Contamination of Heavy Metals, Polychlorinated Dibenzo-\(p\)-Dioxins/Furans and Dioxin-Like Polychlorinated Biphenyls in Wharf Roach \textit{Ligia} spp. In Japanese Intertidal and Supratidal Zones

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Abstract: We investigated the pollution levels of 6 heavy metals and 29 dioxins (polychlorinated dibenzo-\(p\)-dioxins (PCDDs), polychlorinated dibenzofurans (PCDFs), and dioxin-like polychlorinated biphenyls (DL-PCBs)) in intertidal and supratidal zones by using wharf roaches (\textit{Ligia} spp.) collected from 12 sampling sites on the coast of Northeast Japan from November 2011 to June 2012. The total concentrations of heavy metals ranged from 177 to 377 \(\mu\)g/g-dry weight (dw), and the predominant metals were copper, zinc, and aluminum. The order of the detected level of heavy metals was zinc > aluminum > copper > cadmium > lead > chromium, and this trend was similar to a previous report. The total toxic equivalent (TEQ) value of the PCDD/Fs ranged from less than the limit of detection (<LOD) to 2.33 pg-TEQ/g-dw, and the predominant congener was octachlorodibenzodioxin (<LOD to 110 pg/g-dw). Compared with PCDD/Fs, DL-PCBs were detected at a predominantly higher level (total TEQ value: 0.64–27.79 pg-TEQ/g-dw). Detected levels of dioxins, especially DL-PCBs in the wharf roach, were like those in the bivalves. These results indicate that the wharf roach could reflect heavy metals and dioxin pollution in the supratidal zones and is a suitable environmental indicator for these environmental pollutants. This is the first study to investigate heavy metals, PCDD/Fs, and DL-PCBs pollution in coastal isopods in Japan.

Keywords: supratidal zone; \textit{Ligia} spp.; heavy metals; polychlorinated dibenzo-\(p\)-dioxin/furan; dioxin-like polychlorinated biphenyl

1. Introduction

The marine environment is highly exposed to natural compounds and anthropogenic products and chemicals that are released daily and/or accidentally [1–3]. Coastal areas, including intertidal/supratidal zones, are highly polluted even though they are important in marine ecosystems and provide reproduction zones for the larval growth of many species. This critical issue is prevalent in many coastal areas globally [1,4,5]. A wide variety of pollutants is present, such as pesticides [6], antibiotics [7], polycyclic aromatic hydrocarbons [8], heavy metals, and persistent organic pollutants. These environmental pollutants affect and disrupt marine ecosystems, and biomonitoring is necessary to understand these environmental issues.
Heavy metal pollution is widespread and is studied traditionally in the field of environmental pollution. Researchers have focused on several highly toxic elements, such as cadmium (Cd), chromium (Cr), mercury, selenium, lead (Pb), arsenic, fluorine, and boron. These toxic heavy metals can be released from natural sources [9–11] and can be discharged from anthropogenic sources, such as industrial activity [12,13]. Mining provides one of the sources of heavy metals of most concern [14,15]. In the 1910s–1970s, wastewater that contained Cd was released from a mining area in Toyama, Japan, into a surrounding river. The release resulted in a severe health disease, known as “Itai-itai disease”, and soil contamination [16,17]. Some heavy metals are essential to organisms, but excess amounts cause health issues [18,19]. Heavy metals tend to be detected in aquatic wildlife, fish [20], mussels [21,22], sea mammals [23], and isopods [24–27].

Dioxins, including 75 polychlorinated dibenzo-p-dioxins (PCDDs), 135 polychlorinated dibenzofurans (PCDFs), and 12 dioxin-like polychlorinated biphenyls (DL-PCBs), are one of the most known persistent organic pollutant categories that have been regulated by the Stockholm Convention since 2001. These compounds are of great concern as marine wildlife [28,29] and human [30,31] environmental pollutants. These pollutants are ubiquitous marine environmental pollutants globally and have even been detected in polar areas [32,33]. The PCDD/F and DL-PCB toxicity depends on each congener chemical structure. The World Health Organization established the “Toxic Equivalency Factor (TEF)”, which sets 2,3,7,8-TCDD as the highest toxic dioxin to 1 and converts the toxicity of each dioxin compound as a relative value to assess the total dioxin toxicity (toxic equivalent, TEQ) in 2005. These dioxins are generated as byproducts from the incomplete combustion of substances that contain carbon, oxygen, hydrogen, and chlorine. Due to their high hydrophobicity (water octanol partition coefficient, log Pow: 6.3–8.2), dioxins are highly accumulated in organisms. However, a few studies have reported environmental pollution with these pollutants, especially PCDD/Fs and DL-PCBs, using organisms’ habitats in the supratidal zone around Japan.

