Original Article

Determination of Polychlorinated Naphthalenes in Landfill Leachates and its Removal in Wastewater Treatment Processes

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ABSTRACT

Di-through octachlorinated naphthalenes (DiCNs, TrCNs, TeCNs, PeCNs, HxCNs, HpCNs and OcCN) in landfill leachates can be analyzed with the same cleanup procedure as dioxins. In this study, a modified method was developed by confirming the elution order of each isomer of polychlorinated naphthalenes (PCNs) in a chromatogram of a gas chromatograph/high resolution mass spectrometer (GC/HRMS) using an Rh-12 ms capillary column, which is mainly employed for the analysis of dioxins. Using this method, PCNs in landfill leachate samples and treated leachate samples from each corresponding wastewater treatment process were quantified. PCN concentrations in landfill leachates ranged from 8.0–12,000 pg/L, and DiCNs-PeCNs were detected in almost all samples. Removal of PCNs by flocculation and active carbon treatment was greater than that achieved by activated sludge treatment in wastewater treatment plants. Furthermore, by changing the temperature of the H2SO4 coated silica gel column in the modified method, it became possible to measure monochlorinated naphthalenes (MoCNs), which could not be measured previously. MoCN concentrations detected were much higher compared to concentrations of DiCNs-OcCN in most of landfill leachates. Unlike PCNs, MoCNs are not POPs, but they are nevertheless important for understanding the properties of landfill leachates.

Keywords: polychlorinated naphthalenes, landfill leachate, gas chromatography, high resolution mass spectrometry

INTRODUCTION

Polychlorinated naphthalenes (PCNs) are chemically and thermally stable and show electrical insulation properties, with a structure similar to that of polychlorinated biphenyls (PCBs) [1–4]. PCNs are persistent, bioaccumulative, transported over long ranges, and widely present in the environment [5,6]. Numerous serious health issues, such as chloracne and liver disease mortality for industrial workers caused by exposures to PCNs, have been reported [3]. The structural similarity of PCNs to the highly toxic 2,3,7,8-tetrachlorodibenzo-p-dioxin (2378-TCDD) indicates an aryl hydrocarbon receptor-mediated mechanism of toxicity (dioxin-like toxicity). Among PCNs, 1,2,3,6,7,8-hexachlorinated naphthalene showed the highest toxicity, and its relative potency has been detected by a CALUX test as 0.0051 compared to 2378-TCDD [7–9]. This relative toxicity value corresponds to the third highest for the Co-PCB in the WHO’s 2005 reevaluation of toxic equivalency factors (TEFs) for dioxins and dioxin-like compounds [9]; therefore, the behaviors, toxicity, and removal of PCNs in aquatic environments should be investigated.

The manufacture of PCNs began in around 1910, and they were sold as waxes with typical product names such as Halowax (Koppers, USA), Nibren waxes (Bayer, Germany), and Seekay waxes (ICI, UK) [1]. The industrial applications of PCNs comprise mainly cable insulation, automotive capacitors (dielectric), engine oil additives, and as stopoff compounds for the electroplating industry. PCNs cause industrial hygiene addiction, and their production declined in the late 1970s [3,4]. However, PCNs are persistent and continue to be present in previously manufactured products.
Moreover, PCNs are derived from technical PCB byproducts (technical biphenyl contains naphthalene) and unintentionally from thermal processes in the presence of chlorine [1]. Since PCN-containing products are hardly available now, the main sources of PCNs in the environment that should be considered are leachate from landfill sites where PCN-containing products have been disposed of and exhaust gases from waste incinerators or other thermal processes [3–6].

