Isotopic signature and nanotexture of cesium-rich microparticles: Release of uranium and fission products from the Fukushima Daiichi Nuclear Power Plant

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Highly radioactive cesium-rich microparticles (CsMPs) released from the Fukushima Daiichi Nuclear Power Plant (FDNPP) provide nano-scale chemical fingerprints of the 2011 tragedy. U, Cs, Ba, Rb, K, and Ca isotopic ratios were determined on three CsMPs (3.79–780 Bq) collected within ~10 km from the FDNPP to determine the CsMPs’ origin and mechanism of formation. Apart from crystalline Fe-pollucite, CsFeSi2O6·nH2O, CsMPs are comprised mainly of Zn–Fe-oxide nanoparticles in a SiO2 glass matrix (up to ~30 wt% of Cs and ~1 wt% of U mainly associated with Zn–Fe-oxide). The 235U/238U values in two CsMPs: 0.030 (±0.005) and 0.029 (±0.003), are consistent with that of enriched nuclear fuel. The values are higher than the average burnup estimated by the ORIGEN code and lower than non-irradiated fuel, suggesting non-uniform volatilization of U from melted fuels with different levels of burnup, followed by sorption onto Zn–Fe-oxides. The nano-scale texture and isotopic analyses provide a partial record of the chemical reactions that occurred in the fuel during meltdown. Also, the CsMPs were an important medium of transport for the released radionuclides in a respirable form.

Radionuclides with ~520 PBq initial total activity were released from the Fukushima Daiichi Nuclear Power Plant (FDNPP) as a result of the nuclear disaster that occurred after the Tohoku earthquake on March 11, 20111. The released radionuclides, including noble gases (Xe and Kr) and volatile fission products (I, Cs, Te, Sb, and Ag), contaminated the surface over ~14,000 km2 surrounding the FDNPP1–5, requiring the evacuation of some 100,000 residents. Radioactive Cs, 134Cs and 137Cs, are the most important radionuclides contributing to the high radiation in the environment near the FDNPP at present because of their relatively short half-lives, 2.06 and 30.07
years, respectively. About 1–7% of the Cs inventory of three reactor cores was released. Previous studies of the
distribution and migration of radioactive Cs in the surface environment around Fukushima indicated that
an initially soluble form of radioactive Cs was released from the damaged reactors and spread over the Fukushima
Prefecture and the surrounding area through dry and wet deposition. Subsequently, a soluble form of Cs was
tightly bound to the interlayers of clays, such as vermiculite, and remained within the top ~5 cm of the soil. However,
Cs in the contaminated soils is heterogeneous and concentrated locally at the micron scale as hot spots,
as revealed through autoradiography, and this heterogeneity has not been fully described. A possible cause
of the heterogeneity is the formation of Cs-rich microparticles (CsMPs), with a high Cs radioactivity per unit
mass, found at a range of distances from the FDNPP. This is another important route of Cs migration in the
environments. Different from the soluble Cs, the CsMPs are sparingly soluble in water. The $^{134}$Cs/$^{137}$Cs radioac-
tivity ratio of ~1 indicates that they originated from the FDNPP. CsMPs were initially considered as amorphous
glass particles containing various elements derived from the reactors that were melted into the glass matrix. However,
recent studies have shown that the Cs concentrations are associated with discrete Zn–Fe-oxide nano-
particles embedded in a pure SiO2 glass matrix, as well as numerous nano-scale inclusions with a variety of fission
products. The nano-texture within the CsMPs records the chemical reactions that took place during the melt-
downs inside the reactors. The sparingly soluble CsMPs were identified as dominant Cs carriers during the initial
Cs fallout in Tokyo, Japan, and they are expected to differ from soluble Cs in terms of environmental mobility and
health impacts. In addition to the reaction sequence of the CsMPs formation inside the reactors, trace amounts
of U are present in CsMPs. The origin of U and the other nuclides remains uncertain, as insulating material in the
reactors also contains trace amount of natural U. The occurrence of U derived from inside the reactors is poten-
tially useful for understanding the reactions that nuclear fuels experienced during meltdown and even the status
of the melted nuclear fuels in the damaged reactors. This is crucial for developing an appropriate decommission-
ing strategy. The hypothesis of this study is that the isotopic ratio analysis in combination with atomic-resolution
analysis reveals the source of U and the release processes associated with CsMPs from the FDNPP. The CsMPs are a very unique form of condensed matter that was created during the meltdown events inside the reactors at the FDNPP, very different from general concept of Cs release as a soluble Cs species, such as CsI and CsOH. Isotopic analyses of the individual CsMPs are reported for the first time. Isotopic analysis of the other nuclides, stable and radiogenic, also allow us to identify their sources, natural or fissionogenic.

