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A Minimalist Fluorescent Probe for Differentiating Cys, Hcy and GSH in Live Cells

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A simple molecule, tetrafluoroterephthalonitrile (4F-2CN), was discovered to be an efficient fluorescent probe for detecting biological thiol species. The probe responds to Cys and emits strong green fluorescence, whereas it reacts with Hcy/GSH and generates blue fluorescence. Addition of CTAB (cetyl trimethylammonium bromide) was observed to alter the fluorescent color of the reaction product of 4F-2CN and Hcy (from blue to green), but no alteration of fluorescent color occurred for Cys and GSH. For the very first time, cell imaging experiments showed that the three commonly occurring thiols (Cys/Hcy/GSH) could be differentiated using single fluorescent probe. In addition, the reaction product of 4F-2CN and Cys exhibits two-photon property, offering potentially useful tools for tissue imaging studies. To our best knowledge, 4F-2CN is currently the smallest fluorescent probe for thiol detection. We envision this new and versatile probe will add useful tools for further elucidating the roles of thiols in biology.

Introduction

Small molecule thiols play crucial roles in maintaining cellular redox environment and mitigating damages from free radicals and toxins. Three low-molecular-weight thiols are commonly found in biological systems, namely cysteine (Cys), homocysteine (Hcy) and glutathione (GSH). These thiols are closely involved in regulating various physiological and pathological processes. For example, Cys is an essential amino acid for protein synthesis. Abnormal level of Cys is related to slowed growth, edema, lethargy, liver damage, etc. Hcy elevation in plasma, on the other hand, is a risk factor for cardiovascular disease, Alzheimer’s disease and osteoporosis, while GSH is closely linked to leucocyte loss, cancer, HIV infection, etc. The important biological roles of thiols has thus spurred strong interest in developing useful chemical tools for detecting thiols. Fluorescent probes, owing to its simplicity and non-invasiveness, have become a popular approach for thiol detection in living cells. Over the last decade, a large number of fluorescent probes have been developed for detecting thiols. Most of these probes are based on thiol-selective chemical reactions, including Michael addition reactions, nucleophilic substitution, cyclization reactions between aldehyde and aminothiols, cleavage reactions of 2,4-dinitrobenzenesulfonyl (DNBS) with thiols, disulfide exchange reaction, and others. Similarity among the structures and the reaction activities of Cys, Hcy and GSH has posed substantial difficulty in discriminating one thiol species from another. Despite this challenge, several fluorescent probes that allow for selective detection of Cys or GSH have been reported, e.g., Strongin’s seminal work on selective detection of Cys/Hcy over GSH using a cyclization reaction between Cys/Hcy and acrylate. Yang and coworkers also designed a GSH-selective probe by employing specific thiol-halogen reaction between chlorinated BODIPY and GSH. Further effort has been devoted recently to develop single probe systems for simultaneous detection of two or three thiol species. For example, Guo and coworkers has developed a chlorinated coumarin-hemicyanine probe for simultaneous detection of Cys and GSH. Nevertheless, single probes that are able to fully discriminate the three thiols from each other are still quite rare. In this study, we report a remarkably simple but versatile probe 4F-2CN (Fig. 1), which is capable of simultaneously detecting Cys and Hcy/GSH using dual emission channels. Moreover, the fluorescent color of Hcy and the probe can be altered by adding a surfactant called CTAB, thereby allowing all three thiols to be differentiated from each other.

Results and Discussion

In a screening experiment, we identified a commercial compound, 2,3,5,6-tetrafluoroterephthalonitrile (4F-2CN), which was shown to react with Cys and produce bright green fluorescence in PB buffer (10 mM phosphate buffer, pH = 7.4) (Fig. 1A). It was noted that when the CN groups were altered to other electron withdrawing groups like CHO/COOMe, no obvious fluorescence was observed with these analogues. TLC experiments also showed that no reaction has occurred between

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the analogues and Cys under PBS/DMF = 1:1 after 2h (Fig S1†). The possible reason for different reactivity of these compounds is likely due to the difference in the electron-withdrawing ability of CN, CHO and COO Me. With non-thiol amino acids (e.g., histidine and lysine), no reaction with 4F-2CN occurred as indicated by fluorescence assays and mass spectrometry analysis.

