IAEA Review of Safety Related Aspects of Handling ALPS-Treated Water at TEPCO's Fukushima Daiichi Nuclear Power Station

First Interlaboratory Comparison on the Determination of Radionuclides in the Marine Environment



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Published by the International Atomic Energy Agency in January 2024

EXECUTIVE SUMMARY

In 2021, the IAEA started its review of safety related aspects of handling ALPS (Advanced Liquid Processing System) treated water at TEPCO's Fukushima Daiichi Nuclear Power Station (FDNPS). Consistent with the request from the Government of Japan, the IAEA statutory functions and the mandate of the Task Force, the scope of the IAEA review is tailored to assessing safety related aspects of the implementation of Japan's *Basic Policy on Handling of ALPS Treated Water at the Tokyo Electric Power Company's Holdings' Fukushima Daiichi Nuclear Power Station* against the IAEA's Safety Standards. The current approach outlined in the Basic Policy is to conduct a series of controlled discharges of ALPS treated water into the sea ('batch discharges') over a period of approximately 30 years.

Consistent with the relevant IAEA's Safety Standards, TEPCO is required to determine the characteristics and activity of the ALPS treated water (e.g., through the radiological environmental impact assessment) to be discharged into the sea, and to establish and implement monitoring programmes to ensure that public exposure due to the discharges is adequately assessed and that the assessment is sufficient to verify and demonstrate compliance with the authorization granted by the NRA.

To conduct its safety review, the IAEA has organized the work of the Task Force into three main components, the assessment of protection and safety; regulatory activities and processes; and sampling, independent analysis and data corroboration. The latter activities include three elements:

- Sampling, analysis and interlaboratory comparison for ALPS treated water from the FDNPS.
- Sampling, analysis and interlaboratory comparison for environmental samples (e.g., seawater, fish) from the surrounding environment of FDNPS.
- Assessment of the capabilities of dosimetry service providers involved in the monitoring of internal and external radiation exposure of workers at FDNPS.

The IAEA's sampling, independent analysis and data corroboration activities also include a review of sampling and analytical methods used by TEPCO and any other relevant technical institutions.

The corroboration of source and environmental monitoring conducted by TEPCO and relevant Japanese authorities is based on interlaboratory comparisons (ILCs). ILCs, along with proficiency tests (PTs), are standard methods for laboratories to assess the quality of their measurement results in comparison with those of other participating laboratories, and to identify any potential improvements. PTs involve the evaluation of performance against pre-established criteria whereas ILCs involve the organization, performance, and evaluation of measurements on the same or similar items by two or more laboratories in accordance with predetermined conditions.

For this first ILC to corroborate the results of environmental monitoring under the IAEA's ALPS safety review, samples of seawater, sediment, fish and seaweed were taken in November 2022 from offshore locations and a fish market close to FDNPS.

Extensive monitoring of the marine environment around the FDNPS is conducted according to the Comprehensive Radiation Monitoring Plan (CRMP). TEPCO and other relevant Japanese authorities including the Nuclear Regulation Authority, the Ministry of the Environment, and the Japan Fisheries Agency, have responsibilities under the CRMP. Enhancements to the CRMP to specifically address the discharge of ALPS treated water were introduced in April 2022. The resulting data provide a baseline of activity concentrations in the marine environment against which the any consequences of the discharge can be assessed. This ILC is based on samples collected in November 2022; as this was also

before the start of discharges of ALPS treated water, the ILC serves to corroborate the results of the baseline monitoring.

This publication reports the results of this first ILC. It describes the joint sampling mission undertaken to collect seawater, sediment, fish and seaweed samples; the analytical techniques used by participating laboratories – from Japan (participating on behalf of the Japanese authorities); the IAEA and a third party member laboratory of the IAEA ALMERA network (Analytical Laboratories for the Measurement of Environmental Radioactivity)¹; the measurement results and the statistical evaluation of the results.

In total, eight laboratories undertook analyses and reported results for the ILC. Analyses were undertaken by Japanese laboratories participating in marine monitoring relevant to the ALPS discharges within the CRMP and by the following two IAEA Nuclear Sciences and Applications Laboratories:

- IAEA Marine Environment Laboratories, Radiometrics Laboratory (RML), Monaco;
- Isotope Hydrology Laboratory (IHL), Vienna, Austria.

Additionally, under the coordination of the participating IAEA laboratories, a selected third-party laboratory, Korea Institute of Nuclear Safety (KINS), Republic of Korea, a member of the network of Analytical Laboratories for the Measurement of Environmental Radioactivity (ALMERA) with demonstrable competence in the methods required, also conducted analyses of samples as an ILC participant.

The results of the analyses undertaken at each laboratory were reported to the IAEA. For results that could be intercompared (i.e. for radionuclides for which activity concentrations above detection limits were reported by at least two laboratories) a statistical evaluation to assess agreement was carried out by the IAEA. The results are presented in tables and charts in this report.

Although some discrepancies were found, this was to be expected given that several of the analytical methods required were complex (e.g. for OBT, TFWT and ¹⁴C in fish) and relatively new to some participating laboratories, being implemented specifically to assess the ALPS treated water discharges. Furthermore, higher uncertainties are usually associated with complex analyses of low, and close to detection limit, levels. However, no order of magnitude variations were identified, and no systemic deviations between the results reported by Japanese laboratories and those reported by the IAEA and the ALMERA member laboratory. Therefore, despite the discrepancies, the key findings of this ILC are:

- Japanese laboratories have demonstrated a high level of accuracy in their measurements and technical competence.
- Sample collection procedures follow the appropriate methodological standards required to obtain representative samples.

The IAEA notes that these findings provide confidence in Japan's capability for undertaking accurate and precise measurements related to the discharge of ALPS treated water.

Additional ILCs will be conducted in the future for ALPS treated water and environmental samples, as well as occupational radiation protection². Future ILCs, now that discharges of ALPS treated water have started, will allow for an assessment of Japan's capability to evaluate any changes in the levels of relevant radionuclides in the marine environment, relative to the baseline.

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¹ More information on the ALMERA network is available from the following website: https://nucleus.iaea.org/sites/ReferenceMaterials/Pages/ALMERA.aspx

² The IAEA is currently in discussions with other laboratories to participate in future ILCs.

Furthermore, the corroboration of environmental monitoring complements a separate project – NA3/38 Marine Monitoring: Confidence Building and Data Quality Assurance – addressing the quality of data from marine monitoring undertaken in Japan following the accident at FDNPS. Through project NA3/38, which has been implemented since 2014, the IAEA is assisting the Government of Japan in ensuring that sea area monitoring carried out under the regularly updated CRMP is comprehensive, credible and transparent and is helping to build confidence of the stakeholders in the accuracy and quality of the marine monitoring data. Within project NA3/38, the IAEA has organized a series of ILCs and PTs to test the sampling and analytical performance of the Japanese laboratories for the analysis of radionuclides in seawater, sediment, fish and seaweed samples. Data from this project are available online at:

https://www.iaea.org/about/organizational-structure/department-of-nuclear-sciences-and-applications/division-of-iaea-marine-environment-laboratories/marine-monitoring-confidence-building-and-data-quality-assurance.

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1. INTRODUCTION

1.1. BACKGROUND

The corroboration of a representative subset of the radioactivity measurement results reported by TEPCO and relevant Japanese authorities during both the pre-operational and the operational phases of discharge of ALPS treated water to the sea, and a review of the methods for related sampling and analysis used by TEPCO and relevant Japanese authorities is being undertaken by the IAEA. This corroboration provides an independent check of the veracity of the radiological data resulting from source and environmental monitoring programmes related to the ALPS discharges upon which the safety related aspects of the discharges of ALPS treated water are being evaluated. The scope of the corroboration includes an independent check of the radiological characterization of the discharges. A primary objective is to promote transparency and provide sound information to enable interested parties to evaluate the radiological data used as the basis for planning the discharges of ALPS treated water into the sea.

The IAEA corroboration of source and environmental monitoring related to discharges of ALPS treated water from FDNPS is comprised of three distinct elements (see also Figure 1):

- 1. Review of sampling and analytical methods for source and environmental monitoring related to ALPS treated water at FDNPS used by TEPCO and relevant Japanese authorities.
- 2. Corroboration of source monitoring undertaken by TEPCO, including a comprehensive radiological characterization of ALPS treated water samples.
- 3. Corroboration of environmental monitoring undertaken by TEPCO and relevant Japanese authorities.

