Multifunctional Quaternary Phosphorus/Bromoargentate Hybrids: The Achievement of Greenish Blue Luminescence, Repeatable Photocurrent Responses and Durable Antimicrobial Activities with Enhanced Water Stability

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Background: The realization of multifunction in one bulk material is fascinating for developing a new generation of devices. Quaternary phosphorus salts were seldom utilized as templates in halogen systems, and the hybridization of alkyl(triphenyl)phosphonium with halometallate will be a good strategy for the development of multifunctional material, especially for biological material.

Methods: Under the template of (triphenyl)phosphonium-based quaternary phosphorus salts with different spacer lengths (n=2, 3, 4), three bromoargentate hybrids were constructed via the solution method, i.e., (1,2-DBTPP)(AgBr) (1), {[1,3-DBTPP](AgBr)}CH₃CN·H₂O·n (2), and {[1,4-DBTPP](AgBr)}CH₃CN·H₂O·n (3), 1,2-DBTPP²⁻=butane-1,2-diylbis (triphenyl)phosphonium, 1,3-DBTPP²⁻=propylene-1,3-diylbis (triphenyl)phosphonium, 1,4-DBTPP²⁻=butene-1,4-diylbis (triphenyl)phosphonium).

Results: The (Ag₄Br₁₆)n⁻ chain in 2 is a new type of 1-D bromoargentate chain constructed from cubane-like Ag₄Br₄ nodes, AgBr₄ tetrahedrons and AgBr₃ triangles. Interestingly, by elongating spacer n from 2 to 4, argentophilicity interactions are weakened, and the hydrogen bonds are strengthened. Consequently, their water stabilities and photocurrents are improved, in which the Ag₄d/Br⁻⁴p to π* anti-bonding orbital of the quaternary phosphorus transfer is facilitated. Furthermore, the greenish blue emissions can be detected. Finally, high inhabitation rates against Streptococcus mutans and Candida albicans can be observed in 2 and 3.

Conclusion: In all experiments, by elongating the spacer lengths of quaternary phosphorus salts, multifunctions were integrated in the quaternary phosphorus/bromoargentate hybrids, including greenish blue luminescence, repeatable photocurrent responses and durable antimicrobial activities with enhanced water stability. This work could provide a theoretical guide for the design of new biologically multifunctional materials.

Keywords: quaternary phosphorus, bromoargentates, nanoparticles, luminescence, antimicrobial

Introduction
In the past decades, halometallate-based functional materials have attracted great attention due to their versatile structure and unique properties, based on which many novel applications have emerged, such as perovskite solar cells, luminescence, molecular switches, nonlinear-optic devices and chromisms. Among the
halometallate hybrids, $d^{10}$ silver(I) halides have captured special interest for the following reasons: first, various anion dimensionalities based on rich coordination geometries of the Ag(I) ion (from linear MX$_2$, triangular MX$_3$, tetrahedral MX$_4$, and octahedral AgX$_6$) and variable connection modes; second, unique physical properties of electron-rich haloargentates.\textsuperscript{8} To date, in the haloargentate-based hybrids, the templates include organic cations, metal ions and metal-organic complexes.\textsuperscript{9,10} In the organic templated haloargentate hybrids, most of them are N/S-bearing organic cations with different ligand spacers and terminal substituent groups and hydrophobicity. For example, the increased number of methyl substituents in the organic templates can result in more inorganic/organic interactions, which is favorable for the formation of complicated haloargentate chains.\textsuperscript{11} However, quaternary phosphorus salts were seldom utilized as templates in haloargentate system, and only limited haloargentate/quaternary phosphorus hybrids were reported.\textsuperscript{12-14} The hybridization of alkyl(triphenyl)phosphonium with halometallate will be a good strategy for the development of multifunctional material.\textsuperscript{15} First, the tetrahedral geometry of alkyl(triphenyl)phosphonium cations limits benzene-benzene $\pi-\pi$ stacking interactions in the lattice; consequently, long persistence phosphors might be achieved. Second, in alkyl(triphenyl)phosphonium/halometallate hybrids, the LUMOs were commonly localized on the phosphate ligands, while the HOMOs were occupied by halometallates.\textsuperscript{16-18} The cationization of phosphate ligands can promote the LUMO energy levels. Third, the alkyl modification on phosphate ligands can improve the stabilities, which will obviously decrease the fabricating costs and elongate the working life of related devices.\textsuperscript{19,20} Finally, quaternary phosphorus salt and silver-bearing compounds are traditional antibacterial materials, their hybridization will help improve antibacterial performance.\textsuperscript{21,22} In this work, alkyl-bis-(triphenyl)phosphoniums with different spacer lengths (spacer $n=2$, 3, 4) were used as organic templates to obtain three new bromoargentate hybrids, i.e., (1,2-DTPB)AgBr$_2$ (1), [(1,3-DTPB)$_2$(Ag$_2$Br$_1$)]$_1$CH$_3$CN$_2$H$_2$O$_n$ (2), and [(1,4-DTPB)(AgBr)$_2$][CH$_3$(CN)$_2$H$_2$O]$_n$ (3). (1,2-DTPB)$_2$=ethane-1,2-diylbis (triphenyl)phosphinum, 1,3-DTPB$^2$=propane-1,3-diylbis (triphenyl)phosphinum, 1,4-DTPB$^2$=butane-1,4-diylbis (triphenyl)phosphinum). Their greenish blue luminescence, repeatable photocurrent responses and durable antibacterial activities correspond to different spacer lengths, which could provide a theoretical guide for the design of new multifunctional materials.

