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A novel self-assembled supramolecular sensor based on thiophene-functionalized imidazophenazine for dual-channel detection of Ag+ in an aqueous solution†

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Herein, a novel self-assembled supramolecular sensor (S1) based on thiophene-functionalized imidazophenazine for Ag^+ was designed and synthesized. It showed dual-channel detection properties for Ag⁺ based on the competitive mechanism of supramolecular self-assembly with high sensitivity and selectivity even in the presence of other metal ions. Its detection limit for Ag⁺ is 8.18 \times 10⁻⁹ M. Upon the addition of Ag⁺, the solution changes from yellow to light purple and the fluorescence is quenched. Furthermore, the sensing mechanism between Ag^+ and S1 is investigated via IR and ^{1}H NMR spectroscopy, mass spectrometry, and density functional theory calculations. PAPER

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Introduction

With the development of modern industry, increasing industrial emissions containing metal ions have caused serious water pollution, which affects food and agriculture and further threatens human health and the environment.¹ Among the important metal ions, silver has attracted extensive attention due to its wide applications in the fields of electricity, photography, imaging, and pharmacy.² In fact, trance amounts of silver ions can control bacteria breeding or rehabilitate and reconstruct essential human tissue.³ However, as is well known, excess silver ions can inactivate sulfhydryl enzymes; on the other hand, silver ions can combine with amine, imidazole, and carboxyl groups of various metabolites, such as high molecular weight protein and metallothionein, in the tissue of the cytosol fraction, resulting in harmful effects to humans.⁴ Thus, the maximum bearable concentration of $Ag(i)$ issued by the World Health Organization (WHO) is 0.5 μ M in drinking water.⁵

Considering the importance of $Ag(i)$ in industrial production and in daily life, the development of techniques for sensing and monitoring $Ag(i)$ is in great demand. There are various available methods to monitor Ag⁺, such as electrothermal atomic absorption spectrometry (ETAAS), voltammetry, inductively coupled plasma atomic emission spectrometry (ICP-AES), inductively coupled plasma mass spectroscopy (ICP-MS), and

potentiometry.⁶ However, most of them are expensive, timeconsuming, or unsuitable for on-site and real-time monitoring. Therefore, the development of fast, efficient, sensitive, and selective methods for the detection of $Ag⁺$ is necessary for both the environment and human health.⁷ In comparison, fluorescence techniques exhibit attractive advantages, such as high selectivity and sensitivity, real-time detection, practicalsimple sample preparation, and great potential in high throughput detection of $Ag⁺$ on-site. However, some of these probes suffer from serious drawbacks such as complicated synthesis procedures and poor solubility, sensitivity, and selectivity in aqueous media.⁸ Due to the efforts of researchers, numerous Ag⁺ sensors with good properties have been reported. However, most of them exhibit ultraviolet or fluorescent single channel recognition or are less selective.⁹ Moreover, it is worthwhile to note that sensors for the detection of $Ag⁺$ in aqueous systems are relatively scarce although $Ag⁺$ usually exists in water or sewerage. Consequently, the design and synthesis of colorimetric and fluorescent chemosensors with high sensitivity and selectivity for the quantification of $Ag⁺$ in aqueous media via a simple reaction is still an intriguing area of research.¹⁰

In view of this and based on our previous research in host– guest chemistry,¹¹ we designed and synthesized a sensor molecule IM (Scheme 1) based on thiophene-functionalized imidazophenazine via a simple reaction; this sensor could selfassemble into a supramolecular system and form a supramolecular chemosensor S1. This sensor combines the phenazine fluorescent signaling subunit, which has attracted extensive attention due to its excellent photophysical properties. In addition, among the different fluorogens, it is very sensitive to

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Scheme 1 Synthesis of the compound IM and the proposed possible mechanism of **S1** bonding with Ag⁺.