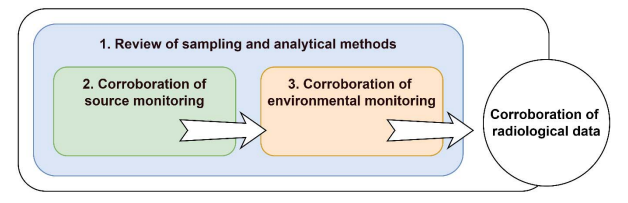


FIG. 1: A schematic overview of the elements of the corroboration being undertaken by the IAEA laboratories and the links between these elements.

The corroboration of source and environmental monitoring is based on interlaboratory comparisons (ILCs). ILCs, along with proficiency tests (PTs), are standard methods for laboratories to assess the quality of their measurement results in comparison with those of other participating laboratories, and to identify any potential improvements. PTs involve the evaluation of performance against pre-established criteria whereas ILCs involve the organization, performance and evaluation of measurements on the same or similar items by two or more laboratories in accordance with predetermined conditions [1].

This publication reports the results of the first ILC to corroborate environmental monitoring. It describes the joint sampling mission undertaken to collect seawater, sediment, fish and seaweed samples; the analytical techniques used by participating laboratories – from Japan (participating on behalf of the Japanese authorities); the IAEA and a third -party member laboratory of the IAEA ALMERA network

(Analytical Laboratories for the Measurement of Environmental Radioactivity)³; the measurement results and the statistical evaluation of the results.

Extensive monitoring of the marine environment around the FDNPS is conducted according to the Comprehensive Radiation Monitoring Plan (CRMP) [2]. TEPCO and other relevant Japanese authorities have responsibilities under the CRMP. This plan defines sampling locations, frequency of sampling, target detection limits and responsibilities of the organizations involved. The monitoring comprises sampling and analysis of seawater to different depths, sediment and marine biota (fish, shellfish and seaweed) The aim of this plan includes ensuring a comprehensive overview of the radiological situation in the marine environment and providing an adequate basis for assessments of radiation exposures from marine pathways. In practice, sampling and analysis are often carried out by contracted laboratories but TEPCO and the other relevant Japanese authorities, as defined in the CMRP, have responsibility for reporting the results of the monitoring with which they have been tasked.

Enhancements to the CRMP to specifically address the discharge of ALPS treated water were introduced in March 2022. These included the monitoring of ³H in seawater at increased frequencies plus monitoring quarterly for 'seven major radionuclides', namely ⁶⁰Co, ⁹⁰Sr, ¹⁰⁶Ru, ¹²⁵Sb, ¹²⁹I, ¹³⁴Cs and ¹³⁷Cs. Monitoring of organically bound tritium (OBT), tissue free-water tritium (TFWT) and ¹⁴C in fish and ¹²⁹I is also being undertaken.

This monitoring started more than one year before the first discharges of ALPS treated water. The results provide a baseline of activity concentrations in the marine environment against which the any consequences of the discharge can be assessed. This ILC is based on samples collected in November 2022; as this was also before the start of discharges of ALPS treated water, the ILC serves to corroborate the results of the baseline monitoring.

The samples were taken at sampling locations defined in the CRMP, with IAEA observation, using the same techniques used for routine monitoring. Similarly, the radionuclides analysed for each sample are defined in the CRMP and the analyses were carried out by the participating Japanese laboratories using the same methods as those used for routine monitoring.

The IAEA wishes to thank all the participating laboratories that took part in this interlaboratory comparison and the Japanese organisations that provided logistical support. The IAEA is also grateful to the Government of Monaco for its support. The IAEA personnel responsible for this publication was P. McGinnity of the IAEA Nuclear Science and Applications Department.

1.2. OBJECTIVE

This publication reports the results of the first ILC to corroborate environmental monitoring related to the discharge of ALPS treated water at FDNPS.

1.3. SCOPE

This publication reports all aspects of the first ILC to corroborate environmental monitoring related to the discharge of ALPS treated water at FDNPS including: the ILC design; participating laboratories; the methods employed for sampling and for distribution of the samples between participating laboratories; the analytical methods used by each participating laboratory to determine activity concentrations of

³ More information on the ALMERA network is available from the following website: https://nucleus.iaea.org/sites/ReferenceMaterials/Pages/ALMERA.aspx

radionuclides in the samples; the methodology employed for the statistical evaluation of the results; and the results and conclusions.

1.4. STRUCTURE

This publication contains reports the participating laboratories (Section 2); the methods employed for the sample collection and pre-treatment, and for the distribution of the samples between participating laboratories (Section 3), and the analytical methods used by each participating laboratory to determine activity concentrations of radionuclides in the samples (Section 4). The methodology employed for the statistical evaluation of the results is described in Section 5. The results of the ILCs are presented in Section 6 and conclusions in Section 7. The results of the ILCs are also presented in charts in an Appendix.

2. PARTICIPATING LABORATORIES

In total, eight laboratories participated in the ILC: Six from Japan (participating on behalf of TEPCO and other relevant Japanese authorities having responsibilities under the CRMP); the IAEA; and a laboratory from the Republic of Korea, a member laboratory of the IAEA ALMERA network (Analytical Laboratories for the Measurement of Environmental Radioactivity).

ALMERA is a network comprising more than 190 member laboratories globally. It provides a platform for maintaining and developing capability on the determination of radionuclides in air, water, soil, sediment and vegetation that can be used for both routine and environmental emergency monitoring in the IAEA Member States.

IAEA analyses were undertaken at two participating IAEA Nuclear Sciences and Applications Laboratories:

- IAEA Marine Environment Laboratories, Radiometrics Laboratory (RML), Monaco;
- Isotope Hydrology Laboratory (IHL), Vienna, Austria.

The participating laboratories are presented in Table 1, and participation of each in specific analyses in Table 2.

TABLE 1. LABORATORIES PARTICIPATING IN ILC 2022

Identifier	Participant
IAEA	IAEA Laboratories, Austria and Monaco
JCAC	Japan Chemical Analysis Center, Chiba, Japan
KAKEN ¹	KAKEN Co. Ltd., Ibaraki, Japan
KANSO	KANSO TECHNOS Co. Ltd., Osaka, Japan
KEEA	Kyushu Environmental Evaluation Association, Fukuoka, Japan
KINS	Korea Institute of Nuclear Safety, Daejeon, Republic of Korea
MERI	Marine Ecology Research Institute, Chiba, Japan
TPT ¹	Tokyo Power Technology Ltd., Fukushima, Japan

Note:

¹ Laboratories conducting analysis under contract from TEPCO.

TABLE 2. OVERVIEW OF ILC 2022

				H					
Sample type	Nuclide	< IAEA	JCAC	KAKEN	KANSO	KEEA	KINS	MERI	TPT
	³ H	7	√	√	√	√	√	×	×
	⁶⁰ Co	✓	✓	×	×	×	✓	×	×
	⁹⁰ Sr	✓	✓	×	×	×	✓	×	×
C 4	¹⁰⁶ Ru	✓	✓	×	×	×	✓	×	×
Seawater	¹²⁵ Sb	✓	✓	×	×	×	✓	×	×
	129I	✓	✓	×	×	×	✓	×	×
	¹³⁴ Cs	✓	✓	×	×	×	✓	×	×
	¹³⁷ Cs	✓	✓	×	×	×	✓	×	×
Cadimant	¹³⁴ Cs	✓	✓	×	×	×	✓	×	✓
Sediment	¹³⁷ Cs	✓	✓	×	×	×	✓	×	✓
	³ H (OBT)	✓	×	×	✓	✓	✓	×	×
Fish (market)	³ H (TFWT)	✓	×	×	✓	✓	✓	✓	×
	¹⁴ C	✓	✓	×	×	×	✓	×	×
	³ H (OBT)	✓	✓	✓	×	✓	✓	×	×
Fish (fixed net)	³ H (TFWT)	✓	✓	✓	×	✓	✓	×	×
	¹⁴ C	✓	✓	×	×	×	✓	×	×
Seaweed	¹²⁹ I	✓	✓	×	×	×	✓	×	✓

Note:

¹ The symbol \checkmark indicates that the laboratory participated in the specific analysis (sample type and radionuclide), the symbol * indicates that it did not participate.

3. SAMPLE COLLECTION, PRETREATMENT AND DISTRIBUTION

Samples of seawater, sediment, fish and seaweed were collected in November 2022 from offshore locations and a fish market close to FDNPS. The sampling locations are shown in Figure 2. The coordinates of these sampling locations are provided in Table 3.