Results

Shape, composition, and radioactivity of CsMPs. Four CsMPs were found at three localities, which are hereafter labeled as OTZ3, OTZ10, KOI2, and OMR1 (Fig. 1). Detailed information on the sampling is provided in the method section. Figure 2 shows secondary electron scanning electron microscopy (SEM) images and energy-dispersive X-ray analysis (EDX) elemental maps of major elements of the CsMPs. In previous studies, the shape of CsMPs has been described as spherical. In contrast, these particles appear as irregularly shaped aggregates rather than spheres. In addition, the radioactivity of the presently examined CsMPs ranges from 3.79
to 780 Bq (Table 1). These values are approximately thirty three times higher than that of spherical CsMPs. The $^{134}$Cs/$^{137}$Cs radioactivity ratios of the OTZ3, KOI2, and OMR1 used for the SIMS analysis are 1.06–1.08 (average 1.07), which approximately corresponds to ~26 GWd/tU according to OrigenArp calculations. Although the $^{134}$Cs/$^{137}$Cs isotopic ratio values are close to those of the reactor Units #2 and #3 in the FDNPP, the specific reactor source could not be determined based only on the isotopic ratios, because burnups of the irradiated fuels in each reactor are heterogeneous depending on the positions of the fuel assemblies.

**Isotopic ratio based on secondary ion mass spectrometry (SIMS) and γ spectrometry.** The results of the isotopic analysis of the three CsMPs, OTZ3, KOI2, and OMR1, are summarized in Table 2. In general, the $^{235}$U/$^{238}$U isotopic ratio in nature is 0.00729, whereas the ratio in non-irradiated nuclear fuel is typically >~0.03 due to enrichment of $^{235}$U. As shown in Table 2, U isotopes were successfully determined to be 0.029584 and 0.029341 in OTZ3 and KOI2, respectively, whereas U was not detected in OMR1. The U isotopic ratio for the standard specimen NIST SRM610 is comparable to the value in a previous report, and the deviation of the

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**Figure 2.** SEM images of CsMPs associated with the elemental maps.
The large Cs-rich particle was subsequently identified as an Fe-rich pollucite structure (CsFeSi$_2$O$_6$·nH$_2$O, $<_{16}$), with a high Cs-content (Fig. 3e) as compared with the franklinite nanoparticles found in the low Cs-zone. The high Cs zone (represented by edx2 in Fig. 3b) appears to contain relatively large nanoparticles, ~50 nm in size, with a high Cs-content (Fig. S1a) as compared with the franklinite nanoparticles found in the low Cs-zone. The large Cs-rich particle was subsequently identified as an Fe-rich pollucite structure (CsFeSi$_2$O$_6$·nH$_2$O, $<_{16}$) based on the SAED pattern (Fig. S1b). Pollucite is a zeolite with the ideal formula (Cs,Na)$_2$Al(Si$_2$O$_5$)$_2$·4H$_2$O. The Fe-pollucite is unstable under electron beam irradiation and becomes amorphous (Fig. S1c), possibly due to the loss of water. In addition, the HAADF-STEM image of the boundary between the high- and low-Cs zones displays different levels of contrast after electron beam irradiation (Fig. 3d). The magnified HAADF-STEM image of the low Cs zone (dark contrast) is composed of nanoparticles that consist of Zn and Fe, as revealed by the elemental maps (Fig. 3e). Cesium is associated with these nanoparticles. The distribution of silicon could not be resolved due to its small particle size, with a high Cs-content (Fig. 3e, Table S1). The area with dark contrast (edx1) contains a lower Cs-content as compared with the zone with bright contrast (edx2) (Fig. 3d). The CsMP is decay daughters of radioactive Cs. The iron-bearing phases rather than forming a separate Ba-rich phase, which is expected since almost all Ba isotopes in the CsMP are decay daughters of radioactive Cs. The iron-bearing phases rather than forming a separate Ba-rich phase, which is expected since almost all Ba isotopes in the CsMP are decay daughters of radioactive Cs. The iron $L$-edge confirms that Fe is present in the form of oxides.

