

A fluorophore ligase for site-specific protein labeling inside living cells

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Biological microscopy would benefit from smaller alternatives to green fluorescent protein for imaging specific proteins in living cells. Here we introduce PRIME (PProbe Incorporation Mediated by Enzymes), a method for fluorescent labeling of peptide-fused recombinant proteins in living cells with high specificity. PRIME uses an engineered fluorophore ligase, which is derived from the natural *Escherichia coli* enzyme lipoic acid ligase (LplA). Through structure-guided mutagenesis, we created a mutant ligase capable of recognizing a 7-hydroxycoumarin substrate and catalyzing its covalent conjugation to a transposable 13-amino acid peptide called LAP (LplA Acceptor Peptide). We showed that this fluorophore ligation occurs in cells in 10 min and that it is highly specific for LAP fusion proteins over all endogenous mammalian proteins. By genetically targeting the PRIME ligase to specific subcellular compartments, we were able to selectively label spatially distinct subsets of proteins, such as the surface pool of neuexin and the nuclear pool of actin.

fluorescence microscopy | biotechnology | enzyme engineering

Techniques for posttranslational labeling of proteins in living cells address some of the shortcomings of green fluorescent protein (GFP) by expanding the repertoire of chemical probes available for protein visualization (1). However, most of these methods, such as HaloTag (2), SNAP/CLIP (3), and DHFR (4), still use large protein tags that sterically interfere with protein trafficking and function, as GFP is known to do (5, 6). FIAsh (7) is the only peptide-based posttranslational labeling method that currently works inside living cells. At 12 amino acids, the FIAsh tag is much smaller than GFP, but poor labeling specificity (7–9), cellular toxicity, and undesired palmitoylation (7) and oxidation (8) of the Cys₄ recognition motif limit its utility.

Here we introduce a method for protein labeling that utilizes a peptide tag while preserving high sequence specificity inside living cells. Our method, called PRIME (PProbe Incorporation Mediated by Enzymes), is based on a “fluorophore ligase” that is engineered from the *Escherichia coli* enzyme lipoic acid ligase (LplA). LplA's natural function is to ligate lipoic acid onto three *E. coli* proteins involved in oxidative metabolism (10). We previously used LplA for labeling of cell-surface proteins by demonstrating that the wild-type enzyme can ligate an azidoalkanoic acid instead of lipoic acid (11) (Fig. 1A, *Middle*). Ligated azide could then be chemoselectively derivatized using cyclooctyne-fluorophore conjugates.

In this work, we wished to extend LplA-mediated labeling to intracellular proteins but recognized the challenges associated with our two-step labeling scheme. First, labeling sensitivity is limited by the kinetics of strain-promoted [3 + 2] cycloaddition, which has a rate constant of $4.3 \times 10^{-3} \text{ M}^{-1} \text{ sec}^{-1}$ (12). Second, for intracellular applications, two washout steps would be needed: first to remove excess azide, and second to remove excess fluorophore. We sought to create a simpler, one-step fluorophore ligation strategy as an alternative.

Through LplA active site mutagenesis, we created a ligase capable of directly attaching the blue fluorophore 7-hydroxycoumarin to proteins fused to a 13-amino acid LplA recognition sequence (Fig. 1A, *Bottom*). We characterized the kinetics and specificity

of this fluorophore ligation in vitro. We then demonstrated coumarin PRIME labeling on different proteins in the cytosol and nucleus of various mammalian cell types. A side-by-side comparison with FIAsh showed that PRIME labeling is faster, more specific, and less toxic. We also used PRIME to label protein subpopulations in specific cellular compartments and to probe the biology of nuclear actin.

Results

Structure-Guided Mutagenesis and Screening for a Coumarin Fluorophore Ligase. We selected 7-hydroxycoumarin as our fluorophore for this study because its excitation and emission wavelengths are suitable for live-cell imaging (excitation 387–405 nm; emission 448 nm) (13), it is brighter than Enhanced Blue Fluorescent Protein (EBFP) [$\epsilon = 36,700 \text{ M}^{-1} \text{ cm}^{-1}$ and QY 0.7 for the anionic form (13), compared to $\epsilon = 31,500 \text{ M}^{-1} \text{ cm}^{-1}$ and QY 0.2 for EBFP (14)], and it is small and hydrophobic, which increases the likelihood that it will be accepted by LplA. We synthesized four 7-hydroxycoumarin structures (Fig. 1B) with varying linkers separating the fluorophore and the carboxylic acid that is activated by LplA to the adenylate ester.

Using a high-performance liquid chromatography (HPLC) readout of ligation onto E2p, a domain from LplA's natural protein substrate pyruvate dehydrogenase (15), we found that wild-type LplA, not surprisingly, does not use any of the coumarin substrates. We then examined the crystal structure of *E. coli* LplA bound to lipoic acid (16), and the related structure of *Thermoplasma acidophilum* LplA bound to lipoyl-AMP (17), to identify candidate positions for mutagenesis. Thirteen residues lie within 7.5 Å of the dithiolane moiety of lipoic acid in either LplA structure: N16, L17, V19, E20, F35, W37, S71, S72, H79, T87, R140, F147, H149 (*E. coli* numbering; Fig. S1A and B). We mutated each of these positions to alanine one by one and tested for uptake of the two smaller coumarin probes ($n = 3,4$). Only W37A-LplA gave detectable ligation product.

