**Variation of atmospheric 137Cs and possible carriers in aerosol samples obtained in Namie in a heavily contaminated area of Fukushima prefecture in 2019**

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# Abstract

A lot of radionuclides were released into the environment from Fukushima Daiichi Nuclear Power Plant (FDNPP) accident on March 11, 2011. Because of the long half-life (30.12 years) and high-concentration deposition about 137Cs, the study regarding on the distribution of 137Cs in aerosol samples and the understanding carriers of 137Cs became a hot topic in the recent decade. However, even nine years after FDNPP accident, the fluctuations of atmospheric 137Cs can be still observed, and explanations about the fluctuations and their carriers remain elusive. In this study, a small fluctuation within 0.0002 Bqm-3 from January to April and a slightly higher level of atmospheric 137Cs from May to September was still observed in the aerosol samples obtained in Namie in a heavily contaminated area of Fukushima prefecture in 2019. Therefore, new observations, obtaining by fluorescent upright microscope and scanning electron microscopes (SEM) equipped with an energy dispersive X-ray spectrometer (EDS), quantitatively demonstrated thatthe carriers of 137Cs were the combination of C-particles and Al-particles (Al-particles was dominated with the percentage of 68%) in early May; meanwhile the predominate carriers of 137Cs were carbonaceous particles with the average percentage of 88% in late May and September. Significantly, small particles (less than 2 μm) and medium particles (2-8 μm) of carbonaceous particles had a higher level in the aerosol samples of May and September. Specially, little particles (less than 1 μm), bacteria (1-1.8 μm), and spores (1.8-10 μm) had a linear relationship with the distribution of atmospheric 137Cs in the aerosol samples of September. In addition, the temperature and the precipitation were the main impact factors on the distribution of 137Cs and its carriers.

# 1 Introduction

## 1.1 The Fukushima Daiichi Nuclear Power Plant accident

The Fukushima Daiichi Nuclear Power Plant (hereinafter referred to as FDNPP, 37°25′ N, 141°02′ E, which is located on the north-eastern Pacific Ocean coast of Honshu, about 200 km far from north-east of Tokyo) was one of the nuclear power plants of the Tokyo Electric Power Holding Company (TEPCO). At 14:46 on March 11, 2011, a large-scale earthquake (also named as the Great East Japan Earthquake) with a magnitude of 9.0 occurred in the Tohoku region (Herp, 2021). At the moment of the disaster, Units 1 to 3 were normally operating, and Units 4 to 6 were shut down in the scheduled maintenance plan (Tepco, 2011a). The shaking caused by the earthquake and the following disaster of flooding caused by the tsunami resulted in the loss of the electricity of FDNPP, which was needed to run and cool the reactors and spent-fuel-pools normally (Tepco, 2011b). Therefore, a large amount of hydrogen gases was generated by the reaction of uncontrollable residual heat with metal in the units. Although the venting and water-injecting operations were performed in Units 1-3, the hydrogen explosions were not avoided in Unit 1 and Unit 3. Thus, a great many of the radionuclides were released into the atmosphere and deposited on the land and into the pacific ocean (Iaea, 2015).

## 1.2 Deposition and distribution of 137Cs

There were various radionuclides released into the atmosphere and deposited in the terrestrial and oceanic environment, which could cause health and environmental pollution, such as contamination problems of soil surfaces, water, agricultural products, and animals by-products, etc. The main radioactive radionuclides were iodine 131 (131I), cesium 134 (134Cs), cesium 137 (137Cs), and xenon 133 (133Xe). As shown in Table 1, the estimated amount of radionuclides released from the FDNPP accident were published by JAEA on April 12, 2011, and NISA on May 12, 2011, respectively (Ohara et al., 2011). Especially, 137Cs has a longer half-life of approximately 30 years, which has attracted much more attention of researchers than other short half-life radionuclides, for example, 131I and 133Xe (which has a short half-life of 8 days and 5 days, respectively) (Christoudias and Lelieveld, 2013). Additionally, the physicochemical properties of 137Cs are similar to potassium. Therefore, the soluble 137Cs was readily absorbed by animals and plants. When 137Cs was entered into the body of animals, it was mainly retained in bone and muscle tissue (Sato et al., 2016). Thus, it is particularly significant to conduct the long-term monitoring and analysis of 137Cs for understanding the fate of 137Cs in the aerosol samples.

Table 1. Radionuclides released from FDNPP accident modified from report of Ohara et al., 2011.

|  |  |  |  |
| --- | --- | --- | --- |
| **Radionuclide** | **Half-life** | **Released amount (Bq)** | |
| Japan Atomic Energy Agency (JAEA) 2011/5/12 | Nuclear and Industrial Safety Agency (NISA) 2011/4/12 |
| 131I | 8.040 ± 0.001 days | 1.5 × 1017 | 1.6 × 1017 |
| 132I | 2.30 ± 0.03 hours | — | 4.7× 1014 |
| 133I | 20.8 ± 0.2 hours | — | 6.8 × 1014 |
| 134Cs | 2.062 ± 0.005 years | — | 1.8 × 1016 |
| 137Cs | 30.17 ± 0.05 years | 1.3 × 1016 | 1.5 × 1016 |
| 106Ru | 368.0 ± 1.6 days | — | 2.1 × 109 |
| 129mTe | 33.61 days | — | 3.3 × 1015 |
| 132Te | 78.2 ± 0.8 hours | — | 7.6 × 1014 |
| 144Ce | 284.4 ± 0.3 days | — | 1.1 × 1013 |
| 133Xe | 5.25 ± 0.02 days | — | 1.1 × 1019 |

