Detection of sub-pptv benzene, toluene, and ethylbenzene via low-pressure photoionization mass spectrometry

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ABSTRACT

This paper reports on the advanced development of an ultrasensitive method for the detection of benzene, toluene, and ethylbenzene (or BTE) by low-pressure photoionization mass spectrometry (LPPI-MS). The LPPI source is composed of a laboratory-assembled krypton lamp and a stainless steel cylindrical ionizer. A compact V-shaped mass spectrometer is coupled to the LPPI source with a set of ion immigration optics under dc bias. The fixed standard concentration (FSC) and fixed standard volume (FSV) method are employed to calibrate the sensitivities of the instrument. The corresponding detection sensitivity toward BTE is 4–7 counts/pptv and the 2σ limit of detection (LOD) is 0.5–0.8 part per trillion (pptv). In addition, the measurement accuracy is 95%–105%, and the corresponding precision ranges from 3% to 15% and from 9% to 31% for the FSC and FSV methods, respectively. The stability (standard deviation) of LPPI-MS for a 1 ppbv BTE mixture is less than 0.025 (/>12 h). In the detection of BTE, water in ambient air is the most significant interfering factor, leading to the increased background, and inferior LODs of 1–2 pptv for BTE under an RH of ~90% is observed. Experimental results indicated that LPPI-MS is reliable for the detection of sub-pptv levels of BTE under laboratory conditions.

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1. Introduction

Benzene, toluene, and ethylbenzene, collectively known as BTE, are common toxic volatile organic compounds (VOCs) existing in
the production of petroleum, plastics, synthetic fibers, and pesticides [1]. In addition, BTE ubiquitous in urban atmosphere, posing substantial toxic health effects (e.g., mutagenic, carcinogenic stress and neurological, respiratory, excretory system damage) [2] and tremendously contribute to the smog photochemical reactions, influencing ozone in the troposphere [3]. For diagnostic purposes, BTE are considered as xenobiotic biomarkers in the breath of smokers [4].

Various real-time continuous analytical approaches have been developed to detect BTE. Ion mobility spectrometry (IMS) is the most portable and widespread method for the trace detection of aromatic compounds [5]. IMS usually equipped with a radioactive ionization source or an UV-lamp [6]. Although the ionization methods of these two kinds of ionization sources are different, their limit of detections (LODs) are similar, as low as several part per billion by volume for BTE [7–9]. The sensitivity ofIMS is susceptible to the ambient moisture level and diverse constituents due to the influence of water and competing ion–molecule reactions [10]. The same situation occurs in atmospheric-pressure chemical ionization mass spectrometry (APCI-MS). The sensitivity of analytes with a lower proton affinity (PA) (e.g., PA benzene = 750 kJ/mol) is subject to be suppressed by higher ones (e.g., PA toluene = 784 kJ/mol, PA ethylbenzene = 799 kJ/mol), along with high background signals from protonated solvent molecules [11]. When employing corona discharge (CD) as an atmospheric ionization source, the generation of H$_3$O$^+$ (H$_2$O)$_n$ and NO$^+$ (H$_2$O)$_m$ reactant ions can be markedly enhanced, leading to the LODs of ~400 part per trillion by volume (pptv) for BTE [12]. Proton-transfer-reaction mass spectrometry (PTR-MS) is another kind of chemical ionization based technique. In spite of the ionization efficiency in PTR is slightly lower than in APCI, PTR-MS could achieve superior LODs of ~10 pptv for BTE, by using high-density of H$_3$O$^+$ [13–15]. For higher mass resolution, time-of-flight (TOF) mass spectrometer used to be coupled with PTR. However, inferior LODs are observed, attributed to the loss of ions [13,16].

