1,3,4-Thiadiazol derivative functionalized-
Fe₃O₄@SiO₂ nanocomposites as a fluorescent
probe for detection of Hg²⁺ in water samples

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5-Amino-1,3,4-thiadiazole-2-thiol was used to synthesize a novel fluorescent functionalizing group on a Fe₃O₄@SiO₂ magnetic nanocomposite surface for detection of heavy metal ions in water samples. The prepared probe was characterized by using X-ray diffraction, transmission electron microscopy, Fourier transform infrared spectroscopy, and a vibrating sample magnetometer. Among various tested ions, the new nanocomposite responded to Hg²⁺ ions with an intense fluorescence "turn-on". The limit of detection of the probe shows that it is sensitive to the minimum Hg²⁺ concentration of 48.7 nM. Theoretical calculations were done for estimating binding energies of the three possible bonding modes and the visualized molecular orbitals were presented.

Introduction

The widespread toxicological properties of mercury have made it one of the most dangerous pollutants in the environment. Mercury released from natural sources or human activities ends up in the atmosphere and in water. According to the U.S. Environmental Protection Agency (EPA), the anthropogenic mercury emission in the US was measured to be 1.2 tons in 2011.1,2 Methyl mercury, which usually bioaccumulates in fish, can cause serious neurotoxic damage to the body. On the other hand, the possible form of mercury in drinking water is inorganic mercury. The main target of inorganic mercury in the body is the kidneys and especially the proximal tubule where mercury is accumulated by either luminal uptake of Cys–S–Hg–S–Cys or by basolateral uptake.3 The limit of mercury(ii) concentration in drinking water has been determined by the EPA to be 2 ppb.4

In order to ascertain the exact amount of mercury ions in drinking water, there are various analytical methods including atomic absorption spectroscopy,5 inductively coupled plasma-mass spectrometry,6 high performance liquid chromatography,7 and electrochemical sensing.8 However, these techniques suffer from complex and unhandy equipment which limits their on-site applications. Recently, fluorescent probes have emerged as a novel, selective, clean, simple, and portable alternative for detection of various chemical compounds in different mediums.9–11 Designing metal fluorescent probes is a challenging field of research since it needs a vast knowledge of metal coordination chemistry as well as photophysical mechanisms. Although designing new organic fluorophores is still of great importance, development of nanostructured substrates is more appealing since they serve a porous structure for hosting large amounts of fluorophore molecules and are conveniently separated after metal adsorption.14

The immobilization of organic fluorophores has been reported on certain nanosized substrates such as graphene quantum dots,15 carbon dots,16 Ag@SiO₂,17 SiO₂,18 MCM-41,19 SBA-15,20 and Fe₃O₄@SiO₂.21 The privilege of easy separation with external magnetic field has made Fe₃O₄@SiO₂ a superior choice. Many reports have already presented application of different fluorescence probes based on Fe₃O₄@SiO₂ nanocomposites in detection and removal of heavy metals.11,12,15–28 Both conventional and novel organic ligands with two important characteristics are potential for this purpose: (i) possessing suitable functional groups on one side to react and attach to SiO₂ surface either directly or through spacers such as [3-aminopropyl]triethoxysilane (APTES) or N-(2-aminoethyl)-3-aminopropyl trimethoxysilane (AEAPTES) (ii) having a fluorophore group which tends to coordinate with heavy metal and its fluorometric response changes after coordination. 1,3,4-thiadiazole has been known as a widely applied chemical compound in agricultural, pharmaceutical, and materials chemistry.29 The broad spectrum of its biological activities in terms of antibacterial, anticancer, antidepressant, etc. has made it an attractive compound for biologists and chemists. The special structure of thiadiazole class of components enables keto/enol tautomerism which is the origin of fluidity of biological membranes,30,31 crystal polymorphism, solvatomorphism effects, and interactions in lipid membranes.32,33 Moreover, the dual fluorescence effect of 1,3,4-thiadiazole is an interesting phenomena which is very important in understanding protein interactions of this molecule in biological systems.34

Recently, our group reported the successful design of two novel Fe₃O₄@SiO₂ fluorescence probes with precise detection ability of mercury ion in water. In continuation of our previous works, here, a novel fluorescent probe is designed and grafted on the Fe₃O₄@SiO₂ surface. A thia-diazole derivative is synthesized and applied as the fluorophore functionalizing group on Fe₃O₄@SiO₂ nanocomposites and is used for detection of Hg²⁺ in drinking water. 5-Amino-1,3,4-thiadiazole-2-thiol (ATT) ligand which is used as a starting material in this work, has already been reported as a potential silica modifier for sorption of different heavy metals including Hg²⁺. The –SH site on the aromatic ring in this molecule has provided an excellent coordination ability to soft heavy metals.

