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A near-infrared ratiometric fluorescent probe for cysteine detection over glutathione indicating mitochondrial oxidative stress in vivo



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ABSTRACT

We establish a near-infrared (NIR) ratiometric fluorescent probe Cy-NB for the selective detection of cysteine (Cys) over glutathione (GSH) and homocysteine (Hcy) in mitochondria to indicate oxidative stress. Heptamethine cyanine dye is chosen as the fluorophore of Cy-NB whose emission locates in NIR region. And p-nitrobenzoyl is employed as the fluorescent modulator due to its capability of selective-Cys response. Once triggered by Cys, the uncaged p-nitrobenzoyl rearranges the polymethine π -electron system of the fluorophore, which leads to a remarkable spectrum shifts in absorption and emission profiles. Taking advantage of these spectroscopic properties, we construct a ratiometric fluorescent signal for the detection of Cys with a detection limit of 0.2 μ M within 5 min. Our probe Cy-NB can sensitively detect the mitochondrial Cys pool changes under different oxidative stress status in HepG2 cells. We also successfully employ Cy-NB to imaging Cys level changes in living mice. It suggests that mitochondrial Cys can be used as an oxidative stress biomarker with simple potential clinical applications. And our probe Cy-NB is of great potential for further utilizing in exploring the physiological function of Cys in biological systems.

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1. Introduction

As one of the intracellular biothiols, cysteine (Cys) plays vital roles in various metabolic processes of human beings, such as catalysis, heavy metal binding, protein turnover, and signal transduction (Reddie and Carroll, 2008). Along with glutathione (GSH), Cys holds important redox pool to regulate reactive oxygen species (ROS) homeostasis, which mainly occurs on mitochondria respiratory chain (Balaban et al., 2005; Schwarzländer and Finkemeier, 2013). The mitochondrial dynamic antioxidant defenses network by Cys is independent on GSH system. Compared with the GSH, Cys is more effective to be oxidized (Jones et al., 2004). And Cys can translate to CySS for defending oxidative stress preferentially when cells are confronted with oxidative stress and ROS can be tightly regulated (Winterbourn, 2008). Furthermore, Cys is also a limiting factor for GSH synthesis which also plays a significant role in the maintenance of redox status (Vincent et al., 1999). All the above aspects reveal that mitochondrial Cys is a key factor for mitochondrial oxidative stress defense. And the

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concentration of Cys is very significant and sensitive to mitochondrial oxidative stress (Armstrong et al., 2004). Once ROS are over-produced and over-whelm the antioxidant defenses capacities, there will be a rapidly depletion of mitochondrial Cys, and then the mitochondrial oxidative stress occurs, which further leads to variety of diseases including Alzheimer's disease, Parkinson's disease and neurodegenerative diseases (Markesbery, 1997; Yee et al., 2014). That is, the levels of mitochondrial Cys can be applied to indicate oxidative stress status potentially. Therefore, it is crucially and urgently to develop specific fluorescence probes for the detection of mitochondrial Cys.

To date, various methods have been established to detect Cys such as colorimetry, electrochemical techniques, and fluorescence methods (Zhang et al., 2007; Lee et al., 2008; Pu et al., 2010; Jung et al., 2012; Kong et al., 2013; Dai et al., 2014; Su et al., 2014; Wang et al., 2015). Compared with other technologies, the fluorescent methods possess high sensitivity, less invasiveness and real-time imaging, which are of great advantage to analyze Cys in cells (Lin et al., 2008; Li et al., 2009, 2013; Long et al., 2011; Lu et al., 2011; Yuan et al., 2011; Wang et al., 2012a,2012b; Yin et al., 2013; Zhu et al., 2014). However, there still remains quite a challenge to distinguish mitochondrial Cys from GSH, because the

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concentrations of mitochondrial Cys ($\sim500~\mu M)$ are much lower than those of GSH ($\sim10~mM$) (Ubuka et al., 1992). Additionally, even the low concentration of mitochondrial Hcy ($\sim50~nM)$ in cells, the similar structure of Hcy with Cys makes it a competitive interference which cannot be ignored in the detection of mitochondrial Cys. Therefore, the desirable probe should function exclusively to Cys detection over GSH and Hcy in cells.

In this work, we report a ratiometric near-infrared (NIR) fluorescent probe Cy-NB for the detection of mitochondrial Cys over GSH and Hcy. The probe Cy-NB exhibits a rapid colorimetric and ratiometric fluorescence response to mitochondrial Cvs by modulating the polymethine π -electron system. The ratiometric signal allows the determination at two emission channels and provides a built-in correction to avoid environment interference (Ueno and Nagano, 2011). Utilizing these advantages, we employ Cy-NB to detect mitochondrial Cys in living cells. The probe Cy-NB can be used to monitor mitochondrial Cys level changes successfully under different oxidative stress status in living cells. Additionally, as a NIR fluorescent probe, Cy-NB owns great advantages for bioimaging for its ability to avoid the autofluorescence of organisms and detect deeply into tissues (Escobedo et al., 2010). We also utilize Cy-NB to image Cys level changes in living mice.

2. Materials and methods

2.1. Chemicals and instruments

Unless stated otherwise, all chemicals used in synthesis are analytical reagent grade. Milli-Q water (18.2 M Ω cm) was used. Thin-laver chromatography (TCL) was performed on silica gel plates. Silica gel P60 (SiliCycle) was used for column chromatography (Hailang, Yantai) 200-300 mesh. UV-vis spectra were measured on a µ-Quant microplate reader Nanodrop 2000C (Thermo Scientific, USA) with a 1 cm quartz cell. Fluorescence spectra were quantitatively measured by FluoroMax-4 spectrofluorometer with a xenon lamp and 0.5 cm quartz cells. High-resolution mass spectra were carried on LCQ Fleet LC-MS System (Thermo Fisher Scientific). ¹H NMR, ¹³C NMR spectra were carried on a Bruker spectrometer. The fluorescence images of HepG2 cells were taken by a confocal laser scanning microscope (Japan Olympus Co., Ltd) with an objective lens (\times 60). The fluorescence images of living BALB/c mice were obtained by in vivo imaging system (Bruker). All pH were determined by ph-3 m (Lei Ci Device Works, Shanghai, China).

