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A unique approach toward near-infrared fluorescent probes for bioimaging with remarkably enhanced contrast†

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Near-infrared (NIR) fluorescent probes are attractive molecular tools for bioimaging because of their low autofluorescence interference, deep tissue penetration, and minimal damage to sample. However, most previously reported NIR probes exhibit small Stokes shift, typically less than 30 nm, and low fluorescence quantum yield, strictly limited contrast and spatial resolution for bioimaging. Herein, by expanding the π -conjugated system of rhodamine B, while, at the same time, keeping its rigid and planar structure, we reported an efficient NIR dye, HN7, with large Stokes shift of 73 nm and fluorescence quantum yield as high as 0.72 in ethanol, values superior to those of such traditional cyanine NIR dyes as Cy5. Using HN7, living cells, tissues and mice were imaged, and the results showed significantly enhanced contrast, improved spatial resolution, and satisfactory tissue imaging depth when compared to Cy5. Moreover, the nonfluorescent spirocyclic structure of rhodamine B is an inherent component of HN7; therefore, our strategy provided a universal platform for the design of efficient NIR turn-on bioimaging probes for various targets. As a proof-of-concept, two different NIR probes, HN7-N2 and HN7-S for NO and Hg²⁺, respectively, were designed, synthesized, and successfully applied for the imaging of NO and Hg²⁺ in living cells, tissues and mice, respectively, demonstrating the potential bioimaging applications of the new probes. In sum, this new type of dye may present new avenues for the development of efficient NIR fluorescent probes for contrast-enhanced imaging in biological applications.

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Introduction

The application of fluorescence imaging through staining with a fluorescent probe is characterized by high sensitivity, real-time spatial imaging and detection of targets in living cells or tissues with minimal damaging effects. As such, fluorescence imaging has become a facile and powerful tool for the detection of biologically relevant species with the ability to visualize morphological details and monitor various physiological processes in living systems.^{1–8} However, the imaging quality of this technique is strictly dependent on the photophysical properties of fluorescent dyes used for sample staining. It has recently been reported that NIR fluorescent imaging can minimize photodamage to biological samples, increase tissue penetration depth, and minimize interference from

autofluorescence in living systems.^{9–15} Accordingly, many NIR fluorescent dyes have been developed in the past few years.^{16–24} However, most previously reported NIR fluorescent dyes suffer from serious self-quenching and excitation light backscattering as a result of their small Stokes shift, typically less than 30 nm, and most of them bear low fluorescence quantum yields, commonly much less than 0.1 in aqueous solution. Since these photophysical properties afford high background signal and low fluorescence signal for these dyes, low contrast for bioimaging can be expected. However, in order to obtain enhanced image contrast and optimal apparent brightness for bioimaging, the development of NIR dyes with both large Stokes shift and high quantum yield would be the most efficient approach.

Cyanine dyes are classic NIR fluorophores with an odd number of carbons in a conjugated polymethine framework between two nitrogen-containing heterocycles.²⁵ The maximal emission wavelengths of pentamethine cyanines (Cy5) and heptamethine cyanines (Cy7) reach well into the NIR region (>650 nm). However, based on their flexible molecular structure, traditional cyanine dyes suffer from several disadvantages, such as small Stokes shift, limited fluorescence quantum yield, poor photostability and undesirable aggregation.²⁶ Moreover, cyanine dyes exhibit relatively high-lying occupied molecular orbital (HOMO) energy levels,^{27,28} making it difficult to develop

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fluorescence in comparison with traditional rhodamine dyes, were designed and synthesized. Compounds with a flexible structure, including **HN1**, **HN3**, and **HN5** with various amino substituents, were first synthesized (Scheme 1), using 4-aminoacetophenone derivatives as starting materials. By utilizing *N*-substituent 6-amino-3,4-dihydro-1(2*H*)-naphthalenone instead of 4-aminoacetophenone derivative as the starting reactant, we also designed and easily synthesized rigid dyes **HN2**, **HN4**, and **HN6** with an alkyl-ring to limit molecular rotation and vibration (Scheme 1).³⁸ In order to achieve even more improved fluorescence properties, rigid dye **HN7** with a π -conjugated system expanded over that of **HN1–6** was further designed and synthesized by employing a β -diketone compound **7** as an intermediate, which was achieved by Claisen condensation between 6-amino-3,4-dihydro-1(2*H*)-naphthalenone and dimethyl phthalate.³⁹ The structures of dyes **HN1–7** were characterized by ¹H NMR, ¹³C NMR and MS (ESI).

