Levels and distribution patterns of short chain chlorinated paraffins in sewage sludge of wastewater treatment plants in China

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ABSTRACT

Short chain chlorinated paraffins (SCCPs) are listed as persistent organic pollutant candidates in the Stockholm Convention and are receiving more and more attentions worldwide. In general, concentrations of contaminants in sewage sludge can give an important indication on their pollution levels at a local/regional basis. In this study, SCCPs were investigated in sewage sludge samples collected from 52 wastewater treatment plants in China. Concentrations of total SCCPs (ΣSCCPs) in sludge were in the range of 0.80–52.7 μg/g dry weight (dw), with a mean value of 10.7 μg/g dw. Most of SCCPs in the sludge samples showed a similar congener distribution patterns, and C11 and Cl7,8 were identified as the dominant carbon and chlorine congener groups. Significant linear relationships were found among different SCCP congener groups (r² ≥ 0.9). High concentrations of SCCPs in sewage sludge imply that SCCPs are widely present in China.

1. Introduction

As a class of highly complex technical mixtures, chlorinated paraffins (CPs) contain thousands of isomers, diastereomers and enantiomers (Bayen et al., 2006; Pellizzato et al., 2007; Feo et al., 2009). The commercial mixtures of CPs are produced by chlorination of n-alkane feedstocks with carbon-chain lengths from 10 to 30 (Tomy et al., 1998). Up to now, more than two hundreds of commercial CP products have been used for different industrial applications, such as extreme pressure lubricant additive, plasticizers, flame retardants and paint additives, etc (Santos et al., 2006). In general, CPs are divided into short chain CPs (C10–13, SCCPs), medium chain CPs (C14–17, MCCPs) and long chain CPs (C18–30, LCCPs) according to their carbon-chain lengths (Zencak and Oehme, 2006).

Due to widespread and unrestricted use in the past decades, CPs have been found widespread in the environment and have been detected in different environmental matrices, such as air (Peters et al., 2000; Barber et al., 2005), water (Coelhan, 2010), soil (Nicholls et al., 2001; Zeng et al., 2011a), sediments (Tomy et al., 1999; Marvin et al., 2003; Iozza et al., 2008), sludge (Stevens et al., 2003), biota (Tomy et al., 2000), terrestrial wildlife (Jansson et al., 1993) and human foods (Iino et al., 2005), even in the environment of remote areas (Tomy et al., 1999). Of all the CP mixtures, SCCPs tend to behave in a similar way to persistent organic pollutants (POPs), leading to some countries and organizations to impose regulation on the production and use of these compounds (Pellizzato et al., 2009). Because of their persistence (Iozza et al., 2008), toxicity (Cooley et al., 2001; Warnasuriya et al., 2010), bioaccumulation (Fisk et al., 1996; Houde et al., 2008; Zeng et al., 2011b) and potential for long-range transport abilities (Tomy et al., 1999), SCCPs have been listed as POP candidate by the Stockholm Convention. However, data on the distributions, environmental fate and toxicity of SCCPs are still scarce compared to other POPs, mainly due to the complexity of analytical methodologies (Eljarrat and Barcelo, 2006; Santos et al., 2006).

Release of SCCPs into the environment can occur during production, incorporation into polymers or products, and through usage and disposal or recycling of these products. The released SCCPs can finally enter the municipal wastewater network through atmospheric deposition or non-point source discharge, and accumulate in the wastewater treatment plants (WWTPs). Due to their high lipophilicity and environmental persistency, a large proportion of SCCPs tend to be preferentially partitioned or concentrated to high organic carbon-containing sewage sludge during wastewater treatment (Zeng et al., 2011b). Therefore, the concentrations of SCCPs in sewage sludge from WWTPs can give an important indication of the general exposure and usages of these contaminants and their pollution status in the region. However, up to now,
levels and distribution of SCCPs in sewage sludge are only found in very limited works (Rieger and Ballschmiter, 1995; Nicholls et al., 2001; Stevens et al., 2003).

China is the largest producer of chlorinated paraffins in the world (De Boer et al., 2010), with most of the manufacturing plants located in the east of China. Since 1990, the annual production volumes of CPs are increasing rapidly (Zeng et al., 2011b). However, studies on SCCPs in the environment in China are still very limited. Only several studies on the distributions of SCCPs in e-waste dismantling areas, wastewater irrigated regions, and sediments have been reported recently (Yuan et al., 2010; Zeng et al., 2011a, 2011b).