Experimental

Characterization

X-ray diffraction (XRD) patterns were recorded by a Philips-X’PertPro, X-ray diffractometer using Ni-filtered Cu Kα radiation at scan range of 10° < 2θ < 80°. Scanning electron microscopy images were obtained on LEO-1455VP. Room temperature photoluminescence properties were studied on a Perkin-Elmer LS 55 Mv fluorescence spectrophotometer. Fourier transform infrared (FT-IR) spectra were recorded on Magna-IR, spectrometer 550 Nicolet with 0.125 cm⁻¹ resolution in KBr pellets in the range of 400–4000 cm⁻¹. TEM images were taken on an EM208S Philips transmission electron microscope with an accelerating voltage of 100 kV. For TEM analysis, the nanoparticle powder was dispersed in ethanol by ultrasonic bath. Then a drop of the dispersion was placed on the TEM grid and was dried in the vacuum oven and then it was used for TEM imaging. Room temperature magnetic properties were investigated using a vibrating sample magnetometer (VSM), made by Meghnatis Daghigh Kavir Company (Iran) in an applied magnetic field sweeping between ±10 000 Oe.

Synthesis of thia-diazol-based ligand

5-(4-Vinylbenzylidene)amino-1,3,4-thiadiazole-2-thiol (VBYT) (3) was prepared using ATT by the classical Schiff base reaction. First, terephthalaldehyde (TPAL) (1 mmol) was suspended in 10 mL of methanol and added to 5-amino-1,3,4-thiadiazole-2-thiol (1 mmol) and the solution was stirred under reflux at 100 °C for 1 h (VBYT, see Scheme 1). Then, AEAAPTES (1 mmol) was added to this solution and the reaction mixture was stirred under reflux at 60 °C for another 1 h. The precipitate was filtered off and recrystallized in DMF. The red product (VBYT–AEAPTES, see Scheme 2) was then dried at room temperature. The progress of the reaction was checked by thin layer chromatography (TLC).

Synthesis of magnetic Fe₃O₄@SiO₂ nanoparticles (NPs)

The Fe₃O₄-coated SiO₂ nanoparticles were synthesized by a similar procedure developed by Dong et al. At first, for degassing the flask, the deionized water was exposed to nitrogen gas for 10 min. Then, Fe(NO₃)₃·9H₂O (8.12 g, 0.020 mol) and FeCl₂·4H₂O (2.00 g, 0.010 mol) and 1 mL oleic acid were dissolved in 30 mL deionized water at 90 °C under nitrogen atmosphere and vigorous mechanical stirring (1000 rpm). Then, 20 mL of NH₄OH (28 wt%) was quickly injected into the reaction mixture in one portion and it immediately changed to black. The mixture continued under this condition (90 °C) for 2.5 h and the colloidal solution was cooled to room temperature. After cooling, it was suspended in chloroform.

For preparation of Fe₃O₄@SiO₂ nanoparticles, a modified method reported by Zhu et al. was used. 25 mg of Fe₃O₄ nanoparticles was added into 40 mL of cyclohexane at room temperature by ultrasonic dispersion. Triton X-100 (9 g), octanol (8 mL) and H₂O (1.7 mL) were then added with stirring to form water-in-oil microemulsion. The reaction mixture was vigorously stirred for 15 min. Then, 1 mL of TEOS was added slowly into the solution. The reaction vessel was stirred on a shaker for 1.5 h at room temperature and then, 1 mL aqueous ammonia (28 wt%) was added into the reaction. The stirring continued for 24 h at room temperature to form the Fe₃O₄@SiO₂ core–shell nanoparticles. The Fe₃O₄@SiO₂ nanoparticles were isolated via centrifugation at 4000 rpm and washed with ethanol for five times. The obtained dark grey product was allowed to dry at 80 °C.