Fig. S1B shows that W37 is 8.5 Å from the dithiolane ring in the *E. coli* structure, whereas the analogous residue in *T. acidophilum* LplA, Y39, is much closer, at 3.9 Å (Fig. S1A). A new structure of *E. coli* LplA with lipoyl-AMP bound (18) instead of lipoic acid suggests that the enzyme undergoes a conformational change upon formation of the adenylate ester. Interestingly, in this new structure, the W37 position is much closer to

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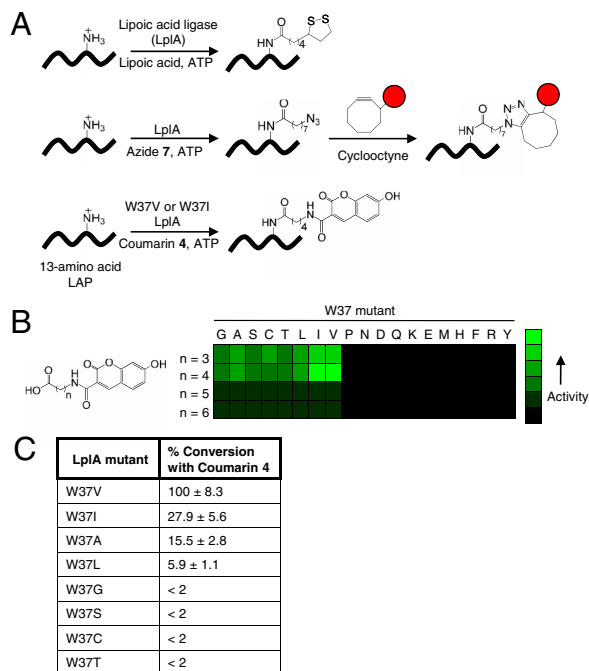


Fig. 1. Engineering a coumarin ligase. (A) Natural and engineered ligation reactions catalyzed by lipoic acid ligase (LplA). The middle row shows two-step probe targeting via alkyl azide 7 ligation followed by [3 + 2] cycloaddition (12), and the bottom row shows direct fluorophore ligation by an LplA mutant. LAP = LplA Acceptor Peptide (20). The red circle represents any probe. (B) Relative activities of W37 LplA mutants with 7-hydroxycoumarin derivatives. Assays were performed under two conditions: 12 hours with LplA's natural protein substrate E2p (15) or 40 min with LAP4.3D (20) (see *SI Methods*). No product was detected for the black-shaded quadrants. (C) Percent of LAP4.3D peptide converted to the coumarin ($n = 4$) conjugate for the eight most active W37 LplA mutants. Enzymes were used at 1 μ M and reaction times were 40 min. Values are normalized to the percent conversion obtained with the best mutant, W37V, which is set to 100%. Measurements were performed in triplicate. Errors, ± 1 s.d.

the dithiolane ring (3.6 Å, Fig. S1C). We note that our previous work to engineer an aryl azide ligase also identified W37 as the position most important (out of nine positions tested) for expanding LplA substrate specificity (19).

Encouraged by the coumarin ligation activity of ^{W37A}LplA, we prepared a panel of all possible W37 LplA mutants and screened

them against all four coumarin probes (Fig. 1B). We found that 11 mutants were inactive, although most of these were still able to ligate lipoic acid. Of the eight active mutants, little product was detected with the longer coumarin probes ($n = 5, 6$), whereas the most product was seen with coumarin 4 ($n = 4$). We therefore reassayed these eight mutants under milder conditions with coumarin 4 to differentiate among them. Instead of E2p, the HPLC assay was performed with an “LplA acceptor peptide” (LAP). Specifically, we used the yeast display-evolved sequence “LAP4.3D” (20) due to its superior chromatographic properties on HPLC when fused to the carrier protein HP1 (11).

Fig. 1C compares the amount of coumarin 4-LAP4.3D conjugate obtained with the eight active W37 mutants. Intermediate-sized hydrophobic sidechains—such as valine and isoleucine—were best able to complement the shape and size of coumarin 4. The preference for hydrophobic residues suggests that the ligase binds the neutral form of 7-hydroxycoumarin, rather than its anionic form ($pK_a = 7.5$) (13). This idea is supported by the observation that a fluorinated isostere of coumarin 4 called Pacific Blue, with a pK_a of 3.7 (13), is ligated by ^{W37V}LplA and ^{W37I}LplA with much lower efficiency than coumarin 4.

Coumarin Ligation Kinetics and Sequence Specificity. We proceeded to characterize the two most active coumarin ligases identified in our screen: ^{W37V}LplA and ^{W37I}LplA. First, HPLC assays were performed, this time using our most kinetically efficient LAP called “LAP2” (20). Controls in Fig. 2A show that coumarin ligation to LAP2 requires ATP and is not catalyzed by wild-type LplA. The HPLC product peaks from the reactions with both ligases were collected and analyzed by mass spectrometry, confirming the identity of the covalent coumarin 4-LAP2 adducts (Fig. S2).