Environmental surveys of coastal areas tend to focus on intertidal zones and use mussels for biomonitoring [4,34,35]. Scientific effort has been expended over several decades to accumulate numerous baseline data. Environmental pollution with many compounds is thought to be distributed in the broad range of coastal areas, including intertidal and supratidal zones [8,36,37]. Environmental research conducted to date has focused mainly on the intertidal zone. One of the most difficult survey points in supratidal zones is the selection of suitable biomonitoring species. To address this scientific gap, our research group used the wharf roach Ligia spp. as an environmental biomonitoring species. Wharf roaches have a global distribution in temperate–tropical areas [38], occur in relatively small habitat areas [39], are omnivorous [40], and may be exposed to environmental pollutants via feeding in land above the tidal line. Previously, we used the wharf roach as an environmental indicator in supratidal zones and investigated the pollution of radio-cesium and radio-silver that was released from the Fukushima Daiichi Nuclear Power Plant [36], tributyltin and dibutyltin [37], polycyclic aromatic hydrocarbons (PAHs), and alkylated PAHs [8]. The wharf roach is a promising environmental indicator in supratidal zones and could fill the gap in coastal environmental surveys.

To investigate environmental pollution in coastal areas, especially in supratidal zones, we collected wharf roaches from 12 sampling coastal sites around Central to Northern Japan in 2011-2012 and analyzed heavy metals, PCDD/Fs, and DL-PCBs. We accumulated environmental data on these pollutants in the wharf roach and estimated the ecological risk from these pollutants with regards to their habitat.

2. Materials and Methods

2.1. Sampling

Details on wharf roach collection are provided in previous studies [8,36]. Briefly, 12 sampling sites on the central and northern coasts of Japan (Table 1) were selected and wharf roaches were collected manually and/or by using traps from October 2011 to June
2012. After collection, samples were frozen immediately and kept in storage at −40 °C. After being frozen, randomly selected samples were placed in a glass beaker and freeze-dried using a FreeZone 2.5 L benchtop freeze-dry system (Labconco Corp., MO, USA). After freeze-drying, samples were pooled and homogenized for further analysis.

Table 1. Sampling date and location (city, prefecture).

| Site Number | Sampling Date   | Sampling Site (City, Prefecture) |
|-------------|-----------------|----------------------------------|
| 1           | 21 January 2012 | Tateyama, Chiba (TAT)            |
| 2           | 20 October 2011 | Hachinohe, Aomori (HAC)          |
| 3           | 19 October 2011 | Usushiri, Hokkaido (USU)         |
| 4           | 23 October 2011 | Sado, Niigata (SAD)              |
| 5           | 14 October 2011 | Ishinomaki, Miyagi (ISH)         |
| 6           | 26 October 2011 | Aomori, Aomori (AOM)             |
| 7           | 19 October 2011 | Noroshi, Ishikawa (NOR)          |
| 8           | 19 October 2011 | Noto, Ishikawa (NOT)             |
| 9           | 27 October 2011 | Gamou, Miyagi (GAM)              |
| 10          | 27 October 2011 | Sasu, Miyagi (SAS)               |
| 11          | 27 October 2011 | Onagawa, Miyagi (ONA)            |
| 12          | 07 June 2012    | Matsukawaura, Fukushima (MAT)    |

2.2. Heavy Metals Analysis

An analytical method was followed, “General Rules for Atomic Absorption Spectrochemical Analysis” (JIS K 0121:1993), with slight modification. Briefly, approximately 0.1 g of dried specimen was weighed and transferred into a Teflon container. The specimen was spiked with 3 mL of sulfuric acid and 3 mL of pure water and was decomposed under pressure and at a high temperature. The decomposed specimen was transferred into a 100 mL Teflon beaker. The decomposition container was rinsed with pure water and the rinse water was added into a specimen beaker. This combined specimen was evaporated to dryness, and a constant volume was obtained by adding 1% sulfuric acid (v/v). After sample preparation, the heavy metal concentration was analyzed by inductively coupled plasma atomic emission spectroscopy (ICP-AES) (ICP-AES-9000; Shimadzu, Kyoto, Japan). We analyzed six heavy metals: Pb, Cd, Cr, copper (Cu), zinc (Zn), and aluminum (Al).