conformational changes.¹² Consequently, phenazines are ideal platforms for the development of cation, anion, and neutral molecule recognition platforms. Moreover, there are only a few reports on ion recognition in host–guest chemistry. Compared to some $Ag⁺$ sensors reported in previous studies, the $Ag⁺$ sensor reported herein is a self-assembled supramolecular sensor (S1). The $Ag⁺$ sensing mechanism is based on a competition between self-assembly of the sensor molecules and $Ag⁺$ coordination with the sensor molecules. This novel competitive mechanism provides the $S1$ sensor with better Ag⁺ detection sensitivity. The detection limit of **S1** for Ag⁺ is 8.18 \times 10⁻⁹ M, which is lower than that reported in some previous studies.⁸ Therefore, S1 is utilized as a supramolecular self-assembled sensor to study the recognition performance of Ag⁺.

Results and discussion

To investigate the $Ag⁺$ recognition abilities of the S1 sensor in an aqueous solution, we performed a series of host–guest recognition experiments. The recognition profiles of the chemosensor **S1** towards various metal ions, including Fe $^{3+}$, Hg $^{2+}$, Ag $^+$, Ca^{2+} , Cu^{2+} , Co^{2+} , Ni^{2+} , Cd^{2+} , Pb^{2+} , Zn^{2+} , Cr^{3+} , and Mg^{2+} , were primarily investigated using UV-vis spectroscopy and fluorescence spectroscopy in a DMSO/water (v: $v = 1 : 1$, buffered with HEPES $pH = 7.2$) solution.

As shown in Fig. 1, the sensor immediately responded with obvious color changes (from yellow to light purple) when $Ag⁺$ was added to the S1 solution at room temperature. In the corresponding UV-vis spectra, the strength of the absorption peak at 413 nm decreased, and the peak red-shifted to 376 nm (Fig. $S1\dagger$). However, yellow fluorescence with one emission band centered at 547 nm appeared when the S1 sensor solution was excited at 400 nm. The fluorescence intensity of the peak at 547 nm sharply decreased with the addition of the $Ag⁺$ solution. No significant UV-vis or fluorescence spectra changes were observed when solutions of other cations (such as Fe^{3+} , Hg^{2+} , Ca²⁺, Cu²⁺, Co²⁺, Ni²⁺, Cd²⁺, Pb²⁺, Zn²⁺, Cr³⁺, and Mg²⁺) were added to the S1 sensor solution.

Fig. 1 An image of S1 upon the addition of 10 equiv. of various cations in DMSO/H₂O (v:v = 1 : 1, buffered with HEPES pH = 7.2), which was taken under (a) visible light and (b) UV-lamp (365 nm) at room temperature. (c) Fluorescence spectral responses in DMSO/H₂O (v:v = 1 : 1, buffered with HEPES pH = 7.2) of **S1** (2.0 \times 10⁻⁵ M) upon the addition of different cations (10 equiv.).

Subsequently, we further investigated the supramolecular sensor **S1** recognition for Pd²⁺, as shown in Fig. S2 and S3;[†] however, **S1** could not sense Pd^{2+} . These results confirmed that the self-assembly supramolecular sensor $S1$ could sense $Ag⁺$ with high sensitivity and selectivity.

The specific selectivity of a sensor towards an analyte in the presence of other competitive species is important. To explore the utility of the S1 sensor as an ion-selective chemosensor for Ag⁺, competitive experiments were carried out in the presence of 10 equiv. of Ag⁺ and 10 equiv. of Fe³⁺, Hg²⁺, Ca²⁺, Cu²⁺, Co²⁺, Ni^{2+} , Cd^{2+} , Pb^{2+} , Zn^{2+} , Cr^{3+} , and Mg^{2+} metal ions in DMSO/water $(v:v = 1 : 1$, buffered with HEPES pH = 7.2) binary solutions of the S1 sensor (Fig. 2 and S4†). From the bar diagram, it was observed that none of these ions induced any signicant change in the fluorescence and UV-vis absorbance spectrum of the sensor. Therefore, it is clear that the interference of other ions is negligible during the detection of Ag⁺. These results further suggested that **S1** could be used as a sensor for $Ag⁺$ over a wide range of cations. Thus, S1 could selectively and instantly (dualchannel) detect Ag⁺ in DMSO/water (v:v = 1 : 1, buffered with HEPES $pH = 7.2$).