Second, we used HPLC to measure the kinetics of coumarin ligation onto LAP2 (Fig. S3A). For ^{W37V}LplA, we obtained a k_{cat} of $0.019 \pm 0.004 \text{ sec}^{-1}$ and a K_m of $56 \pm 20 \mu\text{M}$. ^{W37I}LplA had a similar k_{cat} ($0.016 \pm 0.002 \text{ sec}^{-1}$) but a much higher K_m of $261 \pm 69 \mu\text{M}$. These values can be compared to previously determined kinetic constants for natural and unnatural LplA-catalyzed ligations (Fig. S3B). Whereas the k_{cat} of coumarin ligation is 10–30-fold poorer than previous k_{cat} values determined for lipoic acid (11, 15, 16), azidoalkanoic acid (11), and aryl azide (19), the coumarin K_m for ^{W37V}LplA is better than previous K_m values for unnatural substrates. In contrast, the coumarin K_m for ^{W37I}LplA is much poorer. For live-cell labeling, we were uncertain whether low coumarin K_m would confer an advantage, because intracel-

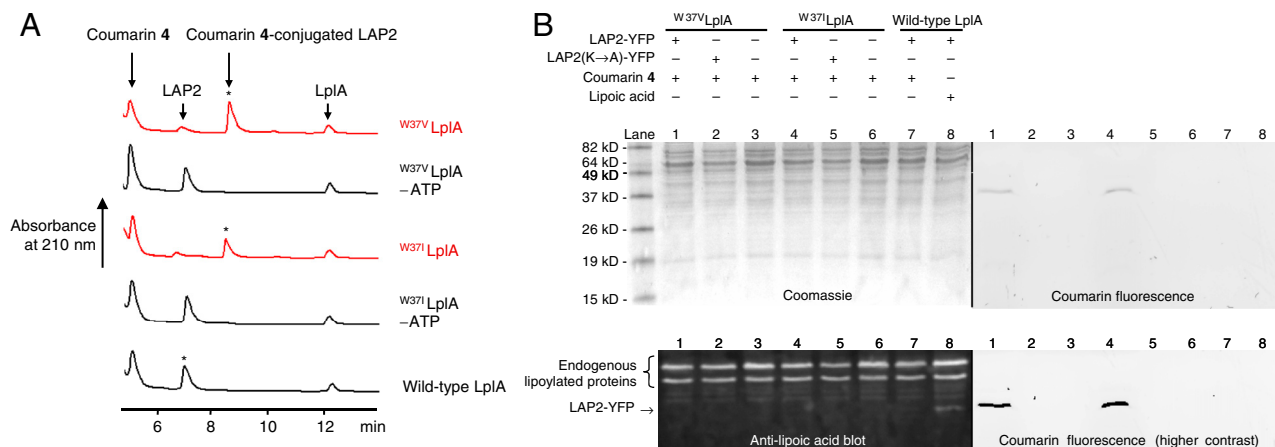


Fig. 2. In vitro characterization of coumarin ligase. (A) HPLC traces showing conversion of LAP2 peptide (20) to the coumarin 4 conjugate by ^{W37V}LplA or ^{W37I}LplA (red traces). Negative controls with wild-type LplA, or ATP omitted, are shown in black. Starred peaks were collected and analyzed by mass spectrometry (Fig. S2). (B) Coumarin labeling of mammalian cell lysate to determine sequence-specificity. Lanes 1 and 4 show coumarin 4 ligation to LAP2-YFP catalyzed by ^{W37V}LplA and ^{W37I}LplA, respectively. Negative controls are shown with wild-type LplA (lane 7), an alanine mutation in LAP2 (lanes 2 and 5), and untransfected lysate (lanes 3 and 6). The same samples were also analyzed by blotting with anti-lipoic acid antibody. Lysate in lane 8 was subjected to in vitro lipoylation with exogenous wild-type LplA.

lular coumarin concentrations following probe loading may exceed 300 μM (vide infra). We therefore tested both LplA mutants for coumarin ligation activity in living cells.

Third, we characterized the sequence specificity of both coumarin ligases. It was unclear if remodeling of the lipoic acid binding pocket would affect LplA's recognition of peptide and protein substrates. In addition, mammalian cells express protein substrates for their own mitochondrial lipoic acid ligase that might cross-react with *E. coli* LplA (15). To test for these possibilities, we performed coumarin labeling of LAP2-YFP (yellow fluorescent protein) in mammalian cell lysate and analyzed the samples by in-gel coumarin fluorescence imaging and anti-lipoic acid Western blotting. Fig. 2B shows that the endogenous protein substrates of mammalian lipoic acid ligase are present in the lysate (see anti-lipoic acid blot) but are not labeled by coumarin. In addition, no other endogenous proteins are labeled by coumarin, even though many are expressed at much higher levels than LAP2-YFP according to the Coomassie stain. It is possible that mitochondrial proteins are not labeled by coumarin because they are already saturated with lipoic acid. This idea is supported by the observation that in vitro lipoylation of lysate does not increase the lipoylation level of these endogenous proteins (lane 8 of Fig. 2B).