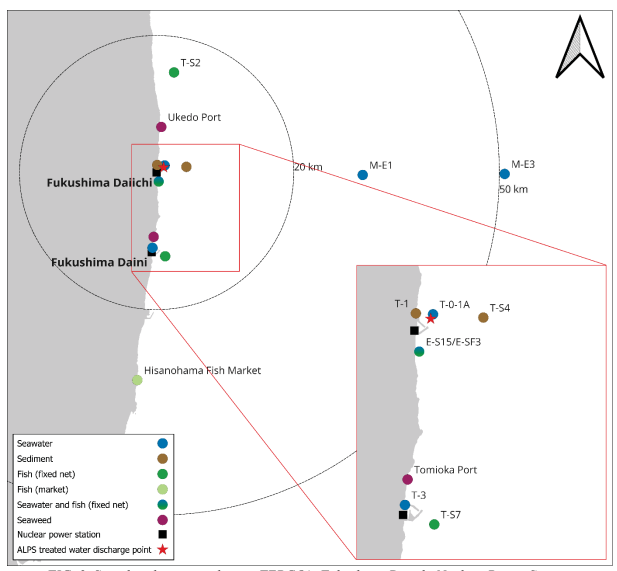


FIG. 2. Sampling locations close to TEPCO's Fukushima Daiichi Nuclear Power Station.

TABLE 3. COORDINATES OF SAMPLING LOCATIONS

Sampling location	Latitude (N)	Longitude (E)	Samples
E-S15/E-SF3 ¹	37.4094	141.0368	Seawater and fish
			(fixed net)
T-0-1A	37.4306	141.0467	Seawater
T-3	37.3222	141.0264	Seawater
M-E1	37.4170	141.3730	Seawater
M-E3	37.4170	141.6070	Seawater
T-1	37.4311	141.0344	Sediment
T-S4	37.4286	141.0825	Sediment
T-S2	37.5528	141.0625	Fish (fixed net)
T-S7	37.3111	141.0472	Fish (fixed net)
Hisanohama Fish	37.1484	141.0010	Fish (market)
Market			
Ukedo Port	37.4811	141.0413	Seaweed
Tomioka Port	37.3367	141.0283	Seaweed

Note:

3.1. SEAWATER

Surface seawater samples were collected at five sampling locations (E-S15, T-0-1A, T-3, M-E1 M-E3) offshore TEPCO's Fukushima Daiichi Nuclear Power Station (Figure 2 and Table 3). The samples were collected between 7 and 10 November 2022 from each sampling location for subsequent analysis for ³H and, for E-S15, for 'seven major radionuclides', namely ⁶⁰Co, ⁹⁰Sr, ¹⁰⁶Ru, ¹²⁵Sb, ¹²⁹I, ¹³⁴Cs and ¹³⁷Cs. Separate samples from each location were provided to participating laboratories for ³H and ¹²⁹I; for radiocaesium (¹³⁴Cs and ¹³⁷Cs); for ⁹⁰Sr and for other gamma-emitting radionuclides (⁶⁰Co, ¹⁰⁶Ru, ¹²⁵Sb).

For ³H, four laboratories planned to participate in the analyses of samples from each sampling location, except for E-S15 for which the number was three.

For T-0-1A, T-3, M-E1 and M-E3 the sample collection and distribution methods were:

- Separate 2 L containers were filled, one at a time, from a Niskin sampler re-filled several times, resulting in a total of four 2 L samples from each sampling station.
- One 2 L sample was provided to each laboratory.

For E-S15 the sample collection and distribution methods were:

- A 400 L plastic container with four valves was first filled with seawater. This container was also
 used to sample seawater from this location for the other radionuclides. As this container cannot be
 filled to full capacity, two separate fills were required to facilitate provision of the required sample
 volume to all participants.
- Separate 2 L containers were filled, two at a time, from the four valves, resulting in a total of four 2 L samples from this sampling location.
- One 2 L sample was provided to each laboratory.
- The same sample was planned to provide for ¹²⁹I analyses.

The seawater sampling procedure and the distribution matrix for ³H for E-S15 are shown in Table 4.

¹ The sampling location is the same. However, for seawater (E-S15) it is a point while for fish (E-SF3) the sampling extends along the length of the fixed net.

TABLE 4. SAMPLE DISTRIBUTION BETWEEN FOUR LABORATORIES (3H)

Valve number	1	2	3	4
Seawater sample codes	TR-1	TR-2	TR-3	TR-4
Distribution pattern of the participating laboratories	IAEA (RML)	IAEA (IHL)	KINS	JCAC

Three laboratories planned to participate in the analyses for ⁹⁰Sr; ¹³⁴Cs and ¹³⁷Cs; and other gamma-emitting radionuclides (⁶⁰Co, ¹⁰⁶Ru, ¹²⁵Sb) for sampling location E-S15. The collection and distribution methods were:

- From the same 400 L plastic container from which the samples to be analysed for ³H and ¹²⁹I were taken, separate 20 L cubitainers were filled simultaneously from each of the four valves. Six cubitainers were filled from each valve, resulting in a total of 24 20 L samples from each sampling station.
- Each sample was acidified to pH 1–2 with concentrated HCl.
- Four 20 L samples from each sampling location were provided to each laboratory planning to participate in analyses for ⁹⁰Sr; ¹³⁴Cs and ¹³⁷Cs; and other gamma-emitting radionuclides (⁶⁰Co, ¹⁰⁶Ru, ¹²⁵Sb).

The seawater sampling procedure and distribution matrix, meant to ensure the homogenisation of the samples, are shown in Table 5.

TABLE 5. SAMPLE DISTRIBUTION BETWEEN FOUR LABORATORIES (90Sr; 134Cs AND 137Cs; AND OTHER GAMMA-EMITTING RADIONUCLIDES)

Valve number	1	2	3	4
	1-1-1	1-2-1	1-3-1	1-4-1
Seawater sample codes	1-1-2	1-2-2	1-3-2	1-4-2
	2-1-1	2-2-1	2-3-1	2-4-1
	2-1-2	2-2-2	2-3-2	2-4-2
Distribution pattern of				
the participating	IAEA (RML)	Archive sample	KINS	JCAC
laboratories				

3.2. SEDIMENT

Sediment samples were collected using a grab sampler on 7 November 2022 offshore from TEPCO's Fukushima Daiichi Nuclear Power Station at sampling locations T-1 and T-S4 (Figure 2 and Table 3). The samples were subsequently oven-dried at 105°C on large stainless-steel trays, crushed using stainless-steel spatulae, and sieved through a 2-mm mesh sieve at the JCAC laboratory in Chiba. No grinding was required prior to sieving due to the sandy nature of the sediments. The fraction with grain size <2 mm was sieved to ≤250 μm, then placed in a plastic bag and mixed thoroughly to ensure homogeneity. An incremental division method was used for the provision of samples to participating laboratories. Each sample was divided into two aliquots using a splitter; one aliquot was archived and the second one was further divided until the required sample weight for each laboratory was attained. The sequence of division of each sample depended on the total mass of the sieved material. The samples were then bottled in 500 mL plastic bottles and shipped to the IAEA Marine Environment Laboratories in Monaco where their ¹³⁷Cs homogeneity was checked using gamma-ray spectrometry with High Purity Germanium (HPGe) detectors. Approximately 350 g of homogeneous dried sediment from each station was then shipped to each participating laboratory for analysis for ¹³⁴Cs and ¹³⁷Cs.

3.3. FISH (MARKET)

Six batches of frozen fish, one each of olive flounder (*Paralichthys olivaceus*), whitespotted conger (*Conger myriaster*), crimson sea bream (*Evynnis tumifrons*), redwing searobin (*Lepidotrigla microptera*), shotted halibut (*Eopsetta grigorjewi*) and willowy flounder (*Tanakius kitaharai*), were collected from the fish market at Hisanohama Port on 7 and 10 November 2022. The fish species were caught by pole and line fishing or bottom trawling on the same date in the vicinity of TEPCO's Fukushima Daiichi Nuclear Power Station at depths between 40 and 130 m. Each batch of fish was divided into two sub-batches: one for provision of samples to be analysed for ³H – organically bound tritium (OBT) and tissue free water tritium (TFWT) – and one for ¹⁴C. Special care was taken to ensure that the fish to be analysed for tritium were isolated from ice used for cooling during shipment and, subsequently, from tap water during pre-treatment.

Each sub-batch of fish of each species was prepared separately by fileting, homogenising the muscle tissue and then dividing into separate samples at the MERI laboratory in Onjuku on 10 and 11 November 2022. These were frozen and shipped to the participating laboratories. Each laboratory received approximately 1 kg of each species of fish for analysis for ³H and 0.5 kg for analysis for ¹⁴C.

3.4. FISH (FIXED NET)

Fish samples were collected at three sampling locations (E-SF3, T-S2 and T-S7) offshore TEPCO's Fukushima Daiichi Nuclear Power Station (Figure 2 and Table 3). The fish were caught using fixed gill nets at each location. Nets were installed on 8 November 2022 and hauled in on 9 November 2022. One batch of olive flounder (*Paralichthys olivaceus*) was collected at each location. As for the fish collected from market, each batch was divided into two sub-batches: one for provision of samples to be analysed for ³H (OBT and TFWT) and one for ¹⁴C.