| Sample | Particle size (μm) | Radioactivity of $^{134}$Cs (Bq) | Radioactivity of $^{137}$Cs (Bq) | Ratio of $^{134}$Cs/$^{137}$Cs radioactivity |
|--------|------------------|-------------------------------|-------------------------------|---------------------------------|
| OTZ3   | 17.3             | 401±0.37                      | 379±0.50                      | 1.06                            |
| OTZ10  | 2.2              | 1.85±0.080                    | 1.94±0.031                    | 0.955                           |
| OMR1   | 5.2              | 21.5±0.50                     | 20.3±0.19                     | 1.06                            |
| KOI2   | 4.4              | 18.2±0.16                     | 16.8±0.05                     | 1.08                            |

Table 1. Radioactivity of four CsMPs described in the present study. The radioactivity was decay-corrected to March 12, 2011, 15:36 JST. The uncertainties in the radioactivity analyses are given in the parenthesis. The particle size was measured using the SEM image.

In order to determine the presence of U, a FIB-prepared cross-section of the OTZ10 CsMP was additionally investigated by TEM (Fig. 5a). Typically, the SAED exhibits broad diffraction maxima that correspond to diffuse scattering from amorphous domains (Fig. 5a inset). The elemental map of the entire cross-section displays a homogeneous distribution of the major constituents at the bulk scale (Fig. 5b). The Cs-concentration is 7–10 wt% (Fig. 5c, Table S1). Despite the apparently homogeneous distribution and the diffuse scattering halo in the SAED, the magnified HAADF-STEM image shows phase separation as evidenced by differences in contrast (Fig. S2a).
Table 2. Summary of isotope analysis for three CsMPs using SIMS and γ spectrometry. A NIST standard, SRM610, which contains depleted U, was also analyzed as a reference. Ten scans were conducted for each analytical spot on the CsMPs, meaning that the sequence of ten analyses represents a depth profile of the variation in the isotopic ratios; thus, SIMS analyses provide the isotope signatures inside the CsMPs. The average values are given in the table with the standard deviation calculated for the ten analyses and the standard deviations are given in the parenthesis as σ. The isotopic ratios recalculated from the results of γ spectrometry are annotated as (γ). Conversion of radioactivity to isotopic ratio was conducted by decay-correcting to the time of SIMS analysis. n/d stands for not detected.

| Isotopic ratio | OTZ3 (1σ) | KOI2 (1σ) | OMR1 (1σ) | Natural abundance ratio | SRM610 (1σ) |
|----------------|------------|------------|------------|-------------------------|-------------|
| 235U/238U      | 0.029584 (0.0004967) | 0.029341 (0.0003032) | n/d        | 0.00729                 | 0.002388 (0.000001) |
| 134(Ca + Ba)/134(Ca + Ba) | 0.022488 (0.000219) | 0.03184 (0.00107) | 0.03167 (0.00037) | —                       | —           |
| 134Ca/136Ca (γ) | 0.0131      | 0.0135      | 0.0132     | —                       | —           |
| 134Ba\textsubscript{fissionogenic}/134Ba\textsubscript{fissionogenic} (γ) | 0.515       | 0.517       | 0.520      | —                       | —           |
| 134Ca/136Ba\textsubscript{fissionogenic} (γ) | 0.190       | 0.190       | 0.191      | —                       | —           |
| 134Ba/138Ba (γ) | 7.47        | 7.46        | 7.47       | —                       | —           |
| 134(Ca + Ba\textsubscript{fissionogenic})/137(Ca + Ba\textsubscript{fissionogenic}) (γ) | 0.0724      | 0.0742      | 0.0727     | —                       | —           |
| 127Cs/127Ba (γ) | 0.3929 (0.00889) | 0.4007 (0.0050) | 0.4011 (0.0020) | —                       | —           |
| 136Ba/138Ba (γ) | 0.3961 (0.0121) | 0.3685 (0.0062) | 0.3777 (0.0025) | —                       | —           |
| 134Ba/136Ca (γ) | 0.9961 (0.0543) | 0.9199 (0.0246) | 0.9417 (0.0102) | —                       | —           |
| 134Ca/136Ba (γ) | 36.6 (3.7)  | 35.3 (5.9)  | 35.3 (5.9) | —                       | —           |
| 138Ba/136Ba (γ) | 0.1523 (0.0124) | 0.1638 (0.0191) | 0.4293 (0.0697) | 0.1095                 | 0.1097 (0.0002) |
| 87Sr/86Sr (γ)   | 2.407 (0.014) | 2.284 (0.017) | 2.236 (0.120) | 0.3856                 | 0.3857 (0.00009) |
| 87Sr/86Sr (γ)   | n/d         | n/d         | n/d        | —                      | —           |
| 140K/86Sr (γ)   | 0.007217 (0.000001) | —       | —       | 0.07216                 | 0.07214 (0.00003) |
| 140K/86Sr (γ)   | 0.000127 (0.0000041) | —       | —       | 0.000125               | —           |
| 133Ba/134Ba (γ) | 0.06465 (0.00115) | —       | —       | 0.06472                 | 0.06469 (0.000005) |