Optical properties

We first tested the optical properties of **HN1–7**, rhodamine B and Cy5 in various solvents, including buffered aqueous solution (pH 7.4 PBS, containing 10% EtOH), EtOH, CH₂Cl₂, DMSO, and CH₃CN. The absorption and emission profiles are illustrated in Fig. 1 and S1–5,† and the corresponding photophysical data are listed in Table 1 and S1–4.† In buffered aqueous

solution, **HN1–7** display broad and intense maximum absorption bands in the 571–600 nm region, which is assigned to the 0–0 band of the S₀ → S₁ transition. Broad and weak absorption bands were found around 420 nm, which could be attributed to the S₀ → S₃ transition. Notably, the absorption of **HN7** exhibited an obvious blue-shift in comparison to Cy5, resulting in a larger Stokes shift in spite of its π -conjugated system which was expanded over that of rhodamine. On the other hand, the molar extinction coefficients (ϵ_{\max}) of all our new dyes are in the range of 120 000 M⁻¹ cm⁻¹ and thus superior to the classic rhodamines.^{33–36} Therefore, as shown in Fig. 1c (top), the rhodamine B solution displays a pink color to the naked eye, while the colors of the **HN1–7** solutions are all much deeper than the classic rhodamine B by their long absorption wavelengths and large molar extinction coefficients.

Since their π -conjugated systems are expanded beyond those of the classic rhodamines, **HN1–6** display maximum fluorescence emission in the far-red region (615–638 nm) in PBS buffered solution. Remarkably, **HN7** displays Cy5-like fluorescence emission with maximum at 673 nm in PBS buffer and covers a significant part of the NIR region. Furthermore, according to the absorption and emission data, the Stokes shift of **HN7** is 73 nm in neutral buffered solution, which is superior to that of the classic rhodamine, fluorescein, **BODIPY**®, and cyanine dyes, all typically less than 30 nm. Such large Stokes shift could reduce background fluorescence for **HN7** and avoid



Fig. 1 Absorption (a) and emission (b) spectra of **HN6** (navy-blue triangle), **HN7** (purple triangle) in pH 7.4 PBS buffer. The spectra of rhodamine B (wine hexagon) and Cy5 (olive-green star) are also shown for comparison. (c) Color and fluorescence images of rhodamine B, **HN1–7**, and Cy5 in pH 7.4 PBS buffer (3.0 μ M, containing 10% EtOH).

Table 1 Photophysical properties of dyes **HN1–7**, rhodamine B and Cy5 in pH 7.4 PBS buffer (containing 10% EtOH)

Dye	$\lambda_{\text{abs}}^a/\text{nm}$	$\lambda_{\text{em}}^b/\text{nm}$	SS ^c	$\epsilon_{\text{max}}/\text{M}^{-1} \text{cm}^{-1}$	Φ_f	$\epsilon_{\text{max}}\Phi_f/\text{M}^{-1} \text{cm}^{-1}$	τ/ns
HN1	571	615	44	117 900	0.06 ^d	7074	0.79
HN2	577	617	40	119 200	0.43 ^d	51 256	0.87
HN3	589	626	37	125 300	0.05 ^d	6265	0.80
HN4	592	624	32	122 100	0.41 ^d	50 061	0.74
HN5	598	638	40	129 500	0.02 ^d	2590	0.66
HN6	599	635	36	123 300	0.37 ^d	45 621	0.71
HN7	600	673	73	137 300	0.25 ^e	34 325	0.88
RB	554	573	19	106 000	0.31 ^f	32 860	0.94
Cy5	649	670	21	250 000	0.03 ^e	7500	0.87

^a The maximal absorption of the dye. ^b The maximal emission of the dye. ^c Stokes shift. ^d Φ_f is the relative fluorescence quantum yield estimated by using cresyl violet as a standard: $\Phi_f = 0.56$ in EtOH.⁴⁰ ^e Φ_f is the relative fluorescence quantum yield estimated by using sulfoindocyanine dye Cy5 as a standard: $\Phi_f = 0.20$ in PBS.⁴¹ ^f Φ_f is the relative fluorescence quantum yield estimated by using rhodamine B as a standard: $\Phi_f = 0.72$ in MeOH.⁴⁰



self-quenching and false-positive signals from excitation back-scattering effect.