Each sub-batch of fish of each species was prepared separately by fileting, homogenising the muscle tissue and then dividing into separate samples. The samples from T-S2 and T-S7 were processed at TEPCO's laboratory in Fukushima on 9 November 2022 and the samples from E-SF3 at the JCAC laboratory in Chiba on 11 November 2022. These were frozen and shipped to the participating laboratories. Each laboratory received approximately 1 kg of each species of fish for analysis for ³H and 0.5 kg for analysis for ¹⁴C.

3.5. SEAWEED

Seaweed samples were collected at Ukedo Port and Tomioka Port offshore TEPCO's Fukushima Daiichi Nuclear Power Station (Figure 2 and Table 3). At Ukedo Port (sampling location of the Ministry of the Environment), one sample of arame (*Eisenia bicyclis*) was collected by a diver. The seaweed was prepared by first removing the leaves and discarding the stems, homogenising and then dividing into separate samples at the JCAC laboratory in Chiba on 11 November 2022. These were frozen and shipped to the participating laboratories. Each laboratory received approximately 1 kg for analysis for ¹²⁹I.

At Ukedo Port and Tomioka Port (sampling locations of TEPCO) no seaweed was available to sample at the scheduled time during the sampling mission, due to hot weather in the preceding weeks. The decision was taken to use two seaweed samples that had been collected previously, sargassum (hondawara, *Sargassum Fuvellum*) from Tomioka Port and kelp (konbu, *Laminariacase*) from Ukedo Port. These samples had been freeze-dried soon after collection. Each laboratory received approximately 100 g of each dried seaweed. In order to facilitate reporting in Bq/kg fresh weight, the participating laboratories were provided with the masses of the seaweeds before and after drying.

4. ANALYTICAL METHODS

4.1. SEAWATER

Radionuclides in seawater were analysed by 7 laboratories participating in this ILC: from Japan, JCAC, KAKEN, KANSO and KEEA; IAEA; and KINS (see Tables 1 and 2).

4.1.1. ³H analysis

The IAEA used two methods of analysis for ³H in seawater, a low-level method employing mass spectrometry whereby the activity concentration of ³H in a sample is determined through the ingrowth of a decay product ³He under controlled conditions, and liquid scintillation counting (LSC) following purification and electrolytic enrichment of the seawater samples. The ³He ingrowth method was carried out at the IHL. The second method was performed at both IHL and MEL. The results reported were determined using the first method as the detection limit achievable was lower, typically of the order of 10 mBq/L.

For the ³He ingrowth method, a 100 mL aliquot of each seawater sample was added to an all-metal water sample container and de-gassed for one to two hours with a dedicated degassing unit to remove all pre-existing ³He. The degassed samples were stored for a period of five to six weeks to allow sufficient ³He-ingrowth and then counted on a Thermo Fisher Helix SFT split flight tube noble gas mass spectrometer. To ensure accurate measurements, the sensitivity of the mass spectrometer, as well as its temporal drift and linearity, were calibrated by analysing gas samples with known amounts of ³He and water samples with known ³H content [3]. The background of the mass spectrometer system was also determined.

In the second method, the samples were measured by liquid scintillation counting after purification by distillation and electrolytic enrichment. An ultra-low level liquid scintillation counter was used for the counting of an aliquot of the enriched and distilled sample mixed with a scintillation cocktail.

The other participating laboratories all analysed the seawater samples in a similar manner: purification by distillation followed by electrolytic enrichment, mixing with scintillation cocktail and measurement by LSC. Enrichment factors were typically 30 to 50 times the initial ³H activity concentration in the seawater. For the Japanese laboratories – JCAC, KAKEN, KANSO and KEEA – the specifics of the methods implemented complied with the "Tritium Analysis Method (Radiation Measurement Method Series 9)" published by the Japanese Ministry of Education, Culture, Sports, Science and Technology [4].

4.1.2. ⁹⁰Sr analysis

The IAEA used liquid-liquid extraction with di-(2-ethylhexyl)phosphoric acid (HDEHP) for the separation of yttrium from the seawater sample, while caesium was precipitated from the same sample by using ammonium molybdophosphate (AMP). The 90 Sr activity concentration was determined through the measurement of 90 Y (yttrium oxalate source) β activity using a proportional counter with an efficiency of up to 44%.

JCAC employed a cation-exchange resin column for pre-concentration of strontium from the seawater sample, followed by precipitation of carbonates and an additional cation exchange resin column for separation of calcium. 90 Y was removed by scavenging and, once the sample reached secular equilibrium, 90 Y was co-precipitated with iron hydroxide and then was measured using a low background β counter.

At KINS strontium pre-concentration of the 40 L seawater sample was carried out using a cation exchange resin. Eluted strontium was then recovered using strontium carbonate precipitation and then

strontium was purified again using fuming nitric acid. ⁹⁰Y and ⁹⁰Sr were determined by liquid scintillation counting in Cerenkov mode after allowing two weeks for the sample to reach secular equilibrium. The chemical yield was determined by ICP-OES (Inductively Coupled Plasma Optical Emission Spectroscopy).

4.1.3. ¹³⁴Cs and ¹³⁷Cs analysis

All participating laboratories (IAEA, JCAC and KINS) used AMP for the chemical separation of caesium from the seawater sample, followed by gamma-ray spectrometry using a HPGe (High Purity Germanium) detector.

4.1.4. Other gamma-emitting radionuclides (60Co, 106Ru and 125Sb) analysis

In all participating laboratories the samples were prepared in Marinelli beakers (1 L for the IAEA and KINS; 2 L for JCAC) and analysed without further treatment by gamma-ray spectrometry using HPGe detectors. ¹⁰⁶Ru was determined by gamma-ray spectrometry via decay of its progeny ¹⁰⁶Rh.

4.1.5. ¹²⁹I analysis

The IAEA analysed for ¹²⁹I in the seawater sample using gamma-ray spectrometry. The method is as described in the previous section for ⁶⁰Co, ¹⁰⁶Ru and ¹²⁵Sb.

At JCAC the sea water sampled was filtered, aliquoted and diluted for measurement of ¹²⁹I by ICP-MS (Inductively Coupled Plasma Mass Spectrometry). Rhenium was employed as an internal standard.

At KINS, after oxidation and reduction processes, the seawater sample was chemically separated using anion exchange resin. The elute was precipitated as PdI₂ using PdCl₂ solution, and the PdI₂ precipitate was filtered. Dried PdI₂ mounted on a Teflon planchette was measured by X-ray spectrometry for the determination of ¹²⁹I.

4.2. SEDIMENT

Radionuclides of interest in sediment samples were analysed by four laboratories participating in ILC 2022: from Japan, JCAC and TPT; IAEA; and KINS (see Tables 1 and 2).

4.2.1. ¹³⁴Cs and ¹³⁷Cs analysis

All participating laboratories analysed for ¹³⁴Cs and ¹³⁷Cs in sediment by gamma-ray spectrometry using HPGe detectors, following preparation of the samples in cylindrical containers.

4.3. FISH

Radionuclides of interest in fish samples collected from the fish market were analysed by six laboratories participating in ILC 2022: from Japan, JCAC, KANSO, KEEA and MERI; IAEA; and KINS (see Tables 1 and 2).

For the fish sampled at sea using fixed nets, radionuclides of interest were also analysed by five laboratories: from Japan, JCAC, KAKEN and TPT; IAEA; and KINS.

4.3.1. ³H (OBT and TFWT) analysis

The samples were vacuum freeze-dried at all laboratories. The tissue free water and dry materials were recovered, weighed and the drying rate (%) calculated.

At IAEA, the tissue free water was analysed for ³H using the ³He ingrowth method as described in section 4.1.1 for seawater.

At KINS, KANSO, KEEA and MERI the tissue free water recovered from the samples by vacuum freeze-drying was purified by reflux decomposition and distillation, mixed with scintillation cocktail, and counted by LSC to determine the ³H activity concentration.

JCAC and KAKEN implemented the same method but with additional steps: following distillation the ³H was concentrated by electrolytic enrichment and the samples were distilled again prior to counting by LSC. This method complied with the "Analysis Method of Tritium (Radiation Measurement Method Series 9)" [4].

For OBT, the dry fish material was first ground using a food processor at IAEA. It was then analysed for OBT using two separate analytical methods: the ³He ingrowth method and high-pressure combustion followed by LSC.

