet 1 of 1)

Inventory 1990
Submission 2012 v1.1
JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,071,021.21	32,038.69	31,740.24	17,930.00	5,670.00	38,240.00	1,196,640.14
1. Energy	1,068,296.26	3,927.31	6,751.65				1,078,975.22
A. Fuel Combustion (Sectoral Approach)	1,068,259.64	890.17	6,751.54				1,075,901.34
1. Energy Industries	324,253.21	29.72	922.25				325,205.19
2. Manufacturing Industries and Construction	371,311.49	355.49	1,350.07				373,017.05
3. Transport	211,053.69	297.54	4,205.72				215,556.95
4. Other Sectors	161,641.24	207.42	273.50				162,122.16
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	36.62	3,037.14	0.11				3,073.88
1. Solid Fuels	NE,NO	2,806.43	NE,NO				2,806.43
2. Oil and Natural Gas	36.62	230.71	0.11				267.45
2. Industrial Processes	59,934.01	357.58	8,266.95	17,930.00	5,670.00	38,240.00	130,398.54
A. Mineral Products	55,368.85	NA,NO	NA,NO				55,368.85
B. Chemical Industry	4,209.07	338.22	8,266.95	NA	NA	NA	12,814.24
C. Metal Production	356.09	19.36	NO	IE,NE	IE,NA,NE	IE,NA,NE	375.45
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				NE,NO	NE,NO	NE,NO	NE,NO
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				17,930.00	5,670.00	38,240.00	61,840.00
G. Other	NO	NO	NO	NE,NO	NE,NO	NE,NO	NE,NO
3. Solvent and Other Product Use	NA,NE		287.07				287.07
4. Agriculture		17,831.10	13,429.93				31,261.03
A. Enteric Fermentation		7,676.61					7,676.61
B. Manure Management		3,094.12	5,533.01				8,627.13
C. Rice Cultivation		6,959.68					6,959.68
D. Agricultural Soils ⁽³⁾		NA	7,864.27				7,864.27
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		100.68	32.65				133.32
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-70,174.84	8.51	90.88				-70,075.44
A. Forest Land	-78,592.34	8.51	0.86				-78,582.97
B. Cropland	2,513.21	NE,NO	90.02				2,603.23
C. Grassland	-444.03	NE,NO	NE,NO				-444.03
D. Wetlands	85.84	NE,NO	NE,NO				85.84
E. Settlements	4,158.35	NE,NO	NE,NO				4,158.35
F. Other Land	1,553.92	NO	NO				1,553.92
G. Other	550.22	NA,NE	NA,NE				550.22
6. Waste	12,965.78	9,914.19	2,913.75				25,793.73
A. Solid Waste Disposal on Land	NA,NE,NO	7,645.06					7,645.06
B. Waste-water Handling		2,143.81	1,295.25				3,439.06
C. Waste Incineration	12,262.95	13.48	1,519.44				13,795.87
D. Other	702.83	111.85	99.06				913.74
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	30,829.18	43.15	279.35				31,151.68
Aviation	13,189.32	7.84	130.44				13,327.60
Marine	17,639.86	35.31	148.92				17,824.08
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	18,747.30						18,747.30
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,266,715.59
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,196,640.14

⁽¹⁾ For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

⁽²⁾ Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

⁽³⁾ Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

⁽⁴⁾ See footnote 8 to table Summary I.A.

¹ Potential emissions of HFCs, PFCs and SF₆ are reported due to the generation of CRF Reporter

A9.2. Emissions² and Removals in 1991SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS
(Sheet 1 of 1)Inventory 1991
Submission 2012 v1.1
JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,072,690.10	31,792.66	31,220.34	18,070.00	6,370.00	43,498.00	1,203,641.10
I. Energy	1,076,104.87	3,691.08	7,030.93				1,086,826.88
A. Fuel Combustion (Sectoral Approach)	1,076,051.20	896.32	7,030.77				1,083,978.29
1. Energy Industries	326,986.60	31.16	955.50				327,973.27
2. Manufacturing Industries and Construction	366,282.86	356.15	1,426.53				368,065.53
3. Transport	222,466.79	299.95	4,368.84				227,135.59
4. Other Sectors	160,314.95	209.06	279.89				160,803.90
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	53.67	2,794.76	0.16				2,848.59
1. Solid Fuels	NE,NO	2,538.33	NE,NO				2,538.33
2. Oil and Natural Gas	53.67	256.43	0.16				310.26
2. Industrial Processes	61,027.71	347.49	7,539.75	18,070.00	6,370.00	43,498.00	136,852.96
A. Mineral Products	56,520.30	NA,NO	NA,NO				56,520.30
B. Chemical Industry	4,184.37	329.15	7,539.75	NA	NA	NA	12,053.27
C. Metal Production	323.04	18.34	NO	IE,NE	IE,NA,NE	IE,NA,NE	341.38
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				NE,NO	NE,NO	NE,NO	NE,NO
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				18,070.00	6,370.00	43,498.00	67,938.00
G. Other	NO	NO	NO	NE,NO	NE,NO	NE,NO	NE,NO
3. Solvent and Other Product Use	NA,NE		356.85				356.85
4. Agriculture		17,955.09	13,237.81				31,192.90
A. Enteric Fermentation		7,787.92					7,787.92
B. Manure Management		3,089.18	5,501.83				8,591.01
C. Rice Cultivation		6,977.75					6,977.75
D. Agricultural Soils ⁽³⁾		NA	7,703.32				7,703.32
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		100.24	32.66				132.90
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽⁴⁾	-77,427.05	6.36	84.82				-77,335.87
A. Forest Land	-85,945.90	6.36	0.65				-85,938.90
B. Cropland	1,641.15	NE,NO	84.17				1,725.32
C. Grassland	-523.32	NE,NO	NE,NO				-523.32
D. Wetlands	77.52	NE,NO	NE,NO				77.52
E. Settlements	5,060.95	NE,NO	NE,NO				5,060.95
F. Other Land	1,735.28	NO	NO				1,735.28
G. Other	527.29	NA,NE	NA,NE				527.29
6. Waste	12,984.57	9,792.64	2,970.18				25,747.39
A. Solid Waste Disposal on Land	NA,NE,NO	7,573.36					7,573.36
B. Waste-water Handling		2,103.57	1,318.10				3,421.68
C. Waste Incineration	12,298.12	13.08	1,561.19				13,872.39
D. Other	686.45	102.62	90.89				879.96
7. Other (as specified in Summary 1.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	32,531.98	45.53	294.78				32,872.29
Aviation	13,919.12	8.27	137.65				14,065.05
Marine	18,612.86	37.25	157.13				18,807.24
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	18,870.94						18,870.94
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,280,976.97
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,203,641.10

(1) For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

(2) Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

(3) Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

(4) See footnote 8 to table Summary 1.A.

² Potential emissions of HFCs, PFCs and SF₆ are reported due to the generation of CRF Reporter

A9.3. Emissions³ and Removals in 1992SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS

(Sheet 1 of 1)

Inventory 1992

Submission 2012 v1.1

JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,081,439.99	31,526.24	31,374.88	19,750.00	6,370.00	47,800.00	1,218,261.11
1. Energy	1,083,526.98	3,438.48	7,227.06				1,094,192.52
A. Fuel Combustion (Sectoral Approach)	1,083,470.03	911.14	7,226.89				1,091,608.06
1. Energy Industries	333,717.45	31.85	929.83				334,679.13
2. Manufacturing Industries and Construction	358,404.85	352.17	1,541.66				360,298.67
3. Transport	226,859.69	303.05	4,460.92				231,623.66
4. Other Sectors	164,488.04	224.07	294.48				165,006.59
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	56.95	2,527.34	0.17				2,584.46
1. Solid Fuels	NE,NO	2,267.52	NE,NO				2,267.52
2. Oil and Natural Gas	56.95	259.82	0.17				316.94
2. Industrial Processes	61,026.54	322.22	7,452.41	19,750.00	6,370.00	47,800.00	142,721.16
A. Mineral Products	56,600.40	NA,NO	NA,NO				56,600.40
B. Chemical Industry	4,101.09	304.45	7,452.41	NA	NA	NA	11,857.96
C. Metal Production	325.05	17.76	NO	IE,NE	IE,NA,NE	IE,NA,NE	342.81
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				NE,NO	NE,NO	NE,NO	NE,NO
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				19,750.00	6,370.00	47,800.00	73,920.00
G. Other	NO	NO	NO	NE,NO	NE,NO	NE,NO	NE,NO
3. Solvent and Other Product Use	NA,NE		413.01				413.01
4. Agriculture		18,044.60	13,108.79				31,153.39
A. Enteric Fermentation		7,830.18					7,830.18
B. Manure Management		3,061.96	5,457.83				8,519.79
C. Rice Cultivation		7,059.04					7,059.04
D. Agricultural Soils ⁽³⁾		NA	7,620.17				7,620.17
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		93.42	30.79				124.21
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-77,137.76	4.39	79.83				-77,053.53
A. Forest Land	-86,301.81	4.39	0.45				-86,296.97
B. Cropland	1,736.08	NE,NO	79.39				1,815.47
C. Grassland	-467.11	NE,NO	NE,NO				-467.11
D. Wetlands	246.52	NE,NO	NE,NO				246.52
E. Settlements	5,746.82	NE,NO	NE,NO				5,746.82
F. Other Land	1,424.64	NO	NO				1,424.64
G. Other	477.11	NA,NE	NA,NE				477.11
6. Waste	14,024.24	9,716.55	3,093.77				26,834.55
A. Solid Waste Disposal on Land	NA,NE,NO	7,536.99					7,536.99
B. Waste-water Handling		2,063.19	1,302.96				3,366.15
C. Waste Incineration	13,325.34	13.43	1,699.63				15,038.41
D. Other	698.90	102.94	91.17				893.01
7. Other (as specified in Summary 1.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	32,937.28	45.92	298.63				33,281.83
Aviation	14,216.76	8.45	140.60				14,365.81
Marine	18,720.51	37.47	158.04				18,916.02
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	18,419.27						18,419.27
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,295,314.64
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,218,261.11

⁽¹⁾ For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

⁽²⁾ Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

⁽³⁾ Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

⁽⁴⁾ See footnote 8 to table Summary 1.A.