The elemental map and the EDX analysis indicate that the phase in bright contrast contains Fe, Zn, Cs, and Sn; whereas, the area of dark contrast corresponds to SiO₂ -domains (Fig. 5b and c). The HRTEM image also reveals the presence of many nanocrystals (Fig. 5d). Based on the d-spacing and the FFT image, these nanocrystals are franklinite. Another magnified HAADF-STEM image with an elemental map reveals that the U distribution is closely associated with Zn–Fe-oxide nanoparticles (Fig. 5e). The U concentration is 0.8 wt% (including Si) or 1.1 wt% (excluding Si), assuming U is in the form of U₃O₈ for the quantitative EDX analysis (edx2, Fig. 5f).
ratios, $^{134}\text{Cs}/^{134}\text{Ba}_{\text{radiogenic}}(\gamma)$ and $^{137}\text{Cs}/^{137}\text{Ba}_{\text{radiogenic}}(\gamma)$ at the time of the SIMS analysis, were ~0.19 and ~7.47, respectively (Table 2).

Since the amount of volatilized Ba is negligible and the Ba in the CsMPs is mostly radiogenic, $^{135}\text{Cs}/^{133}\text{Cs}$ determined by SIMS represents the initial isotopic ratio, because $^{133}\text{Cs}$ is a stable isotope and $^{135}\text{Cs}$ has a long half-life, 2.3 million years. The previous ORIGEN calculation reported $^{135}\text{Cs}/^{133}\text{Cs}$ to be 0.388, 0.344, and 0.353 for Units # 1, 2, and 3, respectively, which are close to the present SIMS results, 0.39, 0.37, and 0.38 for OTZ3, KOI2, and OMR1, respectively. The isotopic ratios $^{135}\text{Cs}/^{137}\text{Cs} + ^{137}\text{Ba}_{\text{radiogenic}}$ and $^{137}\text{Cs} + ^{137}\text{Ba}_{\text{radiogenic}})/^{133}\text{Cs}$ calculated by SIMS are 0.39–0.40 and 0.92–0.99, respectively, which are comparable to the initial $^{135}\text{Cs}/^{137}\text{Cs}$ and $^{137}\text{Cs}/^{133}\text{Cs}$ values calculated by ORIGEN, 0.40 and 0.98, 0.34 and 1.01, 0.35 and 1.01, for Units #1, 2, and 3, respectively. Moreover, $^{135}\text{Cs}/^{137}\text{Cs} + ^{137}\text{Ba}_{\text{radiogenic}}$ is also comparable to $^{135}\text{Cs}/^{137}\text{Cs}$ of the bulk soil samples previously analyzed.

**Figure 3.** (a) A HAADF-STEM image of FIB-TEM specimen of OTZ3-1 CsMP. (b) Magnified HAADF-STEM image of the area indicated by the arrow in (a). (c) HAADF-STEM image (labeled as ADF) and the elemental map of the area displayed in (b). (d) STEM-EDX spectrums of the area indicated by the yellow square in (b). (e) Magnified HAADF-STEM image of the dark contrast zone represented as edx1 in (b) associated with elemental maps. (f) HRTEM image of the Zn–Fe-oxide nanoparticle. The lattice fringe of 2.9 Å corresponds to (220).
using thermal ionization mass spectrometry (TIMS), 0.36–0.38. Because of the high $^{137}\text{Cs}/^{137}\text{Ba}_{\text{radiogenic}}$, the majority of 137-mass derives from $^{137}\text{Cs}$, and the efficiency of secondary ions formation for $^{137}\text{Ba}$ should be minimal.

