Combining High-Resolution Gas Chromatographic Continuous Fraction Collection with Nuclear Magnetic Resonance Spectroscopy: Possibilities of Analyzing a Whole GC Chromatogram

Louise M. van Mourik,* Elwin Janssen, Robin Breeuwer, Willem Jonker, Jacco Koekkoek, Arif Arrahman, Jeroen Kool, and Pim E. G. Leonards

Cite This: Anal. Chem. 2021, 93, 6158–6168

ABSTRACT: This study presents, for the first time, the successful application of analyzing a whole gas chromatography (GC) chromatogram by nuclear magnetic resonance (NMR) spectroscopy using a continuous repeatable and stable (n = 280) high-resolution (HR) GC fractionation platform with a 96-well plate. Typically with GC− or liquid chromatography−mass spectrometry analysis, (isomer) standards and/or additional NMR analysis are needed to confirm the identification and/or structure of the analyte of interest. In the case of complex substances (e.g., UVCBs), isomer standards are often unavailable and NMR spectra too complex to achieve this. This proof of concept study shows that a HR GC fractionation collection platform was successfully applied to separate, purify, and enrich isomers in complex substances from a whole GC chromatogram, which would facilitate NMR analysis. As a model substance, a chlorinated paraffin (CP) mixture (>8,000 isomers) was chosen. NMR spectra were obtained from all 96 collected fractions, which provides important information for unravelling their full structure. As a proof of concept, a spectral interpretation of a few NMR spectra was made to assign sub-structures. More research is ongoing for the full characterization of CP isomers using multivariate statistical analysis. For the first time, up to only a few CP isomers per fraction were isolated from a highly complex mixture. These may be further purified and certified as standards, which are urgently needed, and can also be used for persistency, bioaccumulation, or toxicity studies.
chromatographic fractionation using GC can be superior to LC in terms of separation efficiency and analysis time.\textsuperscript{15,24} GC fractionation is less commonly applied largely due to its difficult setup. An even more complicated setup and thus less commonly applied is preparative GC in combination with NMR, an analyte purification and enrichment tool to elucidate the structure of organic compounds.\textsuperscript{14,17,18} Prior attempts and marketed GC fractionation systems required complex systems involving cryotrapping the analyte or trapping the analyte on a sorbent, which only allowed the collection of a small number of fractions.\textsuperscript{19–21} This is suitable when only a few target compounds are to be analyzed, however unsuitable for complex mixtures such as UVCBs.

Recently, a high-resolution GC fractionation platform was developed\textsuperscript{22–24} that was able to fractionate on 96–384 well plates and post-column off-line analyze all fractions collected from full GC chromatograms, by having, via a split, post-column parallel flame ionization detection (FID) or MS detection. Due to its high-resolution fractionation, capability of fractionating complete GC chromatograms, and the possibility to increase the yield of each fraction collected by straightforward automated multiple fraction collection on the same well plate (or in vials for fraction collection, which is also possible), this platform has been successfully applied to cell-based bioactivity analysis\textsuperscript{23} and may be a promising tool for enabling the analysis of whole GC chromatograms by NMR spectroscopy.

This proof of concept study demonstrates the use of the continuous GC fraction collection platform with repeatable and stable fractionations (n = 280) on a 96-well plate to purify and enrich isomers of complex substances. In addition, it shows the possibilities of analyzing a whole GC chromatogram by NMR using this platform to potentially identify or confirm their structures. Two other analytical methods [i.e., comprehensive GC × GC and quadrupole time of flight MS in the negative atmospheric pressure chemical ionization mode (APCI-QToF-MS)] were also applied to further explore the separation abilities (i.e., the purification and enrichment of isomers) of the GC fractionation platform. As a model substance, arguably one of the most complex organochlorine substances, CP–C\textsubscript{14} mixture with 41.3% chlorine (Cl) by weight, was chosen.\textsuperscript{25}

## Experimental Section

**Chemicals.** n-Hexane (pesticide residue analysis) was purchased from Fluka (Landsmeer, The Netherlands), chloroform-d, n-octane (>99%) from Sigma-Aldrich (Zwijndrecht, The Netherlands), and acetonitrile (99.8%) and iso-octane (+99%) from Thermo Fisher (Geel, Belgium). The polychlorinated n-alkane mixture tetradecane with an average chlorine content of 41.3% by weight (CP–C\textsubscript{14}) was manufactured by Quimica del Cinca (Barcelona, Spain) and kindly donated by the Chlorinated Paraffins Industry Association (Washington DC, United States).

