Fluorogenic Label for Biomolecular Imaging

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ABSTRACT

Traditional small-molecule fluorophores are always fluorescent. This attribute can obscure valuable information in biological experiments. Here, we report on a versatile “latent” fluorophore that overcomes this limitation. At the core of the latent fluorophore is a derivative of rhodamine in which one nitrogen is modified as a urea. That modification enables rhodamine to retain half of its fluorescence while facilitating conjugation to a target molecule. The other nitrogen of rhodamine is modified with a “trimethyl lock”, which enables fluorescence to be unmasked fully by a single user-designated chemical reaction. An esterase-reactive latent fluorophore was synthesized in high yield and attached covalently to a cationic protein. The resulting conjugate was not fluorescent in the absence of esterases. The enzymatic activity of esterases in endocytic vesicles and the cytosol educed fluorescence, enabling the time-lapse imaging of endocytosis into live human cells and thus providing unprecedented spatiotemporal resolution of this process. The modular design of this “fluorogenic label” enables the facile synthesis of an ensemble of small-molecule probes for the illumination of numerous biochemical and cell biological processes.

Fluorescent molecules are critical tools in the study of biochemical and cell biological processes (1). In many studies, however, only few of the fluorescent molecules experience a phenomenon of interest. Because traditional fluorophores, such as rhodamine and fluorescein, are always fluorescent, bulk fluorescence can obscure valuable information. To overcome this limitation, molecules can be designed such that a chemical reaction elicits a change in their fluorescence. Such “latent” fluorophores are at the core of common methods, including the enzyme-linked immunosorbent assay (ELISA), high-throughput screening of enzyme inhibitors, detection of reporter genes, and evaluation of cell viability (1). We reasoned that the use of a latent fluorophore as a “fluorogenic label” could overcome limitations of traditional fluorescent labels and thereby improve the spatial and temporal resolution of bioimaging.

Recently, our laboratory reported on a new class of latent fluorophores based on the “trimethyl lock” (2, 3). The rapid lactonization (4, 5) of the trimethyl lock had been exploited previously to prepare stable pro-drugs that were unmasked by an enzyme-catalyzed reaction (6, 7). We first used the trimethyl lock to shroud the fluorescence of a xanthene dye, rhodamine 110 (Rh110) (2), and then an oxazine dye, cresyl violet (3). This approach afforded highly stable bis(trimethyl lock) “pro-fluorophores” that were labile to esterase catalysis in vitro and in cellulo.

Our bis(trimethyl lock) pro-fluorophores had two problematic attributes. First, two chemical reactions were necessary to unveil the vast majority of their fluorescence, decreasing the rate of fluorescence manifestation and limiting the linear range of assays (8, 9). Analogous fluorogenic protease substrates based on a rhodamine diamide display complex hydrolysis kinetics (10–12), as we observed with our bis(trimethyl lock) pro-fluorophores (2, 3). The second problematic attribute was the absence of a...
handle for target–molecule conjugation. Such a handle is available in derivatives, such as 5- or 6-carboxyrhodamine, that are accessible only from low-yielding synthetic routes.

We suspected that we could solve both problems by capping one of the amino groups of Rh110. The capping of rhodamine dyes with an amide (13–15), carbamate (16), or urea (17) can preserve much of their fluorescence. We were especially intrigued by the attributes of urea–rhodamine, which according to recent reports in the scientific (17) and patent (18, 19) literature appears to retain significant fluorescence intensity relative to Rh110.

Here, we report on a versatile fluorogenic label for biomolecular imaging. First, we describe the synthesis of a complete set of ureated and amidated derivatives of Rh110, as well as a characterization of their fluorescent properties. Then, we show that imposing our trimethyl lock strategy upon a urea–rhodamine yields a stable latent fluorophore with a high rate of enzymatic hydrolysis. Finally, we demonstrate the power of our modular approach by using the urea moiety as a handle for protein conjugation and subsequent continuous imaging of endocytosis by live human cells.