**Experimental Section**

**Materials and Physical Measurement**

All chemicals were commercial products and used without further purification. Alkyl-bis-(triphenyl)phosphonium (1,2-DTPB)Br$_2$, (1,3-DTPB)Br$_2$, and (1,4-DTPB)Br$_2$ were self-prepared. IR spectra were recorded on a Perkin-Elmer Spectrum-2000 FTIR spectrophotometer (4000–400 cm$^{-1}$) with powdered samples spread on a KBr plate. Elemental analyses for C, H and N were performed on a Vario MICRO elemental analyzer. Powder XRD patterns were obtained using a Philips X’Pert-MPD diffractometer with CuK$\alpha$ radiation ($\lambda=1.54056$ Å). The absorption spectra were measured on a Perkin-Elmer Lambda 900 UV/VIS spectrophotometer equipped with an integrating sphere at 293 K, and BaSO$_4$ plates were used as a reference. The optical gaps were calculated from reflectance spectra using the Kubelka–Munk function, $\alpha/s=(1-R)^2/2R$, where $\alpha$ is the absorption coefficient, $s$ is the scattering coefficient, and R is the reflectance.\textsuperscript{23} $^1$H NMR and $^{13}$C NMR spectra were recorded on a Bruker AVII 400 spectrometer whose chemical shifts were reported in parts per million (ppm) downfield from tetramethylsilane. Fluorescence spectra were carried out on an Edinburgh FLS 980 Red PMT. The photocurrent experiments were performed on a CHI660 electrochemistry workstation with three-electrode systems using a Pt plate as the counter electrode and an Ag/AgCl electrode (3 M KCl) as the reference electrode. The morphologies of the devices were investigated by field emission scanning electron microscopy (SEM) spectrometer (Verios G4 UC), and the FE-SEM EDX is recorded by the EDAX Inc. Octane Elect Super.

**Synthesis**

**Synthesis of Quaternary Phosphorus Salts**

Three alkyl-bis-(triphenyl)phosphonium bromides ($n=2$: (1,2-DTPB)Br$_2$, $n=3$: (1,3-DTPB)Br$_2$, $n=4$: (1,4-DTPB)Br$_2$) were prepared by a one-step alkylated reaction of triphenylphosphine with 1,2-dibromoethane, 1,3-dibromopropane and 1,4-dibromobutane in the DMF solvent according to the literature (Scheme S1).\textsuperscript{24} The detailed synthesis process, $^1$H NMR and $^3$P NMR spectra can be seen in Figure S1 in the supplementary data.