2.2. Synthesis

2.2.1. Synthesis of ketone-CY

Heptacyanine Cy.7.Cl (230 mg, 0.45 mmol) which has already been synthesized in our laboratory (Wang et al., 2012a, 2012b) and sodium acetate (100 mg, 1.2 mmol) which dissolved in anhydrous N,N-dimethylformamide (15 mL) were heated at 90 °C for 6 h under Ar atmosphere. After the mixture cooled to the room temperature, the crude mixture was filtered, concentrated and evaporated under reduced pressure and subsequently dried with vacuum pumping to obtain red product. Then the ketone-Cy (150 mg) was purified by silica chromatography eluted with CH_2Cl_2/CH_3OH (8:2, v/v). ¹H NMR (500 MHz, $CDCl_3-D_1$) δ (ppm): 7.65–7.63 (d, 2H), 7.46–7.44 (d, 2H), 6.66–6.94 (m, 2H), 6.90–6.76 (t, 2H), 6.50-6.46 (m, 2H), 4.24-4.22 (m, 2H), 3.70-3.67 (m, 4H), 2.53-2.51 (t, 4H), 1.74-1.66 (s, 12H), 1.41-1.35 (m, 2H), 1.29-1.26 (t, 6 H). ¹³C NMR (125 MHz, CDCl₃-D₁) δ (ppm): 190.37, 178.05, 139.02, 131.23, 130.92, 128.84, 127.57, 122.03, 121.86, 109.30, 97.60, 65.57, 30.58, 29.66, 29.36, 24.36, 22.69, 14.12, 13.73. LC-MS (ESI+): $m/z C_{34}H_{40}N_2O$ calcd. 492.3141, found $[M+H]^+$ 493.3217. Elemental analysis calculated (%) for $C_{34}H_{40}N_2O$: C, 82.9; H, 8.2; N, 5.7; O, 3.2; found: C, 88.8; H, 8.3; N, 5.6; O, 3.3.

2.2.2. Synthesis of CY-NB

Compound 1 (100 mg, 0.19 mmol) and triethylamine (278 µL, 0.6 mmol) were dissolved in 15 mL anhydrous CH₂Cl₂ at 0 °C, then p-nitrobenzoyl chloride (10 mL) was added dropwise and kept stirring at 0 °C for 30 min. Then the mixture was warmed to room temperature and stirred overnight. The solution was diluted with CH₂Cl₂ (30 mL) for extraction operation. The solvent was removed in vacuo to obtain a crude mixture deep green solid. Finally, Cy-NB was isolated by silica chromatography eluting with CH₂Cl₂/CH₃OH (8:2, v/v) as a green solid (40 mg, 38% yield). ¹H NMR (500 MHz, $C_2D_6SO-D_6$) δ : 8.58–8.54 (m, 3H), 7.72–7.17 (m, 8H), 6.23–6.21 (d, 1H), 4.91 (s, 1H), 4.32-4.29 (m, 3H), 4.14-4.11 (m, 2H), 3.47 (m, 2 H), 2.04 (t, 4H), 1.43-1.24 (m, 20H). ¹³C NMR (125 MHz, $C_2D_6SO-D_6$) δ (ppm): 171.16, 162.21, 158.99, 151.63, 141.65, 140.81, 139.81, 133.36, 131.30, 128.93, 128.81, 125.36, 124.55, 122.12, 122.09, 110.82, 100.77, 65.53, 60.36, 48.94, 39.86, 30.54, 27.87, 24.61, 21.02, 20.76, 19.16, 14.17, 13.70, 12.41. LC-MS (ESI+): m/z C₄₁H₄₄N₃O₄⁺ calcd 642.3326, found [M+] 642.3326. Elemental analysis calculated (%) for C₄₁H₄₄N₃O₄⁺: C, 76.6; H, 6.9; N, 6.5; O, 10.0; found: C, 76.7; H, 6.8; N, 6.6; O, 9.9.

2.2.3. Synthesis of CY-B

Compound 1 (100 mg, 0.19 mmol) and triethylamine (278 µL, 0.6 mmol) were dissolved in 15 mL anhydrous CH_2Cl_2 at 0 °C, then benzoyl chloride (10 mL) was added dropwise and kept stirring at 0 °C for 30 min. Then the mixture was warmed to room temperature and stirred overnight. The solution was diluted with CH₂Cl₂ (30 mL) for extraction operation. The solvent was removed in vacuo to obtain a crude mixture deep green solid. Finally, Cy-B was isolated by silica chromatography eluting with CH₂Cl₂/CH₃OH (8:2, v/v) as a green solid (40 mg, 42% yield). ¹H NMR (500 MHz, $C_2D_6SO-D_6$) δ : 8.35–8.33 (*m*, 1H), 7.72–7.22 (*m*, 11H), 6.13–6.10 (*m*, 1H), 4.31–4.28 (*m*, 5H), 4.15–4.06 (*m*, 3H), 2.61 (*m*, 4H), 1.74–1.25 (m, 20H). ¹³C NMR (125 MHz, $C_2D_6SO-D_6$) δ : 167.66, 141.59, 140.91, 140.76, 135.05, 132.27, 130.91, 130.13, 128.80, 125.29, 122.17, 110.59, 100.16, 65.51, 50.40, 48.98, 41.00, 30.53, 27.79, 24.52, 23.71, 23.36, 19.14, 13.70. LC-MS (ESI+): $m/z C_{41}H_{45}N_2O_2^+$ calcd 597.3476, found [M+] 597.3479. Elemental analysis calculated (%) for C₄₁H₄₅N₂O₂⁺: C, 82.4; H, 7.6; N, 4.7; O, 5.3; found: C, 82.3; H, 7.7; N, 4.5; O, 5.5.