³ Potential emissions of HFCs, PFCs and SF₆ are reported due to the generation of CRF Reporter

A9.4. Emissions⁴ and Removals in 1993SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS
(Sheet 1 of 1)Inventory 1993
Submission 2012 v1.1
JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,070,920.05	31,274.03	31,120.70	21,310.00	8,860.00	45,410.00	1,208,894.78
1. Energy	1,077,164.28	3,269.31	7,275.57				1,087,709.17
A. Fuel Combustion (Sectoral Approach)	1,077,111.06	930.09	7,275.41				1,085,316.56
1. Energy Industries	315,598.93	31.64	938.35				316,568.92
2. Manufacturing Industries and Construction	357,499.46	352.95	1,579.56				359,431.97
3. Transport	231,727.93	295.90	4,433.85				236,457.68
4. Other Sectors	172,284.75	249.59	323.64				172,857.98
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	53.21	2,339.23	0.16				2,392.61
1. Solid Fuels	NE,NO	2,075.76	NE,NO				2,075.76
2. Oil and Natural Gas	53.21	263.46	0.16				316.84
2. Industrial Processes	59,959.49	320.55	7,302.85	21,310.00	8,860.00	45,410.00	143,162.89
A. Mineral Products	55,733.90	NA,NO	NA,NO				55,733.90
B. Chemical Industry	3,894.83	303.85	7,302.85	NA	NA	NA	11,501.53
C. Metal Production	330.76	16.70	NO	IE,NE	IE,NA,NE	IE,NA,NE	347.46
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				NE,NO	NE,NO	NE,NO	NE,NO
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				21,310.00	8,860.00	45,410.00	75,580.00
G. Other	NO	NO	NO	NE,NO	NE,NO	NE,NO	NE,NO
3. Solvent and Other Product Use	NA,NE		411.66				411.66
4. Agriculture		18,127.86	12,952.84				31,080.70
A. Enteric Fermentation		7,781.42					7,781.42
B. Manure Management		3,002.79	5,364.14				8,366.93
C. Rice Cultivation		7,247.60					7,247.60
D. Agricultural Soils ⁽³⁾		NA	7,556.74				7,556.74
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		96.05	31.97				128.02
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-79,977.77	24.22	76.19				-79,877.35
A. Forest Land	-86,650.41	24.22	2.46				-86,623.73
B. Cropland	927.58	NE,NO	73.73				1,001.31
C. Grassland	-540.11	NE,NO	NE,NO				-540.11
D. Wetlands	138.47	NE,NO	NE,NO				138.47
E. Settlements	3,908.55	NE,NO	NE,NO				3,908.55
F. Other Land	1,756.59	NO	NO				1,756.59
G. Other	481.56	NA,NE	NA,NE				481.56
6. Waste	13,774.05	9,532.08	3,101.59				26,407.72
A. Solid Waste Disposal on Land	NA,NE,NO	7,403.56					7,403.56
B. Waste-water Handling		2,012.23	1,306.88				3,319.11
C. Waste Incineration	13,093.30	13.36	1,703.54				14,810.19
D. Other	680.75	102.94	91.17				874.85
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	34,935.20	50.42	314.96				35,300.58
Aviation	13,856.19	8.23	137.03				14,001.45
Marine	21,079.01	42.19	177.93				21,299.12
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	17,568.73						17,568.73
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,288,772.14
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,208,894.78

(1) For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

(2) Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

(3) Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

(4) See footnote 8 to table Summary I.A.

⁴ Potential emissions of HFCs, PFCs and SF₆ are reported due to the generation of CRF Reporter

A9.5. Emissions⁵ and Removals in 1994SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS

(Sheet 1 of 1)

Inventory 1994

Submission 2012 v1.1

JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,128,989.90	30,600.69	32,290.68	28,840.00	12,274.00	45,410.00	1,278,405.27
1. Energy	1,133,210.28	2,908.20	7,570.44				1,143,688.92
A. Fuel Combustion (Sectoral Approach)	1,133,159.13	928.67	7,570.28				1,141,658.08
1. Energy Industries	356,359.51	33.79	1,010.41				357,403.72
2. Manufacturing Industries and Construction	365,878.17	361.90	1,717.85				367,957.92
3. Transport	243,681.03	297.61	4,515.10				248,493.74
4. Other Sectors	167,240.42	235.36	326.91				167,802.69
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	51.15	1,979.53	0.16				2,030.84
1. Solid Fuels	NE,NO	1,712.96	NE,NO				1,712.96
2. Oil and Natural Gas	51.15	266.57	0.16				317.88
2. Industrial Processes	61,189.78	320.85	8,298.10	28,840.00	12,274.00	45,410.00	156,332.74
A. Mineral Products	56,698.93	NA,NO	NA,NO				56,698.93
B. Chemical Industry	4,145.10	303.40	8,298.10	NA	NA	NA	12,746.60
C. Metal Production	345.76	17.45	NO	IE,NE	IE,NA,NE	IE,NA,NE	363.21
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				NE,NO	NE,NO	NE,NO	NE,NO
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				28,840.00	12,274.00	45,410.00	86,524.00
G. Other	NO	NO	NO	NE,NO	NE,NO	NE,NO	NE,NO
3. Solvent and Other Product Use	NA,NE		438.02				438.02
4. Agriculture		17,990.78	12,677.09				30,667.88
A. Enteric Fermentation		7,691.88					7,691.88
B. Manure Management		2,942.69	5,250.91				8,193.60
C. Rice Cultivation		7,263.40					7,263.40
D. Agricultural Soils ⁽³⁾		NA	7,395.03				7,395.03
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		92.82	31.15				123.97
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-81,679.07	17.88	69.80				-81,591.38
A. Forest Land	-86,995.18	17.88	1.81				-86,975.48
B. Cropland	878.27	NE,NO	67.99				946.26
C. Grassland	-508.08	NE,NO	NE,NO				-508.08
D. Wetlands	115.95	NE,NO	NE,NO				115.95
E. Settlements	2,881.14	NE,NO	NE,NO				2,881.14
F. Other Land	1,656.09	NO	NO				1,656.09
G. Other	292.73	NA,NE	NA,NE				292.73
6. Waste	16,268.90	9,362.97	3,237.23				28,869.11
A. Solid Waste Disposal on Land	NA,NE,NO	7,290.56					7,290.56
B. Waste-water Handling		1,945.32	1,269.76				3,215.08
C. Waste Incineration	15,566.99	14.49	1,867.73				17,449.21
D. Other	701.91	112.60	99.73				914.25
7. Other (as specified in Summary 1.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	36,093.69	51.04	326.49				36,471.22
Aviation	15,066.49	8.95	149.00				15,224.44
Marine	21,027.20	42.08	177.50				21,246.78
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	17,803.39						17,803.39
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,359,996.65
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,278,405.27

⁽¹⁾ For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

⁽²⁾ Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

⁽³⁾ Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

⁽⁴⁾ See footnote 8 to table Summary 1.A.