RESULTS AND DISCUSSION

Synthesis of Model Compounds. To gain a comprehensive understanding of the urea and amide derivatives of rhodamine, we undertook the synthesis of compounds 1–5 (Table 1). Rhodamine itself and these five derivatives encompass the ensemble of possible ureated and amidated derivatives. We were especially interested in those properties of 1–5 with biological implications, such as the extinction coefficient and quantum yield in aqueous solution. Previous reports (10, 11, 17–20) of similar derivatives did not provide a complete listing of relevant fluorescent characteristics.

Installation of the urea moiety to produce urea 1 proved to be surprisingly difficult. In our hands, the reported conditions (17) involving the reaction of Rh110 with a carbamoyl chloride using Hüning's base gave an intractable mixture of products. In contrast, we found that Rh110 was deprotonated effectively with NaH and that the resulting anion reacted with dimethylcarbamyl chloride to yield the desired urea 1. This deprotonation strategy also proved useful for the synthesis of amide 2 and diurea 3. The additional acetamide group in urea–amide 4 and diamide 5 were installed by reaction with acetyl chloride in the presence of a base.

Fluorescence Properties. The absorbance and fluorescence spectra of Rh110 and each derivative are shown (Figure 1). The corresponding values of λmax, extinction coefficient at λmax (ε), λem, and quantum yield (Φ) are listed (Table 1). We determined the relative fluorescence intensity of these compounds by calculating the product of extinction coefficient and quantum yield and then normalizing these values to those of Rh110. In our measurement, urea 1 retained 35% of the fluorescence intensity of Rh110 with a quantum yield value of 0.49. Amide 2 is only 12% as fluorescent as Rh110, which is consistent with earlier reports (10, 11). The fluorescence of the bis-substituted dyes was largely quenched in aqueous solution. Diurea 3 did, however, possess significant absorbance and fluorescence compared to the urea–amide 4 or diamide 5. These latter two rhodamine derivatives are essentially nonfluorescent.

We also determined the pH dependence of the fluorescence of urea 1 and amide 2. The fluorescence of Rh110 is relatively insensitive to pH values between 4 and 10 (1). This property is beneficial in biological assays, where unknown variations in pH can hamper quantitative measurements. Like Rh110, urea 1 and amide 2 show no significant spectral change between pH values of 4 and 10; details may be seen in Supplementary Figure 1.
Substituent effects on the fluorescent properties of rhodamine dyes are challenging to predict or interpret due to the complexity of the rhodamine system (21). In solution, rhodamine derivatives exist in equilibrium between a zwitterion that absorbs visible light and is fluorescent and a lactone that is colorless and nonfluorescent. Substitution on nitrogen can affect both this open–closed equilibrium and the spectral characteristics of the fluorescent zwitterions (22, 23). We suspected that the differences in optical properties seen in compounds 1–5 could be rationalized, in part, through examination of the electron-donation capability of the different nitrogen substituents. In agreement with this reasoning, weakly donating substituents would favor the colorless lactone as well as decrease the intrinsic absorptivity of the zwitterions and, hence, the extinction coefficient. Weakly donating substituents could also reduce the quantum yield by decreasing the C–N bond order and thereby enhancing nonradiative decay of the excited state through vibrational relaxation processes (24, 25).

We explored the relationship between the values of extinction coefficient and quantum yield and the Hammett $\sigma_p$ substituent constants (26). An unprotonated amino group is a good electron donor ($\sigma_p = -0.66$), whereas an amide group is a relatively poor donor ($\sigma_p = 0.00$), due to amidic resonance. A urea group is peculiar; its carbonyl group is cross-conjugated and both of its nitrogens participate in amidic resonance. This cross-conjugation attenuates its electron-donating ability, as reflected in an intermediate Hammett constant ($\sigma_p = -0.26$). A plot of both extinction coefficient and quantum yield versus $\sigma_p$ substituent constant for Rh$_{110}$ and monosubstituted rhodamines 1 and 2 is shown (Figure 2). The correlation indicates that both spectral properties are affected by electron donation from the nitrogens. A similar trend in quantum yields has been observed in substituted phenoxazinone dyes (27).