2.3. Spectroscopic data

The probe Cy-NB, Cy-B, Rhodamine123 dye, Hoechst 33258 dye were solute in dimethyl sulfoxide (DMSO) and maintained in refrigerator at 4° C. Stock solutions (1.0 mM) of amino acids including cysteine (Cys), homocysteine (Hcy), glutathione (GSH), cysteine (CySS), alanine (Ala), arginine (Arg), glycine (Gly), isoleucine (Ile), lysine (Lys), proline (Pro), serine (Ser), and valine (Val) were prepared in twice-distilled water. Absorption spectra were obtained with 1.0-cm cuvette cells. The probe Cy-NB was added to a 10.0-mL color comparison tube. After dilution to 0.5 µM with 5 mM HEPES buffer, different concentrations of Cys were added. The mixture was incubated at 37° C for 5 min before measurement. Fluorescence spectra were obtained with a 1.0-cm quartz cells by Xenon lamp. The probe Cy-NB was added to a 10.0-mL color comparison tube. After diluted to 0.5 μ M with 5 mM HEPES buffer, different concentrations of Cys were added. The mixture was incubated at 37 °C for 5 min before measurement.

2.4. Cell culture and confocal imaging

HepG2 cells were seeded at a density of 1×10^6 cells mL⁻¹ in RPMI 1640 Medium supplemented with 10% fetal bovine serum (FBS), NaHCO₃ (2 g/L), and 1% antibiotics (penicillin/streptomycin, 100 U/ml). Cultures were maintained at 37 °C under a humidified atmosphere containing 5% CO₂/95% air. The cells were sub-cultured by scraping and seeding on 18 mm glass coverslips in culture dish. The fluorescence images of HepG2 cells were taken by a confocal laser scanning microscope (Japan Olympus Co., Ltd) with an objective lens (× 60). Excitation of HepG2 cells at 559 nm and 635 nm were carried out with a HeNe laser and the emission was collected from 580 nm to 680 nm and 700 nm to 800 nm, respectively. In the case of mitochondria and nucleus staining, excitation at 405 nm and 515 nm the emission was collected from 420 nm to 480 nm and 520 nm to 550 nm, respectively.

2.5. Fluorescent imaging in living mice

All the BALB/c mice were selected and divided into two groups. In the control group a, the peritoneal cavities of BALB/c mice were injected with 50 µL solution (DMSO/saline=1:9, v/v). And the BALB/c mice in group b were injected into their peritoneal cavities with 50 µL 1 mM Cys (DMSO/saline=1:9, v/v). Thirty minutes later, the probe Cy-NB (1 µM, 50 µL in 1:9 DMSO/saline v/v) was injected into peritoneal cavities of the two group BALB/c mice for another 30 min. Fluorescence images were constructed from fluorescence collection channel 1 (600 nm to 700 nm, λ_{ex} =530 nm) and channel 2 (750 nm to 850 nm, λ_{ex} =735 nm) using in vivo imaging system (Bruker). Additionally, we merged the fluorescence image with the corresponding X-ray image to clearly display the reaction site of the mice.

3. Results and discussion

3.1. Design of CY-NB

We have developed a ratiometric near-infrared fluorescent probe Cy-NB for mitochondrial Cys detection with a high signalto-noise (S/N) ratio. We choose heptamethine cyanine as fluorophore for its NIR emission, high molar absorption coefficient, low biological toxicity, good biocompatibility and mitochondria-selectivity. And p-nitrobenzoyl is equipped as the modulator which selectively responds to Cys. Cyanine dye mainly exists as ketone-cy under physiological conditions. While integrated with p-nitrobenzoyl, the ketone-cy changed to enol form and the spectrum properties is changed obviously by modulating the intramolecular polymethine π -electron system (Guo et al., 2012). Once triggered by Cys, the cyanine dye can be released from Cy-NB and comes back to ketone-cy form which recovers the polymethine π -electron system (Scheme 1). Therefore, this spectrum property of cyanine makes the probe Cy-NB exhibit a remarkable emission shift and sensitive to Cys.