⁵ Potential emissions of HFCs, PFCs and SF₆ are reported due to the generation of CRF Reporter

A9.6. Emissions and Removals in 1995

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS
(Sheet 1 of 1)Inventory 1995
Submission 2012 v1.1
JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,141,636.45	29,736.33	32,717.98	20,260.17	14,240.36	16,961.45	1,255,552.74
1. Energy	1,145,820.01	2,647.81	8,284.79				1,156,752.61
A. Fuel Combustion (Sectoral Approach)	1,145,769.09	1,037.94	8,284.63				1,155,091.66
1. Energy Industries	344,948.18	34.41	1,413.32				346,395.92
2. Manufacturing Industries and Construction	370,539.38	437.59	1,871.31				372,848.29
3. Transport	251,166.53	308.82	4,651.65				256,126.99
4. Other Sectors	179,115.00	257.11	348.35				179,720.46
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	50.92	1,609.87	0.16				1,660.95
1. Solid Fuels	NE,NO	1,344.68	NE,NO				1,344.68
2. Oil and Natural Gas	50.92	265.19	0.16				316.26
2. Industrial Processes	61,338.27	322.37	8,212.71	20,260.17	14,240.36	16,961.45	121,335.34
A. Mineral Products	56,761.48	NA,NO	NA,NO				56,761.48
B. Chemical Industry	4,219.57	304.45	8,212.71	NA	NA	NA	12,736.73
C. Metal Production	357.22	17.92	NO	IE,NE	69.74	119.50	564.38
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				17,445.12	762.85	4,708.30	22,916.27
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				2,815.05	13,407.78	12,133.65	28,356.48
G. Other	NO	NO	NO	NO	NO	NO	NO
3. Solvent and Other Product Use	NA,NE		437.58				437.58
4. Agriculture		17,676.22	12,363.00				30,039.22
A. Enteric Fermentation		7,606.43					7,606.43
B. Manure Management		2,893.04	5,151.97				8,045.01
C. Rice Cultivation		7,082.74					7,082.74
D. Agricultural Soils ⁽³⁾		NA	7,179.42				7,179.42
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		94.01	31.61				125.62
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-82,056.24	8.73	61.59				-81,985.92
A. Forest Land	-87,340.67	8.73	0.89				-87,331.05
B. Cropland	822.78	NE,NO	60.71				883.49
C. Grassland	-481.07	NE,NO	NE,NO				-481.07
D. Wetlands	360.41	NE,NO	NE,NO				360.41
E. Settlements	2,799.60	NE,NO	NE,NO				2,799.60
F. Other Land	1,479.20	NO	NO				1,479.20
G. Other	303.50	NA,NE	NA,NE				303.50
6. Waste	16,534.40	9,081.20	3,358.32				28,973.92
A. Solid Waste Disposal on Land	NA,NE,NO	7,075.92					7,075.92
B. Waste-water Handling		1,884.04	1,251.96				3,136.00
C. Waste Incineration	15,866.57	14.87	2,012.15				17,893.59
D. Other	667.83	106.37	94.22				868.42
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	38,179.77	52.60	346.77				38,579.14
Aviation	16,922.99	10.06	167.36				17,100.41
Marine	21,256.78	42.54	179.42				21,478.73
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	18,487.35						18,487.35
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,337,538.66
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,255,552.74

(1) For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

(2) Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

(3) Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

(4) See footnote 8 to table Summary I.A.

A9.7. Emissions and Removals in 1996

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS

(Sheet 1 of 1)

Inventory 1996

Submission 2012 v1.1

JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,149,855.13	28,994.01	33,700.94	19,906.20	14,783.02	17,535.35	1,264,774.64
1. Energy	1,157,958.90	2,522.36	8,417.17				1,168,898.43
A. Fuel Combustion (Sectoral Approach)	1,157,909.53	961.87	8,417.02				1,167,288.42
1. Energy Industries	345,134.72	36.20	1,444.43				346,615.35
2. Manufacturing Industries and Construction	378,811.73	380.74	1,931.29				381,123.77
3. Transport	256,750.56	314.64	4,738.70				261,803.89
4. Other Sectors	177,212.53	230.29	302.59				177,745.40
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	49.37	1,560.49	0.15				1,610.01
1. Solid Fuels	NE,NO	1,297.15	NE,NO				1,297.15
2. Oil and Natural Gas	49.37	263.34	0.15				312.86
2. Industrial Processes	61,696.11	312.02	9,220.07	19,906.20	14,783.02	17,535.35	123,452.75
A. Mineral Products	57,112.69	NA,NO	NA,NO				57,112.69
B. Chemical Industry	4,203.43	293.80	9,220.07	NA	NA	NA	13,717.30
C. Metal Production	379.99	18.22	NO	IE,NE	65.88	143.40	607.48
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				16,052.32	1,007.80	4,182.50	21,242.62
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				3,853.88	13,709.34	13,209.45	30,772.67
G. Other	NO	NO	NO	NO	NO	NO	NO
3. Solvent and Other Product Use	NA,NE		420.94				420.94
4. Agriculture		17,294.20	12,089.89				29,384.09
A. Enteric Fermentation		7,551.46					7,551.46
B. Manure Management		2,859.09	5,089.03				7,948.13
C. Rice Cultivation		6,793.69					6,793.69
D. Agricultural Soils ⁽³⁾		NA	6,970.63				6,970.63
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		89.96	30.22				120.18
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-86,750.73	28.68	54.49				-86,667.56
A. Forest Land	-91,316.25	28.68	2.91				-91,284.66
B. Cropland	656.98	NE,NO	51.58				708.56
C. Grassland	-464.64	NE,NO	NE,NO				-464.64
D. Wetlands	647.00	NE,NO	NE,NO				647.00
E. Settlements	2,061.14	NE,NO	NE,NO				2,061.14
F. Other Land	1,372.34	NO	NO				1,372.34
G. Other	292.70	NA,NE	NA,NE				292.70
6. Waste	16,950.85	8,836.76	3,498.39				29,286.00
A. Solid Waste Disposal on Land	NA,NE,NO	6,874.74					6,874.74
B. Waste-water Handling		1,847.84	1,273.88				3,121.72
C. Waste Incineration	16,310.38	15.24	2,136.87				18,462.50
D. Other	640.47	98.94	87.63				827.04
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	30,958.25	36.01	288.02				31,282.28
Aviation	18,441.91	10.96	182.38				18,635.25
Marine	12,516.34	25.05	105.64				12,647.03
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	18,547.51						18,547.51
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,351,442.21
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,264,774.64

⁽¹⁾ For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

⁽²⁾ Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

⁽³⁾ Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

⁽⁴⁾ See footnote 8 to table Summary I.A.

A9.8. Emissions and Removals in 1997

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS
(Sheet 1 of 1)Inventory 1997
Submission 2012 v1.1
JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,144,452.82	27,931.85	34,364.06	19,905.11	16,164.62	14,998.12	1,257,816.58
1. Energy	1,154,948.65	2,228.39	8,643.71				1,165,820.75
A. Fuel Combustion (Sectoral Approach)	1,154,900.68	951.14	8,643.56				1,164,495.38
1. Energy Industries	342,054.20	38.03	1,490.10				343,582.33
2. Manufacturing Industries and Construction	381,142.92	361.98	2,061.00				383,565.90
3. Transport	258,734.10	315.74	4,786.90				263,836.74
4. Other Sectors	172,969.46	235.38	305.56				173,510.41
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	47.97	1,277.25	0.15				1,325.37
1. Solid Fuels	NE,NO	1,006.86	NE,NO				1,006.86
2. Oil and Natural Gas	47.97	270.39	0.15				318.51
2. Industrial Processes	59,024.03	260.90	9,792.47	19,905.11	16,164.62	14,998.12	120,145.24
A. Mineral Products	54,495.36	NA,NO	NA,NO				54,495.36
B. Chemical Industry	4,144.19	242.58	9,792.47	NA	NA	NA	14,179.23
C. Metal Production	384.48	18.33	NO	IE,NE	59.43	191.20	653.44
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				15,077.99	1,416.80	2,581.20	19,075.99
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				4,827.12	14,688.39	12,225.72	31,741.22
G. Other	NO	NO	NO	NO	NO	NO	NO
3. Solvent and Other Product Use	NA,NE		404.60				404.60
4. Agriculture		16,847.98	11,897.61				28,745.59
A. Enteric Fermentation		7,505.45					7,505.45
B. Manure Management		2,816.67	5,031.14				7,847.81
C. Rice Cultivation		6,440.28					6,440.28
D. Agricultural Soils ⁽³⁾		NA	6,837.81				6,837.81
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		85.58	28.66				114.24
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-87,067.08	34.51	46.02				-86,986.55
A. Forest Land	-91,160.76	34.51	3.50				-91,122.75
B. Cropland	540.56	NE,NO	42.52				583.08
C. Grassland	-446.84	NE,NO	NE,NO				-446.84
D. Wetlands	123.86	NE,NO	NE,NO				123.86
E. Settlements	1,710.78	NE,NO	NE,NO				1,710.78
F. Other Land	1,861.71	NO	NO				1,861.71
G. Other	303.61	NA,NE	NA,NE				303.61
6. Waste	17,547.22	8,560.06	3,579.65				29,686.94
A. Solid Waste Disposal on Land	NA,NE,NO	6,645.54					6,645.54
B. Waste-water Handling		1,800.45	1,283.33				3,083.78
C. Waste Incineration	16,891.99	14.71	2,208.32				19,115.02
D. Other	655.23	99.36	88.01				842.60
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	35,432.29	43.98	326.77				35,803.04
Aviation	19,134.37	11.37	189.23				19,334.97
Marine	16,297.92	32.61	137.54				16,468.07
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	19,107.10						19,107.10
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,344,803.12
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,257,816.58

(1) For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

(2) Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

(3) Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

(4) See footnote 8 to table Summary I.A.

