October 15, 2024 Okuma Analysis and Research Center Fukushima Research and Engineering Institute Japan Atomic Energy Agency

## Analysis Results of ALPS Treated Water (Sampling at September 4, 2024, 8:29 JST)

Analysis was performed on the ALPS treated water in K4-B tank group in measurement and confirmation facility sampled at September 4, 2024, 8:29 JST.

The analysis results of <sup>3</sup>H and other nuclides are as follows:

The confirmation of nuclides other than <sup>3</sup>H which have been purified to below the regulatory limit by ALPS treatment:

The sum of ratios to the regulatory concentration limits of nuclides other than <sup>3</sup>H (30 nuclides<sup>\*1</sup>) was 0.084(less than 1), confirming that regulatory standard is satisfied.

The nuclides that confirmed not to be significantly present in ALPS treated water (38 nuclides<sup>\*2</sup>): All target radionuclides were confirmed not to be significantly present.

The confirmation of <sup>3</sup>H concentration in ALPS treated water:

<sup>3</sup>H concentration was 3.1E+05 Bq/L<sup>\*3</sup>.

The nuclides which confirmed to be less The nuclides which confirmed not to be than regulatory limit (30 nuclides) 90**Y** <sup>134</sup>Cs 234 J <sup>241</sup>Am  $^{14}C$ <sup>244</sup>Cm <sup>238</sup>U <sup>54</sup>Mn <sup>99</sup>Tc <sup>137</sup>Cs <sup>106</sup>Ru <sup>144</sup>Ce <sup>237</sup>Np <sup>55</sup>Fe 113mCd <sup>238</sup>Pu 60**Co** 147**Pm** <sup>63</sup>Ni <sup>125</sup>Sb <sup>151</sup>Sm <sup>239</sup>Pu <sup>79</sup>Se <sup>125m</sup>Te <sup>154</sup>Eu <sup>240</sup>Pu <sup>90</sup>Sr 129**T** 155**F**U <sup>241</sup>Pu

The analyzed 68 radionuclides other than <sup>3</sup>H are shown in Fig.1

significantly present (38 nuclides) <sup>103</sup>Ru <sup>126</sup>Sn 135Cs <sup>146</sup>Pm <sup>243</sup>Am <sup>59</sup>Fe <sup>58</sup>Co <sup>103m</sup>Rh <sup>124</sup>Sb 136Cs <sup>148</sup>Pm <sup>242</sup>Cm <sup>137m</sup>Ba <sup>243</sup>Cm <sup>106</sup>Rh <sup>123m</sup>Te <sup>148m</sup>Pm <sup>65</sup>Zn <sup>110m</sup>Ag <sup>127</sup>Te <sup>140</sup>Ba <sup>152</sup>Eu <sup>86</sup>Rb <sup>89</sup>Sr 115mCd 127m**Te** <sup>141</sup>Ce <sup>153</sup>Gd <sup>160</sup>Tb 91**Y** <sup>119m</sup>Sn <sup>129</sup>Te <sup>144</sup>Pr <sup>242m</sup>Am 123**Sn** 129m**Te** 144mPr <sup>95</sup>Nb

Fig.1 Classification of radionuclides other than <sup>3</sup>H

- \*1: The nuclides that are selected by TEPCO HD based on the flow defined in the implementation plan and should be confirmed to satisfy the discharge standard (below regulatory standards).
- \*2: From the viewpoint of preventing the adverse impacts on reputation, the nuclides which independently confirmed by TEPCO HD not to significantly present in ALPS treated water.
- \*3: Dilute more than 100 times so that the <sup>3</sup>H concentration after seawater dilution is less than the maximum tritium concentration, 1,500 Bq/L/

1. The confirmation of nuclides other than <sup>3</sup>H which have been purified to below the regulatory limit by ALPS treatment

Analysis results for the sum of ratios to the regulatory concentration limits of nuclides other than  ${}^{3}$ H (30 nuclides) that are confirmed to be less than regulatory limit (sum total of 1) are shown in Table 1. As the result of analysis, the sum of ratios to the regulatory concentration limits of nuclides other than  ${}^{3}$ H was <u>8.4E-02 (less than 1)</u>, confirming that regulatory standard is satisfied.

