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1	Cherenkov counting of ⁹	⁹⁰ Sr and ⁹⁰ Y	in bark and	leaf samples	collected	around	Fukushima	Daiichi

- 2 Nuclear Power Plant
- 3

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- 12

13 Abstract

The radioactivity of ⁹⁰Sr and ¹³⁷Cs in environmental samples, bark and leaf, collected around the 14 Fukushima Daiichi Nuclear Power Plant in May 2013 was determined with the aim of investigating the 15 migration of both nuclides using their radioactivity ratio. The radioactivity of ⁹⁰Sr was determined by 16 using Cherenkov counting of ⁹⁰Y after purification using Sr resin and that of ¹³⁷Cs was determined by 17 γ -spectrometry. Quench correction in Cherenkov counting was investigated by measurements of samples 18 spiked with purified ⁹⁰Y revealed that the radioactivity could be evaluated without quench correction. The 19 radioactivity ratio of 90 Sr to 137 Cs in bark samples of 4.2×10^{-3} and 1.2×10^{-2} was compared with the results 20 from soil samples collected in July 2011 to show that the migration of ⁹⁰Sr was slower than ¹³⁷Cs in bark 21 22 and tree.

23

Keywords: strontium-90, cesium-137, Cherenkov counting, Fukushima nuclear accident, vegetation
 samples

26

27 Introduction

The accident at the Fukushima Daiichi Nuclear Power Plant led to the release of a large amount of 28 29 various radioactive materials, which migrated through the atmosphere, hydrosphere, and biosphere and a 30 part of which were chemically and physically trapped in the environment. Among the radioactive nuclides 31 released as a result of the Fukushima accident, the environmental contamination from radioiodine and 32 radiocesium has been well-studied previously [1-6]; these studies have reported that radioiodine spread 33 globally while radiocesium was mainly deposited in East Japan and released into the Pacific Ocean. 34 Radioactive strontium, not well-studied in the wake of the Fukushima accident, is still important to investigate for its migration into vegetation and accumulation in foodstuff in terms of internal exposure. 35

⁸⁹Sr and ⁹⁰Sr, unlike ¹³⁷Cs and ¹³¹I, are pure beta emitter nuclides, and their radioactive measurements should be conducted after the purification of strontium from the sample matrix. Methods for purification include the use of a specific resin [7-11], solvent extraction [12], and anion exchange resin [13,14]. Radioactivity levels are then often determined using a gas flow proportional counter [7,8,11,15,16] and a liquid scintillation counter that employs scintillation fluid [10,17-21] or Cherenkov 41 radiation [13,14,22]. Methods that utilize Cherenkov radiation are generally more practical because the 42 sample solution can be measured without mixing extra fluids, which prevents the recovery of the 43 strontium fraction.

We determined the radioactivity of ⁹⁰Sr and ¹³⁷Cs in vegetation samples collected in the vicinity of the Fukushima Daiichi Nuclear Power Plant, where agricultural activities are prohibited due to high-level radioactive contamination. Thus, we collected bark and leaf samples instead of agricultural samples and investigated the migration of both nuclides in bark and tree in comparison to the radioactivity of soil samples [23]. In order to validate the Cherenkov counting for the determination of ⁹⁰Sr, we evaluated the extent of the quench effect by measuring and comparing the samples with and without the spike of ⁹⁰Y solution.

51

52 Materials and Methods

We collected vegetation samples around the Fukushima Daiichi Nuclear Power Plant from 25–28 May 2013. Bark and fresh leaves from *Cryptomeria japonica* and fresh leaves from *Artemisia indica var. maximowiczii* were collected from plants 3 km north of the power plant's Unit No. 1, and bark from *Metasequoia glyptostroboides* was collected from plants 2 km south of the unit. Air dose rates at 1 m height were measured at 1 µSv/h and 30 µSv/h from the north and south locations, respectively.