Classic rhodamine dyes possess high fluorescence quantum yields, typically larger than 0.5, as a result of their rigid and planar molecular structures. As expected, compounds **HN1**, **HN3**, **HN5** with their flexible structure showed fluorescence quantum yields less than 0.22 in EtOH and less than 0.06 in aqueous solution.⁴⁰ On the other hand, **HN2**, **HN4**, **HN6**, and **HN7** all possess rigid and planar molecular structures similar to rhodamine dyes, but they exhibit higher fluorescence quantum yields than the classic cyanine NIR dyes (e.g., Cy5: $\Phi_f = 0.03$ in buffered aqueous solution), ranging from 0.72 to 0.93 in EtOH and from 0.25 to 0.43 in aqueous solution.⁴¹ To evaluate a newly designed biological imaging reagent, brightness ($\epsilon_{\max}\Phi_f$) is another important parameter to test, as it involves imaging contrast and spatial resolution. Accordingly, the brightness of **HN7** was calculated to reach $34\,325\text{ M}^{-1}\text{ cm}^{-1}$ in buffered aqueous solution, which is superior to that of traditional rhodamine B and Cy5. All these results illustrate that our NIR dye, **HN7**, is suitable for bioimaging applications with enhanced brightness and contrast.

Still another important parameter is photostability. By using continuous irradiation, as described above, we investigated the photostability of **HN2**, **HN4**, **HN6**, **HN7** and Cy5 in PBS buffer. Exactly as anticipated and as shown in Fig. 2a, over 98% of the initial fluorescence intensity of our **HN** dyes was retained after 120 min irradiation, while only 62% of Cy5 was retained, indicating that our rigid **HN** dyes have sufficient photostability for prolonged biological imaging. On the other hand, it is well known that many organic NIR dyes are easily bleached by reactive oxygen species (ROS).⁴² To evaluate the chemical stability of **HN** dyes under these conditions, the fluorescence profiles of **HN2**, **HN4**, **HN6**, **HN7** and Cy5 in the presence of various ROS (HOCl, H₂O₂, TBHP, HO·, *t*BuO·) were also investigated. As shown in Fig. 2b, more than 93% of the initial fluorescence intensity was retained after treatment of **HN** dyes with ROS in excess of 120 min, values much improved of those of Cy5. All these data suggest that our **HN** dyes have excellent photostability and the chemical stability required to resist

photobleaching and common ROS in biological imaging applications.

The pH value around the rhodamine analogues and their spirolactam derivatives typically affects performance because of protonation reaction in acidic conditions or cyclization reaction in alkaline conditions. Therefore, to study the practical applicability of the newly designed **HN** dyes, the effects of pH on their fluorescence response and a representative spirolactam derivative were investigated. As shown in Fig. 3, stable and high fluorescence were observed for pH values ranging from 5.0 to 11.0 for free **HN7**. However, fluorescence did decrease from protonation reaction in strong acidic conditions or the development of nonfluorescent spirocyclization in alkaline conditions, with pK_a at 2.7 and pK_{cycl} reaching 12.0. Moreover, the spirolactam derivative **HN7-N2** showed pK_{cycl} of about 4.1 and predominately existed in the nonfluorescent spirocyclic form with a broad range of pH values over 6.0. Therefore, just like the classic rhodamines and their spiro-ring derivatives,^{33–36} the fluorescence on–off switching of **HN** dyes and their spiro-ring derivatives would seem to have unlimited potential applications based on the robust rhodamine sensing platform.

