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Observation of inconsistent carbon isotope compositions of chlorine-isotopologue pairs of individual organochlorines on gas chromatography-high resolution mass spectrometry

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ABSTRACT

Relative abundances of chlorine and carbon isotopologues of individual organochlorines (e.g. polychlorinated organic pollutants) are generally recognised to comply with stochastic distribution, and thus can be calculated by the binomial theorem. As a result, carbon isotope ratios derived from different chlorine-isotopologue pairs, e.g. $^{12}\text{C}_2^{35}\text{Cl}_4$ vs. $^{12}\text{C}^{13}\text{C}^{35}\text{Cl}_4$, should be consistent. Yet this recognition may not exactly reflect the reality. This study investigated the consistency/inconsistency of carbon isotope compositions of chlorine-isotopologue pairs of individual organochlorines including two chloroethylenes, three polychlorinated biphenyls, methyl-triclosan and hexachlorobenzene. The raw carbon isotope ratios were measured by gas chromatography-high resolution mass spectrometry. Data simulations in terms of background subtraction, background addition, dual $^{13}\text{C}$-atoms substitution, deuterium substitution and hydrogen-transfer, along with measurements at different injection concentrations were conducted to confirm the validity of measured carbon isotope ratios and their differences. Inconsistent carbon isotope ratios derived from chlorine-isotopologue pairs of individual organochlorines were observed, and the isotopologues of each organochlorine were thus inferred to be non-randomly distributed. Mechanistic interpretation for these findings was tentatively proposed according to a basic principle in clumped-isotope geochemistry, reaction thermodynamics and kinetics, along with isotope effects occurring in electron ionisation mass spectrometry. This study sheds light on the actual carbon isotope compositions of chlorine-isotopologue pairs of organochlorines, and yields new insights into the real distributions of carbon and chlorine isotopologues. The inconsistent carbon isotope compositions of chlorine-isotopologue pairs are anticipated to benefit the exploration of formation conditions and source identification of organochlorine pollutants.

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1. Introduction

Organochlorines, produced by both anthropogenic and natural activities [1–3], have been greatly impacting human beings and the natural environment, either positively or negatively [4,5]. Numerous organochlorines are notorious environmental pollutants, such as highly toxic polychlorinated dioxins/furans, polychlorinated biphenyls (PCBs) and di(chloro)phenyltrichloroethanes [6–8]. Conceptually, organochlorines contain at least two elements, i.e. carbon and chlorine. Both carbon and chlorine have two natural stable isotopes ($^{12}\text{C}$ vs. $^{13}\text{C}$ and $^{35}\text{Cl}$ vs. $^{37}\text{Cl}$) with certain isotope ratios in the nature, leading to characteristic isotopologue distributions of organochlorines [9]. However, little is known about the exact distributions of carbon and chlorine isotopologues of organochlorines presently.

Historically, relative abundances of carbon and chlorine isotopologues of individual organochlorines are considered to be randomly distributed, which means the relative abundances can be calculated with the binomial theorem [10,11]. In this context, if the bulk carbon/chlorine isotope ratios of an organochlorine from different sources are identical, then the relative abundances of individual isotopologues of the organochlorine from these sources are identical. However, as reported in a large number of studies involving ‘clumped isotopes’, the relative abundances of multiply-substituted isotopologues of some simple compounds such as carbon dioxide, oxygen, methane and nitrogen are not stochastically distributed both in principle and in practice [12–17]. It can be speculated that the chlorine isotopologues containing more than one $^{37}\text{Cl}$ atom are analogous to the multiply-substituted isotopologues in clumped-isotope geochemistry, if carbon isotopes are not taken into account. Therefore, the chlorine isotopologues may not comply with binomial distribution (stochastic distribution) theoretically. As a consequence, the carbon isotope ratio ($^{13}\text{C}/^{12}\text{C}$) derived from a chlorine-isotopologue pair (CIP), of which the two isotopologues contain 0 and 1 $^{13}\text{C}$ atom, respectively and have the same number of $^{35}\text{Cl}$ and $^{37}\text{Cl}$ atom(s) (e.g. $^{12}\text{C}_2^{35}\text{Cl}_4$ vs. $^{12}\text{C}_2^{13}\text{C}^{35}\text{Cl}_4$), may not exactly equal those of other CIPs (e.g. $^{12}\text{C}_2^{35}\text{Cl}_3^{37}\text{Cl}$ vs. $^{12}\text{C}_2^{13}\text{C}^{35}\text{Cl}_3^{37}\text{Cl}$). Yet the potentially varied carbon isotope compositions of CIPs of individual organochlorines have not been reported. In addition, revelation of the inconsistent carbon isotope compositions of CIPs may be useful to unravel the formation conditions of organochlorines and further track their sources, and thus of important significance in source identification and apportionment for organochlorine pollutants.