A9.9. Emissions and Removals in 1998

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS

(Sheet 1 of 1)

Inventory 1998

Submission 2012 v1.1

JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,109,059.67	27,108.79	32,834.39	19,415.96	13,411.82	13,624.11	1,215,454.75
1. Energy	1,125,032.90	2,059.34	8,513.34				1,135,605.57
A. Fuel Combustion (Sectoral Approach)	1,124,990.17	921.36	8,513.21				1,134,424.73
1. Energy Industries	332,405.28	39.82	1,515.31				333,960.41
2. Manufacturing Industries and Construction	357,838.95	324.31	1,985.64				360,148.90
3. Transport	257,853.86	304.72	4,688.42				262,847.00
4. Other Sectors	176,892.07	252.52	323.83				177,468.42
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	42.73	1,137.98	0.13				1,180.84
1. Solid Fuels	NE,NO	872.46	NE,NO				872.46
2. Oil and Natural Gas	42.73	265.52	0.13				308.38
2. Industrial Processes	53,376.38	243.52	8,577.87	19,415.96	13,411.82	13,624.11	108,649.67
A. Mineral Products	49,443.45	NA,NO	NA,NO				49,443.45
B. Chemical Industry	3,639.82	227.37	8,577.87	NA	NA	NA	12,445.07
C. Metal Production	293.11	16.15	NO	IE,NE	49.40	406.30	764.96
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				14,053.43	1,389.50	2,103.20	17,546.13
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				5,362.53	11,972.92	11,114.61	28,450.06
G. Other	NO	NO	NO	NO	NO	NO	NO
3. Solvent and Other Product Use	NA,NE		377.05				377.05
4. Agriculture		16,548.59	11,757.25				28,305.84
A. Enteric Fermentation		7,466.79					7,466.79
B. Manure Management		2,770.83	4,986.39				7,757.21
C. Rice Cultivation		6,229.14					6,229.14
D. Agricultural Soils ⁽³⁾		NA	6,743.48				6,743.48
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		81.84	27.39				109.23
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-86,869.79	10.73	39.80				-86,819.27
A. Forest Land	-91,002.78	10.73	1.09				-90,990.97
B. Cropland	548.02	NE,NO	38.71				586.73
C. Grassland	-421.24	NE,NO	NE,NO				-421.24
D. Wetlands	501.97	NE,NO	NE,NO				501.97
E. Settlements	1,702.19	NE,NO	NE,NO				1,702.19
F. Other Land	1,502.08	NO	NO				1,502.08
G. Other	299.97	NA,NE	NA,NE				299.97
6. Waste	17,520.19	8,246.62	3,569.08				29,335.89
A. Solid Waste Disposal on Land	NA,NE,NO	6,373.83					6,373.83
B. Waste-water Handling		1,755.01	1,266.22				3,021.23
C. Waste Incineration	16,911.07	14.53	2,211.41				19,137.01
D. Other	609.12	103.25	91.45				803.81
7. Other (as specified in Summary 1.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	37,361.08	46.63	344.33				37,752.04
Aviation	20,001.55	11.89	197.80				20,211.24
Marine	17,359.53	34.74	146.53				17,540.80
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	17,556.58						17,556.58
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,302,274.02
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,215,454.75

⁽¹⁾ For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

⁽²⁾ Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

⁽³⁾ Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

⁽⁴⁾ See footnote 8 to table Summary 1.A.

A9.10. Emissions and Removals in 1999

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS
(Sheet 1 of 1)Inventory 1999
Submission 2012 v1.1
JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,143,783.80	26,490.02	26,398.61	19,934.46	10,395.49	9,309.93	1,236,312.32
1. Energy	1,160,147.36	2,078.81	8,729.73				1,170,955.89
A. Fuel Combustion (Sectoral Approach)	1,160,109.30	950.39	8,729.61				1,169,789.29
1. Energy Industries	349,785.30	42.67	1,615.94				351,443.92
2. Manufacturing Industries and Construction	365,074.78	328.47	2,073.06				367,476.31
3. Transport	260,017.18	303.45	4,681.87				265,002.50
4. Other Sectors	185,232.04	275.79	358.74				185,866.57
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	38.06	1,128.42	0.12				1,166.60
1. Solid Fuels	NE,NO	865.69	NE,NO				865.69
2. Oil and Natural Gas	38.06	262.73	0.12				300.91
2. Industrial Processes	53,400.15	236.22	2,000.86	19,934.46	10,395.49	9,309.93	95,277.12
A. Mineral Products	49,180.61	NA,NO	NA,NO				49,180.61
B. Chemical Industry	3,965.06	220.14	2,000.86	NA	NA	NA	6,186.06
C. Metal Production	254.49	16.08	NO	IE,NE	29.12	645.30	944.99
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				14,260.55	1,270.88	1,529.60	17,061.03
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				5,673.90	9,095.49	7,135.03	21,904.42
G. Other	NO	NO	NO	NO	NO	NO	NO
3. Solvent and Other Product Use	NA,NE		362.53				362.53
4. Agriculture		16,228.80	11,665.50				27,894.30
A. Enteric Fermentation		7,407.75					7,407.75
B. Manure Management		2,717.58	4,933.09				7,650.67
C. Rice Cultivation		6,024.77					6,024.77
D. Agricultural Soils ⁽³⁾		NA	6,706.24				6,706.24
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		78.70	26.17				104.87
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-87,093.55	5.25	36.08				-87,052.21
A. Forest Land	-90,846.46	5.25	0.53				-90,840.67
B. Cropland	509.84	NE,NO	35.55				545.39
C. Grassland	-408.21	NE,NO	NE,NO				-408.21
D. Wetlands	478.14	NE,NO	NE,NO				478.14
E. Settlements	1,313.68	NE,NO	NE,NO				1,313.68
F. Other Land	1,565.94	NO	NO				1,565.94
G. Other	293.52	NA,NE	NA,NE				293.52
6. Waste	17,329.84	7,940.95	3,603.91				28,874.70
A. Solid Waste Disposal on Land	NA,NE,NO	6,116.88					6,116.88
B. Waste-water Handling		1,706.21	1,229.03				2,935.24
C. Waste Incineration	16,677.27	14.04	2,282.92				18,974.22
D. Other	652.58	103.82	91.96				848.35
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	36,022.49	44.55	332.43				36,399.47
Aviation	19,576.46	11.63	193.60				19,781.70
Marine	16,446.03	32.92	138.83				16,617.77
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	18,260.06						18,260.06
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,323,364.53
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,236,312.32

(1) For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

(2) Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

(3) Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

(4) See footnote 8 to table Summary I.A.

A9.11. Emissions and Removals in 2000

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS

(Sheet 1 of 1)

Inventory 2000

Submission 2012 v1.1

JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,163,777.08	25,899.87	28,997.29	18,800.43	9,519.49	7,188.49	1,254,182.66
1. Energy	1,180,079.82	2,006.45	8,787.66				1,190,873.94
A. Fuel Combustion (Sectoral Approach)	1,180,043.79	963.30	8,787.55				1,189,794.65
1. Energy Industries	357,574.13	43.63	1,709.09				359,326.85
2. Manufacturing Industries and Construction	376,777.84	351.50	2,126.30				379,255.64
3. Transport	259,076.39	298.36	4,589.32				263,964.07
4. Other Sectors	186,615.43	269.81	362.85				187,248.08
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	36.03	1,043.15	0.11				1,079.29
1. Solid Fuels	NE,NO	769.13	NE,NO				769.13
2. Oil and Natural Gas	36.03	274.02	0.11				310.16
2. Industrial Processes	53,983.02	195.78	4,690.09	18,800.43	9,519.49	7,188.49	94,377.30
A. Mineral Products	49,841.59	NA,NO	NA,NO				49,841.59
B. Chemical Industry	3,893.01	178.95	4,690.09	NA	NA	NA	8,762.04
C. Metal Production	248.42	16.84	NO	IE,NE	17.78	1,027.70	1,310.74
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				12,659.84	1,359.00	860.40	14,879.24
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				6,140.59	8,142.70	5,300.39	19,583.69
G. Other	NO	NO	NO	NO	NO	NO	NO
3. Solvent and Other Product Use	NA,NE		340.99				340.99
4. Agriculture		16,044.72	11,584.57				27,629.29
A. Enteric Fermentation		7,369.97					7,369.97
B. Manure Management		2,677.89	4,884.82				7,562.71
C. Rice Cultivation		5,919.76					5,919.76
D. Agricultural Soils ⁽³⁾		NA	6,674.25				6,674.25
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		77.10	25.50				102.60
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-87,779.62	7.78	32.70				-87,739.14
A. Forest Land	-90,689.18	7.78	0.79				-90,680.61
B. Cropland	355.68	NE,NO	31.91				387.60
C. Grassland	-405.80	NE,NO	NE,NO				-405.80
D. Wetlands	451.43	NE,NO	NE,NO				451.43
E. Settlements	947.44	NE,NO	NE,NO				947.44
F. Other Land	1,227.94	NO	NO				1,227.94
G. Other	332.87	NA,NE	NA,NE				332.87
6. Waste	17,493.86	7,645.14	3,561.27				28,700.27
A. Solid Waste Disposal on Land	NA,NE,NO	5,878.27					5,878.27
B. Waste-water Handling		1,657.37	1,216.19				2,873.57
C. Waste Incineration	16,837.95	13.33	2,259.90				19,111.18
D. Other	655.91	96.16	85.17				837.25
7. Other (as specified in Summary 1.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	36,731.88	46.01	336.87				37,114.77
Aviation	19,542.61	11.61	191.78				19,746.00
Marine	17,189.28	34.40	145.09				17,368.76
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	18,846.04						18,846.04
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,341,921.79
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,254,182.66

⁽¹⁾ For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

⁽²⁾ Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

⁽³⁾ Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

⁽⁴⁾ See footnote 8 to table Summary 1.A.