2.3. PCDD/Fs and DL-PCBs Analysis

The analytical method followed the “Analytical Guideline of Dioxins in Food” with slight modifications. Briefly, after analyzing the total length and body mass of the freeze-dried specimens, six specimens were mixed and powdered. 13C-labeled internal standards (17 congeners of 13C-2,3,7,8-PCDD/Fs, 50 pg each; 12 congeners of 13C-DL-PCB, 200 pg each) were spiked into the weighed specimen. Specimens were extracted using a high-speed solvent-extraction device (ASE-350; Thermo Fisher Scientific, Waltham, MA, USA) with extraction solvent acetone/hexane (1:2) at 150 °C. The extract was concentrated to near dryness by using a rotary evaporator (R-200; BUCHI, Sankt Gallen, Switzerland), and the concentrate was transferred into a weighed clean beaker. After additional drying at 105 °C for 1 h, the specimen was dissolved with ~10 mL of hexane, transferred into a 50 mL glass tube that contained 5 mL of sulfuric acid, and left overnight. The hexane layer was purified by using a silver nitrate silica gel column and an activated carbon column. A separated layer was loaded onto two connected glass tubes that were filled with silver nitrate silica gel (0.5 g) and activated carbon (0.5 g) and eluted with 15 mL of hexane. The silver nitrate silica gel column was removed and 10% of dichloromethane in hexane (v/v) was added to the activated carbon column to elute most PCBs, except the non-ortho PCBs (PCB fraction). Then, 25 mL of toluene was added to the activated carbon column to elute the PCDDs, PCDFs, and non-ortho PCBs (DXN fraction). The PCB fraction was evaporated to near dryness and reconstituted with 1 mL of hexane. The reconstituted specimen was purified with a sulfoxide column. After concentration by N2 gas and transferal into a 1.5 mL gas chromatograph (GC) vial, the specimen was spiked with a
were higher than the Cu concentration in all samples except for the SAS sample. Cr had a higher concentration (1.46–5.60 \(\mu g/g-dw\)) at a secondary lower level (0.22–1.46 \(\mu g/g-dw\)), and the HAC sample had the highest concentration (13.5 \(\mu g/g-dw\), AOM) and the highest maximum concentration (212 \(\mu g/g-dw\), Zn concentrations). Pb was detected at a secondary lower level (0.22–1.46 \(\mu g/g-dw\)), and the HAC sample had the highest concentration (13.5 \(\mu g/g-dw\), AOM) and the highest maximum concentration (212 \(\mu g/g-dw\), AOM). The NOT samples contained the highest Cd concentration (5.60 \(\mu g/g-dw\)). These three metals were detected in a relatively wide range of concentrations.

2.4. Statistical Analysis

The R software (ver. 4.0.2) was used to conduct the statistical analysis. Statistical significance was set at \(p \leq 0.05\). As a nonparametric statistical test, a Spearman’s rank-order correlation coefficient was used to examine significant correlations of target analyte concentrations between samples. A Wilcoxon rank sum exact test was used to discern differences among analyte concentrations from each sampling sites. A hierarchical cluster analysis was conducted to estimate grouping patterns, using square Euclidean distances.

3. Results and Discussion

3.1. Heavy Metal Concentrations

All analyzed samples contained high concentrations of Cu, Zn, and Al (Figure 1). The other three metals, Pb, Cd, and Cr, existed in lower concentrations. Cu, Zn, and Al had a similar concentration range (33.1–129, 42.6–149, and 13.5–212 \(\mu g/g-dry weight\)), respectively. Al had a broader concentration range with the lowest minimum concentration (13.5 \(\mu g/g-dw\), AOM) and the highest maximum concentration (212 \(\mu g/g-dw\), GAM). Compared with Al, Cu and Zn had a relatively constant concentration range. The level of these three heavy metals was similar to previously reported levels in freshwater trichoptera in Japan [41] and the wharf roach in China [27]. Their average Zn concentration level of these three heavy metals was similar to previously reported levels in freshwater systems.
range; however, their detected levels were lower than those of the other three metals and significant differences in concentrations among the sampling sites were not detected. *Ligia italicica* that was collected in Eastern Sicily, Italy, on September 2001 [26] contained a different pattern of heavy metals with a lower level of Cd (0.22–1.26 µg/g-dw) and a higher level of Pb and Cr (1.74–9.30 and 0.18–1.68 µg/g-dw, respectively), compared with this study. These differences probably reflect pollution differences among the sampling sites. The order of the detected level of the heavy metals was Zn > Al > Cu > Cd > Pb > Cr, and this trend was similar to a previous report that showed Cu > Cd > Pb in *Ligia cinerascens* [27] and a previous report on coastal water and sediment [20,42]. The correlations in the three metal couples (Cu–Zn, Cu–Al, Zn–Al) were not significant (Spearman’s rank correlation test, p = 0.92, 0.45, 0.69, respectively). Additionally, the total concentrations of heavy metals were not distributed widely among the 11 sampling sites (177–377 µg/g-dw). The sum of Cu, Zn, and Al was dominant in six metal total concentrations (Figure 2, 97.2%–99.7%).