**Instruments.** The GC fraction collector platform has been described in previous work\textsuperscript{22–24} and a schematic overview is presented in S1. Briefly, a PAL system (CTC Analytics AG, Zwingen, Switzerland) was used for sample injection on an Agilent HP 6890 GC oven, equipped with a split/splitless injection port (Agilent Technologies, Santa Clara, CA, USA) and an Agilent Ultra 2 (5%-phenyl)-methylpolysiloxane (25.0 m × 0.20 mm × 0.11 μm) column for separation. The column eluate was split toward a FID (Agilent Technologies, Santa Clara, CA, USA) for peak detection and an inverted y-piece where the preheated/vaporized trap solvent was infused. The trap solvent was delivered via a Shimadzu LC-10Ai pump and preheating was facilitated using a modified FID. At the inverted y-piece, eluting compounds mix with the vaporized trap solvent and are directed outside the GC oven via a capillary while the trap solvent condenses at ambient room temperature, enabling fraction collection. The capillary exit was connected to a smart grip unit (Da Vinci Laboratory Solutions, Rotterdam, The Netherlands) that was maneuvered over a 96-well plate for fraction collection. The instrument parameters were controlled with Agilent ChemStation 7.0 software, while the PAL system was controlled with Da Vinci Europe Smart PAL interface version 0.0.9.

The CP–C\textsubscript{14} mixture was injected (1 μL) with an injection temperature of 275 °C in the splitless mode with a 2 min split time. The purge flow was set to 50 mL/min for 2.0 min, and after that, the gas saver was switched on. The oven temperature started at 150 °C, increased by 10 °C/min to 250 °C, then 35 °C/min to 320 °C and at 320 °C ended with a holding time of 2 min. The helium carrier gas flow was set to 2.0 mL/min. The FID was set to 275 °C, with a hydrogen flow of 40 mL/min, an air flow of 400 mL/min, and a nitrogen makeup flow of 45 mL/min. The solvent’s flow rate was set to 0.3 mL/min, and was preheated to 250 °C before being introduced to the y-piece. Fraction collection started 1.0 min after injection in 96 700 μL glass inserts located on a 96-well plate (Waters, US) with a 7.0 s interval per fraction and was performed in a serpentine motion: A1–A12 followed by B12–B1 and so forth to H1 (graphical abstract). After every fractionation trap, the solvent was evaporated and discarded. Injection of the CP–C\textsubscript{14} mixture was repeated 280 times, with a total run time of 65 h. Two types of solvents were tested for their applicability as the trap solvent: n-hexane and n-octane. For GC analysis, 200 μL of each fraction was transferred to a vial containing 800 μL of n-octane. For NMR analysis, each fraction was gently blown down with nitrogen, re-dissolved in 0.6 mL of chloroform-d and transferred to a NMR glass tube. After NMR analysis, chloroform-d was gently evaporated in air and the tube was refilled with ca. 2.5 mL of pentane, of which 40 μL was taken out, blown down, and re-dissolved in acetoni-trile for MS analysis.

NMR spectra were recorded on a Bruker Avance II 500 (Fällanden, Switzerland) at a 1H frequency of 500.20 MHz and equipped with a 5 mm inverse triple resonance (1H, 13C, and 15N) cryoprobe with an active shield (Z gradient and a Bruker Avance III HD 600 at a 1H frequency of 600.13 MHz and equipped with a 5 mm inverse triple resonance (1H, 13C, and 15N) cryoprobe with an active shield Z gradient. The residual solvent peak was used as an internal standard (1H: δ 7.26 ppm for chloroform). A standard proton experiment (zz30) was used at 25 °C. Data were collected with 65k time domain points and 20 ppm spectral width. The frequency offset was set to 6.175 ppm. For each 1H NMR spectrum, 1024 transients were collected with a recycle delay of 1 s. Spectra were processed with an exponential multiplication equivalent of 0.3 Hz line broadening.

