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Detection of Surface-Linked Polychlorinated Biphenyls using Surface-Enhanced Raman Scattering Spectroscopy

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Highlights

- Analytical detection of toxic polychlorinated biphenyls (PCB) was possible using surface-enhanced Raman scattering (SERS) spectroscopy.
- PCB77 compounds were chemically modified with a –SCH3 (PCB77-SCH3) group in order to improve the detection limit of PCBs.
- Our results point towards more efficient path for detecting different PCB congeners from real-life samples.

Abstract

We present an improved procedure for analytical detection of toxic polychlorinated biphenyls (PCB) using surface-enhanced Raman scattering (SERS) spectroscopy. A gold-capped silicon nanopillar substrate was utilized to concentrate PCB molecules within an area of high electromagnetic fields through formation of microsized nanopillar clusters, and consequently, so-called “hot spots” can be formed. In order to improve PCB detection limit, 3,3',4,4'-tetrachlorobiphenyl (PCB77) compounds were chemically modified with a –SCH3 (PCB77-SCH3) group. Experimental and numerical analysis of vibrational modes showed only minor differences between standard PCB77 and PCB77-SCH3. Consequently, we observe significantly increased SERS signals for –SCH3 modified PCB77 while retaining most vibrational modes that characterize standard PCB77. Results point towards more efficient path for detecting different PCB congeners from real-life samples. We interpret the result as PCB77-SCH3 link to gold surface via sulfur atoms that facilitates accumulation of the modified PCB molecules on the metal surface. For similar SERS experimental conditions most spectral characteristics of PCB77 are identifiable down to concentrations of ~10^{-5} M while PCB77-SCH3 spectral fingerprint is retained in ~10^{-8} M range.
Keywords: Polychlorinated biphenyls, PCB, toxic molecules, surface-enhanced Raman scattering, SERS detection, hot spots, gold nanopillars

1. Introduction

Polychlorinated biphenyls (PCBs) are a group of organic compounds that were manufactured by industry back in 1926 [1]. Today it is well known that PCBs cause a number of harmful effects [2] including cancer [3]. Although this toxic group of chemicals was banned in the 1970s, they are still frequently found worldwide in environment [4, 5], animal-source food [6] and natural waters [7].

Conventional PCB detection methods are usually based on gas chromatography (GC) [8, 9] or mass spectroscopy (MS) [10] with reported detection limits for single PCB congeners down to 0.05-0.1 ngL\(^{-1}\) [11]. Even though these methods display excellent PCB discrimination and detection capabilities, they require considerable effort and time.

Recent progress towards new PCB detection schemes include electrochemical impedance sensing using a single-walled carbon nanotube [12], electrochemical magneto-immunosensor [13], changes in surface photovoltage of porous ZnO [14], assay based on immuno-polymerase chain reaction [15], attenuated total reflection Fourier-transform infrared (ATR-FTIR) spectroscopy [16], fluorescence [17, 18] and surface plasmon resonances (SPRs) in thin gold films [19, 20]. In terms of detection speed, ease of handling and reliability there is still room for better PCB sensors. The aforementioned techniques have either practical limitations or target pure PCB congeners instead of realistic PCB mixtures compared to more powerful GC/MS methods [9]. On the other hand, the latter techniques require considerable effort, time and have limited mobility. Chobtang and co-workers have specifically outlined the need for real-time and on-site PCB sensing systems in order to make timely decisions [21].
Surface-enhanced Raman scattering (SERS) [22, 23] spectroscopy has extraordinary potential to discriminate target species down to single molecule level [24]. A number of studies have demonstrated that SERS-based detection of PCBs is feasible using silver nanorods [25], dendrites [26, 27], nanoparticle decorated ZnO nanorods [28] and nanosheet-assembled micro-hemispheres [29]. Zhou et al. showed that PCBs can be detected by SERS using Ag nanorods down to $10^{-8}$ mol/L [30]. However, since all PCB SERS studies utilize pure, isolated PCB compounds results might be different when dealing with real-life samples. It is therefore important to also focus on techniques that allow to filter or concentrate PCBs on the metal surface. For example, Bantz and co-workers have utilized alkanethiol space-layers to concentrate PCBs at the surface of a SERS substrate [31]. Although partition-layers introduce background signal, the authors argued that PCB detection and discrimination can be significantly improved. Zhu and co-workers [32] have also utilized decanethiol self-assembled monolayers for detection of PCB77 and specifically outlined that the detection limit can be further improved if better bonding of PCB77 to the metal surface is achieved.