The aim of this study is to assess the SCCPs load in sewage sludge of WWTPs collected from different cities in China. Congener group patterns were also studied to investigate potential geographic distributions. It is hoped that the results in this study can provide valuable information to indicate the pollution levels and distribution patterns of SCCPs in the human living environment in China.

2. Materials and methods

2.1. Materials, standards and reagents

All reagents (cyclohexane, dichloromethane, n-hexane) were pesticide residue grade purchased from Fisher Scientific (Hampton, NH). Silica gel and Florisil (60–100 mesh) were obtained from Merck (Whitehouse Station, NJ). Commercial standards of SCCPs (C10, C11, 51.5%, 55.5%, and 63.0% chlorine content, 100 ng/L; C10, 50.2%, 55.0%, and 60.0% chlorine content, 10 ng/L; C11, 50.2%, 55.0%, and 60.0% chlorine content, 10 ng/L) and MCCPs mixtures (C14, 42.0%, 52.0%, and 57.0% chlorine content, 100 ng/L) in cyclohexane as well as p-hexachlorocyclohexane (p-HCH, solution in cyclohexane, 10 ng/L) were purchased from Ehrenstorfer GmbH (Augsburg, Germany). 13C10-trans-chlordane (99%) in n-nonane was supplied by Cambridge Isotope Laboratories (Andover, USA).

2.2. Sampling site and sample collection

A total of 52 WWTPs were selected for collection of sewage sludge samples from 25 cities in China during the period of August 2010 to March 2011. The sampling sites cover 13 Chinese provinces and most of them are located in eastern China. Fig. 1 shows the detailed distribution of the Chinese manufacturing plants of CPs and the sampling locations from different provinces in this study. All sludge samples were freeze-dried and homogenized, then stored in a freezer at -20 °C until analysis. A survey form was also associated with each sample. Details of each WWTPs in regards to treatment capacity, serving population, sewage sources (domestic, industrial, or combined), and type of sludge treatment are supplied in Table SI-1 in Supplementary material.

2.3. Sample pretreatment

Sample pretreatment for SCCP analysis were based on the procedure described previously with some modifications (Zeng et al., 2011a). Prior to extraction, 0.2 g of sludge sample was mixed with 10 g anhydrous sodium sulfate, spiked with 10 ng 13C10-trans-chlordane surrogate standard and extracted by accelerated solvent extractor (ASE, Dionex ASE 350, Canada) with mixture solvent of dichloromethane/n-hexane (1:1, v/v). After the extraction, the extract was rotary-evaporated to about 2 mL and then subjected to clean-up using a multilayer silica-Florisil composite column, which was filled with 3 g Florisil, 2 g activated silica gel, 5 g acid silica gel (30%, w/w) and 4 g anhydrous sodium sulfate from bottom to top. The column was pre-cleaned with 50 mL hexane, and then eluted in sequence with 40 mL of hexane (F1) and 100 mL of dichloromethane/hexane (1:1, v/v) (F2). CPs eluted in the second fraction F2 and the solvent was exchanged to cyclohexane and concentrated to a final volume 200 μL. Finally, 10 ng p-HCH was added as an internal standard to determine the recoveries before instrument analysis.

2.4. Identification and quantification

All analyses were performed on high-resolution gas chromatography/electron capture detector (HRGC/ECD) or low-resolution mass spectrometry (HRGC/ECNI-LRMS, Agilent, USA) using selected ion monitoring (SIM) mode. Under ECNI conditions, the most and second most abundant isotope ions of [M–Cl]− ions from isotopic abundance and response factors. The sum of SCCP concentration (ΣSCCPs) were calculated based on the quantification procedure described by Reth and Oehme (Reth et al., 2005).

2.5. Quality assurance and quality control (QA/QC)

Strict quality assurance and control measures were implemented to ensure the identification and accurate quantification of the analysis. All glassware and sodium sulfate were solvent rinsed and heated overnight at 450 °C before usage. Each batch of 10 samples included one procedural blank to check potential contamination. Method blanks were near or below the limit of detection (LOD), which was estimated at 100 ng/g for total SCCPs. The quantitative results were not blank corrected in this study. The recoveries of standards in spiked samples were within 80–105%. Repeatability of the GC–MS measurement was tested and relative standard deviation was in the range of 5–10% (n = 8).
3. Results and discussion

All concentrations were reported on a dry weight (dw) basis. The results indicated that the CP mixtures in sewage sludge were mainly composed by SCCPs. MCCPs could be detected in only some samples at low levels. Therefore in this work, only the concentrations of SCCPs in sewage sludge were presented and discussed.

