A BODIPY-based fluorescent probe for ratiometric detection of gold ions: utilization of Z-enynol as the reactive unit†

Muhammed Üçüncü, Erman Karakuş and Mustafa Emrullahoğlu*

Using an irreversible intramolecular cyclisation pathway triggered by gold ions, a boron-dipyrrromethene (BODIPY) based fluorescent probe integrated with a reactive Z-enynol motif responds selectively to gold ions. With the addition of gold(aq), the probe displays ratiometric fluorescence behaviour clearly observable to the naked eye under both visible and UV light.

Though primarily known for its monetary value, gold has recently attracted attention for its impressive catalytic features as well. In synthetic chemistry, the use of gold species has opened new avenues in the synthesis of complex molecular structures. However, the increasing use of gold species in chemical synthesis is concomitantly raising hard-to-ignore health and environmental risks due to their potential toxicity. The intake of gold ions in living systems has been widely found to damage vital human organs, including the kidney and liver, as well as the peripheral nervous system. It is therefore crucial for the scientific community to be able to assess the levels of gold species in certain chemical, environmental, and biological matrices.

In that context, research continues to focus on developing analytical tools for probing gold species, among which fluorescent-based assays have attracted particular attention for allowing the real-time visualisation of target species in cellular milieus. In recent years, numerous types of fluorescent gold ion probes have appeared in literature on the topic, most of which involve using specific chemical reactions to exploit the alkynophilic nature of gold ions. By extension, various fluorescent molecules—rhodamine, boron-dipyrrromethene (BODIPY), fluorescein, and naphthalimide—have been used as signal-reporting fluorophores to recognise gold species by colorimetric or fluorometric changes, if not both, in which optical output is usually recognised as an increase in or activation of emission intensity. However, measurements based on intensity changes are easily influenced by a host of environmental factors, including concentration variations, photo-bleaching, and excitation intensity.

To overcome the barriers generally associated with intensity-based sensors, measuring optical signals as intensity ratios at two different wavelengths seems quite promising, for it would allow for a built-in correction for environmental effects. Until now, examples of fluorescent probes based on the ratiometric detection of gold ions have been inadequate, due to the absence of effective strategies and guidelines for designing ratiometric fluorescent molecules. In a recent contribution, we introduced a new strategy for ratiometrically sensing gold ions by exploiting the catalytic behaviour of gold ions to transform a highly conjugated probe structure into a non-conjugated one, by which the presence of gold species became detectable as an emission ratio of two distinct wavelengths. However, cross-sensitivity towards mercury(II) ions is still a challenge to be addressed. Other examples of ratiometric gold ion probes available in recent literature are extensions of intensity-based sensors that draw upon a principle of fluorescence resonance energy transfer to achieve ratiometric response.

It is a well-known fact that the extent of π-conjugation within the fluorophore–chromophore structure exerts a dramatic effect on the frontier orbital energy levels of the molecules. With the conjugation of fluorescent dye the energy difference between the orbitals decreases, which often results in a redshift of absorption and emission wavelengths.

Based on these considerations, we designed and constructed BOD-Z-EN by integrating a Z-enynol motif into a BODIPY-based fluorophore scaffold, with the expectation of generating a highly π-conjugated BODIPY derivative emitting at a wavelength distinctly longer than its unmodified scaffold (Scheme 1). We selected a BODIPY-based fluorophore as the signal reporter for not only its outstanding photo-physical features, but also its easy chemical modification. In the presence of gold species, as reported in the literature, substituted Z-enynols can efficiently cyclize to their corresponding furan derivatives.