3.2. Proposed detection mechanism of CY-NB

The proposed detection mechanism of probe Cy-NB for the detection of Cys is shown in Scheme S1. After Cys added into the solution, the ester of Cy-NB is selectively attacked by the sulfydryl of Cys, which releases the ketone-cy fluorophore. The cleavage of p-nitrobenzoyl behaves intramolecular rearrangement to yield N-substituted product and followed by the intramolecular cyclization to complete the reaction (Tanaka et al., 2004; Yang et al., 2012; Lv et al., 2014). It is known that sulfydryl and amino group are both nucleophilic groups in aminothiols. In order to verify that the selectively detection of Cys is caused by sulfydryl, several control experiments are implemented from different aspects. As shown in Fig. S1, the probe Cy-NB could response to $100 \,\mu\text{M}$ β mercaptoethanol rapidly but could not response to 100 µM ethanolamine. The result confirms that the cleavage of ester is induced by sulfydryl but not amino group. And the sulfydryl of Cys can elevate the nucleophilic ability of amino group (about 10³ times) (Burchfield, 1958), which results in intramolecular rearrangement of compound 3 (Scheme S1). Furthermore, the absorption spectra of Cy-NB were examined in the presence of Cys, CySS, Hcy and GSH. An obvious absorption hypsochromic shift was observed with adding 100 µM Cys but the absorption spectra of Cy-NB showed no differences in the presence of equal CySS, Hcy and GSH (Fig. S2). In the case of CySS, the sulfydryl is hidden in disulfide bond, which cannot induce the reaction with Cy-NB. As for Hcy and GSH, which are also aminothiols, but the pK_a of Hcy (10.00) and GSH (9.20) are higher than that of Cys (8.53) (Benesch and Benesch, 1955). The nucleophilic ability of sulfydryl in Hcy and GSH is much weaker than that of Cys, which agrees with our previous research (Yin et al., 2015). In addition, the bulkiness of tripeptide of GSH would significantly hinder the reaction activity of sulfydryl. Therefore, the response of Cy-NB to Cys will not be interferenced by Hcy and GSH. These results further indicate that the sulfydryl is the key point that triggers Cy-NB response to Cys



Scheme 1. Synthesis route of Cy-NB and its reaction with Cys.

selectively. Furthermore, to investigate the function of nitro group in our probe, a control probe Cy-B was synthesized (Scheme S2). Both the probe Cy-NB and Cy-B were incubated with 50 μ M Cys at 37 °C for 5 min. The spectrum property of probe Cy-NB was changed obviously but that of Cy-B showed almost no changes (Fig. S3). We suggest that the nitro group which performed as a strong electron-withdrawing group can facilitate the ester cleavage and the cyanine dye can be uncaged easily from Cy-NB after Cys attacked. Therefore, the probe Cy-NB achieves rapidly response of Cys with the assistance of nitro group. And the fast response of Cy-NB to Cys is expected to detect mitochondrial Cys level changes and be able to assess oxidative stress in living cells.

3.3. Parameter optimization of the probe for Cys detection

We optimized the reaction conditions of probe Cy-NB for the detection of Cys. Buffer solution, pH and reaction time were taken into consideration. As buffer solution plays an important role in the performance of Cy-NB, HEPES, MOPS and Tris–HCl were investigated for their effect on the detection of Cys by Cy-NB. As shown in Fig. S4, the response of Cy-NB to Cys was the best in 5 mM HEPES. In addition, the stability of Cy-NB was good from pH 4.0 to pH 8.0 but glided sharply when pH exceeds 8.0 for the deprotonation of nitrogen atom. As the probe worked well under physiological conditions so pH 7.4 was chosen. And the complete reaction of Cys and Cy-NB could be almost obtained within 5 min. Furthermore, the fluorescence property of probe Cy-NB could remain stable over 60 min under the reaction condition (Fig. S5). Therefore, the detection of Cys by probe Cy-NB was reacted in 5 mM HEPES, pH 7.4 and incubated at 37°C for 5 min.

3.4. Spectral properties of CY-NB towards Cys

3.4.1. Absorbance spectral analysis

We examined the absorption spectral properties of Cy-NB in 5 mM HEPES solutions, pH 7.4. As shown in Fig. 1a, the UV-vis absorption of probe Cy-NB was centered at 780 nm ($\varepsilon = 1.2 \times 10^6 \,\text{M}^{-1} \,\text{cm}^{-1}$), which displayed a green color. However, a new absorption peak at 500 nm ($\varepsilon = 2.4 \times 10^5 \,\text{M}^{-1} \,\text{cm}^{-1}$) appeared and increased gradually after the probe Cy-NB incubated with different concentrations of Cys at 37 °C for 5 min. The color of the solution changed to red with the generation of the ketone-cy. The obvious absorption hypsochromic shift (about 280 nm) indicated that the polymethine π -electron system of Cy-NB was changed after triggered by Cys. These results show that the probe can be employed for Cys detection by naked eyes.

3.4.2. Fluorescence spectral analysis

We next investigated the fluorescence spectra of the probe Cy-NB. The emission peak of probe Cy-NB is 785 nm (λ_{ex} =720 nm). After Cy-NB incubated with Cys at 37 °C for 5 min, a new emission peak at 640 nm (λ_{ex} =560 nm) was observed. The large hypsochromic shift confirms that the π -electron system of Cy-NB is reassigned by the transformation from enol form to ketone form. Utilizing the principle, the NIR ratiometric detection of Cys can be accomplished with Cy-NB by the ratio of $F_{640 \text{ nm}}$ (λ_{ex} =560 nm) to $F_{785 \text{ nm}}$ (λ_{ex} =720 nm). Compared to those probes with single emission wavelength, the result achieved by Cy-NB is more sensitive and accurate because the signal of ratiometric probe is independent of probe concentration, photobleaching, and illumination intensity in the biological samples. These advantages inspire us to explore the property of Cy-NB for the detection of Cys by ratiometric signal. We incubated our probe Cy-NB with 0, 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 75 and 100 µM Cys in 5 mM HEPES at pH



Fig. 1. The spectra properties of Cy-NB (0.5 μ M) with increasing concentrations of Cys (0, 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 75 and 100 μ M) in 5 mM HEPES solution, pH 7.4 and incubated at 37 °C for 5 min. (a) Absorbance spectra (the inset showed the color changes of Cy-NB in the presence and absence of 100 μ M Cys). (b) Fluorescence spectra at 785 nm (λ_{ex} =720 nm). (c) Fluorescence spectra at 640 nm (λ_{ex} =580 nm). (d) The linearity. The linearity was seperated to two parts: the first part was from 0 μ M to 35 μ M and the linear equation was *y*=0.258*x*+0.843 with *r*=0.988; the second part was from 35 μ M to 100 μ M and the linear equation was *y*=1.08*x*-24.8 with *r*=0.995 (*y* means *F*_{640 nm}/*F*_{785 nm} and *x* means the concentration of Cys). The detection limit was calculated 0.2 μ M by signal-to-noise (*S*/*N*) ratio=3. Data were the means for three independent experiments and were presented as the mean \pm standard error. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