To evaluate the Ag $^{\texttt{+}}$ -responsive nature of **S1**, fluorescence and UV-vis absorbance titrations with $Ag⁺$ at varying concentrations were conducted. As shown in Fig. 3, upon the addition of $Ag⁺$ to the S1 sensor, the emission peak at 547 nm gradually diminished, and the fluorescence of S1 was almost quenched by 18.0 equiv. of $Ag⁺$ ions. In the UV-vis spectra (Fig. 4), the initial absorption peak at 413 nm gradually decreased, and simultaneously, a broad shoulder peak appeared at 530 nm and an isosbestic point at 495 nm was clearly observed. This should be attributed to the damage of intramolecular hydrogen-bonding

Fig. 2 Fluorescent emission (λ_{ex} = 400 nm) spectra of S1 (2 \times 10⁻⁵ M) and Ag^+ (10 equiv.) in the presence of various competing cations in DMSO/H₂O (v:v = 1 : 1, buffered with HEPES pH = 7.2) solution. From 1 to 13: **S1**, **S1** + Ag⁺, **S1** + Ag⁺ + Fe³⁺, **S1** + Ag⁺ + Ca²⁺, **S1** + Ag⁺ + Cu²⁺, $S1 + Ag^+ + Co^{2+}$, $S1 + Ag^+ + Ni^{2+}$, $S1 + Ag^+ + Cd^{2+}$, $S1 + Ag^+ + Pb^{2+}$, $S1$ + Ag⁺ + Zn²⁺, S1 + Ag⁺ + Hg²⁺, S1 + Ag⁺ + Cr³⁺, and S1 + Ag⁺ + Mg²⁺, respectively.

Fig. 3 Fluorescence spectra of $S1$ (20 μ M) in the presence of different concentrations of Ag⁺ (0–18 equiv.) in DMSO/H₂O (v:v = 1 : 1, buffered with HEPES $pH = 7.2$) solution. Inset: A plot of the fluorescence intensity depending on the concentration of $Ag⁺$ in the range from 0 to 18 equiv.

interactions in the sensor and confirms the coordination of the sensor and Ag⁺. Based on the equation LOD = $K \times \delta/S$,¹³ the detection limits obtained from the UV-vis and the fluorescence spectra were 8.75 \times 10⁻⁷ M and 8.18 \times 10⁻⁹ M for silver ions, respectively.¹⁴ This data indicates that the sensor can detect Ag+ at very low concentrations in the environment with high selectivity and sensitivity. This result also shows that S1 has higher sensitivity for silver ions as compared to other reported $Ag⁺$ sensors (Table S1†). Therefore, S1 can be used as a potential colorimetric and fluorescent sensor for $Ag⁺$ recognition.

Since the pH value of the environment around a chemosensor usually influences its detection toward target metal ions due to the protonation or deprotonation reaction of the sensor, the influence of pH value on the absorbance and emission bands of S1 in a DMSO/water (1 : 1, v:v, buffered with HEPES pH

Fig. 4 Absorbance spectra of $S1$ (20 μ M) in the presence of different concentrations of Ag⁺ (0–18 equiv.) in the DMSO/H₂O (v:v = 1 : 1, buffered with HEPES $pH = 7.2$) solution. Inset: A plot of the variation in absorbance as a function of concentration of Aq^+ in the range from 0 to 18 equiv.

Fig. 5 pH response of S1 and S1-Ag⁺ in the DMSO/H₂O (v:v = 1 : 1; buffered with HEPES) solution.

