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Thin-layer chromatography coupled with high performance liquid chromatography for determining tetrabromobisphenol A/S and their derivatives in soils

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As brominated flame retardants (BFRs), tetrabromobisphenol A/S (TBBPA/S) and their derivatives have raised wide concerns owing to their widely usage, distributions and adverse effects on human health, thus monitoring these BFRs was urgently needed. In this study, a rapid and cost-effective method based on thin-layer chromatography (TLC) sample pre-treatment coupled with high performance liquid chromatography-diode array detector (HPLC-DAD) (UV = 214 nm) was developed for determining TBBPA/S and their derivatives in soils, including TBBPA, TBBPA bis(allyl ether) (TBBPA-BAE), TBBPA bis(2,3-dibromopropyl ether) (TBBPA-BDBPE), TBBPS bis(allyl ether) (TBBPS-BAE) and TBBPS bis(2,3-dibromopropyl ether) (TBBPS-BDBPE). The method detection limits (MDLs) and the method quantification limits (MQLs) for these BFRs ranged from 0.023 to 0.087 μg g−1 dw and 0.076–0.29 μg g−1 dw, respectively. The recoveries were 41–108% and both RSD of repeatability and intermediate precision were less than 11%. The developed method presented good performance for analyzing natural soil samples collected from BFRs industrial park, suggesting its great application potential for monitoring environmental TBBPA/S and their derivatives.

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1. Introduction

Tetrabromobisphenol A (TBBPA), tetrabromobisphenol S (TBBPS) and their derivatives are the most widely used brominated flame retardants (BFRs), accounting for 60% of total BFRs market [1,2]. As additive products in plastics, ABS resins, etc., TBBPS and their derivatives are easily released to the environment during their production and usage processes and then cause potentially adverse environment and health risks [1–3]. Their universality in various environmental media have been revealed, such as TBBPA bis(2,3-dibromopropyl ether) (TBBPA-BDBPE) in the atmosphere [4] as well as in indoor dust [5], and TBBPA-BAE, TBBPA bis(allyl ether) (TBBPA-BAE) and TBBPS-BDBPE) in herring gull eggs [6]. The toxicological effects related to these BFRs, such as endocrine disruption effects, neurotoxicity and reproductive development toxicity, indicate their potential threats to organisms [7]. Therefore, monitoring these compounds by using a convenient and effective method has become a critical issue for studying their environmental behaviors, fate and toxicities [7–10].

Gas chromatography-mass spectrometry (GC–MS) [1,2,11] and high performance liquid chromatography-mass spectrometry (HPLC–MS) [6,11–13] are the most common proposed methods for determining TBBPA/S and their derivatives. Due to the thermal decomposition for directly analyzing TBBPA and derivatives, GC–MS methods are always not preferred for determining TBBPA/S and their derivatives [14]. For HPLC–MS methods, a variety of ion sources have been introduced to meet the analysis demand of complex samples, such as electrospray ionization-mass spectrometry (ESI–MS) [12], atmospheric pressure chemical ionization-mass spectrometry (APCI–MS) [13] and atmospheric pressure photoionization-mass spectrometry (APPI–MS) [11,15]. But their popularization for monitoring these BFRs is often limited partly owing to ionization difficulties resulted from the prop-
erties of the instruments and compounds. For example, the ESI source could not produce any precursor ions for TBBPA-BAE and TBBPA-BDBPE under full-scan optimization [10,16], the APCI source often leads to poor sensitivity [16], the unpopular APPI source needs an additional and appropriate dopant agent for ionization [11,17] and the extractive electrospray ionization-mass spectrometry (EESI–MS) needs a series of hard modifications of equipment [12].

Beyond that, cumbersome sample pretreatments for TBBPA/S and their derivatives are inevitable when using the methods based on MS techniques. Overall, these pretreatments, such as accelerated solvent extraction (ASE) [18], soxhlet extraction [14], liquid–liquid extraction [12,17], solid phase extraction (SPE) [11], gel permeation chromatography (GPC) [19], etc., are time-consuming, solvent-consuming and costly. Additionally, extra effort-consuming derivation is necessary for analyzing TBBPA by GC–MS because of its thermolability [11]. The controversially extreme conditions including the usage of concentrated sulfuric acid have also been introduced to eliminate the matrix interference, though they can lead to the positive decomposition of TBBPA under strongly oxidized condition [20]. Therefore, developing a rapid and cost-effective method for monitoring TBBPA/S and their derivatives is still urgently needed.