Table 1 Analysis results of nuclides other than <sup>3</sup>H in ALPS treated water (The nuclides which confirmed to be less than regulatory limit)

Nuclide	Concentrations	Expanded Uncertainty <sup>*1</sup>	Detection Limit	Ratios to Regulatory Concentration Limit	Regulatory Concentratio n Limit <sup>*2</sup>	Measurement/ Evaluation method <sup>**3</sup>
140	[Bq/L]	[Bq/L]	[Dq/L]	[-]	2 000	Maagunamant
54Mn	ND	± 2.5E+00	1 1E 02	1.1E.05	1,000	Measurement
55Ee	ND	-	6.2E.01	3 1E 04	2,000	Measurement
60C		-	0.2E-01	1.2E.02	2,000	Measurement
63 II	2.3E-01	± 3.1E-02	9.4E-03	1.2E-03	200	Measurement
<sup>03</sup> N1	ND	-	1.2E+01	2.0E-03	6,000	Measurement
<sup>79</sup> Se	ND	-	2.0E+00	1.0E-02	200	Measurement
<sup>90</sup> Sr	8.5E-01	± 1.5E-01	6.4E-02	2.8E-02	30	Measurement
<sup>90</sup> Y	8.5E-01	-	6.4E-02	2.8E-03	300	Radiative equilibrium evaluation
<sup>99</sup> Tc	ND	-	1.0E-01	1.0E-04	1,000	Measurement
<sup>106</sup> Ru	ND	-	1.0E-01	1.0E-03	100	Measurement
<sup>113m</sup> Cd	ND	-	1.3E-01	3.3E-03	40	Measurement
<sup>125</sup> Sb	9.9E-02	± 3.2E-02	4.3E-02	1.2E-04	800	Measurement
<sup>125m</sup> Te	2.4E-02	-	1.1E-02	2.7E-05	900	Radiative equilibrium evaluation
<sup>129</sup> I	1.9E-01	± 9.0E-02	1.3E-01	2.1E-02	9	Measurement
<sup>134</sup> Cs	ND	-	4.2E-02	7.0E-04	60	Measurement
<sup>137</sup> Cs	5.5E-02	± 1.1E-02	1.2E-02	6.1E-04	90	Measurement
<sup>144</sup> Ce	ND	-	1.4E-01	7.0E-04	200	Measurement
<sup>147</sup> Pm	ND	-	1.5E-01	5.0E-05	3,000	Relative ratio evaluation

					-	
<sup>151</sup> Sm	ND	-	9 2E-03	1.2E-06	8.000	Relative ratio
5111			9.21 05		0,000	evaluation
<sup>154</sup> Eu	ND	-	4.1E-02	1.0E-04	400	Measurement
<sup>155</sup> Eu	ND	-	7.8E-02	2.6E-05	3,000	Measurement
234 <b>T</b> T					20	Measurement, Total
0					20	α (U, Np group)
238 <b>T</b> I	ND		1 3E 02	3 3E 03 <sup>%4</sup>	20	Measurement, Total
0	ND	-	1.5E-02	5.512-05	20	α (U, Np group)
237 <b>N</b> In					0	Measurement, Total
мр					7	α (U, Np group)
						Measurement,
<sup>238</sup> Pu				1.8E-03 <sup>**4</sup>	4	Total α (Pu, Am,
						Cm group)
	ND	JD -	7.3E-03		4	Measurement,
<sup>239</sup> Pu						Total α (Pu, Am,
						Cm group))
						Measurement,
<sup>240</sup> Pu						Total α (Pu, Am,
						Cm group)
					5	Measurement,
<sup>241</sup> Am						Total α (Pu, Am,
						Cm group)
						Measurement,
<sup>244</sup> Cm					7	Total α (Pu, Am,
						Cm group)
241 <b>P</b> 11	ND		2 2E 01	1 1E 02	200	Relative ratio
ru		-	2.213-01	1.112-05	200	evaluation
Sums o	of the Ratios to Reg	gulatory Concenti	8.4E-02		less than 1	