In this study, we used gas chromatography-high resolution mass spectrometry (GC-HRMS) to investigate the carbon isotope ratios derived from CIPs of individual organochlorines including two chloroethylenes, three PCBs, methyltriclosan (Me-TCS) and hexachlorobezene (HCB). Differences among the carbon isotope ratios of CIPs were confirmed and evaluated, and mechanism explanation for the differences was proposed. This study reveals the actual carbon isotope ratios of CIPs of organochlorines, and gains new understanding towards the real distributions of carbon and chlorine isotopologues.

2. Experimental

2.1. Chemicals and materials

Standards of trichloroethylene (TCE, purity ≥ 99.0%) and perchlorethylene (PCE, purity ≥ 99.0%) were purchased from Tianjin Fuyu Chemical Co. Ltd. (Tianjin, China). The
mixed standard solution of polychlorinated biphenyls containing PCB-18, PCB-28 and PCB-52 (10.0 μg/mL in isooctane) was bought from Accustandard Inc. (New Haven, USA). Methyl-triclosan (Me-TCS, purity: 99.5%) and hexachlorobenzene (HCB, purity: 99.5%) were bought from Dr. Ehrenstorfer (Augsburg, Germany). Full names, abbreviations, structures, and other information of the chemicals are documented in Table S1. Isooctane and n-hexane were of chromatographic grade and obtained from CNW Technologies GmbH (Düsseldorf, Germany) and Merck Corp. (Darmstadt, Germany), respectively. Reference standard perfluorotributylamine for GC-HRMS calibration was bought from Sigma-Aldrich LLC. (St. Louis, MO, USA).

The standards of TCE, PCE, Me-TCS and HCB were weighed and subsequently dissolved with n-hexane, isooctane or nonane to obtain stock solutions at 1.0 mg/mL. These stock solutions and the purchased standard solution of PCBs were further serially diluted with n-hexane, isooctane or nonane to prepare working solutions at 1.0 or 5.0 μg/mL (Table S1); specifically, PCB-28 was investigated at both 1.0 and 2.0 μg/mL. All the standard solutions were kept at −20 °C before use.

2.2. Instrumental measurement

The GC-HRMS system consisted of dual gas chromatographs (Trace-GC-Ultra) coupled with a double focusing magnetic-sector high resolution mass spectrometer and a Triplus auto-sampler (GC-DFS-HRMS, Thermo-Fisher Scientific, Bremen, Germany). The system control and data acquisition were performed with Xcalibur 2.0 (Thermo-Fisher Scientific). A capillary GC column (DB-5MS, 60 m × 0.25 mm, 0.25 μm thickness, J&W Scientific, Folsom, CA, USA) was utilised, and helium was used as the carrier gas with a constant flow rate at 1.0 mL/min. The GC temperature programmes are detailed in Table S1. The GC inlet and transfer line were set at 260 °C and 280 °C, respectively. The injection volume was 1.0 μL, and splitless injection mode was used.