## 1.3 Resuspension of radiocesium

In the report (Onda et al., 2020), there were about 2.7 PBq of 137Cs deposited on the ground, and 60%-67% of them were deposited in the forest. These deposited 137Cs and their carriers were readily the secondary contamination sources. Specifically, in the report (Hirose, 2013), the monthly deposition speed of 137Cs decreased with an apparent half-life of 11-14 days during the period of March-June 2011. Also, the second peak of monthly 137Cs deposition was observed in February-April 2012, which may be ascribed to the resuspension of 137Cs-bearing-particle (CsMPs) (Tang et al., 2022). In addition, most of FDNPP-deriving 137Cs deposited on the topsoil, which remained in the soil surface layer as a potential secondary source of atmospheric 137Cs (Hirose, 2020). The resuspension process of radiocesium could be defined as the redistribution of deposited radioactive cesium into the atmosphere by wind or anthropogenic processes (Igarashi et al., 2003; Igarashi, 2009; Kajino et al., 2016). For bare soil surfaces and forest ecosystems, the respective resuspension rates of 137Cs were estimated as 1 × 10−6 day−1 and 2 × 10−6 day−1, which reported by Kajino et al. (Kajino et al., 2016), and revealed a seasonal change that the high level of 137Cs could be observed in warm seasons and the low could be obtained in cold seasons. Also, the resuspension of 137Cs may be derived from the decontamination process in heavily contaminated areas (Steinhauser et al., 2015). However, based on above-mentioned studies, the resuspension of radiocesium and the carriers in the atmosphere has not been fully or accurately understood. Even nine years after FDNPP accident, it was still found a small fluctuation within 0.0002 Bqm-3 from January to April and a slightly higher level of atmospheric 137Cs from May to September in the aerosol samples obtained in Namie in a heavily contaminated area of Fukushima prefecture in 2019. This work, for the first time, quantitively demonstrated thatthe carriers of 137Cs in early May were the combination of C-particles and Al-particles (Al-particles was dominated with the percentage of 68%); meanwhile the predominate carriers of 137Cs in late May and September were carbonaceous particles with the average percentage of 88%. In addition, the effect of weather conditions (precipitation, air temperature, relative humidity, wind speed, and gust wind speed) on the concentration of 137Cs and their carriers was also discussed. Obviously, the temperature and the precipitation were the main impact factors on the distribution of 137Cs and its carriers.

# 2 Experimental sections

## 2.1 Sampling site

The sampling site was a school ground (37°33'44"N, 140°46'07"E, about 30 km far from FDNPP) located in Namie in a heavily contaminated area of Fukushima Prefecture, as shown in Figure 1. It was surrounded by forests, mainly dominated by deciduous forest. Due to the heavy contamination, residents have been evacuated from these contaminated areas. Therefore, the sampling site was free of residential activities, except for decontamination activities and regular research monitoring. As shown in Figure 1, this research site was located at the boundary of the most contaminated area, about 1 MBqm−2.

地图

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**FDNPP**

**Sampling site in Namie**

Figure 1. Distribution of deposited 137Cs obtained from the Ministry of Education, Culture, Sports, Science and Technology (MEXT) converted on March 11, 2013 (Mext, 2013), also the sampling site was marked in a red cross.

## 2.2 Sampling

Table 2. The information of samples in May and September 2019 giving the name and the sampling time.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **Sample name** |  | **Starting time** | |  | **Ending time** | |
|  | **Year/Month/Day** | **a.m./p.m.** |  | **Year/Month/Day** | **a.m./p.m.** |
| NHVA-20190501-D-Q |  | 2019/4/26 | 6:00 p.m. |  | 2019/4/27 | 6:00 a.m. |
| NHVA-20190501-L-Q |  | 2019/4/30 | 6:00 p.m. |  | 2019/5/1 | 6:00 a.m. |
| NHVA-20190511-G-Q |  | 2019/5/7 | 6:00 p.m. |  | 2019/5/8 | 6:00 a.m. |
| NHVA-20190511-J-Q |  | 2019/5/10 | 6:00 p.m. |  | 2019/5/11 | 6:00 a.m. |
| NHVA-20190523-A-Q |  | 2019/5/11 | 6:00 p.m. |  | 2019/5/12 | 6:00 a.m. |
| NHVA-20190523-J-Q |  | 2019/5/15 | 6:00 p.m. |  | 2019/5/16 | 6:00 a.m. |
| NHVA-20190923-H-Q |  | 2019/9/20 | 6:00 p.m. |  | 2019/9/21 | 6:00 a.m. |
| NHVA-20190923-J-Q |  | 2019/9/21 | 6:00 p.m. |  | 2019/9/22 | 6:00 a.m. |
| NHVA-20190923-L-Q |  | 2019/9/22 | 6:00 p.m. |  | 2019/9/23 | 6:00 a.m. |
| NHVA-20190929-B-Q |  | 2019/9/23 | 6:00 p.m. |  | 2019/9/24 | 6:00 a.m. |
| NHVA-20190929-H-Q |  | 2019/9/26 | 6:00 p.m. |  | 2019/9/27 | 6:00 a.m. |
| NHVA-20190501-G-Q |  | 2019/4/28 | 6:00 a.m. |  | 2019/4/28 | 6:00 p.m. |
| NHVA-20190511-L-Q |  | 2019/5/10 | 6:00 a.m. |  | 2019/5/10 | 6:00 p.m. |
| NHVA-20190523-K-Q |  | 2019/5/12 | 6:00 a.m. |  | 2019/5/12 | 6:00 p.m. |
| NHVA-20190523-I-Q |  | 2019/5/15 | 6:00 a.m. |  | 2019/5/15 | 6:00 p.m. |
| NHVA-20190923-G-Q |  | 2019/9/20 | 6:00 a.m. |  | 2019/9/20 | 6:00 p.m. |
| NHVA-20190923-K-Q |  | 2019/9/22 | 6:00 a.m. |  | 2019/9/22 | 6:00 p.m. |
| NHVA-20190929-K-Q |  | 2019/9/28 | 6:00 a.m. |  | 2019/9/28 | 6:00 p.m. |