3.1. SCCPs levels in sewage sludge

The results showed that SCCPs could be detected in all sludge samples. The concentrations for the different carbon chain homologues (C_{10-13}) and total SCCPs ($\Sigma$SCCPs) are summarized in Table SI-2. The levels of $\Sigma$SCCPs in sewage sludge ranged from 0.80 to 52.7 µg/g, with the average concentration at 10.7 µg/g, which are much higher than other contaminants such as polychlorinated biphenyls (PCBs) and organochlorine pesticides (OCPs) in our previous studies (Wang et al., 2007). The highest concentrations of $\Sigma$SCCPs were found in two municipal WWTPs located at Xi'an national high-tech development zone in Shanxi Province, which were up to 52.7 and 42.8 µg/g, respectively. High levels of SCCPs found in this district might originate from several sources. Based on the background information of the two WWTPs (Table SI-2), we found about half of the influents came from industrial effluents. Several CP manufacturing plants were located in the industrial zone of this city. Other potential sources could be from the large number of high-tech enterprises associated with CPs in this area.

Among all the sampling sites, Shandong province should be paid more attention, since it represents one of the major production bases of CPs (Tang and Yao, 2005). In this study, 21 sewage sludge samples of WWTPs were collected from 10 different cities in Shandong (Table SI-1). There was a slight variation of SCCP concentrations among the 21 sewage sludge except two samples with high levels of SCCPs (WWTP No 21 and 26). The maximum concentration of SCCPs in the sludge from this province was 14.1 µg/g in Jining City and the minimum was 0.8 µg/g in Dezhou City (Table SI-2). Generally, no significant differences of SCCP levels can be found in sludge collected from different WWTPs in the same city. Comparing to other provinces, the samples from Shandong Province did not show high levels of SCCPs although it is the main manufacturing base of CPs in China. On the contrary, it is noticeable that, the sludge samples in other districts, such as Beijing, where almost no CP manufacturing plants exist, contained relatively higher concentrations of SCCPs. This phenomenon suggests that release of CPs into the environment during CP production might not be a main source for SCCPs in sewage sludge of WWTPs. No clear-cut geographic distribution characteristics were observed among the sampling sites, but samples from densely populated area or/high-tech industrial zone generally exhibited relatively higher SCCP levels.

Fig. 2 illustrates the frequency distributions of concentrations for five dominating homologues (C_{10}, C_{11}, C_{12}, C_{17}, and C_{18}) and $\Sigma$SCCPs in sewage sludge. The concentrations of C_{10} ranged from 0.20 to 13.9 µg/g (mean 2.80 µg/g). 96% of C_{10} concentrations were in the range of 0.20--6.00 µg/g. The C_{11} homologue was found to be the most abundant carbon homologue in sludge samples. The concentrations of C_{11} ranged from 0.30 to 15.8 µg/g (mean 3.80 µg/g). 94% of C_{11} concentrations were in the range of 0.30--8.00 µg/g, and 6% of them were higher than 10.0 µg/g. The concentrations of C_{12} ranged from 0.20 to 12.8 µg/g (mean 2.40 µg/g), while 90% of C_{12} concentrations were centered in the range of 0.20--4.00 µg/g. Moreover, C_{17} and C_{18} were the predominant chlorine homologues, which were in the concentration range of 0.30--17.3 µg/g and 0.20--18.1 µg/g, respectively. 92% of C_{17} concentrations were in the range of 0.30--8.00 µg/g, with 40% of them higher than 4.00 µg/g. Similarly, 96% of C_{18} concentrations were in the range of 0.20--8.00 µg/g, but only 27% of them were higher than 4.00 µg/g. As shown in Fig. 2, more than 90% of $\Sigma$SCCPs were in the range of 0.80--20.0 µg/g. The distribution of concentrations for $\Sigma$SCCPs in sewage sludge samples were also summarized as medians, quartiles and ranges with box-whisker plots (Fig. SI-1). The statistical results clearly indicated the concentrations of $\Sigma$SCCPs in 75% samples...
were lower than 12.5 μg/g, and about 20% of the samples were in
the range of 12.5–22.5 μg/g. Only 5% of samples showed SCCP
concentrations higher than 22.5 μg/g. These results implied that
concentrations lower than 20.0 μg/g could be representative of the
typical SCCP levels in most sewage sludge from WWTPs in China.