HPLC-DAD, a practical method for determining polar compounds, has also been proposed as a supplementary method for determining TBBPA/S and their derivatives [21–23], such as TBBPA-BDBPE and TBBPA-BAE in soil [9] and biological samples [7,23,24]. TBBPA in food [25], water [26] and biological samples [21,22]. But it often suffers from the interfering substances due to its identification mainly depending on the retention time of target compounds [9]. Elimination of interferences is one of the most important improvements for HPLC-DAD methods. Thin-layer chromatography (TLC) technique, a useful separation technique in pharmaceutical and biomedical field [27–34], possesses higher sample throughput, shorter analysis time and less extract and clean-up procedure than conventional pretreatment techniques [35]. In this study, a novel method coupled HPLC-DAD with TLC technique has been developed for determining TBBPA-BAE, TBBPA-BDBPE, TBBPS-BAE and TBBPS-BDBPE in soil samples. The method performances for environmental samples have been evaluated by analyzing these five TBBPA/S derivatives in natural soil samples.

2. Experimental

2.1. Chemicals and materials

Acetone, methanol, dichloromethane (DCM), n-hexane, ethyl acetate and acetonitrile, purchased from Merck (Darmstadt, Germany) and Fisher Scientific (Trinidad, UK), are HPLC or pesticide grade. Formic acid is obtained from Dikma technologies (USA). TBBPA, TBBPA-BAE, TBBPA-BDBPE, TBBPS-BAE and TBBPS-BDBPE are from Sigma-Aldrich (Steinheim, Germany). Ultrapure water is obtained from a Milli-Q advantage A10 system.

TLC plate (7.5 cm × 2.5 cm) is from Qingdao Haiyang Chemical Co., Ltd in China. The glass plates are precoated with silica gel GF254 and the coating thickness is 0.20–0.25 mm. Glass capillary (5 µL) is purchased from Hirschmann Laborgerate.

2.2. Soil sample extraction

Soil samples were collected from a large BFRs industrial park located in Weifang, Shandong Province, China. The soil samples were air dried, grinded, sieved (100 mesh) and stored in refrigerator at −20 °C before analysis.

Soil samples (1 g), mixed with 5 g of anhydrous Na2SO4, were ultrasonically extracted with 15 mL of a mixture DCM:n-hexane (1:1, v/v) for 15 min after blending sufficiently. The extraction process was repeated twice, and the extracts were combined and concentrated using rotary evaporator. The residues were then solvent-exchanged to 0.1 mL of acetone before TLC sample pretreatment. For the recovery experiments, each sample was spiked with the standards, mixed sufficiently and kept overnight before extraction. The extraction process was followed the procedure described above.

2.3. TLC separation

TLC separation was performed on a TLC plate (7.5 cm × 2.5 cm) precoated with silica gel GF254. Concentrated extracts from soil samples and standard solution (20 µL) were applied on one plate (5 µL of each for four equal sample points) as 7 mm bands, 7 mm from the bottom edge by 5 µL of capillary. The plates were developed with 0.5 mL of n-hexane: ethyl acetate: DCM: formic acid (30:6:1:1, v/v) as the mobile phase (migration distance, 5 cm) in a vitreous chamber at room temperature. After development and drying, visual detection of the plates was carried out at 254 nm in ZF-20C camera obscura ultraviolet analyzer. Referring to standards, the zones belong to TBBPA, TBBPA-BAE, TBBPA-BDBPE, TBBPS-BAE, TBBPS-BDBPE (2.7–4.4 cm off the baseline) were then marked and scraped from the surface of the plate layer. Target compounds were extracted from the scraped layer by acetone (20 mL), solvent-changed to methanol (500 µL) and further filtered through a 0.45 µm Nylon Syringe filters for HPLC analysis.

2.4. HPLC analysis

HPLC (Waters 1250 series) equipped with a photodiode array detector and Chemstation software was used for the instrument analysis. Pretreated samples (20 µL) were injected by automatic sampler, separated on an Agilent Zorbax ODS column (5 µm, 150 mm × 3 mm) and detected by photodiode array detector (UV 214 nm). The best separation was achieved at a flow rate of 600 µL min⁻¹ with acetonitrile and water (containing 0.1% formic acid) as the mobile phase. The linear gradient started isocratic at 60% acetonitrile for 3 min, then increased to 90% in 5 min and to 100% in another 8 min respectively. After remaining at 100% acetonitrile for 2 min, the gradient returned to 60% acetonitrile in 2 min and kept for 3 min.
2.5. Statistical analyses

Statistical comparisons of several mean values were performed by using one-way analyses of variance (ANOVA), taking the appropriate condition as a single factor. When the ANOVA leads to significant results, least significant difference (LSD) test was performed to identify where the differences occur. Statistically significant difference was defined as $p < 0.05$. The analyses were carried out by SPSS Data Editor (version 20.0). Waters Empower in the work station of Waters 1250 HPLC was used for peak purity tests of the analytes. A calculated purity angle less than purity threshold indicated a pure analyte peak.