High-volume aerosol samplers (HV-1000R, Sibata, Japan) equipped quartz fiber filter (2500QAT-UP, Pallflex, USA) were used to collect atmospheric aerosol samples. The sampling flow rate was set to 1000 Lmin-1, and the sampling period was a short-term of 12 hours, as given in supplyment. The daytime and nighttime samples were respectively collected in May and September 2019, with day-time sampling from 6:00 a.m. to 6:00 p.m. and night-time sampling from 6:00 p.m. to 6:00 a.m. of the next day, as detailed given in Table 2. The absent samples in late May and early September 2019 were ascribed to the sampling plan and summer vacation. The quartz filter samples collected by HV aerosol samplers in Namie were stored in the laboratory.

## 2.3 Meteorological monitoring

Regular meteorological monitoring of the sampling location was also carried out, which was located approximately 800 m far from HV aerosol samplers. The pressure (mbar), solar radiation (Wm−2), wind direction (ø), moisture content (m−3m−3), precipitation (mm), air temperature (°C), relative moisture (RH, %), wind speed (ms−1), and speed of gust wind (ms−1) were simultaneously measured per minute. The detailed instrumental settings are shown in Table 3.

Table 3. Several instrumental specifications of meteorological monitoring modified from Appendix of Ishizuka et al., 2017.

|  |  |  |  |
| --- | --- | --- | --- |
| **Monitoring Object** | **Instrument** | **Manufacture** | **Model** |
| **Precipitation** | Tipping bucket rain gauge | Takeda Keiki Kougyou | TKF-1 |
| **Humidity (RH)** | Capacitive chip | Vaisala Corp. | HMP155D |
| **Wind speed** | Three cups anemometer | R. M. Young | 3102 |
| **Gust wind speed** | Sonic anemometer | R. M. Young | 81000 |
| **Air temperature** | Pt resistance thermometer | Vaisala Corp. | HMP155D |
| **Moisture** | ADR sensor | Delta-T Devices Ltd. | Theta probe ML2x |

# 3 Radioactivity measurement and observations of samples

## 3.1 Radioactivity measurement

The radioactivity of 134Cs, 137Cs was measured at the peak gamma-ray at 605 keV and 662 keV, respectively, using coaxial Ge semiconductor equipped with computerized spectrum analyzer in the Meteorological Research Institute (MRI) and the University of Tsukuba. The atmospheric radioactive concentration of 137Cs was calculated by

(1)

where *C*filter (Bqm−3) was the atmospheric radioactive concentration of 137Cs, *Q*filter (Bq) was the radioactive intensity of 137Cs in quartz fiber measured by coaxial Ge semiconductor, and *V*filter (m3) was the volume of the sampling air.

## 3.2 Microscope observations

Two pieces of Φ33 mm were taken out from the HV aerosol filter sample (8 × 10 inches) for DAPI (4’,6-diamidino-2-phenylindole) staining: one piece (a) was obtained in the middle area and the other piece (b) was obtained in the edge area from the same aerosol filter sample. The rest of the aerosol filter sample was sealed and stored for other experiments. Then, one piece of Φ12 mm was taken out from each piece of Φ33 mm for further treatment. Finally, two pieces of Φ12 mm were obtained and the rest of each piece of Φ33 mm was sealed and stored for backup. For all samples, the two pieces of Φ12 mm filters were firstly fixed by formaldehyde solution, and then dried for 2 hours. Formaldehyde solution was used for preserving or fixing tissues or cells due to its functions of embalming, fixing cadavers, disinfection, and bleaching. DAPI for fluorescent staining, because DAPI could penetrate the cell membrane and strongly fix the DNA in the nucleus (Maki et al., 2013). After staining, the samples were rinsed with ultrapure water and dried, in the end, the samples were stored in dark light. For fluorescence observation, the DAPI-stained cells were able to be labeled with blue fluorescence, after excitation by UV light with the wavelength of 360-400 nm, using the fluorescent upright microscope (BS-2040TF, Bio Tools Inc., Gunma, Japan). The dark particles (all particles) could be observed with the reflected-light mode, and the colored particles (carbon-containing particles) could be observed in the fluorescence-mode because the DAPI-stained particles could be shown blue or yellow fluorescence illuminated by UV light under the fluorescent upright microscope. Five sites of each Φ12 mm sample were used for observations using fluorescent upright microscope. Each site was photographed with a CCD camera in reflected-light mode and in fluorescent-light mode, respectively. Finally, all images were saved for the analysis of the size and morphology of aerosol particles. A total of 10 images were taken for each Φ12 mm sample, and a total of 20 images were taken for each sample collected in our sampling site. The images (the number and Feret’s diameter of particles) were analyzed by a free professional software of Image J (Kita et al., 2020). The morphology, the elemental compositions and distribution of the aerosol particles was characterized by a SEM (SU3500, Hitachi High-Technologies Co., Tokyo, Japan) equipped with an EDS (X-max, Horiba Ltd., Kyodo, Japan) in MRI.