For the ³He ingrowth method, approximately 90g of dry material was placed in an all-metal sample container (diameter 13 cm x 21 cm) and degassed using a Turbo Molecular Pump. It was then stored for ³He ingrowth for a period of 7 to 9 weeks. At the end of this time, the sample container was connected to the ultra-high vacuum helium purification line attached to a Thermo Fisher Helix SFT split flight tube noble gas mass spectrometer to measure the ³He produced from the OBT over the ingrowth period.

For the second method, approximately 10 g of dry material was combusted in a Parr high pressure oxygen combustion vessel and a cold trap in the vacuum line was used to recover the combustion water. The process was repeated three times to collect approximately 15 mL of combustion water in total. The combustion water was distilled, mixed with a scintillation cocktail and counted using by LSC.

KINS used the same high pressure combustion method as IAEA for OBT.

At JCAC the dried component of each fish sample was burned under a flow of oxygen gas. The resulting water vapor was collected by the cooling trap as combustion water. At KANSO, KAKEN, KEEA and TPT combustion water was obtained from the dry material using a quartz tube furnace. The dried samples were inserted into quartz tubes and combusted at 650°C to 750°C in an electric furnace. the combustion water was collected in a cold trap. Copper oxide was used as a combustion catalyst.

Subsequent steps at all three laboratories were essentially the same: organic matter present in the combustion water was decomposed with refluxing. The sample was then distilled mixed scintillator cocktail and counted by LSC to determine the ³H activity concentration.

At all laboratories, conversion from Bq/L to Bq/kg fresh weight for the OBT measurements was based on the dry content calculated in the vacuum freezing process and the hydrogen content obtained by elemental analysis of dried sample.

4.3.2. ¹⁴C analysis

At KINS, freeze-dried samples were combusted to CO₂ which was then collected by bubbling into ammonium water (1:1). Calcium chloride was added to the solution and the precipitate CaCO₃ (calcium carbonate) retrieved. CaCO₃ (6 g) was reacted with hydrochloric acid water (1:1) on nitrogen purging to generate CO₂. The CO₂ was absorbed using Carbo-Sorb E, mixed with a scintillation cocktail (PermaFluor E+) in a Teflon vial and CO₂ counted by LSC.

The IAEA's samples were analysed using a similar technique by the Low-Level Radioactivity Measurements (LRM) Laboratory of the Belgian Nuclear Research Centre (SCK CEN), under contract. A distinction was that the scintillation cocktail was Pico-Fluor Plus. This method is accredited to ISO/IEC 17025 at SCK-CEN (accreditation number 015-TEST under the Belgian National Accreditation Body, BELAC).

JCAC used different methods for the fish sampled from market and for fish caught by fixed net. For the former, fish muscle was freeze-dried, ground to a powder and homogenised. Then, 5-10 g was thermally decomposed with pure oxygen in a combustion apparatus at 1000°C. CO₂ was collected by bubbling into ammonium water (1:1). Calcium chloride was added to the solution and the precipitate CaCO₃ (calcium carbonate) retrieved. CaCO₃ (25-27 mg) was reacted with pure phosphoric acid (4 mL) on a vacuum line to generate CO₂. Subsequently, a graphite target (>2 mg) was retrieved by flowing pure hydrogen into the CO₂ and loaded with a pre-purified Fe catalyst at 600°C for 6 hours. After removal of the Fe, pure graphite was measured by AMS for calculation of the ¹⁴C specific activity (Bq/g carbon) in the sample.

For fish caught by fixed net, JCAC a dry matter sample was burned under high oxygen pressure in a combustion device. The generated carbon dioxide was synthesized into benzene using a vacuum line. Then, 2 mL of the synthesized benzene was mixed with 0.5 mL of scintillation cocktail and counted by LSC.

4.4. SEAWEED

Radionuclides of interest in seaweed samples were analysed by four laboratories participating in ILC 2022: from Japan, JCAC and TPT; IAEA; and KINS (see Tables 1 and 2).

4.4.1. ¹²⁹I analysis

IAEA and KINS analysed the freeze-dried seaweed samples by gamma spectrometry using HPGe detectors.

At JCAC, the sample was freeze-dried, ground to a powder and homogenised. Then 0.5-1 g was added to a ceramic boat and mixed with V_2O_5 (Vanadium Oxide). Then the sample was placed in a quartz tube and heated at 1000°C under a flow of oxygen gas. The iodine released by heating was collected with a trap solution containing TMAH (tetramethyl ammonium hydroxide) and Na_2SO_3 (sodium sulfite). The trap solution was diluted for measurement of ^{129}I by ICP-MS. Rhenium was employed as an internal standard.

TPT freeze-dried each seaweed sample after washing with seawater. It was then crushed and dissolved in 25% TMAH solution. The TMAH solution was centrifuged and the resulting supernatant water filtered. This was weighed and the ¹²⁷I concentration measured to determine the total iodine content of the wet seaweed sample.

The TMAH solution was exchanged to 1M HNO₃ using an anion-exchanging resin. The ¹²⁹I/¹²⁷I isotopic ratio was then measured by triple quadrupole ICP-MS (ICP-QQQ) and multiplied by the total iodine content of the wet seaweed sample.

5. STATISTICAL EVALUATION OF THE RESULTS

The IAEA collected and evaluated the results reported by all ILC participants. The method used for the statistical evaluation depended on the number of results received for each sampling location, sample type and radionuclide.

If two or three measurement results above the detection limit were received, then one or three zeta tests [5] were performed. The zeta $\zeta_{i,j}$ test is defined as:

$$\zeta_{i,j} = \frac{x_i - x_j}{\sqrt{u_i^2 + u_j^2}} \tag{1}$$

where:

 x_i is the value of laboratory *i* (Bq *unit*⁻¹);

 x_i is the value of laboratory j (Bq $unit^{-1}$);

 u_i is the standard uncertainty for the value of laboratory i (Bq unit⁻¹);

 u_j is the standard uncertainty for the value of laboratory j (Bq $unit^{-1}$); and

unit is the unit of volume or mass, L or kg, as appropriate for the particular sample type.

If two results were received, $\zeta_{1,2}$ was calculated, while for three received results $\zeta_{1,2}$, $\zeta_{1,3}$ and $\zeta_{2,3}$ were calculated.

Following the current ISO standard for statistical methods for use in proficiency testing [5], ζ scores are interpreted as follows:

A ζ score between -2 and 2 indicates that the reported result is accepted at a 95.4% confidence level;

A ζ score between 2 and 3 or between -2 and -3 is considered to give a "warning signal"; and

A ζ score greater than 3 or less than -3 indicates that the reported result is not accepted at a 99.7% confidence level, an "action signal".

A standard approach, according to the same ISO standard, is that an "action signal" can be taken as evidence that an anomaly requiring investigation has occurred.

For the purposes of this ILC, for ζ scores between -3 and 3, the corresponding result was evaluated as agreeing with the reference value at a 99.7% confidence level. (Therefore, results corresponding to "warning signals" were considered to agree with the reference value.) For ζ scores greater than 3 or less than -3, the corresponding result was evaluated as being discrepant at the same confidence level.

If the data set contained four or more results, the statistical evaluation consisted of a method for calculating a comparison reference value as a power-moderated mean of the combined results [6], which is currently being used by the Consultative Committee for Ionizing Radiation, Section II: Measurement of radionuclides, CCRI(II). After calculating a reference value, a relative degree of equivalence (DoE) was calculated for each submitted result and if this relative DoE was significantly different from zero, the corresponding result was evaluated as being discrepant. The relative DoE (%) was calculated according to:

DoE (%) =
$$\frac{x_{\text{lab}} - X_{\text{ref}}}{X_{\text{ref}}}$$
. 100 (2)

where:

 x_{lab} is the individual laboratory result; and

 x_{ref} is the reference value calculated as the power-moderated mean of the combined results.

The standard uncertainty of the relative DoE, u_{DoE} , was calculated according to reference [5]. If the absolute value of the relative DoE exceeded 3 times u_{DoE} , the corresponding result was evaluated as being discrepant (at a 99.7% confidence level), as the relative DoE in this case would be significantly different from zero.

6. RESULTS

6.1. GENERAL

The results are presented in Tables 6 - 15 and Figures 3 - 9.

6.1.1. Uncertainties

Uncertainties quoted in this report are combined standard uncertainties, i.e. with a coverage factor of k = 1. The numerical result of a measurement is stated in the format $xxx \pm yyy$, where the number following the symbol \pm is the numerical value of the combined standard uncertainty and not a confidence interval, unless otherwise indicated (i.e. in Tables 7 and 9).

6.1.2. Reference time

All activity concentrations of radionuclides in seawater, sediment, fish and seaweed samples were reported for a common reference time of 7 November 2022 12:00 UTC.