As comprehensive GC × GC is one of the few approaches known for its ability to separate MCCPs to individual components,\textsuperscript{26} it is a promising tool to evaluate the fractionation process. All fractions, as well as the CP–C\textsubscript{14} mixture itself, were injected on the GC × GC–µECD system. Single blobs in the obtained chromatograms could possibly indicate that fractionation to individual isomers was achieved. Each fraction and the CP–C\textsubscript{14} mixture were analyzed using the GC × GC system, consisting of an Agilent HP7890 gas chromatograph (Palo Alto,
CA, USA) equipped with μECD of which instrument settings and data analysis are reported in previous work.26

All fractions (and the CP−C14 mixture) were also analyzed on a (APCI-QToF-MS) by direct injection using high-performance liquid chromatography (HPLC). The APCI-QToF-MS method is described elsewhere.27 In brief, the APCI-QToF-MS consisted of a Bruker Compact (Bremen, Germany) and a HPLC system Agilent 1290 infinity on which 5 μL was injected and acetonitrile

Figure 1. Schematic overview and workflow of the continuous high-resolution GC fractionation platform in combination with NMR spectroscopy. First, the continuous high-resolution GC fractionation platform is applied to purify and enrich isomers of complex substances by multiple (n = 280) repeatable and stable fractionations on a single 96-well plate. The solvent of each fraction is then evaporated, after which the fraction is re-dissolved in chloroform-d for NMR spectroscopy. To evaluate the fractionation process and to understand the NMR data, each fraction is also analyzed by GC × GC and (LC)-APCI-TOF-MS.

Figure 2. GC × GC-μECD chromatogram (panel A) and single GC chromatogram (horizontal slice at 3.4 min on the second dimension, panel B) of CP−C14 41.3% Cl mixture of which the black squares illustrate the overlap of the congener group C14Cl6 with C14Cl5 and C14Cl7.
with 10% dichloromethane (v/v) was used as an eluent with an isocratic flow of 250 μL/min. The 10% dichloromethane solution was used as the dopant to enhance the formation of [M + Cl]⁻ fragments of CPs. For integration, strict isotope pattern criteria were applied. The m/z ratios for CP−C₁₄ applied in this study can be found in Table S1, which are related to the two most abundant [M + Cl]⁻ ions of the CP isotope cluster corresponding to the congener groups (CPs with the same molecular formula), expressed as CₙClₙ₋₃ to C₁₄Cl₁₂.²⁷

### RESULTS AND DISCUSSION

The continuous high-resolution GC fraction collection platform was applied to analyze a whole GC chromatogram by NMR spectroscopy as a proof-of-concept study. Other analytical methods such as GC × GC and APCI-QToF-MS were applied to investigate the separation possibilities of the platform further and as a tool to understand the NMR data (Figure 1).

**High-Resolution GC Fraction Analysis by GC-FID.**

Because the injection volume of the GC fractionation platform was limited to 1 μL and taking into account the relative low sensitivity of (proton) NMR, some test injections on a similar GC system with different CP concentrations (0.1, 1, and 5 mg/mL in isooctane) were performed to find out the highest concentration that did not compromise the resolution. Although the signal intensity differs, the characteristic peaks are similar between the concentration of 0.1 and 1 mg/mL (Figure S1A,B). However, with a concentration of 5 mg/mL, there was an obvious loss of resolution (Figure S1C). Based on these results,
the choice was made to proceed with the 1 mg/mL concentrated solution.

Two types of solvents (n-hexane and n-octane) were tested for their applicability as the trap solvent. Although n-octane has superior trapping efficiency compared to n-hexane, also discussed in previous work, n-hexane was chosen because of its higher volatility: for the NMR measurements, the trap solvent needs to be blown down to dryness and the collected analytes re-dissolved in chloroform-d. The volatility of n-octane is so low that there is a risk of losing MCCPs during evaporation. Furthermore, when using n-octane, the well plates need to be switched regularly to prevent overflowing, while n-hexane is volatile enough to perform a continuous fractionation without the risk of flooding the glass inserts in the well plate.