For SERS-based PCB detection methods concentration of PCB molecules in the vicinity of the noble metal surface is key [32]. The SERS enhancement factor approximately scales as $|E_p/E_0|^4$, where $E_p$ is the near-field generated by noble metal particles at the position of the molecular emitter, and $E_0$ is the electric component of the incident radiation. Since the SERS effect is a local phenomenon, large inelastic scattering signals can only be obtained when the target molecule is situated within narrow gaps between plasmonic particles - the so-called hot spots [33]. In this contribution, we suggest a procedure to effectively concentrate PCB molecules at a noble metal surface. In order to enhance affinity of PCB77 molecules towards nanostructured gold surface, the molecules were modified with a $\text{–SCH}_3$ group. We compare strikingly similar spectral characteristics of standard PCB77 and modified PCB77 molecules and analyze vibrational modes using density functional theory (DFT) computational modelling method, which is in good agreement with the experimentally recorded spectra. We investigate evidence for formation of PCB77-$\text{SCH}_3$ molecular layers on the Au metal surface using gold-capped silicon nanopillar SERS substrates. Results indicate that it is considerably easier to detect $\text{–SCH}_3$ modified PCB77 molecules compared to
standard PCB77 when acquiring SERS spectra under identical experimental conditions. We show that the PCB detection limit can be significantly improved, which is of high practical importance not only for detecting different PCB congeners using SERS, but also for the development of new types of label-free nanoplasmonic sensors.

2. Experimental

2.1. Synthesis of PCB77–SCH₃

First, we aim to employ a method to perform a controlled substitution of one –Cl by the –SCH₃ group. This is important in order to systematically evaluate the obtained molecule using both theoretical and experimental methods. Due to this, we synthesized a building block already containing this SMe-group and perform a coupling to obtain the PCB77-SCH₃. This molecule was isolated in pure form and structure was investigated and confirmed by NMR prior to Raman and SERS analyses. Modified PCB77 molecules in this work were synthesized using 4-chloro-3-methylthioiodobenzene and 3,4-dichloroboronic acid blocks, see illustration in Figure 1(a). 4 g – 0.0141 Mol 4-chloro-3-methylthioiodobenzene, 3 g – 0.0155 Mol 3,4-dichloroboronic acid and 4.1 g – 0.0388 Mol Na₂CO₃ were dissolved in 22 ml of water, 22 ml of methanol and 90 ml of toluene. After 10 min under N₂-atmosphere 0.9 g – 0.00078 Mol tetrakis(triphenylphosphin)palladium (Pd(PPh₃)₄) was added and stirred for 30 h at 80°C. After cooling down this mixture was poured into 200 ml of water and extracted 3x with ethyl acetate (EtOAc). Organic phase was washed with 5% NaHCO₃, water and NaCl solution. After drying over Na₂SO₄ and removal of all solvents the residue was purified by chromatography using silica gel and hexane as eluent to get 4,4’3’-trichloro-3-methylthiobiphenyl as white crystals. Sublimation at 88°C. Yield: 3.3 g = 77 %. ¹H NMR (CDCl₃): δ = 7.61 (d, J = 2.0 Hz, 1H), 7.51 (d, J = 8.3 Hz, 1H), 7.41 (d, J = 8.3 Hz, 1H), 7.36 (dd, J = 8.4, 2.1 Hz, 1H), 7.25 (s, 1H), 7.21 (dd, J = 8.4, 2.1 Hz, 1H), 2.53 (s, 3H) ppm.
In Figure 1(b) we suggest another, a more practical procedure that can be potentially used with real life samples. In this procedure the treatment of PCB77 with large excess of MeSNa under harsh conditions is shown. Note, it is not clear at this point how many and which chlorine atoms can be substituted. In order to establish this, one needs to perform a purification and analysis of the mixture which is beyond the scope of this work. We suggest the following process guidelines for working with real life samples: a mixture of e.g. 3,3,4,4-tetrachlorobiphenyl and MeSNa (~3.5:1) in sulfolane, heated up to ~110 °C for 10-20 h under N₂ atmosphere. After cooling down, the mixture can be poured into water and filtrated to get a crystalline powder.