6.2. SEAWATER

Table 6 contains the results reported by the participating laboratories (IAEA, JCAC, KAKEN, KANSO, KEEA and KINS) for activity concentrations of ³H, ⁶⁰Co, ⁹⁰Sr, ¹⁰⁶Ru, ¹²⁵Sb, ¹²⁹I, ¹³⁴Cs and ¹³⁷Cs in the seawater samples. Figures 3 and 4 present these results visually.

TABLE 6. ACTIVITY CONCENTRATIONS (mBq ${\rm L^{-1}}$) IN SEAWATER SAMPLES

Nuclide	Sample	IAEA	JCAC	KAKEN	KANSO	KEEA	KINS	Reference value
	E-S15	62.3 ±	94 ± 20	_	_	_	115 ± 16	_
		5.7						
	T-0-1A	49 ± 3.1	75 ± 19	162 ± 30	_	_	<83	_
$^{3}\mathrm{H}$	T-3	96 ± 19	64 ± 14	154 ± 29	_	_	<81	_
11	M-E1	81.7 ±	80 ± 19	_	63 ± 8.7	_	<82	_
		9.3						
	M-E3	15.5 ±	57 ± 19	_	_	$44.0 \pm$	99 ± 16	52 ± 18
		4.1				9.0		
⁶⁰ Co		<34	<78	_	_	1	<110	_
⁹⁰ Sr		1.133 ±	$0.66 \pm$	_	_	_	$0.53 \pm$	_
		0.069	0.12				0.13	
¹⁰⁶ Ru		<260	<670	_	_	_	<1200	_
¹²⁵ Sb	E-S15	<100	<180	_	_	_	<310	_
¹²⁹ I	L-313	<180	<180	_	_	_	<88	_
¹³⁴ Cs		0.398 ±	< 0.79	_	_	_	< 0.94	_
		0.038						
¹³⁷ Cs		16.41 ±	17.1 ±	_	_	_	16.6 ±	_
		0.75	1.0				0.81	

Table 7 contains the degrees of relative equivalence for the activity concentrations of ³H, ⁶⁰Co, ⁹⁰Sr, ¹⁰⁶Ru, ¹²⁵Sb, ¹²⁹I, ¹³⁴Cs and ¹³⁷Cs in the seawater samples.

TABLE 7. DEGREES OF EQUIVALENCE (%) IN SEAWATER SAMPLES

Nuclide	Sample	IAEA	JCAC	KAKEN	KANSO	KEEA	KINS
	E-S15	Note 1	Note 1	_	_	_	Note 1
	T-0-1A	Note 2	Note 2	Note 2	_	_	DL
³ H	T-3	Note 3	Note 3	Note 3	_	-	DL
	M-E1	Note 4	Note 4	_	Note 4	-	DL
	M-E3	-70 ± 87	8 ± 111	_	_	-16 ± 92	88 ± 102
⁶⁰ Co		DL	DL	_	_	_	DL
⁹⁰ Sr		Note 5	Note 5	_	_	_	Note 5
¹⁰⁶ Ru		DL	DL	_	_	_	DL
¹²⁵ Sb	E-S15	DL	DL	_	_	-	DL
¹²⁹ I		DL	DL	_	_	_	DL
¹³⁴ Cs		Note 6	DL	_	_	_	DL
¹³⁷ Cs		Note 7	Note 7	_	_	_	Note 7

Notes:

The numerical results in this table are stated in the format $xx \pm yy$, where the number following the symbol \pm is the 99% confidence interval.

Note 1: Values of -1.52, **-3.19** and -0.84 for $\zeta_{1,2}$, $\zeta_{1,6}$ and $\zeta_{2,6}$, respectively.

Note 2: Values of -1.35, **-3.80** and -2.47 for $\zeta_{1,2}$, $\zeta_{1,3}$ and $\zeta_{2,3}$, respectively.

Note 3: Values of 1.38, -1.74 and -2.85 for $\zeta_{1,2}$, $\zeta_{1,3}$ and $\zeta_{2,3}$, respectively.

Note 4: Values of 0.10, 1.47 and 0.79 for $\zeta_{1,2}$, $\zeta_{1,4}$ and $\zeta_{2,4}$, respectively.

Note 5: Values of **3.46**, **4.33** and 0.72 for $\zeta_{1,2}$, $\zeta_{1,6}$ and $\zeta_{2,6}$, respectively.

Note 6: No evaluation was possible as only one value above the detection limit was submitted.

Note 7: Values of -0.56, -0.17 and 0.40 for $\zeta_{1,2}$, $\zeta_{1,6}$ and $\zeta_{2,6}$, respectively.

DL: As a value less than the detection limit was submitted, no evaluation was performed.

 $\zeta_{i,j}$ indexes: number 1 refers to IAEA, number 2 refers to JCAC, number 3 refers to KAKEN, number 4 refers to KANSO, number 5 refers to KEEA and number 6 refers to KINS.

6.3. SEDIMENT

Table 8 contains the results reported by the participating laboratories (IAEA, JCAC, KINS and TPT) for activity concentrations of ¹³⁴Cs and ¹³⁷Cs in the sediment samples. Figures 5 and 6 present these results visually.

TABLE 8. ACTIVITY CONCENTRATIONS (Bq kg⁻¹ d.w.) IN SEDIMENT SAMPLES

Nuclide	Sample	IAEA	JCAC	KINS	TPT	Reference value
	T-1	1.99 ± 0.14	1.47 ± 0.26	2.15 ± 0.31	2.12 ± 0.35	1.92 ± 0.16
¹³⁴ Cs	T-S4	0.238 ±	< 0.70	< 0.58	< 0.86	_
		0.052				
¹³⁷ Cs	T-1	78.5 ± 3.7	85.4 ± 4.3	74.2 ± 2.4	88.8 ± 2.6	81.7 ± 3.5
is/Cs	T-S4	7.56 ± 0.37	8.23 ± 0.47	6.90 ± 0.32	7.47 ± 0.37	7.5 ± 0.27

Table 9 contains the degrees of relative equivalence for the activity concentrations of 134 Cs and 137 Cs in the sediment samples.

TABLE 9. DEGREES OF EQUIVALENCE (%) IN SEDIMENT SAMPLES

Nuclide	Sample	IAEA	JCAC	KINS	TPT
¹³⁴ Cs	T-1	4 ± 23	-23 ± 32	12 ± 39	11 ± 43
CS	T-S4	Note 1	DL	DL	DL
¹³⁷ Cs	T-1	-4 ± 14	5 ± 15	-9 ± 12	9 ± 13
	T-S4	1 ± 13	10 ± 16	-8 ± 12	0 ± 13

Notes:

The numerical results in this table are stated in the format $xx \pm yy$, where the number following the symbol \pm is the 99% confidence interval.

Note 1: No evaluation was possible as only one value above the detection limit was submitted.

DL: As a value less than the detection limit was submitted, no evaluation was performed.

 $\zeta_{i,j}$ indexes: number 1 refers to IAEA, number 2 refers to JCAC, number 3 refers to KINS and number 4 refers to TPT.

6.4. FISH

Table 10 contains the results reported by the participating laboratories (IAEA, KANSO, KEEA, KINS and MERI) for activity concentrations of ³H – organically bound tritium (OBT) and tissue free water tritium (TFWT) – in the fish samples collected from the fish market. No evaluation of these results was possible as, for OBT values less than detection limits were submitted in all cases and, for TFWT, one value above the detection limit was submitted for each sample.

TABLE 10. ACTIVITY CONCENTRATIONS OF ³H (OBT, TFWT) IN FISH SAMPLES

Nuclide	Sample number: Species	IAEA	KANSO	KEEA	KINS	MERI
	T22FA0001: Olive	< 0.0053	< 0.046	< 0.039	< 0.30	_
	flounder					
	T22FA0002:	< 0.0087	< 0.059	< 0.053	< 0.38	_
	Whitespotted conger					
	T22FA0003: Crimson	< 0.0062	< 0.053	< 0.042	< 0.31	_
³ H (OBT, (Bq	sea bream					
kg^{-1} f.w.)	T22FA0004: Redwing	< 0.0065	< 0.051	< 0.04	< 0.29	_
	searobin					
	T22FA0005: Shotted	< 0.0044	< 0.05	< 0.038	< 0.36	_
	halibut					
	T22FA0006: Willowy	< 0.0053	< 0.045	< 0.038	< 0.28	_
	flounder					
	T22FA0001: Olive	0.091 ±	< 0.31	< 0.28	<1.6	< 0.31
	flounder	0.004				
	T22FA0002:	$0.019 \pm$	< 0.31	< 0.28	<1.7	< 0.31
	Whitespotted conger	0.009				
	T22FA0003: Crimson	$0.063 \pm$	< 0.30	< 0.28	<1.7	< 0.34
³ H (TFWT,	sea bream	0.005				
Bq L ⁻¹)	T22FA0004: Redwing	$0.067 \pm$	< 0.30	< 0.28	<1.6	< 0.34
	searobin	0.005				
	T22FA0005: Shotted	$0.069 \pm$	< 0.30	< 0.28	<1.7	< 0.34
	halibut	0.006				
	T22FA0006: Willowy	$0.040 \pm$	< 0.30	< 0.28	<1.7	< 0.34
	flounder	0.007				

Table 11 contains the results reported by the participating laboratories (IAEA, JCAC and KINS) for activity concentrations of ¹⁴C in the fish samples collected from the fish market. Figure 7 presents these results visually.

