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Proximal Molecular Probe Transfer (PROMPT), a new approach for identifying sites of protein/ nucleic acid interaction in cells by correlated light and electron microscopy

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The binding and interaction of proteins with nucleic acids such as DNA and RNA constitutes a fundamental biochemical and biophysical process in all living organisms. Identifying and visualizing such temporal interactions in cells is key to understanding their function. To image sites of these events in cells across scales, we developed a method, named PROMPT for PROximal Molecular Probe Transfer, which is applicable to both light and correlative electron microscopy. This method relies on the transfer of a bound photosensitizer from a protein known to associate with specific nucleic acid sequence, allowing the marking of the binding site on DNA or RNA in fixed cells. The method produces a fluorescent mark at the site of their interaction, that can be made electron dense and reimaged at high resolution in the electron microscope. As proof of principle, we labeled in situ the interaction sites between the histone H2B and nuclear DNA. As an example of application for specific RNA localizations we labeled different nuclear and nucleolar fractions of the protein Fibrillarin to mark and locate where it associates with RNAs, also using electron tomography. While the current PROMPT method is designed for microscopy, with minimal variations, it can be potentially expanded to analytical techniques.

There are limited methods for visualizing the specific interaction of proteins-of-interest with DNA and RNA in cells by correlated light and electron microscopy (CLEM), that enable their identification, localization, and quantification of the bound fraction of protein to DNA/RNA. This is crucial for understanding the behavior of transcription factors, DNA or RNA polymerases, DNA repair enzymes, or mRNA editing enzymes. Potential applications include an ultrastructural snapshot of the interacting RNA binding proteins and mRNA in neuronal axons and dendritic spines that enable the spatially limited protein synthesis necessary for synaptic changes during neuronal plasticity. A key outcome and one goal of the project reported is to propel capabilities for localization of specific nucleic acid sequences without the inherent degradation of structure associated with methods now used for in situ hybridization-thus providing localizations of specific sequences in the context of the cell, but now with extremely well-preserved fine structure.

A plethora of methods is available for detecting and imaging protein-protein interactions using FRET between proteins tagged with fluorescent proteins^{1, 2}. Comparable methods for imaging protein-DNA/RNA interactions require fluorescent labeling of specific nucleotide sequences involved in their binding but the harsh conditions required for Fluorescent In Situ Hybridization (FISH) are incompatible with optimal cellular ultrastructure preservation for EM³. FP-tagging for proteins and for RNA or DNA can be challenging as overexpression masks the bound fraction⁴.

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The goal of this work was to develop and apply a new class of dynamic reporters to allow for correlated light and electron (CLEM) imaging of protein interaction with molecular species not amenable to genetically encoded tags, such as DNA, RNA, or lipids. This strategy builds upon our click-EM method for labeling such macromolecular structures in cells with photosensitizing fluorophores⁵. Metabolic substrates containing biorthogonal labels are incorporated into these macromolecules by endogenous enzymes^{6,7}. A click reaction labels these azido or alkyne groups with a fluorescent dye for imaging and can also generate a localized osmiophilic precipitate visible to EM by photosensitization of DAB.

This new extension of this method, which we call "PS-PROMPT", entails the chemical transfer of a photosensitizing fluorophore from a protein to a clickable metabolite that has been incorporated into cellular macromolecules or lipids by pulse chase. The requirement for close physical proximity of protein and macromolecule for photosensitizer transfer ensures only interacting fractions are so marked with untransferred removed before imaging. The PS-PROMPT probe has a modular structure composed of a protein-self labeling ligand e.g., a chloroalkane chain (HaloTag substrate, labeled HTL in Fig. 1a), a linker of variable length (zigzag line, L in Fig. 1a), a cleavable group such as disulfide bond (SS), a fluorescent photosensitizer dye (PS, Fig. 1a, b) and a click-ready group (e.g., N3 or azido).

An example of using PS-PROMPT to detect interaction of a specific protein to DNA is depicted in Fig. 1c. Cells expressing the DNA binding protein of interest fused to a HaloTag⁸ protein are incubated with EdU (5-ethynyl-2'-deoxyuridine) to incorporate alkyne groups in newly synthesized DNA⁹. Following fixation, the protein-HaloTag target is labeled with the PS-PROMPT probe. The probe is then covalently cross-linked by a click reaction⁹ of the azido group to an alkyne group in bound DNA that is within the reach of its flexible linker. Unclicked probe is removed by reductive cleavage of the disulfide bond by dithiothreitol (DTT) and subsequent washes. Thus, the fluorescent photosensitizer is transferred from the protein to nearby DNA revealing the sites of interaction by LM through its fluorescence and by EM after photosensitization of DAB precipitation and subsequent osmification. In proof-of-principle tests, this new PS-PROMPT probe system revealed the interaction sites on DNA of the histone H2B fused with a HaloTag. Application of the method to the RNA binding protein, fibrillarin revealed its ultrastructural localization when bound to RNA.

Results

Design and synthesis of PS-PROMPT

The PROMPT method requires the synthesis of a multimodal compound, generically named PS-PROMPT for PhotoSensitizer-PROMPT. The PS-PROMPT includes a chloroalkane group as a HaloTag ligand (HTL), an azido group (N3) compatible with click chemistry, a disulfide bond (SS) that can be cleaved by reduction, a linker of variable length (L), and a fluorescent photosensitizer (PS) (Fig. 1a). For application to correlative light and electron microscopy approaches, we chose the Janelia fluorophores, JF525 and JF570 as potent photosensitizers for DAB photooxidation¹⁰, as well as tetramethylrhodamine (TMR)¹¹(Fig. 1b). The fluorescence emission peaks of these dyes range from 549 to 599 nm (Fig. S1).

