# A SIMPLE AND RELIABLE METHOD FOR SEPARATION OF MINERAL OIL / POLYCHLOROBIPHENYL MIXTURES

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#### Abstract

Polychlorinated biphenyls (PCBs) were broadly applied world-wide as electrical insulators in transformers and power capacitors, due to their high dielectric constant and non-flammability. They were often added to mineral oils used as dielectric fluids, which are nowadays classified as hazardous waste. Indeed, the Stockholm Convention aims to eliminate the use of equipment with PCB content greater than 0.005 weight-% (= 50 ppm) by 2025 Accurate identification and quantification of small traces of PCBs contained in , neral oil thus represent a great analytical challenge. In order to achieve this goal, a cost-effective nple and fast chromatographic process was developed to separate PCBr from mineral oil, allowing to obtain reliable data to determine the concentration of down to 2-3 ppm. Experimental and analytical methods such as thin layer chromatography, column with mass spectroscopy (GC-MS) chromatography as well as gas chromatography were applied to acquire a high-level of qualitativ and quantitative determination of PCBs in transformer mineral oils.

**Keywords**: Polychlorinated bipliony's (PCBs); mineral oil; chromatographic separation; gas chromatography; mais spectroscopy.



## Introduction

Polychlorinated biphenyls (PCBs) are synthetic aromatic compounds known by the trade names Pyralene, Aroclor and others (Ericson & Kaley, 2011). They refer to a family of 209 organochlorinated compounds consisting of two benzene rings substituted with 1 to 10 chlorine atoms in different positions (Ballschmiter & Zell, 1980). Their chemical and physical stability and non-flammability have allowed these compounds to be used mainly an electrical insulators in power transformers and capacitors, but also in some heaters and other electrical equipments (Ericson & Kaley, 2011; Rossberg et al., 2006).

PCB molecules have been revealed to be toxic, as they promote carcinogenic effects (Knerr & Schrenk, 2006). They have thus been classified as persistent organic pollutants (POPs) according to the Stockholm Convention (Stockholm Convention; Weber et al., 2011). Therefore, all transformer fluids containing PCB and considered hazardous waste and therefore must be monitored (Xu et al., 2013; PCB-elimination-network). In fact, it is prohibited to hold devices containing PCBs bove 50 ppm in the European Union (Council Directive 96/59/EC) pending them disposed by 2025 (Directive 2012/19/Eu; Regulation (Eu) 2019/1021).

Since the phaseout of PCBs mineral oil (MO) has been used as an alternative in power transformers, especially because of its electromagnetic insulating power equivalent to that of PCBs (Rouse, 1998), its wide distribution and its low cost compared to other fluids available in the market such as silicone and synthetic or vegetable ester oils (Borsi & Gockenbach, 2005). However, transformer oils often contain variable amounts of PCBs (Pelitli et al., 2015).

Generally, PCB molecules are analyzed using equipment with a capillary column usually filled with fused silica, this technique known as gas chromatography (GC) is often coupled with an electron capture detector (ECD) (Bunert et al., 2019; Barcauskait, 2019). Other

detectors can be coupled to GC to detect PCBs, including the flame ionization detector (FID) which is sensitive to the detection of Volatile Organic Compounds (VOCs) (Huang et al., 2021), the flame photometry (FP) detector mainly used for compounds containing sulfur or phosphorus compounds (Clark & Thurbide, 2015), the thermoionic detector, also called nitrogen-phosphorus detector (NPD) used for nitrogen or phosphorus compounds (Wang et al., 2016), the photoionization detector (PID) adapted to the detection of ionizable mpounds (Liaud et al., 2014), or the coupling with mass spectrometry (MS) which is nsi live for the detection of POPs (Ballschmiter et al., 1992; Li et al., 2021). However, ite their high des sensitivity, GC detectors show a reduced efficiency of detection of PCBs in the presence of mineral oils of molar masses of around 1000 kg kmol<sup>-1</sup>. At present ECD detector remains the most used reference for the analysis of halogenated compounds (De Kok et al., 1982). This detector captures some electrons from the same which reduces the measured current. The compensation for this reduction is recorded a positive peak. As it is often the case for other GC techniques, a carrier gas should co tain few oxygen and water impurities that can interact with the stationary phase and cause considerable problems such as high reference noise or column bleed in the outp t gas chromatogram, which reduces analyzer sensitivity and column life. Impurit es can a oxidize the radioactive nickel source of ECD detectors and eix reactivity and lifetime, making it a major drawback of this therefore decrease conventional analytical technique (Poole, 2013; Bunert et al., 2017), justifying the broader use ometry for the detection of PCB. Nevertheless, other analytical difficulties can na spe be encountered such as the detection and quantification of PCBs in the presence of mineral oils with molar masses exceeding 1000 kg kmol<sup>-1</sup>. Indeed, the detection of PCBs in such media is considered to be the most difficult, especially for low chlorinated PCBs which have physical and chemical characteristics very similar to those of MO from electrical transformers (Takada et al., 2001). Moreover, in order to analyze all PCBs and to avoid interference

problems between PCBs and MO matrix derivatives, as well as to prevent pollution of detector ionization sources, it is necessary to combine a pretreatment technique capable of extracting PCBs from mineral oils before their injection into the GC-MS.