Density functional theory (DFT) studies

To understand the optical properties of **HN7**, we carried out DFT calculations with the B3LYP exchange functional employing 6-31G* basis sets using a suite of Gaussian 09 programs.³⁷ Fig. 4 shows the representative optimized structure and the HOMO/LUMO molecular orbital plots of **HN7**. The C–C (N or O) bond lengths (in pm) of **HN7**, as determined by DFT calculations, are displayed in Table 2. As shown in Table 2, between the typical C–C single (154 pm) and double (134 pm) bonds, all C–C bond lengths of the conjugated fluorophore core are around 140 pm, which can be attributed to strong electronic delocalization and partial decrease in carbon–carbon bond length alternation (BLA, difference between consecutive single and double bonds) along the π -conjugated system. Such phenomenon can easily be proved by investigating the terminal C–N and C–O bond lengths in the conjugated system. For example, according to the



Fig. 2 (a) Photostability of **HN2** (black), **HN4** (red), **HN6** (blue) and **HN7** (magenta) in pH 7.4 PBS buffer (containing 10% EtOH). The samples were continuously irradiated by a Xe lamp (150 W) at 5 nm slit width at the maximal absorption wavelength of **HN2**, **HN4**, **HN6**, and **HN7**. (b) Chemical stability of **HN2**, **HN4**, **HN6**, and **HN7** in pH 7.4 PBS buffer (containing 10% EtOH). The samples were reacted with various ROS (100 μM) for 120 min (black: HOCl, red: H₂O₂, blue: TBHP, olive-green: HO·, cyan: *t*BuO·).





Fig. 3 (a) The proposed structures and the putative transformations of HN7 and HN7-N2 at various pH values. (b) pH-dependence of the normalized fluorescence intensity of HN7 (red circle) and HN7-N2 (black square).

calculated result based on C–N as a single bond, the bond length of C1–N2 is 147.1 pm, which is close to that of the typical C–N single bond (147 pm). Meanwhile, the bond lengths of N2–C3 and C21–N22 are 136.9 and 136.8 pm, respectively, which are intermediates between the lengths of C–N bond, as a single bond calculated above, and C=N, a double bond around 128 pm. Obviously, these bond length calculations support the strong electronic delocalization along the π -conjugated system of HN7.

Notably, Fig. 4a shows that the benzoic acid moiety is highly twisted and approximately vertical to the fluorophore core with



Fig. 4 (a) DFT optimized structure of HN7. In the ball-and-stick representation, carbon, nitrogen, oxygen, and hydrogen atoms are colored in gray, blue, red, and white, respectively. (b) Molecular orbital plots (LUMO and HOMO) and HOMO/LUMO energy gaps of HN7.

a dihedral angle of 78.7°, whereas the planar angle of the fluorophore core is confirmed to be 175°. Meanwhile, the C11–C12 length is 150.0 pm (Table 2), which is close to the typical C–C single bond (154 pm). All these data demonstrate that the benzoic acid moiety and the fluorophore core are effectively electronically decoupled. Thus, the absence of electronic conjugation between benzoic acid moiety and fluorophore core agrees very well with the finding that benzoic acid moiety makes no contribution to the HOMO/LUMO of HN7 (see Fig. 4). But interestingly, the benzoic acid moiety carries up some electron density of the LUMO molecular orbital plots of HN7, which will greatly stabilize the LUMO and reduce the energy bandgap of the fluorophore core. In other words, this may interpret the large Stokes shift of HN7. Furthermore, the fluorophore core could maintain its planar configuration with a planar angle of 175°, thus leading to such superior optical properties as high fluorescence quantum yield. Taken together, the quantum chemical calculations fit well with the optical performance of HN7.

Biological imaging studies

Our new NIR HN7 dye possesses several excellent photophysical properties favorable for bioimaging applications. First, NIR fluorescence emission is always more suitable for biological imaging by its avoidance of autofluorescence effects of biological matrix. Second, the large Stokes shift could resist potentially serious self-quenching and false-positive signals from excitation backscattering. Third, its high fluorescence quantum yield

Table 2 Representative X–X (C, N or O) bond lengths (pm) of HN7 as determined by DFT calculations

X–X bond	Bond lengths (pm)	X–X bond	Bond lengths (pm)	X–X bond	Bond lengths (pm)
C1–N2	147.1	C8–C9	143.0	C14–C16	142.8
N2–C3	136.9	C9–C10	140.0	C16–C17	142.8
C3–C4	142.7	C10–C11	141.9	C16–C18	141.7
C3–C5	142.9	C11–C12	150.0	C18–C19	137.9
C5–C6	138.0	C11–C13	141.4	C19–C21	143.0
C6–C8	141.6	C13–C14	140.0	C20–C21	142.7
C8–C7	142.6	C14–O15	138.3	C20–C17	138.3
C4–C7	138.4	C9–O15	138.1	C21–N22	136.8



should be appropriate for biological imaging with enhanced brightness and contrast. Therefore, to test these properties, **HN7** was applied for bioimaging in living systems.

