Fly ash-mediated formation of polychlorinated naphthalenes during secondary copper smelting and mechanistic aspects

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HIGHLIGHTS

- High concentrations of PCNs were formed via thermal reactions on fly ash.
- More chlorinated homologs were dominant in the formed PCNs.
- Chlorination of naphthalene is identified as an important PCN formation mechanism.
- A congener-specific pathway from naphthalene to octachloronaphthalene is proposed.

ABSTRACT

Thermal experiments (at 250–450 °C for 10–240 min) on fly ash from secondary copper smelting process (SeCu) were performed to study the polychlorinated naphthalene (PCN) formation in the post-zone of a secondary copper smelter. Unexpectedly high concentrations of PCNs were formed. Total PCN concentrations and toxic equivalents were 47–104 and 44–80 times higher than the initial concentration and toxic equivalent, respectively. The thermal disposal of SeCu fly ash should therefore be reconsidered. The kinetic of each homolog was determined under different thermal conditions. Less chlorinated homologs favored 350 °C and more chlorinated homologs favored higher temperature. Most of the homologs reached an equilibrium of formation and degradation within 30 min, except octachloronaphthalene which did not appear to reach such an equilibrium even after 240 min. Chlorine substitution of the formed PCNs was identified and a similar pattern was found in chlorination products starting with naphthalene and chlorine. Furthermore, inorganic chlorine and unsubstituted naphthalene were found in the reaction products, confirming that the formation of naphthalene and the chlorination of that naphthalene could occur and could be suggested to be an important PCN formation route. A detailed formation pathway from naphthalene through octachloronaphthalene is proposed.

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1. Introduction

Polychlorinated naphthalenes (PCNs) are candidates for inclusion in the Persistent Organic Pollutant Protocol of the United Nations Economic Commission for Europe (Lerche et al., 2002) and it has been proposed that they be included in Annexes A, B, and/or C of the Stockholm Convention. There is growing public concern about the formation and emission of PCNs. Unintentional emissions from industrial activities are currently considered to be the main sources of PCNs because industrial applications of PCNs have ceased in many countries (Falandysz, 1998). PCNs are always emitted from thermal-related industries along with polychlorinated dibenzo-p-dioxins (PCDDs), polychlorinated dibenzofurans (PCDFs), and polychlorinated biphenyls, and these chemicals all have similar physicochemical properties, biological effects, and toxic mechanisms (Hanberg et al., 1990).

In municipal incineration processes, PCNs may be formed from polycyclic aromatic hydrocarbons through similar de novo pathways to those that produce PCDFs (Iino et al., 1999; Imagawa and Lee, 2001; Weber et al., 2001; Oh et al., 2007). Kim and coworkers found that PCNs could be formed by the coupling of chlorinated phenoxy radicals (formed from chlorophenol) at high temperatures (Kim et al., 2005, 2007). It has also been suggested that chlorination reactions are involved in PCN formation during thermal-related process (Imagawa and Lee, 2001; Jansson et al., 2008; Liu et al., 2010). The industrial production of PCNs was achieved by
chlorinating molten unsubstituted naphthalene using chlorine gas in the presence of transition metal chlorides (Schneider et al., 1998). The further chlorination of the resulting PCNs is of particular interest because the most toxic congeners are chlorinated at the lateral positions. Determining the factors that govern the addition of chlorine to PCNs may facilitate the control of these toxic chemicals in industrial exhausts and residues. Chlorination can be catalyzed by transition metal species, especially copper compounds, which are known to have strong catalytic effects (Ryu et al., 2013). In a recent study, Ryu et al. (2013) determined the congener patterns of naphthalene was chlorinated using a mixture of copper(II) chloride and silicon dioxide as a matrix.

In addition to municipal solid waste incinerators (MSWIs) (Abad et al., 1999; Jansson et al., 2008; Hu et al., 2013), various metallurgical industries, particularly SeCu, are becoming increasingly important sources of PCNs (Ba et al., 2010; Nie et al., 2011; Liu et al., 2012a,b). Liu et al. (2014) summarized PCN emission factors and annual emissions for 14 industrial thermal processes, and suggested that SeCu potentially contributes a large proportion of PCN emissions in China, having comparable emission factors to those found for MSWIs. However, no attempt was made currently to determine the detailed PCN patterns and formation pathways that led to the observed PCN emissions from SeCu or other metallurgical processes. Despite the intensive presence of PCNs and toxic effects PCNs can have, the PCN emissions from SeCu have been studied much less than MSWI emissions in the past.

