Highly specific C–C bond cleavage induced FRET fluorescence for in vivo biological nitric oxide imaging†

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A novel Förster resonance energy transfer (FRET) fluorescence "off–on" system based on the highly specific, sensitive and effective C–C bond cleavage of certain dihydropyridine derivatives was reported for real-time quantitative imaging of nitric oxide (NO). 1,4-Dihydropyridine was synthesized as a novel linker which could connect customized fluorophores and their corresponding quenchers. The specific and quantitative response to NO is confirmed using fluorescence spectrometry with the classical example of fluorescein isothiocyanate (FITC) and [4′-(N,N′-dimethylamino)phenylazo] benzoyl (DABCYL). The fluorescence intensity increased linearly with the increase in the amount of NO. Cells incubated with an exogenous NO donor emitted fluorescence as expected. A high fluorescence intensity was detected in macrophages which generate NO when incubated with lipopolysaccharide (LPS). The in vivo imaging shows about an 8-fold contrast between Freund’s adjuvant stimulated feet and normal feet in mice after intravenous injection, which was the first example of in vivo semiquantitative fluorescence imaging of NO in mammals.

Introduction

Nitric oxide (NO), a very active radical signaling molecule, has been found to exist and play a significant role in living body pathophysiology.1 It has been gradually realized that NO is involved in different physiological processes including vasodilation, wound healing, immune responses and nerve cell communication.2 Therefore, NO detection in vitro or, more ideally, 3D imaging the distribution and concentration of NO in vivo would be of a great help in understanding the metabolism of NO.

Various methods, such as electrochemistry,3 optical imaging (fluorescence, chemiluminescence and bioluminescence imaging),4 electron paramagnetic resonance (EPR),5 and magnetic resonance (MR), have been developed to image NO.6 Among these methods, optical probe-based techniques are the most available for NO imaging. With intrinsic molecular sensitivity, high resolution, repeatability, a high level of safety, and a relatively low instrumentation cost, optical probes have become important tools in the study of cells and small animals. Both fluorescence and chemiluminescence imaging have been widely used for elucidating the function of NO in different biological systems.7

Among the existing ratiometric probes, most of them are designed according to a photo-induced electron transfer (PET) mechanism, during which donor–π–acceptor structures are formed after the reaction between NO and the probe. Numerous efforts have been made on o-diamino aromatic compounds which could be oxidized by NO into triazole derivatives in the presence of oxygen.8 Although fluorescein based probes have been effective in zebrafish,9 there are no results regarding mammalian models yet. The speculated reason is the deficiency of oxygen at site for NO detection. Furthermore, o-diamino aromatics are easily disturbed by some oxidative species secreted in cells, such as H2O2 and OONO−.8 A series of copper complexes were tested in mouse models, which could obviously distinguish a normal liver from a liver with acute severe hepatic injury (ASHI).10 Yet, the fluorescence images were obtained from excised liver slices harvested from a living animal using an ex vivo method. There are other limitations of PET probes:11 (1) the measured excitation and emission wavelengths are often different from the expected values; (2) relatively broad fluorescence spectra are often detected for PET probes; (3) although remarkable advances have been made in computational chemistry, the quantum yield of fluorophores cannot be accurately predicted.
On the other hand, FRET mechanism based fluorescent probes have been used in the fields of chemistry and biology for decades. The commercially available FRET fluorophores have been proven to have certain excitation and emission wavelengths and an exact quantum yield. With different fluorophores, the available detection range could be extended to the near-infrared region, which makes FRET probes more promising for in vivo imaging than PET probes. A typical FRET quencher–fluorophore pair is FITC and DABCYL. This pair can directly indicate if the bonds between the two moieties are broken. A probe based on coumarin decorated 1,4-dihydropyridine was synthesized and evaluated in vitro, which was designed based on stoichiometry. The probe transforms into a conjugated fluorophore after NO stimulation, but the in vivo application was largely restricted by the short emission wavelength. Therefore, we designed and synthesized a FRET based probe to conquer the previous obstacles.

Herein, we present a fluorophore–quencher FRET “off–on” system for detecting and imaging NO in vivo in a non-invasive and real-time way. The 1,4-dihydropyridine derivatives can react with NO specifically. Furthermore, as the C4-position is substituted with a benzyl group, the C–C bond between 1,4-dihydropyridine and the benzyl group can be cleaved through NO stimulation with a high yield (Scheme 1). The reaction is very specific and sensitive.

Results and discussion

In our probes, FITC and DABCYL were linked via 1,4-dihydropyridine within a distance of 10 nm. Because of the FRET mechanism, DABCYL quenches the fluorescence emitted by FITC in the absence of NO. The cleavage of the C–C bond leads to the release of DABCYL and the fluorescence can be detected again (Fig. S1†). To achieve an image in vivo, the emission wavelength is tuned to increase the signal–noise ratio through using different fluorophores (Scheme 2).