To evaluate the imaging performance of **HN7**, it was first utilized in living cells, with rhodamine B and Cy5 chosen as reference compounds. Selectable confocal microscopy lasers were limited to 405, 458, 488, 515, 543, 635 nm excitation; therefore, by selecting excitation wavelength 543 nm for rhodamine B, **HN7** and Cy5, some exciting light energy was unavoidably lost. It should be noted that 635 nm was also selected as the excitation wavelength to fit the absorption of Cy5 for comparison. Meanwhile, the emission windows for rhodamine B (560–630 nm) and **HN7**, Cy5 (650–720 nm) were also confirmed. As a proof-of-concept, HeLa cells were incubated with **HN7**, rhodamine B and Cy5 (four groups: 0, 1, 3, and 10 μM , respectively) at 37 $^{\circ}\text{C}$ for 60 min, and the fluorescent imaging results were shown in Fig. 5a–t. Obviously, all the applied fluorescent dyes proved to be membrane permeable, emitting their fluorescence in the cytoplasm, rather than nuclear regions. Furthermore, using the same parameters, different levels of imaging brightness and contrast were observed for these compounds at various concentrations. Relative fluorescence intensity data were shown in Fig. 5u. First, weak fluorescence belonging to either cellular autofluorescence or excitation light backscattering could be observed in Fig. 5a, where it was particularly serious, as well as Fig. 5i and q.⁴³ Thus, rhodamine B showed satisfactory brightness, even at the low concentration of 1 μM . However, the imaging contrast, as determined by the relative fluorescence intensity enhancement ratios to blank, was critically limited by 7-, 12.8-, and 22.3-fold. In contrast, because of its large Stokes shift and high

fluorescence quantum yield, under two different excitation wavelengths of 543 nm and 635 nm, **HN7** not only showed brightness comparable to that of rhodamine B, but also considerable relative fluorescence intensity enhancement ratios as high as 85-fold, with a high enhancement ratio of 24.6-fold, even at the low concentration of 1 μM . While the relative fluorescence intensity enhancement ratios of Cy5 were tolerable, imaging brightness was very weak and not suitable for visual observation. Therefore, compared to classic rhodamine B and Cy5, our new **HN7** bore excellent imaging properties, including NIR emission, high brightness and enhanced contrast. As such, these results demonstrate the potential *in vivo* imaging applications of **HN7**, especially for trace fluorescence imaging.

The cytotoxicity of **HN7** in HeLa cells was also examined using the MTS (3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium) assay. Fig. S5† showed that **HN** dyes exhibited low cytotoxicity after incubation in HeLa cells for 48 h. Such desirable property may also render the new **HN** dyes suitable for applications in living systems.

To evaluate the imaging depth of the new NIR dye, **HN7** was further applied for tissue imaging of rat liver frozen slices, and the changes of fluorescence signal intensity with scan depth were then determined by confocal microscopy (Olympus, FV1000) in z-scan mode. As shown in Fig. 6, experimental results showed that **HN7** possessed satisfactory tissue imaging capability at depths of 40–120 μm . Meanwhile, Cy5 was capable of tissue imaging at depths of 75–100 μm , indicating that the tissue penetration capability of **HN7** was slightly superior to that of Cy5. It is worth noting that **HN7** showed obviously enhanced brightness and contrast when compared to Cy5 for



Fig. 5 Confocal fluorescence images of HeLa cells incubated with various dyes at 37 $^{\circ}\text{C}$ for 60 min (from left to right: 0, 1, 3, 10 μM): (a–d) excitation at 543 nm for red emission of rhodamine B (560–630 nm); (e–h) excitation at 543 nm for red emission of **HN7** (650–720 nm); (i–l) excitation at 635 nm for red emission of **HN7** (650–720 nm); (m–p) excitation at 543 nm for red emission of Cy5 (650–720 nm); (q–t) excitation at 635 nm for red emission of **HN7** (650–720 nm). Scale bar = 30 μm . (u) Relative fluorescence intensity obtained from the selected ellipse region of (a–t), respectively, averaged and plotted as a ratio to blank (a, e, i, m, q).