![Figure 1. Detected concentration of individual heavy metals in wharf roach (µg/g-dw) collected from 11 sampling sites (n.a.: not available).](image1)

![Figure 2. Proportion of each heavy metal concentration in total concentration in 11 sampling sites (n.a.: not available).](image2)

### 3.2. PCDD/F and DL-PCB Concentrations

Most PCDDs were not detected except by GAM and MAT (Table 2). Only OCDD was detected in most sampling sites at higher concentrations (<LOD–110 pg/g-dw). In GAM and MAT’s specimens, most PCDDs were detected (6 of 7 and 5 of 7, respectively). Compared with the PCDDs, PCDFs were detected frequently, especially in specimens from HAC, GAM, and MAT (7 of 10, 7 of 10, and 6 of 10, respectively). Twelve DL-PCBs were detected frequently (83–100%) at high concentrations (<LOD–5900 pg/g-dw) among 11 sampling sites. Mono-*ortho* PCBs had a higher concentration compared with non-*ortho* PCBs. The trends in the detected level of DL-PCBs among the 11 sampling sites differed slightly from those of the PCDD/Fs. The HAC, GAM, and ONA specimens had relatively

![Figure 1. Detected concentration of individual heavy metals in wharf roach (µg/g-dw) collected from 11 sampling sites (n.a.: not available).](image1)

![Figure 2. Proportion of each heavy metal concentration in total concentration in 11 sampling sites (n.a.: not available).](image2)
higher concentrations of DL-PCB congeners (13–4500, 12–5900, and 5.6–1400 pg/g-dw, respectively).

Table 2. Detected concentrations of individual PCDD/Fs and DL-PCBs (pg/g-dw) and total TEQ value (pg-TEQ/g-dw) in wharf roach collected from 11 sampling sites (sample from SAS, Miyagi, was not available).