To test the functionality of each module of PS-PROMPT we first synthesized TMR-PROMPT (Fig. S2). 5⁽⁶⁾-carboxy-tetramethylrhodamine (TMR) was reacted with the 3-azidopropyl amide of S-trityl cysteine, prepared by reaction of Fmoc-NH-cys(S-Trityl)-OH and 3-azidopropyl-1-amine followed by removal of the N-terminal Fmoc. Cleavage of the trityl group and subsequent reaction of the thiol with the SPDP conjugate of HaloTag linker amine afforded TMR-PROMPT. JF570-PROMPT and JF525-PROMPT were synthesized (Fig. S3) by a similar procedure to TMR-PROMPT except the SPDP-HaloTag linker conjugate was reacted before final addition of the photosensitizer. Reaction of JF570-PROMPT and JF525-PROMPT with purified HaloTag protein in vitro was confirmed by LC–MS, giving the expected mass increase for the labeled protein.

Validating the PS-PROMPT method

We first tested whether TMR-PROMPT can bind to HaloTag in tissue culture cells. For this, we expressed the histone H2B fused with HaloTag in U2OS cells and incubated them with TMR-HaloTag ligand (TMR-HTL) or TMR-PROMPT (Fig. 2a). TMR-HTL binds efficiently to HaloTag-H2B whether the incubation is on live cells or following aldehyde fixation, whereas TMR-PROMPT labels its target only when added after fixation. Because of this result and the potential for premature cleavage of the disulfide group by intracellular cysteines and glutathione, any PROMPT assay was performed following aldehyde fixation. The next set of experiments challenged the capability of dithiothreitol (DTT) to reduce the bound TMR-PROMPT disulfide bond and release the TMR fluorophore (Fig. 2b). Fixed U2OS cells transfected with HaloTag-H2B were incubated with TMR-HTL or TMR-PROMPT, then treated with 10 mM DTT and heavily washed. With or without DTT, cells incubated with TMR-HTL display a constant TMR fluorescent signal, verifying that DTT does not affect the fluorescent properties of TMR. However, cells treated with TMR-PROMPT and reduced with DTT, completely lose the nuclear TMR fluorescence, presumably by reductive cleavage of TMR-PROMPT and TMR release from HaloTag-H2B. Lastly, we verified the functionality of the azido group of TMR-PROMPT by performing Cu(I)-catalyzed azide-alkyne cyclization (CuAAC) click chemistry with TMR-PROMPT on EdU treated cells (Fig. 2c). Only when cells are labeled with EdU, is TMR fluorescence observed in the nucleus, and as expected, subsequent treatment with DTT on clicked TMR-PROMPT does not trigger TMR release.

Next, we tested the feasibility of the PROMPT approach for CLEM using again as a model system the interaction between histone H2B with DNA. For CLEM, we used JF570-PROMPT, as JF570 is a potent photosensitizer¹⁰. For the JF570-PROMPT procedure (Fig. 3a), following overnight incubation with 5μ M EdU, cells transfected with HaloTag-H2B are fixed with 4% paraformaldehyde (PFA) and 0.1% glutaraldehyde (GA) and briefly incubated with JF570-PROMPT in the presence of saponin to facilitate probe penetration. After washing, we verified that only HaloTag-H2B expressing cells display JF570 fluorescence in the nucleus, thereby



Figure 1. Principal components of a PS-PROMPT probe and the PROMPT procedure for detecting protein-DNA binding. (a) Chemical structure and schematic of PS-PROMPT modules. N3: azido group; PS: Photosensitizers; SS: disulfide bond; HTL: HaloTag ligand, i.e., chloroalkane, a: methylene or polyethyleneglycol chain (L) of variable length. (b) Chemical structures of the fluorescent photosensitizers in PROMPT probes. R: site of attachment in PROMPT probe. (c) PROMPT Procedure overview for detection of binding between a protein of interest (POI) and DNA.

demonstrating the covalent bond formation between JF570-PROMPT and HaloTag-H2B (Fig. 3b, top panel, Fig. S4). Next, applying CuAAC click chemistry resulted in the reaction of the JF570-PROMPT immobilized on HaloTag-H2B with the nearest clickable substrate, an alkyne-uridinyl residue incorporated in DNA. After washing to remove the click chemistry solution and terminate the reaction, we treated the cells with DTT to cleave the disulfide link between JF570 and HaloTag-H2B. After intense washing to remove unclicked but cleaved JF570-PROMPT, we imaged the JF570 fluorescence (Fig. 3b, bottom panel). Gratifyingly, only in the condition where HaloTag-H2B is expressed in EdU-treated cells, do the majority of cells display a JF570 fluorescent signal



Figure 2. Testing functionality of the PROMPT probe modules. (a) Confocal micrographs of U2OS cells transfected with HaloTag-H2B construct and incubated with 1 μ M of TMR-HaloTag ligand (TMR-HTL) or 1 μ M of TMR-PROMPT live for 1H or overnight at 4 °C after 4% PFA fixation. (b) Confocal micrographs of U2OS cells transfected with HaloTag-H2B construct and incubated overnight with 1 μ M of TMR-HTL or 1 μ M of TMR-PROMPT after 4% PFA fixation. Cells are subsequently treated with or without 10 mM DTT. (c) Confocal micrographs of untransfected U2OS cells pretreated overnight with or without 5 μ M EdU. Following fixation in 4% PFA, the cells were labeled with TMR-PROMPT as above, covalently linked to incorporated EdU in DNA by click chemistry, and cells then treated with or without DTT as in B.