The working parameters and conditions of the mass spectrometer are documented as follows: electron ionisation (EI) source in positive mode was used; EI energy was 45 eV; ionisation source was kept at 250 °C; filament current of the EI source was 0.8 mA; multiple ion detection (MID) mode was applied to data acquisition; dwell time of each isotopologue was around 20 ms; mass resolution was ≥ 10,000 (5% peak-valley definition) and the MS detection accuracy was within ±0.001 u. The mass spectrometer was real-time calibrated with perfluorotributylamine during MID operation. As EI energy around 45 eV is commonly used on GC-HRMS to achieve relatively high signal intensities for molecular ions of compounds [18], therefore the EI energy at 45 eV was applied to the implementation of this study.

Chemical structures of the investigated compounds were drawn with ChemDraw (Ultra 7.0, Cambridgesoft), and the exact masses of isotopologues were calculated with mass accuracy of 0.00001 u. For isotopologues with too low relative abundances to be displayed by ChemDraw, their exact masses were manually calculated by replacing the exact relative atomic masses of the light isotopes (35Cl and 12C) with those of the heavy ones (37Cl and 13C) on the isotopologues whose relative abundances can be displayed by ChemDraw.

Only the chlorine isotopologues containing 0 or 1 13C atom were taken into account. For a compound with n Cl atoms, all the chlorine isotopologues (n + 1) containing no 13C atom were selected. In addition, all the chlorine isotopologues containing one 13C atom were chosen except for TCE and PCE, of which only the first three were chosen (number of 37Cl...
atom(s): 0–2), due to the very low relative abundances of the last one isotopologue of TCE and the last two isotopologues of PCE. The mass-to-charge ratio \((m/z)\) of each isotopologue ion was obtained through subtracting the mass of an electron from that of the corresponding isotopologue. The \(m/z\) values were imported into the MID module for setting up monitoring method. The detailed data including retention times, isotopologue formulas, exact masses and exact \(m/z\) values of the investigated compounds are provided in Table S2, and the representative chromatograms along with mass spectra are shown in Figure 1. The working solutions were directly injected onto the GC-HRMS with six analysis replicates.

2.3. Data processing

The comprehensive carbon isotope ratio (IR) derived from all measured isotopologues of each compound was calculated by

\[
IR = \frac{\sum_{i=0}^{n} l_{bi}}{\sum_{i=0}^{n} [m \times l_{ai} + (m - 1) \times l_{bi}]}
\]

where \(n\) is the number of Cl atoms of an organochlorine; \(i\) represents the number of \(^{37}\)Cl atoms in an isotopologue; \(l_{ai}\) denotes the MS signal intensity of the isotopologue \(i\) containing no \(^{13}\)C atom; \(l_{bi}\) represents the signal intensity of the isotopologue \(i\) containing one \(^{13}\)C atom; \(m\) is the number of C atoms of the organochlorine. This calculation scheme originates from a previously reported scheme for calculating chlorine/bromine isotope ratios \([19,20]\). For TCE and PCE, because only the first three chlorine isotopologues containing one \(^{13}\)C atom were monitored, the calculation scheme of the comprehensive carbon isotope ratio was thus altered to

\[
IR = \frac{\sum_{i=0}^{2} l_{bi}}{\sum_{i=0}^{2} (2l_{ai} + l_{bi})}
\]

The carbon isotope ratio derived from each CIP (IR\(_i\)) was calculated with

\[
IR_i = \frac{l_{bi}}{m \times l_{ai} + (m - 1) \times l_{bi}}
\]

The definition of CIPs is illustrated in Figure 2 with a simulated mass spectrum of a hypothesised organochlorine, and the details of CIPs of all the investigated compounds are provided in Table S2. The average MS signal intensity of each isotopologue derived from each whole chromatographic peak was used for calculating isotope ratios. Background subtraction was carried out prior to exporting MS signal intensities by subtracting baseline intensities neighbouring both ends of the corresponding peak. Data from replicated measurements were applied to calculating the mean isotope ratios and standard deviations (1σ). As isotopic standards with known carbon isotope ratios of the investigated compounds were unavailable in this study, all the measured isotope ratios were raw values without being calibrated to the VPDB (Vienna Pee Dee Belemnite).
standard carbon scale. Due to that the carbon isotope ratios derived from different CIPs of individual organochlorines were synchronously measured, these isotope ratios can be