At present, the best-known pretreatment procedure is solid phase extraction (SPE) (Dmitrovic et al., 2002) combined with liquid-liquid extraction. This method requires the addition of DMSO for SPE, water to immobilize the PCB mixture, hexane for liquid-liquid extraction which must be later dried by sodium sulfate (Takada et al., 2001; Wittsiepe et al., 2014; Crucello et al., 2020). This method is expensive and requires the use of many solvents.

This work aims at describing a simple, reliable, and inexpensive chromatographic analytical separation method to extract PCBs from transformer MO, to reduce the analysis time and to increase the sensitivity of detection and quantification of high and low chlorinated PCBs, without using many additional chemicals.

For this purpose, combined separation and chematographic characterizations were first performed on model MO/PCB systems in the presence of 18 PCB molecules including the 7 most common ones called indic tor POBs (Afful et al., 2013), and then applied to an industrial sample of mineral of contaminated with unknown PCBs provided by Maxei company. The chematographic separation protocol was approved by thin layer chromatography, common chromatography filled with silica gel material under liquid petroleum ether new and nitrogen flow in order to accelerate the separation of MO/PCB systems.

The residting fractions, free of mineral oil molecules, can then be injected and analyzed by GC-EI/MS gas chromatography. This analytical and pretreatment combination improves the detection sensitivity of MO/PCB systems without any pollution concern for the detector ionization source.

#### Materials and methods

## **Chemical products**

#### Transformer mineral oil

The transformer MO samples studied were supplied by a french company (Arras Maxei), and were mainly composed of 3 families of hydrocarbons: paraffins, naphthenes, and non-chlorinated aromatics. A PCB-free virgin MO named *MO virgin* and a PCB-comminated MO denominated *MO contaminated* were considered here. The latter has been used for long-time in an industrial environment and contains *MO virgin* as MO as cell as an unknown number and quantity of PCBs.

## Polychlorinated biphenyls

The following PCB molecules were selected for task k (Figure 1) : Biphenyl (PCB N°0), 2-Chlorobiphenyl (PCB N°1), 3-Chlorobipheny (PCB N°2), 2,2'-Dichlorobiphenyl (PCB 2,4-Dichlorobiphenyl N°4). 2,3-Dichlorobiphenyl (PC) (PCB N°7), 3,4-N 5-Dichorobiphenyl (PCB N°14), 4,4'-Dichlorobiphenyl Dichlorobiphenyl (PCB N°12 (PCB N°15), 2,4,4'-Tricklorobip eryl (PCB N°28), 2,4',5-Trichlorobiphenyl (PCB N°31), 2,2',5,5'-Tetrachlorolinhenvl PCB N°52), 3,3',4,4'-Tetrachlorobiphenyl (PCB N°77), 2.2'.4.5.5'-Pentachlore hiphenyl (PCB N°101), 2,2',3,4,4',5'-Hexachlorobiphenyl (PCB 4,4',5,5'-Hexachlorobiphenyl (PCB N°153), 2,3,3',4,4',5-Hexachlorobiphenyl N°138 2,2',3,4,4',5,5'-Heptachlorobiphenyl (PCB N°180), and Decachlorobiphenyl (PKB (PCB 209), which were provided by Sigma-Aldrich (St. Quentin Fallavier, France). The choice of PCBs was mainly focused on the so-called indicator PCBs accounting for nearly 80% of all PCBs: most often PCBs No. 118, 138, 153 and 180, but also PCBs No. 28, 52 and 101. In 2017, these 7 indicator PCBs among the 209 congeners were selected by the Community Reference Office of the European Commission in Brussels as being the compounds to be searched for in priority in the analysis of organic matrices (sediment, blood, flesh, fat) due to their persistence and abundance in the environment as well as their toxicological properties (Turrio-Baldassarri et al., 1993). Other congeners of the same family with the same chlorine numbers have been added in this study in order to obtain more information on the chromatographic separation parameters.



**Figure 1.** Generalized chemical structure of polychlorinated bohenyls (PCBs). The case n=m=0 corresponds to the chlorine-free PCB N°0 (Biphenyl).

## **Organic** solvents

Petroleum Ether (40-60) and Acetone vere perchased from Aldrich (p.a. grade solvents). These reagents were used as received.

# Determination of the mover mass of transformer mineral oil (MO)

The molar mass of MO was determined by a MALDI-TOF/MS (Matrix Assisted laser Desorption Ionisation-Time of Flight/mass spectroscopy) using a nitrogen laser beam of wavelength 337 nm and pulse time 3 ns. MO was diluted to 1/100 in methanol and analyzed with a DHB matrix (2.5 Dihydroxybenzoic acid). The result obtained was recorded over an interval from 99 to 1500 Da. The determination of the molar mass of MO was repeated three times.