This study builds on previous work, and was conducted to gain a first insight into the PCN formation mechanisms during SeCu. A laboratory-scale system using SeCu fly ash as a reaction matrix was used to simulate the thermal reactions that produce PCNs during SeCu under controlled, realistic conditions. The main goal of this work was to characterize the homolog and congener patterns of the PCNs that are generated during simulated thermal processes using SeCu fly ash and to identify the underlying mechanisms that are responsible for producing the PCNs. The kinetics involved in the formation of the total PCNs and the mono- to octa-chlorinated naphthalenes (mono- to octa-CNs) were also determined in detail. The results may allow methods for controlling PCN emissions during metallurgical processes to be developed.

2. Materials and methods

2.1. Experimental design

It is widely recognized that fly ash is a matrix on which heterogeneous catalyzed reactions occur, so it was used as the matrix for the thermal experiments described here, offering the advantage that as close as possible to a “real” active surface (Milligan and Altwicker, 1993; Addink and Olie, 1995; Pekárek et al., 2007). Fly ash was collected from a reverberatory furnace of a secondary copper smelter in eastern China. The production capacity of each furnace in the smelter was 110 t. Fly ash has a complex composition, so to allow us to understand the nature of the fly ash used in the study, its inorganic and organic components and surface properties were determined using inductively coupled plasma optical emission spectrometry (using an Optima 8300 instrument; Perkin Elmer, Waltham, MA, USA), scanning electron microscopy with energy dispersive X-ray spectrometry (using a S-3000N instrument; Hitachi, Tokyo, Japan), and an ASAP 2400 system (Micromeritics, Norcross, GA, USA) and the specific surface area of the materials were calculated using the BET model. High concentrations of carbon, chloride, oxygen, and metals such as Zn, Ca, Pb, Al, and Fe were found on the fly ash surface, and the Cu concentration was particularly high (262.4 mg g⁻¹). The SeCu fly ash was compared with fly ash from other thermal-related processes, and it was found that the Cu concentration was much higher in the SeCu fly ash than in fly ash from a MSWI plant (0.47–0.33 mg g⁻¹) (Cobo et al., 2009) and fly ash from an integrated iron and steel facility (0.12–7.95 mg g⁻¹) (Tsai et al., 2007). PCNs are usually formed along with PCDDs and PCDFs, and the formation of all of these chemicals is probably favored by the presence of carbon, chlorine, and transition metals, such as Cu, on the surfaces of fly ash particles (Addink and Olie, 1995; Hatanaka et al., 2004; Stanmore, 2004; Chin et al., 2011). Therefore, we speculated that the fly ash we had prepared could lead to the formation of PCNs under favorable conditions. The fly ash had a high surface area obviously mean that the surface and external reactants involved in reactants could easily come into contact with each other (Milligan and Altwicker, 1993). The detailed fly ash characterization information is given in Tables S1–S3.

The PCN formation process was modeled at the laboratory scale. The fly ash was placed as a fixed bed to simulate collected ash particles in a SeCu plant, and an air stream was passed through the fixed bed. The PCN concentrations found at different temperatures and/or times have been determined in some previous studies, whereas the PCN concentrations found after only one or two selected reaction times have been determined in other studies. The temperatures 250 °C, 350 °C, and 450 °C were used in this study because that range covers the temperatures at which PCNs could form in the cooling zone of a real plant. PCDDs and PCDFs can, theoretically, form rapidly (Altwicker et al., 1992), and the residence time of fly ash particles in the post-combustion zone can be hours (Addink and Olie, 1995), so laboratory tests with reaction times of several minutes to hours were used to validate the kinetic models and identify relationships between PCN formation and residence time.

The simulation system was a quartz tube reactor (60 cm long and 4.5 cm in diameter) within a tube furnace that provided even heating and precise temperature control. A schematic of the reaction system used is shown in Fig. S1. An even layer of fly ash (0.2 g) was placed on a porcelain boat in the reactor and the system was heated to 250, 350, or 450 °C for 10, 30, 120, or 240 min (as shown in Table S4). The air stream was kept at a constant flow rate of 50 mL/min to simulate the open system in a SeCu, and was left running for 30 min after the end of the heating to trap products diffused into the gas phase. The outlet gas was quenched as soon as it left the reactor by passing it through three absorption bottles each containing 50 mL of toluene, which were cooled in ice.