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Figure 3. PROMPT protocol. (a) Detailed schematic of the PROMPT protocol. (b) Confocal micrographs of U2OS cells transfected with or without the HaloTag-H2B construct, pretreated overnight with or without 5 μM of EdU, and processed with the PROMPT protocol. In the top panel, the imaging is performed after JF570-PROMPT incubation and washing. In the bottom panel, the imaging is performed after the complete procedure, i.e., after DTT treatment and washing. See Fig. S4 for images adjusted to brighten the dimmer cells. (c) Graph bars plot the average percentage of cells positive for JF570-PROMPT fluorescent signal after PROMPT protocol as in 2b bottom panel. No. of cells counted is 97 (for + EDU, – HaloTagH2B), 55 (for + EDU, + HaloTagH2B), 74 (for – EDU, -HaloTagH2B), and 31 (for – EDU, – HaloTagH2B). (d) Quantification of the mean JF570-PROMPT fluorescence signal per nucleus in cells expressing HaloTag-H2B and pretreated with EdU as in 2b. Imaging is performed after the JF570-PROMPT incubation or after the complete PROMPT procedure. No. of cells counted is 41 (after Halotag binding) and 37 (after PROMPT procedure). Error bars are box-and-whiskers plots containing the mean (X), quartiles (box), and minimum and maximum observations (whiskers).

above background in the nucleus (Fig. 3c). Interestingly, when comparing the JF570 fluorescence before the click chemistry and after the DTT treatment, we observed a significant reduction (5.2-fold) in mean intensity, suggesting that only a fraction of the JF570 bound to HaloTag-H2B is transferred to EdU (Fig. 3d).

PROMPT dependence on molecular distances in cellular ultrastructure

Considering the structural constraints on the PS transfer success rate, we reasoned that elongating the linker chain of PS-PROMPT (L in Fig. 1a) would increase its efficiency. We therefore added a PEG4 linker to TMR-PROMPT (Fig. 4a) and tested the TMR transfer efficiency. In cells expressing HaloTag-H2B and labeled with EdU, the TMR fluorescence intensity is significantly greater after PROMPT when using TMR-PEG4-PROMPT (Fig. 4b) compared to TMR-PROMPT. Elongating the probe not only compensates for unfavorable orientations of partners but also increases the radius in which the PS-PROMPT can find a clickable target.

Visualizing Histone H2B-DNA interaction in U2OS cells with PROMPT and CLEM

Next, we performed photooxidation of DAB through the illumination of the JF570-PROMPT after the PROMPT procedure in cells expressing HaloTag-H2B and labeled with EdU (Fig. 5a). After osmification, embedding, and sectioning, we verified the overall darkening of the correlated nucleus by electron microscopy. At higher magnification, we visualized a high concentration of puncta that were more opaque to electrons. To gain confidence regarding the identity of the puncta, we performed multi-tilt axis tomography, combining 1936 images of the same field of view taken at multiple angles¹². We obtained a volume of the isometric pixel size of 1.4 nm (Fig. 5b, Supplementary movie 1). The size and shape of some of the electron-dense puncta (highlighted by the arrowheads) are consistent with the expected nucleosomes. These show substructures that possibly represent the DNA wound around the nucleosome structure where it contacts the Halo-H2B dimers or mixed Halo-H2B/H2B dimers but additional future studies would be required to demonstrate this speculated specific highlighting.

Interestingly, whereas some chromatin regions are enriched with these dark puncta, others are devoid of those (pink area, Fig. 5b). The lack of DAB precipitates indicates the failure of JF570-PROMPT transfer. The dilution of HaloTag-H2B by endogenous H2B, partial replacement of thymidines by EdU, and non-favorable orientations of the HaloTag active pocket and the clickable nucleotide bases within its reach could explain the inefficient transfer.

Visualizing fibrillarin-RNA interaction in U2Os cells with PS-PROMPT and CLEM

As we have demonstrated the feasibility of PROMPT to pinpoint protein-DNA binding partners by CLEM, replacing EdU with 5-Ethynyl Uridine (5EU) should make PROMPT amendable to the study of protein-RNA interaction. As a study system, we tested PROMPT on the interaction of fibrillarin with RNAs. As a component of the C/D box small nucleolar ribonucleoproteins, fibrillarin is involved in the 2'-O-methylation of rRNAs^{13, 14}. We transfected U2OS cells with HaloTag-Fibrillarin and labeled overnight newly synthesized RNAs with 5EU. After PROMPT using JF570-PROMPT, the JF570 fluorescent signal is only detected in cells treated with both HaloTag-Fibrillarin and 5EU (Fig. 6a, b). Similar observations were made using JF525-PROMPT (Fig. S5) where JF570 has been replaced with the yellow photosensitizer, JF525. Completing the protocol with DAB photooxidation by JF570, the extent of DAB polymerization was dependent on the JF570 fluorescence intensity as expected (Fig. 6c). After correlation, we imaged at high magnification the nucleolus and surroundings by EM (Fig. 6d,