A9.12. Emissions and Removals in 2001

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS
(Sheet 1 of 1)Inventory 2001
Submission 2012 v1.1
JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,148,504.98	25,127.50	25,557.05	16,168.06	7,902.31	5,962.42	1,229,222.32
1. Energy	1,167,416.32	1,772.13	8,797.10				1,177,985.55
A. Fuel Combustion (Sectoral Approach)	1,167,383.88	933.95	8,797.00				1,177,114.84
1. Energy Industries	349,730.24	43.68	1,931.00				351,704.92
2. Manufacturing Industries and Construction	366,480.21	325.89	2,084.72				368,890.82
3. Transport	261,120.73	292.66	4,412.32				265,825.71
4. Other Sectors	190,052.70	271.72	368.97				190,693.38
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	32.44	838.18	0.10				870.72
1. Solid Fuels	NE,NO	570.30	NE,NO				570.30
2. Oil and Natural Gas	32.44	267.88	0.10				300.42
2. Industrial Processes	52,758.23	147.50	1,414.89	16,168.06	7,902.31	5,962.42	84,353.40
A. Mineral Products	48,948.92	NA,NO	NA,NO				48,948.92
B. Chemical Industry	3,598.60	131.66	1,414.89	NA	NA	NA	5,145.14
C. Metal Production	210.71	15.84	NO	IE,NE	15.73	1,147.20	1,389.48
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				9,713.43	1,082.60	788.70	11,584.73
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				6,454.63	6,803.99	4,026.52	17,285.13
G. Other	NO	NO	NO	NO	NO	NO	NO
3. Solvent and Other Product Use	NA,NE		343.60				343.60
4. Agriculture		15,863.11	11,497.59				27,360.71
A. Enteric Fermentation		7,325.24					7,325.24
B. Manure Management		2,652.15	4,839.23				7,491.39
C. Rice Cultivation		5,810.23					5,810.23
D. Agricultural Soils ⁽³⁾		NA	6,633.61				6,633.61
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		75.49	24.75				100.24
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-87,915.51	12.42	29.84				-87,873.26
A. Forest Land	-90,534.44	12.42	1.26				-90,520.77
B. Cropland	295.23	NE,NO	28.58				323.81
C. Grassland	-393.95	NE,NO	NE,NO				-393.95
D. Wetlands	414.48	NE,NO	NE,NO				414.48
E. Settlements	727.90	NE,NO	NE,NO				727.90
F. Other Land	1,327.95	NO	NO				1,327.95
G. Other	247.31	NA,NE	NA,NE				247.31
6. Waste	16,245.95	7,332.34	3,474.03				27,052.31
A. Solid Waste Disposal on Land	NA,NE,NO	5,618.48					5,618.48
B. Waste-water Handling		1,603.75	1,198.15				2,801.90
C. Waste Incineration	15,615.42	12.61	2,189.52				17,817.54
D. Other	630.53	97.50	86.36				814.40
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	33,571.42	40.84	309.05				33,921.32
Aviation	18,721.34	11.13	183.72				18,916.19
Marine	14,850.08	29.72	125.33				15,005.12
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	17,203.99						17,203.99
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,317,095.58
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,229,222.32

(1) For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

(2) Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

(3) Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

(4) See footnote 8 to table Summary I.A.

A9.14. Emissions and Removals in 2003

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS
(Sheet 1 of 1)Inventory 2003
Submission 2012 v1.1
JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,180,429.99	23,673.98	24,508.52	13,761.68	7,181.45	5,253.91	1,254,809.54
1. Energy	1,213,922.14	1,294.65	8,233.99				1,223,450.78
A. Fuel Combustion (Sectoral Approach)	1,213,887.66	905.27	8,233.88				1,223,026.81
1. Energy Industries	395,368.37	36.29	1,901.98				397,306.63
2. Manufacturing Industries and Construction	373,172.66	347.73	2,079.77				375,600.16
3. Transport	252,947.16	270.14	3,879.77				257,097.07
4. Other Sectors	192,399.48	251.11	372.36				193,022.94
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	34.48	389.38	0.11				423.97
1. Solid Fuels	NE,NO	93.86	NE,NO				93.86
2. Oil and Natural Gas	34.48	295.52	0.11				330.11
2. Industrial Processes	49,127.25	133.88	1,259.55	13,761.68	7,181.45	5,253.91	76,717.72
A. Mineral Products	45,757.07	NA,NO	NA,NO				45,757.07
B. Chemical Industry	3,128.60	117.37	1,259.55	NA	NA	NA	4,505.53
C. Metal Production	241.57	16.50	NO	IE,NE	15.21	1,125.53	1,398.81
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				5,459.50	965.60	812.60	7,237.70
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				8,302.18	6,200.65	3,315.79	17,818.61
G. Other	NO	NO	NO	NO	NO	NO	NO
3. Solvent and Other Product Use	NA,NE		320.83				320.83
4. Agriculture		15,516.76	11,370.63				26,887.39
A. Enteric Fermentation		7,163.24					7,163.24
B. Manure Management		2,594.79	4,778.58				7,373.37
C. Rice Cultivation		5,690.55					5,690.55
D. Agricultural Soils ⁽³⁾		NA	6,569.95				6,569.95
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		68.18	22.10				90.27
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-98,191.21	3.93	21.90				-98,165.38
A. Forest Land	-99,129.26	3.93	0.40				-99,124.93
B. Cropland	269.46	NE,NO	21.51				290.97
C. Grassland	-358.71	NE,NO	NE,NO				-358.71
D. Wetlands	68.67	NE,NO	NE,NO				68.67
E. Settlements	-263.00	NE,NO	NE,NO				-263.00
F. Other Land	975.24	NO	NO				975.24
G. Other	246.37	NA,NE	NA,NE				246.37
6. Waste	15,571.81	6,724.77	3,301.62				25,598.20
A. Solid Waste Disposal on Land	NA,NE,NO	5,092.09					5,092.09
B. Waste-water Handling		1,507.51	1,189.15				2,696.66
C. Waste Incineration	15,055.29	16.79	2,016.48				17,088.56
D. Other	516.53	108.37	95.99				720.89
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	37,506.71	46.37	344.55				37,897.63
Aviation	20,387.64	12.12	200.08				20,599.83
Marine	17,119.07	34.25	144.47				17,297.79
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	18,296.50						18,296.50
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,352,974.92
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,254,809.54

(1) For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

(2) Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

(3) Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

(4) See footnote 8 to table Summary I.A.

A9.15. Emissions and Removals in 2004

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS

(Sheet 1 of 1)

Inventory 2004

Submission 2012 v1.1

JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,180,298.73	23,244.49	24,551.69	10,552.49	7,478.30	5,095.89	1,251,221.57
1. Energy	1,214,020.86	1,273.19	7,971.09				1,223,265.13
A. Fuel Combustion (Sectoral Approach)	1,213,985.86	900.22	7,970.98				1,222,857.06
1. Energy Industries	390,980.48	35.27	1,902.22				392,917.97
2. Manufacturing Industries and Construction	378,733.43	354.85	2,119.28				381,207.57
3. Transport	252,413.86	250.33	3,572.83				256,237.02
4. Other Sectors	191,858.09	259.78	376.64				192,494.51
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	34.99	372.96	0.11				408.07
1. Solid Fuels	NE,NO	66.51	NE,NO				66.51
2. Oil and Natural Gas	34.99	306.45	0.11				341.56
2. Industrial Processes	48,959.48	143.54	1,657.60	10,552.49	7,478.30	5,095.89	73,887.29
A. Mineral Products	45,529.84	NA,NO	NA,NO				45,529.84
B. Chemical Industry	3,171.80	126.53	1,657.60	NA	NA	NA	4,955.94
C. Metal Production	257.84	17.01	NO	IE,NE	14.80	1,111.02	1,400.67
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				1,469.74	866.84	764.80	3,101.38
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				9,082.75	6,596.66	3,220.06	18,899.47
G. Other	NO	NO	NO	NO	NO	NO	NO
3. Solvent and Other Product Use	NA,NE		297.54				297.54
4. Agriculture		15,392.03	11,304.43				26,696.46
A. Enteric Fermentation		7,063.68					7,063.68
B. Manure Management		2,549.70	4,750.12				7,299.81
C. Rice Cultivation		5,712.00					5,712.00
D. Agricultural Soils ⁽³⁾		NA	6,532.78				6,532.78
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		66.65	21.54				88.19
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-97,705.95	12.13	19.06				-97,674.77
A. Forest Land	-98,594.30	12.13	1.23				-98,580.94
B. Cropland	234.52	NE,NO	17.83				252.35
C. Grassland	-347.74	NE,NO	NE,NO				-347.74
D. Wetlands	62.33	NE,NO	NE,NO				62.33
E. Settlements	-234.73	NE,NO	NE,NO				-234.73
F. Other Land	937.70	NO	NO				937.70
G. Other	236.27	NA,NE	NA,NE				236.27
6. Waste	15,024.34	6,423.60	3,301.97				24,749.91
A. Solid Waste Disposal on Land	NA,NE,NO	4,825.80					4,825.80
B. Waste-water Handling		1,469.22	1,196.08				2,665.30
C. Waste Incineration	14,517.64	15.38	2,005.62				16,538.65
D. Other	506.70	113.20	100.27				720.17
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA	NA	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	39,113.12	48.45	359.20				39,520.78
Aviation	21,190.20	12.59	207.95				21,410.75
Marine	17,922.92	35.86	151.25				18,110.03
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	18,188.60						18,188.60
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,348,896.34
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,251,221.57