## Separation techniques

Thin Layer Chromatography

Thin Layer Chromatography (TLC) analysis of MO and MO/PCB mixtures was carried out on aluminum supports (20 cm x 20 cm) covered with uniform layers of silica gel as stationary phase. The thickness of this layer was about 0.2 mm (200  $\mu$ m). The TLC plates were placed in a rectangular tank containing the eluent which was a solvent mixture composed of 99 vol-% Petroleum Ether 40-60/1 vol-% Acetone) (Nikolaev et al., 2019; Ciura et al., 2017). The stains from the fractions deposited at 1 cm from the base of the layer migrate on the TLC plate more or less quickly according to interactions between the mobile phase (eluent and MO/PCB) and the stationary phase. The visualization of the different spots on the TLC plates was performed under a UV lamp with a wavelength of 254 nm. The plate appears fluorescent green and the products that absorb UV radiation appear as dark spots. The obtained TLC result of a given compound was expressed by calculating the "ratios to the front" value, R<sub>f</sub>, corresponding to the ratio of the distance traveled by the compound to that of the solvent front. Each TLC analysis was repeated six times.

## Silica column chromatography

Silica column chromatography (1CC) was carried out on a standard silica column (Silica 60 M, 0.04 - 0.063 mm, provided by Macherey-Nagel GmbH & Co. KG, Düren, Germany) (length : 33 cm, liam ter : 3 cm), applying the solvent mixture used by TLC analysis (99 vol-% Petroleum Ether 60-40 / 1 vol-% Acetone) as mobile phase. The obtained separation effects were controlled by TLC analysis. Indeed, the constituents of the MO/PCB mixtures migrated at different speeds on the support and were thus separated. Migration of MO/PCB mixtures on the silica column was done under a controlled compressed nitrogen flow, in order to accelerate separation. Each chromatographic column run was repeated six times.

## Gas chromatography coupled with mass spectroscopy

The recovered separated fractions containing PCB compounds were analyzed by GC-MS after drying in a vacuum oven at 70°C. The GC chromatograms and their associated MS spectra were obtained using a GC/MS apparatus (Clarus 680/Clarus 600T from Perkin Elmer, United States), equipped with a fused silica capillary column, which was maintained at 70°C (Elite-5 (5% Diphenyl) Dimethylpolysiloxane, 30 m × 0.53 mm (internal diameter ID), film thickness  $d_f = 0.5 \mu m$ ). The MS appliance was equipped with an EI (electronic ionization) onization source and a quadrupole filter (ion separator) at DC (U) and AC (V) voltages set by the equipment (Chamkasem et al., 2016).

## **Operating conditions of the GC-MS apparatus**

The thermal conditions of the GC oven were as follows. The c lumn has been maintained at a  $\mathbf{Q}^{\circ}\mathbf{C}$  min<sup>-1</sup> up to 120°C was applied. temperature of 70°C initially for 4 min then a rate The temperature has then been raised to  $250^{\circ}$  C are rate of  $30^{\circ}$  C min<sup>-1</sup>. The oven remained at erature was increased at a rate of 20°C min<sup>-1</sup> up this temperature for 20 min, and then the temp to 300°C. The latter temperature is manytained for another 20 min. The temperatures of the injector and the transfer line 0°C. The carrier gas used in our analysis was Helium. W The characteristic quantities the chromatographic analysis ( $t_R$  (min): retention time,  $t'_R$ tion time, k': capacity factor,  $\alpha$ : selectivity factor, N: number of (min): reduced rete theoretical plates,  $\mathbf{x}_s$ : column resolution) were measured following the method described in ngs, 2000). The reduced retention time  $t'_R$  is given by equation (1):

$$t'_R = t_R - t_M \tag{1}$$

where  $t_M$  is the time required for a species that is not retained by the stationary phase to pass through the column. The capacity factor k' is given by equation (2),

$$k' = \frac{t'_R}{t_M} \tag{2}$$

describing the rate of progression of PCBs in the column. This is the ratio of the quantities of a PCB analyte present at equilibrium in the 2 adjacent stationary and mobile phase volumes. The selectivity factor  $\alpha$  is given by equation (3),

$$\alpha = \frac{t'_R(B)}{t'_R(A)} = \frac{k'_A}{k'_B}$$
(3)

and represents the ability of the chromatographic system to ' distinguish 'chemically between sample components. It is usually measured as a ratio of the retention (capacity) factors (k') of the two peaks and can be visualized as the distance between their apices. The number of theoretical trays N is given by equation (4) :

$$N = 16 \left(\frac{t_R}{\omega}\right)^2 \tag{4}$$

The efficiency of the column increases when N increases. Note that  $\omega = 4 \sigma$  (the width at the base of the peak). The resolution of a column (R<sub>s</sub>) is given by equation (5), and gives a

$$R_{S} = 2 \frac{t_{R}(B) - t_{R}(B)}{\omega_{R} - \omega_{R}}$$
(5)

quantitative measure of its ability to separate two PCBs analytes A and B.

