Click-EM for imaging metabolically tagged nonprotein biomolecules

John T Ngo^{1,9}, Stephen R Adams¹, Thomas J Deerinck², Daniela Boassa², Frances Rodriguez-Rivera³, Sakina F Palida¹, Carolyn R Bertozzi^{4,5}, Mark H Ellisman^{2,6} & Roger Y Tsien^{1,2,7,8*}

EM has long been the main technique for imaging cell structures with nanometer resolution but has lagged behind light microscopy in the crucial ability to make specific molecules stand out. Here we introduce click-EM, a labeling technique for correlative light microscopy and EM imaging of nonprotein biomolecules. In this approach, metabolic labeling substrates containing bioorthogonal functional groups are provided to cells for incorporation into biopolymers by endogenous biosynthetic machinery. The unique chemical functionality of these analogs is exploited for selective attachment of singlet oxygen-generating fluorescent dyes via bioorthogonal 'click chemistry' ligations. Illumination of dye-labeled structures generates singlet oxygen to locally catalyze the polymerization of diaminobenzidine into an osmiophilic reaction product that is readily imaged by EM. We describe the application of click-EM in imaging metabolically tagged DNA, RNA and lipids in cultured cells and neurons and highlight its use in tracking peptidoglycan synthesis in the Gram-positive bacterium *Listeria monocytogenes*.

M has long been the main technique to image cell structures with nanometer resolution but has lagged behind light microscopy in the crucial ability to make specific molecules stand out. Antibodies have traditionally been used to introduce biomoleculespecific contrast to cells through direct conjugation with electrondense secondary labels (such as gold nanoparticles); however, conventional immuno-EM is technically demanding because cells must be cryoprotected and ultracryosectioned for large macromolecules to reach intracellular sites. Detergent permeabilization, which facilitates antibody penetrance for immunofluorescence staining, degrades EM-visible cellular landmarks ('ultrastructure') and is therefore generally avoided during sample preparation. Replacement of gold particles with labels capable of polymerizing diaminobenzidine (DAB) enables catalytic amplification^{1,2}, but use of antibody conjugates of such labels still requires cryosectioning or detergent permeabilization to stain intracellular epitopes.

Genetically encoded EM tags that polymerize DAB can overcome these challenges because labeling occurs before fixation and the auxiliary reagents required for generating contrast (O2, DAB and osmium tetroxide (OsO₄)) are small molecules that readily permeate cell membranes. Tags such as miniSOG³ and the tetracysteine motif bound to ReAsH⁴ polymerize DAB through photogeneration of singlet oxygen (1O₂), a short-lived excited state of oxygen that in the presence of DAB, produces, a localized polymeric precipitate that can be stained with osmium and therefore readily distinguished by EM (Fig. 1a). We have previously shown that photooxidationbased polymerization of DAB occurs within nanometers of the fluorophore and is capable of outlining the surfaces of a labeled structure with the osmiophilic product^{3,4}. DAB can also be polymerized enzymatically using a genetically encoded peroxidase-based tag such as APEX^{5,6}. These tags represent a versatile toolkit for imaging genetically tagged proteins by EM.

What remains missing from the EM toolkit, however, are analogous tools for imaging nonprotein biomolecules such as glycans, nucleic acids and lipids. Nonprotein biomolecules comprise a significant fraction of living matter, and a simple and generalizable method for visualizing them by EM would substantially enhance efforts to dissect cellular biochemistry at the nanometer scale. Although genetically encoded tags are powerful tools for imaging proteins, their direct application to other classes of biomolecules can be challenging; for instance, fusion of a fluorescent protein to the bacteriophage MS2 coat protein can be used to detect transcripts tagged with a cognate stem-loop RNA sequence, but imaging studies using such probes can be confounded by signals arising from both RNA-bound and unbound reporter proteins⁷.