2.2. Analytical methods

The toluene from the traps and the fly ash residues were analyzed for PCNs by isotope dilution high resolution gas chromatography and high resolution mass spectrometry (HRGC/HRMS) (Ba et al., 2010; Li et al., 2010). The toluene from the traps and residues were spiked with a mixture of 13C10-labeled PCN internal standards (ECN-5102, 13C10-CN-27, -42, -52, -67, -73, and -75; Cambridge Isotope Laboratories, Andover, MA, USA). Each residue sample was digested in hydrochloric acid, freeze-dried, then Soxhlet extracted with 250 mL of toluene for 24 h. Each residue extract and sample of toluene from the traps was cleaned by passing it through a series of chromatographic columns (an acidified silica gel column, a multilayer silica gel column, then a basic alumina column). The cleaned extract was then concentrated, by rotary evaporation and then under a gentle stream of nitrogen, to about 20 μL. The initial PCN concentration in the fly ash had to be determined so that the PCNs formed in the reaction system could be determined, so the original fly ash was analyzed using the same method as was used for the solid residues.
The amounts of PCNs in the vapor phase (i.e., in the toluene trap) and the residue were compared, and more than 99.5% of the total PCN concentration was found in the vapor phase. This confirmed that the PCNs volatilized during the tests (that is, the PCNs could be transferred from the solid matrix to the gas phase when the fly ash was heated). It is, therefore, reasonable to use the vapor phase PCN concentration, rather than the sum of the PCN concentrations in the two phases, in the subsequent assessments related to the formation of PCNs.

The PCN concentrations found in the vapor phase produced during the thermal reaction tests under the different test conditions are shown in Figs. 1 and 2. Unexpectedly large amounts of PCNs, 33.40–73.94 µg g⁻¹, were found to be produced during the thermal treatments, and the highest PCN yield was more than two orders of magnitude higher than the PCN concentration in the initial fly ash (0.708 µg g⁻¹). These results indicate that the SeCu fly ash strongly promoted the formation of PCNs during the thermal reactions.

The dioxin-like toxicities of PCNs have been investigated previously (Hanberg et al., 1990; Blankenship et al., 2000; Villeneuve et al., 2000), and the relative potency factors that relate individual PCN congeners to 2,3,7,8-tetrachlorodibenzop-α-dioxin that were estimated in those studies were used to evaluate the toxicity of the PCNs found in this study (Noma et al., 2004; Liu et al., 2010). The PCN toxic equivalent (TEQ) for a sample was defined, in this study, as the sum of the concentration of each congener multiplied by the relative potency factor for that congener, determined by Noma et al. (2004). The PCN TEQs in the products were 36.38-48.41 TEQ ng/g, which were 44–80 times higher than the initial PCN TEQs. Taking this amplification of the toxicity of PCNs in SeCu fly ash into consideration, the thermal disposal of fly ash (such as recycling in furnaces) in some SeCu plants and in other metallurgical plants should be reconsidered and reevaluated to decrease the potential for contaminating the environment.

The PCN concentrations were different when different reaction conditions were used. The PCN yield increased remarkably, from 33.40 to 73.94 µg g⁻¹, as the temperature was increased from 250 to 450 °C. Fig. 1 shows the relationship between the temperature and the total PCN concentration and the degree of chlorination (the average number of chlorine atoms in each molecule). In general, the degree of chlorination increased as the temperature was increased, which indicated that the chlorination reaction was more favored at higher temperatures within the range 250–450 °C. The amount of PCNs formed as a function of time (Fig. 2) followed a similar trend to that found using fly ash from a municipal waste incinerator (Schneider et al., 1998). It can be seen from Fig. 3 that the amount of PCNs formed had increased after a short reaction time (less than 30 min), and the amount formed was the same (52.29–58.04 µg g⁻¹) even after longer reaction times (up to 240 min). Notably, the homologs with different degrees of chlorination peaked at different temperatures and times, the details of which are presented and discussed next.

3.2. Homolog profiles

The dominant homologs in the initial fly ash were the pentachloronaphthalenes (penta-CNs), heptachloronaphthalenes (hepta-CNs), and hexachloronaphthalenes (hexa-CNs), which contributed 21.8%, 19.0%, and 18.1% of the total PCN concentration, respectively. However, the PCN profile had changed significantly during the thermal reaction tests.