# 4 Results and discussion

## 4.1 Variations of 137Cs in aerosol filters sampled in 2019

### 4.1.1 Annual variations of atmospheric 137Cs

The annual variations of 137Cs in the HV aerosol filters sampled in 2019 were shown in Figure 2 (the absent samples were ascribed to the sampling plan and vacations). It could be found a small fluctuation of atmospheric 137Cs within 0.0002 Bqm-3 from January to April. It was obvious that a slightly higher level of 137Cs could be observed from May to September. Moreover, it can be obtained the seasonal variation of the atmospheric 137Cs that the level of 137Cs was higher in the warm season (from May to September) and lower in the cold season (from January to April). Significantly, there were two peaks that appeared in May (about 0.00072 Bqm−3) and September (about 0.00052 Bqm−3). These two peaks may be caused by the resuspension of aerosol particles carrying 137Cs, which will be discussed later in detail. Thus, in the following section, variations of atmospheric 137Cs for HV filter samples collected in May and September were mainly discussed.



Figure 2. Annual variations of 137Cs in aerosol filters sampled in 2019.



Figure 3. Diurnal variations of 137Cs in the aerosol filters sampled in the day-period/night-period in 2019. The sampling date defined as mm/dd. Blackline represents variations of 137Cs concentration in the day-period. Redline was the variations of the samples obtained in the night-period.

### 4.1.2 Diurnal variation of 137Cs

The comparison of daytime and nighttime variation of atmospheric 137Cs was shown in Figure 3. There were 22 groups of 137Cs in HV filter samples collected in the day-period and night-period, respectively. Clearly, it can be found that the concentration of 137Cs in the daytime samples were slightly higher than that sampled in the nighttime. Specifically, among the 22 groups of 137Cs, 15 groups had a positive differences in atmospheric 137Cs between daytime and nighttime samples. Also, it can be noticed that the concentrations of 137Cs sampled in the day-period on the 12 May and on 28 September were about 3 times and 2 times higher than that of those samples collected in the night-period, respectively. Especially, the diurnal variations of 137Cs were similar to the seasonal variation of 137Cs as mentioned above in the annual variations of 137Cs. Specifically, the maximum of 137Cs concentration was 0.00072 Bqm−3 sampled in a day-period on 12 May and the minimum was 0.00002 Bqm−3 sampled in the night-period on 26 April and 27 April. In Autumn (late September), the maximum of 137Cs concentration was 0.00052 Bqm−3 sampled a day-period on 28 September, and the minimum was 0.00011 Bqm-3 sampled a day-period on 22 September.

## 4.2 Possible carriers of 137Cs in May and September

### 4.2.1 Possible carriers of 137Cs for the HV samples collected in May 2019

SEM observations of the HV filter sample (#NHVA2019-0511-J-Q) were shown in Figure 4. Comparative analysis of the observations was shown in Figure 4a in backscattered electrons mode, BSE, and Figure 4b in low-vacuum mode. Clearly, there were some grey-white particles with diameters of 20-35 μm, which could be easily identified as pollen particles. Meanwhile, there were also some small, clear, white, elliptic-shaped particles and they were mapped by EDS, as shown in Figure 4c, 4b, and 4d. Significantly, the several large carbon-containing particles could be pollens as shown in Figure 4c, and some small particles may be organic matters (such as fungal cells and/or debris, sporangia or ascospores or other microorganisms). In Figure 4d and 4e, it can be found that there were a lot of small aluminum-containing and iron-containing particles, which may be mineral particles or soil dusts. Overall, in Spring, more iron/aluminum-containing mineral particles of 2-5 μm and some scattered pollens and/or organic particles can be observed, indicating that main possible carriers of 137Cs in HV filters collected were the mineral particles.

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中度可信度描述已自动生成树林里

描述已自动生成

**a**

**b**

A picture containing light, dark, comet, night sky

Description automatically generatedA picture containing purple, dark, night sky

Description automatically generatedA picture containing green, night sky

Description automatically generated

**e**

**d**

**c**

Figure 4. SEM images and elemental distribution maps of the HV filter named #NHVA2019-0511-J-Q: a) SEM image obtained in backscattered electrons mode; b) SEM image got in low-vacuum mode; c) Elemental distribution of carbon; d) Elemental distribution of aluminum; e) Elemental distribution map of iron.

In the typical optical microscopy photograph (Figure 5a) and fluorescent micrograph (Figure 5c) of 4,6-diamidino-2-phenylindole (DAPI) staining particles in the HV filter sample (#NHVA2019-0523-J-Q), it can be found a lot of bioaerosol particles. After treatment by Image J software, equivalent projected images were used for counting and classifying particles in the reflected-light mode (Figure 5b) and fluorescent-light mode (Figure 5d), respectively. It is easy to distinguish some pollen particles with a size larger than 20 μm (Figure 5a). Based on the fluorescent color and morphology of the lighted particles, the fluorescent aerosol particles could be classified into different bioparticles. Specifically, in Figure 5c, it could be found that the most abundant fluorescent aerosol particles were 1) big elliptic blue particles (diameter >20 μm, indicating pollens or aggregated particles), 2) spindly yellow and blue particles (10 μm < diameter <20 μm, microbial particles of sporangia or ascospores), 3) elliptic yellow and blue particles (diameter <10 μm, identified as bacteria or basidiospore), and 4) white particles indicated other organics.