To ascertain that no retention time shift occurred during fractionation, which could cause mixing of different fractions, frequent chromatogram comparisons were performed. No retention time shift was observed when using n-hexane as the

Figure 4. Highlighted chlorine-optimized APCI-TOF-MS spectra of CP–C_{14} 41.3% Cl mixture (panel A), as well as fractions E10 (fractionation time 466 s, panel B) and F11 (494 s, panel C). In the spectra of the CP–C_{14} 41.3% Cl mixture [M + Cl]^{-} of congener groups, expressed as C_{m}Cl_{n}, C_{14}Cl_{12} can be observed, with m/z values of the most abundant ion per congener group 337.0844, 371.0454, 405.0064, 438.9674, 474.9255, 508.8866, 542.8476, 576.8086, 612.7667, and 646.7277, which are according to previous work. In the spectra of fraction E10, [M + Cl]^{-} of congener groups C_{14}Cl_{4} (m/z value 371.0454) and C_{14}Cl_{5} (m/z value 405.0064) can be observed, while in F11, C_{14}Cl_{4} (m/z value 371.0454) can be observed. The full m/z ratio list can be found in Table S1.
Figure 5. Highlighted $^1$H NMR spectra of CP–C$_{14}$ 41.3% Cl mixture and fractions (B) E3 (fractionation time 417 s), (C,D) E6–E7 (438–445 s), and (D) E10 (466 s), with highlighted areas of peaks that represent certain fragments of CP–C$_{14}$ (see panel A), of which the gray-colored areas represent the water peak (1.54 ppm) and chloroform (7–7.5 ppm). Full-sized spectra can be found in S5.
Figure 6. Highlighted $^1$H NMR spectra of fractions A9 (fractionation time 123 s), B4 (207 s), C9 (291 s), D4 (365 s), E9 (459 s), F4 (543 s), G9 (627 s), and H4 (711 s). Highlighted areas of peaks represent certain fragments of CP–C$_{14}$ (see legend), while the gray-colored areas represent the water peak (1.54 ppm) and chloroform (7–7.5 ppm).
High-Resolution GC Fraction Analysis by GC × GC–ECD. To evaluate the fractionation process, the whole C14 mixture and each fraction were analyzed by GC × GC–ECD. The complexity of the mixture and the power of GC × GC separation are clearly shown in the GC × GC chromatogram of CP−C14 mixture (Figures 2 and 3). Single dots are present on the GC × GC chromatogram, which could indicate single CP isomer separation. The dots elute consecutively as diagonal lines, representing the CP congener groups (i.e., isomers with the same molecular formula, expressed as CnClm in Figure 3A), according to previous work.28 The power of GC × GC separation is especially shown when comparing the GC × GC and the single chromatogram (Figure 2) for the congener groups C14Clk−7. For example, even with an extremely long oven temperature program (90 min), single GC cannot distinguish C14Cl9 from C14Cl10 and C14Cl12, resulting in overlap on 50−53 and 57−61 min in the single GC chromatogram (black squares in Figure 2B), while GC × GC can, resulting in different blobs vertically above each other around the same time (Figure 2A). This is very interesting as the Clk and Cls congener groups are usually the most dominant in environmental samples.28 Congener groups C14Cl3 to C14Cl12 were observed, while the hump of peaks around the retention time 20 min (Figure 2A) might indicate C14Cl2 presence. Thus far, it remains uncertain whether C14Cl1 is present in technical mixtures as these do not ionize and/or their response is too low to be detected by the current analytical methods used. To our knowledge, only one study29 discussed the presence of CPs with Cl1−2, stating that these CPs could be present in mixtures with <50% Cl. In general, isomer information (i.e., chlorine configuration alongside the carbon chain length) of the blobs detected by GC × GC–ECD remains unknown and standards are needed for identification and confirmation.