Monochlorinated naphthalenes (MoCNs) were used as raw material for dyes and as a wood preservative in the 1940s and 1950s [4]. 1-MonoCN is designated as a class 2 chemical substance for their thorough Safety Data Sheet (SDS) system under the Act on Confirmation of Release Amounts of Specific Chemical Substances in the Environment and Promotion of Improvements to the Management in Japan [10]. Di-through octachlorinated naphthalenes (DiCNs, TrCNs, TeCNs, PeCNs, HxCNs, HpCNs, and OcCN) are listed in Annexes A and C of the Stockholm Convention on Persistent Organic Pollutants (POPs), due to their degrees of toxicity, stability, bioaccumulation, and their long-range transportation [5]. Parties that have signed the POPs convention must take measures to eliminate the production and use of the chemicals listed under Annex A and to reduce the unintentional releases of chemicals listed under Annex C with the goal of continuing minimization and, where feasible, ultimate elimination [5]. In Japan, TrCNs-OcCN were designated as a Class 1 Specified Chemical Substance under the Chemical Substances Control Law in 1979, and its manufacture and use have been prohibited. Because of the properties of MoCNs and PCNs, their usage is currently restricted; however, they could still be present in the environment.

In the 1970s, PCNs were measured at nd–5.5 µg/L in water near two capacitor manufacturing sites [11]. A Swedish study reported that PCN concentrations in PCB-polluted rivers and percolating water at city dump sites were 0.89 and 2.6 ng/L, respectively [12]. In a recent monitoring survey in Japan, the MoCNs and PCNs in environmental water were cumulatively nd–260 pg/L [13]. In addition, various results have been reported regarding their presence in the atmosphere, sediments, aquatic and terrestrial organisms, and biota in and outside the polar region [5]. However, few reports have examined the state of PCN concentrations in leachate from landfills where PCN-containing products might have been discarded [14]. Few investigations have been performed because of the need for a complex analytical method that must cover all PCN isomers as target analytes. Our previous report suggested that PCNs listed as POPs in leachate at the final disposal site could be determined with the same pretreatment as for dioxin analysis [14]. In the previous research, DB-5 ms or (5%-Phenyl)-methylpolysiloxane-based similar non-polar capillary column for which the elution behavior of PCNs in GC-HRMS analysis was known [2,15,16] was applied, but these column is not commonly used in dioxin analysis [17]. Ninety one percent of dioxin analyses conducted by laboratories that participated in the ISO/IEC 17043 proficiency test in 2019 had adopted an Rh-12 ms column [18]. In Japan, it is common to measure dioxins by combining an Rh-12 ms and a BPX-DXN capillary column [18]. Therefore, if PCNs can also be analyzed with this column, dioxin and PCNs can be efficiently analyzed without replacing the column.

The purpose of this study was to enable GC-HRMS analysis of PCNs by fitting various parameters of the Rh-12 ms capillary column, and to determine PCN concentrations in landfill leachates, and PCNs behavior in the wastewater treatment processes. We also examined MoCN measurement conditions that could not be quantified in the previous report because D7-labeled 2-MoCN cleanup spikes could not be detected.

MATERIALS AND METHODS

Standards and other reagents
Analytical standards were prepared using certified standards solutions, Mono-Octa PCN Calibration Solutions (Cambridge Isotope Laboratories Inc., Tewksbury, USA). The isotope-labeled analytical solution, which is used as a clean-up and syringe spike, was prepared using certified Mono-Octa PCN Cleanup Spike (Cambridge Isotope Laboratories Inc.) and Mono-Octa PCN Syringe Spike (Cambridge Isotope Laboratories Inc.). Mono-Octa PCN Native PAR Solution (Cambridge Isotope Laboratories Inc.) was used for recovery tests and method detection limits (MDL) tests. Each spike solution was diluted with nonane (dioxin-analysis grade, FUJIFILM Wako Pure Chemical Corporation, Osaka, Japan) before use. Toluene, hexane, and acetone (dioxin-analysis grade, FUJIFILM Wako Pure Chemical Corporation) were used for the extraction and cleanup processes during the analysis.