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Figure 5. Microscope images (BS-2040TF) and processed equivalent projected images by Image J in the same site of a HV filter sample (#NHVA2019-0523-J-Q) collected in May 2019: (a) microscope image in reflected-light mode; (b) equivalent projected area image of Figure 5a; (c) microscope image in fluorescent-light mode; (d) equivalent projected area image of Figure 5c.

The classifications of fluorescence highlighted particles were consistent with that of the reference (Igarashi et al., 2019b). Additionally, only particles with sizes larger than 0.65 μm were counted in this study. Particularly, it is observed that numerous particles with multiple septa, which were most possible the fungal spores of the phylum Ascomycota. More bioaerosol particles are observed in September than in May (based on the comparation in Figure 5c and 8c). It was possibly due to a seasonal change in the bioaerosol source or rainy weather on the sampling days in September, which has also been discussed in the report of Kita et al. (Kita et al., 2020). As a consequent, after analyzing all particles, large particles (such as ascospores, pollens, fragments, aggregated particles), small particles (such as bacteria), and medium particles (such as basidiospores) could be observed and classified. Therefore, the abovementioned six types of particles were discussed in the following section.



Figure 6. Variation of 137Cs, C% and Al%.

Based on the SEM observations (Figure 4), it could be obtained that many small mineral particles were also found in HV filter samples collected in May. Meanwhile, the optical microscope images (Figure 5) also exhibited many dispersed organic large particles, such as pollen particles. Thus, it was assumed that the carriers of 137Cs in the filter samples collected in May can be alternated during this sampling period, because May was just located in the alternate period of spring and summer. In addition, four HV filter samples collected in May were analyzed by SEM/EDS for estimating the elemental mass percentage. The variation of 137Cs and elemental mass percentage of Al and C with sampling time were shown in Figure 6. The black points were the concentration of 137Cs, the red and the blue symbol represented the percentage of C and Al, respectively. It was obvious that the variation of 137Cs concentration was consistent with the trend of Al%. Specifically, the gradual increase of 137Cs concentration and Al% was observed from 26 April to 12 May, reaching the peak on 12 May. After 12 May, Al% showed a clear downward trend. In contrast, the variation of C% was always in a slowly increasing trend. There would be a period for 137Cs carrier’s transition from mineral particles (aluminum-containing particles) to carbonaceous particles. Therefore, it could be assumed that the mineral particles or soil dusts could be the main carriers of 137Cs in early Spring. On the other hand, the carbon-containing particles may be the dominated carriers of 137Cs in the late Spring, but more data and further discussion are still needed.



Figure 7. Comparative percentage variations of carbon- and aluminum- containing particles.

In Figure 7, the comparative variation of carbon-containing particles and aluminum-containing particles estimated from SEM-EDS observations was provided. It could be noticed that the percentage of carbon-containing particles gradually increased, on the contrary, the percentage of aluminum-containing particles gradually decreased from 10 May to 15 May, which is consistent with the assumption that the mineral particles or soil dusts could be the main carriers of 137Cs in early Spring and the carbon-containing particles could be the dominated carriers of 137Cs in the late Spring. Moreover, a close higher-level percentage of carbon-containing particles was found in the samples collected on 15 May and 23 September with the percentage of carbon-containing particles of 92%, and 82%, respectively, which indicated that the main carriers of 137Cs may be carbon-containing particles. This result was consistent with our previous master’s research (Kimura, 2019a) that the bio-particles gradually became the dominant carriers of 137Cs. These results were also consistent with the speculation that the mineral particles or soil dusts could be the main carriers of 137Cs in early Spring and the carbon-containing particles could be the gradually dominated carriers of 137Cs in the late Spring.

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Figure 8. Microscope images (BS-2040TF) and processed equivalent projected images by Image J in the same site of a HV filter sample (＃NHVA2019-0923-L-Q) collected in September 2019: (a) microscope image in reflected-light mode; (b) equivalent projected area image of Figure 8a; (c) microscope image in fluorescent-light mode; (d) equivalent projected area image of Figure 8c.

### 4.2.2 Possible carriers of 137Cs for the HV samples collected in September 2019

As shown in the typical optical microscopy photograph (Figure 8a) and fluorescent micrograph (Figure 8c) of DAPI staining particles in the HV filter sample (#NHVA2019-0923-L-Q), it was easily observed that there were more bio-particles with different morphology and size, which were consistent with report that bio-particles could be the main carriers of 137Cs in autumn (Kimura, 2019b; Igarashi et al., 2019b).

草の上に置かれている

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低い精度で自動的に生成された説明

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**a**

夜空に光っている

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自動的に生成された説明夜空の星

自動的に生成された説明

**e**

**d**

**c**

Figure 9. SEM images and elemental distribution maps: a) SEM image obtained by BSE model; b) SEM image got by UVD; c) Compositional map of C; d) Compositional map of Al; e) Compositional map of Fe. Sample name is #NHVA2019-0929-B-Q. The bar is 100 μm.

SEM observations of the HV filter sample (#NHVA2019-0929-B-Q) were shown in Figure 9. Compared with Figure 9c, 9d, and 9e, there were a higher level of carbonaceous particles and fewer aluminum-containing and iron-containing particles, which indicates carbon-containing particles could be the dominated carriers of 137Cs in September. It was also consistent with the results of microscope observations, as shown in Figure 8.