Substantial progress toward fluorescence imaging of nonprotein biomolecules has been made using metabolic labeling followed by bioorthogonal ligation with imaging probes. Through this approach, cell-surface glycans⁸, nucleic acids^{9,10}, lipids¹¹ and proteins¹² have been imaged in various biological settings. These successes suggested that bioorthogonal labeling could be extended to other imaging modalities, such as EM. Here we describe click-EM, an EM imaging technique for direct visualization of nonprotein biomolecules via DAB polymerization. In this approach, metabolic analogs containing bioorthogonal functional groups (including artificial monosaccharides, nucleotides, fatty acids or other metabolites) are provided to cells and incorporated into biopolymers by endogenous biosynthetic machinery. The unique chemical functionality of these analogs is exploited for selective attachment of ¹O₂-generating fluorescent dyes via bioorthogonal click chemistry reactions¹³⁻¹⁷. Dyelabeled biomolecules can be imaged by fluorescence and correlated to EM observations through detection of photogenerated DAB precipitates. We describe the application of click-EM in imaging metabolically tagged DNA, RNA and lipids in cultured cells and

¹Department of Pharmacology, University of California, San Diego, La Jolla, California, USA. ²National Center for Microscopy and Imaging Research, University of California, San Diego, La Jolla, California, USA. ³Department of Chemistry, University of California, Berkeley, California, USA. ⁴Department of Chemistry, Stanford University, Stanford, California, USA. ⁵Howard Hughes Medical Institute, Stanford University, Stanford, California, USA. ⁶Department of Neurosciences, University of California, San Diego, La Jolla, California, USA. ⁷Department of Chemistry & Biochemistry, University of California, San Diego, La Jolla, California, USA. ⁸Howard Hughes Medical Institute, University of California, San Diego, La Jolla, California, USA. ⁹Present address: Department of Biomedical Engineering, Boston University, Boston, Massachusetts, USA. *e-mail: rtsien@ucsd.edu

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neurons and highlight its use in tracking peptidoglycan synthesis in the Gram-positive bacterium *L. monocytogenes*.

RESULTS

Screening of dyes for click-EM DAB photooxidation

We sought dyes that are capable of efficiently photooxidizing DAB and could also be selectively conjugated to metabolically tagged biomolecules in fixed cells. In an initial screen we tested 13 azidefunctionalized dyes with excitation wavelengths ranging from blue to far-red (**Supplementary Results**, **Supplementary Fig. 1**), including dyes known to photogenerate substantial levels of ¹O₂, such as eosin¹⁸, methylene blue¹⁹ and IRDye700DX²⁰. Eosin, a tetrabrominated derivative of fluorescein, has previously been used for DAB photooxidation as a conjugate to antibodies² or phalloidin²¹. The heavy atoms (bromines) on eosin increase the likelihood that the photoexcited chromophore will undergo an intersystem crossing (ISC) to the triplet state, from which ¹O₂ can be generated from ground state O₂. We also tested dyes not expected to generate significant amounts of ¹O₂, such as Alexa Fluor 647, Cy5.5 and sulforhodamine-101 (SR101). Alexa Fluor 647 and Cy5.5, both of which are derived from Cy5, seemed unlikely to generate substantial levels of ${}^{1}O_{2}$ given that cyanines generally exhibit very low ISC quantum yields ($\Phi_{ISC} < 0.003$ for Cy5 (ref. 22)) in the absence of heavy atom substitution^{22,23}. We thought that SR101 was also unlikely to produce sufficient amounts of ${}^{1}O_{2}$ for DAB photooxidation, given that nearly all photons absorbed by SR101 result in fluorescence emission²⁴.

To assess the performance of the selected dyes, we labeled HeLa cells with 5-ethynyl-2'-deoxyuridine (EdU) (**Fig. 1b**), an alkynecontaining thymidine analog that is incorporated into DNA during replication⁹. After an overnight pulse with EdU, cells were fixed and subjected to Cu(1)-catalyzed azide–alkyne cycloaddition (CuAAC), a reaction referred to as click chemistry^{13–15}, with azide-functionalized derivatives of the selected dyes. Individual dyes were then tested for their ability to photooxidize DAB by immersion of labeled cells in a solution of DAB followed by intense illumination through an appropriate bandpass filter. After 5 min of illumination, successful DAB photooxidation was judged by the appearance of