**Synthesis of (1,2-DTPB)(Ag$_3$Br$_4$) (1):** 1 was prepared by solution evaporation method. (1,2-DTPB)Br$_2$ (0.0768 g, 0.1 mmol) and AgBr (0.0183 g, 0.1 mmol)
were dissolved in 20 mL acetonitrile with the suspension obtained. Afterwards, a small amount of NaBr solid was added slowly until the solution became clear. In addition, the resultant solution was kept stirring for 3 h at room temperature. The resultant solution was filtered, and the filtrate liquor was covered with a cling film at room temperature for slower evaporation. Colorless transparent block crystals were obtained after the growth period of 5 days. Yield: 40.3% (0.0210 g, based on Ag). Analy. Cald. for C_{33}H_{44}Ag_{2}Br_{5}P_{2} (1087.93): calcd. C 41.92, H 3.13%; found C 41.72, H 3.16%. IR (cm⁻¹): 3050(w), 2855(w), 1580(w), 1476(m), 1098(s), 996(w), 733(s), 693(s), 490(s).

Synthesis of {(1,3-DBTPP)₂(Ag₂Br₅)}(CH₃CN-H₂O)ₙ (2): The synthesis process of 2 was similar to that of 1, except (1,3-DBTPP)Br₂(0.0726 g, 0.1 mmol) was used as starting material. Colorless transparent block crystals were obtained after 5 days. Yield: 51.2% (0.0206 g, based on Ag). Analy. Cald. for C_{36}H_{73}Ag₂Br₁₁NOP₄ (2824.26): calcd. C 34.02, N 0.49, H 2.60%; found C 34.34, N 0.54, H 2.66%. IR (cm⁻¹): 3060(w), 2920(w), 1580(w), 1493 (w), 1440(m), 1110(s), 992(w), 719(s), 690(s), 500(s).

Synthesis of {(1,4-DBTPP)₂(Ag₄Br₉)}(CH₃CN-H₂O)ₙ (3): 3 was prepared using the same procedure, except that (1,4-DBTPP) Br₂ (0.0837 g, 0.1 mmol) was used as the starting material. Colorless crystals were obtained after 6 days. The SEM diagrams of 1–3 can be seen in Figure S2. Yield: 38.5% (0.0139 g, based on Ag). Anal. Cald. for C_{44}H_{66}Ag₄Br₉N₂OP₂ (1779.48): calcd. C 29.70, N 1.57, H 2.61%; found C 30.36, N 1.63, H 2.71%. IR (cm⁻¹): 3060(w), 2920(w), 1580(s), 1443(s), 1120(s), 995(w), 854 (w), 720(w), 688(s), 490(s).

Electrode Preparation and Photocurrent Measurement

The solution coating method was adopted to prepare the electrodes of 1–3 in the photocurrent measurements. The freshly synthesized powders (5 mg) were dissolved in DMF (0.3 mL) to obtain suspensions, which were dispersed evenly to produce slurries. The slurries were then spread onto precleaned ITO glasses (0.6×0.6 cm, 14 Ω per cm²), whose side parts were previously covered using scotch tape. The working electrode was dried overnight under ambient conditions. A copper wire was connected to the side part of the working electrode using conductive tape. Uncoated parts of the electrode were isolated with epoxy resin. A 150 W high-pressure xenon lamp, located 15 cm away from the surface of the ITO electrode, was employed as a full-wavelength light source. The photocurrent experiments were performed on a CHI660 electrochemistry workstation in a three-electrode system, with the sample-coated ITO glass as the working electrode mounted on the window with an area of 0.25 cm², a Pt wire as the auxiliary electrode, and a Ag/AgCl electrode as the reference electrode. The supporting electrolyte solution was a 0.2 mol·L⁻¹ sodium sulfate aqueous solution. The applied potential was 0.5 V for all measurements. The lamp was kept on continuously, and a manual shutter was used to block exposure of the sample to the light. The sample was typically irradiated at intervals of 10 s.