Fig. 6 Depth fluorescence images of Cy5 (a), 5 μM and HN7 (b), 5 μM in tissues. The changes of fluorescence intensity with scan depth were determined by spectral confocal microscopy (Olympus, FV1000) in the z-scan mode (from 0 to 150 μm ; step size: 1 μm). The images were collected at 650–720 nm (red channel). Scale bar = 120 μm .

imaging frozen rat liver tissue slices. These data demonstrated the potential of HN7 in tissue imaging applications.

NIR light causes only minimal photodamage to samples and has the advantage of deep tissue penetration and reduced interference from autofluorescence in living animals; therefore, NIR dyes are more favorable for *in vivo* imaging. Compared to the traditional NIR dye Cy5, our unique HN7 dye exhibits a similar fluorescence emission spectrum, but has additional advantages, such as larger Stokes shift, higher fluorescence quantum yield, greater photostability and satisfactory aqueous solubility, all of which could provide low background, high fluorescence signal and enhanced contrast for bioimaging

applications in living animals. As confirmation, we further tested the analytical characteristics of Cy5 and HN7 *in vivo*. Fluorescence images of four-week-old male BALB/c nude mice were first taken with two well-separated wavelengths of an excitation filter 605 nm and a Cy5.5 emission filter. Significantly, as shown in Fig. 7a, without any intraperitoneal injection, the blank image showed extremely weak endogenous fluorescence intensity, which could be attributed to weak biological autofluorescence minimal excitation backscattering effect. Next, by intraperitoneal injection with HN7 and Cy5 (5 nmol), compared to the blank group, a considerable enhanced fluorescence intensity of 42-fold was observed for the HN7-



Fig. 7 Fluorescent images of mice (pseudocolor). Thirty minutes after dye injection, mice were imaged using a Caliper VIS Lumina XR small animal optical *in vivo* imaging system with an excitation filter 605 nm and a Cy5.5 emission filter. (a) The blank group; (b) the Cy5-incubated group; (c) the HN7-incubated group; (d) quantification of fluorescence emission intensity from the selected circle region of (a–c) was averaged and plotted as a ratio to blank.



incubated group (Fig. 7c), whereas fluorescence enhancement of the Cy5-incubated group compared to the blank group was estimated to be only 9.2-fold by its low fluorescence quantum yield (Fig. 7b). These results demonstrate that this novel **HN7** dye is superior to the traditional NIR Cy5 dye for bioimaging applications in living animals.

Application of the HN dye for design of fluorescent probes

Apart from its excellent photophysical properties, **HN7** also possesses a rhodamine-like spirocyclic structure, which provides a universal platform for the design of efficient NIR turn-on bioimaging probes for various targets. To demonstrate the feasibility of this probe design, two different NIR probes, **HN7-N2** and **HN7-S**, were designed and synthesized for sensing NO and Hg²⁺ (Scheme 2), by utilizing a phenylenediamine unit for sensing NO⁴⁴ and a thiolactone for sensing Hg²⁺,^{45,46} respectively. NO and Hg²⁺ were chosen as model targets. NO has been discovered to play a critical role in the cardiovascular

system, the central and peripheral nervous systems and the immune systems. It has also been recognized as an important molecule in several physiological and pathological processes.^{47,48} Hg²⁺ is one of the most toxic heavy metal ions, as demonstrated by even a small amount of Hg²⁺ ion which can cause permanent deterioration of brain, heart, kidney, stomach, intestine, central nervous and endocrine systems.^{49,50} Therefore, considerable attention could be devoted to our new fluorescent probes for the detection of NO and Hg²⁺ with sufficient selectivity, sensitivity and imaging performance.

As shown in Fig. 8a, similar to classic rhodamine spirocyclic derivatives, the **HN7-N2** free probe in buffered aqueous solution was nearly colorless and fluorescence-inactive, indicating predominance of the spirocyclic form. However, with the introduction of NO (from diethylamine NONOate sodium salt), **HN7-N2** instantly exhibited blue color, as determined by the naked eye, accompanied by considerable red fluorescence (Fig. 8a inset). A significant turn-on absorption (Fig. S6†) and fluorescence response (Fig. 8a) was observed following the NO-



Scheme 2 (a) NO-induced ring opening of **HN7-N2**. (b) Hg²⁺-induced ring opening of **HN7-S**.