## 4.3 Bioaerosol particles and their size distributions

According to the observations of optical, fluorescent microscope, and SEM-EDS, aerosol particles in the aerosol filter samples collected in May and September 2019 were mainly analyzed. Diameter was described as the Feret diameter (along the selection boundary, the longest distance between any two points, also known as maximum caliper), which was obtained from the microscope images using Image J. Figure 10a and 10b were the size distributions of bioaerosol particles in the HV filter samples collected in May 2019. It could be easily noticed that the particles with the diameter (*d* < 1 μm) were predominant. The second peak represented the particles with a diameter less than 2 um. Similarly, the size distribution of bioaerosol particles in the HV filter samples collected in September 2019 was given in Figure 10c. A wider diameter range of the bioaerosol particles could be found in three peaks (*d* < 1 μm, *d* < 2 μm, and *d* < 8 μm). In addition, bioaerosol particles with a diameter less than 1 μm were also predominant. Quantitatively, from Figure 10a (early May) to Figure 10b (late May), the normalized number of particles had increased nearly 3 times (obtained by the value of first peak), which was consistent with the SEM observations as shown in Figure 7. Similarly, from Figure 10b (late May) to Figure 10c (September), the normalized number of particles had no apparent change. Overall, although the size distribution appeared bimodal or multi-peaked in Figure 10b and 10c, the bioaerosol particles with the diameter (< 2 μm) could be the predominant possible carriers of 137Cs.





Figure 10. Normalized particle number size distributions of bioaerosol particles in the HV filter samples collected in May (a: ＃NHVA2019-0501-G-Q, b: ＃NHVA2019-0523-J-Q) and September (c, ＃NHVA2019-0923-J-Q) in 2019.

## 4.4 Relation between aerosol particles and 137Cs

In Figure 11a, the unstained particles represented mainly mineral particles and some bioaerosol particles, which were difficult to stain in the DAPI staining experiment. In Figure 11b, the stained particles referred to mainly bioaerosol particles, which could be observed in blue/yellow/white lighted particles under fluorescent light. The blue points and the red points represented the correlation between the concentration of atmospheric 137Cs and the concentration of aerosol particles, which were respectively estimated from the HV aerosol filter samples collected in May 2019 (there were ten samples, n=10) and in September 2019 (there were eight samples, n=8), respectively. It was obviously found that the unstained particles of mainly mineral particles had a strong positive correlation with the concentration of atmospheric 137Cs both in May and September 2019, as shown in Figure 11a. As mentioned in Figure 7-10, it was consistent with the assumption that the combination of mineral particles and bio-particles could be the main possible carriers of 137Cs in May 2019.

In Figure 11b, the stained particles also had a positive correlation distribution with 137Cs in September 2019, reconfirming that the bioparticles could be predominant carriers of 137Cs (which was also consistent with the results in Figure 8 and 9). In contrast, in May, the 137Cs concentration did not have a good linear relation with concentration of colorless particles, which may be caused by the fact that the combination of mineral particles and bioparticles could be the main carriers of 137Cs in May 2019. Namely, the dominant carriers of 137Cs could be mineral particles in early May and bio-particles in late May (as shown in Figure 5 and 7). It seems to be consistent with other studies (Kinase et al., 2018; Igarashi et al., 2019b) that there were different resuspension mechanisms in May and September. Namely, it was generally believed that there was a much lower concentration of 137Cs, and the main carriers could be mineral particles in the Spring. In Summer and Autumn, there were relatively higher concentration of 137Cs, and the bioaerosol particles could be predominant in the aerosol particles, which also implied that the bioaerosols were more possible to be the carriers of 137Cs in September (Kita et al., 2020).





Figure 11. The concentration variations of atmospheric 137Cs with the concentration of aerosol particles, which were respectively estimated from the HV aerosol filter samples collected in May 2019 (there were ten samples, n=10, highlighted in blue points) and in September 2019 (there were eight samples, n=8, highlighted in red points).

Table 4. Classification of bioaerosol particles based on different sizes. AR (Aspect Ratio) represented the ratio of the major to the minor. The major and the minor were the primary and secondary axises of the best fitting ellipse. Circularity was defined as 4π×(area) / (squared circumference). When the value of circularity was 1, representing the particle that could be regarded as a perfect circle. If the value approached 0, it indicated that the shape of the particle was elongated. When the AR was greater than 4 and the circularity was less than 0.45, the particle was defined as a fragment.

图片包含 形状

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**30 μm**

**30 μm**

**20 μm**

**10 μm**

**2 μm**

**50 μm**

**20 μm**

Igarashi et al. reported a strong relationship between carbon-bearing particles and 137Cs concentration (Igarashi et al., 2019b). Also, combining the abovementioned discussion, the main carriers of 137Cs were mineral particles in early May, and the predominant carriers of 137Cs were bioaerosol particles in late May and in September 2019. Therefore, it was necessary to quantify the relation between bioaerosol concentration and 137Cs concentration to estimate the predominant contribution of the specific type of bioaerosol particles to the atmospheric 137Cs. According to the classification, as shown in Table 4, there were several common types of bioaerosol particles (Stanley and Linskens, 2012; Kelly et al., 2002; Yamamoto et al., 2012; Hoorman, 2011). Then, a multiple linear regression was used to estimate the kinds of bio-particles variated with the dominant carriers of 137Cs, as follows:

(2)

Where*I* (Bq) was the total radioactivity of 137Cs in the HV aerosol filter samples, iwas a serial number, i = 1, 2, 3, 4, 5, 6 (little particle, bacteria, spore, ascospore, pollen, fragment) and so on, *ai* was the coefficient (Bq μm-2), *A*i is the area of the HV filter samples (183.2 mm × 234 mm), and *b*0 was residual radioactivity of 137Cs of the HV filter (Bq).