The detection and identification of POB molecules was performed with the help of a reference standard which was PC-43 (Nepracosa). The masses obtained by MS were limited in the range between 30 m z<sup>-1</sup> and 620 m z<sup>-1</sup> due to the type of electron ionization source which is not able to fragment molecules with masses higher than 620 m z<sup>-1</sup>. The ionization source temperature has been set at 280°C, and the voltage applied to the filament was 430 V.

## Quantification of PCBs by GC-MS

Calibration curves were established allowing independent quantitative determination of individual PCB concentrations. For this purpose, each indicator PCB was analyzed as reference standard by GC-MS using a variety of concentrations ranging from 10 ppm to 200 ppm, and signal area (*SA*) calculations were performed. The accuracy of this analytical

method was verified by graphical representation applying the following linear function: Area- $PCB = P^*C$ , where Area-PCB represents the peak area of a given PCB molecule as function of its concentration C, and P stands for the slope of this linear relationship. The  $R^2$ coefficients of determination were obtained by linear regression analysis using Origin version 8 software (Figure S1). Limits of detection (LOD) and of quantitation (LOQ) were calculated from each calibration curve, according to international regulatory agencies. OD was obtained by 3.3 times the ratio of the standard deviation of their response (s) nd he slope of the calibration curve (S),  $LOD = 3.3 * (SD \text{ of intercept x slope}^{-1})$ . LQS obtained from was  $LOQ = 10 * (SD \text{ of intercept x slope}^{-1})$ . The precision was obtained considering the relative standard deviation (RSD) for both samples and standard solutions e accuracy was assessed using certified samples.

## **Results and discussions**

## Molar mass of transformer mineral *sil*

The Maldi-TOF spectrum of virgin MO, shown in Figure 2, clearly proves that the molar mass of this oil exceeds 1000 kg kmol<sup>-1</sup>, which does not allow its direct injection and analysis into GC instruments coupled with a mass spectrometer or other detectors. Indeed, the spectrum shows fragments with masses of 600, 800, 1200 m  $z^{-1}$ , thus confirming the need to separate PCBs from MO before carrying out the GC analysis.





# Approval of the chromatographic protocol

The protocol for chromatographic suparation of MO/PCB mixtures is gathered in Figure 3. The different steps of the period process can be described as follows: first of all, an initial qualitative evaluation want. C of model systems composed of *MO virgin* in the absence and presence of one or more PCB molecules made it possible to separate qualitatively PCB and MO components. In order to implement quantitative and more sophisticated methods, a separate or *MO virgin* / PCB mixtures by SCC was necessary. After obtaining MO and PCB fractions separately, a primary vacuum distillation at 70°C was carried out. In this step, the organic solvents (petroleum ether and acetone) were vaporized and then recycled by condensation.

During the different steps of chromatographic separation, TLC analysis was systematically applied. It should be mentioned that the concentration effect of the deposited stains could hide

the appearance of other fractions. Once distilled, the fractions were dried in a vacuum oven at 70°C to remove traces of solvent. The penultimate step consists in elaborating the mass balance. The mass of *MO virgin* / PCB introduced into the column and then separated must correspond to the sum of the masses of the MO / PCB fractions ( $\Sigma \text{ m MO/PCB} \approx \Sigma$  mass of the MO/PCB fractions). This mass balance allowed to approve this protocol. The last step represents the most important one; it consists in studying the fractions containing PCBs by GC-MS analysis.



Figure 3. Chromatographic separation process of MO / PCB mixtures.

#### Thin Layer and Silica Column chromatography of MO/PCB mixtures

In order to check for the feasibility of the separation protocol for MO / PCB mixtures, TLC analysis was performed on three pure PCBs, PCB N°0 (non-chlorinated biphenyl), PCB N°15, which has 2 chlorines in para position, and PCB N°209, the most chlorinated congener (10 chlorines) of the PCB family. It can be seen that the stain of PCB N°0, i.e. in the absence of chlorines, migrates less quickly than the stains of PCB N°15 and PCB N°20 It c **n** be deduced that the migration of PCBs on the thin silica layer is not related the molecular weight of the PCB species, but rather to the polarity of the solvent and e Interactions of the chlorines with the stationary phase. The chosen mobile phase stroleum ether - acetone solvent mixture) is apolar: increasing the polarity of the PCR compound due to further chlorination leads to an increase of the migration rate. The results of the calculation of R<sub>f</sub> values of PCBs N°0, N°15 and N°209 were f d as 0.55, 0.65 and 0.75, respectively (Table 1).

Similarly, *MO virgin*, MO / PCB N°0 and MO / PCB N°209 mixtures were fractionated by SCC and then qualitatively analyzed by TLC. As a result, *MO virgin* presents three fractions while the other two samples exhibit each four fractions. Calculation of  $R_f$  of all samples were subsequently performed. The corresponding results of fractions 1, 2, 3 of *MO virgin* were found as 0.98, e.02, and 0.89 respectively. As expected, the same data were obtained for the MO part of the MO / PCB N°0 and MO / PCB N°209 mixtures. Moreover, the retention factor of the forth fractions yield almost the same values as those calculated for PCB N°0 and PCB N°209.