Compared to other POPs, such as polybrominated diphenyl
ethers (PBDEs) and PCBs (Clarke et al., 2008; Kupper et al., 2008;
Diaz-Cruz et al., 2009; Guo et al., 2009), information on SCCP
concentrations in sewage sludge are very scarce. In U.K., a survey
of the digested sludge from 14 WWTPs revealed that SCCP
concentrations ranged from 7 to 200 μg/g dw (Stevens et al.,
2003), and the range of concentrations is consistent with those
previous measured in sewage sludge (1.8–93 μg/g dw) from
England and Wales (Nicholls et al., 2001). Our work indicated that
SCCP concentrations in sewage sludge in China were comparable
to the reported data, but the average levels were lower than those
measured in the U.K.

3.2. Congener distribution patterns in sewage sludge

The SCCP congener distribution patterns of sewage sludge
samples from 52 WWTPs in different cities were also analyzed.
Fig. 3 shows both the carbon homologue composition and chlorine
homologue composition profiles. In general, the congener composi-
tions in all sewage sludge were similar among different WWTPs,
although some of them are located in different districts and receive
different types of sewage sources (domestic, industrial, or
combined) (Table SI-1). Composition analysis indicated that the
most abundant group of SCCPs in sludge samples was C11, which
accounted in average for 34% of SCCPs. Other abundant carbon
homologues were C10 and C12, which accounted for 27% and 23%
of total SCCPs, respectively. The concentration distributions for four
carbon chain homologues by box-whisker plots (Fig. SI-1a) also
indicated that their median concentrations were in the following
order: C11 > C10 ≈ C12 > C13.

The chlorine homologue composition profiles in most of sludge
samples also showed similar distribution pattern (Fig. 3). The Cl7
(mean contribution of 37%) and Cl8 (29%) were the most dominant
congener groups in all samples except one sample from Ningbo city
(No 32), which mainly receives dyeing wastewater discharged from
textile industry. Other abundant chlorine homologues were Cl6 and
Cl9, which accounted for 16% and 10% of total SCCPs, respectively.
Both the low chlorinated congener Cl5 and the high chlorinated conge-
ners C10 accounted for less than 10% of total SCCPs. The homologue
distribution pattern in sewage sludge were consistent with the
composition of commercial standards and technical SCCP mixtures
(Reth and Oehme, 2004). The concentration distributions for six
chlorine atom homologues in Fig. SI-1b indicated that their median
concentrations were in the order of Cl7 > Cl8 > Cl6 > Cl9 >
Cl5 > Cl10. The main congeners Cl7 and Cl8 showed obviously higher
mean concentrations than other chlorine atom homologues. To our
knowledge, this is also the first study on the distributions of
different homologues in sewage sludge.

Figs. 4 and 5 shows the representative congener group abun-
dance profiles categorized by carbon and chlorine atom numbers in
four sludge samples, which were collected from Beijing (No. 3),
Shandong (No. 17), Guangdong (No. 45) and Shanxi Province (No.
46). These four sampling sites respectively represent locations from
the northern, eastern, southern and eastern parts of China. Among
the four provinces, Beijing and Guangdong are characterized by
densely population and high industrialization, while Shandong
is a major production base of CPs in China. As shown in the carbon
congener group abundance profiles (Fig. 4), the composition
distributions of congeners were obviously predominated by

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**Fig. 3.** Composition profiles of carbon homologues (C10–C13) and chlorine homologue (C15–C10) for total 52 sewage sludge samples in China.
Fig. 4. The representative carbon congener group abundance profiles of SCCPs for four sludge samples collected from Beijing (No.3), Shandong (No.17), Guangdong (No.45) and Shanxi Province (No.46), respectively.

Fig. 5. The representative chlorine congener group abundance profiles of SCCPs for four sludge samples collected from Beijing (No.3), Shandong (No.17), Guangdong (No.45) and Shanxi Province (No.46), respectively.
medium chlorinated congeners (Cl3–8) within the individual carbon chain groups (C10, C11, C12, and C13). Furthermore, this congener distribution pattern was more clearly reflected in the chlorine congener group abundance profiles (Fig. 5). These specific congener distributions of SCCPs in sewage sludge may be influenced by the difference in composition of technical CP mixtures produced and used in China.

3.3. Correlation analysis

The relationships among different SCCP congener groups were evaluated using linear regression analysis. Significant linear relationships were found among different carbon chain groups (C10, C11, C12, and C13). Besides, significant linear relationships were also found among different chlorine congener groups. The correlation coefficient (R²) among these homologues varied from 0.90 to 0.97 (p < 0.05). Furthermore, in order to examine the potential factors that affect the SCCP levels in sewage sludge, principal component analysis (PCA) was executed using SPSS 16.0. The results indicated that affect the SCCP levels in sewage sludge, principal component analysis and environmental occurrence. Environ. Int. 32, 915–924.

References