2.2. Fabrication of SERS substrates

Gold metal coated silicon nanopillars were fabricated following the protocol in ref. [34]. The SERS substrate is produced using maskless lithography and the fabrication process essentially involves only two fabrication steps. First, silicon nanopillars were produced using SF₆ and O₂ reactive ion etching process. After the silicon etching cycle, the surface was further treated with the O₂-plasma to remove fluorine and sulfur related products [41]. Second, 200 nm thick gold film was deposited using electron beam evaporation and the surface exhibited dark red color. Inspection in a scanning electron microscope (SEM) showed that the SERS substrate consisted of ~500 nm in height gold-capped nanopillars that are roughly perpendicular to the silicon surface, see Figure 4(c). The gold cap is approximately ellipsoidal with average dimensions a ≈ 300 nm and b ≈ 100 nm in height and width, respectively.

2.3. UV-Vis, Raman and SERS measurements

PCB No 77 compound (3,3’,4,4’-tetrachlorobiphenyl, analytical standard) utilized in all the PCB77 related experiments was purchased from Sigma-Aldrich. The UV-Vis absorption spectra of PCB77 and PCB77-SCH₃ compounds were recorded in the range of 200-400 nm using Shimadzu UV-1800 UV-Vis spectrophotometer. Both PCB samples were prepared in isopropanol (IPA, C₃H₈O) in the concentration of 10 µM. A baseline correction was made for both sample and reference cells, and the obtained UV-Vis spectra were normalized to unity for comparison, see results in Figure 1(c).
Raman and SERS measurements were performed using Thermo Scientific DXR Raman Microscope. An optical microscope is coupled to a single grating spectrometer that delivers 5.0 cm\(^{-1}\) FWHM spectral resolution and 2.0 cm\(^{-1}\) accuracy. Frequency-stabilized single mode diode laser operated at 780 ± 0.2 nm. All solid-state Raman and SERS spectra were recorded using 10x long working distance objective and laser powers of 5 mW (~70 kW/cm\(^2\)) and 0.1 mW (~1.3 kW/cm\(^2\)), respectively. Signal collection time was set to 5 seconds, unless stated otherwise. Solid-state Raman spectra were recorded from randomly oriented powdered samples deposited on a glass slide. For SERS measurements both PCB77 and PCB77-SCH\(_3\) compounds were dissolved in IPA using magnetic stirring (12 h). The PCB concentrations ranged from 5 \(\times\) 10\(^{-3}\) down to 5 \(\times\) 10\(^{-8}\). The SERS substrates were then immersed into different PCB solutions of varying concentrations for 3 h, rinsed with IPA and left for drying.

IPA solvent drying groups Au nanopillars together into microsized clusters which is important for obtaining high SERS signals [34], see schematic illustration and corresponding SEM images in Figure 4. Since larger Au nanopillar clusters increase SERS signal intensity [41], IPA solvent was found to be more effective in clustering Au nanopillars compared to e.g. tetrahydrofuran (THF) which is an organic liquid with lower viscosity. However, PCB77 molecules were also dissolved in THF for comparison (1 mM) and the recorded SERS spectral features were similar to the ones obtained using IPA.

2.4. Computational methods

All numerical calculations were performed using the program package Gaussian 09 [36]. The vibrational frequencies and Raman intensities were obtained numerically. The ground-state molecular structures were optimized utilizing density functional theory (DFT) calculations using the hybrid functional B3LYP [35, 37, 38] and the standard 6-31G* basis set. In order to correct for systematic errors, all computed frequencies were scaled by a factor of 0.9614 according to ref. [39]. The GaussSum 3.0 program was used to produce theoretical spectra [40].