 $(2, 7.2)$ solution has been investigated. As shown in Fig. 5 and S7,† when the pH increased from 5 to 10, the absorbance and emission values remained almost unchanged. These results indicated that $Ag⁺$ could be clearly detected via UV-vis and fluorescence spectral measurement using S1 over a wide pH range.

The self-assembly and $Ag⁺$ recognition mechanism of the S1 sensor were investigated using concentration titration, ESI-MS, ¹H NMR titration, and DFT methods. To establish the supramolecular self-assembly mechanism, we performed the concentration titration in the absence and presence of different concentrations of $S1$. As shown in Fig. 6, the fluorescence intensity of the peak at 547 nm sharply increased with the addition of S1; this suggested that with the gradual increase in the concentration of S1, the self-assembled system was formed by $\pi-\pi$ stacking and H-bonding in the aqueous solution. Moreover, the XRD spectra (Fig. S8†) support the

Fig. 6 Fluorescence spectra in the presence of different concentrations of **S1** (0–2.86 \times 10⁻⁵ M) in the DMSO/H₂O (v:v = 1 : 1, buffered with HEPES $pH = 7.2$) solution.

abovementioned competitive mechanism. The free sensor S1 exhibits a peak at $2\theta = 23.5^{\circ}$ corresponding to the d spacing of 3.82 Å, which suggests that $\pi-\pi$ stacking exists. In addition, compared to S1–Ag, this peak disappeared; this is attributed to the coordination of $Ag⁺$ with the **S1** sensor. Based on all the abovementioned tests, we confirmed that IM self-assembly formed S1 via π – π stacking and H-bonding in the aqueous solution.

The reversibility of the sensor function was tested by the addition of I^- to the **S1**–Ag complex. With the alternative addition of Ag^+ and I⁻ to the aqueous solution of the selfassembled S1 sensor, the fluorescence emission of the tested solution exhibited alternating reviving and quenching processes (Fig. S9†). This off–on–off switching process was repeated several times. Therefore, the S1 sensor can be considered a good ON-OFF fluorescence switch.

The $2:1$ stoichiometry of the S1 and Ag⁺ complex was further confirmed by the results of the ESI-MS studies. The free probe shows a main peak at $m/z = 301.09$, corresponding to $\left[\text{IM} - \text{H}^{\dagger}\right]$ (ESI-MS negative ion mode) (Fig. S14†), and the complex is confirmed by the appearance of a peak at $m/z =$ 731.12, assignable to $[2IM + Ag⁺ + H₂O + H⁺]$ (ESI-MS positive ion mode) (Fig. $S15\dagger$). The Job plot between **S1** and Ag⁺ was implemented, demonstrating a 2 : 1 stoichiometry for S1 and Ag⁺ , as shown in Fig. S16.†

The FTIR spectra of the receptor **IM** and its complex with $Ag⁺$ were obtained to examine the binding sites (Fig. S10†). In the FTIR spectrum of the free probe **IM**, the characteristic absorption band of **S1**–Ag $^+$ for N–H (3428 $\rm cm^{-1})$ of the imidazole group was absent, and an obvious enhancement was observed for the C-N absorption bands from 1322 to 1421 cm^{-1} ; this confirmed the binding of S1 with silver.

To gain more information about the complexation between **S1** and Ag⁺, we performed an ¹H NMR titration in the absence and presence of different concentrations of Ag⁺. As shown in Fig. S11,† the 1 H NMR titration of **S1** in DMSO- $d_{6}\left(10\text{ mmol}\right)$ with $Ag⁺$ in D₂O (1 mol) established the stoichiometry and binding site towards the sensor response. Most interestingly, the N–H peak $(H¹)$ of the imidazole group in IM disappeared after the

addition of 2.0 equiv. of Ag⁺ to D_2O ¹⁵, this proved that **S1** was strongly bound with Ag⁺. Moreover, the signal of the hydrogen atoms in the phenazine $(H^2, H^4, H^5, H^7,$ and $H^8)$ and thiazole $(H³, H⁶, and H⁹)$ rings showed a significant downfield shift; this suggested that the intermolecular π - π stacking was destroyed after the addition of Ag^+ , and it further caused the deshielding for the hydrogen protons and resulted in downfield shifts.