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 $F_{^{785\mathrm{nm}}}$

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7.4 for 5 min. As shown in Fig. 1b, the emission peak at 785 nm (λ_{ex} =720 nm) of probe Cy-NB was decreased with the increase of Cys concentration. Meantime, the emission peak at 640 nm (λ_{ex} =560 nm) was increased gradually (Fig. 1c). The results showed that the fluorescence ratio $(F_{640 \text{ nm}}/F_{785 \text{ nm}})$ increased from 1.9 to 84.9, with a 45-fold enhancement. In addition, good linearity of emission ratios ($F_{640 \text{ nm}}/F_{785 \text{ nm}}$) in the range of 0–35 μ M and 35– 100 µM was exhibited respectively (Fig. 1d). The detection limit was 0.2 μ M, which was calculated by *S*/*N* ratio=3. The high ratio signal of Cy-NB to Cys is benefited from the modulation of polymethine π -electron system which results a large hypsochromic shift of emission band. Then the analyses of Cvs in spiked cell lysis samples were carried out by our probe, high performance liquid chromatography (HPLC) and liquid chromatography tandem mass spectrometry (LC-MS) (Hoogerheide and Campbell, 1992; Huang et al., 2011). As shown in Table S1, the accuracy of quantitative analysis by the probe Cy-NB are lower than HPLC/LC-MS methods. But the results of our probe agreed well with those obtained by HPLC and LC–MS for the detection of Cys in cell lysis samples. Compared with the probes have already be established (Table S3), our probe Cy-NB provides rapid response time of Cys with satisfactory sensitivity. These advantages make Cy-NB an efficient candidate to detect endogenous mitochondrial Cys level changes.

3.5. Selectivity of CY-NB to Cys

3.5.1. Selectivity of CY-NB to Cys in water solution

To investigate the selectivity of Cy-NB towards Cys, the probe was incubated with various species such as Cys, Hcy, N-Acetyl-Lcysteine (NAC), 2-Methyl-L-cysteine (MC), Cys-Gly, GSH and other amino acids in 5 mM HEPES buffer at pH 7.4. As shown in Fig 2a, 100 µM Hcv. 10 mM GSH and other amino acids showed little interference in the detection of Cvs by Cv-NB. Even though the concentration of mitochondrial GSH is much higher than that of Cys, the probe Cy-NB owns the ability to distinguish mitochondrial Cys from GSH. The concentration of mitochondrial Hcy (\sim 50 nM) is much lower than that of Cys (\sim 500 μ M), which means that Hcy cannot make any interference in the detection of mitochondrial Cys. The ratiometric response ($F_{640 \text{ nm}}/F_{785 \text{ nm}}$) of probe Cy-NB was observed in presence of NAC. Although an acetyl group is attached to the amino group of NAC, the structure of NAC is quite similar to Cys. There exists free thiol in NAC. And NAC has been widely utilized as antioxidant in vivo (Sandstrom et al., 1994; Bilodeau et al., 2001). However, the reactivity of the thiol in NAC is reduced by the acetyl group. Therefore, the probe Cy-NB can respond to NAC, but the response of NAC by the probe Cy-NB is weaker than that of Cys. The phenomenon was further verified that the detection of Cys by Cy-NB was induced by sulfydryl but not amino group. Considering that the concentrations of Cys in cell lysis samples were much higher than those of NAC (Table S2), which were consistent with the results obtained by Wu et al. (2005), the NAC would not influence the detection of Cys at cell lysis level (Fig. S6). Therefore, the detection of Cys by our probe Cy-NB would not be influenced by NAC in living cells. All the results indicate that our probe Cy-NB exhibits excellent selectivity for Cys over other kinds of biothiols and amino acids. Furthermore, we examined the time-dependent emission ratio ($F_{640 \text{ nm}}/F_{785 \text{ nm}}$) changes of Cy-NB (0.5 μ M) in the presence of Cys (100 µM), Hcy (100 µM), GSH (10 mM), NAC (100 μ M), MC (100 μ M) and Cys-Gly (100 μ M), respectively (Fig. 2b). There are almost no signal obtained from Cy-NB in the presence of GSH, MC and Cys-Gly with almost no emission ratio change. The emission ratio of Cy-NB toward Hcy displayed a little variation from 1.86 to 6.11 during 10 min. By comparison, the response of the probe Cy-NB to Cys exhibited an obvious emission ratio $(F_{640 \text{ nm}}/F_{785 \text{ nm}})$ leap from 1.9 to 84.9 within 5 min. Even though the response of the probe Cy-NB to $100\,\mu\text{M}$ NAC also

 $F_{\rm ^{640\,m}}$ 4 (20 Cys Hey GSH NAC MCCys-GlyAla b Cys 80 - Hey GSH 640nm/ $F_{
m 785nm}$ NAC 60 MC Cys-Glv 40 20 Time (min) Fig. 2. (a) The ratiometric response ($F_{640 \text{ nm}}/F_{785 \text{ nm}}$) of Cy-NB (0.5 μ M) to Cys and