The moderate electron-donating character of the urea moiety provides an explanation for the advantageous properties of urea 1. Substitution with the cross-conjugated urea suppresses the fluorescence intensity of urea 1 relative to Rh$_{110}$. This decrease is not, however, as severe as seen in amide 2, due to the greater electron-donating properties of the urea moiety. Still, the attenuated electron-donation allows complete suppression of fluorescence upon amidation of the remaining nitrogen in urea–amide 4. Finally, the effect of electron-rich substituents on the rhodamine system is apparent again in the fluorescence of diurea 3, as it is greater than that of diamide 5.

**Synthesis of Urea–Rhodamine Trimethyl Lock.**

Having affirmed the desirable properties of urea–rhodamine, we next sought to apply our trimethyl lock strategy to this dye. The synthetic route to the fluorogenic substrate, which employs rhodamine morpholinourea 6 (17), is shown (Scheme 1). Again, we found that the use of Hünig’s base in the synthesis afforded a mixture of products. Deprotonation of Rh$_{110}$ with NaH followed by dropwise addition of 4-morpholinecarbonyl chloride furnished rhodamine morpholinourea 6. This compound exhibited similar fluorescent characteristics to urea 1 (Table 1), and it had an extinction coefficient of
51 700 M$^{-1}$ cm$^{-1}$ and quantum yield of 0.44. Carbodiimide coupling of rhodamine morpholino-urea 6 with acid 7 (28) afforded the desired pro-fluorophore 8.

**Chemical Stability.** Pro-fluorophore 8 must be stable in aqueous solution to be useful in biological assays. Such stability can be problematic for hydrolase substrates, as spontaneous hydrolysis can compete effectively with enzymatic activity and raise background levels. As shown (Figure 3), pro-fluorophore 8 showed remarkable stability in both phosphate-buffered saline (PBS) and Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% (v/v) fetal bovine serum (FBS). In contrast, fluorescein diacetate, which is a widely used esterase substrate (29), suffered relatively rapid hydrolysis in both solutions. This dramatic increase in stability arises from the large difference in $pK_a$ values between the conjugate acids of the two leaving groups. Specifically, fluorescein ($pK_a$ 6.32 (30)) is a much better leaving group than is the electron-rich trimethyl-lock phenol ($pK_a$ of 10.28 (31)).

**Enzymatic Reactivity.** An objective in the design of pro-fluorophore 8 was to improve its reactivity as an esterase substrate relative to the original bis(trimethyl lock) rhodamine substrate. The appearance of fluorescence upon reaction of porcine liver esterase (PLE) with pro-fluorophore 8 was indicative of single-hit kinetics (Figure 4). The kinetic constants were calculated to be $k_{cat}/K_M = 8.2 \times 10^5$ M$^{-1}$ s$^{-1}$ and $K_M = 0.10$ $\mu$M. Comparison with the apparent kinetic constants from the original bis(trimethyl lock) rhodamine substrate (2) ($k_{cat}/K_M = 1.9 \times 10^3$ M$^{-1}$ s$^{-1}$ and $K_M = 0.47$ $\mu$M) shows a 430-fold increase in the $k_{cat}/K_M$ value. A more appropriate comparison takes into account the expected 65% decrease in fluorescence of urea 6 (Table 1), which is the hydrolysis product of pro-fluorophore 8, relative to Rh110. After this adjustment, latent fluorophore performance is still enhanced by 150-fold.

The substantial increase in catalytic efficiency is likely due to the change from the double-hit kinetics observed for the bis-substituted substrate to the single-hit kinetics of pro-fluorophore 8. Hydrolysis of the bis-substituted fluorogenic substrate progresses from diamide to free Rh110 via a monoamide intermediate, with the unmasking of the second amino group producing the majority (~90%) of the fluorescence (10, 11). In contrast, the urea–rhodamine substrate requires only a single cleavage event for the complete manifestation of fluorescence.

**Cellular Imaging.** Once the high chemical stability and enzymatic reactivity of pro-fluorophore 8 were established, we next evaluated the behavior of this compound in live human cells. Pro-fluorophore 8 was incubated with HeLa cells and imaged using confocal fluorescence microscopy. The substrate was activated in cellulo.

