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A two-photon fluorescent probe for colorimetric and ratiometric monitoring of mercury in live cells and tissues[†]

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Owing to the extreme toxicity of mercury, methods for its selective and sensitive sensing in solutions, and in live cells and tissues are in great demand. In this study, we developed a naphthalimide-based diphenylphosphinothioyl group-containing fluorescent and colorimetric probe that selectively detects mercury (Hg²⁺). Upon addition of mercury (Hg²⁺) to a solution of the probe, both a colorimetric change from colorless to yellow and a fluorescence change from blue to green (under a 365 nm hand-held UV lamp) occur, both of which can be observed using the "naked-eye". Furthermore, the probe possesses the capability of sensing intracellular mercury in both live cells and tissues using dual-emission channels and twophoton microscopy.

Owing to its wide use in the chemical industry and its extreme toxicity, mercury is a hazardous threat to both human health and the environment.¹ The toxicity of the mercuric ion is caused by its strong binding affinity to thiols in enzymes and proteins, which results in severe and irreversible damage to physiological processes. Consequently, novel strategies for selective and sensitive detection of mercury in the environment and living systems are in great demand. In contrast to several traditional methods that have been developed for this purpose thus far, including those that rely on absorption spectroscopy, gas chromatography-mass spectrometry, voltammetry and atomic absorption/emission spectrometry, those that utilize fluorescence methods possess significant advantages, including instantaneous response, operational simplicity, high selectivity and sensitivity, and real-time detection capability.² Thus, considerable attention

has been given to the development of fluorescent probes for monitoring mercury in aqueous solution and in biological samples. These studies have led to a number of fluorescent chemosensor systems that utilize various mercury mediated chemical processes such as cyclization,³ desulfurization,⁴ oxymercuration⁵ and others.⁶ Though effective, these detection methods have important drawbacks. For example, most of these methods detect mercury in an "off-on" manner. In contrast, because changes in interconnected fluorescent signals are monitored, ratiometric fluorescent probes are capable of detecting analytes with high accuracies because interferences arising from instrumental issues, probe concentrations and ambient environmental changes are minimized.⁷ Notably, Coskun and co-workers reported a significantly ratiometric fluorescent probe with the advantage of an internal charge transfer mechanism utilizing boradiazaindacene dyads.⁶ The probe efficiently detected mercury with larger intensity ratio changes and excellent selectivity. Regrettably, its application in live cells has not been explored. Besides that, though some fluorescent probes have successfully been applied to monitor mercury in vivo, the majority of them developed to date employ one-photon microscopy to detect mercury inside the cells or tissues. In contrast, two-photon microscopy (TPM), in which two near-infrared (NIR) photons serve as the excitation source, possesses significant advantages including reduced photodamage and greater tissue imaging depths.8

Thus, in the investigation described below, we developed a new colorimetric and ratiometric fluorescence probe, which contains a naphthalimide fluorophore and which can be utilized to monitor mercury in both live cells and tissues using two-photon microscopy.

Guided by the results of previous studies,⁹ we designed the naphthalimide containing fluorescent probe, **NAP-PS**, for the detection of mercury. We anticpated that **NAP-PS** would possess several beneficial features, including high stability and cellpermeability (Scheme 1 and Scheme S1, ESI†). Moreover, the diphenylphosphinothioyl moiety was incorporated into a familiar 4-hydroxy-1,8-naphthalimide fluorophore in order to achieve ratiometric detection of mercury. It was envisaged that mercury

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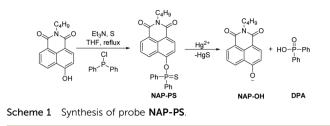
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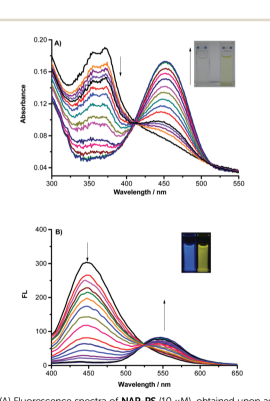
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would promote cleavage of the thiophosphinate ester P–O bond in **NAP-PS** to produce NAP-O⁻, which is an intramolecular charge transfer (ICT) system that displays long emission wavelength owing to the strong electron-donating oxy-anion site and the strong electron-withdrawing naphthalimide group (Scheme 1).¹⁰ To assess the validity of the expectations stated above, we prepared **NAP-PS** by using a four-step sequence, starting with commercially available 4-bromo-1,8-naphthalic anhydride and butylamine (Scheme S1, ESI[†]).