⁽¹⁾ For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

⁽²⁾ Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

⁽³⁾ Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

⁽⁴⁾ See footnote 8 to table Summary I.A.

A9.16. Emissions and Removals in 2005

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS
(Sheet 1 of 1)Inventory 2005
Submission 2012 v1.1
JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,191,514.71	22,863.86	24,080.53	10,518.22	7,002.07	4,807.94	1,260,787.33
1. Energy	1,217,733.79	1,283.59	7,912.99				1,226,930.36
A. Fuel Combustion (Sectoral Approach)	1,217,696.19	887.85	7,912.87				1,226,496.91
1. Energy Industries	406,038.52	37.23	2,119.24				408,194.99
2. Manufacturing Industries and Construction	371,228.70	350.70	2,093.24				373,672.65
3. Transport	247,009.69	237.50	3,320.13				250,567.32
4. Other Sectors	193,419.28	262.42	380.25				194,061.95
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	37.60	395.74	0.12				433.46
1. Solid Fuels	NE,NO	73.56	NE,NO				73.56
2. Oil and Natural Gas	37.60	322.18	0.12				359.90
2. Industrial Processes	50,031.45	133.87	1,299.94	10,518.22	7,002.07	4,807.94	73,793.48
A. Mineral Products	46,902.66	NA,NO	NA,NO				46,902.66
B. Chemical Industry	2,886.85	116.98	1,299.94	NA	NA	NA	4,303.77
C. Metal Production	241.93	16.89	NO	IE,NE	14.80	1,157.31	1,430.93
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				816.01	837.49	975.12	2,628.62
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				9,702.21	6,149.78	2,675.51	18,527.50
G. Other	NO	NO	NO	NA,NO	NA,NO	NO	NA,NO
3. Solvent and Other Product Use	NA,NE		266.41				266.41
4. Agriculture		15,309.32	11,212.04				26,521.36
A. Enteric Fermentation		7,001.90					7,001.90
B. Manure Management		2,502.84	4,747.79				7,250.63
C. Rice Cultivation		5,739.10					5,739.10
D. Agricultural Soils ⁽³⁾		NA	6,443.07				6,443.07
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		65.48	21.17				86.65
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-90,741.81	9.18	15.67				-90,716.96
A. Forest Land	-92,010.94	9.18	0.93				-92,000.83
B. Cropland	277.08	NE,NO	14.74				291.82
C. Grassland	-335.58	NE,NO	NE,NO				-335.58
D. Wetlands	15.63	NE,NO	NE,NO				15.63
E. Settlements	125.87	NE,NO	NE,NO				125.87
F. Other Land	954.87	NO	NO				954.87
G. Other	231.25	NA,NE	NA,NE				231.25
6. Waste	14,491.29	6,127.90	3,373.48				23,992.68
A. Solid Waste Disposal on Land	NA,NE,NO	4,568.68					4,568.68
B. Waste-water Handling		1,419.01	1,165.77				2,584.77
C. Waste Incineration	13,984.48	14.27	2,096.16				16,094.90
D. Other	506.81	125.95	111.56				744.32
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA	NA	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	41,564.88	53.16	380.11				41,998.14
Aviation	21,336.33	12.68	209.39				21,558.39
Marine	20,228.55	40.48	170.72				20,439.75
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	21,743.33						21,743.33
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,351,504.29
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,260,787.33

(1) For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

(2) Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

(3) Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

(4) See footnote 8 to table Summary I.A.

A9.17. Emissions and Removals in 2006

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS

(Sheet 1 of 1)

Inventory 2006

Submission 2012 v1.1

JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,178,092.59	22,499.94	24,063.25	11,742.22	7,315.75	4,910.86	1,248,624.61
1. Energy	1,199,312.92	1,325.13	7,707.39				1,208,345.44
A. Fuel Combustion (Sectoral Approach)	1,199,277.03	916.88	7,707.28				1,207,901.18
1. Energy Industries	394,358.50	39.16	2,117.05				396,514.71
2. Manufacturing Industries and Construction	373,287.05	364.96	2,075.96				375,727.97
3. Transport	243,632.49	221.80	3,139.41				246,993.71
4. Other Sectors	187,998.99	290.95	374.85				188,664.80
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	35.89	408.25	0.11				444.26
1. Solid Fuels	NE,NO	68.12	NE,NO				68.12
2. Oil and Natural Gas	35.89	340.14	0.11				376.14
2. Industrial Processes	50,102.06	133.09	1,624.72	11,742.22	7,315.75	4,910.86	75,828.70
A. Mineral Products	47,005.76	NA,NO	NA,NO				47,005.76
B. Chemical Industry	2,918.74	115.93	1,624.72	NA	NA	NA	4,659.40
C. Metal Production	177.55	17.16	NO	IE,NE	14.82	1,091.08	1,300.62
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				938.25	879.14	1,366.36	3,183.75
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				10,803.97	6,421.79	2,453.41	19,679.17
G. Other	NO	NO	NO	NA,NO	NA,NO	NO	NA,NO
3. Solvent and Other Product Use	NA,NE		242.34				242.34
4. Agriculture		15,211.28	11,222.29				26,433.56
A. Enteric Fermentation		6,999.93					6,999.93
B. Manure Management		2,438.80	4,756.40				7,195.20
C. Rice Cultivation		5,707.49					5,707.49
D. Agricultural Soils ⁽³⁾		NA	6,444.86				6,444.86
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		65.06	21.03				86.09
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-84,977.56	2.44	13.12				-84,961.99
A. Forest Land	-86,436.29	2.44	0.25				-86,433.60
B. Cropland	295.00	NE,NO	12.87				307.88
C. Grassland	-338.54	NE,NO	NE,NO				-338.54
D. Wetlands	23.40	NE,NO	NE,NO				23.40
E. Settlements	510.88	NE,NO	NE,NO				510.88
F. Other Land	737.66	NO	NO				737.66
G. Other	230.34	NA,NE	NA,NE				230.34
6. Waste	13,655.17	5,828.00	3,253.39				22,736.56
A. Solid Waste Disposal on Land	NA,NE,NO	4,301.21					4,301.21
B. Waste-water Handling		1,383.85	1,165.75				2,549.60
C. Waste Incineration	13,132.81	13.29	1,972.81				15,118.91
D. Other	522.36	129.65	114.83				766.84
7. Other (as specified in Summary 1.A)	NA,NO	NA,NO	NA,NO	NA	NA	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	38,991.92	49.94	356.50				39,398.36
Aviation	19,964.61	11.87	195.93				20,172.40
Marine	19,027.31	38.07	160.58				19,225.96
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	21,976.71						21,976.71
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,333,586.60
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,248,624.61

⁽¹⁾ For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

⁽²⁾ Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

⁽³⁾ Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

⁽⁴⁾ See footnote 8 to table Summary 1.A.

