Receptor-mediated Targeting of Fluorescent Probes in Living Cells*

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A strategy was developed to label specified sites in living cells with a wide selection of fluorescent or other probes and applied to study pH regulation in Golgi. cDNA transfection was used to target a single-chain antibody to a specified site such as an organelle lumen. The targeted antibody functioned as a high affinity receptor to trap cell-permeable hapten-fluorophore conjugates. Synthesized conjugates of a hapten (4-ethoxymethylene-2-phenyl-2-oxazolin-5-one, phOx) and fluorescent probes (Bodipy FL, tetramethylrhodamine, fluorescein) were bound with high affinity (~5 nM) and specific localization to the single-chain antibody expressed in the endoplasmic reticulum, Golgi, and plasma membrane of living Chinese hamster ovary cells. Using the pH-sensitive phOx-fluorescin conjugate and ratio imaging microscopy, pH was measured in the lumen of Golgi (pH 6.25 ± 0.06). Measurements of pH-dependent vacuolar H⁺/ATPase pump activity and H⁺ leak in Golgi provided direct evidence that resting Golgi pH is determined by balanced leak-pump characteristics rather than the inability of the H⁺/ATPase to pump against an electrochemical gradient. Like expression of the green fluorescent protein, the receptor-mediated fluorophore targeting approach permits specific intracellular fluorescence labeling. A significant advantage of the new approach is the ability to target chemical probes with custom-designed spectral and indicator properties.

Small-molecule fluorescent probes have been widely used to study protein localization, cytoplasmic ionic content (1), and solute diffusion (2). Probes are available with high intrinsic brightness and excitation and emission peaks from ultraviolet to infrared wavelengths; however, in general these chemical probes cannot be targeted to specific sites in living cells. The green fluorescent protein (GFP)1 is a genetically targetable probe that has been used extensively to study gene expression and protein localization in living cells (3, 4). However, GFP fluorescence is limited to blue, low green, and green/yellow variants, which have relatively low intrinsic brightness (4, 5). pH-based indicators are currently limited to measuring pH (6–8), Ca²⁺ (9, 10), and membrane potential (11).

A cell labeling method is reported here that combines the site specificity conferred by genetically encoded targeting sequences with the excellent spectral and indicator properties of small chemical fluorophores. The strategy is to express a high affinity “receptor” at a specified intracellular location to trap a conjugate of a fluorophore linked to a receptor “ligand” (Fig. 1a). We chose a single-chain antibody (sFv) (12) as the receptor and a hapten (phOx) as the ligand. Although many receptor-ligand pairs are possible, the antibody-hapten pair was selected because of the simple ligand-probe chemistry and high affinity interaction without interference from cellular factors. For sFv targeting, cells are transfected with cDNAs encoding sFv in fusion with targeting sequences. Fluorophore-hapten conjugates are added to the extracellular solution at low concentrations, diffuse to sites of sFv expression, and bind to the sFv. Conjugates of different indicator and spectral properties were synthesized (Fig. 1b), including phOx-Bodipy FL (green fluorescent), phOx-fluorescein (green fluorescent, pH-sensitive), and phOx-tetramethylrhodamine (red fluorescent). The flexible linkers were designed to permit stacking of the unbound hapten with its covalently attached fluorophore to form a dark complex and reduce background fluorescence.

The ability to label cellular sites with fluorescent probes with varied spectral and indicator properties was demonstrated, and the phOx-fluorescein conjugate was applied to measure pH in the Golgi lumen. Here we report the first in vivo measurement of the regulation of the pump rate of the vacuolar H⁺/ATPase by Golgi luminal pH to test the thermodynamic model (13) of how the resting pH is set in organelles such as the Golgi.

EXPERIMENTAL PROCEDURES

Plasmids and Cell Transfection—The cDNA encoding the sFv (25-kDa protein) and two e-Myc epitopes was amplified by polymerase chain reaction using the plasmid pHook1 (12) (Invitrogen) as template, with sense primer (5′GGGATCCCGAGGCCTCAAGGTGAGGAGG) containing an EcoRI site (underlined) and antisense primer containing an XbaI site (underlined) (5′GCTCTAGATGGCCCACAGGATCAGTCTCAGGAGG) containing specific targeting sequences as described in Ref. 6. Plasma membrane targeting was achieved with the pHook1 plasmid (12). CHO cells (ATCC CRL 9618) were transfected with plasmids encoding targeted sFv using LipofectAMINE (Life Technologies, Inc.) as described previously (6).