The first peaks were observed in fractions B5−B3 (fractionation time 200−214 s). This is in line with the peaks observed around 3.5 min in the GC-FID chromatogram (Figure S2). After that, peaks remained below detection limits until C5 (263 s). From fraction C6 at 270 s, peaks were detected, of which examples are shown in Figure 3A−C (fraction E6−E7, 438−445 s). Each consecutive fraction contains a separate peak eluting ca. 45 s later than the peak of the fraction before, illustrated in Figures 3A−C and S3 by the blue dotted lines, which indicates that different isomers are collected in each fraction. Interestingly, a certain pattern was identified in the fractions by inspecting the single chromatograms of the GC × GC–ECD, of which an example of fraction E6 and F11 is shown in Figure 3B,D: each fraction contains two peaks with a 5 min interval on the first column retention time, indicating that two different isomers are present. The two peak combination gradually moves with each consecutive fraction alongside the retention time of the first column to the right maintaining the 5 min interval. Furthermore, the second eluting peak of a fraction is eluting just right after the first eluting peak (ca. 7 s) in eight fractions later, indicating that these are two different isomers. An example is shown in Figure 3B,D, where the second eluting peak in fraction E6 is eluting just right after the first eluting peak in fraction F11, illustrated by the blue dotted line. As these are different isomers, it rejects the argument that something went wrong during fractionation. Rather, the difference in oven temperature program could explain this observation. Although the column separated the isomers during GC fractionation to a good degree (indicated by only having two peaks in one fraction rather than an uncountable number of peaks in a 5 min interval on the GC × GC chromatogram, Figure 3D), it was unable to separate the two remaining isomers due to the relatively fast oven temperature program (15 min). With the rather long oven temperature program of the GC × GC–ECD (90 min), the column (same type as GC fractionation) was able to separate these two peaks. This observation clearly shows how complex CP mixtures are and how many isomers are present. Nonetheless, in fractions E3−E7 (Figure S4), the second peak is of such low intensity that those fractions contain (as good as) isolated isomers, which might be used as standards.

High-Resolution GC Fraction Analysis by APICI-TOF-MS. To further evaluate the fractionation process, the unfraccionated mixture and fractions were analyzed by the chlorine-optimized APICI-TOF-MS method. The APICI-TOF-MS method omits chromatographic separation and solely focuses on mass separation by using only a guard column in the LC. How and which CP isomers form [M + Cl]− ions by this method remains unknown. CP congener groups C14Cl3 to C14Cl12 were detected in the unfraccionated mixture (Figure 4A), although C14Cl3 (zoomed-in panel Figure 4A) and C14Cl11−12 were in low abundance. The CP congener groups C14Cl4 (m/z 303.12344) and C14Cl12 remained undetected, indicating that they do not ionize (C14Cl2) or that they are absent in the mixture (C14Cl5 and/or C14Cl13). The absence of C14Cl12–14 congener groups is in line with the GC × GC–ECD results and reasonable, as CP−C14 is a low-chlorinated CP mixture (41.3% Cl). In line with the low chlorination degree, C14Cl3 is the most dominant congener group in the mixture.

In the first 53 fractions (A1−ES, fractionation time 0−431 s), peaks remained below detection limits. This included fractions C6−ES, in which peaks were observed by GC × GC–ECD, which indicates that GC × GC–ECD is more sensitive in detecting CPs than APICI-TOF-MS. Two or more m/z spectral clusters representing different congener groups were identified in most fractions, of which an example is shown in Figure 4B, which is consistent with GC × GC–ECD results. In fractions E9−E12 (459−480 s), F12 (487 s), and F9 (508 s), only one spectral cluster was identified (of which an example is shown in Figure 4C), which could indicate that one isomer or isomers with the same molecular formula (i.e., congener groups) were present in these fractions. To our knowledge, congener group standards of CP−C14 are commercially unavailable. Response factors between congener groups are unknown, let alone for different isomers. For APICI-TOF-MS, these fractions could be suitable as standards.

High-Resolution GC Fraction Analysis by NMR. The unfraccionated mixture (Figure 4A) and all 96 fractions were analyzed by NMR (Figure S5), of which the mixture and 12 fractions [i.e., A9 (123 s), B4 (207 s), C9 (291 s), D4 (375 s), E3 (4157 s), E6−E7 (438−445 s), E9−E10 (459−466 s), F4 (543 s), G9 (627 s), and H4 (7111 s)] were selected for interpretation.

In the first 49 fractions (A1−E1, 0−403 s), peaks remained below detection limits by spectral interpretation. From fraction E2 and onward, peaks were detected, of which examples are given in Figures 5 and 6. This shows that these fractions were enriched enough to be analyzed by NMR. To the best of our knowledge, this is the first time that a whole GC chromatogram could be analyzed by NMR. Given below are the first steps of structure assignments, by describing what kind of structures are
expected and which structures the observed peaks highly likely represent.