Antimicrobial Activity on Streptococcus mutans and Candida albicans

A Streptococcus mutans (S. mutans) strain (ATCC, 700610) and a Candida albicans (C. albicans, ATCC, 90029) strain, both from the American Type Culture Collection, were selected as the model pathogens for the antimicrobial activity experiments. After recovery on Brain Heart Infusion (BHI, for S. mutans) agar or Sabouraud dextrose agar (Sda, for C. albicans) plates, the microbes were cultured in 4 mL of BHI broth medium (for S. mutans) or Yeast Extract Peptone Dextrose (YPD) medium (for C. albicans) at 37°C for 24 h. For colony formation assay, 20 µL of microbial solution containing approximately 1 × 10⁴ colony-forming units per mL (CFU/mL) of S. mutans or C. albicans was uniformly spread on a BHI agar plate or SDA plate, respectively, containing different concentrations of compounds. After incubation at 37°C for 48 h, the plates were photographed, and the number of colonies was counted. For growth curve assay, 200 µL of the suspension of S. mutans (10⁴ CFU/mL) in BHI broth medium or C. albicans in YPD medium were added to each well of a 96-well plate, and different concentrations of the compounds were added to the wells. The growth of S. mutans or C. albicans was assessed using a Bioscreen C Automated Microbiology Growth Curve Analysis System (Turku, Finland) at 37°C under shaking at 200 rpm. The OD values at 600 nm were measured every hour for up to 36 or 48 h.

Computational Methods

First-principle density-functional theory (DFT) calculations were performed with the CASTEP program.\(^{26,27}\)
The exchange correlation functional was described by a generalized gradient approximation (GGA) with Perdew–Burke–Ernzerhof functional for solids (PBE) scheme. A $3\times2\times2$ Monkhorst-Pack grid with a total number of 6 k points in the irreducible Brillouin zone and 45 empty bands was adopted. The number of plane waves included in the basis was determined by a cutoff energy $E_c$ of 435.4 eV. The other parameters and convergence criteria were the default values of CASTEP code.

**X-Ray Crystallography**

The intensity data of 1–3 were collected on a Bruker APEX II diffractometer using graphite-monochromated MoKα radiation ($\lambda = 0.71073$Å) at room temperature, during which the $Lp$ factor corrections and multiscan absorption corrections were applied. The structure was solved by direct methods with SHELX-97 program and refined by full-matrix least-squares techniques on $F^2$ with SHELXL-97 program. For all nonhydrogen atoms, anisotropic refinements were carried out. All hydrogen atoms attached to carbon atoms were geometrically placed. Crystal parameters of three compounds are summarized in Table 1. Selected bond lengths and angles are listed in Table S1. π-π stacking interactions and C-H···π interactions are given in Table S2 and S3. Crystallographic data have been deposited with the Cambridge Crystallographic Data Center as supplementary publication number CCDC-1818452 (1), 1818423 (2), and 1590456 (3). These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via [www.ccdc.cam.ac.uk/data_request/cif](http://www.ccdc.cam.ac.uk/data_request/cif).

**Results and Discussion**

**Structural Description**

Compounds 1–3 are composed of bromoargentate anions ((Ag$_2$Br$_3$)$_2$)dimers for 1, (Ag$_7$Br$_{11}$)$_{4n}$ chains for 2 and (Ag$_9$Br$_7$)$_{2n}$ chains for 3) and corresponding alkylbis-(triphenyl)phosphonium cationic templates, among which hydrogen bonds contribute to the structural stabilization. The (Ag$_2$Br$_3$)$_2$ in 1 is defined by two edge-sharing AgBr$_3$ triangles, giving two terminal (Br(2), Br(2)$^{\#1}$, #1: -x, 1-y, -z) and two bridged Br ions (Br(1), Br(1)$^{\#1}$, Figure 1A). The distance of terminal bromine and silver (2.5356(11) Å) is slightly shorter than that of bridging atoms (2.6632(11) Å, Table S2). Furthermore, the parallelograms defined by four atoms Ag(1), Br(1), Ag(1)$^{\#1}$ and Br(1)$^{\#1}$ are coplanar. The (Ag$_2$Br$_4$)$_2$- dimers were seldom documented. The Ag-Ag distance of 3.0253(14) Å is shorter than the sum of the van der Waals radii of silver (3.44 Å), hinting at the presence of an argentophilic interaction. The configuration of the (1,2-DBTPP)$_2^+$ cation in 1 is unfolding based on the P(1)-(19)-(19)$^{\#1}$ angle (115.99°) (Figure 1B). C-H···Br hydrogen bonds between (1,2-DBTPP)$_2^+$ cations and (Ag$_2$Br$_4$)$_2^-$ dimers contribute to the formation of a 2-D layer (Figure 1C, Table S3). Due to the tetrahedral geometry of the PPh$_3$ moiety, no benzene-benzene π-π stacking interactions can be detected.