Synthesis of Hapten-Fluorophore Conjugates—A flexible linker was added to phOx (Sigma) by reaction of 50 mg of phOx with 14.3 μl of 1,5-diaminopentane (Aldrich) in 2.5 ml of acetone for 1 h. The disubstituted amine was precipitated by the addition of 2 volumes of 50 mM borate buffer (pH 9.2), leaving the product in solution. A more rigid linker was added to phOx (Sigma) by reaction of 50 mg of phOx with 14.3 μl of 1,5-diaminopentane (Aldrich) in 2.5 ml of acetone for 1 h. The disubstituted amine was precipitated by the addition of 2 volumes of 50 mM borate buffer for 2 h. The product was obtained as a precipitate.

The abbreviations used are: GFP, green fluorescent protein; phOx, 4-ethoxymethylene-2-phenyl-2-oxazolin-5-one; CHO, Chinese hamster ovary.
equimolar amounts of phOx (0.5 mg in 25 \( \mu l \) of acetone) and fluorescein cadaverine (1 mg in 50 \( \mu l \) of dimethylformamide; Molecular Probes) for 1 h. Phosphate-buffered saline was added, unreacted phOx was removed by hexane extraction, and the product was extracted with butanol. phOx-ethanolamine was prepared by reaction of 1 mg of phOx with 0.3 \( \mu l \) of ethanolamine in 10 ml of ethanol for 1 h. Reactions were conducted at room temperature. Products were judged to be >95% pure by TLC, and structures were confirmed by mass spectrometry.

Fluorescence Measurements—Cells were labeled at 2 days after transfection by incubation with low concentrations (<100 nm) of the conjugates. Unless otherwise indicated, cells were observed in the absence of the conjugate in the bathing media. Images were recorded at room temperature on a K2 BIO microscope (Technical Instruments) equipped with a 60\( \times \) PlanApo objective (Nikon, N.A. 1.4), coaxial-confocal attachment, and cooled CCD camera. Dual excitation ratio images of fluorescein were acquired using 440- and 490-nm excitation filters and a 520-nm long pass emission filter. Continuous recordings of the fluorescence time course were obtained on a Nikon Diaphot epifluorescence microscope equipped with a 100\( \times \) PlanApo objective (Nikon, N.A. 1.4), a photomultiplier using a 530-nm bandpass emission filter, and an optical filter changer (model 10-C, Sutter Instrument Co.) containing 440- and 490-nm excitation filters. Cuvette fluorescence measurements were conducted on an SLM 8000c fluorometer (SLM Aminco). Spectra were recorded with 4-nm slit widths; time courses were acquired with 495-nm excitation and 510-nm long pass emission filters.

RESULTS

For effective targeting, the requirements of the sFv/hapten-fluorophore system include: bright fluorescence of the bound conjugate, high affinity binding of the conjugate to sFv, stability of the conjugate, minimal cellular toxicity, strong cellular expression of functional sFv, and membrane permeability of the conjugate. These requirements were fulfilled for cellular sFv expression (using the Golgi, ER, and plasma membrane vectors) and binding of the conjugates in Fig. 1c. Fluorescence spectra of phOx-fluorescein bound to sFv and of unbound phOx-fluorescein in solution had similar spectral shapes (Fig. 1c). As intended, the fluorescence of the unbound conjugate was decreased considerably (by 5-fold) over that of the bound conjugate. Images of CHO cells expressing sFv at the plasma membrane were acquired with increasing concentration of phOx-fluorescein. The fluorescence from sFv-bound phOx-fluorescein gave a dissociation constant \((K_d)\) of 6.8 nm (Fig. 1d). This agrees with the value of 5.5 nm obtained in CHO cell suspensions expressing sFv at the plasma membrane (not shown). At 10 nM phOx-fluorescein, fluorescence from free dye was 75 times lower than that of bound dye. No significant fluorescence from non-sFv-expressing cells was seen.

The toxicity and stability of the conjugates were investigated. There were no differences in cell growth as assessed by cell counting and viability as assessed by trypan blue exclusion between control cells and cells incubated for 24 h with 200 nM of each conjugate. The stability of the imine bond in the conjugates was determined in cells. In freshly prepared phOx-fluorescein and phOx-fluorophore incubated with cell suspensions for up to 7 h, the imine bond was hydrolyzed in 0.5 M NaOH, resulting in increased fluorescence. The fluorescence increase after treatment with NaOH was the same in both samples, indicating that the imine bond was not hydrolyzed in cells.