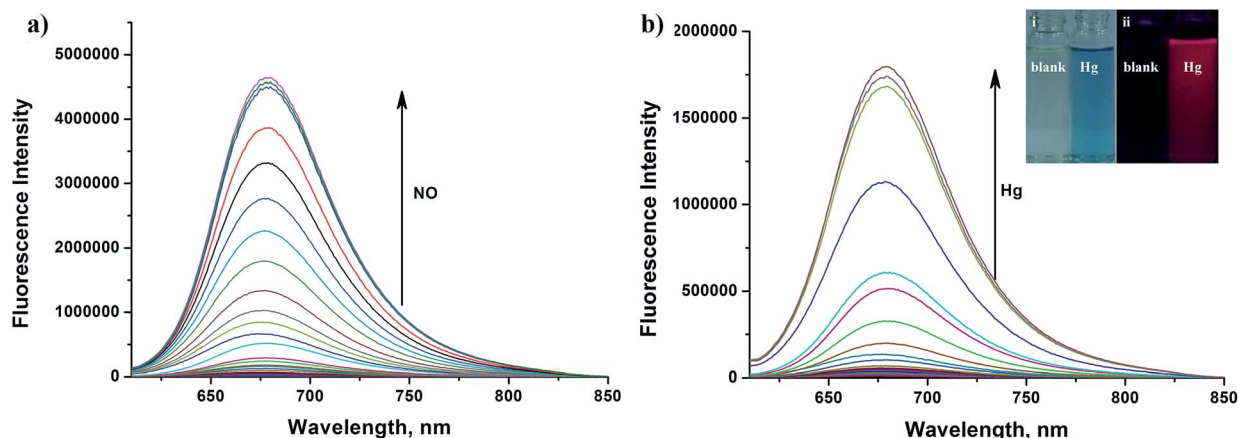


Fig. 8 (a) Fluorescence spectra of **HN7-N2** (5 μM) in the presence of various concentrations of NO (from diethylamine NONOate sodium salt, 0–100 μM) in PBS buffer (0.01 M, pH 7.4, containing 10% EtOH) with excitation at 600 nm. Inset: change in color (i) and fluorescence (ii) in the absence of NO (left) and in the presence of 100 μM NO (right). (b) Fluorescence spectra of **HN7-S** (5 μM) in the presence of various concentrations of Hg²⁺ (0–10 μM) in PBS buffer (0.01 M, pH 7.4, containing 10% EtOH) with excitation at 600 nm. Inset: change in color (i) and fluorescence (ii) in the absence of Hg²⁺ (left) and in the presence of 10 μM Hg²⁺ (right).



induced ring-opening reaction of **HN7-N2** over other biologically relevant species (Fig. S7†). Interestingly, upon the addition of 20 equiv. of NO, a 1500-fold fluorescence enhancement at 676 nm was observed. Fig. S8† shows the calibration curve of the **HN7-N2** probe, and it exhibits a linear range from 20 nM to 1 μ M, with a detection limit of 8 nM (3s/slope). Similar results were also obtained when using the **HN7-S** probe for Hg^{2+} detection (Fig. 8b and S10–12†). Obviously, our analytical data *in vitro* is as well as the reported rhodamine-based probes (Tables S5–S6†).^{44–46} Thus, these results indicate that **HN7** could be employed as a universal platform to design efficient NIR bioimaging probes for various analytes.

The excellent photophysical properties of **HN7** make both **HN7-N2** and **HN7-S** probes especially favorable for bioimaging applications. To demonstrate the practical applicability of the probes in biological systems, we chose **HN7-N2** as the model probe and carried out fluorescence imaging experiments in living cells, tissue slices and mice. First, intracellular NO was monitored using confocal fluorescence imaging. In Fig. 9, HeLa cells incubated with **HN7-N2** (10 μ M) for 30 min showed no observable fluorescence signal, indicating that the **HN7-N2** probe maintained its spirocyclic form in cells. When HeLa cells preincubated with **HN7-N2** were further treated with different concentrations of NO donor, diethylamine NONOate (10, 20, 50 μ M), a dramatic increase in fluorescence brightness was observed, suggesting that the new probe was cell permeable and could be used as an imaging tool to quantify the concentration of NO in living cells.