![Scheme 1. Synthetic Route to Pro-Fluorophore 8](image-url)

![Figure 3. Stability of pro-fluorophore 8 and fluorescein diacetate in aqueous solution. a) Time course for the generation of fluorescence ($\lambda_{ex} = 496$ nm, $\lambda_{em} = 520$ nm) of pro-fluorophore 8 (25 nM) and fluorescein diacetate (25 nM) in PBS. b) Time course for the generation of fluorescence ($\lambda_{ex} = 496$ nm, $\lambda_{em} > 520$ nm) of pro-fluorophore 8 (25 nM) and fluorescein diacetate (25 nM) in DMEM containing FBS (10%, v/v).](image-url)
Fluorogenic Label. The high chemical stability and rapid in cellulo unmasking of pro-fluorophore 8 prompted us to develop a derivative for bioconjugation. We reasoned that such a fluorogenic label would be stable enough to survive conjugation and purification protocols while still providing a strong signal for continuous biological experiments. It is noteworthy that simple fluorescein diesters have found only limited use as fluorogenic labels (32–35). As fluorescein diesters suffer from toxic reactivity, we attached it to a thiol-reactive fluorogenic label 9. This fluorogenic label 9 reacted cleanly with thiol groups of rhodamines has been used previously in cellulo (46, 47). Deprotection of maleimidourea–rhodamine–t-Boc 11 with TFA afforded fluorescent urea–rhodamine 12. Condensation with 7 using EDC gave thiol-reactive fluorogenic label 13.

Bioconjugation. To test the utility of fluorogenic label 13 in a biological experiment, we attached it to a thiol-containing variant of bovine pancreatic ribonuclease (RNase A (48)). RNase A is a cationic protein that is internalized by mammalian cells via endocytosis (49). This internalization is critical to the action of cytotoxic RNase A variants and homologues (50). Fluorogenic label 13 reacted cleanly with the A19C variant of bovine pancreatic ribonuclease (RNase A) to give a mono-substituted conjugate as determined by MALDI mass spectrometry. This protein conjugate was stable to purification by cation-exchange chromatography at pH 5.0 and showed a 1200-fold increase in fluorescence upon incubation with PLE (data not shown).

At physiological pH, the protein conjugate was less stable than unconjugated pro-

Figure 4. Kinetic traces (λex = 496 nm, λem = 520 nm) and Michaelis–Menten plot (inset) for a serial dilution of pro-fluorophore 8 (0.5 μM → 2 nM) with PLE (2.5 μg mL⁻¹).
fluorophore 8. Spontaneous hydrolysis of the acetate ester was slow but significant in PBS, presumably because conjugation to the protein places the probe in close proximity to nucleophilic functional groups of the protein. Storage at pH 5.0 did, however, extend the stability of the conjugate, allowing multiple experiments to be performed with one preparation.

