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Suraj U. Hettiarachchi  
*Louisiana State University*

Bijeta Prasai  
*Louisiana State University*

Robin L. McCarley  
*Louisiana State University*

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Detection and Cellular Imaging of Human Cancer Enzyme Using a Turn-On, Wavelength-Shiftable, Self-Immolative Profluorophore

Suraj U. Hettiarachchi, Bijeta Prasai, and Robin L. McCarley*

Department of Chemistry, Louisiana State University, Baton Rouge, Louisiana 70803-1804, United States

Supporting Information

ABSTRACT: A frontier area in the development of activatable (turn-on) fluorescence-based probes is that concerned with rapid and selective stimulus triggering of probe activation so as to allow for biomarker identification and cellular imaging. The work here is concerned with a cloaked fluorophore composed of a reporter whose fluorescence is efficiently quenched by it being bound to an activatable trigger group through a novel self-immolative linker. Highly selective and rapid activation of the trigger group is achieved by chemical and enzymatic means that result in activated trigger group detachment from the self-immolative linker, with the latter subsequently cleaved from the reporter autonomously, thereby unmasking intense, red-shifted fluorescence emission. To achieve this success, we used a trimethyl-locked quinone propionic acid trigger group and an N-methyl-p-aminobenzyl alcohol self-immolative linker attached to the reporter. Delineated here are the synthesis and characterization of this cloaked fluorophore and the evaluation of its triggered turning on in the presence of an up-regulated enzyme in human cancer cells, NAD(P)-H:quinone oxidoreductase-1 (NQO1, DT-diaphorase, EC 1.6.99.2).

The success of the ever-growing field of in vivo and ex vivo fluorescence imaging of diseased cells for diagnosis, pathology, and treatment applications rests upon the existence of disease-specific molecules whose spectroscopic signal is readily differentiated from that of the background of surrounding normal cells. Molecular probes whose cloaked fluorescence reporter signal is turned on by endogenous enzymes offer distinct advantages over always-on reporters, in particular, higher signal-to-background (positive-to-negative) values. Critical for cancer imaging is the development of libraries of turn-on probes that will allow for the collection of real-time information on the diseased tissue microenvironment and the high-fidelity definition of diseased and healthy tissue borders during fluorescence-assisted surgical resections.

Unfortunately, the number of pro-fluorogenic probes that can have their fluorescence output selectively and quickly revealed by the presence of cancer-associated enzyme is extremely small, as is the type of disease-associated activating enzyme. Also, routes to improving the positive-to-negative value are limited. There are three extant probes that passively target cancer cells and rapidly (<30 min) identify the diseased cells by reporter fluorescence that is initiated by intracellular enzyme action; only three distinct cancer-linked enzymes have been targeted to date. Enzyme-initiated dequenching of probe fluorescence has been used to improve the signal-to-background value of the resulting reporter in diseased (positive) cells in comparison to that of unactivated probe or unaffected (negative) cells. Probe fluorescence is quenched via fluorescence resonance energy transfer (FRET) or photo-induced electron-transfer (PeT) mechanisms, and reporter fluorescence comes about by enzyme-initiated disassociation of the FRET quencher/donor pair upon enzyme binding or cleavage of the quencher (trigger group) from the probe. However, modest signal-to-background (SBR)/positive-to-negative (PNR) or tumor-to-background (TBR) ratios have been reported.

Ideal probe characteristics include rapid cell uptake and highly selective activation to yield the reporter, low fluorescence efficiency (Φreport) of the unactivated probe but high fluorescence efficiency of the reporter (Φprobe), and high-quality retention of the reporter inside the diseased cells. Furthermore, the probe–reporter pair will exhibit large differences in energies of absorption maxima (λabs) and emission maxima (λem) so significantly enhanced PNR/TBR values arise from the low absorbivity of the probe at the reporter’s maximum excitation wavelength. A highly promising approach that incorporates many of the above desired probe/reporter qualities is one wherein the probe is tripartite, being composed of a self-immolative linker between the enzyme substrate/trigger group and the reporter moiety in the fluoroently silent probe. As a result of enzyme activation, the trigger group is first removed, and then the self-immolative linker is autonomously eliminated to yield the now highly fluorescent reporter. It is anticipated that by the careful selection of linker characteristics, the optical properties of the probe can be made distinct from those of the reporter, and the linker can be quickly cleaved from the reporter subsequent to trigger group activation.