Figure 4. PROMPT for distance estimation. (a) U2OS cells expressing HaloTag-H2B and pretreated with 5 μ M EdU are processed with PROMPT using TMR-PROMPT probe or TMR-PEG4-PROMPT. A higher fluorescence signal is observed by confocal for TMR-PEG4-PROMPT compared to TMR-PROMPT. (b) Quantification of the mean TMR fluorescence signal per nucleus in cells treated as 4a. No. of counted cells is 67 (for TMR-PROMPT) and 82 (for TMR-PEG4-PROMPT). Error bars are box-and-whiskers plots containing the mean (X), quartiles (box), and minimum and maximum observations (whiskers).

b





Figure 5. CLEM of histone H2B binding to DNA using PROMPT. (**a**) The top left panel shows confocal, and DIC imaging (middle left) of EdU treated HaloTag-H2B expressing U2OS cells processed with JF570-PROMPT. The lower left image shows the amplitude of the DAB photooxidation by JF570-PROMPT. The center image shows an EM micrograph of the cell's nucleus framed in the pink box in the left panels. The image on the right shows the enlargement of the area bounded in the white box in the center image. (**b**) Both panels display an electron tomogram slice from a cell treated and photooxidized as in 5a. The image on the right is a close-up view of the area contained in the yellow box on the left panel. The pink shaded area highlights a region lacking the electron-dense puncta. For (**a**) and (**b**), arrowheads point at some of the electron-dense particles.

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e). As expected, we observed a high density of dark particles in the nucleolus (cyan dotted line boundaries) and at the Cajal bodies (purple dotted line boundaries) tentatively identified by their size, localization, and dense coiled fibre ultrastructure (Fig. S6). Interestingly, we also detected evidence of fibrillarin transferring PROMPT to RNAs at discrete puncta of a few nanometers throughout the nucleoplasm (Fig. 6e, arrowheads) and in 200–250 nm granules (Fig. 6d, white arrows) of unknown identity. Electron tomography not only confirms the preponderance of fluorophore transfer sites in the nucleolus and the Cajal body but also highlights Fibrillarin and RNA interaction clusters at the surface of these organelles (Fig. 7, Supplementary movie 2, Fig. S6). Furthermore, the 3D distribution of the fibrillarin-RNA complexes throughout the nucleus is made visible. In summary, the



Figure 6. CLEM of Fibrillarin binding to RNA using PROMPT. (a) Confocal micrographs of U2OS cells transfected with or without the HaloTag-Fibrillarin construct, pretreated overnight with or without 1 mM 5EU, and processed with the PROMPT procedure. (b) Quantification of the mean JF570-PROMPT fluorescence signal per nucleus in cells treated as in 6A. Error bars are box-and-whiskers plots containing the mean (X), quartiles (box), and minimum and maximum observations (whiskers). (c) DAB photooxidation by JF570-PROMPT after the PROMPT procedure in U2OS cell transfected with HaloTag-Fibrillarin construct and treated with 1 mM 5EU. The left micrographs show the fluorescence and the brightfield images pre and post photooxidation. The image on the right shows the electron microscopy micrograph of the cells shown on the left and framed in the pink box. The asterisk points at a speck of dust on the resin section. (d) Electron microscopy micrograph of the area of interest in the black box in (c). The top image shows the original acquisition, and the bottom one is a composite image including the original TEM micrograph layered with the DAB polymers extracted by thresholding and colorized in yellow. The purple dotted lines circle the Cajal bodies. The white arrows point at 250 nm diameter granules of unknown identities. The cyan dotted line surrounds darker areas of the nucleolus. No, Nucleolus; Np, Nucleoplasm; Cb, Cajal body. (e) Electron microscopy micrograph of the area of interest in the orange box in (\mathbf{d}) . Arrowheads point at some of the electron-dense DAB polymers. The dotted white line separates the nucleolus on the right from the nucleoplasm on the left.



Figure 7. Electron Tomography of Fibrillarin/RNA PROMPT. (**a**) Confocal micrographs of U2OS cells transfected with the HaloTag-Fibrillarin construct, pretreated overnight with 1 mM 5EU, and processed with the PROMPT procedure. Brightfield images are acquired before and after pho-oxidation (PO) of DAB by JF570-PROMPT. The lowest image shows the corresponding photo-oxidized cell by transmission electron microscopy. The orange box delineates the area acquired by electron tomography in (**b**). (**b**) Both panels display a minimum projection of 8 consecutive electron tomogram slices from the cell treated and photo-oxidized in 7A. In the lowest image, the most electron dense areas have been segmented by thresholding and colorized in orange. No, Nucleolus; Cb, Cajal Body.

PROMPT method confirmed the known interaction of fibrillarin with rRNAs for their methylation and revealed potential functions in the nucleoplasm.

Discussion

In this study, we developed PROMPT, a new class of molecular probes and method for the visualization of the interaction of proteins with DNA or RNA demonstrating the detection by correlated light and electron microscopy of the binding of DNA with the histone H2B and RNAs with fibrillarin. The chains of nucleosomes visualized using PROMPT for histone H2B and DNA interaction validates the specificity and high-resolution images obtainable with the method. In addition to verifying the known nuclear localization of fibrillarin, PROMPT reveals its interactions with RNAs, presumably rRNAs and small nucleolar RNAs¹⁵. In addition, the method reveals a previously undescribed nuclear but non-nucleolar fraction of fibrillarin is also associated with RNAs upon ribosome maturation, on its entry to or exit from the nucleolus.