**Figure 1.** Representative chromatograms and high resolution mass spectra of the investigated organochlorines. TCE: trichloroethylene, PCE: tetrachloroethylene, PCB: polychlorinated biphenyl, Me-TCS: methyl-triclosan, HCB: hexachlorobenzene, NL: nominal level, m/z: mass to charge ratio.
applied to ascertaining and evaluating the isotope ratio differences among CIPs, despite the measured isotope ratios were not calibrated to the international scale.

2.4. Data simulations for confirming the validity of measured isotope ratios

Theoretical abundances of isotopologues were simulated by the isotope modelling program embedded in MassLynx V4.1 (Waters Corp., Manchester, UK). The simulation was based on the binomial theorem. The simulated abundances were calibrated with the measured comprehensive carbon isotope ratios with the processes detailed in the Supporting Information. In this way, the simulated comprehensive carbon isotope ratios derived from the calibrated abundances of isotopologues equalled the measured. The calibrated abundances were then subjected to simulated background subtraction and addition. The simulated abundances, calibrated abundances, and calibrated abundances with simulated background subtraction/addition are provided in Table S3.

Substitution of two $^{13}$C atoms, deuterium substitution, and hydrogen-transfer (H-transfer) might influence the measured isotope ratios and thereby the observed isotope ratio differences among CIPs. We conducted the data simulations in terms of dual $^{13}$C-atoms substitution, deuterium substitution and H-transfer to obtain simulated isotope ratios along with isotope ratio differences for confirming the validity of measured isotope ratios.
2.5. **Statistical analysis**

Statistical analysis was carried out with SPSS Statistics 19.0 (IBM Inc., Armonk, USA). Paired-samples T test and independent-samples T test were applied to evaluating differences between two groups of carbon isotope ratios or two isotope ratios with p-values (2-tailed) of 0.01 as the cut-off for significance. If a p-value is under 0.01, the null hypothesis (e.g. no difference between two groups of carbon isotope ratios) is declined, demonstrating a significant difference indeed present.

3. **Results and discussion**

3.1. **Validity of measured carbon isotope ratios**

3.1.1. **Impact of background subtraction**

To confirm the validity of measured carbon isotope ratios, we conducted some data processing procedures to examine the possible influences of instrumental uncertainties and artificial errors. We speculated that the background subtraction of MS signal intensities might negatively affect the measured carbon isotope ratios, because it might take away slight real signals of target ions. Therefore, we acquired the carbon isotope ratios with and without background subtraction, and compared them for examining the influence. In addition, we conducted data simulation for the background subtraction, in order to further confirm the validity of measured isotope ratios. The measured carbon isotope ratios with/without background subtraction and the simulated carbon isotope ratios are provided in Table S4.

As for PCB-28, Me-TCS and PCB-52, the measured isotope ratios with and without background subtraction presented increasing discrepancies from the first to the last CIPs (Figure 3(a–c)), and the isotope ratios with background subtraction were significantly lower than those without background subtraction for CIP-4 of individual compounds with the discrepancies ranging from $-0.00111 \pm 0.00016$ to $-0.00075 \pm 0.00007$ ($p \leq 0.00003$, Table S5). The measured isotope ratios with background subtraction were well consistent with those without background subtraction for individual CIPs of HCB (Figure 3(d)). The simulated isotope ratios without background subtraction (i.e. simulated theoretical isotope ratios) of all the CIPs of each compound were consistent in theory. The simulated isotope ratios with background subtraction successively decreased from the first CIPs to the last, whereas those with background addition continually increased from the first to the last (Figure 3). The three types of simulated isotope ratios of the first CIP were very close for each compound, and close to the corresponding comprehensive isotope ratio. Whereas the simulated isotope ratios with background subtraction/addition of the last CIPs of PCB-18, Me-TCS and PCB-52 and the last two CIPs of HCB showed apparently large deviations from the simulated theoretical isotope ratios. If the detected isotopologues were stochastically distributed and the background subtraction indeed caused signal loss for target ions, then the measured isotope ratios with background subtraction ought to match the simulated isotope ratios with background subtraction. On the other hand, the measured isotope ratios without background subtraction were supposed to match the simulated isotope ratios with background addition. In practice, however, the patterns of the measured carbon isotope ratios with/without background subtraction were definitely different from those of the simulated isotope ratios with background subtraction or
addition (Figure 3). Therefore, the differences of measured carbon isotope ratios derived from CIPs were determined to be not caused by instrumental uncertainties and artificial errors but really existent.