The values of $^{134}(\text{Cs} + \text{Ba}_{\text{radiogenic}})/^{137}(\text{Cs} + \text{Ba}_{\text{radiogenic}})$ (γ) determined through gamma spectrometry (0.0724–0.0742), which are the same as those of the initial $^{134}\text{Cs}/^{137}\text{Cs}$, are higher than the values of $^{134}(\text{Cs} + \text{Ba})/^{137}(\text{Cs} + \text{Ba})$ for CsMPs that were determined by SIMS (0.0249–0.0318). This is attributed to the difference in ionization efficiency between Cs and Ba. Introducing the secondary ion production efficiency coefficient, $i_f = \text{Ba ionization efficiency}/\text{Cs ionization efficiency}$, where i stands for the mass, the ratio by SIMS can be expressed as

$$\frac{^{134}\text{Cs} + 134\text{Ba}_{\text{radiogenic}} \times 134f}{137\text{Cs} + 137\text{Ba}_{\text{radiogenic}} \times 137f}$$

= 0.029 (average for three CsMPs)

(1)

Because the difference between $^{134}f$ and $^{137}f$ is negligible, the ratio of the ionization efficiency is simply expressed as $f$, and equation (1) can be rewritten as

$$\frac{^{134}\text{Cs} + 134\text{Ba}_{\text{radiogenic}} \times f}{137\text{Cs} + 137\text{Ba}_{\text{radiogenic}} \times f} = \frac{^{134}\text{Cs}/^{137}\text{Cs}_{\text{initial}} \times (0.16 + 0.84 \times f)}{(0.88 + 0.12 \times f)}$$

= 0.029

(2)

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**Figure 4.** (a) HAADF-STEM image of the FIB-TEM specimen OTZ3-2 associated with the elemental maps. (b) A magnified HAADF-STEM image of the high Cs phase indicated by the arrow in (a). (c) A representative EDX spectrum of the high Cs phase indicated by the arrow in (a). (d) HRTEM image of the high Cs phase indicated by the arrow in (a). (e) The SAED pattern of the high Cs phase from two different major zone axes.
Solving equation (2): \( f = 0.24 \).

This ionization efficiency factor is reasonably close to that determined for rhyolitic glass (SRM610); 0.288. Note, the ionization efficiency of the standard specimen is determined on the polished flat surface on the pure glass, while that for the CsMP was determined on the spherical shaped particle that is a mixture of SiO\(_2\) glass and the Zn–Fe-oxides nanoparticles. Thus, the matrix composition is similar but not identical.

The Rb isotopic ratio, \(^{87}\text{Rb}/^{85}\text{Rb}\), is 2.2–2.4, which is obviously higher than the natural \(^{87}\text{Rb}/^{85}\text{Rb}\) (0.3856), indicating that the Rb isotopes are fissionogenic. Compared with the \(^{87}\text{Rb}/^{85}\text{Rb}\) values calculated by ORIGEN\(^{19}\) (~1.7), \(^{87}\text{Rb}/^{85}\text{Rb}\) obtained by SIMS is relatively high. Nevertheless, our OrigenArp calculation gives ratio values between 2.5 and 2.7, which are closer to the \(^{87}\text{Rb}/^{85}\text{Rb}\) values by SIMS. The difference is not ascribed to the mixture of natural Rb isotopes because the mixing with the natural isotopes would lower the \(^{87}\text{Rb}/^{85}\text{Rb}\) isotopic ratio, but rather due to the local volatilization of Rb.

The nanoscale texture of OTZ10 and the low-Cs zone in OTZ3-1 reveal that franklinite nanoparticles are embedded in a glassy matrix of SiO\(_2\). These textures and the Cs association with franklinite were also reported in previous studies\(^{15,16}\), indicating multiple steps in the formation sequence: radioactive Cs is released to form nanoparticles and/or is present as droplets in a mist during meltdown; subsequently, numerous Zn–Fe-oxide nanoparticles are embedded in the glassy matrix of SiO\(_2\).

