Five models have been used to estimate the oceanic dispersion of $^{137}$Cs from the Fukushima Daiichi nuclear power plant during March and April 2011, following the accident on March 11, 2011. The total discharged activity of $^{137}$Cs is estimated to be 2 to 15 petabequerels. A weak southward direction, while mesoscale eddy-like structures and surface-current systems contributed to dispersion in areas beyond the continental shelf. Most of the discrepancies among the models in April are caused by differences in how the mesoscale current structures off the Ibaraki coast are represented.

**INTRODUCTION**

A devastating earthquake and huge tsunami struck the Tohoku area, Japan, on March 11, 2011, causing major damage to the cooling systems of reactors in the Fukushima Daiichi nuclear power plant, operated by Tokyo Electric Power Company (TEPCO). In order to cool the reactor cores and the spent fuel in storage pools, large amounts of seawater and freshwater were used. A significant part of this radioactivity-contaminated water was discharged into the Pacific Ocean close to the power plant. In addition, several hydrogen explosions between March 12 and 15 resulted in the release of significant radioactivity into the atmosphere, some of which was deposited onto the sea surface over a wide area of the Pacific Ocean. Careful monitoring combined with modeling of the dispersion of the radioactivity provide critical information (1) on the processes responsible for dispersion of the radionuclides, (2) for simulation and prediction of the spread of radioactivity in the seawater, and (3) for the evaluation of the impact on the health of marine ecosystems and humans.

Since the accident at the Fukushima plant, several groups have been conducting numerical dispersion simulations of radionuclides discharged into the ocean, each group having different objectives. Some of the results from the simulations have been used to determine locations of monitoring observations off the east coast of Japan. This article reviews the present status of such simulations, without considering atmospheric deposition, and describes common aspects and discrepancies among the simulated results. The article also points out potential problems and provides guidance for future studies. This is the first attempt to conduct an intercomparison of models for the oceanic dispersion of $^{137}$Cs from the Fukushima plant.

**DISPERSION SIMULATION MODELS**

Numerical simulations of the dispersion of radionuclides in the ocean basically consist of two parts: an ocean circulation model and a radionuclide dispersion model. The ocean circulation model provides evolving circulation patterns for the dispersion model, while the dispersion model calculates the movement and spread of radionuclides in the ocean. In this article, we compare results from five groups: the Central Research Institute of Electric Power Industry (CRIEPI), the Japan Atomic Energy Agency (JAEA), the Japan Coastal Ocean Predictability Experiment (JCOPE) group at the Japan Agency for Marine-Earth Science and Technology (JAMSTEC), the Simulation Réaliste de l’Océan Côtié (Sirocco) group from the Observatoire Midi-Pyrénées, Centre National de la Recherche Scientifique and Toulouse University, and the National Oceanographic and Atmospheric Administration (NOAA) group.

Each group utilized a different set of models (Table 1). All of the models have their finest regional domain resolution focused on the area close to the nuclear plant, with various grid spacings in both the horizontal and vertical directions. Lateral boundary conditions of the circulation models are typically obtained from larger domain ocean circulation models, with relatively coarse resolutions, into which observed data, such as temperature, salinity, and sea-surface height, are assimilated to provide realistic upper-ocean conditions. With these assimilation schemes, mesoscale eddies and meandering of ocean currents, which are crucial

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1. Research Institute for Global Change, Japan Agency for Marine-Earth Science and Technology, Yokohama, Kanagawa 236-0001, Japan
2. Environmental Science Research Laboratory, Central Research Institute of Electric Power Industry, 1646 Abiko-cho, Chiba-ken 270-1194, Japan
3. Japan Atomic Energy Agency, 2-4 Shirane Shirakata, Tokai-mura, Naka-gun Ibaraki 319-1195, Japan
4. CNRS, Toulouse University, Laboratoire d’Aérologie, 14 Avenue Edouard Belin F-31400 Toulouse, France
5. Earth Resources Technology, Inc., Laurel, Maryland 20707, USA
6. National Oceanic and Atmospheric Administration, Silver Spring, Maryland 20910, USA
7. National Centers for Environmental Prediction/National Weather Service/National Oceanic and Atmospheric Administration, Camp Springs, Maryland 20746, USA
8. I.M. Systems Group, Inc., Camp Springs, Maryland 20746, USA

Corresponding author: Yukio Masumoto (masumoto@jamstec.go.jp)
for the radionuclide dispersion in the open ocean, are adequately represented. Tidal currents are also included in several models in order to reproduce realistic spatial-temporal current variations near the coastal regions.