The MO compounds (paraffins, naphthenes...) present a more polar character than PCBs, explaining thus that they were less retained by the stationary phase compared to PCBs, which migrate more slowly.





The unknown sample (*MO contaminated*) was also studied by the same chromatographic separation process; showing six fractions (Table 1). Further analysis by GC-MS was needed on the three fractions that do not belong to *MO virgin*, in order to detect, distinguish and quantify PCBs present in this mixture.

## Detection and identification of PCBs in Mineral Oil by GC-MS

In order to obtain accurate information on the qualitative and quantitative templification of individual PCBs by GC-MS, detailed analysis of model systems based on a series of 18 PCB compounds was undertaken. This preliminary study comprised establishment of discrete calibration data for each PCB molecule as function of concentration, and also investigation of detection limits, which were found around 2-3 ppm per CB molecule.



**Figure 4.** GC chromatograms of model mixtures including 18 PCBs with different concentrations.

The concentration of each PCB molecule in the model mixture is mentioned on the associated chromatogram. For example, in the case of the concentration of 100 ppm, the total PCB concentration is: 100 ppm x 18 = 1800 ppm = 0.18 weight-%.

It will thus be demonstrated that the proposed chromatic separation method allows to determine nature and concentration of PCB molecules far below the actual 50 ppm limitation of PCB content in transformer fluids. It will also been shown later (Table 2) that the retention times of PCB molecules obtained by GC remained constant comparing those firm nodel PCB blends with those from PCB-containing fractions separated from MO / PCB mixtures by SCC.



Figure 5a



Figure 5c



**Figure 5.** Mass spectra of the 8 PCris separated from the model mixture: a) PCBs N °1, 2, 3, 4, 5, 7; b) PCBs N °12, 14, 15, 18, 28; c) PCBs N °31, 52, 70, 77, 101; d) PCBs N °138, 153, 156, 180, 209.

Subsequently, model mixtures comprised of 18 PCB molecules (indicator PCBs, listed in Table 2) were prepared at different concentrations (10, 25, 50, 100, 150, 200 ppm). For example, in the case of the concentration of 25 ppm, the total PCB concentration of the sample is: 25 ppm x 18 = 450 ppm = 0.045 weight-%. The chromatograms of some of these PCB mixtures (25, 50, 100 ppm) were presented in Figure 4, and the mass spectra associated with each molecule exhibiting specific retention times were shown in Figure 5. This method allowed detection, separation and identification of PCB molecules from model mixtures with

different concentrations of PCBs, and also of PCB fractions separated from MO / PCB systems (Figure 6), by comparing the obtained retention times with those of injected PCB molecules (standards) and also with data retrieved from an electronic library (NIST/EPA/NIH Mass Spectral Library (NIST 14)).



ppm) (b) the PCB fraction separated from the model MO / 18 PCB blend via SCC.

Since PCB compounds passess subilar chemical structures and differ only in the degree of chlorination, their vapor pressure was found directly proportional to the retention time on stationary plases of non-polar GC columns such as the one used in this report (Ballschmiter et al., 1993, Chamkasem et al., 2016). The molecular interactions of the PCB molecules with the stationary phase are thus governed by van der Waals forces (dispersive interactions). In addition, there are other effects affecting the separation of PCB on GC columns, like steric effects influencing the shape of the molecules, and dipole-dipole interactions.