58 The vegetation samples were washed with distilled water under an ultrasonic wave to remove soil 59 particles from the surface of the vegetation samples. The wet samples and the resulting suspended 60 solution were dried at approximately 100 °C, packed in plastic bags and bottles, and stored in a desiccator before further treatment. An aliquot of each vegetation sample (2.5 g dry weight) was placed in a quartz 61 test tube and incinerated at 600 °C in a tubular furnace. The ashed samples were then dissolved in hot 62 concentrated nitric acid, fumed to dryness, dissolved in 0.1 M HNO₃, and centrifuged into separate 63 64 supernatants. The resulting solution for each sample was treated with cation exchange resin Dowex 50WX8 (100-200 mesh), Eichrom UTEVA-resin (100-150 μm), and Eichrom Sr resin (100-150 μm) to 65 purify the strontium fraction [24]. The strontium fraction, contained in the supernatant solution in 50 mL 66 of 0.1 M HNO₃, was loaded into 4 mL of cation exchange resin and recovered with 20 mL of 8 M HNO₃ 67 following a resin wash with 0.1 M HNO₃. The recovered strontium was then injected into a column filled 68 69 with 1 mL of UTEVA resin to remove uranium and plutonium. The strontium fraction eluted into the first 70 effluent was fumed to dryness and prepared to 10 mL of 3 M HNO₃ solution. This solution was injected 71 into a column filled with 1 mL of Sr resin, resin washed with 5 mL of 3 M HNO₃, and treated with 20 mL 72 of 0.05 M HNO₃ to recover the strontium fraction. Finally, for each sample, the radioactivity levels of 90 Sr 73 and ⁹⁰Y in the purified strontium solution were measured by Cherenkov counting and the recovery of 74 strontium throughout the purification was evaluated using an inductively coupled plasma quadrupole 75 mass spectrometry (ICP-QMS, HP-4500, Yokogawa).

The Cherenkov radiation from an aliquot of the purified strontium solution (10 mL) was measured with a liquid scintillation analyzer (Tri-Carb 2700 TR, Packard), with Milli-Q water used to obtain the background value. The radioactivity of ¹³⁷Cs, which accounted for dominant levels of radioactivity in the samples collected around the Fukushima Daiichi Nuclear Power Plant, was measured by γ -spectrometry (a coaxial-type Ge detector, Canberra). For this measurement, energy calibration was conducted with a certified calibration standard source (¹⁰⁹Cd, ⁵⁷Co, ¹³⁹Ce, ²⁰³Hg, ¹¹³Sn, ⁸⁵Sr, ¹³⁷Cs, ⁸⁸Y, and ⁶⁰Co; GE
Healthcare), and solid angle correction factors were determined with potassium chloride powder and
solution [25,26].

84 The calibration of the Cherenkov counting, the evaluation of its coefficient, and the correction of quenching in the counting were conducted using ¹³⁴Cs generated at the Kyoto University Research 85 Reactor (KUR) [27] and the standard solution of ⁹⁰Sr was purchased from Eckert and Ziegler Isotope 86 Products. The counting coefficients of ⁹⁰Sr and ⁹⁰Y were obtained from the increase in the radioactivity of 87 ⁹⁰Y in the purified ⁹⁰Sr solution. The calculation is described below. The correction of quenching was 88 conducted by the spiking of purified ⁹⁰Y solution to the sample strontium solution and nitric acid, as 89 90 control for quenching, for comparison after quantitative analysis. The difference of the resulting increase in counting rate between both solutions showed a need to correct for quenching. Purified ⁹⁰Y, absent of 91 ⁹⁰Sr, was spiked to avoid the spoil of sample strontium solution, whose radioactivity was finally regulated 92 by the original ⁹⁰Sr after the decay of the extra radioactivity of ⁹⁰Y. The separation of ⁹⁰Sr and ⁹⁰Y from 93 94 the standard solution was conducted by using the Sr resin method, as described above, and the radioactive 95 purity of both fractions were 99.9%.

96

97 Calculations

We quantified levels of 90 Sr using the radioactivity of its daughter nuclide, 90 Y, which required long periods of time for measuring low-level samples. The change in 90 Y radioactivity during measurements must be accounted for in this process because of its relatively short half-life, and corrections for decay and growth are described in detail below. The radioactivity levels of environmental samples due to 90 Sr were calculated using regression analysis, with the 90 Y and 90 Sr measurement coefficient.

103 When radioactivity changes during measurements, the counting rate C (e.g., count per minute 104 (CPM)) can be expressed according to the following equation:

105
$$C = \frac{1}{t_d} \int_{t_1}^{t_1+t_d} \sum \varepsilon_i A_i(t) dt + BG$$
(1)

where t_d is counting time, t_1 is the start time of counting, ε_i and A_i are the measurement coefficient and the radioactivity, respectively, of nuclide *i* (i.e., $i = {}^{90}$ Y, 90 Sr, 134 Cs, 137 Cs, 40 K, etc.) with the radioactivity of 89 Sr ignored because of its short half life causes it to decay away before the sample collection, and *BG* is the background value derived from the blank measurement. We used this equation for the correction of growth for 90 Y. The net counting rate (net CPM), *Y*, was then expressed according to the following equation:

$$112 \qquad Y = C - BG \tag{2}$$

Both measurement coefficients for ⁹⁰Y and ⁹⁰Sr were evaluated from the measurement of a standard ⁹⁰Sr solution in equilibrium, completely eliminated of ⁹⁰Y. For the measurement of the ⁹⁰Sr solution with the growth of ⁹⁰Y, Eq. (1) can be modified to the following equation:

116
$$Y = \frac{1}{t_d} \int_{t_1}^{t_1 + t_d} \varepsilon_{\text{Y-90}} A_{\text{Y-90}}(t) dt + \varepsilon_{\text{Sr-90}} A_{\text{Sr-90}}^0$$
(3)

117 where A_{Sr-90}^{0} is the initial radioactivity of ⁹⁰Sr and its decay is ignored for the short period. The

118 radioactivity of 90 Y, $A_{Y-90}(t)$, produced through the decay of 90 Sr, can be expressed according to the 119 following equation:

120
$$A_{Y-90}(t) = \frac{\lambda_{Y-90}}{\lambda_{Y-90} - \lambda_{Sr-90}} A_{Sr-90}^{0} \left(e^{-\lambda_{Sr-90}t} - e^{-\lambda_{Y-90}t} \right)$$
(4)

121 where λ_{Y-90} and λ_{Sr-90} are the decay constants of ${}^{90}Y$ and ${}^{90}Sr$, respectively, and the initial 122 radioactivity of ${}^{90}Y$ is zero as ${}^{90}Y$ is eliminated by the separation of ${}^{90}Sr$. In the short period where the 123 decay of ${}^{90}Sr$ can be ignored, the relationship between both decay constants, $\lambda_{Y-90} \gg \lambda_{Sr-90}$, 124 simplifies the equation to the following:

125
$$A_{\rm Y-90}(t) = A_{\rm Sr-90}^0 \left(1 - e^{-\lambda_{\rm Y-90}t} \right)$$
(5)

126 This equation combined with Eq. (3), yields the following:

127
$$Y = \varepsilon_{\text{Sr-90}} A^0_{\text{Sr-90}} \left(1 + \alpha X\right)$$
(6)

128 where
$$\alpha = \frac{\mathcal{E}_{Y-90}}{\mathcal{E}_{Sr-90}}$$
 and $X = 1 - e^{-\lambda_{Y-90}t_1} \left(\frac{1 - e^{-\lambda_{Y-90}t_d}}{\lambda_{Y-90}t_d}\right)$, and X is the ⁹⁰Y saturation ratio corrected

for growth during the measurement time (corrected equilibrium fraction). We conducted a regression analysis to determine the initial radioactivity of 90 Sr. Eq. (6) is transformed by taking the log of both sides to yield the following equation:

132
$$\log Y = \log\left(\varepsilon_{\mathrm{Sr-90}}A_{\mathrm{Sr-90}}^{0}\right) + \log\left(1 + \alpha X\right)$$
(7)

133 This equation is adapted to a non-linear least squares (LSQ) equation to calculate the value of α 134 and $\log(\varepsilon_{Sr-90}A_{Sr-90}^0)$. We use only the value of α to evaluate the absolute values of both

measurement coefficients in the next step, because the value of A_{Sr-90}^{0} is not equal to the initial solution and becomes unknown through the yttrium elimination process where some of ⁹⁰Sr is discarded with ⁹⁰Y. The net CPM of the initial ⁹⁰Sr solution in equilibrium shows that Eq. (3) can be converted to the following:

139
$$Y = \varepsilon_{Y-90} A_{Sr-90}^{0} + \varepsilon_{Sr-90} A_{Sr-90}^{0} = \varepsilon_{Sr-90} (1+\alpha) A_{Sr-90}^{0}$$
(8)

140 and the resulting equation provides the measurement coefficients for 90 Y and 90 Sr.

The evaluation of ⁹⁰Sr concentration in the environmental samples would require the consideration
of coexistence of other radionuclides, even though a careful purification of strontium is conducted. In this
case Eq. (1) is expressed as:

144
$$Y = \left(\varepsilon_{Y-90} A_{Sr-90}^0\right) X + B \tag{9}$$

145 where $B = \sum \varepsilon_i A_i$ with $i = {}^{90}$ Sr, 134 Cs, 137 Cs, 40 K, etc., and *B* is regarded as a constant value. After 146 the purification of strontium, the increase in net CPM with the corrected equilibrium fraction is applied to 147 regression analysis to yield the value of $\mathcal{E}_{Y-90}A^0_{Sr-90}$ as the slope. Then, the value of A^0_{Sr-90} , the 148 concentration of ⁹⁰Sr in the purified solution, can be determined.