By viewing the absorption spectra shown in Fig. 1A, **NAP-PS** (10 μ M) in HEPES buffer (1.0 mM, pH = 7.4) has an absorption band centered at 370 nm. Upon addition of various amounts of mercury (Hg²⁺) to the solution of **NAP-PS**, the absorption peak shifts in a continuous fashion to 450 nm, associated with a



color change from colorless to yellow (Fig. 1A inset). Moreover, **NAP-PS** emits strong blue fluorescence with a maximum at 450 nm. Addition of mercury causes a significant decrease in the intensity of the emission band at 450 nm and a concurrent increase in the intensity of a peak at 560 nm. Also, the gradual change occurring in emission can be observed as a blue to green color change using the naked eye under a hand-held UV lamp (365 nm) (Fig. 1B inset). At mercury concentrations above 60 μ M, the ratio of the fluorescence intensity at 550 and 450 nm (F_{550}/F_{450}) reaches a maximum (Fig. S1, ESI†). Moreover, the F_{550}/F_{450} ratio is linearly dependent on the mercury concentration in the range of 0–12 μ M (Fig. S2, ESI†). Based on the $3\delta/k$ criterion, the detection limit of the probe for mercury is 43 nM.

The excellent selectivity of **NAP-PS** for sensing mercury was demonstrated by the observation that no fluorescence responses are induced by other metal ions including Zn^{2+} , Cu^{2+} , Mn^{2+} , Pb^{2+} , Al^{3+} , Ca^{2+} , Cr^{3+} , K^+ , Fe^{3+} , Fe^{2+} , Cs^+ , Na^+ , Ni^{2+} , Mg^{2+} , Sr^{2+} , Cd^{2+} and Ag^+ . As the results in Fig. 2 show, only the addition of mercury leads to a significant ratiometric response of the fluorescence intensities at 450 nm to 550 nm. Moreover, other metal ions do not interfere with the detection of mercury by **NAP-PS**.

The time dependence of the fluorescence response of **NAP-PS** was measured in order to determine the observed rate

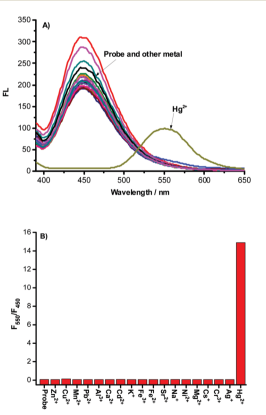


Fig. 1 (A) Fluorescence spectra of **NAP-PS** (10 μ M), obtained upon addition of mercury(II) perchlorate (0–30 μ M) in HEPES buffer (1.0 mM, pH = 7.4); (B) UV-vis spectra of **NAP-PS** (10 μ M) obtained upon addition of mercury(II) perchlorate (0–30 μ M) in HEPES buffer (1.0 mM, pH = 7.4); all spectra were recorded after incubation with different concentrations of mercury(II) perchlorate for 40 min; λ_{exx} = 380 nm, λ_{em1} = 450 nm, λ_{em2} = 550, and slits: 3/5 nm; (A) inset: photograph of solutions of probe **NAP-PS** (10 μ M) before and after the addition of mercury(II) perchlorate (30 μ M); (B) inset: photograph of solutions of probe **NAP-PS** (10 μ M) before and after the addition of mercury(II) perchlorate (30 μ M); (B) inset: photograph of solutions of probe **NAP-PS** (10 μ M) before and after the addition of mercury(II) perchlorate (30 μ M) under a 365 nm hand-held UV lamp.

Fig. 2 (A) Fluorescence spectra of probe **NAP-PS** (10 μ M) before and after the addition of 200 μ M different metal ions in HEPES buffer (1.0 mM, pH = 7.4), $\lambda_{ex} = 380$ nm, slits: 3/5 nm; (B) fluorescence intensity ratios (F_{550}/F_{450}) of probe **NAP-PS** (10 μ M) before and after the addition of 200 μ M different metal ions in HEPES buffer (1.0 mM, pH = 7.4).

constant of its reaction with mercury. As the results in Fig. 3 show, a HEPES buffer (1.0 mM, pH = 7.4) solution of **NAP-PS** (10 μ M) displays very faint fluorescence at 550 nm and a high overall photostability. Following the addition of 2 equivalents of mercury, the intensity of the emission band of **NAP-PS** at 550 nm increases with time and reaches a maximum after 20 min. Pseudo first order kinetic analysis of the time course data leads to an observed rate constant (25 °C) of 9.9 × 10⁻¹ min⁻¹ for the reaction of mercury with **NAP-PS** (Fig. S3, ESI†).