| Sampling Site | TAT | HAC | USU | SAD | ISH | AOM |
|---------------|-----|-----|-----|-----|-----|-----|
| **PCDDs**     |     |     |     |     |     |     |
| 2,3,7,8-TeCDD | <LOD| <LOD| <LOD| <LOD| <LOD| <LOD|
| 1,2,3,7,8-PeCDD| <LOD| <LOD| <LOD| <LOD| <LOD| <LOD|
| 1,2,3,4,7,8-HxCDD| <LOD| <LOD| <LOD| <LOD| <LOD| <LOD|
| 1,2,3,6,7,8-HxCDD| <LOD| <LOD| <LOD| <LOD| <LOD| <LOD|
| 1,2,3,7,8,9-HxCDD| <LOD| <LOD| <LOD| <LOD| <LOD| <LOD|
| 1,2,3,4,6,7,8-HpCDD| <LOD| <LOD| <LOD| 5.8| <LOD| <LOD|
| OCDD          | <LOD| 6.9 | 1.5 | 8.0 | 3.7 | 0.66|
| **PCDD/Fs**   |     |     |     |     |     |     |
| 2,3,7,8-TeCDF | <LOD| 1.5 | 0.41| <LOD| <LOD| 0.22|
| 1,2,3,7,8-PeCDF| <LOD| 1.2 | 0.39| <LOD| <LOD| 0.30|
| 2,3,4,7,8-PeCDF| <LOD| 1.1 | 0.50| <LOD| <LOD| 0.26|
| 1,2,3,4,7,8-HxCDF| <LOD| 0.79| <LOD| <LOD| <LOD| <LOD|
| 1,2,3,6,7,8-HxCDF| <LOD| 0.83| 0.23| <LOD| <LOD| <LOD|
| 2,3,4,6,7,8-HxCDF| <LOD| 0.33| 0.34| <LOD| <LOD| <LOD|
| 1,2,3,7,8,9-HxCDF| <LOD| <LOD| <LOD| <LOD| <LOD| <LOD|
| OCDF          | <LOD| <LOD| <LOD| <LOD| <LOD| <LOD|
| **Dioxins TEQ (pg-TEQ/g-dw)** | <LOD| 0.72| 0.26| 0.06| 0.00| 0.11|
| **non-ortho** |     |     |     |     |     |     |
| 3,4,4′,5-TeCB(#81)| <LOD| 31  | 16  | <LOD| 11  | 6.0 |
| 3,3′,4,4′-TeCB(#77)| 100 | 730 | 310 | 74  | 250 | 100 |
| 3,3′,4,4′,5-PeCB(#126)| 18  | 110 | 47  | 14  | 40  | 21  |
| 3,3′,4,4′,5,5′-HxCB(#169)| 3.3 | 13  | 7.7 | <LOD| 5.0 | 3.3 |
| **DL-PCBs**   |     |     |     |     |     |     |
| 2′,3,4,4′,5-pentaCB(#123)| 7.2 | 55  | 7.9 | 1.7 | 7.1 | 2.4 |
| 2′,3,4,4′,5-pentaCB(#118)| 310 | 4500| 740 | 170 | 570 | 260 |
| 2′,3,4,4′,5-pentaCB(#114)| 7.2 | 120 | 26  | 56  | 17  | 7.2 |
| **mono-ortho**|     |     |     |     |     |     |
| 2′,3,4,4′-pentaCB(#105)| 99  | 2200| 350 | 64  | 230 | 140 |
| 2′,3,4,4′,5,5′-hexaCB(#167)| 19  | 360 | 64  | 14  | 36  | 13  |
| 2′,3,4,4′,5,5′-hexaCB(#156)| 36  | 870 | 110 | 24  | 61  | 25  |
| 2′,3,3′,4,4′,5,5′-heptaCB(#157)| 12  | 200 | 25  | 5.8 | 15  | 6.1 |
| 2′,3,3′,4,4′,5,5′,5′-heptaCB(#189)| 4.1 | 33  | 15  | 2.0 | 3.8 | 2.6 |
| **DL-PCBs TEQ (pg-TEQ/g-dw)** | 1.92| 11.72| 5.01| 1.42| 4.21| 2.22|
| **TOTAL TEQ (pg-TEQ/g-dw)**    | 1.92| 12.44| 5.27| 1.48| 4.21| 2.33|
## Table 2. Cont.