We further used **HN7-N2** for NO imaging in tissue of rat liver frozen slice. The changes of fluorescence intensity with scan depth were determined by confocal microscopy (Olympus, FV1000) in the z-scan mode. As shown in Fig. 10, in the presence of NO, the **HN7-N2** probe was capable of tissue imaging at depths of 45–110 μ m. These data showed that the **HN7** probe had good tissue penetration and staining ability. **HN7-N2** was also used for NO detection *in vivo*. For four-week-old male BALB/c nude mice, the control group was only given an



Fig. 9 Confocal fluorescence imaging of NO in HeLa cells with **HN7-N2** (10 μ M) for 30 min and then NO donor, diethylamine NONOate (from left to right: 0, 10, 20, 50 μ M, respectively) for 30 min. Excitation at 543 nm for red emission of **HN7-N2** (650–720 nm).



Fig. 10 Depth fluorescence images of 10 μ M **HN7-N2** for NO detection in tissues were obtained with spectral confocal microscopy (Olympus, FV1000). The changes of fluorescence intensity with scan depth were determined by spectral confocal microscopy (Olympus, FV1000) in the z-scan mode (from 0 to 150 μ m; step size: 1 μ m). The images were collected at 650–720 nm (red channel). The slice was cultured with **HN7-N2** (5 μ M) for 60 min and then NO (50 μ M) for another 60 min.

intraperitoneal injection of **HN7-N2** (20 nmol). The experimental group was first given an intraperitoneal injection of **HN7-N2** probe (20 nmol), followed by an intraperitoneal injection with NO donor, diethylamine NONOate (50 nmol), after 30



Fig. 11 Fluorescent images of mice (pseudocolor). The mice were imaged using a Caliper VIS Lumina XR small animal optical *in vivo* imaging system with an excitation filter 605 nm and a Cy5.5 emission filter. Left: imaging of control group after intraperitoneal injection of **HN7-N2** probe (20 nanomoles) for 30 min; right: intraperitoneal injection of **HN7-N2** probe (20 nanomoles) for 30 min, followed by intraperitoneal injection of NO donor (50 nanomoles). The mice were imaged 30 min after NO donor injection.



min. The mice were then imaged using a Caliper VIS Lumina XR small animal optical *in vivo* imaging system with an excitation filter 605 nm and a Cy5.5 emission filter. As shown in Fig. 11, the control group showed weak fluorescence, indicating that the probe maintained the fluorescence-inactive spirolactam form *in vivo*. However, mice treated with both **HN7-N2** and **NO** exhibited significantly stronger fluorescence intensity than the control group, indicating that **HN7-N2** could be applied for imaging **NO** *in vivo*.

Fluorescence imaging experiments of **HN7-S** for Hg^{2+} in living cells, tissue slices and mice were also performed, and results were similar to those of **HN7-N2** for **NO** were obtained (Fig. S13–15†). All these results indicated that **HN7** is an efficient and universal platform for the design of NIR bioimaging probes for various analytes with enhanced brightness and contrast.

Conclusions

In summary, by introducing a practical strategy of constructing a rigid molecular structure, we have successfully developed a novel NIR functional fluorescent dye, **HN7**, with excellent photophysical properties, such as large Stokes shift and high fluorescence quantum yield, which could afford significantly enhanced image contrast and apparent brightness in biological imaging applications when compared to traditional rhodamine B and Cy5. Furthermore, the spirocyclic structure of rhodamine B is inherent in **HN7**; as such, our strategy provided a universal platform for the design of efficient NIR fluorescent turn-on bioimaging probes for various targets. As a proof-of-concept, two different NIR probes, **HN7-N2** and **HN7-S** for **NO** and Hg^{2+} , were designed, synthesized, and successfully applied for the imaging of **NO** and Hg^{2+} in living cells, tissues and mice, respectively, demonstrating the potential bioimaging applications of the new probes. Thus, the new type of dye presented herein may open up new avenues for the development of efficient NIR fluorescent probes for contrast-enhanced biological imaging applications.

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