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**Figure 5.** (a) HAADF-STEM image of the FIB-TEM specimen of the OTZ10 CsMP with the SAED as inset. (b) Elemental maps of the same area as the HAADF-STEM image in (a). (c) The EDX spectrum of the area indicated by the yellow square labeled as edx1 in (a). (d) A HRTEM image of the Zn–Fe-oxides nanoparticles accompanied with the magnified lattice image and the FFT image of the selected area indicated by the white square. The lattice fringe of 2.2 Å corresponds to (400). (e) HAADF-STEM image of the enlarged figure of the thin edge in (a) associated with elemental maps. (f) STEM-EDX point analysis on a Zn–Fe-oxide nanoparticle indicated by a red cross labeled as edx2 in (e).
(franklinite) nanoparticles formed during the failure of the reactor pressure vessel (RPV), and Cs in wet form is adsorbed onto the Zn–Fe-oxide nanoparticles; then, the melted fuels break through the RPV, hit the concrete pedestal, and generate SiO gas at ~2,000 K. This process is known as the molten core-concrete interaction (MCCI)\(^{29}\), which immediately leads to the condensation of SiO\(_2\) over the Zn–Fe-oxide nanoparticles and the incorporation of fission product nanoparticles. As shown in a previous study\(^{40}\), a low concentration of U (~1 wt%) is observed associated with franklinite nanoparticles that formed before the CsMP formation. Moreover, U adsorption onto the franklinite nanoparticles possibly occurs through volatilization of the oxidized form of the UO\(_2\) fuel. In fact, UO\(_2\) pellets can be oxidized depending on the H\(_2\)O:H\(_2\) ratio\(^{30}\). In general, U is considered a nonvolatile element; however, the partially oxidized form of U\(_{2+}\) can volatilize by ~10% at ~1,900 K at a heating rate of ~30 K/min; whereas, the unoxidized form U\(_0\) does not become volatile even at 2,700 K\(^{31,32}\). Uranium association with microscale aerosols during a severe accident was also recognized in some experimental studies\(^{33}\). The chemical form of U inside CsMPs has been determined in the OTZ10 CsMP, as well as in the other CsMP separated from the same soil sample\(^{34}\). The U isotopic ratio in another CsMP, OTZ3, found in the same soil sample, was determined. \(^{235}\text{U}/^{238}\text{U} = ~0.030\), as analyzed from the two CsMPs, indicating that the U incorporated in the CsMPs is not from naturally occurring trace U that has a \(^{235}\text{U}/^{238}\text{U}\) ratio of 0.00729, which was initially considered as the source of U because trace amounts of U are included in the insulating materials surrounding reactor pressure vessel. Rather, nuclear fuel is the most likely origin of U in the CsMPs. The original isotope composition of non-irradiated fuel before operation at the FDNPP was 0.0389\(^{19}\). Based on calculations using the ORIGEN code, the U isotopic ratios for the irradiated fuel in Units # 1, 2 and 3 were 0.0172, 0.0193, and 0.0192\(^{18}\), respectively. Thus, the isotopic ratios are between the isotopic ratio calculated from burnup and that of the non-irradiated fuel. The isotopic ratios calculated by the ORIGEN code are based on the average burnup for all fuel assemblies. The intermediate values of \(^{235}\text{U}/^{238}\text{U}\) are due to the fact that the burnup and the temperature are not homogeneous within the reactor core. A fuel assembly will experience different temperatures depending on its position in the core. Even within a single pellet, there is a thermal gradient; the center of the pellet can be heated as high as ~1,973 K, whereas the temperature of the rim is only ~673 K\(^{34}\). Moreover, the burnup within a single pellet is not uniform with low burnup at the center and high burnup at the edge\(^{34}\). The burnup of the fuel assembly is not uniform either, as the fuel rods with different burnup are assembled in a certain manner. Thus, the fuel rods with relatively lower burnup might have been heated at the temperature at which a small amount of U was volatilized, resulting in the high \(^{235}\text{U}/^{238}\text{U}\) isotopic ratios for both CsMPs. The volatilized U is associated with Zn–Fe-oxide, most likely by simple adsorption, as evidenced by the U in the OTZ10 CsMP.

Dispersion of U released from the FDNPP has been reported in previous studies\(^{35–37}\), including \(^{236}\text{U}/^{238}\text{U}\) values of ~10\(^{-9}\) in paddy-field water and ocean water as far as ~30 km from the FDNPP\(^{36}\). \(^{239}\text{U}/^{238}\text{U}\) values of ~10\(^{-7}\) were also reported in black-colored dust samples collected within a ~30-km distance\(^{38}\). This evidence that a small amount of U, about 150 g\(^{36}\), was discharged from the fuels in the reactors. Nevertheless, another study measuring \(^{238}\text{U}/^{235}\text{U}\) failed to identify the FDNPP as the origin due to the dilution by natural U isotopes\(^{37}\). However, all those analyses were performed on bulk soil samples and did not include data from single CsMPs. Thus, the speculation of U was never determined. The present study convincingly identifies the U speciation as associated with Zn–Fe-oxide nanoparticles at ~1 wt%, and the nuclear fuel origin of U in CsMPs is confirmed through isotopic ratio analyses. Yet, quantitative analysis of the amount of U in the form of CsMPs in the total U that was dispersed to environment remains undetermined.