Figure 1 | **Click-EM imaging of EdU-labeled HeLa cells.** (**a**) Schematic of $1O_2$ use to generate EM contrast. (**b**) Schematic of metabolic labeling and CuAAC reaction steps of the click-EM procedure. Structures of EdU, DBF-azide and their CuAAC ligation product are shown. (**c**) Fluorescence and transmitted light images of EdU-labeled HeLa cells after CuAAC ligation with DBF-azide. Transmitted light images before and after photooxidation are shown; polymeric DAB precipitates accumulated during the illumination period appear as optically dense deposits that overlay with DBF-azide emission. (**d**) Photooxidized cells after OsO₄ staining. Osmium-stained DAB precipitates appear brown under white light. The dotted blue line defines the area of illumination. (**e**) Correlated light and EM images of a mitotic cell labeled with EdU and DBF-azide. (**f-m**) EM images of EdU-labeled HeLa cells: a nonphotooxidized cell (**f**), S- or G2-phase cells (**g-i**) and mitotic cells (**j-m**).

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optically dense precipitates that were visible under transmitted light. The photooxidation efficiency and specificity of each dye was estimated by the opacity and localization of the generated precipitates, which we expected to be predominantly nuclear owing to incorporation of EdU into DNA.

Of the 13 dyes tested, 10 generated detectable DAB precipitates after 5 min of illumination (**Supplementary Fig. 2**). Precipitates were not observed in cells labeled with the control dyes Alexa Fluor 633–azide, Cy5.5-azide and SR101-azide. Of the photooxidizing dyes, eosin-azide produced DAB precipitates of the greatest opacity, though precipitates were detected throughout the cell, including in cytosolic areas, where we did not expect EdU-labeled DNA to be present. Eosin has a substantial ${}^{1}O_{2}$ quantum yield ($\Phi_{\Delta} = 0.57$) 18 but a low fluorescence quantum yield ($\Phi_{F} = 0.20$) 25 . Owing to the presence of four bromine atoms, eosin-azide is hydrophobic, and extranuclear DAB precipitation is likely due to nonspecific binding of the chromophore to hydrophobic cellular components.

After eosin-azide, the second most efficient photooxidizer was dibromofluorescein-azide (DBF-azide) (**Fig. 1b**). DBF possesses a slightly lower Φ_{Δ} than eosin (0.42)¹⁸ and contains only two bromine atoms, making it less hydrophobic. After CuAAC ligation with EdU-labeled cells, fluorescence emission from DBF-azide colocalized with Hoechst 33342 staining (**Supplementary Fig. 3**). Dye-labeled control cells that were not exposed to EdU did not exhibit detectable levels of DBF-azide fluorescence. Subsequent DAB photooxidation with DBF-azide produced exclusively nuclear precipitates (**Fig. 1c** and **Supplementary Fig. 2**). The nuclear specificity of DAB precipitation by DBF-azide was maintained over an illumination period of up to 15 min, at which point fully opaque DAB precipitates were generated (**Supplementary Fig. 4**). Thus, DBF-azide is capable of efficiently photooxidizing DAB and can be conjugated to alkyne-labeled biomolecules via CuAAC with excellent specificity.

Click-EM imaging of newly EdU-labeled DNA using DBF-azide

After photooxidation with DBF-azide, EdU-labeled cells were stained with OsO_4 (**Fig. 1d**), embedded in resin and cut to 80-nm sections before imaging by transmission EM (TEM). On EM micrographs, OsO_4 -stained DAB precipitates appeared as discernable contrast in patterns that correlated with DBF-azide fluorescence (**Fig. 1e**) and were consistent with the known localization of cellular DNA. To determine selectivity of DAB deposition at the level of EM, we compared photooxidized and nonphotooxidized cells (those residing outside the area of illumination). Nonphotooxidized cells showed the expected contrast due to OsO_4 (1% w/v) staining of endogenous nuclear components (**Fig. 1f**). Notably, the extranuclear regions of photooxidized and nonphotooxidized cells showed similar intensity, indicating that little or no DAB was nonspecifically precipitated during photooxidation.