Although most of the groups are conducting dispersion calculations for several radionuclides, such as $^{131}$I, $^{134}$Cs, and $^{137}$Cs, we focus only on $^{137}$Cs in this article, since it has a significantly long half-life of ~30 years and is observed in a wide area of the northwestern Pacific Ocean. All five groups are now considering both direct discharge and atmospheric deposition in their calculations, but we discuss here only the results without atmospheric deposition.

**SOURCE ESTIMATION**

The temporal evolution and amount of radioactivity released to the ocean and atmosphere from the plant are key pieces of information for dispersion simulations. In general, the source conditions are not readily available for this kind of accident, leaving a large uncertainty in the simulated results. One of the main purposes of the oceanic dispersion simulations is, therefore, to estimate the source information, using inversion techniques, as accurately as possible. So far, several estimates of the amount of $^{137}$Cs discharged into the ocean have been reported, which are summarized in Table 2 and Figure 1. Note that the values come not only from peer-reviewed scientific papers but also from unreviewed articles.

TEPCO reported that the estimated total amount of $^{137}$Cs discharged directly into the ocean through a crack in the concrete wall near the reactor of Unit 2 during April 1–6 was 0.94 petabecquerels (PBq), which is equivalent to about 25 kilocuries (kCi) (TEPCO 2011a). In Figure 1, this amount is indicated by a gray bar, assuming that the discharge occurred constantly for five days from April 1. On a few other occasions, TEPCO reported discharges of contaminated water into the ocean, but those were two to five orders of magnitude smaller in terms of radioactivity than the amount released during April 1–6.

Kawamura et al. (2011) estimated the amount of $^{137}$Cs discharged into the ocean (red line in Fig. 1) using radioactivity data measured near the power plant by TEPCO (2011b) (black line in Fig. 1); they assumed that the contaminated water of the observed concentration occupied an area of 1.5 km$^2$ in front of the plant and was 1 m deep. After adjustment to the values reported by TEPCO for the period of April 1–6, a value of 4 PBq was obtained. The time series of the source information thus estimated (red line in Fig. 1) shows two peaks of release at the end of March and the beginning of April, with a magnitude of about 0.4 PBq day$^{-1}$. After April 7, the discharge diminished exponentially to about 0.001 PBq day$^{-1}$ at the end of April.

On the other hand, Tsumune et al. (2011) estimated a source function of $^{137}$Cs by multiplying by a factor to adjust their model results, with a unit release of the radionuclide, to the observed values, giving a total of 3.5 PBq of $^{137}$Cs discharged directly into the ocean (Fig. 1, blue line). They provided a simple scenario for the time evolution of the radionuclide discharge, in which they assumed that the time fluctuation of the observed radioactivity is associated with dispersion processes after entering the ocean. Indeed, Tsumune et al. (2011) are successful in reproducing a detailed time evolution similar to that observed at several locations along the coast south of the plant, including $^{137}$Cs maxima observed on March 30 and April 6 near the plant.

![Figure 1](image_url)
Several other estimates, including those by the Sirocco, NOAA, and JCOPE groups, were basically obtained from numerical models using the TEPCO data. The values obtained by Sirocco and NOAA, about 3 to 4 PBq of $^{137}$Cs discharged directly into the ocean, are similar to those derived by Kawamura et al. (2011) and Tsumune et al. (2011). The second report from the Institut de Radioprotection et de Sûreté Nucléaire (IRSN) on the impact of radioactivity released from the plant on the marine environment (IRSN 2011a) estimated the amount of radioactivity discharged until April 11 to be 2.3 PBq, which is somewhat smaller than other estimates. Another IRSN report shows a significantly large value of 27 PBq of $^{137}$Cs (IRSN 2011b), which seems to have been derived by simple interpolation of sparsely observed data and assuming constant radioactivity within a relatively thick surface mixed-layer in March. A relatively large estimate of 14.8 PBq from JCOPE is mainly due to relatively coarse horizontal resolution with a simple boundary condition, in which the simulated $^{137}$Cs concentration at the sea surface in front of the plant is forced to adjust toward the observed value.