3.1.2. Influence from the substitution of two $^{13}$C atoms, deuterium substitution and H-transfer

In addition to background subtraction, substitution of two $^{13}$C atoms, deuterium substitution, and H-transfer in EI-MS might affect the measured carbon isotope ratios, thus needing scrutiny and assessment. To this end, we further performed some data simulations to evaluate these potential impacts. We chose PCE as a model organochlorine to carry out the simulations of substitution of two $^{13}$C atoms and H-transfer, in consideration of that PCE comprises merely two elements (i.e. C and Cl). In addition, the simulation of...
deuterium substitution was implemented with TCE, due to that TCE contains only one H atom in the formula.

As documented in Table S6, the theoretical carbon isotope ratios of the first three CIPs of PCE corrected with the substitution of two $^{13}$C atoms were 0.010997–0.011000, with the differences of $0.000001–0.000003$. While the corresponding measured isotope ratio differences were from $0.00040 \pm 0.00009$ to $0.00095 \pm 0.00009$ ($p \leq 0.00012$), which were over two orders of magnitude higher than the simulated. This result indicates that the substitution of two $^{13}$C atoms can only negligibly affect the isotope ratio differences. On the other hand, the mass difference between $^{13}$C$_2$^{35}$Cl_4$ and $^{12}$C$_2$^{35}$Cl_3^{37}$Cl is 0.00966 u, which can be separated at a MS resolution of 17,172. Although the MS resolution applied in this study (around 10,000) was less than 17,172, partial separation between the two ions could be achieved, further reducing the impact of dual $^{13}$C-atoms substitution on the measured carbon isotope ratios.

In another study, we found the H-transfer ratio of the molecular ion of $^{13}$C$_6$-HCB was around 0.002 [21]. Based on this finding, we set the H-transfer ratio of the molecular ion of PCE at 0.01 to conduct the H-transfer simulation, which is anticipated to trigger larger impacts on carbon isotope ratios in contrast to the observed low H-transfer ratio, thus rendering the simulation results more convicive. As shown in Table S7, the differences among the corrected theoretical carbon isotope ratios of the first three CIPs ranged from $−0.000001$ to $−0.000002$, which accounted for $−3.77\%$ to $−0.36\%$ of the measured isotope ratio differences and were within the analysis uncertainties (standard deviations: 0.00007–0.00009). Therefore, the isotope ratio differences caused by H-transfer were negligible and thereby could not impact the measured isotope ratio differences among CIPs. Furthermore, as provided in Table S7, the simulated theoretical carbon isotope ratios with H-transfer correction were 0.015866–0.015867 for the first three CIPs, which were extremely higher than the measured isotope ratios. This discrepancy demonstrates that the real H-transfer ratio was lower than 0.01, which further indicated the negligible influence of H-transfer on the observed isotope ratio differences.

As shown in Table S8, the theoretical carbon isotope ratios corrected with deuterium substitution of the first three CIPs of TCE were 0.011506–0.011507, and the simulated isotope ratio differences are $−0.000001–0.000001$, which were 2–3 orders of magnitude lower than corresponding measured values ($0.00022 \pm 0.00005–0.00053 \pm 0.00004$), indicating the ignorable effect of deuterium substitution on the observed isotope ratio differences.