We captured images of EdU-labeled cells occupying various stages of the cell cycle (**Fig. 1g–m**). In cells that were likely in S phase at the time of fixation, EdU labeling was distributed throughout the nucleus and staining was observed along the nuclear periphery, at the edges of nucleoli and in subnuclear regions where replication foci are localized during mid–S phase^{26,27} (**Fig. 1g–i**). Mitotic cells in prometaphase and metaphase showed darkly stained condensed chromosomes (**Fig. 1j,k**), and cells in telophase displayed lightly stained extensions radiating from tightly packed and darkly stained regions, which appeared to represent various degrees of chromatin decondensation (**Fig. 1l,m**).

A significant advantage of the photooxidation procedure is that DAB precipitates are photogenerated within intact cells, allowing electron density to be introduced at labeled sites throughout entire cellular volumes. In contrast, immuno-EM of cell cryosections results in sparse labeling of only a fraction of total epitopes (those accessible at the section surface). We exploited the staining uniformity afforded by DAB photooxidation to visualize EdU-labeled



Figure 2 | **SBEM imaging of EdU-labeled HEK293 cells.** (**a**) Montage of selected 2D (45 μ m × 45 μ m) images collected by SBEM. Individual images are numbered according to their position in a series of 385 sequentially recorded micrographs. (**b**) Full 3D volume of an EdU-labeled mitotic cell reconstructed from a sum of 385 images from the area shown in **a**. Images were collected serially along the *z* axis at 60-nm intervals (*z*-dimension distance = 23 μ m). Signal intensity arising from EdU-labeled DNA is shown in purple. (**c**) A 3D rendering of the signal arising from OsO₄-stained DAB deposits.

chromosomes in 3D through stereomicroscopic analysis of individual sections (**Supplementary Fig. 5**). Additionally, entire volumes of EdU-labeled HEK293 cells were reconstructed using serial block-face scanning EM (SBEM), a technique in which sequential surface images are collected from a piece of mounted resin containing labeled cells of interest between rounds of automated, on-stage serial sectioning (**Fig. 2** and **Supplementary Video 1**). Labeling of DNA in HeLa and HEK293 cells demonstrates the potential of click-EM to visualize nonprotein molecules with excellent temporal and spatial resolution.

Click-EM imaging of nascent RNAs

Next we applied click-EM to visualize nascent transcripts using 5-ethynyl-uridine (EU) (**Fig. 3a**), an alkyne-containing uridine analog that is selectively incorporated into RNA¹⁰. Three different RNA polymerase (RNAP) complexes perform nuclear RNA synthesis: RNAP I synthesizes preribosomal RNAs (pre-rRNAs) and is localized to nucleoli; RNAP II is nucleoplasmic and transcribes messenger RNAs (mRNAs); and RNAP III (also nucleoplasmic) synthesizes small RNA molecules such as transfer RNAs (tRNAs) and the 120-nucleotide 5S-rRNA. Fluorescence imaging of EU-labeled cells revealed a predominantly nuclear localization of labeled transcripts with intensely stained nucleoli (**Fig. 3b**). Consistent with previous observations¹⁰, inhibition of RNAP I by actinomycin selectively abolished nucleolar staining, confirming that nucleolar signal arose from EU incorporation by RNAP I transcriptional activity.

EM imaging showed a similar distribution of EU-labeled transcripts, though in addition to strong nucleolar staining and diffuse cytoplasmic labeling, we also detected punctate nucleoplasmic densities that were not observed by fluorescence microscopy (Fig. 3c-e). Previous studies using immunogold staining of cells

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Figure 3 | Imaging of nascent transcripts using EU. (a) Structure of EU. (b) Fluorescence detection of EU-labeled transcripts in HeLa cells after CuAAC ligation with DBF-azide. (**c**-**g**) EM micrographs of EU labeled cells: uninhibited HeLa cells exhibiting darkly stained nucleoli, diffuse nucleoplasmic labeling and stained nucleoplasmic intensities (**c**-**e**); the nucleus of a HeLa pulsed with EU in the presence of α -amanitin (for inhibition of RNAP II and III) exhibiting darkly stained nucleoli without stained nucleoplasmic intensities (**f**) and the nucleus of a nonphotooxidized cell (**g**).