The (Ag$_2$Br$_{11}$)$_{4n}$ of 2 is a straight chain built from cubane-like Ag$_4$Br$_4$ nodes, AgBr$_4$ tetrahedrons and AgBr$_3$ triangles (Figure 2A), which has not been documented in bromoargentate systems. The distorted cubane-like Ag$_4$Br$_4$ core is an iso-structure of Ag$_4$I$_4$ cubane, which has been commonly observed in iodoargentates, for example, 1-D chains ([(APHEN-H)H$_2$(Ag$_4$I$_4$)$_{1n}$]). 2-D layers ([(PC)$_2$(Ag$_4$I$_4$)$_{1n}$]), and 3-D open frameworks ([BCP]$_2$ [Ag$_4$I$_4$])$_n$. In 2, each cubane-like Ag$_4$Br$_4$ core connects with two neighboring AgI$_4$ tetrahedrons via an edge-sharing mode (defined by Br(3)-Br(7)) to give a Ag$_8$Br$_{10}$ unit; furthermore, the Ag$_8$Br$_{10}$ unit links with the Ag(4)Br$_3$ triangle to generate a Ag$_8$Br$_{11}$ building block. Neighboring Ag$_8$Br$_{11}$ building blocks are extended into a 1-D chain via vertex (defined by Br(6))-sharing mode (Figure 2A). Consequently, two kinds of coordinated geometries of Ag centers can be found, including tetrahedral (Ag(1), Ag(2), Ag(3), Ag(5)) and planar triangle (Ag(4)). Specifically, the Ag(3)Br$_4$ tetrahedron is highly distorted with Ag-Br distances of 2.617(3)–905(3) Å. Abundant Ag···Ag interactions with separations of 3.026(3)–3.323(4) Å are observed, indicating the presence of strong argentophilic interactions (Table S2). The bond lengths of the (1,3-DBTPP)$_2^+$ cation are within normal range, and the bond angle in the methane (C(19)-C(20)-C(19)$^{\#1}$) is 107.16°, suggesting its folded configuration (Figure 2B). There are no π-π stacking interactions in the lattice, but C-H···Br hydrogen bonds contribute to the formation of a quasi-3-D network (Figure 2C). In addition, solvent molecules CH$_3$CN and H$_2$O also stack in the lattice.