$t_R$ (min)	$t_R$ '(min)	k'	á	ù	N	Rs
Retention time	Reduced retention time	Capacity factor	Selectivity factor	Width of the base peak	Number of theoretical plates	Column resolution
mod/sf	mod/sf	mod/sf	mod/sf	mod/sf	mod/sf	mod/sf
7.63/7.62	2.63/2.58	0.53/0.51		0.29/0.22	11468/11850	
9.88/9.84	4.84/4.8	0.96/0.95	1.83/1.86	0.18/0.1	46905/50586	9.31/9.76
11.88/11.83	6.88/6.79	1.38/1.35	1.43/1.41	0.10/0.20	53734/55979	10.78/10.61
13.67/13.59	8.67/8.55	1.73/1.70	1.26/1.26	0.51(0.30	31112/32615	6.95/7.03
16.02/15.90	11.02/10.86	2.20/2.15	1.27/1.27	0.62/0.58	10546/1024	5.03/5.24
17.92/17.75	12.92/12.71	2.58/2.52	1.17/1.17	0.41/0.39	29833/32973	3.66/3.81
19.76/19.59	14.76/14.91	2.95/2.89	1.1111.14	0.46/0.43	28893/33054	4.18/4.48
23.47/23.24	18.47/18.2	3.69/3.61	1.257.25	0.51/0.50	33885/34019	7.61/7.81
24.68/24.34	19.68/19.3	3.94/3.83	.0,1.06	0.63/0.62	24554/24344	2.12/2
28.07/28.05	23.07/23.01	4.61/4	1.17/1.19	0.14/0.14	599610/651562	8.75/9.72
29/29.01	24/23.97	4.8/4.1	1.04/1.04	0.13/0.12	861184/919696	6.89/7.38
30.47/30.46	25.47/25.42	5. 9/5.0	1.06/1.06	0.13/0.13	815075/865041	11.31/11.5
30.93/30,93	25.93/25.89	5.19/5.14	1.02/1.09	0.17/0.13	499809/919814	2.97/3.61
31.46/31.32	26.46/26.28	5.29/5.21	1.02/1.02	0.11/0.11	1197407/1297113	3.66/3.59
31.74/31.70	26.74/26.66	5.35/5.28	1.01/1.01	0.12/0.12	1031606/1062743	2.33/3.60
32.22/32.20	27 22/21 16	5.44/5.39	1.02/1.02	0.14/0.14	789034/822725	3.41/3.77
32.41/32.38	2 3727.34	5.48/5.42	1.01/1.01	0.12/0.12	1057306/1091013	1.49/1.35
34.60/34.58	29.6/29.56	5.92/5.85	1.08/1.08	0.14/0.13	1051005/1098049	16.93/17.19
	$t_R$ (min) Retention time mod/sf 7.63/7.62 9.88/9.84 11.88/11.83 13.67/13.59 16.02/15.90 17.92/17.75 19.76/19.59 23.47/23.24 24.68/24.34 28.07/28.05 29/29.01 30.47/30.46 30.93/30,93 31.46/31.32 31.74/31.70 32.22/32.20 32.41/32.38 34.60/34.58	$t_R$ (min) $t_R'$ (min)Retention timeReduced retention time $mod/sf$ $mod/sf$ $nod/sf$ $mod/sf$ $7.63/7.62$ $2.63/2.58$ $9.88/9.84$ $4.84/4.8$ $11.88/11.83$ $6.88/6.79$ $13.67/13.59$ $8.67/8.55$ $16.02/15.90$ $11.02/10.86$ $17.92/17.75$ $12.92/12.71$ $19.76/19.59$ $14.76/14.91$ $23.47/23.24$ $18.47/18.2$ $24.68/24.34$ $19.68/19.3$ $28.07/28.05$ $23.07/23.01$ $29/29.01$ $24/23.97$ $30.47/30.46$ $25.47/25.42$ $30.93/30.93$ $25.93/25.89$ $31.46/31.32$ $26.46/26.28$ $31.74/31.70$ $26.74/26.66$ $32.22/32.20$ $27.22/21.16$ $32.41/32.38$ $29.6/29.56$	$t_R$ (min) $t_R'$ (min) $k'$ Retention timeReduced retention timeCapacity factor $mod/sf$ $mod/sf$ $mod/sf$ $7.63/7.62$ $2.63/2.58$ $0.53/0.51$ $9.88/9.84$ $4.84/4.8$ $0.96/0.95$ $11.88/11.83$ $6.88/6.79$ $1.38/1.35$ $13.67/13.59$ $8.67/8.55$ $1.73/1.70$ $16.02/15.90$ $11.02/10.86$ $2.20/2.15$ $17.92/17.75$ $12.92/12.71$ $2.58/2.52$ $19.76/19.59$ $14.76/14.91$ $2.95/2.89$ $23.47/23.24$ $18.47/18.2$ $3.69/3.61$ $24.68/24.34$ $19.68/19.3$ $3.94/3.83$ $28.07/28.05$ $23.07/23.01$ $4.61/4.57$ $29/29.01$ $24/23.97$ $4.54/4.8$ $30.47/30.46$ $25.47/25.42$ $5.29/5.21$ $31.46/31.32$ $26.46/26.28$ $5.29/5.21$ $31.74/31.70$ $26.74/26.66$ $5.35/5.28$ $32.21/32.38$ $27.22/2.16$ $5.44/5.39$ $32.41/32.38$ $29.6/29.56$ $5.92/5.85$	$t_R$ (min) $t_R'$ (min) $k'$ $\acute{a}$ Retention timeReduced retention timeCapacity factorSelectivity factor $mod/sf$ $mod/sf$ $mod/sf$ $mod/sf$ $nod/sf$ $mod/sf$ $mod/sf$ $mod/sf$ 9.88/9.844.84/4.80.96/0.951.83/1.8611.88/11.836.88/6.791.38/1.351.43/1.4113.67/13.598.67/8.551.73/1.701.26/1.2616.02/15.9011.02/10.862.20/2.151.27/1.2717.92/17.7512.92/12.712.58/2.521.17/1.1719.76/19.5914.76/14.912.95/2.891.11423.47/23.2418.47/18.23.69/3.611.25/2.2524.68/24.3419.68/19.33.94/3.830.91.0628.07/28.0523.07/23.014.61/4.41.04/1.0430.47/30.4625.47/25.425.9/5.141.02/1.0231.46/31.3226.46/26.285.29/5.211.02/1.0231.74/31.7026.74/26.665.35/5.281.01/1.0132.22/32.2072/27.21.165.44/5.391.02/1.0231.46/31.3225.9/2.73.45.48/5.421.01/1.0134.60/34.5829.6/29.565.92/5.851.08/1.08	$t_R$ (min) $t_R'$ (min) $k'$ $\dot{a}$ $\dot{u}$ Retention timeReduced retention timeCapacity factorSelectivity factorWidth of the base peak $mod/sf$ $mod/sf$ $mod/sf$ $mod/sf$ $mod/sf$ $mod/sf$ 7.63/7.622.63/2.580.53/0.510.29/0.299.88/9.844.84/4.80.96/0.951.83/1.860.19/0.1911.88/11.836.88/6.791.38/1.351.43/1.410.10/0.2013.67/13.598.67/8.551.73/1.701.26/1.260.52/0.5817.92/17.7512.92/12.712.58/2.521.17/1.770.62/0.5817.92/17.7512.92/12.712.58/2.521.17/1.170.41/0.3919.76/19.5914.76/14.912.95/2.891.D1/1.140.46/0.4323.47/23.2418.47/18.23.69/3.611.25/2.250.51/0.5024.68/24.3419.68/19.33.94/3.830.91.060.63/0.6228.07/28.0523.07/23.014.61/4.471.17/1.190.14/0.1429/29.0124/23.973.64/3.1.06/1.060.13/0.1330.93/30.9325.93/25.895.9/5.141.02/1.090.17/0.1331.46/31.3226.46/26.185.29/5.211.02/1.020.11/0.1131.74/31.7026.74/26.665.35/5.281.01/1.010.12/0.1232.22/32.2027.22/3.165.44/5.391.02/1.020.14/0.1432.41/32.383.90/29.565.92/5.851.08/1.080.14/0.13	$t_k$ (min) $t_k'$ (min) $k'$ $\hat{a}$ $\hat{u}$ $N$ Retention timeReduced retention timeCapacity factorSelectivity factorWidth of the base peakWhere of theoretical plates $mod/sf$ $mod/sf$ $mod/sf$ $mod/sf$ $mod/sf$ $mod/sf$ $mod/sf$ $mod/sf$ 7.63/7.622.63/2.580.53/0.51 $0.29/0.29$ 11468/118509.88/9.844.84/4.80.96/0.951.83/1.860.1260146905/5058611.88/11.836.88/6.791.38/1.351.43/1.410.00.2053734/5597913.67/13.598.67/8.551.73/1.701.26/1.26 $0.00.20$ 53734/5597913.67/13.5911.02/10.862.20/2.151.27/1.27 $0.62/0.58$ 10546/102417.92/17.7512.92/12.712.58/2.521.17/1.17 $0.41/0.39$ 29833/3297319.76/19.5914.76/14.912.95/2.891.14/1.4 $0.66/0.43$ 28893/305423.47/23.2418.47/18.23.69/3.611.25/2.52 $0.51/0.50$ 33885/3401924.68/24.3419.68/19.33.94/3.83 $0.1/60$ $0.63/0.62$ 24554/2434428.07/28.0523.07/23.01 $4.61/4.54$ $1.04/1.04$ $0.13/0.12$ 861184/91969630.47/30.4625.47/25.42 $5.95/5.14$ $1.06/1.06$ $0.13/0.13$ 815075/86504130.93/30.9325.93/25.89 $5.95/5.28$ $1.01/1.01$ $0.12/0.12$ $10.31606/1062743$ 31.74/31.7026.46/26.28 $5.35/5.28$ $1.01/1.01$ $0.12/0.12$ $10.31606/1062743$