We also found that LAP2-YFP proteins were not detectably lipoylated, even though free lipoic acid may be present in the cytosol (lanes 1, 4, and 7). To confirm site-specificity, we mutated the central lysine in LAP2 to alanine and found that this abolished labeling by both W37V LplA and W37I LplA (lanes 2 and 5). Thus, both coumarin ligases retain the high sequence-specificity of wild-type LplA.

Characterization of PRIME Labeling Inside Living Mammalian Cells. To test coumarin ligation inside cells, we first addressed the issue of probe delivery. Coumarin 4 has 1–2 negative charges at physiolo-

gical pH, making it membrane-impermeant. We protected either the carboxylic acid alone or both the acid and the 7-hydroxyl group with acetoxymethyl (AM) groups (coumarin-AM₂; structure shown in Fig. 3A), which are known to be cleaved inside mammalian cells by endogenous esterases (21). Over time, excess anionic fluorophore is removed from cells via the action of organic anionic transporters (22). We found that one AM protecting group was not sufficient for membrane permeability; however, coumarin-AM₂ entered cells readily (Fig. S4A).

We used the “wedge method” (see *SI Methods*) to measure the intracellular concentration of coumarin probe immediately after loading. With 20 μM coumarin-AM₂, the intracellular coumarin concentration was $\sim 469 \mu\text{M}$ after 10 min of loading (Fig. S4B). This concentration exceeds the coumarin K_m for both W37V LplA and W37I LplA. Empirical optimization showed that coumarin probe could be thoroughly “washed out” of cells by replacing the media several times over 30 min (Fig. S4C).

Next, we evaluated LplA expression in mammalian cells. The wild-type *E. coli* LplA gene was found to express poorly in Human Embryonic Kidney (HEK) and HeLa cells. Fusions to fluorescent proteins helped, but we obtained the best results by resynthesizing the LplA gene with human-preferred codons. Humanized genes for both W37V LplA and W37I LplA gave robust and reproducible expression in a variety of cell lines, and their levels could be titrated by modulating the plasmid concentration used for liposome-mediated transfection (Fig. S5A).

Combining the optimized coumarin loading protocol, humanized LplA gene, and a LAP2-YFP fusion construct, we performed coumarin labeling with both W37V LplA and W37I LplA in live HEK cells (Fig. S5B). We observed specific labeling with both coumarin ligases, evident from comparing the coumarin signal for LAP2-YFP and LAP2(K \rightarrow A)-YFP. W37V LplA gave higher coumarin signal/background ratios on average, consistent with its superior