As suggested previously,^{9*a*,11} the chemical process responsible for the absorption and emission spectroscopic changes of **NAP-PS** involves the mercury promoted cleavage of the P–O bond, which generates the strongly fluorescent NAP-O⁻ anion. To confirm the occurrence of this process, a solution of **NAP-PS** (10 μ M) and mercuric perchlorate (20 μ M) in HEPES was monitored using high-resolution mass spectrometry (HRMS) methods (Fig. S4, ESI†). The observed formation of peaks at *m*/*z* 268.0952 and 217.0399 in the HRMS spectrum is consistent with the formation of **NAP-OH** (calcd *m*/*z* 268.0979 for C₁₆H₁₄NO₃) and diphenylphosphinic acid (**DPA**) (calcd *m*/*z* 217.0424 for C₁₂H₁₀PO₂), respectively. Moreover, both the fluorescence and absorption spectra of **NAP-OH** and the mixture following the reaction of **NAP-PS** with mercuric perchlorate are identical (Fig. S5 and S6, ESI†).

The ability to utilize **NAP-PS** to monitor mercury in the solid state was assessed. The initially white **NAP-PS** solid was mixed with solid mercuric perchlorate. Grinding of this solid mixture gradually promotes a color change to yellow (Fig. S7, ESI[†]). The emission color change caused by grinding this solid mixture was examined using a hand-held UV lamp (365 nm). As shown in Fig. S8 (ESI[†]), upon grinding the strong blue emission of solid **NAP-PS** undergoes a change to green-emission.

Encouraged by the excellent performance by **NAP-PS** in sensing mercury in the solution and solid states, we explored the use of the probe for fluorescence imaging in live cells. MTT assays showed that **NAP-PS** and **NAP-OH** have negligible cytotoxicity under the imaging conditions (Fig. S10, ESI[†]). In order to avoid limitations associated with photodamage and autofluorescence caused by using a short excitation wavelength ($\lambda_{ex} = 380$ nm), we first assessed the use of two-photon microscopy (TPM) for determining mercury concentrations. To evaluate the

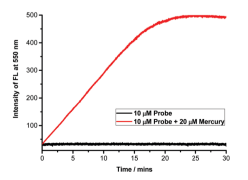


Fig. 3 Time-dependent changes in F_{550}/F_{450} nm for solutions of **NAP-PS** (10 μ M) upon addition of mercuric perchlorate (20 μ M), λ_{ex} = 380 nm, and slits: 3/5 nm.

permeability of NAP-PS in the cells, real-time images were taken at an interval of 5 s (Fig. S11, ESI[†]). No signal was observed in the cells before treatment with NAP-PS, however, the fluorescence intensities of the cells began to increase when loaded with the probe, and the signal was saturated within 1 min. Thus, NAP-PS has high cell permeability, and the fluorescence observed in the cell can be confirmed to be the fluorescence caused by the probe, and not the auto-fluorescence of the cell. To find appropriate twophoton excitation wavelengths, TPEF intensities were assessed by incorporating NAP-PS and NAP-OH in HeLa cells (Fig. S12, ESI⁺). Cells containing NAP-OH display bright emission when exposed to two-photon excitation wavelengths in the 740-800 nm range, while those containing NAP-PS display bright fluorescence when exposed to wavelengths of light below 740 nm (Fig. S12a-c, ESI⁺). The degree of two-photon absorption cross-section was calculated to be 15 and 42 GM respectively (1 GM = 10^{-50} cm⁴ per photon) for NAP-PS and NAP-OH in HEPES buffer (Fig. S9, ESI[†]). Analysis of the results from the TPEF spectroscopy experiments with NAP-PS and NAP-OH containing HeLa cells demonstrated that the detection windows of 400–450 nm (F_{blue}) and 500–600 nm (F_{vellow}), and excitation at 740 nm are optimal for determining $F_{\text{vellow}}/F_{\text{blue}}$ ratios (Fig. S12, ESI⁺). In addition, fluorescence of NAP-PS in HeLa cells is maintained over 1 h with a laser pulse interval of 2 s, indicating that it has the level of robust photostability needed for cell imaging experiments (Fig. S13, ESI⁺).