Leachate samples
Leachate samples collected with a stainless steel bucket from 6 landfills were dispensed into 3 L glass bottles and 500 mL polypropylene (PP) bottles. The leachates were collected five times from these landfills during September 2019 to November 2020. Landfill A-1, A-2, and A-3 are the disposal sites for industrial wastes which are mainly sludge, waste...
extraction of leachate samples by a grass filter (GF/F, Whatman™, Maidstone, UK). PCNs were analyzed as described in the following steps. Extraction and purification for analysis of MoCNs and PCNs

In order to quantify MoCNs and PCNs, the pH of each leachate sample was adjusted to a value from 6 to 9, according to the standard method for dioxin analysis (JIS K 0312) [17]. Then, a cleanup spike (200 pg for each isomer) diluted with acetone was added to each sample. These samples were treated with DioFLOCK® for wastewater samples (Miura Co. Ltd., Matsuyama, Japan), and the flocks formed were collected through glass fiber filter paper (pore size of 0.5 μm). The filter papers were dried in a drying oven heated to 45°C, and a Dean Stark Soxhlet extraction performed with 500 mL toluene was refluxed for 16 h at a rate of approximately 6 cycles per hour. The extracts were concentrated by rotary evaporation, and the solvents changed to 0.5 mL of decane (dioxin-analysis grade, FUJIFILM Wako Pure Chemical Corporation).

The decane solutions were automatically purified using the SPD-600GC system (Miura Co. Ltd.), which can carry out accurate and rapid purification of extracts. This system consists of a multi-layer silica gel column and an active carbon silica gel column. The multi-layer silica gel column was in its usual combination of an upper column packed with 10% AgNO₃ coated silica gel, and a lower column packed with 37% H₂SO₄ coated silica gel. The standard method for dioxin analysis involves putting the decane solution through a multi-layer silica gel column and passing it through 80 mL of hexane heated at 60°C for purification. An activated carbon column will then capture the coplanar compounds. Finally, the activated carbon silica gel column was backflushed with a small amount of toluene and sample solution concentrated to approximately 1.5 mL.

PCNs can be treated in the same way, while MoCNs are tightly bound to the H₂SO₄ coated silica gel and will not elute. Therefore, a 30 cm Teflon tube was connected between the upper and lower columns of the multi-layer silica gel and was cooled by the fan attached to the SPD-600GC system. This procedure allowed hexane in the tube to be cooled at room temperature of 25°C before flowing into the lower column. This modified method promotes elution of MoCNs from the lower column.

Prior to GC-HRMS analysis, 1.5 mL of toluene solution was fortified with a syringe spiked solution containing 13C₁₂ labeled PCB (200 pg each). It was sprayed with N₂ gas and concentrated to less than 20 μL to prepare a sample for GC-HRMS.

GC-HRMS analysis

The samples were analyzed for PCNs using a gas chromatograph/high-resolution mass spectrometer (7890B GC, Agilent, Santa Clara, USA) and JMS-800D Ultra-FOCUS (JEOL, Tokyo, Japan) fitted with an Rh-12 ms column (60 m × 0.25 mm i.d. InventX, Torrance, USA). The GC-HRMS system was tuned to > 10,000 resolving power (10% valley definition) and 1 μL of the sample solution was injected in the splitless mode with a He carrier gas at a flow rate of 1.7 mL/min, with the inlet and transfer line temperatures maintained at 250°C and 270°C, respectively. The GC program was set at an initial temperature of 90°C (hold for 2 min), then ramped to 160°C at a rate of 20°C/min, 220°C at a rate of 3°C/min, 280°C at a rate of 4°C/min, 320°C at a rate of 5°C/min, and finally to 320°C (hold for 1.5 min). The HRMS was operated in electron ionization mode and selected ion monitoring (SIM) mode, as shown in Table 1, at an electron energy of 38 eV and ionizing current of 500 μA. Perfluorokerosene (PFN) was used as a reference compound for lock mass, to adjust for drift. Polychlorinated naphthalene isomers were identified based on the standard method for dioxin analysis. That is, the signal-to-noise ratio (S/N) was > 3, the isotopic ratios of the two isotope peaks were within ±15%, and the coefficient of variation (CV) of the calibration curves was within ±10% for RRcs and within ±20% for RRrs. The relationship between native PCN isomers, clean-up spikes, and
Confirmation of elution order of PCNs using a RH-12 ms capillary column

Mono-Octa PCN Calibration Solutions (Cambridge Isotope Laboratories Inc.), PCN-INC (Wellington Laboratories

syringe spikes are shown in Table 2. The isomers without the corresponding cleanup spikes were quantified using the average RRcs of each chloride.
Inc., Wellington Laboratories Inc., and fly ash samples were analyzed by DB-5 ms capillary column (60 m × 0.32 mm i.d., 0.25 µm film thickness, Agilent) and an Rh-12 ms capillary column, and each peak area was compared to determine the elution order. The chromatogram of the PCNs obtained with the Rh-12 ms capillary column is displayed in Fig. 1, which shows selected ion chromatograms obtained from fly ash samples.