Most of the above estimates rely on radioactivity measurements by TEPCO near the nuclear plant. Errors in the TEPCO data, if any, can propagate into these estimates directly. Another factor affecting the value of estimated source information is the vertical distribution of radionuclides, especially in the oceanic surface layer. The surface mixed-layer defined by temperature or water density can be relatively thick in March due to winter cooling at the sea surface and subsequent vertical convective motion. We do not know at this stage, however, whether or not the radionuclides are also evenly distributed vertically within the surface mixed-layer. In this regard, the values in Table 2 should be considered to have large uncertainties.

### DISPERSSION SIMULATIONS

With the above-mentioned source information for simulating the dispersion of radionuclides in the ocean, time series of the three-dimensional distribution of $^{137}$Cs were obtained from each model. In this section, we compare these results, focusing on the surface distribution of $^{137}$Cs in the coastal and continental shelf regions during the first two months after the accident. We present 10-day averaged surface horizontal-velocity fields and $^{137}$Cs distributions for two periods, from March 22 to 31 (Fig. 2) and from April 21 to 30 (Fig. 3). These periods correspond, respectively, to a time of southward dispersion along the Fukushima coast and to a time of gradual dispersion toward the margin of the continental shelf. Monitoring of the radionuclides was conducted by the Ministry of Education, Culture, Sports, Science and Technology (MEXT) and TEPCO during March and April, 2011 (MEXT 2011; TEPCO 2011b), and comparisons of the simulated results with the observations are also made for the two periods.

#### March 22–31

A lack of observational data prevents us from providing a detailed description of the $^{137}$Cs distribution at the end of March. The monitoring observations, however, indicate high concentrations of $^{137}$Cs along the coast, near the nuclear plant (Fig. 2c). The data along a line 30 km offshore from the coast also show $^{137}$Cs contamination, with a magnitude of about 10 to 15 Bq L$^{-1}$. These observed values are significantly higher than those observed in Japanese coastal waters before the accident, a typical value of which is about 0.003 Bq L$^{-1}$ (Kasamatsu and Inatomi 1998).

In general, the surface current fields in all the models show a strong eastward or northeastward flow—the Kuroshio current—along the coast of Japan south of Inubo Peninsula. The current separates from the coastal area off Inubo Peninsula. The models also show a broad southward flow, with a speed of 0.2 to 0.5 m s$^{-1}$, in the region east of 141.5° E and north of the Kuroshio current (Fig. 2a-i). All models demonstrate a weak southward flow, with a speed of 0.1 m s$^{-1}$ or less, along the coast in front of the Fukushima plant. This southward current along the coast is responsible for the southward distribution of $^{137}$Cs at the end of March. The southward flow along the coast can be traced back up to 38° N in all the models, while the speed of the flow varies among the models and, in the JCOPE model, is partially associated with a cyclonic circulation off the coast of Fukushima. The local flow pattern in this region is susceptible to wind forcing, which shows higher temporal variability associated with synoptic weather disturbances.

All the models fail to simulate the relatively high concentration of $^{137}$Cs along a line 30 km offshore. Since the results shown here are the dispersions of $^{137}$Cs released directly from the plant, it is reasonable to expect that this offshore contamination at the end of March was due to deposition from the atmosphere. This is consistent with the conclusion of Tsumune et al. (2011), who showed the importance of deposition from the atmosphere by checking the $^{131}$I/$^{137}$Cs activity ratio.