To obtain the NO target probe, an asymmetric synthesis strategy was used in the Hantzsch reaction. An equivalent amount of aldehyde, β-keto ester and methyl 3-aminobut-2-enoate ester were combined into an asymmetrical 1,4-dihydropyridine (compound 4) which had only one azido moiety. Compound 6 was designed as a linker (Scheme 2). The amino group on compound 6 is the binding site for the fluorophore, because many commercial fluorophores are designed for labeling the amino groups on peptides and proteins. The azido group would react with an yne group which was linked to a quencher. DABCYL-yne and FITC were selected to react with the linker to obtain the target probe. The reaction between DHPFQ (dihydropyridine–fluorophore–quencher) and NO was evaluated in phosphate buffer using a fluorospectrometer, and the reaction proceeded in a quantitative way. The C–C bond cleavage was confirmed from the increase in the intensity of the fluorescence.

As expected, an enhancement of the fluorescence intensity at certain excitation (490 nm) and emission (525 nm) wavelengths was detected in 2 min after DHPFQ reacted with NO (Fig. 1a). The fluorescence intensity increases with an increasing amount of NO (diluted NO saturated solution). The most small molecule inflammatory factors in the complicated internal environment are reactive oxygen/nitrogen species (ROS/RNS). Therefore, specific recognition and reaction between the probes and NO is essential for in vivo applications. To evaluate the reaction selectivity, compound S (Scheme S2†) as a DHPFQ analog was treated with different reactive species firstly (Fig. S1†). About 98% of compound S was found to be decomposed when treated with NO, which was measured using HPLC.

Furthermore, DHPFQ was treated with NO and many other reactive species (Fig. 1b). The decomposition was measured using a fluorescence spectrometer. DHPFQ showed a significant increase in fluorescence intensity with NO. No significant decomposition was found when DHPFQ and compound S were treated with other reactive species. The results reveal the exclusive response of 1,4-dihydropyridine derivatives to no ROS or RNS but NO. The UV spectra of DHPFQ before and after NO treatment were also investigated.
which showed that activated and inactivated DHPFQ share similar profiles (Fig. S3†).

With excellent responsiveness to NO and exceptional selectivity in vitro confirmed, the ability of DHPFQ to visualize NO in live cells was further tested. Sodium nitroprussiate (SNP) as an exogenous NO donor was used to validate the high contrast. Compared to the control group (Fig. S7a†), the SNP treated group (Fig. S7b†) showed strong fluorescence, which implies that DHPFQ could respond to exogenous NO in a physiological environment.

Since NO is an inflammatory factor generated by macrophages when stimulated with an antigen, in order to test the feasibility of application in vitro, raw 264.7 cells were stimulated with lipopolysaccharide (LPS) to generate endogenous NO.19 A significant enhancement of the fluorescence intensity was observed when the macrophages were incubated with DHPFQ and LPS (Fig. 2, middle). There was a high correspondence between the green fluorescence and the Lytracker Red signal which suggests this probe is metabolized in lysosomes. In contrast, when the macrophages were pretreated with N\(^{G}\)-monomethyl-L-arginine (L-NMA), an inhibitor of nitric oxide synthase (NOS),18 only a weak fluorescence signal above the background was detected (Fig. 2, bottom). This result demonstrated that the fluorescence is indeed from the switch-on reaction between DHPFQ and NO.

DHPFQ has the ability to detect endogenous NO in vitro, suppress interference from other inflammatory factors, and has the potential to detect NO in vivo.

NO, as a significant inflammatory factor, is concentrated in inflamed areas.19 An inflamed mouse model was then examined to test whether DHPFQ is effective in vivo. Freund’s adjuvant was subcutaneously injected into the left rear paws of mice to trigger inflammation.20 After two weeks, another portion of Freund’s adjuvant was subcutaneously injected. At 3 days after the second injection, DHPFQ was intravenously injected (injected dose: 0.5 mg kg\(^{-1}\), 10% ethanol, 90% physiological saline, 100 µL) and the mice were imaged after 10 min to 60 min using an in vivo imaging system (Fig. 3). To reduce the background absorption, depilatory paste was used to partially remove hair from the mice. 470 nm and 600 nm were chosen as
demonstrate the feasibility of the structure. In this study, FITC and DABCYL were used to
inflammation region was about 8-times higher in the in
penetration property of NIR probes is tunable
the immunohistology staining (Fig. 4b) also indicated that
penetration was about 8-times higher in the in
imaging and low specificity. The fluorescence intensity was observed in the
penetration was about 8-times higher in the in
ery injection, DHPFQ was switched on
then 10 min post-injection. After 60
absorptivity of optical imaging, and low specificity can be overcome with the application of the 1,4-dihydropyridine
concentration of NO in physiological
The comparison of existing imaging methods and the probe
demonstrates the feasibility of the fluorescence imaging of NO in
Fluorescence imaging of NO
Conclusions
A novel C-C bond cleavage induced FRET fluorescent probe DHPFQ based on 1,4-dihydro-pyridine was synthesized. In vitro
assessments have shown a ratiometric and specific result in responding to NO over other RNS/ROS. The fluorescence intensity increased linearly with the increasing amount of NO.
Furthermore, semiquantitative results were obtained through ROI analysis. The penetration property of NIR probes is tunable simply through the substitution of the fluorophores and quenchers, which will make the probe more practical for clinical applications.

Table 1  Comparison of molecular imaging methods and probes for NO detection

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Notes and references