We characterized the reaction product of 4F-2CN and Cys. 4F-2CN was mixed with Cys in a DMF solution to give the product 4F-2CN-Cys which was characterized by 

\[ ^{1}H, ^{13}C \text{ and } ^{19}F \text{ NMR, ESI-MS, FT-IR and X-ray single crystallography (CCDC 1402255; Fig. S2-S8†, Tables S1-S3†). We hypothesize that the formation of 4F-2CN-Cys was initiated by the Cys thiol group replacing a fluoro group on 4F-2CN in an aromatic nucleophilic substitution, and the subsequent cyclization was facilitated by the six-membered ring configuration afforded by the Cys substrate (Fig. 1B). The strong fluorescence from the cyclized product can be ascribed largely to the electron donation from the amino and sulfide moieties onto the aromatic core. We also determined the quantum yield of 4F-2CN-Cys to be 0.35 (Table S4†), indicating that it is a good fluorophore.

**Proposed reaction mechanism of 4F-2CN and Cys.**

**Fig. 1** (A) Fluorescence response of 4F-2CN and its analogues after incubation with Cys in PB buffer (Ex/Em at 420 nm/500 nm). (B) Proposed reaction mechanism of 4F-2CN and Cys.

Next we carried out time-dependent absorbance experiments with the probe. 4F-2CN alone in PB buffer did not show any absorption in the range of 340-500 nm (Fig. S9†). Upon addition of Cys, a new absorption peak at 420 nm was immediately observed, and the absorption signal reached equilibrium after around 2h (Fig. 2A and 2D). For Hcy and GSH, an absorption peak at 350 nm could be observed, and it also plateaued after around 2h (Fig. 2B, 2C, 2E and 2F).

**Fig. 2** (A-C) Time-dependent absorption spectra of 4F-2CN (50 μM) incubated with 1 mM of Cys (A), Hcy (B) and GSH (C) in PB buffer (10 mM, pH 7.4). (D-F) Corresponding absorbance changes of 4F-2CN incubated with Cys (D), Hcy (E) and GSH (F).

**Proposed reaction mechanism of 4F-2CN and Cys.**

**Fig. 1** (A) Fluorescence response of 4F-2CN and its analogues after incubation with Cys in PB buffer (Ex/Em at 420 nm/500 nm). (B) Proposed reaction mechanism of 4F-2CN and Cys.

![Diagram](image1.png)

The fluorescence response of 4F-2CN with Cys/Hcy/GSH was then investigated using an excitation wavelength of 350 nm. Similar to the previous study, the probe itself displayed negligible fluorescence in this channel. Upon addition of GSH or Hcy, a new fluorescence emission peak at 450 nm appeared (Fig. 3B and S10†). On the other hand, Cys did not show obvious fluorescence.

![Diagram](image2.png)
increment at 450 nm (Fig. 3D). The fluorescence increments at 450 nm for Hcy, GSH and Cys are 29-, 26- and 2.7-fold respectively. The results demonstrated that the probe was able to differentiate GSH/Hcy from Cys when excited at 350 nm. Concentration-dependent experiment with the probe indicated that the fluorescence signal from Cys was more significant at 450 nm increased with increasing concentrations of Hcy or GSH (Fig. S13 and S14†). The detection limits of Hcy and GSH were determined to be 2.27 and 0.24 μM respectively. It is noticed that the LOD of Hcy/GSH is about ten times higher than that of Cys. The higher sensitivity observed for Cys arises from the strong fluorescence of the reaction product of Cys and 4F-2CN. The enhanced fluorescence in the case of Cys can be attributed to the electron-donating amino group, as well as the rigid cyclized structure in the product. In addition, the reaction with Cys proceeds faster than that with Hcy/GSH (Fig. 3C and 3D), contributing to the higher sensitivity for Cys observed in the given length of time.