Fig. 2. (a) The fatiometric response ($F_{640 \text{ nm}}/F_{785 \text{ nm}}$) of Cy-No (0.5 µM) to Cys and other amino acids in 5 mM HEPES solution, pH 7.4 and incubated at 37 °C for 5 min. C means control (none amino acids exist), Cys (100 µM), Hcy (100 µM), GSH (10 mM), NAC (100 µM), MC (100 µM), Cys-Gly (100 µM) and other amino acids (10 mM). (b) Time-dependent ratiometric response ($F_{640 \text{ nm}}/F_{785 \text{ nm}}$) of Cy-NB (0.5 µM) to Cys (100 µM), Hcy (100 µM), GSH (10 mM), NAC (100 µM), MC (100 µM) and Cys-Gly (100 µM), respectively. Data were the means for three independent experiments and were presented as the mean \pm standard error.

exhibited an emission ratio ($F_{640 \text{ nm}}/F_{785 \text{ nm}}$) leap from 1.98 to 63.7 within 5 min, but NAC would not influence the detection of Cys under physiological condition. Next, we determined the reaction kinetics with excessive amounts of Cys, Hcy, GSH, NAC, MC and Cys-Gly towards Cy-NB in 5 mM HEPES at pH 7.4 and incubated at 37 °C. Herein, the reaction kinetics followed the pseudo-first-order kinetics and the rate constant *k* was calculated by Eq. (1):

$$\ln\left[\left(F_{max}-F_t\right)/F_{max}\right] = -kt\tag{1}$$

where F_{max} means fluorescence intensities at 640 nm at the maximum value and F_t means the fluorescence intensities at time t. The rate constant k for Cys, Hcy, GSH, NAC, MC and Cys-Gly is 9.3×10^{-2} min⁻¹, 1.2×10^{-3} min⁻¹, 2.8×10^{-4} min⁻¹, 6.9×10^{-2} min⁻¹, 2.6×10^{-4} min⁻¹ and 3.3×10^{-4} min⁻¹, respectively. The pseudo-first-order reaction constant k for Cys is almost 330-fold faster than that of GSH, 78-fold faster than Hcy, 1.3-fold faster than NAC, 360-fold faster than MC and 280-fold faster than Cys-Gly. All the results clearly indicate that the probe Cy-NB is an outstanding ratiometric fluorescent probe for the selective detection of Cys over GSH, Hcy, MC and Cys-Gly. The selectivity of our probe Cy-NB is not good for NAC, but considering that the concentration of NAC is much lower than Cys in living cells, it will not influence the detection of Cys.

3.5.2. Selectivity of CY-NB to Cys in living cells

We next studied whether the probe Cy-NB can respond to Cys without interference in living cells or not. The living HepG2 cells were chosen as bioassay model to evaluate the selectivity of Cy-NB toward Cys in living cells. The living HepG2 cells were incubated in



the presence of Cys (200 μ M), Hcy (200 μ M), GSH (10 mM) and other amino acids (10 mM) respectively at 37 °C for 30 min. The pre-treated cells were washed by RPMI-1640 three times and then incubated with 1 μ M Cy-NB at 37 °C for another 5 min. Confocal ratiometric fluorescence images of living HepG2 cells were obtained by the ratio of emission intensities from 580 nm to 680 nm (λ_{ex} =559 nm) and from 700 nm to 800 nm (λ_{ex} =635 nm). As shown in Fig. S6b, when the HepG2 cells pre-incubated with 200 μ M Cys, the ratio image was much brighter than control group (Fig. S7a). However, the ratio images showed almost no difference when HepG2 cells pre-incubated with 10 mM GSH, 200 μ M Hcy and 10 mM other amino acids (Fig. S7c–S7l). These results verified that the response of Cy-NB to Cys is also specific in living HepG2 cells.

3.6. Cytotoxicity of CY-NB to Cys

We also performed the MTT assays to estimate the cytotoxicity of Cy-NB. As shown in Fig. S8, the cellular viability was about 85% even in the presence of 10 μ M Cy-NB at 37 °C for 12 h which indicated the low cytotoxicity of our probe.

3.7. Mitochondrial sub-location of CY-NB

Mitochondria are known as power plant as well as a cellular compartment which proceeds various biosynthetic pathways. However, these important life-supporting mitochondria are also linked to death-promoting activity of cells (Newmeyer and Ferguson-Miller, 2003). As a kind of by-products of oxidative phosphorylation on mitochondria, ROS lead to mitochondrial oxidative stress and play a vital role in apoptosis regulation (Simon et al., 2000). As an important antioxidant defender, mitochondrial Cys level is sensitive to ROS. We next confirmed that our probe Cy-NB can be used to monitor mitochondrial Cys level to access mitochondrial oxidative stress. As mentioned above, the probe Cy-NB displayed excellent properties in the detection of Cys with great sensitivity and selectivity both in solution and living cells. We evaluated the ability of probe Cy-NB to sub-locate on mitochondria as the Cy-NB achieves the ability to target mitochondria for its ammonium cation (Wang et al., 2013). We utilized co-localization experiments which were usually employed for describing the difference of the location between two or more molecular dyes to validate Cy-NB located in mitochondria (Yu et al., 2013). As shown in Fig. 3, the living HepG2 cells were co-stained with 1 µM Cy-NB (5 min), 5 µM Rhodamine 123 (a mitochondria dye, 15 min) and 2 µM Hoechst 33258 (a nuclear dye, 30 min) (Crissman and Hirons, 1994). Fluorescence confocal microscopic image of probe Cy-NB, Rhodamine 123 and Hoechst 33258 was constructed from 580 nm to 680 nm (λ_{ex} =559 nm), from 520 nm to 550 nm (λ_{ex} =515 nm) and from 420 to 480 nm (λ_{ex} =405 nm), respectively. Fig. 3a1 is co-localization image of Cy-NB and Hoechst 33258 and Fig. 3a₂ is co-localization image of Rhodamine 123 and Hoechst 33258. Then we merged Fig. $3a_1$ and a_2 to form co-localization image of three dyes (Fig. 3a₃). Rhodamine 123 has already been confirmed with excellent quality of mitochondria localization in living cells (Johnson et al., 1980). As shown in Fig. 3a₃, the image of Cy-NB merged well with that of staining with the Rhodamine 123 dye in discrete subcellular locations. And we evaluated the Pearson's coefficient $R_r = 0.96$ and the Manders' coefficients $m_1 = 0.99$, $m_2=0.98$ by the Image-Pro Plus software. The intensity distributions of the two co-localization dyes were demonstrated through counting the intensity of color-pair feature pixel to reveal their correlation (Fig. $3b_1-b_3$). As shown in Fig. $3b_3$, the probe Cy-NB displaying a highly correlated plot with Rodamine 123. In addition, the intensity profiles of linear regions of the interest across HepG2 cells (white arrow in Fig. 3a₃) further proved Cy-NB and Rhodamine 123 depicted a good synchrony. All above results verify that the probe Cy-NB offers excellent ability to locate in mitochondria.