A9.18. Emissions and Removals in 2007

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS
(Sheet 1 of 1)Inventory 2007
Submission 2012 v1.1
JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,212,153.52	22,056.88	22,812.39	13,279.24	6,411.99	4,407.45	1,281,121.49
1. Energy	1,232,951.02	1,290.80	7,659.82				1,241,901.64
A. Fuel Combustion (Sectoral Approach)	1,232,913.49	874.60	7,659.70				1,241,447.80
1. Energy Industries	446,853.32	44.88	2,181.76				449,079.96
2. Manufacturing Industries and Construction	370,254.58	370.10	2,119.28				372,743.96
3. Transport	237,830.98	207.63	3,002.14				241,040.75
4. Other Sectors	177,974.62	252.00	356.52				178,583.13
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	37.53	416.20	0.12				453.84
1. Solid Fuels	NE,NO	51.48	NE,NO				51.48
2. Oil and Natural Gas	37.53	364.72	0.12				402.37
2. Industrial Processes	49,344.83	134.15	860.18	13,279.24	6,411.99	4,407.45	74,437.85
A. Mineral Products	46,142.38	NA,NO	NA,NO				46,142.38
B. Chemical Industry	2,990.43	116.85	860.18	NA	NA	NA	3,967.46
C. Metal Production	212.02	17.30	NO	IE,NE	14.69	1,089.34	1,333.35
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				497.61	783.02	1,198.82	2,479.45
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				12,781.64	5,614.28	2,119.29	20,515.20
G. Other	NO	NO	NO	NA,NO	NA,NO	NO	NA,NO
3. Solvent and Other Product Use	NA,NE		159.95				159.95
4. Agriculture		15,066.31	11,028.40				26,094.71
A. Enteric Fermentation		6,974.38					6,974.38
B. Manure Management		2,376.23	4,773.35				7,149.58
C. Rice Cultivation		5,652.17					5,652.17
D. Agricultural Soils ⁽³⁾		NA	6,234.73				6,234.73
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		63.53	20.32				83.85
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-84,150.27	2.04	11.70				-84,136.52
A. Forest Land	-85,228.55	2.04	0.21				-85,226.30
B. Cropland	258.97	NE,NO	11.50				270.47
C. Grassland	-314.91	NE,NO	NE,NO				-314.91
D. Wetlands	27.70	NE,NO	NE,NO				27.70
E. Settlements	228.49	NE,NO	NE,NO				228.49
F. Other Land	553.07	NO	NO				553.07
G. Other	324.96	NA,NE	NA,NE				324.96
6. Waste	14,007.95	5,563.58	3,092.33				22,663.86
A. Solid Waste Disposal on Land	NA,NE,NO	4,053.98					4,053.98
B. Waste-water Handling		1,340.77	1,144.50				2,485.27
C. Waste Incineration	13,446.75	12.40	1,809.28				15,268.43
D. Other	561.20	156.43	138.55				856.18
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA	NA	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	37,259.15	48.74	339.71				37,647.60
Aviation	18,358.58	10.91	180.16				18,549.66
Marine	18,900.57	37.83	159.54				19,097.94
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	22,957.60						22,957.60
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,365,258.01
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,281,121.49

(1) For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

(2) Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

(3) Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

(4) See footnote 8 to table Summary I.A.

A9.19. Emissions and Removals in 2008

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS

(Sheet 1 of 1)

Inventory 2008

Submission 2012 v1.1

JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,134,499.75	21,542.50	22,829.48	15,298.30	4,617.89	3,795.22	1,202,583.13
1. Energy	1,152,455.61	1,273.94	7,354.68				1,161,084.24
A. Fuel Combustion (Sectoral Approach)	1,152,417.77	865.65	7,354.56				1,160,637.99
1. Energy Industries	420,263.03	43.62	2,118.33				422,424.97
2. Manufacturing Industries and Construction	335,619.40	367.12	2,050.95				338,037.48
3. Transport	228,099.17	191.44	2,843.45				231,134.06
4. Other Sectors	168,436.17	263.47	341.84				169,041.47
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	37.85	408.29	0.12				446.26
1. Solid Fuels	NE,NO	45.83	NE,NO				45.83
2. Oil and Natural Gas	37.85	362.46	0.12				400.43
2. Industrial Processes	45,738.97	121.48	1,262.15	15,298.30	4,617.89	3,795.22	70,834.01
A. Mineral Products	43,009.11	NA,NO	NA,NO				43,009.11
B. Chemical Industry	2,574.10	106.46	1,262.15	NA	NA	NA	3,942.70
C. Metal Production	155.77	15.02	NO	IE,NE	14.67	652.47	837.94
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				701.41	523.80	1,288.21	2,513.42
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				14,596.89	4,079.42	1,854.54	20,530.85
G. Other	NO	NO	NO	NA,NO	NA,NO	NO	NA,NO
3. Solvent and Other Product Use	NA,NE		129.10				129.10
4. Agriculture		14,875.64	11,034.46				25,910.10
A. Enteric Fermentation		6,913.14					6,913.14
B. Manure Management		2,302.01	5,019.17				7,321.18
C. Rice Cultivation		5,598.59					5,598.59
D. Agricultural Soils ⁽³⁾		NA	5,995.66				5,995.66
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		61.90	19.63				81.53
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-78,706.71	21.73	10.58				-78,674.40
A. Forest Land	-79,927.73	21.73	2.21				-79,903.79
B. Cropland	224.16	NE,NO	8.37				232.53
C. Grassland	-302.63	NE,NO	NE,NO				-302.63
D. Wetlands	15.67	NE,NO	NE,NO				15.67
E. Settlements	143.84	NE,NO	NE,NO				143.84
F. Other Land	834.33	NO	NO				834.33
G. Other	305.63	NA,NE	NA,NE				305.63
6. Waste	15,011.87	5,249.71	3,038.50				23,300.08
A. Solid Waste Disposal on Land	NA,NE,NO	3,758.54					3,758.54
B. Waste-water Handling		1,322.45	1,160.89				2,483.35
C. Waste Incineration	14,481.46	11.98	1,738.79				16,232.23
D. Other	530.41	156.73	138.81				825.95
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA	NA	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	34,849.64	45.09	318.18				35,212.92
Aviation	17,517.99	10.41	171.92				17,700.32
Marine	17,331.65	34.68	146.27				17,512.60
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	21,597.88						21,597.88
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,281,257.54
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,202,583.13

⁽¹⁾ For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

⁽²⁾ Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

⁽³⁾ Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

⁽⁴⁾ See footnote 8 to table Summary I.A.

A9.20. Emissions and Removals in 2009

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS
(Sheet 1 of 1)Inventory 2009
Submission 2012 v1.1
JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,070,380.54	20,889.20	22,580.45	16,554.17	3,267.84	1,851.27	1,135,523.47
1. Energy	1,089,177.47	1,227.40	7,011.03				1,097,415.91
A. Fuel Combustion (Sectoral Approach)	1,089,142.33	833.14	7,010.92				1,096,986.39
1. Energy Industries	385,896.01	41.92	2,032.04				387,969.97
2. Manufacturing Industries and Construction	318,978.00	363.72	1,977.52				321,319.24
3. Transport	222,768.36	179.41	2,671.62				225,619.40
4. Other Sectors	161,499.96	248.08	329.75				162,077.79
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	35.15	394.26	0.11				429.52
1. Solid Fuels	NE,NO	46.25	NE,NO				46.25
2. Oil and Natural Gas	35.15	348.01	0.11				383.27
2. Industrial Processes	40,313.87	109.60	1,559.50	16,554.17	3,267.84	1,851.27	63,656.24
A. Mineral Products	37,713.68	NA,NO	NA,NO				37,713.68
B. Chemical Industry	2,488.20	96.64	1,559.50	NA	NA	NA	4,144.34
C. Metal Production	111.99	12.96	NO	IE,NE	11.02	239.00	374.97
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				222.14	399.48	260.51	882.13
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				16,332.03	2,857.34	1,351.76	20,541.13
G. Other	NO	NO	NO	NA,NO	NA,NO	NO	NA,NO
3. Solvent and Other Product Use	NA,NE		120.50				120.50
4. Agriculture		14,624.76	10,959.90				25,584.66
A. Enteric Fermentation		6,773.41					6,773.41
B. Manure Management		2,247.27	5,247.45				7,494.72
C. Rice Cultivation		5,544.83					5,544.83
D. Agricultural Soils ⁽³⁾		NA	5,693.76				5,693.76
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		59.25	18.70				77.95
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-71,873.45	8.61	8.48				-71,856.36
A. Forest Land	-73,672.97	8.61	0.87				-73,663.48
B. Cropland	257.51	NE,NO	7.60				265.11
C. Grassland	-276.24	NE,NO	NE,NO				-276.24
D. Wetlands	22.57	NE,NO	NE,NO				22.57
E. Settlements	476.52	NE,NO	NE,NO				476.52
F. Other Land	1,049.04	NO	NO				1,049.04
G. Other	270.12	NA,NE	NA,NE				270.12
6. Waste	12,762.64	4,918.83	2,921.05				20,602.52
A. Solid Waste Disposal on Land	NA,NE,NO	3,517.23					3,517.23
B. Waste-water Handling		1,272.67	1,132.60				2,405.27
C. Waste Incineration	12,248.96	10.64	1,683.67				13,943.27
D. Other	513.69	118.29	104.77				736.75
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA	NA	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	30,686.03	39.77	280.09				31,005.89
Aviation	15,372.73	9.14	150.86				15,532.73
Marine	15,313.30	30.64	129.22				15,473.16
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	19,753.79						19,753.79
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,207,379.83
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,135,523.47

(1) For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

(2) Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

(3) Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

(4) See footnote 8 to table Summary I.A.