### 3.1.3. Effect of abundance differences among isotopologues

Concentration effects on compound-specific carbon isotope analysis are widely recognised in the field of isotope ratio analysis [22]. Therefore, it is not surprising that different concentrations may result in different raw isotope ratios, which should be calibrated with isotopic standards before practical application. To investigate whether the observed inconsistent carbon isotope ratios of CIPs in this study were caused by different abundances of isotopologues, we additionally measured the carbon isotope ratios of CIPs of PCB-28 at 1 and 2 μg/mL. As indicated in Figure S1, both the detected and theoretically simulated relative abundances of CIP-1 and CIP-2 of PCB-28 were very close. Specifically, the theoretical abundance ratio of CIP-2 to CIP-1 of PCB-28 is 96%, while the concentration ratio between 1 and 2 μg/mL is 50%. Therefore, the difference in abundances between CIP-1 and CIP-2 of PCB-28 at a same concentration is much less than that of CIP-1 or CIP-2 between 1 and 2 μg/mL. If the observed carbon isotope ratio differences among CIPs in this study were
caused by different MS signal intensities (or in other words, different concentrations or abundances), then the isotope ratio difference between CIP-1 and CIP-2 of PCB-28 at each concentration should be less than that of CIP-1 or CIP-2 between 1 and 2 μg/mL. However, the measured carbon isotope ratios of individual CIPs of PCB-28 at 1 and 2 μg/mL exhibited no statistically significant difference (p ≥ 0.30, Figure 4). This result confirms that the measured different carbon isotope ratios of CIPs of individual organochlorines were not attributable to different abundances of isotopologues.

In conclusion, after the evaluation of possible impacts from several factors, the validity of the measured carbon isotope ratios was guaranteed, demonstrating that the observed variations of measured carbon isotope ratios of CIPs of each organochlorine were not artificial but real.

3.2. Measured carbon isotope ratios

Currently, no available study has demonstrated whether the carbon isotope ratios derived from different CIPs of individual organochlorines are exactly identical. If they are not identical, yet it is unclear whether the discrepancies can be measured by available analytical techniques. In this study, we used the scheme as expressed by Equation (3) to obtain the carbon isotope ratio of each CIP. As shown in Figure 5, Tables S-9 and S-10, the main measured carbon isotope ratios of CIPs of individual organochlorines could be differentiated, with the differences between two random CIPs several times higher than the standard deviations and the p-values less than 0.01. For instance, the carbon isotope ratios of the first three CIPs (CIP-1 to CIP-3) of PCE were 0.01150 ± 0.00005, 0.01094 ± 0.00003 and 0.01053 ± 0.00008, respectively and those for TCE were 0.01126 ± 0.00004, 0.01106 ± 0.00003 and 0.01074 ± 0.00002 (Table S9), which were significantly different from each other for individual compounds (p ≤ 0.00012, Table S10), with the range of differences from 0.00022 ± 0.00005 to 0.00095 ± 0.00009 (Table S10). For PCBs, Me-TCS and HCB, the carbon isotope ratios of the first CIPs were significantly higher than those of others for individual compounds (Figure 5(b–d)), with the discrepancies ranging from 0.00027 ± 0.00005 to 0.00219 ± 0.00029 (p ≤ 0.0014, Table S10). The CIP-2 and CIP-3 of PCB-18,

![Figure 4](image-url). Measured carbon isotope ratios derived from the CIPs of PCB-28 in two standard solutions at different concentrations (1.0 and 2.0 μg/mL in isooctane).
PCB-28 and Me-TCS showed insignificantly different carbon isotope ratios ($0.019 \leq p \leq 0.13$), and the four middle CIPs (CIP-2 to CIP-5) of HCB also presented insignificant isotope ratio differences ($0.016 \leq p \leq 0.93$), except that between the CIP-2 and CIP-3 ($p = 0.0028$). The carbon isotope ratios between CIP-3 and CIP-4 of PCB-18 and PCB-28 were insignificantly different ($p > 0.01$). The carbon isotope ratios of CIP-4 of PCB-18, PCB-28, Me-TCS and HCB, together with those of the last two CIPs of PCB-52 and HCB showed generally higher standard deviations ($0.00015–0.00029$) compared with others (Figure 5(b–d) and Table S9), possibly due to the relatively low abundances of corresponding isotopologues.