A $^1$H spectrum of 2-chlorobutane was recorded (Figure S6) as a model spectrum to determine the chemical shift of the methyl group adjacent to a chloromethylene group (−CHCl=, green region in Figure S7). Based on the literature $^6,7,29$ and $^1$H NMR spectra of 2-chlorobutane (Figure S6), we would expect at least some of the following substitution patterns. CPs are made by radical chlorination of paraffinic hydrocarbon mixtures that contain straight-chain saturated hydrocarbons, in which secondary and primary carbons are primarily available for chlorine substitution. $^7$ Therefore, Howard et al. $^7$ suggested that chlorine atoms are probably randomly distributed, up to a point, along the methylene (−CH=−) groups of the carbon chain. The reactivity of the aliphatic hydrogens during free-radical chlorination is dependent on the stability of the intermediate radical species, meaning that tertiary carbons are chlorinated faster than secondary ones, followed by primary ones, which was also found by Yuan et al. $^7$ Geminal chlorination (two chlorines on one carbon) is unfavorable. As the inductive effect of chlorine reduces the reactivity of the neighboring C−H bond, geminal dichlorination is highly unfavorable and does not occur in lower-chlorinated CP mixtures such as the one used in this study (41.3% Cl). $^7$ The same inductive effect also inhibits vicinal substitution (−CHCl−CHCl−) whenever more favorable sites are available. Similarly, terminal chlorination is also less likely to occur in CPs present in the mixture CP−C$_{14}$ 41.3% Cl.$^7$

Following the reasoning above, it is expected to find at least peaks in δ 1.6−2.5 (blue region Figures 5 and 6), which are most likely from the (nonchlorinated) methylene groups, as well as peaks between δ 4.0 and 4.5 (yellow region), representing the chlorinated methylene groups (−CHCl−). These peaks are indeed present in the spectra of the mixture (Figure 5A) and fraction E2 and onward (Figures 5, 6, and S5), while absent in earlier fractions. Interestingly, differences are observed in the number of peaks and peak shapes between the fractions. For example, the peaks in the blue region in fraction F4 (Figure 6F) differ from those in fraction H4 (Figure 6H). A clear upfield shift is visible, indicating that the isomer(s) in fraction H4 contains more chlorinated methylene groups and thus more chlorine atoms. For structure assignment, this warrants future research.

As these fractions are derived from a low-chlorinated CP mixture (41.3%), peaks are also expected between δ 0.8 and 1.2 (purple region Figures 5 and 6), which most likely represent a terminal non-chlorinated methyl (−CH$_3$) group attached to one or more non-chlorinated methylene (−CH=−) groups (see legend in Figure 5), as well as between δ 1.2 and 1.6 (green region), most likely from methyl groups adjacent to chlorinated methylene groups (−CHCl−) as seen in the $^1$H spectrum of 2-chlorobutane (Figure S6). These peaks are also present in the spectra of the mixture and fractions, again although in different number and peak shapes. The region δ 1.2−1.6 unfortunately partly overlaps with the water peak.

Peaks at 3.5−4.0 (orange region) come from chloromethyl groups (−CH$_2$Cl), representing in this case chlorinated terminal carbons. These are found in higher abundance in the fractions containing higher chlorinated CP isomers (fractions G9 and F4, Figure 6G, H) compared to the rest of the fractions (Figure 6A−F). This is in line with their higher chlorination degree and thus has a higher likelihood of terminal chlorination. All in all, the results above indicate that there is variation per well and thus that different isomers are successfully collected according to their properties, during fractionation.

Due to the numerous possibilities and combinations of C−H and C−Cl bonds, it remains a challenge to elucidate the whole structure of individual isomers by spectral interpretation in NMR spectra of even single to a few CP isomers. For assigning the structure of individual isomers in each fraction, more advanced methods are needed. An option is to use multivariate data analysis with the GC × GC, MS, and NMR data, which falls outside the scope of this proof-of-concept study, but is ongoing research. With this method, the goal is to assess whether there are similarities in NMR shifts (that might stand for certain bonds) in the spectra obtained by NMR analysis between the fractions, using the information on the number of chlorines present in the isomers per fraction, which is obtained by GC × GC and APCI-TOF-MS analysis.

Concentrations in the fractions were too low to perform $^{13}$C NMR, correlation spectroscopy, and heteronuclear single quantum coherence experiments with a reasonable resolution and S/N ratio. An option to enable such analyses is to extend the number of fractionations for more enrichment. Fractionation may also be extended to 384 wells in future work depending on the GC resolution and peak widths, although the fractionation might be too scattered to enrich enough material needed for NMR spectroscopy. Other suggestion for follow-up studies is to investigate the possibilities of using larger internal diameter columns and large volume injection to increase the loadability on the GC column to reduce the number of injections. This might be challenging due to the required higher flow rates, which require a new setup of the system and optimization of the backpressure.