To that end, we developed probe 1 consisting of a trimethyl-locked quinone propionic acid (Q3PA) trigger group linked to a fluoroently masked naphthalimide (reporter 2) by N-methyl-p-aminobenzyl alcohol, NMPABA. The Q3PA trigger group was selected due to its known rapid and highly selective reduction by NAD(P)H:quinone oxidoreductase-1 (NQO1), an enzyme intimately associated with cancer and overexpressed 2- to 50-fold in the
cytosol of numerous human tumor cells (e.g., colon, breast, pancreas, and nonsmall cell lung). The previously unreported NQO1-selective, turn-on fluorescence probe 1 relies on tuning the push–pull internal charge transfer (ICT) of the naphthalimide scaffold via its attachment to the self-immolative NMPABA linker using an electron-withdrawing carbamate naphthalimide connection. We show here that upon initiation of trigger group removal from the tripartite probe, subsequent rapid cleavage of linker occurs so as to afford reporter 2, which has distinct spectral properties such that NQO1-positive cancer cells can be imaged and identified with a positive-to-negative ratio of ∼500:1.

As seen in Figure 1, the spectroscopic properties of probe 1 and reporter 2 in buffered aqueous media are strikingly different, and these differences are attributed to the presence of the quinone propionic acid trigger group (Q3PA), as well as the carbamate connection between the N-methyl-p-aminobenzyl alcohol, NMPABA, linker, and the naphthalimide fluorophore. The λ\textsubscript{abs} for probe 1 is red shifted roughly 50 nm in comparison to λ\textsubscript{abs} for reporter 2 (432 nm vs 380 nm), as is λ\textsubscript{em} for probe 1 (532 nm vs 480 nm), with both observations being in accord with the presence of the electron-deficient carbamate in probe 1. Interestingly, the absorptivity of probe 1 at the maximum absorption wavelength of reporter 2 is ∼5 times lower than that for probe 1. The \( \Phi \) value of 0.002, which is 95 times less than \( \Phi \) of 0.19, is due to photoinduced electron transfer (PeT)\textsuperscript{15} quenching of the naphthalimide reporter by the Q3PA group. This conclusion is supported by outcomes based on voltammetric and spectroscopic data for probe 1, reporter 2, and Q3PA;\textsuperscript{9} the free energy of the PeT process is −0.98 eV, indicating ready electron transfer from the excited naphthalimide to the electron-poor Q3PA group of probe 1. In comparison to the previously reported probe Q3NI,\textsuperscript{5} there is a 3-fold-larger change in fluorescence efficiency for probe 1 upon NQO1 action to yield the corresponding reporter (\( \Phi \) of reporter/\( \Phi \) of probe), and the fluorescence energy range of reporter 2 is more attractive for possible imaging applications.

We next wanted to demonstrate rapid turning on of reporter 2 by the self-immolative cleavage of the NMPABA linker subsequent to the reduction of probe 1. Dithionate reduction of the Q3PA moiety is complete within 1 s, and the half-life of the lactonization reaction is less than 2 min.\textsuperscript{16} Thus, sodium dithionate was added to aqueous solutions of probe 1, and the resulting fluorescence from reporter 2 was used to determine the extent of probe 1 conversion to reporter 2 (Figure 2).

![Figure 1. Absorbance and emission spectra of 2 μM probe 1 and reporter 2 in pH 7.40, 0.1 M phosphate buffer. T = 25 °C.](image)

Figure 1. Reduction-stimulated turning on of reporter 2. Time-course fluorescence (λ\textsubscript{exc} = 432 nm, λ\textsubscript{em} = 540 nm) from 10 μM probe 1 (pH 7-40, 0.1 M phosphate buffer) after reduction by 16 μM sodium dithionite. The inset contains time-dependent spectra of 2 μM probe 1 upon reduction. T = 25 °C.

Reporter 2 is quickly turned on (\( t_{1/2} \sim 6 \text{ min} \)) under physiologically relevant conditions, pointing to the rapid self-immolative cleavage of the N-methyl-p-aminobenzyl alcohol and CO\textsubscript{2} loss\textsuperscript{8} to yield reporter 2; this occurs because of the electron-rich character of the N-methyl-p-aminobenzyl alcohol vs that of the more typically encountered p-aminobenzyl alcohol linker.\textsuperscript{8,17} Conversion was confirmed by mass spectrometry analysis of probe 1 solutions treated with dithionite.\textsuperscript{9} On the basis of the relatively short time for the turn-on process, the use of probe 1 should allow for detecting the presence of NQO1 enzyme activity.