Scheme 1 Synthetic procedure of VBYT.

Scheme 2 The procedure for synthesis of VBYT–Fe₃O₄@SiO₂.
Preparation of VBYT-Fe₃O₄@SiO₂

First, 30 mg of dried Fe₃O₄@SiO₂ and 30 mg (1 mmol) of VBYT-AEAPTES ligand were suspended in 20 mL of DMF. DMF was used as solvent here, because the solubility test for VBYT-AEAPTES ligand showed that it is completely soluble in it. The mixture was refluxed for 12 h at 60 °C. The products were collected by filtration and repeatedly washed with DMF and then ethanol several times.

Preparation of the fluorometric metal ion titration solution

The fluorometric measurements of metal ions were performed in phosphate-buffered saline (PBS) solutions (0.02 M, pH = 7.0). 0.03 g of the functionalized Fe₃O₄@SiO₂ nanoparticles were dispersed in 3 mL of 1 × 10⁻⁶ M of different metal nitrate (Hg²⁺, Na⁺, Mg²⁺, Ca²⁺, Al³⁺, Cr³⁺, Zn²⁺, Ni²⁺, and Fe³⁺) aqueous PBS solutions for testing the effect of metal type on fluorescence response of the as-prepared probe. The performance of the prepared nanocomposite was also tested in tap water of the chemistry laboratory located in University of Zabol, as a real sample.

Calculations

The molecular structure of VBYT-AEAPTES and its interaction with Hg²⁺ in three different bonding modes in the ground state were optimized on the basis of density function theory (DFT) at the Becke3–Lee–Yang–Parr (B3LYP) (with LanL2DZ basis) and by means of visual inspection using the GAUSSVIEW program (Version 5.0). All the calculations were performed using the GAUSSIAN 09 software package. The population analysis was performed by natural bond orbital (NBO) method using NBO program implemented under Gaussian 09 program package.

Results and discussion

Characterization of as-prepared samples

The phase of the samples were characterized by their X-ray diffraction patterns. Fig. 1a and b show the XRD patterns of Fe₃O₄ and Fe₃O₄@SiO₂ nanoparticles, respectively. Fig. 1a shows that the pattern corresponds to pure cubic Fe₃O₄ with JCPDS card no.: 19-629. The XRD pattern of Fe₃O₄@SiO₂ consists of SiO₂ and Fe₃O₄ phases (Fig. 1b). The broad band at ca. 2θ = 20–30° is attributed to the amorphous phase of SiO₂. The peaks corresponding to Fe₃O₄ in Fe₃O₄@SiO₂ (Fig. 1b) are weaker than those in pure Fe₃O₄ which shows that Fe₃O₄ is covered with SiO₂ shell.

In order to study the surface chemistry of the samples, FT-IR spectroscopy was used. Fig. 2a–c show the FT-IR spectra of VBYT–AEAPTES ligand, Fe₃O₄@SiO₂, and VBYT-Fe₃O₄@SiO₂, respectively. In Fig. 2a, the weak peak at 712.86 cm⁻¹ is referred to C–S stretching vibration. The peak at 1060.18 cm⁻¹ is attributed to Si–O stretching vibration. The characteristic bands at 1610.43 and 1554.68 cm⁻¹ are attributed to C=O and C=C stretching vibrations, respectively. The extremely weak band at 2598.26 cm⁻¹ is for S–H stretching vibration. The appeared peak at 2925.59 cm⁻¹ is assigned to sp³ C–H stretching of alkane chain. A single vibration at 3342.17 cm⁻¹ is assigned to secondary amine in AEAPTES counterpart.