The various sFv targeting constructs and phOx conjugates were studied in CHO cells. Fig. 2a shows a fluorescence image of living cells expressing the sFv at the plasma membrane and labeled with phOx-rhodamine. A plasma membrane staining pattern was found. Fig. 2b shows staining of sFv in the same cells with a fluorescein-labeled anti-c-Myc antibody. Comparison with Fig. 2a demonstrates that only sites of sFv expression

![Diagram](image-url)
were significantly labeled with phOx-rhodamine. There was no significant staining of adjacent cells that did not express sFv (Fig. 2c). Fig. 2d shows specific phOx-Bodipy staining of Golgi. Staining was reversed by addition of 1 μM phOx-ethanolamine (Fig. 2e) but not by 1 μM ethanolamine. Fig. 2f shows phOx-Bodipy staining of ER, seen as a characteristic reticular pattern. The high expression level of the sFv, the relatively high affinity of the hapten/sFv, and the low fluorescence of the unbound conjugate allowed images to be obtained in the presence of <10 nM of unbound conjugate with little contribution from free conjugate. Leakage out of the Golgi, which required dissociation from the sFv and diffusion through lipid membranes and unstirred layers, had a half-time of tens of minutes.

The subcellular location of expressed sFv was confirmed by immunofluorescence. Cells transfected with the Golgi-sFv construct showed perinuclear staining by a fluorescein-labeled, anti-c-Myc antibody (Fig. 2g), which colocalized with staining by antibodies against the Golgi marker 58-kDa protein (14) (Fig. 2h). Cells transfected with the ER-sFv construct showed a reticular staining pattern with the c-Myc antibody (Fig. 2i), which colocalized with staining by fluorescein-labeled concanavalin A, an ER marker (15) (Fig. 2j). The membrane permeability of the conjugates was high enough to load cells by incubation at 37 °C for 4 h for phOx-fluorescein, 2 h for phOx-rhodamine, or 10 min for the less polar phOx-Bodipy. Cells could be loaded at 4 °C, indicating that the conjugate entered the cells primarily by transmembrane diffusion and not by endocytosis. These results demonstrate the selective targeting of fluorescent probes to expressed sFv in living cells.

Organelle-specific sFv targeting was applied to measure Golgi pH using phOx-fluorescein as the probe. Ratio images were calculated from images of Golgi labeled with phOx-fluorescein acquired at 440- and 490-nm excitation wavelengths. To convert ratios to absolute pH, cells were perfused with “calibration buffers” at different pH values containing high K+ and the ionophore monensin to equalize extracellular and Golgi lumenal pH. The dependence of the fluorescence ratio on pH was measured for Golgi and plasma membrane-expressed sFv (Fig. 3a). The apparent pKₐ of 6.56 of bound phOx-fluorescein was not different from that of unbound phOx-fluorescein in solution (pKₐ = 6.54). The average Golgi fluorescence ratio of 1.05 ± 0.05 corresponds to a pH of 6.25 ± 0.06, in agreement with previous estimates (7, 16, 17).

The Golgi-targeted phOx-fluorescein was used to detect continuous changes in lumenal pH in individual cells. Fig. 3b shows that the fluorescence ratio increases upon addition of the vacuolar H⁺ pump inhibitor bafilomycin A₁. The ratios measured using calibration buffers were used to convert fluorescenes to pH.
ence ratios to pH (scale at right). Golgi pH initially at \(-6.3\) promptly alkalinized after the addition of bafloycin A1.

It has been proposed that the steady-state Golgi pH is determined thermodynamically by the free energy of ATP hydrolysis used by the vacuolar H\(^+\)/ATPase to pump H\(^+\) against an electrochemical gradient (13). To test the model prediction that the H\(^+\) pump rate is 0 at steady-state pH, the rates of Golgi H\(^+\) pump and leak were measured as a function of Golgi lumenal pH. After Golgi alkalinization by a 20 mM sodium acetate prepulse, H\(^+\) pumping into the Golgi restores steady-state pH (Fig. 3c). The initial rate of pH change (dashed line) is the difference between pump and leak rates: \(d\text{pH}/dt = (d\text{pH}/dt)_{\text{back}} - d\text{pH}/dt_{\text{pump}}\). The buffer capacity, \(\beta\), was measured by the NH\(_4\)Cl pulse method (18) to be constant (38 ± 3 mV/pH units) in the pH range 6–7 (not shown). The H\(^+\) leak rate, \(d\text{pH}/dt_{\text{leak}}\), was measured from the pH change (Fig. 3c, solid line) after an identical 20 mM sodium acetate prepulse with the H\(^+\) pump inhibited by bafloycin A1. Similar prepulse measurements were done at different Golgi pH by varying sodium acetate and NH\(_4\)Cl concentrations. There should be little effect of acetate or NH\(_4\)\(^+\) on H\(^+\) transport in the prepulse method because these ions have left the cell by the time of the H\(^+\) transport measurements (18). Fig. 3d shows that the computed H\(^+\) pump rate increases sharply with Golgi pH and that the pump rate is not 0 at the Golgi steady-state pH of \(-6.25\). Although it is recognized that the prepulse method used to alter Golgi pH also alters cytoplasmic pH, it has been reported that the H\(^+\) pump rate is relatively insensitive to cytoplasmic pH in both mammalian (17) and plant (19) systems.