**Cellular Imaging with a Bioconjugate.** Fluorescently labeled biomolecules have been used to image endocytic events (51). We sought to determine the efficacy of our fluorogenic label approach by comparing endocytosis of HeLa cells incubated with Oregon Green-labeled RNase A (49) to that of cells incubated with the protein conjugated with fluorogenic label 13. The Oregon Green conjugate showed intense extracellular background signal that obscures the fluorescence from endocytosed material (Figure 6, panel a). This background could be eliminated only with many vigorous washing steps (Figure 6, panel b). In contrast, the pro-fluorophore conjugate allowed imaging without intermediate washing steps. Unwashed HeLa cells incubated with the RNase A conjugate showed intense extracellular background signal that obscures the fluorescence from endocytosed material (Figure 6, panel a). This background could be eliminated only with many vigorous washing steps (Figure 6, panel b). In contrast, the pro-fluorophore conjugate allowed imaging without intermediate washing steps. Unwashed HeLa cells incubated with the RNase A conjugate showed intense extracellular background signal that obscures the fluorescence from endocytosed material (Figure 6, panel a). This background could be eliminated only with many vigorous washing steps (Figure 6, panel b). In contrast, the pro-fluorophore conjugate allowed imaging without intermediate washing steps. Unwashed HeLa cells incubated with the RNase A conjugate showed intense extracellular background signal that obscures the fluorescence from endocytosed material (Figure 6, panel a). This background could be eliminated only with many vigorous washing steps (Figure 6, panel b). In contrast, the pro-fluorophore conjugate allowed imaging without intermediate washing steps.
gate have bright, punctate staining (Figure 6, panel c), indicative of the conjugate being localized in small vesicles. Counterstaining with LysoTracker Red shows a large degree of colocalization (Figure 6, panel d), suggesting that the latent conjugate is internalized via endocytosis and activated by endosomal or lysosomal esterases (52–54). Images with the protein conjugate (Figure 6, panel c) are less diffuse and more punctate that are images with free pro-fluorophore 8 (Figure 5), which has much more ready access to the cytosol. To ensure that the signal is due to unmasked fluorophore attached to RNase A (Figure 6, panel c), we fixed cells incubated with our latent conjugate and counterstained them with a primary antibody to RNase A and a secondary antibody labeled with AlexaFluor 594. In a fluorescence microscopy image, we observed a significant overlap of the green and red fluorescent signals to produce a yellow signal, indicating that the unmasked RNase A conjugate is largely intact; details may be seen in Supplementary Figure 3.

The high chemical stability and low background fluorescence of the fluorogenic label conjugate allowed for the time-lapse imaging of its endocytosis. Cells were incubated with the fluorogenic label RNase A conjugate at room temperature, and images were recorded without washing during the next 90 min. The compilation of these images into a movie revealed that internalization of the conjugate occurred continuously and that vesicular fluorescence increased monotonically; details may be seen in the Supplemental Movie.

Envoi. We have demonstrated how a common fluorophore, Rh110, can be elaborated into a powerful new tool for biochemistry and cell biology. The use of a trimethyl lock provides a latent fluorophore with high chemical stability while maintaining enzymatic reactivity (2, 3). The use of a urea group (rather than a second trimethyl lock) improves enzymatic reactivity markedly while preserving desirable fluorescence properties, as in pro-fluorophore 8. The elaboration of the urea to include an electrophile outfits the latent fluorophore for conjugation, as in fluorogenic label 13. Conjugation of this fluorogenic label to a target molecule enables, for example, the continuous imaging of the endocytosis of a target molecule by live human cells.

We note that the urea–rhodamine–trimethyl lock probe is modular and, hence, can be tailored to suit a variety of applications (Scheme 3). For example, alteration of the bioconjugative group on the urea moiety could be used to change conjugation chemistry, enhance cellular internalization, or target a conjugate to a specific subcellular location. Modification of the enzyme-reactive group on the trimethyl lock could enable the detection of a conjugate in a particular organelle. The use of fluorogenic labels could even transcend cultured cells, allowing for continuous imaging in tissues or in vivo. These applications would be facilitated by extant comprehensive inventories of the enzymes in numerous organs and organelles (55, 56). Accordingly, the fluorogenic label strategy could enable the development of specific probes for biological experiments of ever-increasing sophistication.

METHODS

General Spectroscopic Methods. HEPES (2-[4-(2-hydroxyethyl]-1-piperazine)ethanesulfonic acid) was from Research Products International. Fluorescein (reference standard grade) was from Molecular Probes. Other reagents were from Sigma-Aldrich or Fisher Scientific. Phosphate-buffered saline, pH 7.4 (PBS), contained (in 1.00 L) KCl (0.20 g), KH2PO4 (0.20 g), NaCl (8.0 g), and Na2HPO4·7H2O (2.16 g). All measurements were recorded at ambient temperature (23 ± 2°C), and buffers were not degassed prior to measurements. Compounds were prepared as stock solutions in DMSO and diluted such that the DMSO concentration did not exceed 1% (v/v). Porcine liver esterase (PLE, MW = 163 kDa (57)) was obtained from Sigma Chemical (product number E2884) as a suspension in 3.2 M (NH4)2SO4 and was diluted to appropriate concentrations in PBS before use. In pH dependency studies, the pH of PBS was adjusted by addition of 1.0 M HCl or 1.0 M NaOH and measured using a Beckmann glass electrode that was calibrated prior to each use. Graphs were manipulated and parameters were calculated with Microsoft Excel 2003 and GraphPad Prism 4.