3. Results and discussion
3.1. UV-Vis absorption and Raman scattering of PCB77 and PCB77-SCH₃

The obtained UV-Vis absorption spectra of PCB77 and modified PCB77-SCH₃ are shown in Figure 1(c). Both PCBs display characteristic UV-Vis spectra dominated by a main-band at \( \lambda_{1,\text{max}} = 213 \) nm which is attributed to resonance in the aromatic ring, i.e. the benzenoid skeleton, and is generally observed for all 209 PCB congeners [42]. The second strong absorption band found at \( \lambda_{2,\text{max}} = 261 \) and 288 nm for PCB77 and PCB77-SCH₃, respectively. The band is attributed to conjugation between the phenyl groups and the position of the band varies for different PCB congeners. The ~30 nm shift is consistent with UV-Vis spectral changes observed for adjacent PCB congeners, e.g. compare absorption spectra of PCB76 and PCB77 in ref. [42].

Before we analyze vibrational modes of modified PCB77 molecule, it is worthwhile to identify six distinct experimentally observed bands of PCB77 in the 1600-600 cm\(^{-1}\) spectral region. The calculated and experimental Raman spectra of standard PCB77 are shown in Figure 2(a) and the insets show molecular structures. Results are in good agreement with the previously published data [31]. The 400-700 cm\(^{-1}\) spectral region mainly contains Raman modes associated with C-Cl stretch and relative motions of two benzene rings, where the strongest band is at 676 cm\(^{-1}\). Based on our calculations, this experimentally observed band is dominated by deformation of benzene rings and C-Cl bond stretching. The 1030 cm\(^{-1}\) band is related to motions of benzene rings (ring breathing), the 1136 and 1245 cm\(^{-1}\) are dominated by C-H bending motions, and 1297 cm\(^{-1}\) band is attributed to biphenyl C-C bridge stretching motion. The strongest vibrational mode at 1597 cm\(^{-1}\) is related to symmetric in-plane stretching of benzene rings, see calculated vibrational amplitudes and frequencies in Figure 3.

Whereas PCB77 molecule has been studied experimentally and theoretically [31, 41], to the best of our knowledge there are no previous Raman investigations of –SCH₃ modified PCB77 that enables link to the noble metal surface via sulfur atoms. As can be seen from Figure 2(b), the experimental solid state spectrum of PCB77–SCH₃ shows no pronounced differences in comparison to standard PCB77. In the 1600-400 cm\(^{-1}\) spectral region all vibrational modes characteristic to PCB77 are preserved. One noticeable change is the reduced relative intensity of the Raman band at 1297 cm\(^{-1}\) (biphenyl C-C bridge stretching) which is also
well reflected in the calculated spectrum. In the 1600-400 \text{ cm}^{-1} range, positions of the most intense Raman bands of PCB77 and PCB77-SCH$_3$ are within ~3 \text{ cm}^{-1}, Figure 2. Our calculations show that the 400-700 \text{ cm}^{-1} region of PCB77-SCH$_3$ molecule is dominated by deformation of benzene rings, C-Cl and C-S (643 \text{ cm}^{-1}) bond stretching motions. Specifically, the experimentally observed 677 \text{ cm}^{-1} band is related to deformation of one benzene ring and C-Cl bond stretching, see Figure 2(b). In case of PCB77 molecule, the 676 \text{ cm}^{-1} mode involves deformation of both benzene rings and C-Cl stretching. The 1031 and 1134 \text{ cm}^{-1} bands are attributed to stretching motions of benzene rings, and the atomic displacement in these modes is similar to the standard PCB77. The 1242 and 1297 \text{ cm}^{-1} vibrational patterns are also comparable to PCB77, and are dominated by C-H wagging and biphenyl C-C bridge stretching motions. Finally, the most pronounced vibrational band at 1595 \text{ cm}^{-1} is attributed to symmetric stretching of benzene rings, and the motion is particularly dominated by one benzene ring, see summarized data in Figure 3(c).