3.1. PCN concentrations

The reactor was kept at 350 °C for 6 h and time were used, was performed in triplicate. The PCN recoveries of the two most intense characteristic m/z ratios were monitored for each PCN homolog group. The recoveries to be calculated. All 75 PCN congeners were analyzed using an Agilent 6890 high resolution gas chromatograph (Agilent Technologies, Santa Clara, CA, USA), coupled to a DFS MS instrument (Thermo Fisher Scientific, Waltham, MA, USA), which was operated in selective ion monitoring mode. The recoveries of the labeled PCN standard (ECN-5260, containing 13C10-CN-64; Cambridge Isotope Laboratories) was added, to allow the PCN concentrations in the two phases, in the subsequent assessments related to the formation of PCNs.

2.3. Quality control and assurance

A blank test was performed after every three sample tests to determine the PCNs produced when reactants were not present. The PCN peaks were identified from their retention times compared with available individual PCN standards, ion ratios and by taking into consideration the PCN elution order on the DB-5 column. The recoveries of the 13C10-labeled PCN congeners relative to the labeled injection standard were 30–123% for both matrices.

3. Results and discussion

3.1. PCN concentrations

Before each extract was analyzed by HRGC-HRMS, a 13C10-labeled PCN standard (ECN-5260, containing 13C10-CN-64; Cambridge Isotope Laboratories) was added, to allow the PCN recoveries to be calculated. All 75 PCN congeners were analyzed using an Agilent 6890 high resolution gas chromatograph (Agilent Technologies, Santa Clara, CA, USA), equipped with a DB-5 MS fused-silica column (60 m long, 0.25 mm i.d., 0.25 µm film thickness; Agilent Technologies, Santa Clara, CA, USA), coupled to a DFS MS instrument (Thermo Fisher Scientific, Waltham, MA, USA), which was operated in selective ion monitoring mode. The two most intense characteristic m/z ratios were monitored for each PCN homolog group.

Fig. 1. Total polychlorinated naphthalene concentrations (bars) and degree of chlorination (line) in the products of thermal experiments at 250–450 °C and in the original fly ash (OA: original fly ash).

Fig. 2. Total polychlorinated naphthalene concentrations after different thermal treatment times.
after the fly ash had been treated, and octachloronaphthalene (octa-CN) was the most abundant homolog in all of the tests, accounting for 35.4–59.1% of the total PCN concentrations. Hepta-CN was the second most abundant homolog in the tests, contributing 23.8–34.7% of the total PCN concentrations. The penta-CN and hexa-CN contributions to the total PCN concentrations decreased to 4.2–7.1% and 7.8–11.3%, respectively, after the tests. These results demonstrate that heating the fly ash caused the more chlorinated homologs to become dominant, even though the less chlorinated homologs had initially been dominant in the fly ash. This obvious transformation from less chlorinated homologs dominating to more chlorinated homologs dominating suggests that a chlorination reaction, changing less chlorinated congeners into more chlorinated congeners, may contribute to the formation of PCNs in thermal treatments.

The homolog profiles changed noticeably within the temperature range 250–450 °C. Fig. 3 shows that the concentration of the less chlorinated homologs (mono-CNs to hexa-CNs) peaked at 350 °C but that the concentrations of the more chlorinated homologs (hepta-CNs and octa-CN) increased continuously as the temperature was increased. These results indicate that the production of the less chlorinated homologs was favored at 350 °C, which may have been caused by the mechanism involved having its lowest activation energy at 350 °C. The production of the more chlorinated homologs was favored at higher temperatures (possibly even higher than 450 °C), implying that the mechanism involved may have a lower activation energy at such high temperatures.

The kinetics of the formation of the mono to octa-CNs in terms of the net concentrations of each (the difference between the total concentration and the initial concentration) are presented in Fig. 4. The highest concentrations of most of the homologs (mono-CNs to hepta-CNs) were reached within 30 min of heating, whereas the most chlorinated homolog, octa-CN, continued to be formed until 240 min. These phenomena suggest that PCN formation and degradation processes may have occurred on the fly ash simultaneously.