**Table 1** PL Data of 1–3 at Room Temperature

| Compound | $\lambda_{ex}$ (nm) | $\lambda_{em}$ (nm) | $\Phi$ (%) |
|----------|---------------------|---------------------|------------|
| 1        | 320                 | 488                 | 6.24       |
| 2        | 350                 | 513                 | 6.21       |
| 3        | 340                 | 490                 | 6.26       |
The (Ag$_5$Br$_7$)$_n^{2n-}$ chain takes on a zigzag configuration based on much more complicated Ag···Ag interactions, which is the iso-structure of the (Ag$_3$I$_7$)$_n^{2n-}$ chain in [{Et$_3$N(CH$_2$)$_3$NEt$_3$}[Ag$_3$I$_7$]]$_n$. The building block is a Ag$_5$Br$_{10}$ unit in which all Ag centers adopt an Br$_4$ donor set and tetrahedral geometries. The Ag$_5$Br$_0$ building block could be described as the edge-sharing of five AgBr$_4$ tetrahedrons. Adjacent Ag$_5$I$_{10}$ building blocks are extended into a 1-D zigzag chain along the a-axis via an edge-sharing mode (Figure 3A). Strong Ag···Ag interactions could be found with distances ranging from 3.0289(13) to 3.1918(14) Å. In particular, a μ$_5$-Br(3) is found in this polymeric chain, which is seldom seen in haloargentate compounds. The configuration of (1,4-DTBP$^2$) is determined from the bond angles in methene (111.73 and 116.915°, Figure 3B). Similarly, C-H···Br hydrogen bonds contribute to the formation of quasi-3-D network (Table S3, Figure 3C), and additional CH$_3$CN and H$_2$O molecules stack in the cavities. This work again proves the rigidity/flexibility competition in haloargentates/organic hybrids, ie, when the spacer is short, the alkyl flexibility dominates in the structural stabilization, but when the spacer is longer, to some extent, the benzene rigidity plays a more important role. In this work, when spacer n=2, only simple (Ag$_5$Br$_{4}$)$_2$ dimmer is given, but when spacer n=3 and 4, more complicated (Ag$_5$Br$_{11}$)$_n^{4n-}$ and (Ag$_5$Br$_7$)$_n^{2n-}$ chains are produced.

**Stabilities in Water**

The stabilities of functional materials in solvents will determine their real applications. In this work, alkyl-bis-(triphenyl)phosphoniums with different spacer lengths might affect their water stability. Most of the functional properties of this work were conducted in water; therefore, their stabilities in water were studied. During the water stability measurements, the freshly prepared crystalline samples of 1–3 were ground as fine powders and soaked in deionized water at room temperature with different time...
intervals. Afterwards, the samples were centrifuged and dried at 40°C, and PXRD experiments were conducted to check their phases. The phase purities of bulk compounds 1–3 have been verified by powder X-ray diffraction (PXRD), in which the peaks of the as-synthesized samples are in good agreement with the simulated peaks (Figure 4). Furthermore, the water stability measurements under different soaking times (1, 3 and 5 days) imply that the crystal structures of 1–3 can persist after soaking in water for 3 days. After soaking for 5 days, 1 will collapse to some extent judging from its PXRD pattern (Figure 4A). However, 2 and 3 still maintain their structures (Figure 4B and C), implying that they have better water stabilities than 1. This trend suggests that the longer spacer lengths will result in better water stabilities, which might be induced by their better flexibilities when facing the attack of solvents. This phenomenon again proves the conclusion that in the quaternary phosphorus-containing hybrids, a longer alkyl chain can give rise to better water stability. In addition, the thermal stability of 3 was also determined, which could not decompose until 320°C (Figure S3). The good water/thermal stabilities will be beneficial for their long-term antibacterial performances.

Optical Diffuse-Reflection Spectra and Photoluminescence
Solid-state optical diffuse-reflection spectra of 1–3 were recorded from powder samples at room temperature (Figure 5A). They exhibit intense adsorption in the...
ultraviolet zone (281 nm for 1, 301 nm for 2 and 291 nm for 3). Comparably, the adsorption peaks of organic quaternary phosphorus salts generally appear at approximately 330 nm, corresponding to the \( n-\pi^*/\pi-\pi^* \) transitions of phenyl groups in the phosphonium core. Therefore, compared with free phosphonium, the corresponding electronic transitions of 1–3 can definitely be attributed to the ligand-centered \( n-\pi^*/\pi-\pi^* \) transitions of phosphine ligands. There are blanks in the range of 350–800 nm, suggesting the absence of ligand-to-ligand charge transfer (LLCT) and charge transfer between inorganic bromoargentates and organic moieties. These correspond to the absence of \( \pi-\pi \) and \( \text{C–H–Br} \) interactions in their lattices, as indicated by structural analysis. Solid-state UV–Vis diffuse spectra of 1–3 calculated from the diffuse reflectance data by using the Kubelka–Munk function is plotted in Figure S4. The band gaps can be estimated as 3.60, 3.55 and 3.70 eV, which indicate that 1 and 2 are wide-gap semiconductors and 3 is generally an insulator. The wide band gaps are commonly observed in organic/bromoargentate hybrids. Compared with the gap of bulk AgBr (2.81 eV), obvious blueshifts have occurred. More importantly, the conclusion could be drawn that in the phosphonium-containing hybrids, the gaps will be dominated by organic phosphonium cores.