Photoionization (PI) technologies are known for the soft ionization. Recently, the detection performance of PI methods has been largely improved [17–19]. Photoionization mass spectrometry (PI-MS) could be a reliable alternative to IMS and PTR-MS. Laser-based resonance-enhanced multi-photon ionization (REMPI) is highly sensitive and selective for soft ionization of aromatic hydrocarbons, affording the LODs of ~10 pptv for BTE [4,20]. Nevertheless, the specific but limited analyte classes restrained the wide application of REMPI-MS. On the contrary, single-photon ionization (SPI) is able to soft ionize all organic compound classes [21]. The ionization efficiency in SPI is determined by the photon ionization cross-section (σ$_{pi}$) of the analyte, which affords better quantification by SPI as compared with that by other ionization methods [17]. Whereas, traditional SPI is limited to high vacuum condition and higher LODs (~10 ppbv) [22,23]. Atmospheric pressure PI (APPI) is a relatively new technique, allowing direct SPI at ambient pressure [17]. The LODs of APPI-MS for BTE are ~500 pptv [24]. Low-pressure PI (LPPI) is another relatively new method based on SPI, which is conducted under a pressure of hundreds pascals [25]. LPPI-MS has been confirmed as a rapid and ultrasensitive instrument for real-time monitoring of trace VOCs in previous studies, with the LODs of ~5 pptv for BTE [26]. Furthermore, LPPI can be more readily coupled to a TOF analyzer for higher resolution, without excessive loss of ions [17].

In this paper, the latest development in the detection of BTE by LPPI-MS is described. The detection sensitivity toward BTE is 4–7 counts/pptv. The corresponding 2× LODs for BTE are 0.5–0.8 pptv under nitrogen and 1–2 pptv under 90% RH. The accuracy and precision of the instrument are evaluated by two calibration methods. Detailed methods and experimental results are described in following sections.

2. Materials and methods

2.1. Instrumentation

Fig. 1 shows the structure and mechanism of our self-developed LPPI source, which mainly consists of a Kr vacuum ultraviolet (VUV) lamp and a highly efficient photoionizer with an optical baffle. The output of the lamp is approximately 5 × 10$^{14}$ photon s$^{-1}$ with energies of 10.0 eV (~80%) and 10.6 eV (~20%). Through a plano-convex MgF$_2$ lens, VUV light is focused into the ionization zone. The sample is introduced into the photoionizer via a needle valve controlled stainless steel tube. The pressure inside the photoionizer is set at 500–1000 Pa, maintaining a sampling flow rate of ~3.5 cm$^3$ s$^{-1}$. A set of ion immigration optics are aligned between the LPPI source and TOF-MS, serving as an ion guiding system for higher ion transmission efficiency. The ions expelled from the photoionizer are focused by the electrostatic fields of the ion lenses, followed by injection into the TOF mass analyzer. The laboratory-built TOF-MS has a flight distance of 460 mm in the V configuration. The detailed configuration of the LPPI-MS has been described previously [26,27]. Table 1 summarizes the optimized system parameters of the instrument. All experiments are conducted at 25 °C.

2.2. Sample preparation

For the calibration by the fixed standard volume (FSV) method, the signal intensity responses to increasing concentrations of three analytes are obtained with liquid samples, which are prepared by diluting pure BTE 500 times with dichloromethane. 0.5 μL of each diluted sample is extracted using a syringe and injected into a 120 L calibration chamber filled with high-purity nitrogen, corresponding to a concentration of 2 ppbv. For the calibration by the FSV method, a 0.5 μL diluted sample is injected six times, and mass spectra are accordingly acquired after each injection. Five

| Table 1 |
| --- |
| **LPPI-MS operational parameters used in this work.** |
| Parameter | Value selected |
| Extraction field | 600 V, 58000 Hz |
| Acceleration field | 1500 V |
| Reflector | 600 V |
| MCP detector | 2000 V |
| VUV lamp | 60 W |
| Ionizer pressure | 500–1000 Pa |
| Detection cycle | 10 s |
independent measurements are conducted for each chemical. Accordingly, the average and standard deviation are calculated.

The mass spectra shown herein are obtained with gas-phase samples to avoid the formation of impurity ions in solution. The gas-phase samples are prepared by injecting 6 mL of each pure chemical into a 6-L narrow-mouthed bottle filled with high-purity nitrogen and sealed with Parafilm member (Structure Probe, Inc., West Chester, PA). 5 mL of the gas-phase sample is added into the 120 L calibration chamber, ending up with a concentration of 10 ppbv for each sample.