The PROMPT procedure itself is relatively simple and, except for the probe, does not require specialized chemicals or equipment. It can potentially be applied to visualize the interaction of any HaloTag fusion protein with cellular components that can incorporate clickable metabolites such as proteins, nucleotides, sugars, lipids, enzymatic inhibitors, etc. The method is amenable for spatial-temporal studies using pulse-chase to detect lifetimes of HaloTag fusion proteins interaction with newly synthesized macromolecules. Conversely, to measure binding of new HaloTag fusion proteins by pulse chase would require blocking old protein with a conventional HaloTag ligand, followed by its omission for a defined time before fixation. Because the PS-PROMPT probe is

not incorporated in live cells, only newly synthesized proteins would be accessible to post-fixation labeling by the current PS-PROMPT probe.

The inability of PS-PROMPT to label live cells probably reflects a low membrane permeability that could be improved by replacing the polar amide bonds with more hydrophobic groups. Additionally, to prevent reduction of the disulfide bond of the PS-PROMPT probe in the reducing environment of the cytoplasm, replacement with a photocleavable linker¹⁶ or periodate-cleavable vicinal diol-containing linker¹⁷ would be required. One potential indication of this phenomenon is the residual fluorescence in the oxidizing environment of the endoplasmic reticulum on live labeling with TMR-PROMPT (Fig. 2a) that may reflect reaction of its cleaved disulfide group¹⁸ with endogenous ER thiols, and therefore the non-specific anchoring of the fluorophore. The PS-PROMPT cleavage system was inspired by the cleavable SNAP ligands used to study the dynamics of endocytosis¹⁹. In this context, the impermeable probe is always on the luminal side of the cell and therefore not sensitive to cytoplasmic reduction.

The partial lack of transfer of the fluorophores between H2B and DNA, while potentially revealing some biological phenomena, may also highlight certain limitations of the method. These limitations are the coexistence of HaloTag-H2B and endogenous H2B, the partial replacement of thymidines by EdU²⁰, and steric and distance constraints in transfer reaction. Incomplete HaloTag binding and click reaction may also be at play here. Alternative methods for the click chemistry might improve the transfer efficiency²¹. The PROMPT method does not probe the specific native molecular interactions between the protein and nucleic acid directly but depends on fusion to HaloTag and an appropriate linker length to mark binding. It therefore adds limitations on successful ligation with uncertainties dependent on the site of HaloTag attachment, its orientation and mobility on fixation, and ability of PROMPT probe to fully access and react within the surrounding space. Alternative methods for probing macromolecular interactions such as FRET or proximity labeling have different limitations, and do not give EM readout.

The choice of the protein fused to HaloTag could facilitate the identification and subcellular localization of specific DNA or mRNA sequences in cells. In combination with dCas9-HaloTag with a specific guide RNA⁴, one could target only the binding fraction of dCas9 to the chosen DNA sequences. While relatively weak, the signal should be detectable by removal of the excess, unbound fraction of labeled dCas9-HaloTag that loses fluorescence by the PROMPT method. Similar approaches can be applied to mRNA using MS2 protein fused with HaloTag²². The PROMPT method should reduce this background signal and highlight only the mRNA fused with the MS2 sequences and bound to the MS2 protein.

The PROMPT probe is composed of different functional modules, and the modification of one or several modules broadens the range of potential applications of the method. Replacement of the HaloTag ligand module by SNAP or CLIP ligands (benzyl guanine and benzyl cytosine respectively)²³ expands the spectrum of protein targeting systems and enables simultaneous PROMPT for multiple proteins. Replacing alkyne groups incorporated with metabolic intermediates such as EdU with SNAP or CLIP ligands containing alkynes, would potentially allow CLEM applications of the PROMPT method for protein-protein interaction. Probably the most exciting applications of PROMPT may come from the replacement of the module that is being transferred. For instance, the photosensitizer can be replaced by a wide range of fluorophores, including the ones compatible with super-resolution, such as JF549 or JF646²⁴. An interesting modification would be to add an additional fluorophore (of non-overlapping spectral properties or part of a FRET donor-acceptor pair) to the HaloTag side of the disulfide bond, so it is retained when PROMPT transfer does not occur. In this way, one could potentially determine both the total protein pool with this fluorophore and the bound fraction with the transferred fluorophore. By incorporating a biotin group, PROMPT could also enable pulldown of binding partners for sequencing and/or mass spectrometry, in addition to imaging their interaction. Finally, PROMPT can potentially serve as a molecular ruler and highlight structural changes of the protein in situ that positively or negatively affect the transfer rate.

Material and methods

Cell culture, 5EU and EdU labeling, transfection

U2OS cells were cultured on 35 mm MatTek dishes (MatTek Corp) in DMEM supplemented with 10% FBS at 5% CO2. On reaching 70% confluency and two days before the experiment, cells are transfected with 0.5 μ g of DNA with lipofectamine 3000 (ThermoFisher Scientific) following the manufacturer protocol. If needed, cells are incubated overnight in culture medium with 5 μ M of 5-ethynyl 2'-deoxyuridine (EdU, #1149, Click Chemistry Tools) diluted from a 10 mM stock in DMSO. For 5-EU (5-ethynyluridine) incorporation, 200 mM 5-EU (#1261, Click Chemistry Tools) in DMSO is diluted to 1 mM in culture medium and incubated with cells overnight.

DNA constructs

For the construct of HaloTag-H2B (JH1348), the H2B sequence was amplified from pminiSOG-H2B-6²⁵ and substituted EGFP in pHaloTag-EGFP (addgene #86629)²⁶ through the In-Fusion cloning method. For the construct of HaloTag-Fibrillarin (JH1239), the Fibrillarin sequence was amplified from pEGFP-C1-Fibrillarin (addgene #26673)²⁷ and substituted EGFP in pHaloTag-EGFP (addgene #86629) through the In-Fusion cloning method.