| Sampling Site | NOR | NOT | GAM | SAS | ONA | MAT |
|---------------|-----|-----|-----|-----|-----|-----|
| PCDDs         |     |     |     |     |     |     |
| 2,3,7,8-TeCDD | <LOD| <LOD| <LOD| n.a.| <LOD| <LOD|
| 1,2,3,7,8-PeCDD | <LOD| <LOD| 0.81| n.a.| <LOD| <LOD|
| 1,2,3,4,7,8-HxCDD | <LOD| <LOD| 0.31| n.a.| <LOD| 1.3 |
| 1,2,3,6,7,8-HxCDD | <LOD| <LOD| 1.7 | n.a.| <LOD| 4.1 |
| 1,2,3,7,8,9-HxCDD | <LOD| <LOD| 0.75| n.a.| <LOD| 3.0 |
| 1,2,3,4,6,7,8-HpCDD | <LOD| <LOD| 5.7 | n.a.| <LOD| 17 |
| OCDD          | 3.4 | 110 | 21  | n.a.| 7.6 | 51  |
| PCDFs         |     |     |     |     |     |     |
| 2,3,7,8-TeCDF | <LOD| <LOD| 1.1 | n.a.| 1.0 | 2.0 |
| 1,2,3,7,8-PeCDF | <LOD| <LOD| 1.2 | 0.38| n.a.| 1.5 |
| 2,3,4,7,8-PeCDF | <LOD| <LOD| 0.87| 0.73| n.a.| 1.0 |
| 1,2,3,4,7,8-HxCDF | <LOD| <LOD| 1.1 | n.a.| <LOD| 3.1 |
| 1,2,3,6,7,8-HxCDF | <LOD| <LOD| 1.3 | n.a.| <LOD| 3.2 |
| 2,3,4,6,7,8-HxCDF | <LOD| <LOD| 1.1 | n.a.| <LOD| <LOD|
| 1,2,3,7,8,9-HxCDF | <LOD| <LOD| <LOD| n.a.| <LOD| <LOD|
| 1,2,3,4,6,7,8-HpCDF | <LOD| <LOD| 1.6 | n.a.| 1.3 | 3.4 |
| OCDF          | <LOD| <LOD| <LOD| n.a.| <LOD| <LOD|
| Dioxins TEQ (pg-TEQ/g-dw) | 0.00| 0.33| 1.86| n.a.| 0.46| 2.33|
| non-ortho     |     |     |     |     |     |     |
| 3,4,4′,5,5,6-TeCB(#81) | 5.3 | 4.9 | 48  | n.a.| 19  | 9.3 |
| 3,3′,4,4,5′-TeCB(#77) | 94  | 55  | 1100| n.a.| 340 | 100 |
| 3,3′,4,4,5,5,6-HxCB(#126) | 19 | 5.3 | 270 | n.a.| 51  | 40  |
| 3,3′,4,4,5′,5,6′-hexaCB(#169) | 1.3 | 3.1 | 12  | n.a.| 5.6 | 9.9 |
| DL-PCBs       |     |     |     |     |     |     |
| mono-ortho    |     |     |     |     |     |     |
| 2′,3′,4,4,5-pentaCB(#123) | 4.1 | 2.0 | 55  | n.a.| 15  | 12  |
| 2,3′,4,4′,5-pentaCB(#118) | 220 | 97  | 5900| n.a.| 1400| 660 |
| 2,3,4,4′,5-pentaCB(#114) | 6.7 | 3.5 | 130 | n.a.| 42  | 20  |
| 2,3′,3′,4,4′,5-pentaCB(#105) | 100 | 46  | 2100| n.a.| 390 | 270 |
| 2,3′,4,4′,5,5′-hexaCB(#167) | 11 | 9.2 | 510 | n.a.| 96  | 54  |
| 2,3,3′,4,4′,5,6-hexaCB(#156) | 18 | 13 | 1100| n.a.| 160 | 110 |
| 2,3′,3′,4,4′,5,6-hexaCB(#157) | 6.8 | 3.4 | 200 | n.a.| 30  | 24  |
| 2,3′,3′,4,4′,5,5′-heptaCB(#189) | <LOD| 3.0 | 34  | n.a.| 7.4 | 8.4 |
| DL-PCBs TEQ (pg-TEQ/g-dw) | 1.96| 0.64| 27.79| n.a.| 5.37| 4.34|
| TOTAL TEQ (pg-TEQ/g-dw) | 1.96| 0.97| 29.64| n.a.| 5.83| 6.68|

The PCDD/F congener composition differed among the 11 sampling sites. The highest-concentration congener was OCDD in all sampling sites, and this trend was more noticeable in the work by ISH, NOR, and NOT (100%, 100%, and 98%, respectively, Figure 3, Table S1: supplementary materials). In other sites, the OCDD burden was similar (44.6%–61.3%). However, other congeners were different. The SAD sample contained a high percentage of 1,2,3,4,6,7,8-HpCDD (42%), which was different from the other sites. In other sites, the low-chlorinated PCDF concentrations were higher than other congeners, but no other constant trend was observed. These results may reflect a site-specific contamination source.
Compared with the PCDD/Fs, the DL-PCBs had almost the same congener pattern among the 11 sampling sites. The most dominant congener was 2,3′,4,4′,5-pentaCB (#118: 39.5%–54.8%) after 2,3,3′,4,4′-pentaCB (#105: 15.0%–23.9%) (Figure 4, Table S2). This trend indicated that the contamination source of DL-PCBs was almost similar among the 11 sampling sites. This congener concentration composition was similar to that of mussels in previous studies [28,43,44]. However, 3,3′,4,4′-TeCB (#77: 7.6%–22.4%) existed in a relatively higher percentage compared with the previous research. In the study by Ohashi [28], a higher concentration of 3,3′,4,4′-TeCB (#77) was observed from sediment compared with the analyzed fishes. The wharf roach lived on a supratidal zone (terrestrial field); therefore, the exposure pathway may be different from the marine organisms. Exposure via soil has a higher importance with regards to 3,3′,4,4′-TeCB (#77) pollution in terrestrial organisms compared with marine organisms. This constant trend was not correlated with the log $P_{ow}$ value of each DL-PCBs among the 11 sampling sites (Spearman’s rank correlation, $p = 0.215–0.675$).