- $\bigcirc$ . $\bigcirc$ E $\pm$  $\bigcirc$  means  $\bigcirc$ . $\bigcirc$ ×10<sup> $\pm$ </sup> $\bigcirc$
- The results are shown in two significant digits.
- Due to rounding of values, the sums may not exactly match with the actual value.
- ND (Not Detected) in the table indicates that the value is below the detection limit.
- The reference value for the decay correction is the date and time of sampling.
- $\approx$ 1: The uncertainty is the degree of variation in analytical value. Uncertainty is determined by combined all the variations of each step of the analytical procedure from sample collection to measurement. Here, the expanded uncertainty (U = 2 x u) is attached to the analyzed value by doubling the combined standard uncertainty (u).
- \*2: Legally required activity concentration limit established in the Ordinance for Operational Safety and Protection of Specified Nuclear Fuel Materials of the Nuclear Reactors at TEPCO's Fukushima Daiichi (Appendix of first and sixth column: The concentration limit in water outside the peripheral surveillance zone [In this table, Bq/cm<sup>3</sup>

was converted to Bq/L]).

X3: Details of the measurement and evaluation methodology are described below.

Measurement: The radiation of samples is measured then the results are converted into the concentrations of each nuclide.

Total  $\alpha$  (U, Np group): U and Np derived  $\alpha$  rays of the samples were measured and the results are converted to total  $\alpha$  concentration.

Total  $\alpha$  (Pu, Am, Cm group): Pu, Am, Cm derived  $\alpha$  rays of the samples were measured and the results are converted to total  $\alpha$  concentration.

- Radiative equilibrium evaluations: The state in which the ratio of the number of atoms of the parent nuclide to that of its progeny nuclides in a decay series is nearly constant is called radiative equilibrium. The concentration of each nuclide was evaluated based on this radiative equilibrium relationship and the measurement results of the parent (or progeny) nuclide.
- Relative ratio evaluation: The ratio of each nuclide present in the reactor is evaluated by taking into account the generation, decay, impairment loss, etc. of the nuclides. The concentration of each nuclide is calculated by multiplying the measurement results of the reference nuclide by its existence ratio.
- %4: The ratio to the regulatory concentration limits of α nuclides (U-234, U-238, Np-237, Pu-238, Pu-239, Pu-240, Am-241, and Cm-244) was calculated by dividing the total alpha value by the lowest required activity concentration limit (4 Bq/L) of the selected alpha nuclides.

## 2. Analysis results of <sup>3</sup>H in ALPS treated water

Analysis results for the <sup>3</sup>H concentration in ALPS treated water are shown in Table 2. As the result of analysis, <sup>3</sup>H concentration was confirmed to be 3.1E+05 Bq/L.

	Activity	Expanded	Detection	Regulatory	Regulatory	Measurement/
Nuclide	Concentrations	Uncertainty <sup>*1</sup>	Limit	Concentration	Limit <sup>**2</sup>	Evaluation
	[Bq/L]	[Bq/L]	[Bq/L]	Ratio	[Bq/L]	method <sup>**3</sup>
<sup>3</sup> Н	3.1E+05	±2.8E+04	8.6E+01	5.2E+00	60,000	Measurement

Table 2 Analysis results of <sup>3</sup>H in ALPS treated water (sampled at September 4, 2024 8:29 JST)

•  $0.0E \pm 0$  means  $0.0 \times 10^{\pm 0}$ 

• The results are shown in two significant digits.

• The reference value for the decay correction is the date and time of sampling.

- $\approx$  1: The uncertainty is the degree of variation in analytical value. Uncertainty is determined by combined all the variations of each step of the analytical procedure from sample collection to measurement. Here, the expanded uncertainty (U = 2 x u) is attached to the analyzed value by doubling the combined standard uncertainty (u).
- \*2: Legally required activity concentration limit established in the Ordinance for Operational Safety and Protection of Specified Nuclear Fuel Materials of the Nuclear Reactors at TEPCO's Fukushima Daiichi (Appendix of first and sixth column: The concentration limit in water outside the peripheral surveillance zone [In this table, Bq/cm<sup>3</sup> was converted to Bq/L]).
- ※3: Measurement: The radiation of samples is measured then the results are converted into the concentrations of each nuclide.

3. Nuclides other than <sup>3</sup>H to be confirmed as not significantly present in ALPS treated water Analysis results for the target nuclides (38 nuclides) other than <sup>3</sup>H that were confirmed to be as not significantly present in ALPS treated water are shown in Table 3. As the result of analysis, <u>all target</u> <u>radionuclides were confirmed not to be significantly present</u>. Furthermore, the sum of the ratios to the regulatory concentration limits for the 68 nuclides in Tables 1 and 3 is also less than 1.