In the steady state, H\(^+\) pump rate must equal H\(^+\) leak rate. The dependence of the leak rate on Golgi pH was measured from the kinetics of pH change after bafloycin A1 addition as shown in Fig. 3b. The data for different pH are summarized in Fig. 3d, showing decreased H\(^+\) leak as Golgi pH increases. The intersection of the H\(^+\) pump and leak curves predicts correctly the observed steady-state Golgi pH, supporting a balanced pump/leak mechanism for settling Golgi pH.

**DISCUSSION**

The fluorophore targeting method reported here provides a new strategy for cellular labeling that complements existing methods. Of the current methods, GFP is the easiest to use because it does not require exogenously added reagents or cofactors. However, the available GFP mutants are limited in terms of spectral properties (brightness and excitation and emission wavelengths) and indicator sensitivities. Recently, a method was developed for covalently labeling proteins in living cells with a fluorescein derivative (FLASH) (20). Addition of an arseno-fluorescein derivative to the cells leads to covalent attachment of the derivative to a short \(\alpha\)-helix containing 4 cysteines added to the protein of interest. An advantage of FLASH over GFP and the current approach is the significantly smaller size of the protein tag; however, FLASH is currently limited to the use of fluorescein as the probe.

The addition of labeled macromolecules either requires microinjection (21) or is limited to compartments that are accessible by endocytosis (22) or retrograde transport through the secretory pathway (17). GFP, FLASH, and the receptor-mediated targeting methods use genetically encoded targeting sequences to localize fluorophores to virtually any cellular site, provided that the targeted protein is able to fold properly. Although strong sFv expression was found in plasma membranes and various intracellular compartments (Fig. 2, above), preliminary experiments suggest that functional sFv expression is relatively poor in reducing environments (data not shown). If disulfide bond formation is critical to sFv folding, it may be possible to generate sFv mutants lacking disulfide bonds that fold well in reducing environments (23).

The ability to target fluorophores of varied spectral properties is a distinct advantage of the receptor-mediated targeting approach over GFP. The receptor-mediated targeting method utilizes fluorescent probes with a potentially wide range of excitation and emission wavelengths and other optical properties. The use of multiple probes with well separated excitation and emission spectra allows simultaneous labeling of multiple sites. For measurements of important cellular parameters, small chemical probes sensitive to ions (Ca\(^{2+}\), Na\(^+\), K\(^+\), H\(^+\), Cl\(^-\)), viscosity, and membrane potential are available.

Receptor-mediated targeting of a hapten-fluorescein conjugate was used to label the Golgi with a fluorescent pH indicator permitting the measurement of the dependence of the vacuolar H\(^+\)/ATPase pump rate on Golgi pH. The pump rate increased and the leak rate decreased as Golgi pH increased. The steep dependence of pump rate on pH is in agreement with measurements of ATPase activity (24) of the vacuolar H\(^+\)/ATPase made *in vitro* and of the pump rate made in phagosomes (25). In contrast to the predictions of a thermodynamic model of pH regulation in the Golgi, the net pump rate was not 0 at the resting Golgi pH. Thus, the resting pH is determined by the kinetics of proton leak versus pump. Shifts in the leak or pump curves could account for the differences in resting pH in organelles of the secretory pathway.

In summary, the receptor-mediated probe targeting strategy allows the labeling of specified cellular structures with fluorescent or other indicator molecules. This targeting method can readily be extended to deliver conjugates containing magnetic resonance probes, caged compounds, or chemical cross-linkers. Finally, the use of cell-specific promoters and gene transfer should allow the *in vivo* targeting of hapten-probe complexes to specific cell types in multicellular organisms.

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