Ultraviolet–Visible and Fluorescence Spectroscopy. Absorption spectra were recorded in 1-cm path length cuvettes having a volume of 1.0 or 3.5 mL on a Cary model 50 spectrometer from Varian. The extinction coefficients were measured in 10 mM HEPES–NaOH buffer, pH 7.5. Fluorometric measurements were made using fluorescence grade quartz or glass cuvettes from Starna Cells and a QuantaMaster1 photon-counting spectrophuorometer from Photon Technology International equipped with sample stirring. The quantum yields of Rh110 and compounds 1–5 were measured with dilute samples (A < 0.1) in 10 mM HEPES–NaOH buffer, pH 7.5. These values were obtained by the comparison of the integrated area of the emission spectrum of the samples with that of fluorescein in 0.1 M NaOH, which has a quantum efficiency of 0.95 (58). The concentration of the fluoro-
rescin reference was adjusted to match the absorbance of the test sample at the excitation wavelength. Under these conditions, quantum yields were calculated using eq 1.

\[ \Phi_{\text{sample}} = \Phi_{\text{standard}} \frac{F_{\text{em, sample}}}{F_{\text{em, standard}}} \]  

(1)

Protein Purification and Labeling. The TNB-protected A19C variant of RNase A and the Oregon Green-labeled RNase A conjugate were prepared as described previously (49). The TNB-protected protein was deproteinated with a 3-fold molar excess of dithiothreitol (DTT) and desalted by chromatography using a HiTrap Desalting column (Amersham). The protein conjugate was then prepared by reaction with a 10-fold molar excess of thiolic-reactive maleimide 13 for 16 h at 4 °C. Purification by chromatography using a HiTrap HP SP column (Amersham) afforded the desired conjugate (MS (MALDI): m/z 14 468 (expected, 14 475)).

Protein concentration was determined by using a bicinecinonic acid (BCA) assay kit from Pierce with wild-type RNase A as a standard.

Cell Staining: Cells were plated on Nunc Lab-Tek II 8-well Chamber Coverglass (Fisher Scientific) and grown to 70–80% confluence at 37 °C in DMEM (Invitrogen) containing FBS (10% v/v). For static imaging, cells were first washed with Dulbecco’s phosphate-buffered saline (DPBS, Invitrogen). Cells were then incubated with pro-fluorophore 8 (10 μM), RNase A conjugated to maleimide 13 (10 μM), or Oregon Green-labeled RNase A (10 μM) for 1 h at 37 °C prior to imaging. Nuclear staining was accomplished by addition of Hoechst 33342 (2 μg/mL) for the final 5 min of incubation. Lysosomal staining involved washing the cells with DPBS followed by incubation with 100 nM LysoTracker Red (Molecular Probes) in DPBS for 1 min at ambient temperature. For dynamic imaging, cells were incubated with Hoechst 33342 (2 μg/mL) for 5 min at 37 °C, and then washed twice with DPBS. Pro-fluorophore 13–RNase A conjugate (10 μM) was added to the cells at ambient temperature (23 ± 2 °C). Imaging of endocytosis was performed 1 min after the addition of the conjugate.

Cell Imaging. Cells were imaged on a Nikon Eclipse TE2000-U confocal microscope equipped with a Zeiss AxioCam digital camera, unless indicated otherwise. Excitation at 408 nm was provided by a blue-diode laser, and emission light was passed through a filter centered at 515 nm with a 75-nm band-pass. For time-lapse imaging, one image per minute was recorded during the first 30 min of incubation, two images per min were recorded during the next 10 min, and five images per min were recorded during the last 50 min. The resulting movie condenses these 300 images recorded over 90 min into 40 s. Brightfield images indicated that the cells were alive and appeared to have normal physiology, both before and after the time-lapse imaging.

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Supporting Information Available: This material is available free of charge via the Internet.

REFERENCES