		<u> </u>			
Nuclide	Activity Concentrations	Regulatory Concentration Ratio	Regulatory Limit <sup>**1</sup>	Evaluation <sup>**2</sup>	Measurement/ Evaluation method <sup>*3</sup>
	[Bq/L]	[-]	[Bq/L]		
<sup>59</sup> Fe	2.0E-02	5.0E-05	400	0	Measurement
<sup>58</sup> Co	1.1E-02	1.1E-05	1,000	0	Measurement
<sup>65</sup> Zn	1.8E-02	9.0E-05	200	0	Measurement
<sup>86</sup> Rb	2.1E-01	7.0E-04	300	0	Measurement
<sup>89</sup> Sr	7.9E-02	2.6E-04	300	0	Measurement
<sup>91</sup> Y	4.0E+00	1.3E-02	300	0	Measurement
<sup>95</sup> Nb	1.4E-02	1.4E-05	1,000	0	Measurement
<sup>103</sup> Ru	1.5E-02	1.5E-05	1,000	0	Measurement
103m <b>D</b> h	1.5E.02	7.5E 09	200.000	0	Radiative equilibrium
KII	1.5E-02	7.52-08	200,000	0	evaluation
106 <b>D</b> h	1.0F.01	3 3E 07	300.000	0	Radiative equilibrium
	1.02-01	5.52-07	500,000	0	evaluation
<sup>110m</sup> Ag	1.1E-02	3.7E-05	300	0	Measurement
<sup>115m</sup> Cd	5.7E-01	1.9E-03	300	0	Measurement
<sup>119m</sup> Sn	1.2E-02	6.0E-06	2,000	0	Relative ratio evaluation
<sup>123</sup> Sn	1.6E+00	4.0E-03	400	0	Measurement
$^{126}$ Sn	3.7E-01	1.9E-03	200	0	Measurement
$^{124}$ Sb	2.0E-02	6.7E-05	300	0	Measurement
<sup>123m</sup> Te	2.1E-02	3.5E-05	600	0	Measurement
<sup>127</sup> Te	1.3E+00	2.6E-04	5,000	0	Measurement
<sup>127m</sup> Te	1.4E+00	4.7E-03	300	0	Relative ratio evaluation
<sup>129</sup> Te	1.8E-01	1.8E-05	10,000	0	Measurement
<sup>129m</sup> Te	3.6E-01	1.2E-03	300	0	Measurement
<sup>135</sup> Cs	8.3E-08	6.2E-10	600	0	Relative ratio evaluation
<sup>136</sup> Cs	1.3E-02	4.3E-05	300	0	Measurement
<sup>137m</sup> Ba	1.2E-02	6.5E-08	800,000	0	Radiative equilibrium evaluation

Table 3 Analysis results of nuclides other than <sup>3</sup>H in ALPS treated water (Nuclides which confirmed not to be significantly present) (sampling time: September 4, 2024 8:29 JST)

<sup>140</sup> Ba	6.2E-02	2.1E-04	300	0	Measurement
<sup>141</sup> Ce	5.8E-02	5.8E-05	1,000	0	Measurement
<sup>144</sup> Pr	1.4E-01	7.0E-06	20,000	0	Radiative equilibrium evaluation
<sup>144m</sup> Pr	1.3E-03	3.3E-08	40,000	0	Radiative equilibrium evaluations
<sup>146</sup> Pm	1.9E-02	2.1E-05	900	0	Measurement
<sup>148</sup> Pm	7.9E-02	2.6E-04	300	0	Measurement
<sup>148m</sup> Pm	1.2E-02	2.4E-05	500	0	Measurement
<sup>152</sup> Eu	5.7E-02	9.5E-05	600	0	Measurement
<sup>153</sup> Gd	4.8E-02	1.6E-05	3,000	0	Measurement
<sup>160</sup> Tb	3.5E-02	7.0E-05	500	0	Measurement
<sup>242m</sup> Am	3.8E-05	7.6E-06	5	0	Relative ratio evaluation
243 <b>A</b> m			E	0	Measurement, Total $\alpha$
Alli			5	0	(Pu, Am, Cm group)
<sup>242</sup> Cm	7 3E 03	1 8E 03 <sup>%4</sup>	60	0	Measurement, Total $\alpha$
	7.5E-05	1.8E-03 <sup>***</sup>	00	0	(Pu, Am, Cm group)
<sup>243</sup> Cm			6		Measurement, Total α
Cili			0	0	(Pu, Am, Cm group)

•  $\bigcirc.\bigcirc E \pm \bigcirc$  means  $\bigcirc.\bigcirc \times 10^{\pm \bigcirc}$ 

• The results are shown in two significant digits.