Two different occurrences of radioactive Cs have been identified: Fe-pollucite and Cs associated with franklinite nanoparticles embedded in the glassy SiO\(_2\) matrix. The latter has been already described above and in previous studies\(^{15,16}\) and is not discussed further here. On the other hand, Fe-pollucite, CsFeSi\(_2\)O\(_6\)·nH\(_2\)O, is identified for the first time in CsMPs. None of the previous studies reported pollucite during meltdown; however, a recent experimental study reported formation of pollucite and CsFeSi\(_2\)O\(_6\) during the CsOH chemisorption onto stainless steel containing 5% Si\(^{38}\). Both materials have a zeolite structure that can be synthesized by annealing a mixture of gels\(^{39}\) and appropriate elements, such as Cs. Their formation by chemisorption likely occurred at the high-Cs zone during the formation of CsMPs. However, the reaction required for CsMPs formation are apparently different from the simple interaction of CsOH with stainless steel at ~1,273 K\(^{38}\). In fact, the Cs–Cr phases that result from the reaction with stainless steel are not present in CsMPs. Rather, it is probable that CsOH is involved in the chemical reactions with Si and Fe oxide during the MCCI at >2,000 K. The clear boundary between the high- and low-Cs zones (Fig. 3b) indicates that Zn–Fe-oxide nanoparticles formed aggregates prior to the formation of Fe-pollucite, which formed at the time of MCCI.

In summary, the present study successfully determined the origin of U, Cs, Ba, Rb, K, and Ca in the CsMPs, as well as their chemical form based on the isotopic ratios and their nanoscale structure. In particular, the data obtained are critically important for delineating the reactions involving nuclear fuels, which occurred inside the reactor during the formation of the CsMPs. The detailed information on the chemical state and source provide an understanding of the source term at the FDNPP during the meltdown event. This is important information for severe accident (SA) analysis codes, such as MELCOR\(^{41,42}\) and MAAP\(^{43}\), since some of the phases identified in the CsMPs, such as Zn–Fe-oxide and pollucite, are not considered in these severe accident codes\(^{44}\).

Methods

Sampling. We collected the samples at three locations (Fig. 1). As for the sample labels, the capital letters represent the locality and the following number indicate the number in our CsMP database in Kyushu University. The OTZ3 and OTZ10 were separated from the same soil collected from the top ~1 cm of the paddy soil at Ottozawa located ~4 km west from the FDNPP in Okuma Town, Futaba County, Fukushima, on March 16, 2012. The soil was mainly composed of clay minerals, quartz, and feldspar\(^{45}\). Because it is still prohibited to enter the area due to the high radiation dose, the locality remained untouched and was not disturbed by decontamination or restoration. The radiation dose ~1 m above the ground was measured at 84 μSv/h.
The second sample (KOI2) was composed of gravels collected under the drainpipe of the assembly house in Koirino located 2.9 km southwest from the FDNPP during the same sampling campaign. The radiation dose underneath the drainpipe was extremely high compared to the surroundings; the dose at the sampling point was as high as 630 μSv/h. The gravel samples were carefully collected from the ground surface using a hand shovel and placed in plastic bags. The soil was also mainly composed of clay minerals, quartz, and feldspar. The third sample (OMR1) was collected under the drainpipe of the warehouse in Omaru located in Namie Town, Futaba County, Fukushima, ~10.5 km northwest from the FDNPP during the sampling campaign on December 20, 2012. The radiation dose ~1 m above the ground exceeded 30 μSv/h.