The formation of all the homologs was dominant at the beginning of the heat treatment, and the formation and degradation of most of the homologs reached an equilibrium within 30 min. Only octa-CN did not appear to reach such an equilibrium.

The mechanistic relationships between each of the PCN homologs were investigated by determining the Pearson correlation coefficient (R) for the relationships between each of the PCN homologs. It can be seen from Table S5 that adjacent PCN homolog groups were closely related to each other. Similar results were found in a previous study performed by Oh et al. (2007), who found that adjacent PCN homologs were closely related to each other when municipal waste incineration was simulated. This correlation analysis provided another support for the chlorination pathway suggested above being responsible for PCN formation.

3.3. Congener patterns

The concentration of each congener was expressed as a fraction of the concentration of its entire homolog group (see Fig. 5), to make identifying the PCN congener distribution patterns easier to achieve. The substituent position numbers for PCNs are shown in Fig. S2. The congener patterns were used because they offer a detailed “fingerprint” of the PCN pattern, potentially allowing the identification of the sources of the PCNs produced from thermal formation on fly ash from SeCu.

The congener patterns were unexpectedly similar to the patterns that have been found when naphthalene has been chlorinated, as reported by Schneider et al. (1998), who determined the congener patterns in industrial PCN products that had been produced by chlorinating molten naphthalene using chlorine gas in the presence of a transition metal chloride. The dominant congeners in each homolog in industrial PCN products and in the products formed in our study are shown in Table 1. It can clearly be seen that the PCNs in both the industrial products and this study had a specific congener pattern, with chlorine substituents usually present at the 1 and 4 positions and the 2, 5, 6, and 8 positions often being attacked by chlorine atoms. This suggests that the industrial products and the products of this study may share the same formation pathway, which supports the idea that the PCNs found in this study were formed through the chlorination pathway.
starting from naphthalene. The PCN congener patterns found in this study were also very similar to the congener patterns found in PCNs synthesized in the laboratory using the chlorination reaction starting from naphthalene and catalyzed by CuCl₂ (Ryu et al., 2013). This offers greater support for the chlorination pathway being responsible for the formation of PCNs in our study. As mentioned above, the chlorination reaction can be catalyzed by transition metals. An aromatic molecule such as naphthalene can be directly chlorinated through the route shown in Eqs. (1) and (2) (Stiegltz et al., 1989; Takaoka et al., 2005; Ryu et al., 2013). The Deacon reaction can transform HCl into Cl₂ and result in chlorination catalyzed by the transition metal (see Eq. (3)) (Gullett et al., 1992).

\[
\text{ArH} + \text{CuCl}_2 \rightarrow \text{ArHCl}^+ + \text{CuCl}
\]  
\( (1) \)

\[
\text{ArHCl}^+ + \text{CuCl}_2 \rightarrow \text{ArCl} + \text{CuCl} + \text{HCl}
\]  
\( (2) \)

\[
4\text{HCl} + \text{O}_2 \rightarrow 2\text{Cl}_2 + \text{H}_2\text{O}
\]  
\( (3) \)

It has been suggested in several studies that the de novo synthesis of PCNs occurs in MSWIs, from the degradation of polycyclic aromatic hydrocarbons followed by their repeated chlorination (Iino et al., 1999; Weber et al., 2001). A considerable amount of carbonaceous material was found in the fly ash used in this study, so a similar PCN formation pathway may have been feasible. A large amount of chlorine was found in the fly ash (8.97% by weight, shown in Table S2), but we were unable to determine if the chlorine species present in the matrix caused the PCNs to be formed through chlorination reactions. Eqs. (1)–(3) show that chlorination reactions can produce gaseous inorganic chlorine. Therefore, detecting gaseous inorganic chlorine in our reaction system would offer strong evidence for the occurrence of chlorination reactions. We tested this hypothesis by attempting to detect inorganic chlorine in the system. The amount of gaseous inorganic chlorine produced in Experiment 1 was determined by ion chromatography (the analytical method is described in Supporting Information).

Figure 5. Polychlorinated naphthalene congener distribution patterns in the thermal treatment samples (the congener concentration is shown relative to its total homolog concentration, as a wt%). R1: 30 min, 350 °C; R2: 30 min, 250 °C; R3: 30 min, 450 °C; R4: 10 min, 350 °C; R5: 120 min, 350 °C; R6: 240 min, 350 °C.