Synthesis of TMR-PROMPT and TMR-PEG4-PROMPT (Fig. S2)

NH_2 -cys(S-Trt)-CO-NH-(CH₂)₃-N₃ (2)

Fmoc-NH-cys(S-Trt)-OH, (1) (19.8 mg, 33.8 μ mol) and HATU (14.1 mg, 37.2 μ mol) were dissolved in dry DMF (100 μ L) in a plastic screw cap tube and 3-azidopropyl-1-amine (3.7 μ L, 37.2 μ mol, Click Chemistry Tools) followed by DIEA (13 μ L, 74.4 μ mol) were added with mixing. The reaction mix turned yellow and

LC–MS revealed complete reaction in 30 min; ES-MS (m/z) $[M + Na]^+$ for $C_{40}H_{37}N_5NaO_3S$, 690.25; found 689.3. Piperidine (20 µL, 0.2 mmol) was added and the solution evaporated under high vacuum after 1 h, dissolved in DMSO, separated by RP-HPLC and lyophilized to give **2** as a white solid. Yield, 14.3 mg, 76%. ES-MS (m/z) $[M]^+$, $[M + Na]^+$ for $C_{25}H_{27}N_5OS$, 446.2, 468.2; found 446.1, 468.1.

5(6)-TMR-CONH-cys(SH)-CO-NH-(CH₂)₃-N₃ (4)

5(6)-Carboxytetramethylrhodamine, 5(6)-TMR-CO₂H (1.35 mg, 3.14 µmol, Novabiochem) and TSTU (1.3 mg, 4.4 µmol) were dissolved in dry DMSO (25 µL) with TEA (0.96 µL, 6.9 µmol) and kept at room temperature. Reaction was complete in 30 min (by LC–MS) and then added to a solution of NH₂-cys(S-Trt)-CO–NH-(CH₂)₃-N₃ (2) (2.0 mg, 3.6 µmol) in DMSO (10 µL) with NMM (1 µL, 9.1 µmol) and kept at room temperature overnight when LC–MS revealed complete reaction. After acidification with HOAc (2 µL), the desired product, **3** was isolated by RP-HPLC and lyophilized to a red solid. Yield, 2.0 mg (74%) ES-MS (m/z) [M]⁺ for C₅₀H₄₈N₇O₅S, 858.3; found 858.3. The trityl group was removed by dissolving the product (1.89 mg, 2.2 µmol) in TFA:H₂O :Triisopropylsilane:Ethanedithiol (92.5:2.5:2.5 v/v, 0.5 mL) for 30 min, evaporation under high vacuum, purification by RP-HPLC and lyophilization to give **4** as a red solid. Yield, 0.9 mg (66%) ES-MS (m/z) [M]⁺ for C₃₁H₃₄N₇O₅S, 616.2; found 616.1.

SPDP-HaloTag linker

HaloTag linker amine (²⁸ (3.0 mg, 13.5 μ mol) and SPDP (4.7 mg, 15 μ mol) were dissolved in dry DMSO (50 μ L) and NMM (3.3 μ L, 30 μ mol) added. LC–MS revealed reaction was complete after overnight when the reaction mixture was neutralized with HOAc (5 μ L) and the product was purified by RP-HLPC (and lyophilized to a colorless oil. Yield, 3 mg (54%) ES-MS (m/z) [M]⁺ for C₁₈H₃₀ClN₂O₃S₂, 421.1; found 421.1.

TMR-PROMPT; 5(6)-TMR-CONH-cys(S-S-(CH₂)₂CONH-HaloTag ligand)-CO-NH-(CH₂)₃-N₃ (5)

A solution of 5(6)-TMR-CONH-cys(SH)-CO-NH-(CH₂)₃-N₃ (**4**) in DMSO (100 μ L, 6.25 mM measured by absorbance in 0.1 M HCl in 95% ethanol using ϵ_{max} 95,000 M⁻¹ cm⁻¹ at 554 nm, 0.626 μ mol) was mixed with SPDP-HaloTag linker (20 μ L, 49 mM in dry DMSO, 0.98 μ mol) and NMM (1 μ L, 10 μ mol) added. After 1 h, HOAc (5 μ L) was added and the desired product, (**5**) purified by RP-HPLC to give a colorless oil. Yield, 0.42 mg (72%) ES-MS (m/z) [M]⁺ for C₄₄H₅₈ClN₈O₈S₂, 925.4; found 925.4.

SPDP-PEG4-HaloTag linker

Solutions of HaloTag linker amine²⁸ (3.0 mg, 4.5 μ mol,) in dry DMSO (90 μ L) and SPDP-PEG4-NHS (2.5 mg, 4.5 μ mol, Quanta Biodesign) in dry DMSO (90 μ L) were mixed and NMM (1 μ L, 10 μ mol) added. LC–MS revealed reaction was complete after 4 h and the product was used without further purification. ES-MS (m/z) [M]⁺ for C₂₉H₅₁ClN₃O₈S₂, 668.3; found 668.3.