Table 5. Each coefficient in equation (2). *ai* was the coefficient of each bioaerosol, referring to the 137Cs radioactivity accumulated in each type of bioaerosols within each HV aerosol filter sample. The statistical analysis was made by regression analysis for 10 samples. Asterisk (⁎) indicated a significance level.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| *ai* | Coefficient value  (Bq μm-2) |  | *b0* | Residual radioactivity  (Bq) |
| *a1* | 1.11× 10-8 |  | *b0* | 0.19 |
| *a2* | 3.08× 10-8 ⁎ |  |  |  |
| *a3* | 9.14× 10-12 |  |  |  |
| *a4* | 0.00 |  |  |  |
| *a5* | 0.00 |  |  |  |
| *a6* | 0.00 |  |  |  |

In Figure 12, the relationship between each type of bioaerosols and 137Cs was obtained by the multiple linear regression analysis. The black symbol was the measured value of 137Cs obtained from the coaxial Ge semiconductor detector. The red points represented the predicted radioactivity value estimated from equation (2). The other symbols represented the contribution of different bioaerosol particles to the radioactivity of 137Cs.

The distribution of 137Cs for each bioaerosol in aerosol filter samples collected in September 2019 was given in Table 6. As a result of a multivariate analysis (performed by least-squares) under non-negative constraints, the contribution of each species of bioaerosols to the radioactivity of 137Cs and the residuals were calculated. Obviously, bacteria (blue squares) had the highest contribution to radioactivity of 137Cs (Figure 12), which showed the strongest correlation with the concentration of 137Cs, followed by the order of little particles (wine squares), spores (violet squares), and ascospores (magenta squares) and fragment (olive squares). It was strongly consistent with the results of the size distribution of particles (as shown in Figure 10c), in which particles with a diameter (< 2 μm) could be the predominant possible carriers of 137Cs. Therefore, combining the information on the size distribution of the particles (Figure 10), the dominant 137Cs-bearing particles could be bacteria (1-1.8 µm), followed with little particles (less than 1 µm) and medium particles of 1.8-10 µm, possibly deriving from spores.



Figure 12. Bioaerosol particles’ radioactivity contribution to 137Cs radioactivity based on the multiple linear regression equation (2). The black squares are the measured values (*I*) of 137Cs radioactivity in each HV aerosol filter sample. The red squares represent the estimated values (*I*) from equation (2) in each sample. The rest symbols represent their contributed radioactivity of different bioaerosol particles to the radioactivity of 137Cs based on statistical predictions from the multiple linear regression equation, as detailed information is summarized in Table 6.

Table 6. Contribution values of each species of bioaerosols to 137Cs radioactivity. There were ten data (for eight in September, two on 15 May, consideration of bioaerosols could be the main carriers). The observed values (Bq) of 137Cs radioactivity are obtained from the HV aerosol filter samples (183.2 mm × 234 mm). The estimated values (Bq) are based on a prediction of equation (2) in each sample. The predicted radioactivity (Bq) in little particles, bacteria, spores, ascospores, and fragments (without considering pollens because of seasonality) was calculated from the statistical analysis in a multiple linear regression equation (2). The residual values were involved in the difference between the measured value and the estimated value.

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Data number | Measured value  (Bq) | Estimated value  (Bq) | Little particle  (Bq) | Bacteria  (Bq) | Spore  (Bq) | Ascospore  (Bq) | Fragment  (Bq) | Residual value  (Bq) |
| 1 | 0.2538 | 0.1516 | 0.0119 | 0.1395 | 0.0002 | 0.0000 | 0.0000 | 0.1023 |
| 2 | 0.1983 | 0.2812 | 0.0275 | 0.2536 | 0.0002 | 0.0000 | 0.0000 | -0.0829 |
| 3 | 0.1730 | 0.0751 | 0.0083 | 0.0667 | 0.0001 | 0.0000 | 0.0000 | 0.0980 |
| 4 | 0.1490 | 0.1355 | 0.0306 | 0.1048 | 0.0002 | 0.0000 | 0.0000 | 0.0134 |
| 5 | 0.1296 | 0.1442 | 0.0118 | 0.1321 | 0.0003 | 0.0000 | 0.0000 | -0.0146 |
| 6 | 0.0671 | 0.0786 | 0.0139 | 0.0645 | 0.0001 | 0.0000 | 0.0000 | -0.0114 |
| 7 | 0.1045 | 0.1573 | 0.0264 | 0.1305 | 0.0004 | 0.0000 | 0.0000 | -0.0528 |
| 8 | 0.1110 | 0.0980 | 0.0143 | 0.0835 | 0.0002 | 0.0000 | 0.0000 | 0.0130 |
| 9 | 0.1806 | 0.0978 | 0.0161 | 0.0817 | 0.0001 | 0.0000 | 0.0000 | 0.0828 |
| 10 | 0.3158 | 0.3139 | 0.0266 | 0.2870 | 0.0004 | 0.0000 | 0.0000 | 0.0019 |

In addition, to verify the accuracy of equation (2), a correlation analysis between the observed and estimated radioactivity values was shown in Figure 13. The X-axis was the observed value of the atmospheric radioactivity of 137Cs, and Y-axis was the estimated value of the atmospheric radioactivity of 137Cs. It was obvious that they were in a good positive linear relationship, indicating the feasibility and reasonability of equation (2).