Another important surface current pattern is eddy-like structures off the coast of Ibaraki in the region between 36.7° N and the Kuroshio current; in this region, the differences among the models are rather large. A clear example is an anticyclonic circulation centered at 36.4° N, 141° E in the CRIEPI result. A similar eddy structure can also be seen in other model results, but it is relatively weak and shifted slightly to the east in the JCOPE model, and it appears as a part of a strong dipole eddy structure in the

### Table 2

<table>
<thead>
<tr>
<th>Institution</th>
<th>Period</th>
<th>$^{137}$Cs*</th>
<th>Method</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>TEPCO</td>
<td>April 1–April 6</td>
<td>0.94</td>
<td>Based on observed data</td>
<td>Media release</td>
</tr>
<tr>
<td>IRSN</td>
<td>Up to April 11</td>
<td>2.3</td>
<td>Based on observed data</td>
<td>IRSN (2011a)</td>
</tr>
<tr>
<td>IRSN</td>
<td>March 25–July 18</td>
<td>27</td>
<td>Simulated results with observed boundary conditions</td>
<td>IRSN (2011b)</td>
</tr>
<tr>
<td>JAEA</td>
<td>March 21–April 30</td>
<td>3.58</td>
<td>Based on observed data</td>
<td>Kawamura et al. (2011)</td>
</tr>
<tr>
<td>CRIEPI</td>
<td>March 26–May 31</td>
<td>3.54</td>
<td>Unit release experiment with adjustment to observed values</td>
<td>Tsumune et al. (2001)</td>
</tr>
<tr>
<td>JCOPE</td>
<td>March 21–May 6</td>
<td>14.8</td>
<td>Based on observed data</td>
<td></td>
</tr>
<tr>
<td>NOAA</td>
<td>March 10–June 27</td>
<td>3.6</td>
<td>Based on observed data (data from Kawamura et al. 2011)</td>
<td></td>
</tr>
<tr>
<td>Sirocco</td>
<td>March 20–June 30</td>
<td>4.2</td>
<td>Inverse model based on observed data</td>
<td></td>
</tr>
</tbody>
</table>

*In petabecquerels (1 PBq = 10$^{15}$ Bq)
JAEA model. The Sirocco and NOAA models show relatively broad southward flow off Ibaraki. Satellite images of the sea-surface temperature distribution indicate a weak, warm-core, eddy-like feature off the coast of Ibaraki at the end of March and in early April (not shown). The 137Cs concentration had not reached the region of these eddies by the end of March.

An ensemble mean of the five model results, giving equal weight in the averaging (Fig. 2F), captures most of the above-mentioned velocity and 137Cs distributions. It is rather difficult, however, to show that the ensemble mean field is better at representing the observed distribution, since the observational data are so sparse and limited to a region close to the plant.

April 21–30

The high 137Cs concentration had spread offshore by the end of April, while the radioactivity along the line 30 km offshore diminished slightly to a value of around 10 Bq L⁻¹, or less, except for two locations, where values of more than 20 Bq L⁻¹ were detected (Fig. 3C). New monitoring stations were installed in the region off Ibaraki, but the observed values were all under the detection level of about 10 Bq L⁻¹ for this time period.

All the models show southward or southeastward dispersion of the 137Cs in the latter part of April (Fig. 3D–G). The offshore dispersion seems to be associated with a nearshore, northeastward surface flow broadly distributed near the power plant in all models. The southward or southeastward movements of 137Cs in the offshore region south of 37° N, however, show large differences among the models. While the anticyclonic circulation brings the 137Cs distribution southeastward in the CRIEPI and JCOPET models, the JAEA, Sirocco, and NOAA results show southward dispersion along the coast associated with a weak southward flow near the coast. The NOAA results indicate strong eddy features beyond the coastal region, but the radionuclide distribution was not affected by the eddies at the end of April. The differences among the models suggest that the surface circulations in the region between 37° N and the Kuroshio current are susceptible to mesoscale eddy activity and to variability of the Kuroshio, and the modeled Kuroshio current is in turn strongly affected by data-assimilation processes in the larger domain models. In addition, the radionuclide distribution is affected by velocity fields not only during April 21–30 but also before that period. Again, the ensemble mean fields capture reasonably well the main features of the 137Cs distribution and the velocity characteristics off Fukushima and Ibaraki (Fig. 3F).