In Fig. 2b and c, the main appeared peaks correspond to Fe₃O₄ cores and SiO₂ shells. The peaks around 590 cm⁻¹ and 470 cm⁻¹ in Fig. 2b and c are assigned to the Fe–O vibration of magnetite core. The displayed bands around 3400 cm⁻¹ and 1636 cm⁻¹ are due to the stretching and bending vibration of remained H₂O and silanol O–H groups, respectively. In both spectra, the broad and strong peak around 1100 cm⁻¹ is assigned to the overlapped stretching vibration of Si–O–Si bonds. Two characteristic peaks at 2854 cm⁻¹ and 2925 cm⁻¹ are referred to the symmetric and asymmetric stretching vibration of C–H of the alkyl groups. In Fig. 2c, weak peaks in the range of 1000–1700 cm⁻¹ show the attachment of VBYT–AEAPTES ligand on the surface of Fe₃O₄@SiO₂ nanocomposites.

For characterization of the prepared core–shell nano-composites, TEM analysis was performed on images of VBYT-Fe₃O₄@SiO₂ sample and the results were shown in Fig. 3a and b. It can be seen that the formed nanoparticles are very tiny and are

![Fig. 1. XRD patterns of (a) Fe₃O₄ nanoparticles and (b) Fe₃O₄@SiO₂.](image-url)

![Fig. 2. FT–IR spectra of (a) VBYT–AEAPTES ligand, (b) Fe₃O₄@SiO₂, and (c) VBYT-Fe₃O₄@SiO₂.](image-url)
partly agglomerated because of the magnetic properties of the nanocomposites. A higher magnification of one separate nano-particle, shown in Fig. 3b-inset, clearly shows the core–shell structure of the nanocomposite. The diameter of the nanocomposite particle is approximately 32 nm with a core with the diameter of 11 nm. Therefore, the SiO2 coverage layer around Fe3O4 core is about 21 nm.

For studying the magnetic properties of the Fe3O4 and VBYT-Fe3O4@SiO2, a vibrating sample magnetometer was used at 300 K (Fig. 4). The measured saturation magnetizations (Ms) are 50.33 emu g⁻¹ and 2.89 emu g⁻¹ for Fe3O4 and VBYT-Fe3O4@SiO2, respectively that are shown in Fig. 4a. This decrease in magnetic properties is due to coverage of the Fe3O4 cores by SiO2 and VBYT layers. Both samples are superparamagnetic because they almost show no remanence and coercivity. Although the magnetization of VBYT-Fe3O4@SiO2 is small, it is easily separated from the solution by placing a magnet, within 3 minutes (Fig. 4b). This results show that the prepared material is potential for water purification applications since it is efficiently absorbed by a simple magnet.

Testing the fluorescence properties of the nanocomposite in response to metal absorption

In order to test the response of the prepared sample against different metal ions, the fluorescence spectra of the nanocomposite were recorded in presence and absence of some important metal ions including Ni²⁺, Hg²⁺, Zn²⁺, Mg²⁺, Al³⁺, Na⁺, Ca²⁺, Fe³⁺, and Cr³⁺. Fig. 5 shows the results of the fluorescence test after exciting the samples at λ_exc = 268 nm. The λ_max emission (λ_max em) of the free receptor (FR) i.e. VBYT-Fe3O4@SiO2 dispersed in buffer solution in absence of metal ions is at 409 nm. After addition of fixed amounts of Hg²⁺, Cr³⁺, Ni²⁺, Zn²⁺, Mg²⁺, Fe³⁺, Al³⁺, Ca²⁺, and Na⁺ the intensity of the maximum peak was either unchanged or slightly tuned in case of Al³⁺, Ni²⁺ and Ca²⁺. Moreover, almost no peak shift was observed at λ_max em compared to the FR. For Zn²⁺, the peak at 340 nm was intensified that probably could be attributed to its different bonding mode compared with other metal ions which prohibits the photoinduced electron transfer (PET) mechanism and results in a “turn-on” switching. In case of Hg²⁺, a significant quenching (about 1100 units) was observed for λ_max em peak at 409 nm. Therefore, by comparing the response of VBYT-Fe3O4@SiO2 nanocomposite to different ions, it was shown that the prepared material could be a good candidate for detection of Hg²⁺ ions. Similar results have been reported for application of silica gel modified with ATT by Tzvetkova et al. They indicated that among some heavy metals including Cu(II), Co(II), Ni(II), Cd(II), Pb(II) and Hg(II), the modified silica gel shows the highest sorption capacity against Hg(II). Certain factors such as suitable coordination geometry of ligand and ion, nitrogen affinity, and ion size and charge are effective in high selectivity of an ion. In continue, more tests were

![Fig. 3](image-url) (a) and (b) TEM images of VBYT-Fe3O4@SiO2 from two different parts of the samples. Inset: one separate core–shell VBYT-Fe3O4@SiO2 nanoparticle.