Based on the measured carbon isotope ratios, we conclude that not all the measured carbon isotope ratios derived from CIPs of individual organochlorines were exactly equivalent. To the contrary, many CIPs presented significantly different carbon isotope ratios. The observed significant isotope ratio differences support the conclusion that carbon and chlorine isotopologues of individual organochlorines are not stochastically distributed.

### 3.3. Theoretical derivation and mechanistic interpretation

In this study, we applied a basic principle in clumped-isotope geochemistry, reaction thermodynamics and kinetics, and theories relevant to isotope effects in EI-MS to the mechanistic interpretation for the observed inconsistent carbon isotope ratios of CIPs and non-randomly distributed isotopologues of organochlorines.
3.3.1. Interpretation based on the basic principle in clumped-isotope geochemistry

In clumped-isotope geochemistry, a basic principle is that the relative abundances of isotopologues of individual compounds (such as carbon dioxide, nitrogen, methane, oxygen and hydrogen) do not conform to stochastic distribution except that those compounds are generated at extremely high temperatures [12–17]. The theoretical explanation regarding the non-random distribution of isotopologues has been detailed in a previous review [13]. In the present study, the isotopologues containing more than one heavy isotope atom (\(^{37}\)Cl and/or \(^{13}\)C) are analogous to the multiply-substituted isotopologues in clumped-isotope geochemistry. Therefore the carbon/chlorine isotopologues of organochlorines are supposed to be non-binomially distributed if the organochlorines are not produced at extremely high temperatures, resulting in inconsistent carbon isotope ratios of CIPs. The observation of inconsistent carbon isotope ratios derived from CIPs of individual organochlorines in this study is a new evidence for the principle in clumped-isotope geochemistry. In addition, this finding demonstrates that the deviations between actual and theoretical (random) distributions of carbon/chlorine isotopologues of organochlorines are measurable by GC-HRMS.

3.3.2. Inference in light of reaction thermodynamics and kinetics, and isotope effects in EI-MS

The observed inconsistent carbon isotope ratios of CIPs of individual organochlorines in this study may ascribe to the chlorination reactions in synthesis. As derived in the Supporting Information, the carbon isotope ratios of CIPs of synthesised organochlorines are deduced to be inconsistent, no matter the chlorination reactions are thermodynamically or kinetically controlled.

In this study, the investigated organochlorines were analysed by GC-EI-HRMS. Isotope effects occurring in EI-MS can be applied to explaining the inconsistent carbon isotope ratios of CIPs of an organochlorine whose chlorine isotopologues are hypothesised to be binomially distributed. As indicated in a previous study, fragmentation in EI-MS can cause significant hydrogen and chlorine isotope effects [23,24]. Thus, dechlorination of organochlorines in EI-MS are anticipated to generate concurrent carbon and chlorine isotope effects. Due to the isotope effects in EI-MS, the carbon isotope ratios of CIPs of an organochlorine measured by EI-MS cannot be consistent, even though the chlorine isotopologues are postulated to comply with binomial distribution prior to fragmentation. The related theoretical derivation is detailed in the Supporting Information.

