We described the design and synthesis of a molecular sensor based on a rhodamine/BODIPY platform that displayed differential fluorescence responses towards Hg$^{2+}$ and Au$^{3+}$ and demonstrated its utility in intracellular ion imaging.

In recent years, the construction of fluorescent molecular sensors for the detection of metal ion species has received a great deal of attention.$^1$ To date a large number of molecular sensors have been designed and developed, the majority of which are single-ion responsive and present no great challenge to researchers. Compared to single-ion responsive molecular sensors, however, the construction of multi-ion responsive molecular sensors with multiple emission modes are extremely challenging.$^2$ Molecular sensors displaying differential responses towards multiple ions are indispensable for designing molecular logic gates and molecular keypad lock devices.$^{3,4}$

The challenge of multiple analyte recognition presents several detection strategies. Incorporating multiple binding motifs onto a single sensing molecule, or alternatively, combining different transducing units (chromophores/fluorophores), allows for rapid access to molecular sensors with multiple emission modes.$^3$

We envisaged that incorporating both a chemosensor and a chemodosimeter onto a single molecule could provide a suitable sensing platform for the differential detection of metal species. On the basis of this hypothesis, we constructed a molecular sensor possessing two different fluorophore units chemically integrated with each other. Both fluorophore units were elegantly designed to be non-emissive (i.e., “off”) in their initial states and are expected to turn on respectively in response to the metal species of interest. To the best of our knowledge, molecular sensors based on this novel approach have not been covered in the literature.

Ionic species of mercury (Hg$^{2+}$) and gold (Au$^{3+}$) share several similarities in terms of coordination properties. As both metal species show high affinities to thiols, they have the potential to interact with sulfur bearing biomolecules such as enzymes, proteins, and DNA. As a result, these metal species can disturb a series of cellular processes that lead to toxicity in humans.$^5$ In recent years, a variety of well-designed fluorescent probes highly specific for Hg$^{2+}$ and Au$^{3+}$ ions have been developed, most of which are built on the exploitation of the thiophilic and alkynophilic behavior of these metal species.$^7$ Despite impressive advances, many of these probes suffer from cross affinity. Because of their similar coordination properties, it is extremely difficult to construct a molecular sensor that differentiates between the Hg$^{2+}$ and Au$^{3+}$ species. To our knowledge, in the literature there is only one example of a fluorescent probe that can differentiate between Au$^{3+}$ and Hg$^{2+}$. This fluorescent probe, reported by Dong et al., operates through a single emission mode and the differentiation is highly dependent on the sensing conditions.$^2d$

Obviously, there is a high demand for the development of molecular sensors that can differentiate multiple analytes of a similar chemical nature (e.g. Hg$^{2+}$ and Au$^{3+}$). In addition, small-molecule fluorescent sensors allowing the intracellular monitoring of multiple ions via differential responses are of high necessity for real-time cell imaging studies.

Herein, we present the design, synthesis, spectral properties, and cell imaging studies of RhS-BOD, a new “turn-on” multi-fluorescent probe that allows the Hg$^{2+}$ and Au$^{3+}$ species to be differentiated on the basis of distinct fluorescence responses. RhS-BOD constitutes a boron-dipyromethene (BODIPY) dye and a spirocyclic rhodamine dye covalently attached to each other. RhS-BOD was prepared in a reasonable yield (25%, overall) by the synthetic route outlined in Scheme S1 (see ESI†), and the structure of RhS-BOD was confirmed by $^1$H-NMR, $^{13}$C-NMR, and mass spectroscopy.

Importantly, the novel molecular sensor, RhS-BOD, was designed in such a way that both of the fluorophore units are non-emissive before the addition of any metal species. As can be seen from the structure of the probe in Scheme 1, the C—N...
functionality of the BODIPY core diminishes the BODIPY emission potentially because of a non-radiative deactivation process of the excited state through rapid isomerization of the C–N group. Similarly, the rhodamine fluorophore is non-emissive because the rhodamine dye exists in the ring closed isomeric form.

The sensing behavior of RhS-BOD towards the addition of different metal species was studied using UV-Vis and fluorescence spectroscopy. As shown in Fig. S1 (see ESI†), the UV-Vis spectrum of free RhS-BOD (CH3CN/HEPES 1:1, pH 7.0) exhibits a single absorption band at 527 nm, which belongs to the BODIPY core. As the rhodamine core is in the ring closed isomeric form, we expect no absorption bands for the rhodamine derivative. However, the addition of Hg2+ (1 equiv.) to RhS-BOD led to the appearance of a new strong absorption band at 554 nm, which was assigned to a ring opened rhodamine derivative.

The fluorescence spectra of RhS-BOD displayed a similar behavior towards the addition of Hg2+ (Fig. 1). Initially, when excited at 525 nm there were no emission bands in the fluorescence spectrum of RhS-BOD. However, upon the addition of Hg2+, a new emission band with a maximum at 585 nm appeared and the intensity of this band gradually increased with an increasing concentration of Hg2+ (Fig. 1). The increase in emission intensity showed a linear relationship towards the addition of Hg2+ in the range of 0–3 μM. The minimum amount of Hg2+ was evaluated to be 8.0 nM under these conditions (Fig. S10, ESI†). The response of the probe towards the addition of Hg2+ was immediate and the emission intensity became saturated when 1 equiv. of Hg2+ was added, creating an enhancement factor of over 50-fold.

As expected, in the presence of Hg2+, RhS-BOD displayed the optical features of the rhodamine chromophore. During the process of adding Hg2+, no other accompanying emission bands were noticed in the emission spectrum that might belong to the BODIPY dye, indicating that the BODIPY dye was still in a sleep (“off”) mode.