![Fig. 4](image-url) (a) The magnetic hysteresis loops of Fe3O4@SiO and VBYT-Fe3O4@SiO2 (b) the separation of VBYT-Fe3O4@SiO2 in aqueous solution after 3 minutes.

![Fig. 5](image-url) The fluorescent spectra of VBYT-Fe3O4@SiO2 in presence of various metal ions (λ_exc = 268 nm).
conducted in order to show the potential application of the nanocomposite on detection of Hg$^{2+}$ ions in water samples.

To elucidate the detection potential of VBYT-Fe$_3$O$_4$@SiO$_2$ against Hg$^{2+}$, a titration experiment was performed and the results were shown in Fig. 6. For this purpose 0.03 g of nanocomposite was dispersed in 10 mL of 0, 0.1, 0.3, 0.5, 1, 3, 5, 10, 20, and 30 × 10$^{-6}$ M of Hg$^{2+}$ in PBS buffer solution (pH = 7, water). As can be seen in Fig. 6a, with increasing Hg$^{2+}$ concentration, two appeared peaks at 409 nm (denoted as blue emission) and 351 nm (denoted as UV emission) are gradually quenched. It is clearly seen that the blue emission is quenched faster than the UV emission. It proves that the part of the fluorophore molecule which is responsible for the UV emission, is first involved in PET mechanism from Hg$^{2+}$ as the receptor. On the other hand, it seems that the receptor is directly bonded to the group(s) responsible for the blue emission and is transferring electron to this group(s) through PET mechanism$^{44}$ which results in a faster “turn-off”. Fig. 6b shows the graph of fluorescence intensity versus Hg$^{2+}$ concentration. It is seen that in the range of 0.1–0.5 μM of Hg$^{2+}$, there is a good linear relationship in the graph with $R^2 = 0.9833$. The regression equation is $F = -1896.2C (\mu$M Hg$^{2+}) + 1225.5$ for the intensity change of the red shift with increasing Hg$^{2+}$ concentration. The limit of detection (LOD) of the probe was calculated to be 48.7 nM for Hg$^{2+}$ detection which satisfies the U.S. EPA (the U.S. Environmental Protection Agency) limits (~2 ppb) of Hg$^{2+}$ detection in drinking water and is a good record compared to some other reports.$^{38,46–48}$

The selectivity of the synthesized probe was tested in presence of some interfering ions. Fig. 7 shows the fluorescence changes of VBYT-Fe$_3$O$_4$@SiO$_2$ by treatment of 10$^{-6}$ M of the interfering ions in 10 mL PBS buffer solution containing 0.03 g of the dispersed nanocomposite. Moreover, the interfering ions were tested in presence of the same concentration of Hg$^{2+}$. It is shown that the tested metal ions (Na$^+$, K$^+$, Mg$^{2+}$, Al$^{3+}$, Ca$^{2+}$, Cd$^{2+}$, Ni$^{2+}$, Co$^{2+}$, Zn$^{2+}$, Cu$^{2+}$, Fe$^{3+}$, Pb$^{2+}$, Ag$^+$) have small or no interference with Hg$^{2+}$ ions and this ion can be easily detected in presence of either of the shown elements. In order to have an account of the behaviour of the designed probe in the real sample, the fluorescence spectra of FR as well as FR plus Hg$^{2+}$ ions were tested in drinking tap water of University of Zabol as a real sample. The results shown in Fig. 7 (in blue columns) show that very similar behaviour is also observed in the real sample. These results suggest that VBYT-Fe$_3$O$_4$@SiO$_2$ is a good candidate to be applied for detection of Hg$^{2+}$ with high selectivity against other metal ions in a real aqueous samples.

To test the regeneration ability of the probe, VBYT-Fe$_3$O$_4$@SiO$_2$ was tested by using excess amount of NaI which is able to remove Hg$^{2+}$ from the chemosensor surface. Fig. 8 shows the results of the regeneration test which proves that the fluorescence properties of VBYT-Fe$_3$O$_4$@SiO$_2$ is reversible and the...
probe still has its ability after removing the content of Hg^{2+} ions by NaI.