TABLE 11. ACTIVITY CONCENTRATIONS OF ¹⁴C (Bq g⁻¹ C) IN FISH SAMPLES¹

Nuclide	Sample number: Species	IAEA	JCAC	KINS
	22FA0001: Olive flounder	0.226 ± 0.006	0.224 ± 0.004	0.209 ± 0.011
	22FA0002: Whitespotted	0.233 ± 0.006	0.223 ± 0.003	0.201 ± 0.011
	conger			
	22FA0003: Crimson sea	0.247 ± 0.006	0.231 ± 0.002	0.210 ± 0.011
¹⁴ C	bream			
	22FA0004: Redwing	0.234 ± 0.006	0.233 ± 0.003	0.204 ± 0.011
	searobin			
	22FA0005: Shotted halibut	0.243 ± 0.006	0.219 ± 0.002	0.222 ± 0.011
	22FA0006: Willowy	0.270 ± 0.007	0.222 ± 0.002	0.202 ± 0.011
	flounder			

Note:

Table 12 contains the zeta scores for the activity concentrations of ¹⁴C in the fish samples collected from the fish market.

TABLE 12. ZETA SCORES FOR ¹⁴C IN FISH SAMPLES

Nuclide	Sample number: Species	$\zeta_{IAEA,JCAC}$	$\zeta_{IAEA,KINS}$	$\zeta_{JCAC,KINS}$
	22FA0001: Olive flounder	0.31	1.48	1.41
	22FA0002: Whitespotted	1.61	2.76	2.07
	conger			
	22FA0003: Crimson sea	2.53	3.13	2.03
¹⁴ C	bream			
	22FA0004: Redwing	0.16	2.64	2.79
	searobin			
	22FA0005: Shotted halibut	4.10	1.81	-0.26
	22FA0006: Willowy	7.06	5.34	1.78
	flounder			

 $^{^1}$ These results are expressed using the unit Bq g-1 C (Becquerel per gram carbon) whereas the unit requested and specified in the CRMP is Bq kg-1 f.w. (Bequerel per kilogram fresh weight). To convert from the former to latter, the total carbon content of each fish sample is required. Unfortunately, at the time of publication of this report, this parameter was not available to all participating laboratories. Therefore, on an exceptional basis, it was decided to intercompare the results reported using the unit Bq g-1 C. The results will be re-evaluated at a later date when all laboratories have reported the results using the unit Bq kg-1 f.w..

Table 13 contains the results reported by the participating laboratories (IAEA, JCAC, KAKEN and KINS) for activity concentrations of ${}^{3}\text{H} - \text{OBT}$ and TFWT – and ${}^{14}\text{C}$ in the fish samples in fish samples caught by fixed gill nets. Figures 8 and 9, respectively, present the results for TFWT and ${}^{14}\text{C}$ visually.

TABLE 13. ACTIVITY CONCENTRATIONS IN FISH SAMPLES

Nuclide	Sampling location	IAEA	JCAC	KAKEN	KEEA	KINS
Nuclide	(Species)					
³ Ц (ОРТ (Ра	E-SF3 (Olive flounder)	< 0.0052	< 0.06	_	_	< 0.27
³ H (OBT, (Bq kg ⁻¹ f.w.)	T-S2 (Olive flounder)	< 0.0053	_	< 0.038	_	< 0.26
kg i.w.)	T-S7 (Olive flounder)	< 0.0066	_	_	< 0.043	< 0.32
	E-SF3 (Olive flounder)	0.052 ±	0.093 ±	_	_	<2.1
		0.009	0.023			
³ H (TFWT,	T-S2 (Olive flounder)	$0.060 \pm$	_	0.161 ±	_	<2.0
Bq L ⁻¹)		0.007		0.030		
	T-S7 (Olive flounder)	$0.065 \pm$	_	_	$0.069 \pm$	<2.0
		0.006			0.010	
¹⁴ C (Bq g ⁻¹	E-SF3 (Olive flounder)	$0.197 \pm$	0.240 ±	_	_	0.211 ±
C) ¹		0.005	0.006			0.011

Note:

Table 14 contains the zeta scores for the activity concentrations of ¹⁴C in the fish samples in fish samples caught by fixed gill net.

TABLE 14. ZETA SCORES FOR ³H (TFWT) and ¹⁴C IN FISH SAMPLES

Nuclide	Sampling location (Species)	Zeta scores		
	E-SF3 (Olive flounder)	ζ _{IAEA,} JCAC		
	E-31'3 (Offive flounder)	-1.66		
³ H (TFWT)	T-S2 (Olive flounder)	ζ _{IAEA,KAKEN}		
H(IFWI)	1-32 (Office flourider)	-3.27		
	T-S7 (Olive flounder)	ζ _{IAEA,KEEA}		
	1-37 (Office flourider)	-0.34		
¹⁴ C	E-SF3 (Olive flounder)	ζιαεα,jcac/ζιαεα,kins/ζjcac,kins		
	L-313 (Office flounder)	-5.51 / -1.22/ 2.47		

6.5. SEAWEED

Table 15 contains the results reported by the participating laboratories (IAEA, JCAC, KINS and TPT) for activity concentrations of ¹²⁹I in the seaweed samples.

TABLE 15. ACTIVITY CONCENTRATIONS (Bq kg⁻¹ f.w.) IN SEAWEED SAMPLES

Nuclide	Sample	IAEA	JCAC	KINS	TPT
¹²⁹ I	Tomioka Port (Eisenia bicyclis)	< 0.083	< 0.02	< 0.092	_
	Ukedo Port (Laminariacase)	< 0.068	_	< 0.083	< 0.1
	Tomioka Port (Sargassum Fuvellum)	< 0.11	_	< 0.079	< 0.1

¹ These results are expressed using the unit Bq g−1 C (Becquerel per gram carbon) whereas the unit requested and specified in the CRMP is Bq kg−1 f.w. (Bequerel per kilogram fresh weight). To convert from the former to latter, the total carbon content of each fish sample is required. Unfortunately, at the time of publication of this report, this parameter was not available to all participating laboratories. Therefore, on an exceptional basis, it was decided to intercompare the results reported using the unit Bq g−1 C. The results will be re-evaluated at a later date when all laboratories have reported the results using the unit Bq kg−1 f.w..

7. CONCLUSION

A detailed data analysis was performed on the activity concentrations reported for ³H, ⁶⁰Co, ⁹⁰Sr, ¹⁰⁶Ru, ¹²⁵Sb, ¹²⁹I, ¹³⁴Cs and ¹³⁷Cs in seawater samples, for ¹³⁴Cs and ¹³⁷Cs in sediment samples, for organically bound tritium (OBT), tissue free water tritium (TFWT) and ¹⁴C in fish samples, and ¹²⁹I seaweed samples. All samples were collected offshore TEPCO's Fukushima Daiichi Nuclear Power Station in November 2022. The samples were shared between eight laboratories: six from Japan (JCAC, KAKEN, KANSO, KEEA, MERI and TPT); IAEA; and one laboratory from the Republic of Korea, KINS, a member of the IAEA ALMERA network.

IAEA analyses were undertaken at two participating IAEA Nuclear Sciences and Applications Laboratories:

- IAEA Marine Environment Laboratories, Radiometrics Laboratory (RML), Monaco;
- Isotope Hydrology Laboratory (IHL), Vienna, Austria.

From this analysis it can be concluded that the majority of results are not significantly different from each other. A global analysis of the whole data set demonstrated 10 discrepant⁴ values from the 58 statistical tests applied to the data, i.e. more than 82 % were passed with a high level of confidence (99.7%). The exceptions were 10 zeta scores that were higher than the critical limit of 3:

- $\zeta = -3.19$ for ³H in seawater sample E-S15 between IAEA and KINS.
- $\zeta = -3.80$ for ³H in seawater sample T-0-1A between IAEA and KAKEN.
- $\zeta = 3.46$ and $\zeta = 4.33$ for ⁹⁰Sr in seawater sample E-S15 between IAEA and JCAC and between IAEA and KINS, respectively.
- $\zeta = 3.13$ for ¹⁴C in fish sample T22FA0003 between IAEA and KINS.
- $\zeta = 4.10$ for ¹⁴C in fish sample T22FA0005 between IAEA and JCAC.
- $\zeta = 7.06$ and $\zeta = 5.34$ for ¹⁴C in fish sample T22FA0006 between IAEA and JCAC and between IAEA and KINS, respectively.
- $\zeta = -3.27$ for TFWT in the fish sample from T-S7 between IAEA and KAKEN.
- $\zeta = -5.51$ for ¹⁴C in the fish sample from E-SF3 between IAEA and JCAC.