To check the reversibility of the Hg2+ sensing process, the highly emissive probe solution pre-treated with Hg2+ (RhS-BOD/Hg2+) was subsequently treated with a cyanide ion source (KCN or NH4CN) (Fig. S7, ESI†). The probe solution immediately lost its color and its emission, thus showing that the sensing process is based on a reversible metal–ligand coordination process. The binding stoichiometry of the Hg2+/RhS-BOD association was determined by Job’s plot from both the UV-Vis absorption and fluorescence data (Fig. S9, ESI†). Both of the plots revealed that Hg2+ is associating with the probe in a 1:4 ratio.

We further investigated the selectivity profile of RhS-BOD in response to other metal ions. For all other metal cations, such as Cu2+, Ag+, Zn2+, Pb2+, Ni2+, Na+, Mg2+, Li+, K+, Pd2+, Fe2+, Co2+, Cd2+, Ca2+, Ba2+, Fe3+ and Cr3+, no detectable change in the emission intensity for RhS-BOD was observed (Fig. S5, ESI†). The probe was highly selective towards Hg2+ and showed no spectral response to any other metals ions, except for Au3+ ions. Upon the addition of Au3+, the non-emissive probe solution immediately turned to a strong green emissive solution that could be easily monitored by the naked eye under the UV lamp. A green emission was clear evidence of the existence of an emissive BODIPY derivative. This suggestion was also supported from the outcome of the reaction of RhS-BOD mediated by Au3+, as controlled using TLC. The formation of a green emissive compound, BODIPY-AL, could be easily monitored from the spots on the TLC plate (Fig. S21, ESI†).

The fluorescence sensing behavior of RhS-BOD towards Au3+ was comprehensively surveyed upon excitation at 470 nm and 525 nm. As shown in Fig. S11 (ESI†), the fluorescence spectrum of RhS-BOD/Au3+ (1:1) displays an emission band at 506 nm when excited at 470 nm, a characteristic emission band of a BODIPY fluorophore. On the other hand, the same probe solution (RhS-BOD/Au3+) when excited at 525 nm displays a different emission band at 585 nm, which is supposed to belong to the ring opened isomer of the rhodamine core (Fig. S13b, ESI†). The fluorescence emission intensity at both wavelengths increased linearly with an increasing concentration of Au3+ over a wide concentration range (Fig. 2 and Fig. S14, ESI†). The response of RhS-BOD towards Au3+ was fast (<1 min) and the emission intensity became saturated when 2 equiv. of Au3+ was added. In addition, the detection limit measured at both wavelengths was at the nM level (65 nM, λem = 585 nm and 10 nM, λem = 506 nm).

The fluorescence response of RhS-BOD toward Au3+ (1 equiv.) in the presence of other metal ions (10 equiv.) was explored in...
order to assess the possible interference by other metal ions. As shown in Fig. S16 (see ESI†), the tested metal ions displayed no interference with the detection of Au3+ ions.

As discussed earlier, the addition of Hg2+ to RhS-BOD triggers a spiro-ring opening reaction and results in the formation of a highly emissive rhodamine derivative. Throughout the addition of Hg2+ to RhS-BOD, the BODIPY core continues to be non-emissive because the C–N moiety was still preserved. However, the addition of Au3+ to the probe solution pre-treated with Hg2+ (RhS-BOD/Hg2+) resulted in an immediate change in the emission color from orange to green. Evidently, in the presence of Hg2+ and Au3+, RhS-BOD hydrolyzes to give a green emissive BODIPY derivative, BODIPY-AL, which dominates the emission color of the probe solution (Scheme 2).

RhO-BOD, the oxygen bearing derivative of RhS-BOD, was used as the control probe to clarify the nature of the sensing process. Under the same sensing conditions, RhO-BOD displayed no response towards any metal species, indicating the indispensable role of the sulfur functionality in the detection of both metal species.

We next assessed the ability of RhS-BOD to operate within living organisms. To our delight, RhS-BOD showed the same sensing behavior in living cells. Human A549 lung adenocarcinoma cell lines were incubated with the probe (5 μM) for 40 min and then followed by the addition of metal species. With the aid of fluorescence microscopy, the differential turn-on response towards Au3+ and Hg2+ was clearly monitored in the cells (Fig. 3). The images taken before and after the addition of the metal species displayed a distinct fluorescence change consistent with the results observed in the solution.

In conclusion, we have presented the synthesis, spectral properties, and biological applications of RhS-BOD, a new type of fluorescent probe for the differential detection of Hg2+ and Au3+. This novel probe features excellent selectivity for Hg2+ and Au3+. Detection of Hg2+ and Au3+ is realized through two distinct fluorescence changes resulting from Hg2+–ligand coordination or from the hydrolysis of the C–N moiety catalyzed by Au3+. RhS-BOD exhibits a dual emission mode for the detection of Au3+ ions and a single emission mode for the detection of Hg2+ ions.

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Fig. 3 (a) Fluorescence image of A549 cells treated with only RhS-BOD (5 μM); (b) image of cells treated with the probe (5 μM) and Au3+ (5 μM) (λex = 470 nm); (c) image of cells treated with the probe (5 μM) and Au3+ (5 μM) (λex = 525 nm); (d and h) images of cells treated with DAPI for 15 min (control); (d) merged image of frames (b) and (d); (f) merged image of frames (c) and (e); (g) image of cells treated with the probe (5 μM) and Hg2+ (5 μM) (λex = 525 nm); (i) merged image of frames (g) and (h).