3. Results and discussion

3.1. Method optimization

Standard solutions of the target compounds were used to optimize the developing solvents and applied volume for achieving good separation and narrow bands. Various combinations of the developing solvents, comprising those reported in the literature ($n$-hexane: ethyl acetate (5:1, v/v)) and fine-tuning ($n$-hexane: ethyl acetate: DCM (30:6:1, v/v); $n$-hexane: ethyl acetate: formic acid (30:6:1, v/v)) [30,36], were thoroughly tested. Better separation of analytes (fall into target zone, Fig. 1(B)) from co-extracted interferences (Fig. 1(A), (C)) and narrower bands of target zone were obtained with $n$-hexane: ethyl acetate: DCM: formic acid (30:6:1:1, v/v) as the developing solvents. The various applied volume of 2 µL, 5 µL, 10 µL and 20 µL was evaluated to obtain narrow, distinct and regular bands and stable retention factor ($R_f$) value that might decrease recovery on account of artificial operation. It turned out that an application volume of less than 5 µL showed perfect fingerprints.

Compared with conventional pretreatment techniques, TLC presented shorter analysis time, less solvent consumption, simpler clean-up procedure and lower cost, etc., than conventional pretreatment techniques in our study [35]. The solvent consumption and time consumption of TLC ($<30$ mL, $<30$ min) was much less than GPC ($>200$ mL, $>90$ min) [9]. The commercial silica gel plate selected was 10 times cheaper than SPE (e.g. LC-Si) cartridges [21,24]. The chromatographic process only covering physical adsorption and partition process can also effectively avoid the oxidation and degradation of these tested BFRs during pretreatment process.
Table 1
Method performance results of the developed TLC-HPLC-DAD method for TBBPA/S derivatives.

| Spiked concentration (ng g⁻¹) | TBBPA | TBBPS-BAE | TBBPS-BDBPE | TBBPA-BAE | TBBPA-BDBPE |
|------------------------------|--------|-----------|-------------|-----------|-------------|
| Models                       | -      | A = −2.2 × 10⁴ + 1.8 × 10⁴ | A = 3.8 × 10⁴ + 1.2 × 10⁴ | A = 7.7 × 10³ + 7.7 × 10³ | A = 8.9 × 10⁴ + 1.8 × 10⁴ |
| r                            | 0.9999 | 0.9997    | 0.9999      | 0.9999    | 0.9999      |
| Recovery (%)                 | 6.25   | 53 ± 1    | 92 ± 4      | 100 ± 5   | 51 ± 4      |
| Matrix effect (%)            | 2 µg mL⁻¹ | −0.76   | 4.66        | 0.55      | 7.68        |
| Intermediate precision       | Intraday | 2.06   | 0.25        | 0.025     | 0.023       |
| RSD (%)                      | Interday | 2.14   | 0.59        | 1.56      | 0.99        |

Under the optimized instrumental parameters, five target compounds could be baseline separated completely and quantified within 25 min (Fig. 2A). The identification of these compounds in the test solutions was verified by comparing the retention time (tR) and spectrum (Fig. 2) with the standard solutions from 8.24 min (TBBPA) to 22.76 min (TBBPA-BDBPE). Meanwhile, purity angles of the analytes were all less than purity thresholds, indicating peak purity of the analytes.

3.2. Method performance

Due to the probable degradation during the purification process in an open system [32], the stability of the target compounds (Fig. 3) in methanol at different storing temperature and on the plate were evaluated. The standard solutions (10 µg mL⁻¹) were stored at −20 °C and 4 °C for two weeks, and then analyzed. Compared with the freshly prepared standards, the results showed that the solutions were stable below 4 °C in two weeks (p > 0.05). Besides, a standard solution (1.25 mg mL⁻¹) was freshly prepared and applied on new plates. The plates were exposed to air and light for the intervals of 0, 30, 60 and 120 min before development. Compared with the freshly prepared standards, the results showed that the development on the plate should be finished in 30 min (p > 0.05).

The specificity of the method was ascertained according to Rf of fingerprints from the TLC plate images and the retention time (tR) of HPLC chromatogram. Rf, tR, and spectra of the sample solutions were appropriately in agreement with that of standard solutions (Rf, 0.54–0.88, Fig. 1; tR, 8.24–22.76, Fig. 2A). The range of linearity was from 0.1 µg mL⁻¹ to 100 µg mL⁻¹, with the correlation coefficients (r) higher than 0.9993 for all the five target compounds (Table 1).