The percentage of discrepant results in this first ILC to corroborate environmental monitoring related to the ALPS treated water discharges is higher than in the IAEA ILCs organised since 2014 in IAEA's ongoing project "Marine Monitoring: Confidence Building and Data Quality Assurance" [7]. This was to be expected. In this case, the analytical methods are often more complex (e.g. for OBT, TFWT and 14C in fish) and relatively new to some participating laboratories, being implemented specifically to assess the ALPS treated water discharges. Furthermore, higher uncertainties are usually associated with complex analyses of low, and close to detection limit, levels. Notwithstanding these analytical challenges, there were no order of magnitude variations, and no systemic deviations identified between the results reported by Japanese laboratories and those reported by the IAEA and the ALMERA member laboratory. Therefore, despite the above departures, it can be said with confidence that the laboratories are reporting fit-for-purpose reliable and comparable results for the tested radionuclides in seawater, sediment, fish and seaweed samples prepared and analysed according to each laboratory's regularly used methods. The IAEA recommends the organisation of such ILCs annually to improve and maintain the level of confidence in the data reported by the participating laboratories.

⁴ As defined in section 5.

Following the sampling mission, the IAEA can report that Japan's sample collection procedures follow the appropriate methodological standards required to obtain representative samples. The results obtained in ILC 2022 demonstrate a high level of accuracy and competence on the part of the Japanese laboratories involved in the analyses of radionuclides in marine samples for environmental monitoring related to the discharges of ALPS treated water as part of the Government of Japan's Comprehensive Radiation Monitoring Plan.

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⁵ Note: 3rd revision published in October 2023: https://www.kankyo-hoshano.go.jp/wp-content/uploads/2023/10/No09 R510.pdf

APPENDIX: FIGURES

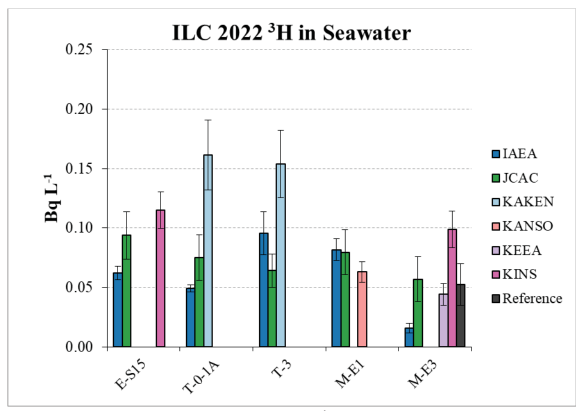


FIG. 3. Activity concentrations of 3H in seawater samples.

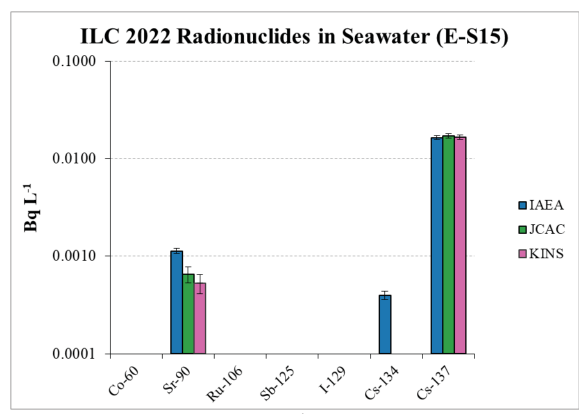


FIG. 4. Activity concentrations of radionuclides⁶ in seawater samples (note logarithmic scale).

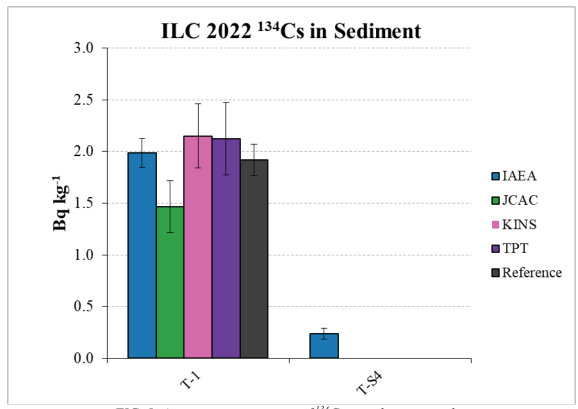


FIG. 5. Activity concentrations of ¹³⁴Cs in sediment samples.

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 $^{^6}$ The "7 major radionuclides" (60 Co, 90 Sr, 125 Sb, 106 Ru, 129 I, 134 Cs and 137 Cs) which, along with 3 H, 14 C and 99 Tc can be routinely detected in samples of ALPS treated water.

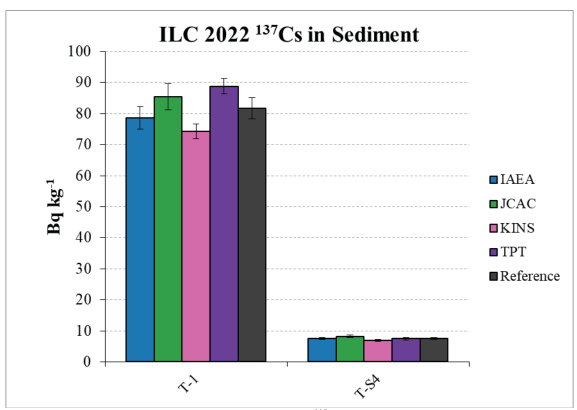


FIG. 6. Activity concentrations of 137 Cs in sediment samples.

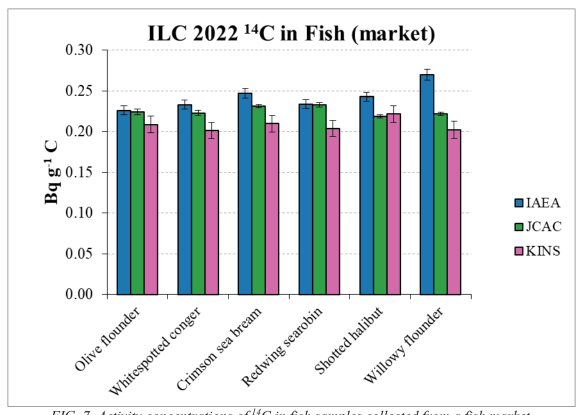


FIG. 7. Activity concentrations of ¹⁴C in fish samples collected from a fish market.

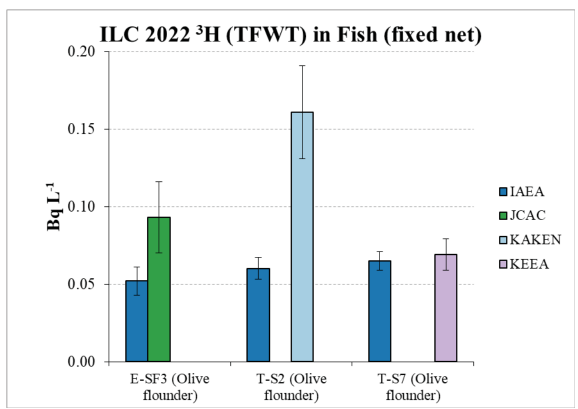


FIG. 8. Activity concentrations of ³H (TFWT) in fish samples caught by fixed gill net.

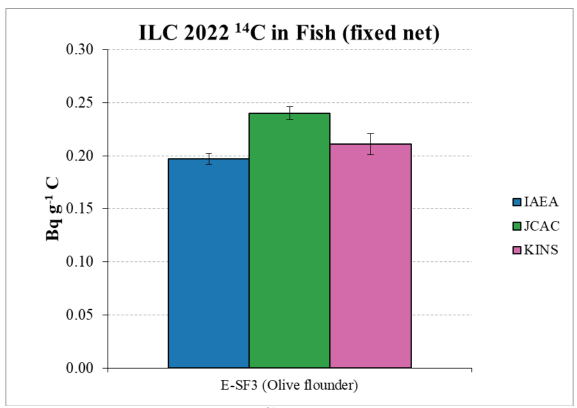


FIG. 9. Activity concentrations of ¹⁴C in fish samples caught by fixed gill net.