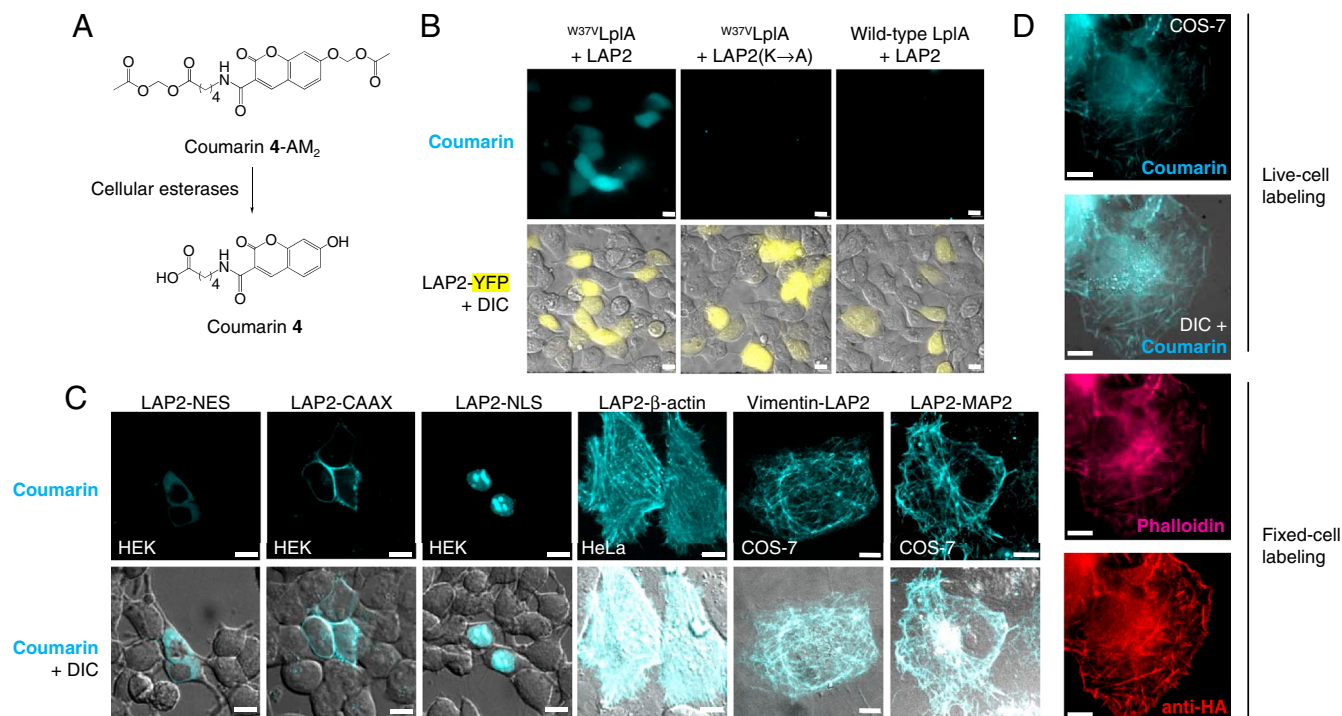


Fig. 3. Coumarin PRIME labeling in living mammalian cells. (A) Structure of membrane-permeant coumarin 4-AM₂ probe used for intracellular labeling. Reaction catalyzed by endogenous cellular esterases produces the parent coumarin 4 structure. (B) PRIME labeling of LAP2-YFP in HEK cells. Negative controls are shown with a Lys \rightarrow Ala mutation in LAP2, and with W37V LplA replaced by wild-type LplA. (C) Labeling of different LAP2 fusion proteins. NES = nuclear export sequence. CAAX = prenylation tag. NLS = nuclear localization sequence. MAP2 = microtubule-associated protein 2. (D) Comparison of methods for actin visualization. Coumarin PRIME labeling was performed on live cells expressing HA-LAP2-β-actin. After fixation, actin was detected by staining with phalloidin-Alexa Fluor 647 and anti-HA antibody. For B–D, coumarin labeling was performed for 10 min with W37V LplA, followed by 1-hour washout of excess probe. All scale bars, 10 μm . Epifluorescence images are shown in B and D; confocal images are shown in C.

k_{cat}/K_m compared to W37V LplA. However, when each ligase was overexpressed by using 200 ng plasmid instead of 20 ng (per 0.95 cm² of cells), some nonspecific labeling was seen with W37V LplA (Fig. S5 B and C). Analysis using targeted ligase constructs showed that this background correlated with W37V LplA localization and expression level. We hypothesize that W37V LplA may retain the coumarin-AMP ester intermediate within its active site, as the homologous enzyme biotin ligase is known to do with biotin-AMP ester (23). W37V LplA does not exhibit this behavior, likely due to its much higher K_m for coumarin.

From these observations, we conclude that W37V LplA is the best coumarin ligase for PRIME labeling, but W37V LplA is preferred when ligase overexpression cannot be avoided. We did not observe background labeling with W37V LplA under any experimental condition that we tested, including six-hour-long incubation with 20 μ M coumarin-AM₂.

PRIME labeling with W37V LplA was characterized in HEK cells along with negative controls (Fig. 3B). After 10-min incubation with probe, followed by 60-min washout, only cells expressing LAP2-YFP retained coumarin fluorescence, whereas neighboring untransfected cells were not labeled. Mutation of LAP2 or replacement of W37V LplA with wild-type LplA eliminated coumarin labeling. PRIME labeling was performed successfully in a variety of cell lines, including HEK, HeLa, and COS-7 (Fig. 3C).

We also tested labeling of various LAP2 fusion proteins (Fig. 3C). LAP2-YFP targeted to the nucleus, cytosol, or plasma membrane gave the expected coumarin labeling patterns. Similarly, LAP2 fusions to the cytoskeletal proteins β -actin and vimentin or to MAP2 (microtubule-associated protein 2) could be specifically labeled with coumarin. Fig. 3D shows a comparison of PRIME labeling of β -actin in live cells to actin visualization by phalloidin or antibody staining after cell fixation. The coumarin signal overlaps well with the other stains but has the benefit of compatibility with live cells. Actin can also be visualized via fusion to fluorescent proteins. Fig. S9A shows mApple-actin imaged in live COS-7 cells; here, actin is clearly excluded from the nucleus, an artifact that probably arises from the mApple tag interfering with nuclear import. We note that LAP2 was recognized by LplA at N-terminal, C-terminal, and internal fusion sites (see Table S1 for LAP2 placement in each construct).

To estimate the intracellular coumarin ligation yield, we used the “wedge method” (see SI Methods) to determine the single-cell concentrations of LAP2-YFP and attached coumarin probe after PRIME labeling. We found that $24 \pm 13\%$ of LAP2-YFP was labeled with coumarin after 10 min by W37V LplA (Fig. S6). The yield increased to $53 \pm 25\%$ after 1 h incubation with coumarin probe.

We also performed complete cellular characterization with W37V LplA, the less sensitive but more specific coumarin ligase. PRIME labeling with this mutant is shown with specificity controls, various LAP2 fusion proteins, and in various mammalian cell lines in Fig. S7A and B. Intracellular labeling yield is reported in Fig. S6.