• Due to rounding of values, the sums of shown results in the table may not match.

• The reference value for the decay correction is the date and time of sampling.

- \* 1: Legally required activity concentration limit established in the Ordinance for Operational Safety and Protection of Specified Nuclear Fuel Materials of the Nuclear Reactors at TEPCO's Fukushima Daiichi (Appendix of first and sixth column: The concentration limit in water outside the peripheral surveillance zone [In this table, Bq/cm<sup>3</sup> was converted to Bq/L]).
- 2: If it is not significantly present, it is marked with a " $\bigcirc$ "; if it is significantly present, it is marked with an " $\times$ ".

The evaluation is made as not significantly present (" $\bigcirc$ ") if any of the following are satisfied:

- The measured nuclide concentration is below the detection limit.
- Nuclides evaluated by radiative equilibrium, etc.: When the evaluated nuclide is detected, the concentration
  is extremely low compared to the regulatory concentration limit, i.e., the evaluated value is less than the set
  detection limit (less than 1/100 of the regulatory concentration limit), and can be judged to be equivalent to
  less than the detection limit.

Evaluated Nuclide	Evaluated Value [Bq/L]	Regulatory Concentration Ratio	Regulatory Limit <sup>**1</sup> [Bq/L]
$^{103\mathrm{m}}\mathrm{Rh}$	ND	-	200,000
$^{106}$ Rh	ND	-	300,000

<sup>119m</sup> Sn	ND	-	2,000
<sup>127m</sup> Te	ND	-	300
<sup>135</sup> Cs	3.7E-07	6.2E-10	600
<sup>137m</sup> Ba	5.2E-02	6.5E-08	800,000
<sup>144</sup> Pr	ND	-	20,000
<sup>144m</sup> Pr	ND	-	40,000
<sup>242m</sup> Am	ND	-	5

• ND (Not Detected) in the table indicates that the value is below the detection limit.

X3: Details of the measurement and evaluation methodology are described below.

Measurement: The radiation of samples is measured then the results are converted into the concentrations of each nuclide.

Total  $\alpha$  (U, Np group): U and Np derived  $\alpha$  rays of the samples were measured and the results are converted to total  $\alpha$  concentration.

Total  $\alpha$  (Pu, Am, Cm group): Pu, Am, Cm derived  $\alpha$  rays of the samples were measured and the results are converted to total  $\alpha$  concentration.

Radiative equilibrium evaluations: The state in which the ratio of the number of atoms of the parent nuclide to that of its progeny nuclides in a decay series is nearly constant is called radiative equilibrium. The concentration of each nuclide was evaluated based on this radiative equilibrium relationship and the measurement results of the parent (or progeny) nuclide.

Relative ratio evaluation: The ratio of each nuclide present in the reactor is evaluated by taking into account the generation, decay, impairment loss, etc. of the nuclides. The concentration of each nuclide is calculated by multiplying the measurement results of the reference nuclide by its existence ratio.

%4: The ratio to the regulatory concentration limits of α nuclides (Am-243, Cm-242, and Cm-243) was calculated by dividing the total alpha value by the lowest required activity concentration limit (4 Bq/L) of the selected alpha nuclides.