3.3. Hierarchical Cluster Analysis

The hierarchical cluster analysis showed different clustering depending on whether they were calculated using heavy metals or dioxins (Figure 5a,b). However, in both cases, HAC and GAM were separated from other sampling sites. In the case of heavy metal concentrations, HAC and GAM were separated from the other nine sites. However, linkage
heights between the two sites is high (100). The NOR sample was also separated from the other sites. The separation of these three sites from other sites was induced by high relative concentrations of Al. The other eight sites were more closely clustered. In particular, there are two small clusters: NOT, SAD, and ONA; and TAT, USU, and ISH. The clustering with dioxins (Figure 5b) also indicate that HAC and GAM were different from the other nine sites. These differences were likely the result of significantly higher concentrations of DL-PCBs (total DL-PCBs: 9222 and 11459 pg/g-dw, respectively). The ONA sample was secondly separated from other eight sites, and completely far from HAC and GAM. Compared with the results of heavy metals, the other eight sites were more clustered, forming two groups: NOT, SAD, TAT, AOM, and NOR; and USU, ISH, and MAT. The different grouping patterns that emerge when the analyses are performed with heavy metals and dioxins suggest differing contaminant sources, accumulation patterns, and mobilities in water. This is particularly the case with the HAC and GAM sites.

Figure 5. Hierarchical dendrogram for 11 sampling sites obtained using hierarchical cluster analysis with an average linkage method: (a) calculated using heavy metal concentrations and (b) calculated using PCDD/F and DL-PCB concentrations.

3.4. Ecotoxicological Risk for Wharf Roach

Normally, heavy metal and PCDD/F toxicities depend on the species. However, significant concentrations of heavy metals and PCDD/F were detected in wharf roach, which suggest a risk of PCDD/F toxicity to this isopod.

Pb, Cd, and Cr (particularly hexavalent chromium) have a high toxicity. However, due to arthropod exoskeleton molting, isopods can reduce the risk from excessive heavy metal contamination [45]. In this study, the concentrations of these three heavy metals were lower compared with the other heavy metals. In particular, Cr was detected at low concentrations (0.02–0.21 µg/g-dw), which is comparable to a previous study that showed 0.18–1.68 µg/g-dw in L. italica [26]. However, the detected concentrations of Cu, Cd, and Pb (33.1–129, 0.46–6.85, and 0.22–1.46 µg/g-dw, respectively) are slightly higher than reported in L. cinerascens (23.1–35.6, 0.319–0.893, and 0.0220–0.218 µg/g-wet, respectively) from China [27]. Prato et al. [46] reported that the 96 h–LC50 of Cu and Cd on Sphaeroma serratum (marine isopod) was 4.60 mg/L and 4.79 mg/L, and on Idotea baltica (marine isopod) it was 1.45 mg/L and 1.29 mg/L. Hong and Reish [47] reported that the 96 h–LC50 in sea water on Jaeropsis sp. and Limnoria tripunctata (both marine isopods) was 0.41 and 7.12 mg/L. These heavy metal concentrations in sea water were in almost the same range as those in wharf roach in this study. These heavy metals can accumulate in crustacean bodies from environmental media [48,49], having high bioconcentration factors (BCFs): Cd 17000, Pb 31000, Cu 120000, and Zn 9200 for freshwater isopod Asellus aquaticus (L.) [50];
and Cu 2060 and Cd 311 for marine amphipod Paramorea walkeri [51]. Even though the sensitivity to heavy metals differs among isopod species, the contamination level of heavy metals in this study poses an ecological risk to the wharf roach.