Comparison of PRIME and FIAsh Labeling. We quantitatively analyzed the specificity, sensitivity, and toxicity of PRIME labeling in conjunction with a side-by-side comparison to FIAsh methodology. For the comparison, we prepared a single nuclear-targeted construct containing LAP2, mCherry, and the optimized 12-amino acid FIAsh recognition motif (7). In live HEK cells, PRIME labeling was performed first with W37V LplA for 10 min, followed by FIAsh for 30 min (Fig. S8A). To assess labeling specificity, we compared the nuclear coumarin or FIAsh signal (representing specific labeling, as the substrate is nuclear-localized) to the cytosolic signal (representing nonspecific labeling) in single cells. Fig. S8B shows that FIAsh gives more cytosolic background than PRIME. We also observed nonspecific FIAsh labeling in untransfected cells. A similar analysis of labeling sensitivity revealed that both methods can detect as little as 10 μ M target protein with signal/background ratio $>2:1$ (Fig. S8C). An assay for mitochon-

drial respiration showed that FIAsh is more toxic than PRIME (Fig. S8D).

Imaging Nuclear Actin with PRIME. We used PRIME methodology to explore the biology of nuclear actin. Cytosolic β -actin has a well-recognized role in controlling cell migration and morphology, but biochemical studies have also suggested several roles for β -actin in the nucleus, including involvement in transcriptional regulation (24) and chromatin remodeling (25). Studies of nuclear actin have been hampered by the lack of methods to noninvasively detect all forms of actin in living cells. For example, antibody detection and phalloidin staining require cell fixation. Fluorescent proteins disrupt nuclear entry of actin, as shown in Fig. S9A. Phalloidin and Lifeact (26) are specific for polymeric actin over monomeric actin. Rhodamine-labeled actin can be visualized by fluorescent speckle microscopy, but this requires invasive microinjection (27).

We tested PRIME for labeling of nuclear actin in living cells. Fig. S9A (Left) shows confocal images of coumarin-labeled LAP2- β -actin in COS-7 cells also expressing a nuclear marker (histone 2B fused to YFP, or H2B-YFP). Coumarin signal is clearly present in the nucleus, which contrasts with confocal images of mApple- β -actin (Fig. S9A, Right). Though actin was overexpressed in these experiments, the diffuse coumarin labeling pattern suggests that nuclear actin may exist as monomers or short oligomers, consistent with previous studies (28).

Previous work has shown that cell stress, such as heat shock, can increase the abundance of nuclear actin (29). We therefore imaged actin after 30–90 min of heat shock (43 °C) and found that coumarin-labeled actin accumulates in the nucleus within 30 min, whereas mApple-actin accumulation is much slower (Fig. S9A). In a small fraction ($<10\%$) of heat-shocked cells, we observed interesting coumarin-labeled nuclear actin structures (Fig. S9B), such as nucleolar clusters and filaments (which also stained with phalloidin, confirming F-actin morphology; Fig. S9C). This heterogeneity contrasted with the uniformly diffuse distribution of nuclear actin in nonstressed cells as well as mApple-actin in the nuclei of stressed cells. Perhaps filamentous actin, which has not previously been observed in the nuclei of living somatic cells, plays a role in transcriptional regulation (30) or nuclear stabilization (31).

The ability of PRIME methodology to detect nuclear pools of polymeric and nonpolymeric actin in living cells differentiates it from antibody, phalloidin, Lifeact, and fluorescent protein visualization techniques. PRIME may prove to be a useful tool for studying the biological function of nuclear actin.

Labeling Spatially Defined Protein Subpopulations Using Genetically Targeted Ligase. A unique feature of PRIME labeling is its ability to selectively highlight protein subpopulations via genetic targeting of the fluorophore ligase. To test this concept, we transfected HEK cells with both nuclear LAP2 and cytosolic LAP2. When cytosolic W37V LplA was coexpressed, we observed specific labeling of the cytosolic subpopulation of LAP2 (Fig. 4A, Left). Conversely, when nuclear-localized W37V LplA was coexpressed, we observed specific labeling of the nuclear subpopulation of LAP2 (Fig. 4A, Right). When enzyme and substrate were not colocalized, no coumarin labeling was observed. Similar results were obtained with genetically targeted W37V LplA (Fig. S7C).

We also tested this concept on neuexin-1 β , a transmembrane adhesion protein that functions in synapse development and plasticity (32). We prepared a LAP2-neuexin-LAP2 double fusion construct to provide access to labeling reagents on both sides of the membrane. Fig. 4B shows PRIME labeling of the cell surface neuexin population by either purified W37V LplA applied to the cell medium or W37V LplA targeted to the inner leaflet of the plasma membrane. In contrast, untargeted W37V LplA labels the total intracellular pool of LAP2-neuexin-LAP2, including LAP2 present on the cytosolic face of the endoplasmic reticulum (ER) and the Golgi apparatus. Nuclear-targeted W37V LplA, as