The increase of the order of the separated PCBs in terms of the retention time followed essentially structure-retention relationships described in literature (Li et al., 2016). For example, the shortest retention time of 7.6 min was observed for the unsubstituted biphenyl (PCB N°0), whereas the longest retention time of 34.6 min was found for the fully chlorinated PCB N°209. It can be seen that the chromatographic method applied also allowed the separation by GC of PCBs possessing the same number of chlorines, such as PCB 10°4, N°5, N°7, N°12, N°14, and N°15 (Table 2).

The capacity (k') and selectivity  $(\alpha)$  factors of PCBs calculated from mal  $(t_R)$  and reduced  $(t_R)$  retention times allow to describe the rate of progression of RCB in the chosen stationary phase and to estimate the extent to which the latter can reparate them. Indeed, three situations have to be distinguished (Chamkasem et al. 2016) 1: elution is too fast; which is the case for PCBs N°0 and N°1; 1 < k' < 5: optimized ention for PCBs N°2, N°4, N°5, N°7, N°12, N°14, N°15, N°28, and N°52; k' > 5; elution is too slow for PCBs N°77, N°101, N°138, N°153, N°156, N°180, and N°209. The obtain ed capacity factors can be directly related to the etermining the molecular interactions with the non-polar chemical structures of the PCP stationary phase. Absence of child rines will thus lead to a high elution rate (k' < 1) as it is the case for PCB N°0, esents an apolar neutral molecule. The presence of only one vhi<u>ch</u> re ostion already increases k' from 0.53 (PCB N°0) to 0.96 (PCB N°1), chlorine in the ortho close to case of ptimal elution). In the case of PCB molecules with an adjacent k' factor the number of chlorine varies between 4 and 10. Indeed, these relatively high greate than generate more molecular interactions between PCBs and the stationary phase. These number "heavy" indicator PCBs, exhibiting long retention times, represent good separation characteristics on the GC column.