The total TEQs of the PCDD/Fs and DL-PCBs in the total dried body weight were discussed to assess the ecological risk (Table 2). MAT’s sample had the highest PCDD/Fs (2.33 pg-TEQ/g-dw), followed by GAM’s sample (1.86 pg-TEQ/g-dw). Compared with these two sampling sites, the other site had a lower PCDD/Fs level (<LOD–0.72 pg-TEQ/g-dw). DL-PCBs were detected at a higher level (0.64–27.79 pg-TEQ/g-dw) compared with the PCDD/Fs. The GAM sample contained the highest amount of DL-PCBs (27.79 pg-TEQ/g-dw), followed by HAC’s sample (11.72 pg-TEQ/g-dw). Other samples contained slightly lower levels (0.64–5.37 pg-TEQ/g-dw). The total TEQs of the PCDD/Fs and those of DL-PCBs in the total dried body weight were correlated significantly (Spearman’s rank correlation test, \( p = 0.04 \)), and this correlation trend was retained in the total TEQ value in the lipid (\( p = 0.03 \)). The TEQ of the DL-PCBs was dominant in the total TEQ (65%–100%, Figure 6). However, the contribution ratios of the PCDD/Fs in the total TEQ in NOT and MAT were higher than in the other nine sampling sites (34% and 35%, respectively). These detected levels of dioxins were higher than those detected previously in mussels [43,52] and small marine arthropods [53]. However, currently detected levels in the wharf roach were similar and/or lower than in marine crabs [54–56]. These differences may be caused by trophic level, species differences on feeding habitat, and metabolism. Due to a lack of toxicological information, it is difficult to estimate the ecotoxicological risk of dioxins on the wharf roach. Michielli et al. [56] detected a relatively higher level of dioxins on crab Carcinus mediterraneus and estimated a significant risk from dioxins. The ecological risk of dioxins on the wharf roach remains unclear, however, and should be studied further.

Figure 6. Proportion of PCDD/Fs and DL-PCBs on total TEQ at 11 sampling sites.

3.5. Case Study: Effect of the Great East Japan Earthquake

Enormous amounts of debris and anthropogenically derived chemicals were released from damaged areas to surrounding coastal areas by the tsunami that followed the Great East Japan Earthquake [57–59]. Related areas were exposed to heavy metals and dioxins. For this reason, we expected that the damaged areas site 2 (HAC), 5 (ISH), 9 (GAM), 10 (SAS), 11 (ONA), and 12 (MAT) had high concentrations of these pollutants. However, no significant difference in total concentration of heavy metals was observed between the damaged areas and others (177–299 µg/g-dw and 241–377 µg/g-dw) (Wilcoxon rank sum exact test: \( p = 0.08 \)). Heavy metal pollution in these areas occurred immediately after the earthquake and tsunami, but this was not a serious problem [60]. The total TEQs of the dioxins in the damaged areas were significantly higher than others (4.21–29.64 pg-TEQ/g-dw and 0.97–5.27 pg-TEQ/g-dw) (Wilcoxon rank sum exact test: \( p < 0.01 \)). Wharf roaches from GAM and HAC contained higher levels of dioxins (29.64 and 12.44 pg-g/g-dry, respectively). Ishiyama et al. [61] reported a significant elevation of PCB concentration in
blue mussel *Mytilus galloprovincialis* from the Miyagi prefecture after the tsunami (collection date: June 2011). Additionally, marine sediment around sampling sites contained a higher level of dioxins compared with the surrounding sea area [62]. Current results showed a relatively higher level of dioxin pollution in these areas; therefore, one pollution exposure factor was the earthquake and ensuing tsunami.

A highly polluted area was not included as a sampling site in this study. Therefore, in the future, the wharf roach should be used as an environmental indicator of multi-class pollution for polluted areas, such as developed ports, discharge ports of industrial waste, and sewage treatment plants.

4. Conclusions

This study presented new data on the level of heavy metals, PCDD/Fs, and DL-PCBs in the wharf roach. The predominant heavy metals were Cu, Zn, and Al. The detected concentrations of PCDDs were higher than those of the PCDFs. Among the PCDD/Fs, the predominant congener was OCDD. Compared with the PCDD/Fs, the DL-PCBs were detected with higher concentrations. The detected levels of dioxins, especially DL-PCBs, in the wharf roach were similar to those in bivalves. These results indicate that wharf roach could reflect heavy metal and dioxin pollution in the supratidal zones and is a suitable environmental indicator for these environmental pollutants. To our knowledge, this is the first study to investigate dioxin pollution in coastal isopods and to report pollution of heavy metals and dioxins in supratidal zones around Japan. However, the contamination source, pathway, and ecological risk of these pollutants on the wharf roach and related coastal habitats remains unclear. Thus, further study is required.

Supplementary Materials: The following are available online at https://www.mdpi.com/2076-3417/11/4/1856/s1, Table S1: Percentage of individual PCDD/Fs for sum of PCDD/Fs at 11 sampling sites, Table S2: Percentage of individual DL-PCBs in the sum of DL-PCBs at 11 sampling sites.

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