### CONCLUSIONS

A continuous high-resolution GC-fractionation platform was successfully applied to analyze, for the first time, a whole GC chromatogram by NMR spectroscopy. In addition, up to a few CP isomers per fraction from a technical mixture were successfully isolated for the first time.

The application provided 96 NMR spectra of one GC chromatogram containing valuable information of the structure of single to a few CP isomers per fraction. Detailed spectral interpretation of a few NMR spectra was made to assign some of the structures present in the isomers, showing that the isomers in the fractions were enriched enough for NMR spectroscopy. To fully characterize the CP isomers, additional in-depth analysis such as multivariate data analysis is needed, which is outside the scope of this article. CPs are known as one of the most complex (i.e., 10,000 isomers and numerous possibilities and combinations of C−H and C−Cl bonds) and thus challenging mixtures for characterization.

The next step is to analyze the NMR, GC × GC, and MS data by multivariate data analysis, which is ongoing. The goal is to identify important spectral ranges that could stand for certain sub-structures present in a C$_{14}$−CP mixture to unravel and confirm the modeled results of the chlorine substitution on CPs in technical mixtures. This can then lay the foundation for assessing quantitative structure−activity relationship predictions. The isolated isomers obtained in this study can be further purified and certified as reference standards, which are urgently needed, or used in persistency, bioaccumulation, and toxicity studies.
Chlorinated Para synthesizing the CP-14 50% mixture and Andrew Jaques of the Da Vinci Laboratory Solutions is thanked for providing the GC-FID chromatograms of CP-C14 41.3% Cl mixture, all 96 fractions, and 2-chlorobutane (PDF)

1H NMR spectra of CP-C14 41.3% Cl mixture, all 96 fractions, and 2-chlorobutane (PDF)

1H NMR spectra of all 96 fractions (MP4)

AUTHOR INFORMATION

Corresponding Author
Louise M. van Mourik — Department of Environment and Health (E&H), Faculty of Sciences, Vrije Universiteit, 1081 HV Amsterdam, The Netherlands; orcid.org/0000-0003-3536-8895; Email: louise.van.mourik@vu.nl

Authors
Elwin Janssen — Department of Chemistry and Pharmaceutical Sciences, Faculty of Science, Vrije Universiteit, 1081 HV Amsterdam, The Netherlands
Robin Breeuwer — Department of Chemistry and Pharmaceutical Sciences, Faculty of Science, Vrije Universiteit, 1081 HV Amsterdam, The Netherlands
Willem Jonker — Department of Chemistry and Pharmaceutical Sciences, Faculty of Science, Vrije Universiteit, 1081 HV Amsterdam, The Netherlands
Jacco Koekkoek — Department of Environment and Health (E&H), Faculty of Sciences, Vrije Universiteit, 1081 HV Amsterdam, The Netherlands
Arif Arrahman — Department of Chemistry and Pharmaceutical Sciences, Faculty of Science, Vrije Universiteit, 1081 HV Amsterdam, The Netherlands
Jeroen Kool — Department of Chemistry and Pharmaceutical Sciences, Faculty of Science, Vrije Universiteit, 1081 HV Amsterdam, The Netherlands
Pim E. G. Leonards — Department of Environment and Health (E&H), Faculty of Sciences, Vrije Universiteit, 1081 HV Amsterdam, The Netherlands

Complete contact information is available at: https://pubs.acs.org/10.1021/acs.analchem.1c00049

Notes
The authors declare no competing financial interest.

ACKNOWLEDGMENTS

Da Vinci Laboratory Solutions is thanked for providing the GC-fractionation platform and Jaap Schap from Da Vinci is acknowledged for his advice and support concerning the GC fractionation platform. Quimica del Cinca is acknowledged for synthesizing the CP-14 50% mixture and Andrew Jaques of the Chlorinated Paraffins Industry Association is thanked for arranging the provision of the samples. Dr. Eelco Ruijter is acknowledged for his input and expertise on the interpretation of the NMR Spectra. L.M.v.M. and P.E.G.L. were financially supported by CEFIC LRI (ECO 38, ECO 42). L.M.v.M. was also financially supported by Eurostars (E!113388—CHLOFFIN).

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