149

150 Results and Discussion

In this study, we used Cherenkov radiation generated by beta particles to detect radionuclides in 151 vegetation collected around the Fukushima Daiichi Nuclear Power Plant. Table 1 lists the characteristics 152 of major beta-emitting nuclides released from the power plant as well as those of ⁴⁰K, a naturally 153 154 occurring radionuclide with a long half-life [28]. The Cherenkov spectra of selected representative radionuclides are shown in Figure 1. The width of each spectrum peak increases with increased beta 155 energy, and the spectra obtained for ⁹⁰Sr and ¹³⁴Cs were derived from similar beta energy profiles. Thus, it 156 is reasonable to suggest that the spectra of ¹³⁷Cs and ⁸⁹Sr resemble those of ⁹⁰Sr and ⁴⁰K, respectively. In 157 158 addition, the Cherenkov spectra obtained in this study support the purity of focal radionuclides as well as 159 the need for further purification of samples.

160 When the same amount of radioactivity is spiked to two measurement samples, the increase in counting rate is expected to be the same for each unless the effect of quenching in measurement occurs. 161 We have investigated the contribution of quenching by the addition of ⁹⁰Y solution of 0.2 mL in 0.1 M 162 HNO₃ to the each original measurement sample where ⁹⁰Y was in radioactive equilibrium and in 10 mL of 163 164 approximately 2 M HNO₃. While this treatment changed the total volume and the resulting acid 165 concentration of the measurement samples, these changes produced no effect to the net CPM, as shown in 166 the following results. Figure 2 shows the effect of sample solution volume on net CPM where the value of 167 the net CPM is normalized to the result for the volume of 10 mL, and exhibits good agreement between 168 sample solutions of volume 5–18 mL. Figure 3 shows the effect of sample acid concentration on net CPM, 169 where the value of the net CPM is normalized to the result for the concentration of 2 M HNO₃, and 170 exhibits good agreement between HNO₃ concentrations ranging from 0.3 M to 5.4 M. These results suggest that the net CPM of ⁹⁰Sr of 10 mL in 2 M HNO₃ will not change by addition of water or nitric 171 acid at most a few mL, which validates the spike method used to correct for quenching. Figure 4 shows 172 173 that the increase in the net CPM of four environmental samples spiked with ⁹⁰Y is normalized to the 174 standard ⁹⁰Sr solution spiked. The normalized value of all environmental samples is consistent among all samples, which means that the measurement coefficient of the resulting solution of ⁹⁰Sr extracted and 175 purified from environmental samples was equal to that of standard ⁹⁰Sr solution. The purification of 176 177 strontium adequately removed impurities and quenching from the environmental samples and therefore 178 the radioactivity in the samples can be evaluated without the need for quench correction.

Figure 5 shows that the spectrum peak area for 90 Sr changed with the increase of 90 Y in samples through time. Although the total radioactivity of the sample increased to twice the initial value at full 90 Y saturation, the peak area was much larger than twice the initial value at full 90 Y saturation. This result suggests the presence of different measurement coefficients between 90 Sr and 90 Y. Table 2 lists the net CPM of the 90 Sr solution after purification to eliminate 90 Y. The increasing values of the net CPM with increasing corrected equilibrium fraction yielded the α value of 16.7, which can be viewed in Figure 6. The net CPM of 5.83 Bq 90 Sr in equilibrium with 90 Y was 208 ± 2. The values of α , net CPM, and radioactivity of 90 Sr were substituted into Eq. (8) to yield measurement coefficients of 90 Sr and 90 Y of 2.01 ± 0.02 and 33.6 ± 0.3 (net CPM/Bq), respectively. These values provided the radioactivity of 90 Sr in the environmental samples.

Radioactivity of ⁹⁰Sr in the environmental samples was evaluated from aging of ⁹⁰Y using Eq. (9). The increase in net CPM of ⁹⁰Y is listed in Table 3, where the detection limit of net CPM is 5, which was evaluated with Curie's equation [29], and correspondingly 0.14 Bq from the measurement coefficient. Although both leaf samples show negligible radioactivity, both bark samples show the increase in net CPM with aging time. The increase in net CPM with aging provided the radioactivity of ⁹⁰Sr, which was analyzed by the regression analysis and can be viewed in Figure 7. Table 4 lists the resulting specific radioactivity of ⁹⁰Sr and ¹³⁷Cs.