**Separation of CsMPs.** Prior to the procedure, both samples were sieved through a 114-μm mesh. The powder samples were dispersed on grid paper and then covered with a plastic sheet. Next, an imaging plate (IP, Fuji film, BAS-SR 2025) was placed on the samples for 5–15 min. Further, using an IP reader, autoradiograph images with pixel sizes of 100 μm were recorded. After identifying the positions of intense radioactive spots, droplets of pure water were added to these positions and then drawn using a pipette to produce suspensions with small amounts of soil particles by dilution with pure water. This procedure was repeated until the suspension did not contain a significant amount of soil particles. Subsequently, the positions containing hot spots were selected using pieces of double-stick carbon tape that were cut as small as possible with a blade. The pieces were checked by autoradiograph imaging to obtain CsMPs with maximum efficiency using scanning electron microscopy (SEM) observations. Prior to SEM analysis, the pieces were placed on an aluminum plate and coated with carbon using a carbon coater (SANYU SC-701C). The CsMPs were found and observed using two SEMs (Shimadzu, SS550 and Hitachi, SU6600) using acceleration voltages of 15–25 kV both equipped with an energy dispersive X-ray spectrometer (EDX, EDAX Genesis).

**Preparation of TEM specimens.** A focused ion beam (FIB) instrument (FEI, Quanta 3D FEG 200i Dual Beam) was utilized to prepare a thin foil of individual CsMPs with diameters of a few μm. Gallium was used as an ion source, and W deposition was used to minimize the damage from the ion bombardment. The current and the semilunar-shaped Cu grid for FIB and further thinned by an ion beam operating at 5 kV.

**TEM analysis.** HRTEM with EDX and high-angle annular dark-field scanning transmission electron microscopy were performed using a JEOL JEM-ARM200F and JEM-ARM200CF with an acceleration voltage of 200 kV. The JEOL Analysis Station software was used to control the STEM-EDX mapping. To minimize the effects of sample drift, a drift-correction mode was used during acquisition of the elemental map. The STEM probe size was ~0.13 nm, generating an ~140-pA current when 40 μm of the condenser lens aperture was inserted. The collection angle of the HAADF detector was ~97–256 mrad. A Gatan imaging filter system was used to conduct electron energy-loss spectroscopy in STEM mode. The convergent and collection semi-angles were ~30 mrad and ~17 mrad, respectively.

**Gamma spectrometry.** The 134Cs and 137Cs radioactivities of the CsMPs were determined using gamma spectrometry. The radioactivity of an additional microparticle with a size of ~400 μm obtained from surface soil in Fukushima was precisely determined at the radioisotope center in Tsukuba University, Japan, and utilized as a standard point specimen for 134Cs and 137Cs. The radioactivity of the point source standard was 23.9 Bq for 134Cs and 94.6 Bq for 137Cs as of September 29, 2015. Measurement of the radioactivity was performed on the CsMPs and the point source standard using a germanium semi-conductor detector, GMX23 (SEIKO E&G), GMX40 (SEIKO E&G) and GX6020 (Canberra) at the center for radioisotopes in Kyushu University, Japan. The acquisition times were 21,238 s for OMR1 and 84,289 s for OTZ10 using GMX23, 4,004 s for OTZ3 using GMX40, and 77, 880.8 s for KOI2 using GX6020.

**Secondary ion mass spectrometry.** Isotopic ratio analysis was performed using secondary ion mass spectrometry (SIMS, SHRIMP-II, Australian Scientific Instruments) at the National Institute of Polar Research, Tokyo. The SIMS specimens were put on Al plates or a Cu grid and fixed on 1-inch slide glass by Cu tape. They were coated with Au at a 13.5-nm thickness prior to the analysis. An O2− primary ion beam of 0.2–0.4 nA was used to sputter the specimen surface with a beam diameter of 3.0–7.0 μm. The typical mass resolution is about ~0.13 nm, generating an ~140-pA current when 40 μm of the condenser lens aperture was inserted. The collection angle of the HAADF detector was ~97–256 mrad. A Gatan imaging filter system was used to conduct electron energy-loss spectroscopy in STEM mode. The convergent and collection semi-angles were ~30 mrad and ~17 mrad, respectively.

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Author Contributions
S.U. conceived the idea, designed all experiments, and wrote the manuscript. J.I. performed measurements, data analysis, and wrote the first draft. A.O. conducted TEM analysis. G.F., K.H., and M.T. performed SIMS analysis. M.S. and R.I. performed SEM analysis. T.O. and K.N. provided navigation during field research in Fukushima. S.Y. performed gamma spectroscopy at Tsukuba University. G.T.W.L., B.G., and R.C.E. participated in the discussion and interpretation of the results and helped to write the paper.

Additional Information
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