TMR-PEG4-PROMPT; 5(6)-*TMR-CONH-cys*(S-S-(*CH*₂)₂*CONH-PEG4-HaloTag ligand*)-*CO-NH-*(*CH*₂)₃-*N*₃ (6) 5(6)-TMR-CONH-cys(SH)-CO-NH-(CH₂)₃-*N*₃ (4) (3 mL of 0.25 mM in DMSO; measured as above) was added to the solution of SPDP-PEG4-HaloTag Linker (15 μL of 50 mM, 0.75 μmol) and NMM (15 μL, 150 μmol) added. After 2 h, LC–MS indicated reaction was complete, HOAc (50 μL) was added, the product, (6) purified by RP-HPLC and lyophilized. Yield, 0.7 μmol (by absorbance in 0.1 M HCl in 95% ethanol using ε_{max} 95,000 M⁻¹ cm⁻¹ at 554 nm) after dissolving in dry DMSO (100 μL). ES-MS (m/z) [M]⁺ for C₅₅H₇₉ClN₉O₁₃S₂, 1172.5; found 1172.4.

Synthesis of JF525-PROMPT and JF590-PROMPT (Fig. S3)

$Fmoc-NH-cys(SH)-CO-NH-(CH_2)_3-N_3$ (7)

Fmoc-NH-cys(S-Trt)-OH (1) (59 mg, 100 μmol) and HATU (42 mg, 110 μmol) were dissolved in dry DMF (200 μL) in a plastic screw cap tube and 3-azidopropyl-1-amine (11 μL, 110 μmol, Click Chemistry Tools) followed by DIEA (38 μL, 220 μmol) were added with mixing. The reaction mix turned yellow and LC–MS revealed complete reaction in 30 min; ES-MS (m/z) $[M + Na]^+$ for C₄₀H₃₇N₅NaO₃S, 690.3; found 690.2. The reaction mixture was evaporated to yellow oil and trityl group removed by dissolving in TFA-H₂O-Triisopropylsilane-Ethanedithiol (92.5/2.5/2.5 v/v, 1 mL) and kept at room temperature for 2 h. Following evaporation to an oily solid, the desired product was purified by RP-HPLC, and lyophilized to give (7) a white solid. ES-MS (m/z) $[M]^+$ for C₂₁H₂₄N₅O₃S, 426.2; found 426.1.

NH₂-cys(S-S-(CH₂)₂CONH-HaloTag ligand)-CO-NH-(CH₂)₃-N₃

Fmoc-NH-cys(SH)-CO-NH-(CH₂)₃-N₃ (7) (1.25 μ mol, 50 μ L of 25 mM solution in dry DMSO) was mixed with SPDP-HaloTag linker (1.25 μ mol, 25 μ L of 50 mM solution in dry DMSO) and NMM (2.5 μ L, 25 μ mol) added. LC-MS revealed complete reaction after 20 min, ES-MS (m/z) [M]⁺ for C₃₄H₄₈ClN₆O₆S₂, 735.3; found 735.3. Piperidine (20 μ L) was added and the solution evaporated after 5 min. The residue was dissolved in DMSO and HOAc (5 μ L), product **8** isolated by RP-HPLC, lyophilized and dissolved in dry DMSO (75 μ L). ES-MS (m/z) [M]⁺ for C₁₉H₃₈ClN₆O₄S₂, 513.2; found 513.2.

JF570-PROMPT; Janelia Fluor 570-CONH-cys(*S-S-(CH₂)₂CONH-HaloTag ligand*)-*CO-NH-(CH₂)₃-N₃* (9) The product, **8** from the previous step was reacted with JF570-NHS ester (0.4 μ mol, 8 μ L of 50 mM solution in dry DMSO) and NMM (10 μ mol, 1 μ L) for 2 days at room temp, the product **9** isolated by RP-HPLC, lyophilized

and dissolved in DMSO (100 μL) to give a 1.4 mM solution, measured by ϵ_{max} 100,000 M^{-1} cm $^{-1}$ in 0.1 M HCl in 95% ethanol at 574 nm. ES-MS (m/z) $[M]^+$ for $C_{46}H_{58}ClN_8O_7S_3$, 965.3; found 965.3.

JF525-PROMPT; Janelia Fluor 525-CONH-cys(S-S-(CH₂)₂CONH-HaloTag ligand)-CO-NH-(CH₂)₃-N₃ (10) Prepared as for JF570-PROMPT. Measured by ε_{max} 80,000 M⁻¹ cm⁻¹ in 0.1 M HCl in 95% ethanol at 532 nm. ES-MS (m/z) [M]⁺ for C₄₆H₅₄ClF₄N₈O₈S₂⁺,1021.3; found 1021.3.

Reaction with HaloTag protein

JF525-PROMPT or JF570-PROMPT ($0.5 \ \mu$ L of 1 mM solution in DMSO) was added to HaloTag protein ($2.5 \ \mu$ L of 100 μ M solution in PBS) diluted in 100 mM Na·MOPS pH 7.2, and kept at room temperature for 2 h. Acetic acid (1 μ L) was added and analyzed by LC–MS using a PLRP-S 1000A column (8 μ m, 50×2.1 mm, Agilent) eluting with linear 20–60% ACN-water with constant TFA (0.05%) in 16 min.

JF570-PROMPT: ES-MS (m/z) $[M]^+$ for adduct with HaloTag protein (loss of HCl), $C_{46}H_{57}N_8O_7S_3$, (965.3–35.98) 929.3; for deconvoluted masses of JF570-PROMPT:HTP complex, 35,729.0 minus HTP, 34,798.6; found 930.4.

JF525-PROMPT: ES-MS (m/z) $[M]^+$ for adduct with HaloTag protein (loss of HCl), $C_{46}H_{53}F_4N_8O_8S_2$, 985.3; for deconvoluted masses of JF525-PROMPT:HTP, 35,783.5 minus HTP, 34,798.6, found 984.9.