196 The radioactivity ratio of ⁹⁰Sr to ¹³⁷Cs changing with time after the accident provides the migration 197 tendency of both radionuclides in the environment. The radioactivity levels for all leaf samples collected 198 from plants 3 km north of the power plant were below the detection limit. The radioactivity levels for ¹³⁷Cs in all samples were significant, with particularly high levels in bark samples (up to 173 Bq/g dry 199 weight). The radioactivity ratios for the bark samples were 4.2 $\times 10^{-3}$ (north) and 1.2 $\times 10^{-2}$ (south), 200 indicating that the fraction of ⁹⁰Sr to total radioactivity levels was one hundredth or less. Table 5 201 compares radioactivity levels of ⁹⁰Sr and ¹³⁷Cs in soil collected near the power plant before and after the 202 203 nuclear accident from a previously published report [23]. The different release rates of ⁹⁰Sr and ¹³⁷Cs from the nuclear reactors caused the radioactivity of ¹³⁷Cs to increase to more than one hundred times to the 204 pre-accident level while the radioactivity of ⁹⁰Sr increased to only several times the pre-accident level, 205 and accordingly the radioactivity ratios decreased from 0.17 to 8.1×10^{-4} and 4.8×10^{-3} . The radioactivity 206 of bark and soil samples collected in the vicinity of the nuclear power plants showed a larger degree of 207 contamination in the south area and a larger isotropic release of ⁹⁰Sr compared to ¹³⁷Cs because between 208 the north and south area there was difference in one order in radioactivity of 137 Cs while the order of 209 radioactivity ⁹⁰Sr was the same between the north and south area. The radioactivity ratios of ⁹⁰Sr to ¹³⁷Cs 210 in the bark samples were at least 2.4 times those of the soil samples. The radioactivity ratio in soil and 211 212 bark samples would be the same value immediately after the accident due to the same contamination history at each location where the samples were collected. However, the radioactivity ratio of soil and 213 214 bark samples, collected 4 months and 26 months after the accident, respectively, were different from each other, showing the different migration rate of ⁹⁰Sr and ¹³⁷Cs. The increase in the radioactivity ratio in bark 215 samples can be ascribed to the accumulation of ⁹⁰Sr to bark or the diffusion of ¹³⁷Cs from bark. The 216 accumulation of ⁹⁰Sr through root uptake is negligible because the migration rate of ⁹⁰Sr in soil is reported 217 218 to be 0.2 - 1.0 cm/yr [30-32]. The rapid translocation of ¹³⁷Cs from bark into the wood was observed in 219 Fukushima forest [33]. This increase in the radioactivity ratio in bark samples with time shows slower migration of ⁹⁰Sr than ¹³⁷Cs in bark and tree. The effect of different migration would be expected to cause 220 considerable variation in radioactivity ratio from the initial value. The determination of 90 Sr in 221 222 environmental samples is required to avoid suffering from unexpected '90Sr hot spots.'

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- 224

225 Conclusions

226	We measured the specific activity of ⁹⁰ Sr and ¹³⁷ Cs in leaf and bark samples collected in the vicinity
227	of the Fukushima Daiichi Nuclear Power Plants in May 2013. The radioactivity of ⁹⁰ Sr was quantitatively
228	measured by using Cherenkov counting, where purified 90 Sr and 90 Y from the standard solution of 90 Sr
229	were used to evaluate the measurement coefficient of Cherenkov counting and to verify the determination
230	without any quench correction. The radioactivity of both radionuclides in the bark samples was larger
231	than that in the leaf samples. The slower migration rate of 90 Sr compared to 137 Cs was shown by
232	comparison to the radioactivity ratios of both radionuclides in soil samples collected in July 2011. The
233	rough tendency of the migration in trees is shown; however, further analysis is required for the discussion
234	of the migration of ⁹⁰ Sr in the environment.
235	
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295	90 Y (i.e., growth of 90 Y)
296	
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298	corrected equilibrium fraction of ⁹⁰ Y
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301	Daiichi Nuclear Power Plant
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310	SD: 2 σ
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313	concentration of 2 M HNO ₃
314	SD: 2 σ
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317	comparison to the standard ⁹⁰ Sr solution spiked for the same amount of ⁹⁰ Y.
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321	fraction of ⁹⁰ Y.
322	
323	Figure 6 Fitting line defined in Eq. (7) to evaluate the ratio of the detection coefficient of 90 Y and 90 Sr, α ,
324	using the values in Table 2
325	
326	Figure 7 Regression analysis for the evaluation of radioactivity of ⁹⁰ Sr in vegetation samples (closed
327	circle: #1 bark <i>Metasequoia</i> ; open circle: #2 bark <i>Cryptomeria japonica</i>)
328	