One-photon photoluminescence spectra of 1–3 measured at room temperature in the solid state are shown in Figure 5B (excited, emission peaks and quantum yields are
summarized in Table 1). Generally, the three hybrids generated the same single greenish blue emissions in the solid state at room temperature when excited at 320–350 nm, with luminescent colors illustrated in the CIE-1931 profiles (Figure 5B-C). As reported by literature, free organic quaternary phosphorus iodide (Ph₃Pn-Bu·I) also exhibits cyan emission (at 485 nm) under excitation at 340 nm, which was attributed to the photoinduced charge transfer from the iodine anion to the phosphonium core.¹⁵ Compared with the emission of phosphonium iodide, compound 2 with spacer length n=3 presents the largest red-shift of approximately 28 nm, and conclusions could be

![Figure 4 The PXRD patterns of 1 (A), 2 (B) and 3 (C) soaked in water for different times.](image-url)
been conducted to study the photoinduced charge transfer from bromoargentates to phosphoniums. Moreover, their quantum yields are generally the same (ranging from 6.21 to 6.26%). Their photoluminescence implies their potential application as biological imaging or cancer therapy compounds.42,43

**Photocurrent Response Performances**

Photoinduced current generation can be potentially used in molecular switches, information storage, and so on, and has been detected in organic/metal halides hybrids.5,44 Photocurrent response performances have also been executed on 1–3 using typical methods, in which three electrode photo/electrochemical cell systems were adopted (a sample-modified ITO electrode as a working electrode, a Pt wire as an auxiliary electrode and a Ag/AgCl electrode as a reference electrode). All the experiments were conducted in Na2SO4 aqueous solutions under the illumination from a 150 W Xe arc lamp with on–off cycles (an on-off interval of 20 s). Under repetitive irradiation, repeatable photocurrents with rapid responses can be detected on all compounds (Figure 6A). The photocurrent of 1 and 2 first decreases and then stabilizes after five cycles, but the current of 3 is steady without decay. Their photocurrents stabilize at approximately 1.10×10^{-7} (for 1 and 2) and 1.30×10^{-7} A (for 3), which are weaker than those of viologen-containing compounds, such as (MV)$_2$[Li$_4$(L)$_2$(H$_2$O)$_6$] and MV[Ni(4-pedt)$_2$]$_2$.45 It is universally accepted that the pure organic-coated system did not show any photocurrent response,46 so the bromoargentate/phosphonium donor–acceptor systems are responsible for the photocurrent generation in this work. Consequently, its photoinduced current generation mechanism is given: first, the photosensitive phosphonium cations are irradiated to generate the (Ph$_3$-R-Ph$_3$)$_2$• radicals, and at the same time, Ag$_{6x}$Br$_y$(y-x)$^+$ donors also transfer electrons to phosphoniums with production of Ag$_{6x}$Br$_y$(y-x)$^+$ radicals. Second, the (Ph$_3$-R-Ph$_3$)$_2$• radicals transfer their electrons to the ITO electrodes with reproduction of the (Ph$_3$-R-Ph$_3$)$_2$• acceptor. As a result, an electron conductive pair of Ag$_{6x}$Br$_y$(y-x)$^+$–(Ph$_3$-R-Ph$_3$)$_2$• is responsible for their repeated currents (Figure 6B). This process can also be verified by the components of the tops of VBs (Ag-4d, Br-4p) and the bottoms of CBs (p-π anti-bonding orbital of the quaternary phosphorus) in the PDOS of 1 (Figure 6C). As indicated by this mechanism, the stronger C-H···Br hydrogen bond in 3 (the shorter H···Br distance of 2.74 Å, Table S3) can facilitate the electronic transfer; as a result, a relative higher current intensity in 3 can be found.