We envisioned that an intramolecular cyclisation triggered by gold ions could break the π-conjugation and generate a new...
The fluorescent probe **BOD-Z-EN** was prepared by way of a straightforward synthetic pathway, as outlined in Scheme 2. The monoiodo derivative of BODIPY (**BODIPY-I**) was prepared individually with 2-enynol \([Z]-3\text{-methylpent-2-en-4-yn-1-ol}\) according to a Sonogashira coupling protocol in order to induce the desired probe structure in a moderate yield. Following chromatographic purification, the identity of the title compound was clearly confirmed by analytical data collected with nuclear magnetic resonance (NMR) spectroscopy and mass spectrometry. Using the synthetic route we prepared **BOD-E-EN**, the \(E\)-isomer of the probe, in order to elucidate the mechanism of gold ion detection. With both compounds in hand, we first investigated the spectral properties of **BOD-Z-EN** and its response to a range of metal ions in a solution buffered to physiological pH (0.1 M phosphate buffer/EtOH \((pH 7.0, v/v, 7 : 3)\)). We commenced spectral investigation by screening the optical behaviour of the probe in response to the addition of gold ions. In the absence of any analyte species, **BOD-Z-EN** displayed a single absorption band at 532 nm (Fig. 1a). Meanwhile, in the fluorescence spectrum of **BOD-Z-EN**, collected upon excitation at 460 nm, a long wavelength emission band at 575 nm was clearly observed. As expected, upon adding AuCl\(_3\) (10 equiv.), we observed a new green emissive compound, as monitored on a thin-layer chromatography plate, was additional clear evidence of the formation of a non-conjugated core BODIPY derivative (Fig. S11, ESI†). Following chromatographic purification, the structure of the new BODIPY derivative was confirmed by NMR spectroscopy and high-resolution mass spectrometry (ESI-TOF) as **BOD-FUR** (quantum yield; \(\Phi_F = 0.90\)), the intramolecular cyclisation product of **BOD-Z-EN** \((\Phi_F = 0.05)\). Our investigation continued with the systematic addition of gold ions to the probe solution. Upon adding Au\(^{3+}\) \((0–10 \text{ equiv.})\), we observed the absorption band at 532 nm decrease gradually, with a concomitant linear increase of a new absorbance band at 503 nm (Fig. 1a). A similar trend occurred in fluorescence emission behaviour; with an increased concentration of Au\(^{3+}\) across a wide concentration range, fluorescence emission intensity at 512 nm increased linearly (Fig. 1b). The probe’s response to AuCl\(_3\) was clearly visible to the naked eye; in the presence of gold species, the probe solution’s orange emission became distinctly green, which we attributed to a structural modification of the dye structure (Scheme 1). A new green emissive compound, as monitored on a thin-layer chromatography plate, was additional clear evidence of the formation of a non-conjugated core BODIPY derivative (Fig. S11, ESI†). Following chromatographic purification, the structure of the new BODIPY derivative was confirmed by NMR spectroscopy and high-resolution mass spectrometry (ESI-TOF) as **BOD-FUR** (quantum yield; \(\Phi_F = 0.90\)), the intramolecular cyclisation product of **BOD-Z-EN** \((\Phi_F = 0.05)\). Our investigation continued with the systematic addition of gold ions to the probe solution. Upon adding Au\(^{3+}\) \((0–10 \text{ equiv.})\), we observed the absorption band at 532 nm decrease gradually, with a concomitant linear increase of a new absorbance band at 503 nm (Fig. 1a). A similar trend occurred in fluorescence emission behaviour; with an increased concentration of Au\(^{3+}\) across a wide concentration range, fluorescence emission intensity at 512 nm increased linearly (Fig. 1b). The response of **BOD-Z-EN** to Au\(^{3+}\) \((e.g., 1 \text{ equiv. of AuCl}_3)\) was exceptionally fast \((< 30 \text{ s})\), and within a couple of minutes, emission intensity \((> 65\text{-fold})\) became completely saturated (Fig. S9, ESI†). An 80-fold enhancement in emission ratio occurred when 10 equiv. of gold was introduced. Under optimum sensing conditions, the detection limit of **BOD-Z-EN** for detecting Au\(^{3+}\) was 293 nM, based on the signal-to-noise ratio \((S/N = 3)\) (Fig. S1, ESI†). The metal ion selectivity test for **BOD-Z-EN** resulted in no obvious spectral changes for competing alkynophilic metal ions.
species, including Ag⁺, Ni²⁺, Pd²⁺, Hg²⁺, Cu²⁺, and other metal species such as Mg²⁺, Mn²⁺, Pb²⁺, Zn²⁺, Cd²⁺, Fe³⁺, K⁺, Li⁺, Ba²⁺, and Ca²⁺. Only the addition of Au³⁺ and, to a lesser extent the addition of Au⁺ resulted in an increase of fluorescence at 512 nm, which obviously implied the high selectivity of BOD-Z-EN to gold ions (Fig. S6, ESI†). Remarkably, other metal ions did not interfere with the detection of gold ions, and the spectral response induced by Au³⁺ ions remained unaffected in the presence of any of those metal species. Such results indicate that BOD-Z-EN can properly detect Au³⁺ ions in mixtures with other related species.¹²