329 Table 1

Nuclide	$Maximum E_\beta$	Emission
	(MeV)	ratio
⁴⁰ K	1.311	
⁸⁹ Sr	1.495	
⁹⁰ Sr	0.546	
⁹⁰ Y	2.280	
¹³⁴ Cs	0.658	70.2%
	0.415	2.5%
	0.089	27.3%
¹³⁷ Cs	1.176	5.6%
	0.514	94.4%

331

333	Table 2
555	14010 2

Aging Time (hr)	Counting Time (hr)	X^*	Net CPM
0.4	0.5	0.007	28.5 ± 1.5
44.4	0.5	0.383	174.0 ± 2.6
81.7	0.5	0.588	260.8 ± 3.2
123.6	0.5	0.738	323.2 ± 3.5
621.4	0.5	0.999	438.2 ± 4.0

335 **X*: Corrected Equilibrium Fraction

336

#1 Bark Metasequo	ia

1		
Aging Time (hr)	X^*	Net CPM
112.0	0.71	23.38 ± 0.98
146.7	0.80	25.00 ± 0.99
216.5	0.91	28.12 ± 1.02
289.1	0.96	29.59 ± 1.03
327.9	0.97	30.17 ± 1.03
371.4	0.98	30.08 ± 1.03

 $A=25.91\pm1.24$

 $B=4.73\pm1.11$

#2 Bark Cryptomeria japonica

Aging Time (hr)	X^*	Net CPM
131.8	0.76	6.34 ± 0.82
170.6	0.85	7.41 ± 0.83
214.1	0.90	8.34 ± 0.83
307.5	0.96	8.43 ± 0.84

 $A=11.08\pm2.13$

 $B=-2.00\pm1.85$

#3 Leaf Cryptomeria japonica

Aging Time (hr)	X^*	Net CPM
136.0	0.78	1.81 ± 0.77
174.8	0.85	1.92 ± 0.77
218.3	0.91	2.02 ± 0.77
311.7	0.97	2.15 ± 0.77

Aging Time (hr)	<i>X</i> *	Net CPM
49.8	0.43	1.51 ± 0.76
103.8	0.68	1.46 ± 0.75
147.5	0.80	0.88 ± 0.76

X*: Corrected equilibrium fraction 340

341 Detection limit: net CPM = 5

342

Table 4

345

Sample	Location*	⁹⁰ Sr (Bq/g)	¹³⁷ Cs (Bq/g)	⁹⁰ Sr / ¹³⁷ Cs
#1 Bark Metasequoia	2 km south	0.73 ± 0.04	172.9 ± 2.0	$(4.2 \pm 0.2) \times 10^{-3}$
#2 Bark Cryptomeria japonica	3 km north	0.31 ± 0.06	26.9 ± 0.5	$(1.2 \pm 0.2) \times 10^{-2}$
#3 Leaf Cryptomeria japonica	3 km north	< DL**	10.4 ± 0.2	
#4 Leaf Artemisia	3 km north	< DL**	4.9 ± 0.2	

346 * Location: Distance from the Fukushima Daiichi Nuclear Power Plant Unit No. 1

347 ** DL: 0.11 Bq/g

348

Location*: 2 km south

Sampling Year	2005	July 2011
⁹⁰ Sr (Bq/g)	ND**	0.0808
¹³⁷ Cs (Bq/g)	ND**	99.7
⁹⁰ Sr/ ¹³⁷ Cs		8.1 ×10 ⁻⁴

352

Location*: 3 km north					
Sampling Year	2005	July 2011			
⁹⁰ Sr (Bq/g)	0.0030	0.0149			
¹³⁷ Cs (Bq/g)	0.0173	3.08			
⁹⁰ Sr/ ¹³⁷ Cs	0.17	4.8 ×10 ⁻³			

353 Location*: Distance from the Fukushima Daiichi Nuclear Power Plant Unit No. 1

354 ND**: Not detected

355

- 357 Figure 1











373 374