To show that **NAP-PS** can be employed to detect mercury in live cells, the average emission intensity ratios $F_{\text{yellow}}/F_{\text{blue}}$ were determined using Hg(ClO₄)₂ treated HeLa cells. The cell calibration test indicated that the $F_{\text{yellow}}/F_{\text{blue}}$ ratio of **NAP-PS** was linearly increased by treatment with various concentrations of Hg(ClO₄)₂ (Fig. S14, ESI†). Upon excitation at 740 nm, the respective $F_{\text{yellow}}/F_{\text{blue}}$ values of untreated HeLa cells labeled with probes **NAP-PS** and **NAP-OH** were found to be 0.39 and 4.01 (Fig. 4a and d). Treatment of **NAP-PS** labeled HeLa cells with Hg(ClO₄)₂ (25 µM) for 30 min leads to an increase in the

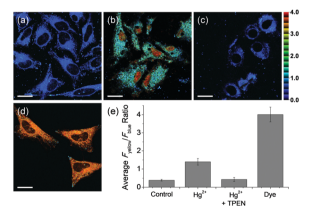


Fig. 4 Pseudo colored ratiometric TPM images of HeLa cells incubated with (a) probe **NAP-PS** (5 μ M) and (d) dye (**NAP-OH**, 5 μ M) for 30 min; HeLa cells were pretreated with (b) Hg(ClO₄)₂ (25 μ M) for 30 min and (c) Hg(ClO₄)₂ and then with TPEN (1 mM) for an additional 30 min before labeling with probe **NAP-PS**; (e) average F_{yellow}/F_{blue} ratios in the TPM images. Images were acquired using 740 nm excitation and emission windows of 400–450 nm (blue) and 500–600 nm (yellow); scale bars = 20 μ M.

Fig. 5 Pseudo colored ratiometric TPM images of rat liver tissues incubated with **NAP-PS** (50 μ M) for 1 h (a) before and (b) after pretreatment with Hg(ClO₄)₂ (250 μ M) for 1 h; (c) average F_{yellow}/F_{blue} ratios in the TPM images; images were acquired using 740 nm excitation and emission windows of 400–450 nm (blue) and 500–600 nm (yellow); scale bars = 20 μ M.

 $F_{\text{yellow}}/F_{\text{blue}}$ ratio to 1.40, which corresponds to a 3.6-fold enhancement (Fig. 4b). Finally, incorporation of the known mercury chelator *N*,*N*,*N*,*N*-tetrakis(2-pyridylmethyl)ethylenediamine (TPEN, 1 mM)¹² reduces the $F_{\text{yellow}}/F_{\text{blue}}$ value to 0.43 (Fig. 4c).

The liver is an important organ for carrying out metabolic and detoxification functions of humans. It has been reported that accumulation of mercury leads to liver dysfunction and related diseases.¹³ In order to further explore the biological value of **NAP-PS**, fluorescence imaging of mercury in live tissues was carried out using TPM. The F_{yellow}/F_{blue} ratio of TPM images of liver tissues stained with **NAP-PS** (50 µM) for 1 h was found to be 0.77 (Fig. 5a). When liver tissues are pretreated with Hg(ClO₄)₂, the F_{yellow}/F_{blue} ratio value increases to 1.70 (Fig. 5b). These results show that **NAP-PS** along with TPM can be employed to directly and quantitatively determine the presence of mercury in both live cells and tissues.

The results presented above show that **NAP-PS** is a highly effective colorimetric and ratiometric fluorescent probe for monitoring mercury. The ratiometric fluorescence response of **NAP-PS** is a consequence of its reaction with mercury, which cleaves the thiophosphinate P–O bond in the probe to form the intramolecular charge transfer (ICT) system, NAP-O⁻. Importantly, the use of this probe as a dual-channel sensor for mercury in both live cells and tissues was demonstrated in this effort.

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Conflicts of interest

There are no conflicts to declare.

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