To evaluate the ability of human NAD(P)H:quinone oxidoreductase-1 (hNQO1) to initiate the decloaking of probe 1 and reveal free reporter 2 upon Q3PA trigger group activation, we utilized the fluorescent product formation technique.\textsuperscript{18} Under in vitro conditions, solutions of probe 1 (2–60 μM) incubated with hNQO1 (40 μg) and its cofactor NADH (100 μM) exhibited steady increases in fluorescence, suggesting a relatively fast rate of reporter 2 production; control experiments with NADH alone did not yield significant changes in fluorescence, a result in accord with that previously shown for the Q3PA trigger group.\textsuperscript{10} In addition, the Q3PA trigger group is stable to reduction/addition reactions in the presence of glutathione and ascorbate as well as dithiothreitol.\textsuperscript{5}
Furthermore, hNQO2 plus its cofactor\textsuperscript{13} is ineffective at releasing reporter 2. From the hNQO1 enzyme experiments, the initial rate of reporter 2 formation, $V$ ($\mu$mol min$^{-1}$ mg hNQO1$^{-1}$), was calculated and then plotted as a function of probe 1 concentration (Figure 3). Apparent kinetic parameters were obtained by fitting the data in Figure 3 to Michaelis–Menten kinetics,\textsuperscript{19} namely, the Michaelis constant ($K_M$) = 10.4 ± 1.0 $\mu$M, maximum velocity ($V_{\text{max}}$) = 0.00225 ± 0.00008 $\mu$mol min$^{-1}$ mg hNQO1$^{-1}$, catalytic constant ($k_{\text{cat}}$) = 0.068 ± 0.002 min$^{-1}$, and catalytic efficiency ($k_{\text{cat}}/K_M$) = 6.52 ± 0.67 $\times$ 10$^3$ M$^{-1}$ min$^{-1}$. Also, there is no apparent inhibition of hNQO1 activity with the [probe 1] values studied. Thus, substantial reporter 2 release occurs upon probe 1 interaction with hNQO1 under physiologically relevant conditions.

To investigate the possibility of differentiating types of cancer cells based on hNQO1 content (positive vs negative), we exposed live hNQO1-positive and hNQO1-negative cells to probe 1 under typical culture conditions (37 °C/CO$_2$ atmosphere). Confocal fluorescence microscopy images of hNQO1-positive colorectal cancer cells (HT29) incubated with probe 1 indicated significant probe 1 uptake and activation, resulting in intense intracellular reporter 2 production. However, hNQO1-negative nonsmall cell lung cancer cells (H596) treated with probe 1 revealed minimal signal (Figure 4), pointing to the intracellular stability of probe 1 in hNQO1-negative cells, similar to what we have observed for another amide-linked naphthalimide reporter, Q,NL.\textsuperscript{5} Wide-field fluorescence micrographs yielded similar results. Upon statistical evaluation of the fluorescence intensities in confocal images of hNQO1-positive (35 samples) and hNQO1-negative (29 samples) cells like those in Figure 4, a value of 11 was found for the positive-to-negative ratio (PNR).

In summary, we have presented the synthesis, properties, and biological evaluation of a novel turn-on fluorescent probe for the cancer-linked enzyme NQO1. The discrimination of human cells possessing NQO1, from those that do not, is achieved in a highly selective fashion, which results from the dramatically increased fluorescence efficiency of the red-shifted reporter 2 that is generated only upon hNQO1 action on the trigger group of probe 1. We are now using the ICT-tunable probe 1 and its variants to develop methods, including multiphoton imaging routes, that allow for the identification of hard-to-detect NQO1-expressing cancerous tissues, as well as the diversification of disease targets\textsuperscript{20,21} for the activation of new trigger groups. With regard to longer-term impacts, enzyme-activatable probes will play important roles as “intelligent” imaging agents applied to the fluorescence-assisted resection of cancerous tissues\textsuperscript{22} and efficacy studies of personalized chemotherapy.\textsuperscript{23}

### ASSOCIATED CONTENT

3 Supporting Information

Synthesis conditions and spectral properties of probe 1 and reporter 2, voltammetric data, and PNR data. This material is available free of charge via the Internet at http://pubs.acs.org.

### AUTHOR INFORMATION

Corresponding Author
tunnel@lsu.edu

Notes

The authors declare no competing financial interest.

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(9) See Supporting Information.