For the calibration by the fixed standard concentration (FSC) method, 17 ppbv of a gas-phase sample is injected into a 0.24 L cylinder. High-purity nitrogen at a flow rate of 0.2 L min⁻¹ is continuously injected into the cylinder to dilute the 17 ppbv sample. Dynamical concentrations of the continuously diluted sample are calculated by an exponential equation described previously [28,29]. LPPI-MS mass spectra are acquired. Each FSC calibration starts from the highest signal intensity of sample. Five replicate batches are measured for each chemical.

Benzene (Shanghai Macklin Bio-chemical Co. Ltd., >99.5%), toluene (Beijing Beihua Fine Chemicals Co. Ltd., >99.5%), ethylbenzene (Shanghai Macklin Bio-chemical Co. Ltd., >99.5%), and dichloromethane (J.T. Baker Co. Ltd., chromatographic grade) are used in the experiments. High-purity nitrogen (>99.999%) is purchased from Beijing Haikeyuanchang Practical Gas Co. Ltd.

2.3. Accuracy and precision

The accuracy and precision of LPPI-MS for BTE (n = 5) are evaluated by analyzing various sample concentrations for the FSV and FSC calibration processes. Accuracy is expressed as the percentage of the target sample concentration using the mean measured concentration (n = 5) from a single detection batch. Accordingly, precision is expressed as the percent relative standard deviation (RSD).

2.4. Stability and interference experiments

The stability of LPPI-MS is assessed by continuously measuring a BTE mixture of 1 ppbv in 14 h. To assess the interference from water and other protonated impurities on BTE signal intensities, the BTE mixture (1 ppbv) is measured with acetaldehyde concentrations of 5, 10, 15, 20, and 25 ppbv in the calibration chamber. Relative humidity (RH) values of 10%, 30%, 50%, 70%, and 90% are set by injecting double distilled water (ddH₂O) into the calibration chamber monitored with a dewpoint monitor. The BTE mixture (1 ppbv) at different RH values is measured by LPPI-MS.

3. Results and discussion

3.1. Mass spectrum and detection limits

During LPPI, nonpolar VOCs (e.g., BTE) undergo direct photoionization, where a molecular ion is produced by the absorption of a single photon and the emission of an electron [30]. Fig. 2a shows the LPPI mass spectrum of benzene (ionization potential, IP = 9.24 eV) under nitrogen, where m/z 78 corresponds to the characteristic peak (M⁺). The signal intensity of 10 ppbv benzene is
as high as $7 \times 10^4$ counts. A mass resolution of 200 (fwhm) is achieved at m/z 78. The mass peaks at m/z 19, 37, and 55 correspond to $\text{H}_2\text{O}^+$ (protonated water), $(\text{H}_2\text{O})_2\text{H}^+$, and $(\text{H}_2\text{O})_2\text{H}^+$ (protonated water clusters), respectively. The protonated water clusters correspond to the photon-induced chemical ionization in the LPPI source. This phenomenon has also been observed in our recently reported measurement of other polar species, e.g., alcohols, aldehydes, ketones, and esters [31]. As shown in Fig. 2b, linear responses ($R^2 = 0.999$) are obtained for ppbv levels of benzene from both FSV and FSC calibration methods. The slopes of the fitted lines are 7.1 and 5.0 counts/ppbv, respectively. The LODs at an S/N = 2 are estimated by LOD = 2σc/h, where σ is the standard deviation of the noise, c is the sample concentration, and h is the ion signal intensity [32]. The noise (σ) value in the no-signal region of high-purity nitrogen background is 1.7. The LODs for LPPI-MS toward benzene (B) are calculated as 0.5 ppbv (FSV method) and 0.7 ppbv (FSC method). However, for ambient measurements, the noise intensities increased by approximately 2–3 times, indicating that the method detection limit (MDL) for ambient benzene samples is 1–2 ppbv.

Table 2 lists the reported LODs of various instruments toward BTE.