The selectivity factor  $\alpha$  (ratio of the distribution coefficients) was calculated in relation to the peaks of neighboring PCBs. This factor makes it possible to estimate the separating power of

the used column. It was found that  $\alpha$  values of low-chlorinated PCBs were slightly higher compared to the corresponding data of highly-chlorinated compounds; this is probably related to the mass effect of the PCBs which depends essentially on the number of chlorines.

The efficiency of the chromatographic column, which is directly proportional to the number of theoretical plateaus (N), depends on the degree of peak broadening of PCB peaks ( $\omega$ ), occurring when the PCB compound moves through the column. This broadening, pends on the residence time of each PCB molecule. The resolution of the column was also calculated in order to measure the ability to separate PCBs that have neighbor peaks. Three orin situations were highlighted (Table 2): If Rs > 1.5: neighboring PCB were completely separated; the situation  $Rs \approx 1$  corresponds to incomplete separation adjacent PCBs, and if Rs < 0.75, bordering PCBs were poorly separated. The results shown in Table 2 of the Rs **PAR** fractions from MO / PCB blends calculated from PCB-model mixtures and separate confirm the correct choice of the stationary phase

It can be deduced that all results obtained from the model mixture (18 PCB molecules) and the PCB fraction separated from the model MO / 18 PCB blend were almost identical, approving thus the separation method described here.

The correlation coefficients of the PCBs calibration curves (Figure S1) show a good linearity (between 0.992 and 0.993). The *LOD* and *LOQ* values were determined from the data of the calibration curves of PCBs (Table 3). These values confirm the reliability of the separation method described above, showing that the pretreatment used is consistent for complex matrice such as transformer oils.

РСВ	Linearity	LOD (ppm)	LOQ (ppm)
PCB N°0	0.997	9	30
PCB N°1	0.988	15	45
PCB N°2	0.999	4	14
PCB N°4	0.998	7	
PCB N°5	0.996	10	32
PCB N°7	0.995	13	40
PCB N°12	0.999	5	17
PCB N°14	0.997		30
PCB N°15	0.998		22
PCB N°18	0.995		38
PCB N°28	0.999	2	7
PCB N°31	0.92	13	40
PCB N°52	0.998	7	21
PCB 1°70	0.999	6	17
PCB 1º77	0.999	6	17
NCE N° 101	0.995	11	35
<b>P</b> CB N°138	0.999	2	7
PCB N°153	0.994	13	39
PCB N°156	0.999	3	10
PCB N°180	0.995	12	38
PCB N°209	0.993	14	45

**Table 3.** Linearity, limit of detection (*LOD*) and limit of quantification (*LOQ*) obtained for each PCB calibration curve.



**Figure 7.** GC chromatograms corresponding to the fractions 4-6 (F4, F5, F6) of a sample containing an unknown number and quantity of TSBs (*MO contaminated*).

The chromatographic separatio cocedure described above was also applied on a sample denominated MO contaminated taining unknown number and quantities of PCBs. In e analyzed (Figure 7), which do not belong to MO virgin. The particular, fractions 4 **-** <u>F</u>6 ntired and quantitatively determined indicator PCBs were presented in results in terms Table 4 The concentrations of the different PCB congeners were determined using multiple ves, following the same procedure as mentioned above. It can be seen that the calion 10nation of PCBs in MO contaminated greatly exceeds the limit imposed by current concen legislation (50 ppm PCB), with a total PCB concentration of 219 ppm. The results presented in Table 4 correspond to the analysis of fraction F6, which contains qualitatively the same indicator PCB molecules but at their highest concentration, compared to the other fractions F4 and F5.

	Type of indicator PCB	$t_R$ (min)	Concentration (ppm)
-	PCB N°156	32.36	52
	PCB N°138	<mark>31.44</mark>	27
	PCB N°77	<mark>29.1</mark>	
	PCB N°52	<mark>28.01</mark>	
	PCB N°31	<mark>24.56</mark>	
	PCB N°12	<mark>19.67</mark>	
	PCB N°7	<mark>16.11</mark>	
	PCB N°4	13.7	9
	PCB N°2	11.8	8
	PCB N°1	10.1	13
	PCB N°0	7.9	11

**Table 4.** Identification and quantification of *indicator PCBs* detected in the unknown *MO contaminated* sample. The results presented here correspond to the analysis of fraction F6.

## Conclusions

A new, simplified and efficient method for separating PCB molecules from contaminated transformer toine allows was established, allowing to identify these toxic molecules and to determine their concentrations. A chromatographic separation method by silica column was developed and applied on model systems composed of known quantities of PCB molecules mixed with PCB-free MO. An unknown sample of PCB-contaminated MO was investigated as well. GC-MS was applied as powerful analytical technique to evaluate the efficiency of the separation method in terms of the qualitative and quantitative analysis of the different PCB containing fractions. Indeed, these data confirmed reliably and efficiency of the proposed separation process.

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