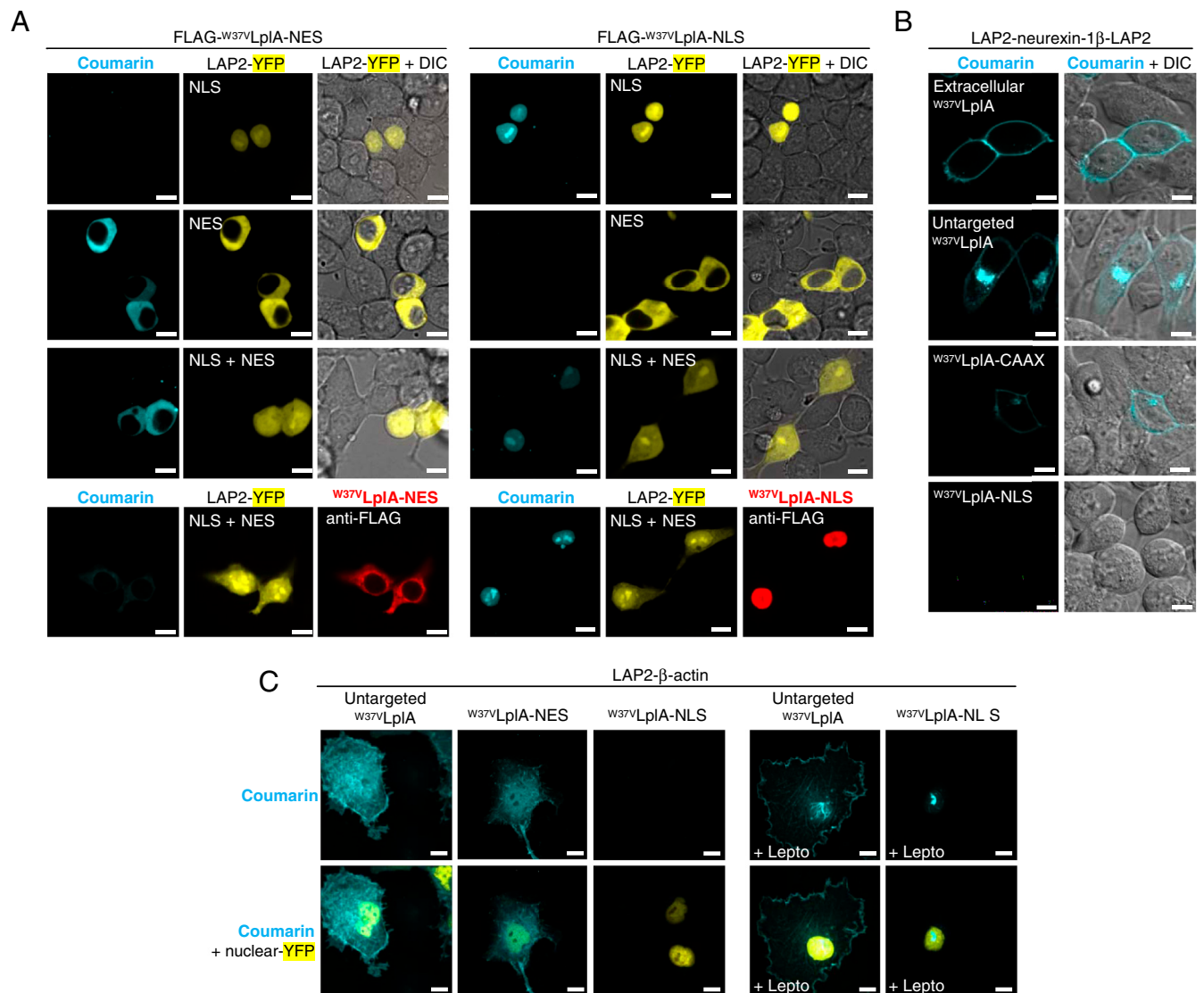


Fig. 4. Compartmentalized PRIME labeling with genetically targeted coumarin ligase. (A) Compartmentalized labeling of cytosolic or nuclear LAP2-YFP. HEK cells expressing both LAP2-YFP-NES and LAP2-YFP-NLS (third row), or either construct alone (first and second rows), were labeled with coumarin using nuclear-excluded (left) or nuclear-targeted (right) ligase. On the bottom, anti-FLAG staining of fixed cells shows the expression pattern of FLAG-^{W37V}LpIA. (B) Compartmentalized labeling of the membrane protein neurexin-1β. HEK cells expressing double-LAP2 tagged neurexin-1β were labeled with ^{W37V}LpIA added to the media (first row), expressed in the cytoplasm (second row), anchored to the inner plasma membrane (third row), or targeted to the nucleus (fourth row). (C) Compartmentalized labeling of actin. COS-7 cells expressing LAP2-β-actin and untargeted, nuclear-excluded, or nuclear-localized ^{W37V}LpIA were labeled with coumarin for 30 min. The right panels show labeling in the presence of 2 ng/ml leptomycin B (Lepto) to inhibit nuclear export. All images in A–C are confocal. Scale bars, 10 μm.

expected, does not label neurexin. Similar results were obtained using targeted forms of ^{W37V}LpIA (Fig. S7D).

We also experimented with ^{W37V}LpIA and ^{W37V}LpIA targeted to the ER using a KDEL sequence. We did not detect coumarin ligation onto LAP2-neurexin-LAP2 or a number of other LAP2 fusion proteins. This may be a consequence of high calcium concentration in the ER (33), which we found inhibits LpIA activity.