Table 1 Dominant polychlorinated naphthalene congeners in Halowax and generated in thermal treatments.

<table>
<thead>
<tr>
<th>Homolog</th>
<th>Halowax (Schneider et al., 1998)</th>
<th>Formation in fly ash</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mono-CN</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Di-CN²</td>
<td>1.4/1.6</td>
<td>1.4/1.6</td>
</tr>
<tr>
<td>Tri-CN¹</td>
<td>1.46/1.45</td>
<td>1.46/1.24/1.45</td>
</tr>
<tr>
<td>Tetra-CN³</td>
<td>1.2,5,7/1.2,4,6/1,2,4,7/1,2,6,8</td>
<td>1.2,5,7/1.2,4,6/1.2,4,7/1,2,5,7/1.3,5,8/1,2,5,8/1,2,6,8</td>
</tr>
<tr>
<td>Penta-CN⁴</td>
<td>1.2,4,6,8</td>
<td>1.2,4,6,8/1.2,4,5</td>
</tr>
<tr>
<td>Hexa-CN⁵</td>
<td>1.2,4,5,6,8/1,2,4,5,7,8</td>
<td>1.2,3,5,7,8/1,2,3,4,5,6</td>
</tr>
<tr>
<td>Hepta-CN⁶</td>
<td>1.2,3,4,5,6,8</td>
<td>1.2,3,4,5,6,7/1,2,3,4,5,6,8</td>
</tr>
</tbody>
</table>

¹ Mono-CN: monochloronaphthalene.  
² Di-CN: dichloronaphthalene.  
³ Tri-CN: trichloronaphthalene.  
⁴ Tetra-CN: tetrachloronaphthalene.  
⁵ Penta-CN: pentachloronaphthalene.  
⁶ Hexa-CN: hexachloronaphthalene.  
⁷ Hepta-CN: heptachloronaphthalene.
Fig. 6. Proposed pathway for the formation of polychlorinated naphthalenes. MoCN: monochloronaphthalene; DiCN: dichloronaphthalene; TrCN: trichloronaphthalene; TeCN: tetrachloronaphthalene; PeCN: pentachloronaphthalene; HxCN: hexachloronaphthalene; HpCN: heptachloronaphthalene; OCN: octachloronaphthalene.
monochloronaphthalene and more chlorinated PCN homologs. We postulated a chlorination pathway from naphthalene to octa-CN based on the dominant congeners found within each homolog, and this pathway is shown in Fig. 6. Despite the results of this study suggesting that the chlorination reaction may be important in the formation of PCNs, the contribution of de novo synthesis cannot be neglected because, as mentioned above, the de novo synthesis of PCNs was found to be possible in previous studies (Benfenati et al., 1991; Iino et al., 1999; Imagawa and Lee, 2001; Weber et al., 2001). It seems likely that the large amounts of PCNs formed in thermal treatments are not formed through a single pathway but rather through a combination of chlorination reactions and de novo synthesis (Oh et al., 2007; Jansson et al., 2008; Liu et al., 2014). Further studies will be required to identify other formation pathways that may be involved.

4. Conclusion

We performed a series of laboratory-scale experiments using fly ash as the reaction matrix to simulate the thermal formation of PCNs in the post-zone of a secondary copper smelter. Under different thermal conditions, unexpectedly high concentrations of PCNs about two orders of magnitude higher than the initial PCNs were detected. Kinetic examination demonstrated that less chlorinated homologs seemed to favor 350 °C and more chlorinated homologs seemed to favored higher temperature. Most of the homologs could be formed within a short time period, while octa-CN did not appear to reach equilibrium of formation and degradation within our investigated reaction time. The de novo synthesis of polychlorinated naphthalenes following a consecutive chlorination of naphthalene molecules was suggested to be an important pathway for the formation of PCNs. This study will provide valuable information that improves our understanding of the mechanism and kinetics by which PCNs are formed during metallurgical processes and offer important improvements in our ability to control and minimize the unintentional formation and emission of PCNs.

Acknowledgments

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Appendix A. Supplementary material

Detailed characterization information of the reaction matrix, setting of the reaction conditions, Pearson coefficients between each of the produced polychlorinated naphthalene homologs, schematic of the reaction system, structure and commonly used numbering system of PCN congeners, and the description of analysis method for inorganic chlorine and naphthalene are provided as supporting information. Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.chemosphere.2014.09.052.

References