3.4. Application prospects

As revealed in clumped-isotope geochemistry, clumping isotope effects strongly correlate with reaction temperatures [13]. When a reaction takes place at a low temperature, the ratios of multiply-substituted isotopologues tend to deviate from stochastic ratios. To the contrary, if a reaction occurs at a high temperature, the ratios of multiply-substituted isotopologues are prone to close to stochastic ratios [13]. We thus conclude that the inconsistency extents of carbon isotope ratios of CIPs are related to chlorination temperatures. The lower the chlorination temperature is, the more inconsistent the carbon isotope ratios of CIPs are. Therefore, the patterns of carbon isotope ratios of CIPs may be able to probe the temperature conditions of chlorination reactions, and further illuminate...
the reaction mechanisms. On the other hand, since organochlorines from different sources may be synthesised at different temperatures, the carbon isotope ratios of CIPs have promising application prospects in source identification and apportionment for chlorinated organic pollutants. For instance, HCB synthesised by UV assisted chlorination of benzene may have a different pattern of carbon isotope ratios of CIPs from that of HCB synthesised by heating chlorination reaction of trichlorobenzene, due to the different temperatures in the two chlorination reactions.

3.5. In comparison with compound-specific carbon isotope analysis using gas chromatography-isotope ratio mass spectrometry

In the last two decades, compound-specific carbon isotope analysis (CSIA) using gas chromatography-isotope ratio mass spectrometry (GC-IRMS) has been increasingly applied to source identification and apportionment, and investigation of environmental behaviours and fate for organic pollutants [25,26]. In our laboratory, Zeng et al. recently developed and utilised CSIA methods by means of GC-IRMS to investigate environmental behaviours and fate of halogenated organic pollutants, such as PCBs and polybrominated diphenyl ethers (PBDEs) [27–30]. In these studies, the standard deviations of the measured $\delta^{13}C$ values were generally less than 0.5‰ ($\delta^{13}C = \frac{IR_{\text{sample}}}{IR_{\text{standard}}} - 1$, where $IR_{\text{sample}}$ and $IR_{\text{standard}}$ denote carbon isotope ratios of environmental samples and the Vienna Pee Dee Belemnite (V-PDB) reference standard, respectively) [27–30]. These precisions obtained with GC-IRMS (<0.5‰) were much better than those achieved in the present study using GC-HRMS (2.0‰-5.0‰). However, CSIA using GC-IRMS can only measure average (bulk) carbon isotope ratios of individual compounds, since analytes have to be converted to $CO_2$ before entering mass analyser. Therefore, CSIA results obtained with GC-IRMS cannot profile carbon isotope compositions among different isotopologue pairs of individual analytes. Nevertheless, in this study, our data attained with GC-HRMS can reveal and differentiate carbon isotope compositions among CIPs of individual organochlorine analytes, and thus providing more details of each analyte in contrast with CSIA data achieved by GC-IRMS.

4. Conclusions

In this study, we systematically investigated whether the carbon isotope ratios derived from CIPs were consistent using seven exemplary organochlorines. The carbon isotope ratios were determined by GC-HRMS with sufficient precisions for evaluating the isotope ratio discrepancies among CIPs. The experimental data were carefully processed, and the validity of measured carbon isotope ratios were confirmed by data simulations. Most of the measured carbon isotope ratios derived from CIPs were found to be significantly different, showing generally declining tendencies from the first to the last CIPs. The relative abundances of carbon and chlorine isotopologues of organochlorines were deduced to be non-randomly distributed, which well coincides with the principle in clumped-isotope geochemistry, reaction thermodynamics and kinetics, and isotope effects occurring in EI-MS. The carbon isotope compositions of CIPs are anticipated to be compound-specific and source-specific, and thus can be used as fingerprint features to trace sources of organochlorines in the future. The experimental methods and data processing approaches applied in this study can be extrapolated to isotopologues containing other elements such as bromine, sulphur
and silicon for revealing actual isotope compositions. The results of this study provide a prospection that the isotologues of brominated, sulphureted and siliciﬁed organic compounds are also non-stochastically distributed. Further studies is worthwhile in terms of applications of the inconsistent isotope ratios of CIPs to source identiﬁcation and apportionment for organochlorine pollutants.

Disclosure statement

No potential conﬂict of interest was reported by the authors.

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