3.8. Assessment of mitochondrial oxidative stress status

We next examined the ability of Cy-NB to assess oxidative stress status in mitochondria. In the control group, the living HepG2 cells were incubated with 1 μ M Cy-NB at 37 °C for 5 min. Then HepG2 cells were washed with RPMI-1640 three times before image acquisition. Confocal ratiometric fluorescence images of living HepG2 cells were obtained by the ratio of emission intensities from 580 nm to 680 nm (λ_{ex} =559 nm) and from 700 nm to 800 nm (λ_{ex} =635 nm). As shown in Fig. 4a, a clear fluorescence ratio image was obtained which indicated that our probe was able



Fig. 3. The co-localization image of probe Cy-NB, Rhodamine 123 dye and Hoechst 33258 dye in living HepG2 cells. The HepG2 cells were stained with 1 μ M Cy-NB for 5 min, 5 μ M Rhodamine 123 for 15 min, and 2 μ M Hoechst 33258 for 30 min at 37 °C in RPMI-1640. (a₁) Fluorescence confocal microscopic images constructed from 580 nm to 680 nm (λ_{ex} =559 nm) for Cy-NB and from 420 to 480 nm (λ_{ex} =405 nm) for Hoechst 33258 dye. (b₁) Co-localization areas displayed of the red and blue channels selected in a₁. (a₂) Fluorescence confocal microscopic images constructed from 520 nm to 550 nm (λ_{ex} =515 nm) for Rhodamine 123 dye and from 420 nm to 480 nm (λ_{ex} =405 nm) for Hoechst 33258 dye. (b₁) Co-localization areas displayed of the red and blue channels selected in a₁. (a₂) Fluorescence confocal microscopic images constructed from 520 nm to 550 nm (λ_{ex} =515 nm) for Rhodamine 123 dye and from 420 nm to 480 nm (λ_{ex} =405 nm) for Hoechst 33258 dye. (b₂) Co-localization areas displayed of the green and blue channels selected in a₂. (a₃) Merged (a₁) and (a₂). (b₃) Co-localization areas displayed of the red and selected in a₂. (c₁) Analysis of intensity profile of chosen regions (white arrow in (a₃) across HepG2 cells). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



Fig. 4. Confocal ratiometric fluorescence images ($F_{580-68 \text{ nm}}/F_{700-800 \text{ nm}}$) of living HepG2 cells with Cy-NB. (a) The living HepG2 cells incubated with 1 μ M Cy-NB at 37 °C for 5 min. (b) The living HepG2 cells were pretreated with 100 μ M NEM for 30 min and then incubated with 1 μ M Cy-NB at 37 °C for 5 min. (c) The living HepG2 cells were pretreated with 100 μ M NEM for 30 min and then incubated with 1 μ M Cy-NB at 37 °C for 5 min. (c) The living HepG2 cells were pretreated with 10 nM PMA for 30 min and then incubated with 1 μ M Cy-NB at 37 °C for 5 min. (d) The living HepG2 cells were pretreated with 10 nM PMA for 30 min and then incubated with 1 μ M Cy-NB at 37 °C for 5 min. (d) The living HepG2 cells were pretreated with 10 nM PMA for 30 min and then incubated with 1 μ M Cy-NB at 37 °C for 5 min. The image (e), (f), (g) and (h) is the bright field image of (a), (b), (c) and (d), respectively. The right figure is the quantification of fluorescence ratio of each group. Data were the means for three independent experiments and were presented as the mean \pm standard error.

to detect endogenous mitochondria Cys. In the group two, the living HepG2 cells were pretreated with 100 µM N-ethyl-maleimide (NEM, a thiol blocking reagent) for 30 min and then incubated with 1 µM Cy-NB at 37 °C for 5 min. As shown in Fig. 4b, the fluorescence ratio image was almost black and the fluorescence ratio changed from 25.0 (Fig. 4a) to 1.5 (Fig. 4b) after mitochondrial Cys consumed by NEM. In the group three, the living HepG2 cells were pre-treated with 200 µM Cys for 30 min and followed incubated with 1 μ M Cy-NB at 37 °C for 5 min. As shown in Fig. 4c, the fluorescence ratio of living HepG2 cells changed to 80.0. In the group four, the living HepG2 cells were pre-treated with 10 nM phorbol 12-myristate 13-acetate (PMA) for 30 min to induce mitochondrial oxidative stress by respiratory burst (Emerit and Cerutti, 1982) and then incubated with 1 µM Cy-NB at 37 °C for 5 min. As shown in Fig. 4d, the fluorescence ratio of HepG2 cells pretreated by PMA was calculated to be 6.0, which was much lower than control group (Fig. 4a). As described above, the detection limit of probe Cy-NB is not as good as traditional methods like LC–MS. It meets big challenge for exactly quantifying the Cys concentration in living cells by the probe Cy-NB. However, the above results demonstrated that the mitochondrial Cys level decreased obviously after intracellular oxidative stress and our probe Cy-NB could monitor this alteration successfully. All these results confirm that Cy-NB is an excellent ratiometric fluorescent probe for mitochondrial Cys detection and can be utilized to assess mitochondrial oxidative stress successfully.