**RESULTS AND DISCUSSION**

**Method application and performance**

In order to assess the application of the modified method, recovery tests (n = 6) were performed using fortified leachate samples. The samples were fortified with native isomers (MoCNs and PCNs) and their stable isotopes for a cleanup spike at concentrations of 97–100 and 183–209 pg/L, respectively. The measured concentrations of each native isomer were 87–112 pg/L with the mean CV of 3.9%. The mean recovery rates of each isomer of the cleanup spike was 72–110% (Table S1). When hexane was heated at 60°C as in the ordinal dioxins analysis, the recovery rate of $^{13}$C$_{10}$ labeled 2-MoCN decreased to 31%, and the recovery rates of Di-Octa cleanup spikes remained at 60 to 106%. These results indicate that the modified method is robust and suitable for the analysis of MoCNs and PCNs in leachate from landfills. The instrumental detection limits (IDLs) were calculated as the standard deviation multiplied by 3, respectively, for five replicate injections of the lowest level standard (native 0.2pg each), the IDLs for the MoCNs and PCNs were 0.02 and 0.01–0.04 pg/L.
The method detection limits (MDLs) were calculated as the standard deviation multiplied by 3, respectively, for six replicate injections of 3,000 mL leachate samples to which 0.5 pg of each native isomer had been added, the MDLs for MoCNs and PCNs were 0.40 and 0.20–0.67 pg/L.

PCNs in landfill leachate samples

The concentration ranges of PCN homologues in landfill leachates at the 6 landfill sites are shown in Fig. 2 (Table S2). DiCNs-OcCN concentrations ranged from 8.0 to 12,000 pg/L in each leachate (Table S3). Among MoCNs and PCNs, MoCNs-PeCNs, relatively high hydrophilic homologues, were predominantly detected in landfill leachates. HxCNs-OcCN homologues were rarely contained in landfill leachates, and their concentrations were less than a few pg/L. The average percentage of homologues of PCNs listed as POPs (DiCNs-OcCN) in landfill leachates were as follows: DiCNs was 27%, TrCNs was 54%, TeCNs was 17%, PeCNs was 2.1%, HxCNs was 0.29%, HpCNs was 0.11%, and OcCN was 0.049%, respectively (Fig. S1). The CVs of DiCNs, TrCNs, and TeCNs were 36%, 47%, and 46%, respectively. The concentration of DiCNs-TeCNs in the leachates from industrial wastes disposal sites were about 10 times higher than those in the leachates from MSW disposal landfill sites.

Table 3 shows the percentage of product-derived isomers in the DiCNs-TeCNs homologues at of each leachate. The leachate from industrial waste sites A-1, A-2, and A-3 had a higher percentage of product-derived DiCNs-TeCNs than the MSW site. This result suggests that the type of PCN isomers contained in the leachate reflects the properties of landfill waste to some extent. The PCNs contents of MSW incineration residues are lower than that of PCN-containing products such as rubber belts and waste woods [19,20]. This might be a reason that higher PCN concentrations were determined in the leachates from industrial waste landfill. Under the GC column conditions of this study, the product-derived isomers 136-TrCN, 1467-TeCN and the incineration-derived isomers 137-TrCN, 1367-TeCN could not be separated on each mass number channel. This separation is a future task to clarify the resources of PCNs in the exudate. At the MSW site, only the incineration residue is landfilled, but at the industrial waste site where various wastes are landfilled, it is difficult to estimate what kind of PCN-containing products in the leachate is derived from. The concentrations of PCNs in the leachates from landfill B-1 were higher than those from B-2, while the percentage of homologues of PCNs were similar between
the landfill sites. Landfill B-1 was constructed earlier than B-2, so advances in a landfill management and incineration technology may have contributed in the decreasing of PCN concentrations in landfill site B-2, but the specific reason still remains unclear.