Previous studies have shown that the surfactant CTAB can help to form micelles and facilitate the intramolecular ring formation. In our study, it was found that the fluorescent property of the reaction between 4F-2CN and Hcy could be altered by adding CTAB. As shown in Fig. 4A, a weak fluorescence signal was observed at 500 nm when 4F-2CN was incubated with Hcy for 1h. After CTAB was added, a gradual fluorescence increase could be readily observed at 500 nm. On the other hand, the reactivity of 4F-2CN with Cys and GSH remains almost unchanged after the addition of CTAB. From these data, we can draw the conclusion that 4F-2CN displayed distinct reactivity patterns towards Cys, Hcy and GSH by using CTAB. The probe can be potentially used to differentiate the three highly similar thiol species. We further characterized the reaction products of 4F-2CN with Hcy and GSH using HPLC, ESI-MS and NMR. Experimental data revealed that disubstituted products were produced in the reaction of 4F-2CN and Hcy/GSH (Fig. 4B and S15-S24†). Addition of CTAB will facilitate the cyclization of 4F-2CN and Hcy, but not the cyclization of 4F-2CN and GSH (Fig. S18†).

Selectivity experiment is instrumental for biological applications of the probe, such as cell imaging experiment. The selectivity experiments were performed by incubating the probe with various biological analytes, including 20 natural amino acids, Hcy, GSH, H$_2$S, RNS, ROS and metal ions. The fluorescence intensity was then measured using two different emission wavelengths. As shown in Fig. 5A, the fluorescence intensity at 450 nm showed that only GSH/Hcy induced significant fluorescence change whereas other biological analytes gave small increment. At the fluorescence intensity of 500 nm, Cys displayed a substantial fluorescence increment; Hcy showed small increment and other samples induced very little increment (Fig. 5B). These results indicated the probe is highly selective toward thiols.

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**Fig. 4** (A) Fluorescence intensity changes of 4F-2CN reacting with Cys, Hcy and GSH with/without addition of CTAB buffer (Ex/Em at 420 nm/500 nm). The red lines represent the reaction without adding CTAB. The black lines denote the event that CTAB was added to the reaction after 1h of incubation. (B) Schemes of 4F-2CN reacting with Cys, Hcy and GSH respectively.

**Fig. 5** (A-B) Selectivity studies of 4F-2CN with various biological analytes at two different emission wavelengths. (A) λ$_{ex}$ = 350 nm, λ$_{em}$ = 450 nm; (B) λ$_{ex}$ = 420 nm, λ$_{em}$ = 500 nm. 1, Control; 2, Cys; 3, Hcy; 4, GSH; 5, K'; 6, Na'; 7, Mg$^{2+}$; 8, Zn$^{2+}$; 9, NO$_3$; 10, Ni$^{2+}$; 11, S$_2$O$_3$$^-$$^2$; 12, S$^-$$^2$; 13, •OH; 14, •OrBu; 15, H$_2$O$_2$; 16, TBHP; 17, Ala; 18, Ile; 19, Leu; 20, Val; 21, Phe; 22, Try; 23, Tyr; 24, Asn; 25, Gin; 26, Met; 27, Ser; 28, Thr; 29, Asp; 30, Glu; 31, Arg; 32, His; 33, Lys; 34, Gly; 35, Pro.