PROMPT method

After EdU or 5EU labeling, transfected cells are washed with culture medium and fixed for 2 min with room temperature fixative (4% EM grade paraformaldehyde (Electron Microscopy Sciences #19202) + 0.1% EM grade glutaraldehyde (Ted Pella, #18426) in 0.1 M Hepes (Sigma-Aldrich #H3375) pH7.4 + 2 mM CaCl₂) followed by a 1-h incubation with 4 °C fixative. Next, the cells are washed three times with PBS and incubated for 10 min with PBS + 10 mM glycine. The cells are then incubated for 30 min with 200 nM of PROMPT probe in PBS + 0.1% BSA (Sigma-Aldrich #A8022) + 0.1% saponin (Sigma-Aldrich #S4521), then washed five times with PBS + 0.1%BSA + 0.1%saponin, and five times with PBS. For the click chemistry step, the cells are incubated for 2 times 30 min with 50 mM Hepes pH7.4 + 100 mM NaCl + 2 mM CuSO₄ + 10 mM of freshly prepared sodium ascorbate (Sigma-Aldrich #A7631). The reaction is terminated with ten washes of PBS. For the disulfide bond reduction, cells are incubated three times 10 min with freshly prepared 10 mM DTT (Sigma-Aldrich #D9779) in PBS. Released photosensitizers are removed with five washes of PBS + 0.1%BSA + 0.1%saponin and five washes with PBS.

Imaging and photooxidation

If not for EM, cells were imaged in PBS with an inverted confocal microscope Leica SPEII. Within the same experimental set, which includes triplicates, the imaging parameters (laser power, gain, and offset) are adjusted on the brightest samples/conditions for optimal use of the microscope detector's dynamic range. For photooxidation, cells were imaged in 2.3 mM DAB (Sigma-Aldrich #D8001) in 0.1 M cacodylate (Sigma-Aldrich #C0250) buffer pH 7.4 with the same microscope. DAB was photo-oxidized by JF570-PROMPT under illumination with a 63× objective for 12 min using the 575/25 line (310 mW) of a Spectra 7 LED light source (Lumencor). Then samples were washed with 0.1 M cacodylate buffer pH 7.4 and processed for EM.

Image analysis

Fluorescence intensities were measured using ImageJ. Mean intensities were calculated from more than 30 cells from 3 independent experiments. In the figures, the brightness and contrast are optimized on the brightest sample/condition and applied to all the images displayed in the same experiment.

Resin embedding, sectioning, and TEM imaging

After photo-oxidation, samples were post-fixed for 30 min with 1% osmium (Electron Microscopy Sciences #19150) + 1.5% potassium ferrocyanide in 0.1 M cacodylate buffer pH 7.4 at 4 °C. Then the samples were dehydrated in 1-min successive 4 °C baths of increasing ethanol concentration (20%, 50%, 70%, 90%, $3 \times 100\%$), and incubated for 1-h in 50% ethanol/50% durcupan resin (Sigma-Aldrich #44611, #44612, #44613, #44614). Samples were then incubated 3×2 -h in 100% durcupan mixture, and the resin was polymerized for 48 h at 60 °C. At the photo-oxidized areas, blocks of hardened resin were cut off from the dish and mounted onto dummy blocks using glue. The glass was popped up by water infiltration, and the blocks were trimmed around the cells of interest to shape a block face of approximatively 400 µm wide and 200 µm tall. Block faces were imaged with an upright light microscope at 10X to facilitate correlation. Ribbons of 70 nm thick sections were produced from these blocks and collected on glow-discharged formvar coated slot grids (Electron Microscopy Sciences #FF2021-Cu-ET). Areas of interest were imaged at 120 keV with an FEI Tecnai Spirit transmission electron microscope.

Multi tilt axis electron tomography

For electron tomography, 250 nm thick sections were produced and collected onto Luxel grids (Luxel, #LuxFilmTEM C-S-M-L), then incubated for 20 s with 20% 10 nm gold fiducials (Ted Pella#15703-20) + 0.05% BSA in water, blotted, and dry. The grids with sections were then imaged with an upright light microscope at 10×. The grids were loaded onto a rotation holder (Fischione Instruments, model 2040). After correlation, a four tilt-axis series (one tilt-series: -60° to $+60^{\circ}$, one image every 0.25°) at different orientation (0°, 45°, 90°, 135°) of the region of interest was acquired using an FEI Titan Halo operated at 300 keV and a DE64 camera, while controlled by SerialEM²⁹. The entire set of 1936 acquired images were aligned and reconstructed using TxBR package^{30, 31}.

Data availability

All data generated or analyzed are included in the article and its Supplementary Information files. Additional datasets and analyses, as well as unique reagents (e.g., plasmids) and relevant information reported in this paper will be shared by the corresponding authors upon request.

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Author contributions

G.A.C., S.R.A, D.B, and M.H.E. designed research; G.A.C., S.P., J.H., and S.R.A. performed research; G.A.C., S.R.A, D.B, S.P., and M.H.E analyzed data; G.A.C., S.R.A, and M.H.E. drafted the paper and all authors revised it.

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Competing interests

A patent application (Provisional Application No. 63/343,469, filed May 18, 2022) covering the PROMPT method has been filed by the University of California, with inventors G.C., S.R.A., D.B., and M.H.E. The authors J.H. and S.P. declare no competing interests.

Additional information

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