3.2. SERS of PCB77 and PCB77-SCH$_3$

In Figure 4(a)-(b) the SERS detection principle utilizing gold-capped silicon nanopillars for standard and -SCH$_3$ modified PCB77 is schematically illustrated. In the PCB77 case, the metal molecule link is not expected to occur, and therefore microsized clusters of Au nanopillars encapsulate fewer random PCB77 molecules, Figure 4(a). SERS spectra of PCB77 are shown in Figure 5. The largest difference due to the presence of the gold metal surface was observed for the Raman mode at 1297 \text{ cm}^{-1}. In the case of SERS measurements, the mode is now found at 1280 \text{ cm}^{-1} (~17 \text{ cm}^{-1} shift), see Figure 3(c) for comparison. The SERS measurements using different concentrations of PCB77 ranging from $5 \times 10^{-3}$ to $5 \times 10^{-6}$ M show that the Raman spectral characteristics diminish and are essentially not visible in the \text{µM} concentration range. Due to low solubility of PCB77 in IPA, we have verified the result using a moderately polar THF solvent [31] that can be used to effectively dissolve PCB77. SERS spectra of PCB77 and detection limit using THF were similar to the result shown in Figure 5.

For the modified PCB77 case, the -SCH$_3$ group is expected to improve the PCB affinity towards the Au metal film, see schematic illustration in Figure 4(b). An increased SERS signal can be observed because
higher number of PCB molecules are now concentrated within the high electromagnetic field area, i.e. close to the Au metal surface. The SERS spectra of PCB77-SCH$_3$ with concentrations varying from $5 \times 10^{-3}$ down to $5 \times 10^{-8}$ M are presented in Figure 6 and are very similar to the solid state vibrational pattern shown in Figure 2. Contrary to the PCB77 case, the entire molecular fingerprint of PCB77-SCH$_3$ is preserved down to the $\sim 10^{-8}$ M regime. Chemical stability of the modified PCB77 molecule on the gold surface was verified by monitoring SERS spectra for 3 min prior to acquiring the signal. No significant signs of transient effects similar to the ones observed e.g. for 2,2′:5′,2″-terthiophene [34,37] were identified.

Comparison of the scattering intensity evolution of most intense Raman bands for standard (1589 cm$^{-1}$) and modified PCB77 (1586 cm$^{-1}$) are shown in Figure 7. The 1586 cm$^{-1}$ PCB77 band is rapidly decreasing in intensity and the detection limit for current experimental conditions is $\sim 10^{-6}$ M. For the same concentration range the PCB77-SCH$_3$ Raman band displays $\sim 4$ times higher intensity and all 6 characteristic vibrational modes are clearly identifiable, see Figure 6. A slight dip in the SERS intensity is observed for modified PCB77 at $\sim 10^{-3}$ M, see Figure 7. This is probably due to a reduced Au NP leaning which decreases the SERS signal intensity. Similar effect was observed for multilayer folic acid surface coverages using Ag NP structures [43]. Results from Figure 7 indicate that by modifying PCB77 with –SCH$_3$ group the detection limit can be significantly improved, and is close to $\sim 10^{-8}$ M which is due to a more effective concentration of PCB molecules on the noble metal surface.

4. Conclusions

According to the present study, an improved procedure based on a gold-capped silicon nanopillar substrate could be successfully employed for detecting toxic polychlorinated biphenyls (PCB) using surface-enhanced Raman scattering (SERS) spectroscopy. Upon binding of S-CH$_3$ modified PCB, the Au-PCB77 interaction significantly increases the SERS signal intensity in comparison to unmodified PCB77. We interpret this as an effective concentration of analyte molecules within the high electromagnetic field area, i.e. hot spots. A complete molecular fingerprint of PCB77-SCH$_3$ was preserved down to $\sim 10^{-7}$ M. The
estimated detection limit is $\sim 10^{-8}$ M while using the same experimental setting the detection limit for standard PCB77 was $\sim 10^{-6}$ M.