Figure 13. Correlation between observed values and estimated values for 137Cs radioactivity.

## 4.5 Effect estimation of weather conditions on 137Cs and its carriers

Several representative items of weather conditions, such as precipitation, air temperature, relative humidity, wind speed, and gust wind speed, were selected and monitored to analyze the influence of these weather factors on the concentration of 137Cs and its carriers. Precipitation was obtained from a 12-hour accumulated value, and the other values were the average value, during the sampling period. Table 7 shows the effect of atmospheric 137Cs concentration and concentration of particles in HV aerosol samples by precipitation, air temperature, RH, wind speed, and gust in May 2019, which was estimated by Pearson correlation and P-value significance tests (ten samples were used for analysis, n=10). It could be found that the precipitation had a negative effect on both the concentration of 137Cs and its carriers. The temperature had a positive effect onthe concentration of 137Cs and its carriers, which was consistent with the abovementioned speculation that the concentration of 137Cs was higher in the warm season and lower level in the cold season. Other weather conditions of RH, wind speed, and gust had no significant correlations in the current research.

Table 7. The effect of atmospheric 137Cs and concentration of particles in HV filter samples by precipitation, air temperature, RH, wind speed, and gust in May 2019, was estimated by Pearson correlation and P-value significance tests (ten samples were used for analysis, n=10).

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The effect of atmospheric 137Cs and concentration of particles in HV filter samples by precipitation, air temperature, RH, wind speed, and gust in September 2019 was shown in Table 8, also estimating by Pearson correlation and P-value significance tests (eight samples were used for analysis, n=8). Obviously, the temperature had a positive effect on both the concentration of 137Cs and its carriers, which was consistent with the results in May and the speculation. In contrast, the precipitation had a negative impact on the concentration of 137Cs and a positive effect on the concentration of the particles, which was inconsistent with that in May.

In the report (Kita et al., 2020), the precipitation was conducive to an increase of the atmospheric 137Cs compared with the non-rainfall sampling period. In September, the precipitation positively affected the concentration of the bioparticles. Because the rainfall was possible and reasonable to promote spores’ multiplication. Moreover, there was no doubt that higher temperature could accelerate microbial colonization, which was able to explain the positive impact on the concentration of 137Cs and its carriers by temperature both in May and September.

Table 8. The effect of atmospheric 137Cs concentration and concentration of particles in HV filter samples by precipitation, air temperature, RH, wind speed, and gust in September 2019, was estimated by Pearson correlation and P-value significance tests (ten samples were used for analysis, n=8).

形状

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# 5 Conclusions

From the analysis of annual atmospheric concentration variation of 137Cs (based on the HV air filter samples collected in 2019), the atmospheric concentration of 137Cs had a small fluctuation within 0.0002 Bqm−3 from January to April and a slightly higher activity level of 137Cs in the air could be observed from May to September. Therefore, it could be obtained that the seasonal variation of 137Cs was higher in the warm season (from May to September) and lower in the cold season (from January to April). Significantly, there were two peaks that appeared in May (about 0.00072 Bqm−3) and September (about 0.00052 Bqm−3). Thus, a lot of attention was given to understanding the difference of concentration variations of 137Cs and its carriers in HV filter samples collected in May and September. By SEM/EDS and optical microscope observations, it could be obtained that the small mineral particles or soil dusts could be the main carriers of 137Cs in early Spring and the carbon-containing particles (some microorganisms, spores, and some other bio-particles) could be the dominated carriers of 137Cs in the late Spring. The bio-particles (bacteria and other small bio-aerosols particles) could be dominant carriers of 137Cs in September.

According to an effect estimation of atmospheric 137Cs concentration and concentration of particles in HV filter samples by measuring precipitation, air temperature, RH, wind speed, and gust in May and September 2019, using Pearson correlation and P-value significance tests, it could be obtained that the temperature and the precipitation were the main impact factors of 137Cs and its carriers. Specifically, the temperature had a positive effect onthe concentration of 137Cs and its carriers both in May and September 2019, which was consistent with the abovementioned speculation that 137Cs concentration was higher in the warm season and lower level in the cold season (Kinase et al., 2018). Moreover, the rainfall had a negative effect on both the concentration of 137Cs and its carriers in May 2019. In contrast, the rainfall had a negative impact on the concentration of 137Cs and a positive effect on the concentration of the particles in 2019, which may be resulted from the fact that the effect of precipitation on the concentration of 137Cs and/or bioparticles may persist for some time. For instance, the rainfall in one day may affect changes of the concentration of 137Cs and/or bioparticles in the following several days. Moreover, there was no decreasing trend in the concentration of 137Cs in 2019, which means long-term monitoring was still necessary for further and deeply understanding the fate and variation of atmospheric 137Cs resuspended from 137Cs bearing microparticles (CsMPs) (Tang et al., 2022; Igarashi et al., 2019a; Higaki et al., 2020; Nihei et al., 2018) or soluble 137Cs (Otosaka et al., 2022) deriving from FDNPP.

# Competing interests

The contact author has declared that none of the authors has any competing interests.

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