A9.21. Emissions and Removals in 2010

SUMMARY 2 SUMMARY REPORT FOR CO₂ EQUIVALENT EMISSIONS

(Sheet 1 of 1)

Inventory 2010

Submission 2012 v1.1

JAPAN

GREENHOUSE GAS SOURCE AND SINK CATEGORIES	CO ₂ ⁽¹⁾	CH ₄	N ₂ O	HFCs ⁽²⁾	PFCs ⁽²⁾	SF ₆ ⁽²⁾	Total
	CO ₂ equivalent (Gg)						
Total (Net Emissions)⁽¹⁾	1,118,759.78	20,445.22	22,073.61	18,256.50	3,405.25	1,862.42	1,184,802.79
1. Energy	1,137,584.04	1,217.89	6,809.57				1,145,611.50
A. Fuel Combustion (Sectoral Approach)	1,137,550.89	842.17	6,809.47				1,145,202.52
1. Energy Industries	406,096.17	45.53	2,005.11				408,146.81
2. Manufacturing Industries and Construction	342,609.09	372.81	1,941.96				344,923.85
3. Transport	224,943.47	167.73	2,526.29				227,637.49
4. Other Sectors	163,902.16	256.10	336.11				164,494.37
5. Other	NO	NO	NO				NO
B. Fugitive Emissions from Fuels	33.15	375.73	0.11				408.98
1. Solid Fuels	NE,NO	44.49	NE,NO				44.49
2. Oil and Natural Gas	33.15	331.24	0.11				364.49
2. Industrial Processes	41,177.28	118.85	1,077.74	18,256.50	3,405.25	1,862.42	65,898.05
A. Mineral Products	38,280.19	NA,NO	NA,NO				38,280.19
B. Chemical Industry	2,737.23	103.98	1,077.74	NA	NA	NA	3,918.95
C. Metal Production	159.86	14.87	NO	IE,NE	10.38	307.90	493.01
D. Other Production	IE						IE
E. Production of Halocarbons and SF ₆				128.34	200.24	198.37	526.96
F. Consumption of Halocarbons and SF ₆ ⁽²⁾				18,128.16	3,194.63	1,356.15	22,678.94
G. Other	NO	NO	NO	NA,NO	NA,NO	NO	NA,NO
3. Solvent and Other Product Use	NA,NE		98.95				98.95
4. Agriculture		14,387.40	11,112.22				25,499.61
A. Enteric Fermentation		6,673.27					6,673.27
B. Manure Management		2,205.06	5,475.35				7,680.42
C. Rice Cultivation		5,451.67					5,451.67
D. Agricultural Soils ⁽³⁾		NA	5,618.74				5,618.74
E. Prescribed Burning of Savannas		NO	NO				NO
F. Field Burning of Agricultural Residues		57.39	18.12				75.52
G. Other		NO	NO				NO
5. Land Use, Land-Use Change and Forestry⁽¹⁾	-73,187.60	2.12	6.40				-73,179.09
A. Forest Land	-76,676.91	2.12	0.22				-76,674.57
B. Cropland	452.41	NE,NO	6.18				458.59
C. Grassland	-215.86	NE,NO	NE,NO				-215.86
D. Wetlands	82.13	NE,NO	NE,NO				82.13
E. Settlements	2,518.29	NE,NO	NE,NO				2,518.29
F. Other Land	382.22	NO	NO				382.22
G. Other	270.12	NA,NE	NA,NE				270.12
6. Waste	13,186.06	4,718.97	2,968.73				20,873.76
A. Solid Waste Disposal on Land	NA,NE,NO	3,270.04					3,270.04
B. Waste-water Handling		1,269.65	1,131.61				2,401.26
C. Waste Incineration	12,657.57	10.46	1,687.59				14,355.62
D. Other	528.50	168.82	149.53				846.84
7. Other (as specified in Summary I.A)	NA,NO	NA,NO	NA,NO	NA	NA	NA,NO	NA,NO
Memo Items:⁽⁴⁾							
International Bunkers	31,214.79	39.58	285.77				31,540.14
Aviation	16,265.18	9.67	159.62				16,434.47
Marine	14,949.61	29.91	126.15				15,105.67
Multilateral Operations	NO	NO	NO				NO
CO₂ Emissions from Biomass	32,841.20						32,841.20
Total CO ₂ Equivalent Emissions without Land Use, Land-Use Change and Forestry							1,257,981.87
Total CO ₂ Equivalent Emissions with Land Use, Land-Use Change and Forestry							1,184,802.79

⁽¹⁾ For CO₂ from Land Use, Land-use Change and Forestry the net emissions/removals are to be reported. For the purposes of reporting, the signs for removals are always negative (-) and for emissions positive (+).

⁽²⁾ Actual emissions should be included in the national totals. If no actual emissions were reported, potential emissions should be included.

⁽³⁾ Parties which previously reported CO₂ from soils in the Agriculture sector should note this in the NIR.

⁽⁴⁾ See footnote 8 to table Summary I.A.

Abbreviations

1. Greenhouse Gasses

Table 1-1 6 Gasses controlled by Kyoto Protocol

Term	Gas
CO ₂	Carbon dioxide
CH ₄	Methane
N ₂ O	Nitrous oxide
HFCs	Hydrofluorocarbons
PFCs	Perfluorocarbons
SF ₆	Sulphur hexafluoride

Table 1-2 Indirect gasses and precursors

Term	Gas
NO _x	Sum of nitrogen oxide and nitrogen dioxide
CO	Carbon monoxide
NMVOC	Non-methane volatile organic compounds
SO ₂	Sulphur dioxide

2. Prefixes and Units

Table 2-1 Prefixes

Term	Prefix	Definition
P	peta	10 ¹⁵
T	tera	10 ¹²
G	giga	10 ⁹
M	mega	10 ⁶
k	kilo	10 ³
h	hecto	10 ²
da	deca	10 ¹
d	deci	10 ⁻¹
c	centi	10 ⁻²
m	milli	10 ⁻³
μ	micro	10 ⁻⁶

Table 2-2 Units

Term	Definition
m ³	cubic metre
l	litter
a	are
ha	hectare
g	gram
t	tonne
J	joule
°C	degree Celsius
yr	year
cap	capita
d.m.	dry matter

3. Notation Keys

Table 3-1 Notation Keys (See Annex5.)

Notation Key	Definition
NO	Not Occurring
NE	Not Estimated
NA	Not Applicable
IE	Included Elsewhere
C	Confidential

4. Other Abbreviations

Table 4-1 Abbreviations

	Terms	Definition	
A	AAU	Assigned Amount Units	
	ARD	Afforestation, Reforestation and Deforestation	
B	BFG	Blast Furnace Gas	
	BOD	Biochemical Oxygen Demand	
C	CFG	Converter Furnace Gas	
	CGER	Center for Global Environmental Research	
	CO ₂ eq.	Gas Emission in CO ₂ equivalent	
	COD	Chemical Oxygen Demand	
	COG	Coke Oven Gas	
	CRF	Common Reporting Format	
	CS-EF	Country-Specific Emission Factor	
	CY	Calendar Year	
	E	EF	Emission Factor
	F	FM	Forest Management
		FY	Fiscal Year
G	GCV	Gross Calorific Value	
	GHG	Greenhouse Gas	
	GIO	Greenhouse Gas Inventory Office	
	GPG	Good Practice Guidance	
	GPG (2000)	Good Practice Guidance and Uncertainty Management in National Greenhouse Gas Inventories (2000)	
	GPG-LULUCF	Good Practice Guidance for Land Use, Land-Use Change and Forestry	
	GWP	Global Warming Potential	
	I	IEA	International Energy Agency
		IEF	Implied Emission Factor
		IPCC	Intergovernmental Panel on Climate Change
J	JNGI	Japanese National GHG Inventory	
K	KP	Kyoto Protocol	
L	L.D.converter	Linz-Donawitz converter	
	LDG	Linz-Donawitz converter Gas	
	LNG	Liquefied Natural Gas	
	LPG	Liquefied Petroleum Gas	
	LTO	Landing and Take-off	
	LULUCF	Land-Use, Land-Use Change and Forestry	
	M	MAFF	Ministry of Agriculture, Forestry and Fisheries
		MDI	Metered Dose Inhalers
		METI	Ministry of Economy, Trade and Industry
		MOE	Ministry of the Environment
MOFA		Ministry of Foreign Affairs of Japan	
MIC		Ministry of Internal Affairs and Communications	
MLIT		Ministry of Land, Infrastructure and Transport and Tourism	
MSW		Municipal Solid Waste	
N		NCV	Net Calorific Value
		NFRDB	National Forest Resource DataBase
	NGL	Natural Gas Liquids	
	NIES	National Institute for Environmental Studies	
	NIR	National Inventory Report	
	Q	QA/QC	Quality Assurance / Quality Control
		QAWG	Quality Assurance Working Group
R	RDF	Refuse Derived Fuel	
	RPF	Refuse Paper and Plastic Fuel	
	RV	Revegetation	
T	THC	Total Hydrocarbon	
	TOE	Tonnes of Oil Equivalent	
U	UNFCCC	United Nations Framework Convention on Climate Change	