**Fig. 6** Fluorescence microscopy experiments of imaging Cys/Hcy/GSH with 4F-2CN using dual emission channels (green channel: emission was collected at 500-530 nm, blue channel: emission was collected at 440-460 nm. Both channels were excited at 405 nm). HeLa cells were first incubated with CTAB for 5 min, then NEM (2 mM) for 20 min. Subsequently Cys/Hcy/GSH (5 mM) was added and incubated for 15 min. 4F-2CN was then added and incubated for another 20 min. (A) Cells with NEM and GSH (blue channel), (B) Cells with NEM and Hcy (blue channel), (C) Cells with NEM and Cys (blue channel).
Encouraged by the above results, we moved forward to study the capability of 4F-2CN to image Cys/Hcy/GSH in living cells. We first treated HeLa cells with CTAB. N-ethylmaleimide (NEM), a common thiol depletion reagent was then added. Cys, Hcy and GSH were subsequently added to the medium respectively. 4F-2CN was finally added to the medium and incubated for 20 min. Confocal imaging results showed that cells with the addition of Cys displayed green fluorescence (Fig. 6C and 6F). GSH treated cells, on the other hand, gave off bright blue fluorescence (Fig. 6A and 6D). Cells treated with Hcy showed both blue and green fluorescence (Fig. 6B and 6E). The green fluorescence is attributed to the cyclized product of the reaction between 4F-CN and Hcy. These results together unambiguously proved that 4F-2CN can be used to differentiate Cys, Hcy and GSH via dual emission channels. We also performed MTT assay to examine the cytotoxicity of 4F-2CN. As shown in Fig S26, at 10 µM (this is the concentration we used for the cell imaging studies in this work), the probe showed relatively low toxicity. Higher toxicity was observed with higher concentrations. Thus it is recommended to keep the probe’s concentration equal or below 10 µM.

In addition, we found that 4F-2CN-Cys possesses two-photon properties when excited at 860 nm (Fig. S27), whereas no two-photon signal from the reaction product of 4F-2CN and Hcy/GSH was observed. Further cell imaging experiment showed that living cells displayed fluorescence under two-photon excitation after 4F-2CN was added. (Fig. 7). This study indicated that 4F-2CN might serve as a useful two-photon probe for selective detection of Cys in tissue imaging studies. The reason that 4F-2CN-Cys has two photon property could be attributed to the following two factors. First, 4F-2CN-Cys has strong electron withdrawing (CN) and electron donating (NH) groups. In general, adding strong electron donors and acceptors to the conjugated π system can enhance two photon signals. For 4F-2CN-Hcy and 4F-2CN-GSH, the electron donating ability of the thiol group is less effective compared with that of amine moiety. Second, it was observed that rigid conformation can also enhance two-photon properties. 4F-2CN-Cys forms a cyclized product which increased the rigidity of the product’s conformation whereas 4F-2CN-Hcy and 4F-2CN-GSH do not.

Finally, we examined the fluorescence response of 4F-2CN with three sets of thiol mixtures, Cys/Hcy, Cys/GSH and Hcy/GSH. As shown in Fig. S28, detection of Cys is highly sensitive. It can be selectively detected even in the presence of 10 eq of Hcy or GSH ($\lambda_{ex} = 420$ nm). On the other hand, in the mixture Cys/Hcy and Cys/GSH sets, Hcy (as well as GSH) can also be detected without much interference from Cys in the mixture ($\lambda_{ex} = 350$ nm). As for the set of Hcy/GSH mixture, the two thiols cannot be differentiated when excited at 350 nm. However, selective detection of Hcy can be achieved by addition of CTAB. And the signal of Hcy was not interfered in the presence of GSH.

**Conclusion**

In this study, we have discovered a novel and remarkably simple probe 4F-2CN, which could undergo selective reactions with thiols under physiological condition. Importantly, 4F-2CN displayed distinct reaction profiles against Cys, Hcy and GSH with the use of CTAB. Our bioimaging experiments proved, for the very first time, that Cys, Hcy and GSH can be differentiated using single fluorescent probe. It should be noted that the probe can be subjected to further chemical modification and produce derivatives with different photophysical properties. The research work for synthesizing different derivatives of 4F-2CN is currently in progress. Interestingly 4F-2CN-Cys was found to possess two-photon property, whereas the reaction product of 4F-2CN and Hcy/GSH does not possess any two-photon property. Cell imaging experiment showed that two-photon fluorescence can be observed with the addition of 4F-2CN. The results indicated that the probe could serve as useful tools for selective detection of Cys in tissue imaging studies. Given the small size and excellent property of 4F-2CN, we envision this new and versatile probe will add useful tools for further elucidating the roles of thiols in biology.

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**Notes and references**

† Electronic Supplementary Information (ESI) available: synthesis, spectroscopic properties, NMR, mass spectra and confocal imaging. See DOI: 10.1039/b000000x/