To provide further evidence for nuclear actin, we performed compartmentalized labeling of LAP2-β-actin using ^{W37V}LpIA-NLS (Fig. 4C). We initially did not detect any nuclear coumarin signal, perhaps because nuclear actin is bound within protein complexes and not sterically accessible to LpIA. The nuclear actin signal seen when labeling with untargeted ^{W37V}LpIA (Fig. S9A and Fig. 4C, Top) probably results from translocation of coumarin-labeled cytosolic actin to the nucleus, as suggested by a labeling experiment with ^{W37V}LpIA-NES (Fig. 4C).

Therefore, we repeated the experiment with ^{W37V}LpIA-NLS in the presence of leptomycin B to inhibit nuclear export of actin. In

this case, we observed nuclear labeling, including actin structures similar to those seen in the nuclei of heat-shocked cells (Fig. 4C). Leptomycin B may cause nuclear actin levels to rise, saturating actin-binding proteins and producing free actin that is accessible to nuclear LpIA.

Discussion

Using structure-guided mutagenesis of *E. coli* LpIA, we have engineered a coumarin fluorophore ligase for PRIME labeling in living cells. Compared to other protein labeling methods, PRIME uniquely offers the combination of a peptide tag, high labeling specificity, and compatibility with the interior of living mammalian cells.

Aside from FAsH, other peptide-based protein labeling methods are restricted to the cell surface either because the labeling reagents are not membrane-permeable [e.g., AcpS/Sfp (34) and sortase (35)] or because the chemistry is not orthogonal to intracellular functional groups [e.g., aldehyde tag (36) and ketone biotin (37)]. Unnatural amino acid mutagenesis stands apart as a

tagless protein labeling method. Though demonstrations of fluorophore incorporation in mammalian cells are promising (38), the prevalence of native amber stop codons in mammalian cells (38) and potential dominant negative effects from truncated protein by-products (39) could limit the utility of this method.

Several other protein labeling methods, like PRIME, use enzymes to catalyze probe conjugation to proteins (40). Here we explored genetic targeting of the labeling enzyme as a method to spatially restrict labeling to protein subpopulations of interest. With suitable sensitivity and labeling speeds, one might be able to use this strategy to study protein trafficking between subcellular compartments and organelles. The utility would be greatest for compartments that are too small to selectively illuminate with focused light, such as neuronal synapses.

Materials and Methods

In Vitro Coumarin Ligation. General reaction conditions: 100–150 μ M LAP or E2p substrate was incubated with 1–2 μ M LpIA, 500 μ M coumarin probe, 5 mM ATP, and 5 mM Mg(OAc)₂ in 25 mM Na₂HPO₄ pH 7.2 at 30 °C for various lengths of time. Reactions were quenched with EDTA (C₇ 100 mM) and analyzed on a Varian Prostar HPLC using a reverse-phase C18 Microsorb-MV 100 column (250 \times 4.6 mm). Chromatograms were recorded at 210 nm. For analysis of LAP2 peptide (GFEIDKVVYDLDA), we used a 10-min gradient of 30–60% acetonitrile in water with 0.1% trifluoroacetic acid at 1 ml/min flow rate. Percent conversions were calculated by dividing the product peak area by the sum of (product + starting material) peak areas.

Coumarin Labeling of Mammalian Cell Lysate. HEK cells expressing LAP2-YFP (or LAP2(K \rightarrow A)-YFP) were lysed under hypotonic conditions in 1 mM HEPES

pH 7.5 with 5 mM MgCl₂, 1 mM phenylmethylsulfonyl fluoride, and protease inhibitor cocktail (Calbiochem). After three cycles of freeze-thaw, cells were mixed by vortexing for 2 min. Lysate was cleared by centrifugation and stored in aliquots at –80 °C. To label with coumarin or lipoic acid, lysates were incubated with 200 nM LpIA, 500 μ M coumarin 4 or 250 μ M lipoic acid, 1.5 mM ATP, and 5 mM Mg(OAc)₂ in 25 mM Na₂HPO₄ pH 7.2. After overnight incubation at 30 °C, reactions were boiled in protein loading buffer containing 2-mercaptoethanol for 7 min, then separated on a 12% SDS/PAGE gel. Coumarin fluorescence was visualized on an Alpha Innotech Chemilmager 5500 instrument using 365 nm UV light for excitation.

General Protocol for Coumarin PRIME Labeling in Living Cells. HEK cells were transfected at ~70% confluency with expression plasmids for ^{w37V}LpIA (20 ng for a 0.95 cm² dish) and the LAP2 fusion protein of interest (400–600 ng) using Lipofectamine 2000 (Invitrogen). 8–24 h after transfection, cells were incubated with 20 μ M coumarin-AM₂ in serum-free DMEM (Dulbecco's modified Eagle medium, Cellgro) for 10 min at 37 °C. 0.1% w/v Pluronic F-127 (Invitrogen) can be optionally added to the labeling solution to give more even coumarin distribution in cells. The media was replaced 3–4 times over 30–60 min at 37 °C, with DMEM with 10% FBS (Fetal Bovine Serum), to wash out excess coumarin. Cells were imaged as described in *SI Methods*.

For further information on cloning, synthetic methods, cell culture, labeling, imaging, and other protocols, see *SI Methods*.

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