3.9. Visualization of Cys levels in living mice

We further tested the ability of Cy-NB to detect Cys in living animals, since the NIR emission of Cy-NB can penetrate deep tissue and avoid background autofluorescence. We chose BALB/c mice as the biological model and divided them into four groups. In the control group a, the peritoneal cavities of BALB/c mice were injected with 50 µL solution (DMSO/saline=1:9, v/v). The BALB/c mice in group b–d were intraperitoneally injected with 50 µL 1 mM Cys, 50 µL 1 mg/mL lipopolysaccharide and 100 µL 10 mg/mL diethylmaleate (DMSO/saline=1:9, v/v), respectively. Thirty minutes later, the probe Cy-NB (1 µM, 50 µL in 1:9 DMSO/ saline v/v) was injected into peritoneal cavities of the four group BALB/c mice for another 30 min. Fluorescence images were constructed from fluorescence collection channel 1 (600 nm to 700 nm, λ_{ex} =530 nm) and channel 2 (750 nm to 850 nm, λ_{ex} =735 nm) using in vivo imaging system (Bruker). Additionally, we merged the fluorescence image with the corresponding X-ray image to clearly display the reaction site of the mice. In the control group a, a recognizable signal was collected in channel 2 (Fig. $5a_2$), which indicated that our probe Cy-NB had reacted with endogenous Cys in peritoneal cavity of BALB/c mice. In the group b, the signal intensity in channel 1 was decreased obviously (mean signal intensity changed from 163.7 to 37.2) after exogenous Cys added into the peritoneal cavity (Fig. $5b_1$). Meanwhile, the mean signal intensity collected in channel 2 was increased from 295.7 to 1435.4, which confirmed that Cy-NB could detect Cys level changes in peritoneal cavity of BALB/c mice (Fig. 5b₂). As shown in Fig. 5c₂, the signal of Cy-NB collected in channel 2 was much weaker than that of group a. The result indicated that the concentration of Cys in the peritoneal cavity of mice was decreased under oxidative stress condition, which induced by lipopolysaccharide (Kheir-Eldin et al., 2001) and the result agreed with previous research (Iyer et al., 2009). In the group d, the signal of Cy-NB collected in channel 2 was also weaker than that of group a, which indicated that the Cys in mice peritoneal cavities were depleted after diethylmaleate treated and the results agreed with Gerard-Monnier et al. (1992). All these results verify that our probe Cy-NB is able to image endogenous Cys level in living animals.

4. Conclusions

In summary, a near-infrared ratiometric fluorescent probe Cy-NB for the detection of mitochondrial Cys has been developed. The probe Cy-NB is able to selectively respond Cys in aqueous solution, in living cells and in vivo. The triggered rearrangement of the conjugated π -electron system in fluorophore will produce a ratiometric fluorescent signal that allows avoiding environment interference. The mitochondria-targeted probe Cy-NB exhibits excellent properties including fast reaction kinetics, good selectivity, high sensitivity, and good photostability. Our probe reveals that the endogenous mitochondrial Cys pool will vary under different oxidative stress status. We also achieve imaging Cys in mice successfully. The promising ratiometric NIR fluorescent probe can be of great potential for further utilizing in investigating the physiological function of Cys in biological systems.



Fig. 5. Fluorescence images of Cys with the probe Cy-NB in the peritoneal cavities of BALB/c mice. In control group a, BALB/c mice were pretreated with 50 μ L solution (DMSO/saline = 1:9, v/v) injecting in the peritoneal cavities. Thirty minutes later, the probe Cy-NB (1 μ M, 50 μ L in 1:9 DMSO/saline v/v) was injected into peritoneal cavity of BALB/c mice for another 30 min. (a₁) the image constructed from fluorescence collection channel 1: 600 nm to 700 nm (λ_{ex} =530 nm). (a₂) The image constructed from fluorescence collection channel 1: 600 nm to 700 nm (λ_{ex} =530 nm). (a₂) The image constructed from fluorescence collection channel 1: 600 nm to 700 nm (λ_{ex} =530 nm). (a₂) The image constructed from fluorescence collection channel 2: 750 nm to 850 nm (λ_{ex} =735 nm). In the group b–d, BALB/c mice were pretreated with 50 μ L 1 mM Cys, 50 μ L 1 mg/mL lipopolysaccharide and 100 μ L 10 mg/mL diethylmaleate (in 1:9 DMSO/saline, v/v) injecting in the peritoneal cavities. Thirty minutes later, Cy-NB (1 μ M, 50 μ L in 1:9 DMSO/saline v/v) was injected in peritoneal cavities of BALB/c mice for another 30 min. (b₁–d₁) The image constructed from fluorescence collection channel 1. (b₂–d₂) The image constructed from fluorescence collection channel 2. (e₁) The quantification of total photon flux from channel 2. Data were the means for three independent experiments and were presented as the mean \pm standard error.

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Appendix A. Supplementary Information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.bios.2015.06.039.

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