The means of water qualities except PCNs were shown in Table S4. Although the number of data is not sufficient (only 5 data set for each site), significant positive correlations between PCN concentrations and turbidity, TOC, DOC, COD, and TN. On the other hand, there was no correlation between the PCNs concentrations and BOD. The tendency might suggest that PCNs move with suspended solids (SS) or dissoluble organic substances which is not BOD, assuming turbidity relates to SS. There are future plans to examine the relationship between PCN concentrations and these parameter or other water qualities, for example aquatic humic substances and SS from more survey sites. Moreover, the leachate at landfill sites B-1, B-2 and C, where only incineration residue was landfilled, had low PCN concentrations. A greater variety of leachate samples should be investigated to clarify whether this tendency is common.

**Behavior of PCNs in wastewater treatment process**

The leachate from the landfill sites is collected as raw wastewater and it undergoes several treatment processes in a wastewater treatment plant before exhausted to the environment. In the three out of five times surveys on the leachate from in the landfill sites, the PCN concentrations in the raw wastewater and water sample collected from the different stage of treatment processes in the wastewater treatment plants were analyzed in addition to the leachate samples. Here, the raw wastewater at each leachate treatment plant A and B stands for the mixture of the leachate from landfill site A-1, A-2, and A-3 and landfill site B-1 and B-2, respectively. Figure 3 shows the concentrations of PCN homologues listed as POPs (DiCNs-OcCN) at each treatment process in the plant A and B. The removal rates of PCNs were calculated by comparing the PCN concentrations before and after each treatment processes. Table 4 shows the average removal rate of DiCNs-PeCNs after each treatment at the plant A and B. In the flocculation process, the average removal rates of DiCNs, TrCNs, TeCNs, and PeCNs were 47, 67, 87, and 80%, respectively. Although the removal rates varied depending on the plants or/and survey day, PCNs were effectively removed by the flocculation treatment. Notably, the removals of TeCNs and PeCNs, which are highly hydrophobic homologues, were higher than those of DiCNs and TrCNs. This suggests that PCNs of highly hydrophobic homologues were removed more effectively than that of hightly hydrophilic homologues by a flocculation process. However, the average removal rate of DiCNs-PeCNs in the biological treatment was ranged from −32–70%. There were large variations in the removal rates depending on the collection day or/and plants, and no particular tendency was observed in the biological treatment. In contrast, TOC, DOC, COD and BOD were gently removed in each treatment process, and TN were removed in a process of activated sludge treatment (Figs. S2, S3, S4, and S5), therefore the removal process of PCNs seems to be different form these parameters and the biological treatment might be ineffective for PCNs removal. Activated carbon treatment resulted in a high removal rate for PCNs (Fig. 3 and Table 4). However, since most of the PCNs had already been removed from the leachate during the water treatment process up to that point, it is difficult

![Table 3 The percentage of product-derived isomers in the DiCNs-TeCNs homologues at of each leachate.](http://example.com/table3)

<table>
<thead>
<tr>
<th>Site</th>
<th>A-1</th>
<th>A-2</th>
<th>A-3</th>
<th>B-1</th>
<th>B-2</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>DiCNs</td>
<td>57%</td>
<td>48%</td>
<td>68%</td>
<td>23%</td>
<td>32%</td>
<td>24%</td>
</tr>
<tr>
<td>TrCNs</td>
<td>70%</td>
<td>66%</td>
<td>76%</td>
<td>34%</td>
<td>48%</td>
<td>41%</td>
</tr>
<tr>
<td>TeCNs</td>
<td>78%</td>
<td>68%</td>
<td>76%</td>
<td>45%</td>
<td>50%</td>
<td>59%</td>
</tr>
</tbody>
</table>

A-1, A-2 and A-3 ; leachate-controlled landfill sites for industrial wastes
B-1, B-2 and C ; leachate-controlled landfill sites for municipal solid wastes

Type of isomers at each homologue; derived from product or incineration [2,15,16].