Reference:	Measurement	and evalu	ation me	thods for	each n	uclide

		in t	the third-party analysis of ALPS treated water
N.	Nice all de		Measurement and evaluation methods for each nuclide
INO.	Nuclide		in the third-party analysis of ALPS treated water
1	311	0	Purify tritiated water by distillation and mixing sample and scintillator
1	<sup>2</sup> H	p-ray	Liquid scintillation counter
0	<sup>14</sup> C	0	Isolation by collection on adsorbent, mixing sample and scintillator
2		p-ray	Liquid scintillation counter
2	541 4		A single sample was prepared in a Marinelli beaker.
3	- Min	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
4	5512-	V	Isolation by resin and sedimentation
4	<sup>55</sup> Fe	A-ray	Low-energy photon detector (Ge-LEPS)
5	5917		A single sample was prepared in a Marinelli beaker.
2	JFe	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
(	5 <sup>58</sup> Co		A single sample was prepared in a Marinelli beaker.
6		o γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
7	60.0	γ-ray	A single sample was prepared in a Marinelli beaker.
/	7 °°Co		Gamma-ray spectrometry using high purity germanium (HPGe) detector
0	63 1	β-ray	Isolation by resin, mixing sample and scintillator
8	8 <sup>03</sup> N1		Liquid scintillation counter
0	657	γ-ray	A single sample was prepared in a Marinelli beaker.
9	<sup></sup> Zn		Gamma-ray spectrometry using high purity germanium (HPGe) detector
10	798 -	ICP-MS	Isolation by resin
10	~Se		Inductively coupled plasma mass spectrometer
11	86101		A single sample was prepared in a Marinelli beaker.
11	<sup>oo</sup> Kb	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
10	890	0	Isolation by resin and sedimentation
12	° Sr	p-ray	Beta-ray spectrometer (plastic scintillator)
12	900	0	Isolation by resin and sedimentation
13	Sr	p-ray	Beta-ray spectrometer (plastic scintillator)
14	<sup>90</sup> Y	Evaluation	Radiative equilibrium evaluation
15	9157		A single sample was prepared in a Marinelli beaker.
15	ŶŶŶ	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
16	958.11		A single sample was prepared in a Marinelli beaker.
16	<sup>33</sup> Nb	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
17	99 <b>-</b>		Isolation by resin
17	<sup>37</sup> Ic	ICP-MS	Inductively coupled plasma mass spectrometer
10	1025		A single sample was prepared in a Marinelli beaker.
18	<sup>105</sup> Ru	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector

10	19 <sup>106</sup> Ru	γ-ray	A single sample was prepared in a Marinelli beaker.
19	Ku		Gamma-ray spectrometry using high purity germanium (HPGe) detector
20	<sup>103m</sup> Rh	Evaluation	Radiative equilibrium evaluation
21	<sup>106</sup> Rh	Evaluation	Radiative equilibrium evaluation
22	110m <b>A</b>		A single sample was prepared in a Marinelli beaker.
22	22 <sup>riom</sup> Ag	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
22	113mc 1	0	Isolation by resin, mixing sample and scintillator
23	"IIImCd	p-ray	Liquid scintillation counter
24	115mC 4		A single sample was prepared in a Marinelli beaker.
24	Ca	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
25	<sup>119m</sup> Sn	Evaluation	Relative ratio evaluation:
26	1230		A single sample was prepared in a Marinelli beaker.
26	<sup>125</sup> Sn	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
27	1260		A single sample was prepared in a Marinelli beaker.
27	<sup>120</sup> Sn	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
29	124 01		A single sample was prepared in a Marinelli beaker.
28	12150	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
20	12501	γ-ray	A single sample was prepared in a Marinelli beaker.
29	29 <sup>125</sup> Sb		Gamma-ray spectrometry using high purity germanium (HPGe) detector
20	123mTo	γ-ray	A single sample was prepared in a Marinelli beaker.
30	Ie		Gamma-ray spectrometry using high purity germanium (HPGe) detector
31	<sup>125m</sup> Te	Evaluation	Radiative equilibrium evaluation
22	127 <b>T</b> 2	AL #011	A single sample was prepared in a Marinelli beaker.
32	Te	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
33	<sup>127m</sup> Te	Evaluation	Radiative equilibrium evaluation
24	129Ta	AL #011	A single sample was prepared in a Marinelli beaker.
54	Ie	γ-lay	Gamma-ray spectrometry using high purity germanium (HPGe) detector
35	129m <b>T</b> e	N 7037	A single sample was prepared in a Marinelli beaker.
35	IC	y-lay	Gamma-ray spectrometry using high purity germanium (HPGe) detector
26	129 <b>т</b>	V rou	A single sample was prepared in a measurement cell.
50	1	A-lay	Low-energy photon detector (Ge-LEPS)
27	134		A single sample was prepared in a Marinelli beaker.
57	Cs	γ-lay	Gamma-ray spectrometry using high purity germanium (HPGe) detector
38	<sup>135</sup> Cs	Evaluation	Relative ratio evaluation:
20	136	0/ 2037	A single sample was prepared in a Marinelli beaker.
37		γ-iay	Gamma-ray spectrometry using high purity germanium (HPGe) detector
40	137 <b>C</b> a	0/ 2037	A single sample was prepared in a Marinelli beaker.
40 <sup>15</sup> Cs	Cs	Cs γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector

41	<sup>137m</sup> Ba	Evaluation	Radiative equilibrium evaluation
42	140 <b>D</b> -	γ-ray	A single sample was prepared in a Marinelli beaker.
42	Ба		Gamma-ray spectrometry using high purity germanium (HPGe) detector
42	1410		A single sample was prepared in a Marinelli beaker.
43	Ce	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
4.4	144 <b>C</b> a		A single sample was prepared in a Marinelli beaker.
44	Ce	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
45	<sup>144</sup> Pr	Evaluation	Radiative equilibrium evaluation
46	<sup>144m</sup> Pr	Evaluation	Radiative equilibrium evaluation
47	146 <b>D</b>		A single sample was prepared in a Marinelli beaker.
4/	Pm	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
48	<sup>147</sup> Pm	Evaluation	Relative ratio evaluation:
40	148 <b>D</b>		A single sample was prepared in a Marinelli beaker.
49	<sup>r</sup> <sup>o</sup> Pm	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
50	148m <b>D</b>		A single sample was prepared in a Marinelli beaker.
50	Pin	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
51	<sup>151</sup> Sm	Evaluation	Relative ratio evaluation:
50	152	γ-ray	A single sample was prepared in a Marinelli beaker.
32	Eu		Gamma-ray spectrometry using high purity germanium (HPGe) detector
52	154		A single sample was prepared in a Marinelli beaker.
55	Eu	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
51	155	γ-ray	A single sample was prepared in a Marinelli beaker.
54	Eu		Gamma-ray spectrometry using high purity germanium (HPGe) detector
55	153 C 4	δd γ-ray	A single sample was prepared in a Marinelli beaker.
33	Ga		Gamma-ray spectrometry using high purity germanium (HPGe) detector
56	160-71-		A single sample was prepared in a Marinelli beaker.
50	10	γ-ray	Gamma-ray spectrometry using high purity germanium (HPGe) detector
57	234 <b>1</b> T	0 101	Separated by resin and evaporated and solidified on a stainless plate
57		u-lay	α-ray scintillation counter (ZnS scintillator)
50	238 <b>t</b> t	0 101	Separated by resin and evaporated and solidified on a stainless plate
58	0	u-lay	α-ray scintillation counter (ZnS scintillator)
50	237	of 40011	Separated by resin and evaporated and solidified on a stainless plate
39	мр	u-lay	α-ray scintillation counter (ZnS scintillator)
60	238 <b>D</b> 11	0 7037	Separated by resin, collected as sedimentation, and evaporated to dryness
00	ru	u-ray	α-ray scintillation counter (ZnS scintillator)
61	2390	0 7017	Separated by resin, collected as sedimentation, and evaporated to dryness
	61 <sup>239</sup> Pu	ru α-ray	α-ray scintillation counter (ZnS scintillator)

62	<sup>240</sup> Pu	α-ray	Separated by resin, collected as sedimentation, and evaporated to dryness
			α-ray scintillation counter (ZnS scintillator)
63	<sup>241</sup> Pu	Evaluation	Relative ratio evaluation:
64	<sup>241</sup> Am	α-ray	Separated by resin, collected as sedimentation, and evaporated to dryness
			α-ray scintillation counter (ZnS scintillator)
65	<sup>242m</sup> Am	Evaluation	Relative ratio evaluation:
66	<sup>243</sup> Am	α-ray	Separated by resin, collected as sedimentation, and evaporated to dryness
			α-ray scintillation counter (ZnS scintillator)
67	<sup>242</sup> Cm	α-ray	Separated by resin, collected as sedimentation, and evaporated to dryness
			α-ray scintillation counter (ZnS scintillator)
68	<sup>243</sup> Cm	α-ray	Separated by resin, collected as sedimentation, and evaporated to dryness
			α-ray scintillation counter (ZnS scintillator)
69	<sup>244</sup> Cm	α-ray	Separated by resin, collected as sedimentation, and evaporated to dryness
			α-ray scintillation counter (ZnS scintillator)