**Table 2**

| Number | BTE    | m/z | IP (eV) | LODs               | References        | FSV method | FSC method |
|--------|--------|-----|---------|--------------------|-------------------|------------|------------|
| 1      | Benzene| 78  | 9.24    | 3 ppbv             | LPPI-MS [26]      | 0.47 pptv  | 0.67 pptv  |
|        |        |     |         |                    | HIPP-MS/RF-only Quadrupole [33] |            |            |
|        |        |     |         |                    | PTR-MS [13]       |            |            |
|        |        |     |         |                    | TD-SPI-MS [4]     |            |            |
|        |        |     |         |                    | TD-REMPI-MS [4]   |            |            |
|        |        |     |         |                    | GC-FID [34]       |            |            |
|        |        |     |         |                    | APCLI-MS [12]     |            |            |
|        |        |     |         |                    | APPI-MS [24]      |            |            |
|        |        |     |         |                    | PI-IMS [8]        |            |            |
|        |        |     |         |                    | MVP-SPI-MS [23]   |            |            |
|        |        |     |         |                    | TD-FTIR-HWG [35]  |            |            |
|        |        |     |         |                    | HiKE-IMS [9]      |            |            |
|        |        |     |         |                    | GC/MPI/TOFMS [36] |            |            |
| 2      | Toluene| 92  | 8.83    | 21 ppbv            | HIPP-MS/RF-only Quadrupole [33] | 0.55 pptv  | 0.63 pptv  |
|        |        |     |         |                    | PTR-MS [13]       |            |            |
|        |        |     |         |                    | TD-SPI-MS [4]     |            |            |
|        |        |     |         |                    | TD-REMPI-MS [4]   |            |            |
|        |        |     |         |                    | GC-FID [34]       |            |            |
|        |        |     |         |                    | APCLI-MS [12]     |            |            |
|        |        |     |         |                    | TG/GC/SPI-MS [38] |            |            |
|        |        |     |         |                    | MVP-SPI-MS [23]   |            |            |
|        |        |     |         |                    | HiKE-IMS [9]      |            |            |
|        |        |     |         |                    | GC/MPI/TOFMS [36] |            |            |
| 3      | Ethylbenzene| 106 | 8.77  | 3 ppbv             | PTR-MS [14]       | 0.59 pptv  | 0.83 pptv  |
|        |        |     |         |                    | GC/MPI/TOFMS [36] |            |            |

**References**

- FSV method: High pressure photoionization mass spectrometer (HPPI-MS).
- FSC method: Direct photoionization mass spectrometry (APPI-MS).
- Other methods: Thermal-desorption single photonionization mass spectrometry (TD-SPPI), medium vacuum pressure single photonionization mass spectrometry (MVP-SPPI), HiKE-IMS (high kinetic ionenergies mobility spectrometry), TD-FTIR-HWG (thermal-desorption fouriertransform infraredhollowwaveguides), GC-FID (gas chromatography with flame ionization detection), GC-DMS (gas chromatography with differential mobility spectrometry).
FSC methods, respectively (Table 2). Accordingly, the MDL for ambient ethylbenzene is evaluated as $1 \text{e}^2 \text{pptv}$. These experimental results revealed that LPPI-MS is significantly sensitive for real-time measurements and can detect BTE samples with concentrations as low as sub-parts per trillion by volume, which exceeds other competitive techniques (e.g., TD-SPIMS and GC-FID). However, without the help of GC, LPPI-MS cannot discriminate isomers, e.g., ethylbenzene and xylene.

3.2. Accuracy and precision

As shown in Table 3, the accuracy of LPPI-MS ranges from 96 to 104% for BTE samples for the FSV calibration method. With increasing concentrations of BTE, the precision (RSD) of the FSV method for BTE is gradually improved from ~30%. The accuracy for the FSC method ranges from 97% to 102% for BTE samples. However, the precision (RSD) for the FSC method is obviously superior to that observed for the FSV method, ranging from 3% to 15%. The better RSDs for the FSC method are caused not only by the higher concentrations of BTE in samples utilized in the approach but also by the superior calibration curves with less error bars (Figs. 2b, 3b and 4b). Calibration results indicated that the FSC method is more stable than the FSV method from experiments, while the sensitivity obtained by the FSV method is greater than that obtained by the FSC method. It is thought that the underestimation of the initial concentration (evaluating the dynamical concentration with an acquisition time of 10 s) in the FSC method leads to lower calibration sensitivities.