To further investigate the binding behavior of $S1$ with $Ag⁺$ ions, we optimized the structure of **IM** and $IM-Ag^+$ complex by applying the B3LYP/6-31G(d, p) method in the gas phase using the Gaussian-09 computational code.¹⁶ The optimized structures of IM and IM-Ag⁺ are shown in Fig. S12 and S13,[†] respectively. The short N34–H49 \cdots N23 distance of 2.041 Å indicates the presence of an intramolecular hydrogen bond in IM. Additionally, some of the relevant optimized geometric parameters are presented in Tables S2–S5.† The relevant occupied molecular orbital HOMO $(-0.295 a.u.)$ of IM was localized mainly to the thiophene groups, whereas the unoccupied orbital LUMO (-0.227 a.u.) was localized over the phenazine and imidazole groups. The HOMO–LUMO energy gap (DE) for IM was found to be 0.068 a.u. (Fig. 7). On the other hand, upon interaction with Ag^+ , the HOMO (-0.291 a.u.) and LUMO $(-0.279$ a.u.) were almost localized on the entire molecule, where the electron density was polarized toward the whole molecule. The HOMO-LUMO energy (DE) for IM with $Ag⁺$ was found to be 0.012 a.u. These results suggested that the fluorescence quenching was most probably attributed to the prevention of **IM** intramolecular charge transfer (ICT) by Ag^+ (Scheme 1). BSC Advances
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To investigate the practical application of the chemosensor S1, test strips were prepared by immersing filter paper in a DMSO/water (v: $v = 1 : 1$, buffered with HEPES pH = 7.2) solution of S1 and then drying them in air. The test strips containing $S1$ were utilized to sense $Ag⁺$ and other metal ions. As shown in Fig. 8, when $Ag⁺$ and other ions were added to the test kits, an obvious color change was only observed for the Ag⁺ solution under a 365 nm UV-lamp. The potential competitive

Fig. 7 DFT-optimized Frontier molecular orbitals and HOMO–LUMO energy gaps for S1 (left) and S1-Ag⁺ (right).

Fig. 8 Images of S1 on test papers. (a) Only S1, (b) after immersion in DMSO/H₂O (v:v = 1 : 1, buffered with HEPES pH = 7.2) solution with Ag⁺, (c) after immersion in DMSO/H₂O (v:v = 1 : 1, buffered with HEPES $pH = 7.2$) solution with others cations, and (d) after immersion in DMSO/H₂O (v:v = 1 : 1, buffered with HEPES pH = 7.2) solution with Ag+ and other ions under irradiation at 365 nm.

ions did not influence the detection of $Ag⁺$ by the test strips. Therefore, the test strips could conveniently detect $Ag⁺$ in the solution.

Conclusions

In summary, we developed a novel self-assembled supramolecular sensor $S1$ that could recognize Ag⁺ over other metal ions with significant UV-vis absorption and fluorescence quenching changes with highly selective and sensitive properties based on the competitive mechanism of supramolecular self-assembly. The obvious color changes and pronounced ON–OFF-type fluorescence signaling behavior can be seen by the naked eyes. The limit of detection for Ag⁺ was as low as 8.18×10^{-9} M in an aqueous solution, which was lower than that in some previously reported research. We believe that the sensor will be regarded as an environmentally friendly material that can sense $Ag⁺$ in an aqueous solution and make a great contribution to the development of silver ion sensors.

Conflicts of interest

There are no conflicts to declare.

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