<table>
<thead>
<tr>
<th>Product (Halowax)</th>
<th>incineration (fly-ash)</th>
<th>Included in both</th>
</tr>
</thead>
<tbody>
<tr>
<td>DiCNs 14/16</td>
<td>13,27,12,23,18</td>
<td>15,17/26</td>
</tr>
<tr>
<td>TrCNs 146,145</td>
<td>127,125,126,137,167/236,123</td>
<td>136/135,124,138,128</td>
</tr>
<tr>
<td>TeCNs 1257/1246,1358/1236,1248,1258/1268,1458</td>
<td>1247,1368/1256/1235,1237,1234,1267,2367,1238</td>
<td>1357,1367/1467,1245,1278</td>
</tr>
</tbody>
</table>

Isomers that cannot be separated into product or incineration are underlined.
to verify the treatment efficiency. Table S5 shows the mean concentrations of PCN homologues in leachate wastewater at each process of the treatment plant. After the activated carbon treatment, DiCNs-OcCN were decreased to below or near MDL. In addition, MoCNs was also decreased to below 6.9 pg/L after the treatment. These results demonstrated that MoCNs and PCNs removal progressed through these water treatment processes. So the leachates were adequately released at a lower concentration than in river water [13]. In order to determine the types of water treatment processes effective in removing PCNs, it is necessary to obtain results from fact-finding surveys at monitoring sites as well as laboratory experiments of unit operations.

MoCN isomers in landfill leachate samples

Figure 4 shows the mean concentrations of MoCN isomers and DiCNs-OcCN at each landfill leachate (Table S4). At some industrial waste disposal sites, the concentrations of MoCNs were much higher than those of PCNs. Similar to metal compounds and pentachlorophenol (PCP), MoCNs
have been used as wood preservatives, especially in its role as an antiseptic \[21,22\]. In a study investigating wood preservative POPs in waste timber from demolished buildings and their recycled products in Japan, chlorden, PCP, and PCNs were detected in wood chips from waste timber \[20\]. In addition, it has been reported that most of the chlorinated naphthalene in recycled products are MoCNs and DiCNs \[20\]. The industrial waste at disposal sites A-1, A-2, and A-3 contains waste wood, and thus, MoCNs and DiCNs that leached out from these sites are probably derived from wood preservatives that had been used a long time ago. Moreover, the leachates in site A contain a high proportion of 1-MoCN in which chlorine is substituted at the α-position. This result is consistent with previous studies that have described that most of the industrially produced MoCNs are considered to be 1-MoCN \[1,2\].

CONCLUSIONS

Isomer-specific analysis of MoCNs and PCNs in landfill leachate was developed in accordance with standard dioxin analysis methods (JIS K 0312). The elution order of PCNs by the Rh-12 ms capillary column was determined. The modified method of cooling hexane on multilayer silica gel made it possible to measure MoCNs in a stable manner. Among the DiCNs-OcCN listed as POPs, most of the PCN homologues detected in the leachates were DiCNs-PeCNs, and their concentrations varied depending on the landfill sites. Although further investigation is required for the removal rate of each water treatment process, flocculation and an active carbon treatment were assumed to be effective for the removal of PCNs, and the PCN concentrations were reduced from MDLs to a few pg/L thorough water treatments, and the leachates were adequately released. MoCNs that may be derived from wood preservatives were detected at much higher concentrations than PCNs in landfill sites containing waste wood.

In order to estimate the potential risk of PCNs in landfill leachates and manage them appropriately over the long term, it is necessary to clarify the actual state of leaching of various leachates and study more effective treatment methods. The modified method developed in this study is practical and will help to elucidate the content of PCNs, including MoCNs, in leachate from landfill sites.

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![Fig. 4](image-url) The mean concentrations of MoCN isomers and DiCNs-OcCN in each landfill leachate.
search Institute of Environment, Agriculture and Fisheries, Osaka, Japan) for their assistance with laboratory analysis.

SUPPLEMENTARY MATERIALS

Supplementary Materials file for this article is available at the link below.
https://www.jstage.jst.go.jp/article/jwet/19/2/19_20-135/_supplement/_download/19_20-135_1.pdf

REFERENCES