Analogous to reports in the literature,¹⁰ we propose that the sensing mechanism occurs when Au³⁺ activates the triple bond and the subsequent intramolecular exo-dig cyclisation of hydroxyl group to the triple bond, which, in forming a gold intermediate and a rapid isomerisation and protonolysis pathway, affords the BODIPY-furan structure (BOD-FUR) (Scheme 3).

In establishing the most efficient conditions for operation, BOD-Z-EN in the absence of Au³⁺ was quite stable, even in strong acidic and basic conditions. Likewise, the response of BOD-Z-EN to the addition of Au³⁺ ions remained unaffected by altering the pH of the sensing media (Fig. S4, ESI†). As such, BOD-Z-EN operates efficiently over a wide pH range (pH 2–12), especially in physiological conditions (pH 7.4), which fulfils a basic requirement for cell bioimaging.

In sharp contrast to BOD-Z-EN, the E-stereoisomer of the probe, BOD-E-EN, was inert to gold, as well as other metal species, despite being in identical sensing conditions. Consistent with the proposed mechanism and in agreement with reports in the literature, adding Au³⁺ did not alter the emission pattern of BOD-E-EN, thereby indicating the sensing process’s high stereoselectivity (Fig. S10, ESI†).

Relying on the promising photochemical and physical properties of BOD-Z-EN, including its rapid response time, unique gold ion specificity, exceptionally low detection limit, and high fold ratiometric change, among other qualities, we next evaluated its potential for tracking Au³⁺ ions in living cells. To that end, A549 human lung adenocarcinoma cells were incubated at 37 °C first with BOD-Z-EN (5.0 μM) for 20 min, followed by incubation with Au³⁺ (10 μM) for another 20 min. Using fluorescence microscopy, we could clearly monitor the ratiometric sensing behaviour of the probe toward Au³⁺ in the cells (Fig. 2). Consistent with results observed in the solution, following incubation with Au³⁺ the orange emissive cells immediately turned green, thereby unambiguously showing that BOD-Z-EN is cell-membrane permeable and has great potential for use in imaging Au³⁺ in living cells. Moreover, there were no indications of cell damage. Cells were intact and showed healthy spread and adherent morphology throughout the cell imaging process. The potential cytotoxicity of the probe was evaluated using a standard MTT assay with A549 cell line. The cellular viability was estimated to be 100% after 24 h, which showed that the probe (<5 μM) has no cytotoxicity (Fig. S2, ESI†).

In sum, we devised a fluorescent probe that shows a ratiometric fluorescence response to gold ion species with high sensitivity and selectivity over other metal ion species. The fluorescent probe is built upon a BODIPY fluorophore scaffold that uses a highly novel reactive unit (e.g., Z-enynol) for Au³⁺ ions. After structural modification triggered by gold ions—namely, an intramolecular cyclisation—the high wavelength-emitting conjugated probe structure transformed into a non-conjugated, low wavelength-emitting structure, thereby allowing us to recognise gold ions as a ratio of distinct emission wavelengths. Apart from the rapid (<30 s), sensitive (293 nM), and highly specific response to gold species in the solution, the probe proved highly successful in imaging gold ions in living cells.

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Notes and references


12. See ESI†.