3.3. Stability and interfering factors

The stability of LPPI-MS is evaluated by continuously measuring a 1 ppbv BTE mixture for 14 h. Fig. 5a shows the standard deviation for measurements. The maximum standard derivation for BTE is 0.025, indicative of the reasonably good stability of LPPI-MS. Fig. 5b and c show the effects of interfering factors of water (under RH values of 10%, 30%, 50%, 70%, and 90%) and acetaldehyde (at 5, 10, 15, 20, 25 ppbv), respectively, on the signal intensities of BTE. Linear regression equations of BTE are consistent to be $y = 0.002x + 1$ from 10% to 90% RH. The slopes of the fitted lines for BTE are $1.1 \times 10^{-4}$, $1.6 \times 10^{-4}$, and $4.8 \times 10^{-4}$, respectively, when doped with 5–25 ppbv acetaldehyde. Hence, the direct effects of water and acetaldehyde on the signal intensities of BTE are not significant.

Moreover, in our previous study, the matrix effect of CH$_2$Cl$_2$ was also not observed for benzene, while a significant doping effect of CH$_2$Cl$_2$ was observed for polar compounds, e.g., water and acetaldehyde [40]. These results might indicate that BTE mainly undergo photoionization in the LPPI source. However, considering multiple collisions occurring in the LPPI source, LPPI-MS is supposed to be valid for the detection of BTE in presence of low impurity concentrations (dozens of ppbv).

The water impurity in the carrier gas affects the S/N, evidenced by the increased baseline noise. The intensities of noise at 90% RH are approximately 2–3 times greater than those at 10% RH. Hence, the LODs for BTE accordingly worsen to $1 \text{e}^2 \text{pptv}$, which are in accordance with the MDLs under the ambient condition. These experimental results revealed that LPPI is suitable for the sub-pptv...
level detection of benzene and its derivatives under dry conditions, but the presence of high-concentration water will partly decrease the LOD of the instrument.

4. Conclusion

LODs on the order of sub-pptv for benzene, toluene, and ethylbenzene (or BTE) are evaluated by the FSV and FSC calibration methods. The detection sensitivity of the instrument toward BTE is 4–7 counts/pptv with excellent linear responses. Measurement accuracy for both methods is 95%–105%; precision ranges from 3% to 15% for the FSC method and from 9% to 31% for the FSV method. These experimental results demonstrated that low-pressure photoionization mass spectrometry provides reliable responses to sub-pptv BTE under nitrogen. Under ambient moisture, water increases the noise level, affording decreased LODs (1–2 pptv). Given

Table 3
Accuracy and precision (RSD) of BTE in different calibration methods.

| Concentration (ppbv) | Benzene | Toluene | Ethylbenzene |
|----------------------|---------|---------|--------------|
|                      | Accuracy (%) | RSD (%) | Accuracy (%) | RSD (%) | Accuracy (%) | RSD (%) |
| FSV calibration      |          |         |              |
| 2                    | 101      | 31      | 101          | 31      | 98           | 28      |
| 4                    | 98       | 22      | 100          | 30      | 100          | 19      |
| 6                    | 99       | 17      | 99           | 23      | 104          | 15      |
| 8                    | 101      | 15      | 98           | 20      | 100          | 17      |
| 10                   | 100      | 12      | 100          | 19      | 96           | 17      |
| 12                   | 100      | 9       | 102          | 18      | 101          | 15      |
| FSC calibration      |      |         |              |
| 5.6                  | –        | –       | 102          | 3       | –            | –       |
| 6.4                  | 100      | 7       | 99           | 3       | 102          | 3       |
| 7.4                  | 101      | 9       | 97           | 4       | 98           | 3       |
| 8.5                  | 99       | 11      | 98           | 4       | 97           | 4       |
| 9.8                  | 98       | 15      | 102          | 5       | 98           | 4       |
| 11.2                 | 99       | 11      | 102          | 4       | 99           | 4       |
| 12.9                 | 101      | 9       | –            | –       | 102          | 3       |
| 14.8                 | –        | –       | –            | –       | 102          | 4       |

Fig. 4. Low-pressure photoionization mass spectra of nitrogen and 10 ppbv ethylbenzene (a); linear responses of ethylbenzene daughter ions (m/z 91) under two calibration approaches in nitrogen (b).
the ultrasensitive detection ability, accuracy, precision, and stability of the LPPI-MS, it is anticipated to be widely applicable in laboratory medicine, environmental monitoring, drugs, foodstuffs, and other promising research fields.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.aca.2017.01.065.

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