**Antimicrobial Performance Against S. mutans and C. albicans**

Silver, quaternary ammoniums are traditional antibacterial materials to inhibit or reduce the growth of detrimental bacteria.47 Streptococcus mutans (S. mutans) is a commensal microorganism found in the human oral cavity and the primary causative agent of dental caries.48 In this work, the antibacterial activity of 2 and 3 were tested against S. mutans according to the standard antibacterial test of insoluble antimicrobials, and the results are shown in Figure 7. Interestingly, the presence of 2 and 3 can effectively inhibit the growth of S. mutans. The antimicrobial rate can reach 90% for 2 and 68% for 3 when the bromoargentate hybrids’ concentrations are 100 μg/mL (Figure 7A and B), and the bacteria did not grow for 36 h with 40 μg/mL of 2 or 50 μg/mL of 3 (Figure 7C and D). The minimum inhibitor concentration (MIC) is approximately 50 μg/mL according to the growth curves, which is comparable to and even better than the MIC of currently accessible silver-loaded zeolites (50–150 μg/mL) and AgBr/NVP composites (approximately 100 μg/mL).49 This improved antibacterial performance of quaternary

![Figure 5](https://example.com/figure5.png)

*Figure 5 (A) Solid state electronic spectra; (B) emission spectra at room temperature with CIE-1931 chromaticity of 1–3. (C) CIE-1931 chromaticity diagram.*
phosphorus/bromoargentate hybrids can be assigned to the following aspects. First, the quaternary phosphorus moieties of hybrids can lead to the disruption of the cytoplasmic membranes of \textit{S. mutans}.\footnote{50} Second, the sustained and slow releases stemming from the exceedingly low solubility of bromoargentates in 2 and 3 are beneficial for efficient inhibition of bacterial growth and are consistent with their good solution stabilities.\footnote{51} The sustained release of biocidal Ag\textsuperscript{+} ions can deactivate cellular enzymes and DNA by coordinating electron-donating groups and causing pits in bacterial cell walls, and as a result, lead to increased permeability and cell death. \textit{Candida albicans} is one of the commonly seen yeasts in the human mouth and the main pathogen of denture stomatitis; the detection rate of dental plaque is 81.7\%.\footnote{52} In the present study, we also tested the antifungal activity of these compounds on \textit{C. albicans}. Interestingly, we found that compounds 2 and 3 showed strong antifungal activity, as shown in Figure 8. The inhibition rate was 80\% for 2 and 57\% for 3 when the bromoargentate hybrids’ concentrations were 100 μg/mL (Figure 8A and B), and the fungi did not grow for more than 40 h with 200 μg/mL of 2 or 3 (Figure 8C and D). These results indicated that the bromoargentate hybrids possess excellent antimicrobial activities by inhibition the growth of \textit{S. mutans} and \textit{C. albicans}.

**Conclusions**

In summary, bromoargentate hybrids directed by (triphenyl) phosphonium-based quaternary phosphorus salts with

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**Figure 6** (A) Photocurrent curves, (B) photocurrent response mechanisms and (C) bond structure (left), projected density of states (pDOS, middle) and frontier orbitals (right) for 1.
different spacer lengths (n=2, 3, 4) have been prepared. The \((\text{Ag}_7\text{Br}_{11})_{\text{n}}^{4-}\) chain in 2 is a new type of 1-D bromoargentate chain constructed from \(\text{Ag}_7\text{Br}_{12}\) units, but the zigzag \((\text{Ag}_5\text{Br}_7)_{\text{n}}^{2-}\) chain in 3 is based on \(\text{Ag}_5\text{Br}_9\) units. Interestingly, with the elongation of spacer \(n\) from 2 to 4, argentophilicity interactions are weakened, and the hydrogen bonds are strengthened. Consequently, better water stabilities and enhanced photocurrents are observed. Furthermore, high inhabitation rates against \(S.\) mutans and \(C.\) albicans can be observed. Overall, the elongation of the spacer lengths can give rise to better functional properties, which could provide a theoretical guide for the design of new multifunctional materials.

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**Disclosure**

The authors declare no competing financial interest.

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