The method detection limits (MDLs) and the method quantification limits (MQLs) were defined as the signal to noise (S/N) ratios of 3:1 and 10:1. The MDLs and MQs for TBBPS/S and their derivatives ranged from 0.023 to 0.087 µg g⁻¹ dw and from 0.076 to 0.29 µg g⁻¹ dw, respectively (Table 1). The MDLs of TBBPA (0.087 µg g⁻¹) and TBBPA-BDBPE (0.036 µg g⁻¹) were comparable with that obtained using HPLC-DAD (0.015 µg g⁻¹ of TBBPA [22], 0.006 µg g⁻¹ of TBBPA-BDBPE [7]) for biological samples.

Compared with some novel methods based on MS, the MDLs of TBBPA-BAE, TBBPA-BDBPE and TBBPS-BDBPE (0.023, 0.036 and 0.049 µg g⁻¹) in this study were higher, eg. the MDLs of TBBPA-BAE (0.03 ng g⁻¹), TBBPA-BDBPE (0.07 ng g⁻¹) and TBBPS-BDBPE (1.28 ng g⁻¹) for herring gull eggs by LC-APPI-MS [11], the MDLs of TBBPA-BAE (0.76 µg L⁻¹), TBBPS-BAE (4.6 µg L⁻¹) for waters by EESI–MS [12,17]. The instrument sensitivity of HPLC-DAD is the bottle-neck for the developed method, but HPLC-DAD can embody the advantage in the instrument price and usability for regular monitoring. Meanwhile, in view of the shorter analysis time and less extract and clean-up procedure of TLC separation, the developed method is proposed to be a supplementary technique for determining TBBPA/S and their derivatives.

The accuracy of the proposed method was evaluated by spiking blank soil sample with the analytes at concentrations of 6.25, 12.5, 25, 50 and 125 µg g⁻¹ [32]. The recoveries were from 41% to 108% and RSD were all less than 11% (Table 1). The intermediate precision was also examined by injection of freshly prepared standard solutions (5 µg mL⁻¹) in triplicate in one day and consecutive 3 days. The results of the intra- and inter-days precision were also shown in Table 1 and the RSD values ranged from 0.59% to 2.14%. These results indicated the stability of the method can meet the demand for monitoring these BFRs in soil samples.

3.3. Assessment of matrix effect

Due to the potential target signal enhancement or suppression caused by interferences, the matrix effects have been evaluated to assess the accuracy of the method. The matrix effects of soil samples were calculated by using the following Eq. (1) and the results were shown in Table 1.

\[
\text{Matrix effect} = \frac{C_2 - C_1}{C_1} \times 100\%
\]

C₁ was the concentrations of standard solutions of the studied BFRs in methanol and C₂ was the measured concentrations of the extracted and pretreated solution from blank soil sample spiked with equal standard solutions, with the same amounts of BFRs to C₁. The matrix effect experiments are composed of three groups of samples and the concentrations of the standard solutions are 2, 5 and 10 µg mL⁻¹. The results were shown in Table 1 and the matrix effects ranged from −5.68% to 8.44%, indicating that the matrix effects of soil samples could be ignored.

3.4. Application in natural soil samples

The performance of the developed TLC-HPLC-DAD method for monitoring environmental TBBPA/S and their derivatives were
evaluated by using two natural soil samples obtained from a large BFRs industrial park located in Weifang City, Shandong Province. Four compounds including TBBPA (2.0 µg·g⁻¹), TBBPS-BDBPE (3.1 µg·g⁻¹), TBBPA-BAE (2.2 µg·g⁻¹) and TBBPA-BDBPE (12.3 µg·g⁻¹) were detected in soil sample of S1 and two compounds including TBBPA (24.6 µg·g⁻¹) and TBBPA-BAE (2.1 µg·g⁻¹) were found in soil sample of S2 (Fig. 5). Comparison of the chromatograms and spectra of the analytes with standards (Fig. 2) and determination of the peaks purity indicated that the chromatographic peaks were pure, and further confirmed the real presence of these analytes in soil samples. The developed method can indeed meet the demand for monitoring these BFRs in the ambient area of BFRs plant.

4. Conclusion

In this study, a novel method based on TLC technique coupled with HPLC-DAD has been developed for analysis of five TBBPAs and their derivatives in soil samples. The proposed method presented simple, time-saving and solvent-saving pretreatment procedures, satisfactory precision, acceptable limits of detection, good recoveries and ignorable matrix effects for monitoring these compounds. Comparing with previous methods, TLC can dramatically simplify the sample pretreatment process without significant matrix effects. Using the developed method, high contents of TBBPAs and their derivatives in soil samples were detected, indicating that this method should be an important supplementary technique for monitoring these compounds in the environment.

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