In the present study, interactions of the Hg^{2+} ions with positions on VBYT–AEAPTES ligand which encompass nitrogen and sulphur atoms were considered. Results indicate that the Hg^{2+} ions interact preferably with the ligand through atoms N_{28}, N_{31}, and S_{37} which are labelled in Scheme 3. The binary complexes which was formed through interactions of the Hg^{2+} ions with the mentioned atoms on the ligand are denoted as A, B, and C, respectively (see Scheme 3). The order of the binding energies of these complexes (in kcal mol^{-1}) is: B (217.61) < A (220.20) < C (227.75). Also, gaps between energies of highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) in ligand and complexes were calculated. The results point out that energy gap in the ligand is 2.857 eV and interactions of Hg^{2+} ions with the ligand lead to decreasing the energy gaps in the complexes. The order of energy gaps in the complexes (in eV) is: B (0.423) < A (0.432) < C (0.478) which is in agreement with the order of the binding energies.

The NBO analysis was performed on the mentioned complexes to obtain better insight regarding the order of binding energies. Donor–acceptor interaction energy (E^2) values between all donors and acceptors were studied. Results indicate that the order of E^2 values (in kcal mol^{-1}) of interactions between lone pair (Lp) of Hg and antibonding (BD*) orbitals of X–Y (X = N_{28}, N_{31}, and S_{37}; Y = C_{11}, C_{32}, and C_{33}) in the complexes is: C (0.36) < A (0.51) < B (0.74). In fact, decrease of

![Scheme 3](image)

Scheme 3 Molecular orbital structure of HOMO and LUMO of different bonding modes including (a) N_{28}, (b) N_{31} and S_{36}, and (c) S_{37}.

anti-bonding feature of X–Y bonds through interaction of Hg^{2+} ions with VBYT–AEAPTES ligand leads to increase of binding energies of the complexes.

Based on the abovementioned theoretical results, the fluorescence “turn-off” behaviour of VBYT-Fe_{2}O_{4}@SiO_{2} nanocomposites is depicted in Scheme 4. As can be seen in Scheme 4a, two different parts of the VBYT molecule including benzyldiene and thiadiazole are responsible for UV and blue emissions, respectively. After addition of low concentrations of Hg^{2+}, S_{37} site coordinated to the entered metal ions due to its higher binding energy. The coordinated Hg^{2+} ion may act as receptor and transfers electron to the fluorophore via PET mechanism causing the quenching of the blue emission (Scheme 4b). Another possible mechanism for quenching the UV band could be the enhanced intersystem crossing (ISC) to the triplet state due to “heavy ion effect”. By increasing the concentration of the Hg^{2+} ions in the solution and saturation of S_{37} sites in VBYT molecule, gradually N_{28}–Hg^{2+} bond with high binding energy is formed and again, by PET mechanism, the UV emission turns off, as well. Quenching of blue emission due to the “heavy ion effect” is less likely since the sensor is already significantly spin–orbit coupled due to the already coordinated Hg^{2+} ion.

**Conclusions**

VBYT-Fe_{2}O_{4}@SiO_{2} nanocomposites were synthesized by using ATT as a starting material. The characterizations showed
The presence of various active sites for coordinating Hg^{2+} ions. The sensitivity of the probe was calculated by showing that the blue fluorescence response of the nanocomposite against different ions showed that the blue fluorescence peak is easily quenched by adding Hg^{2+} ions. The sensitivity of the probe was calculated by titration tests and it was shown that the limit of detection is 48.7 nm and the fluorescence intensity diagram against Hg^{2+} concentration showed a good linear relationship in the range of 0.1–0.5 μM. Theoretical calculations showed that the binding energies of S_{37}, N_{28}, and N_{31} bonds with Hg^{2+} ion has an order of S_{37} > N_{28} > N_{31}. The presence of various active sites for coordination to mercury atoms makes this detector potential adsorbent for removing heavy metal atoms with high capacity. The absorbing application of the designed nanomaterial is the subject of our further research work.

Conflicts of interest

There are no conflicts to declare.

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