The detection limit of standard PCB77 can of course be further improved by optimizing the experimental conditions or employing SERS signal mapping techniques [40]. However, these kind of techniques should also further improve the detection limit of PCB77-SCH$_3$ in comparison to PCB77. Herein we employed a simple experimental set-up for comparison and to demonstrate straightforward PCB detection benefits when forming a PCB-metal link. It is of practical importance to develop on-site PCB detection method capable of concentrating different PCB types because real-life samples contain mixtures of PCB congeners (209 in total). We suggest one PCB modification procedure that can be adopted to simultaneously target different PCB congeners in real-life samples, see process guidelines in the experimental methods section and Figure 1(b). Real-life PCB-containing samples can have several appearances. Due to this an individual workup is required and solids must be powdered. For example, rubberlike material can be powdered using liquid nitrogen treatment, and in general all samples should be extracted with hydrocarbons e.g. in a Soxhlet apparatus with hexane. Removal of solvent gives crude material which can be treated with MeSNa without any further purification. The method could potentially be used on-site to extract different PCBs from relevant samples prior to SERS measurements. The SERS measurements could be performed using handheld Raman spectrometers that progressively are becoming smaller, better and cheaper [44].

Acknowledgments

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References


Figure captions

Fig. 1. (a) Illustration of PCB77-SCH$_3$ synthesis steps. In (b) an alternative method for working with real-life samples is suggested. PCB77 reaction with sodiummethanethiolate is shown where one or more chlorine groups can be substituted with –SCH$_3$ groups.

(c) UV-Vis absorption spectra of PCB77 (10$^{-5}$ M) and PCB77-SCH$_3$ (10$^{-5}$ M), obtained using the method shown in (a)).

Fig. 2. Experimental and calculated Raman spectra for PCB77 (a) and –SCH$_3$ group modified PCB77 molecules, (b). Insets show the corresponding molecular structures.

Fig. 3. Comparison of calculated vibrational frequencies (cm$^{-1}$) and amplitudes for the observed Raman active PCB77 and PCB77-SCH$_3$ modes, (a) and (b) respectively. Optimized molecular structures were calculated at the B3LYP/6-31G$^*$ level. In (c) the most intense Raman modes of PCB77 and PCB77-SCH$_3$ in the solid state, SERS and in theory are shown.

Fig. 4. Schematic comparison of the SERS-based PCB detection principles utilizing ~500 nm tall gold-capped Si nanopillars on 200 nm thick gold film. Average pillar head dimensions are ~300 and ~100 nm in height and width, respectively. Solvent drying forms nanopillar clusters and encapsulates standard PCB77, (a), and –SCH$_3$ modified PCB77 molecules, (b). In (c) representative SEM images illustrate the clustering step of gold-capped silicon nanopillars.

Fig. 5. Representative SERS spectra of PCB77 at concentrations of 5x10$^{-3}$ – 5x10$^{-6}$ M. The spectra were recorded for 5s at a laser power of 0.1 mW. All spectra have been vertically offset.

Fig. 6. Representative SERS spectra of gold metal surface bonded PCB77-SCH$_3$ at concentrations of 5x10$^{-3}$ – 5x10$^{-8}$ M. The spectra were recorded for 5s at a laser power of 0.1 mW. All spectra have been vertically offset.
**Fig. 7.** Comparison of integrated intensities of Raman active modes at varying concentrations for PCB77 (1589 and 1137 cm\(^{-1}\)) and PCB77-SCH\(_3\) (1586 and 1128 cm\(^{-1}\)). Each data point corresponds to an averaged intensity from 10 different measurement positions. All spectra were recorded for 5s at a laser power of 0.1 mW.
Figure 1

(a) 

(b) 

110 °C under N₂ gas

(c) 

[Graph showing absorbance vs. wavelength with peaks at 213 nm, 261 nm, and 288 nm]
Figure 3

(a) PCB77

(b) PCB77-SCH₃

(c)

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