In addition to the mesoscale eddy activity, the magnitude of horizontal and vertical mixing processes in the ocean may strongly influence the 137Cs distribution. For example, the CRIEPI and NOAA results show relatively weak values of less than 10 Bq L⁻¹ in most of the region affected by radionuclide contamination as compared with other model results. One possible reason for this discrepancy could be differences in the magnitude of the vertical diffusivity; in general, the larger the vertical diffusivity, the weaker the surface concentration of the 137Cs. However, it is not straightforward to deduce this effect in a simple comparison among the models in Figures 2 and 3, since each model uses different schemes for advection and diffusion as well as different diffusion coefficients.
Another possible cause for the discrepancy is the degree of horizontal movement of the radionuclides within the 10-day-average window. When the radionuclides move around in a large area, the averaged concentration becomes smaller as compared with a case where radionuclides stay in the same location. This aspect strongly depends on the variability in the current system and should be evaluated in the future in order to clarify the detailed dispersion processes in this region.

At the end of April 2011, the relatively strong southward flow in the offshore region had strengthened as compared with the end of March. A major part of the $^{137}$Cs distribution, however, was confined within the region between the coast and the offshore southward flow. The $^{137}$Cs dispersed to the south or southeast was eventually captured in early May by the northern flank of the Kuroshio current and spread rather quickly to the east into the Pacific Ocean. This eastward movement of $^{137}$Cs can be seen in all the model outputs, and one example for the JCOPET model is shown in Figure 4.

**FUTURE ISSUES**

Comparison of the surface horizontal distributions of $^{137}$Cs among the dispersion models for the Fukushima accident demonstrates general agreement of the flow fields and associated $^{137}$Cs distributions at the end of March and April 2011. However, there are noticeable differences among the models as well, in particular for the region between 37°N and the Kuroshio current, where cyclonic or anticyclonic eddy-like circulations can be seen in some models. Accurate representation of such mesoscale structures and associated radionuclide dispersion near the coast is an important challenge for simulations on the regional scale.

The degree of vertical mixing in a model, as well as the vertical distribution of the source term, may also affect the surface distribution of the radionuclides. During March and April, several low-pressure systems passed through the Fukushima region, and these could have produced relatively large vertical mixing in this region. We need to
complete systematic sensitivity analyses and more detailed model intercomparisons in order to resolve these issues.

In this article, we have focused only on the dispersion of the \(^{137}\text{Cs}\) discharged directly from the Fukushima plant into the ocean. However, \(^{137}\text{Cs}\) was also deposited as atmospheric fallout on the sea surface over a wide region. Investigations of the relative importance of these two sources and the distribution of total \(^{137}\text{Cs}\) concentration in the ocean are now being undertaken by several groups. Kawamura et al. (2011), for example, have discussed the noticeable impact of atmospheric deposition of \(^{137}\text{Cs}\) over a large area of the northwestern Pacific. NOAA’s simulations in a larger area of the northwestern Pacific, including a large area of the northwestern Pacific. The incorporation of better-performing models of clides, including adsorption onto particulates and absorption due to biological processes. At present, only a few groups are trying to integrate these processes into their models. Further incorporation of better-performing models of scavenging mechanisms into dispersion simulations in the ocean will be essential.

**CONCLUDING REMARKS**

We have reviewed the present status of oceanic dispersion simulations of radionuclides discharged from the Fukushima nuclear plant. There are large differences in the models and model settings among the research groups, leading to different results for ocean currents near Fukushima and, hence, for the distributions of radionuclides such as \(^{137}\text{Cs}\). At the moment, we cannot say that one model is better than another. Rather, we need to be able to explain the discrepancies and reduce the overall uncertainty of the dispersion simulations. The International Atomic Energy Agency is now coordinating a more detailed model intercomparison in order to facilitate this research. Insights into oceanic dispersion gained through such international efforts will facilitate implementation of more predictive models.

According to the TEPCO data obtained near the Fukushima plant, \(^{137}\text{Cs}\) radioactivity at the end of April 2012 was of the order of 1 Bq L\(^{-1}\). Results from recent observations and numerical models indicate that the distribution of \(^{137}\text{Cs}\) has expanded and shifted to the east into a large portion of the North Pacific Ocean, while the radioactivity in most of the area is of the order of, or below, 0.01 Bq L\(^{-1}\). In order to describe what happened and is happening in terms of the dispersion of radionuclides from the Fukushima plant, research-based, accurate observations and analyses in a wide area of the North Pacific Ocean for more than a few decades are needed; as well, detailed comparisons and a synthesis of the simulated results and observations are strongly required.

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