labeled with 5-bromouridine-5'-triphosphate (BrUTP) or biotinylated cytidine-5'-triphosphate (biotin-14-CTP) reported clustered gold particles that colocalized with either RNAP II or RNAP III in nucle-oplasmic 'transcription factories'^{28,29}. Labeling of nascent transcripts within transcription factories was sensitive to inhibition of the associated RNAP. Similarly, we found that inhibition of RNAP II and

RNAP III using α -amanitin³⁰ abolished the punctate nucleoplasmic densities we observed by click-EM (**Fig. 3f,g**). Thus, the nucleoplasmic densities observed by click-EM probably represent nascent RNA molecules emerging from RNAP II and RNAP III transcription factories. This observation illustrates that click-EM imaging of EU-labeled transcripts is in agreement with established methods of imaging newly synthesized RNAs²⁸.

Click-EM imaging of choline-containing phospholipids

Choline-containing phospholipids (Chophospholipids) have important roles in various signaling pathways and are the most abundant phospholipids in cellular membranes. Chophospholipids can be tracked via fluorescence microscopy using choline analogs such as propargyl-choline¹¹ and azido-choline (AzCho)³¹ (**Fig. 4a**). Azide-functionalized analogs such as AzCho are especially versatile metabolic labeling substrates because they are susceptible to modification by CuAAC as well as strainpromoted ligation with cyclooctynes—a reaction referred to as copper-free click chemistry that is compatible with live cells^{16,17}. To test the applicability of copper-free click chemistry to click-EM we reacted AzCho-labeled HeLa cells with a dibenzoazacyclooctyne (DIBAC)functionalized³² derivative of the far-red excited dye IRDye700DX (IRDye700DX-DIBAC) (**Supplementary Fig. 1**). Control cells that were not exposed to AzCho showed only minimal levels of background fluorescence after dye labeling (**Supplementary Fig. 6**), whereas AzCho-labeled cells revealed bright staining of intracellular membranes in patterns similar to those previously observed using bioorthogonal choline analogs¹¹.

DAB photooxidation using IRDye700DX-DIBAC in AzCho-labeled cells was specific and generated precipitates in patterns that correlated with dye fluorescence (Fig. 4b,c). By EM we observed continuous staining along intracellular membranes, including those of the endoplasmic reticulum (ER), nuclear envelope and mitochondria (Fig. 4d). Under high magnification, sites of ER-mitochondria contacts and the intricate folds of inner mitochondrial membranes were visible (Fig. 4e). In cultured cortical neurons, we observed AzCho staining on various membranes in neuronal somata, axons, dendrites and synapses (Supplementary Fig. 7). The ER membrane was the most intensely stained structure in both cell types examined, consistent with its high composition of Cho-phospholipids. In mammals, the ER membrane contains more choline than any other cellular mem-

brane and is composed of nearly 60% phosphatidylcholine³³. Mitochondrial membranes contain less Cho-phospholipids than the ER membrane and were less intensely stained. To confirm that the EM contrast observed was due to osmium-stained DAB precipitates deposited during photooxidation and not to intrinsic osmiophilicity of these membranes, we imaged cells at the edge



Figure 4 | Click-EM imaging of AzCho labeled HeLa cells. (a) Schematic of the metabolic incorporation of AzCho into Cho-phospholipids and subsequent detection of labeled membranes using copper-free click chemistry. **(b)** AzCho-labeled HeLa cells imaged by fluorescence (left) and transmitted light before (middle) and after (right) DAB photooxidation. **(c)** AzCho-labeled HeLa cells imaged by white light after DAB photooxidation and OsO₄ staining. The blue dotted line indicates the area of illumination. **(d)** TEM micrograph of an AzCho-labeled HeLa cell after DAB photooxidation and osmium staining. **(e)** A high-magnification TEM image showing the detailed features of AzCho-labeled mitochondria, including sites of ER-mitochondria contacts (black arrowheads).

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Figure 5 | Click-EM imaging of PG in *L. monocytogenes.* (a) Structure of AlkDAla. (b) Extracellular (black arrow) and intracellular (blue arrows) routes of D-amino acid incorporation into *L. monocytogenes* PG. (c) TEM images of AlkDAla-labeled *L. monocytogenes* cells. A nonphotooxidized control cell (left) is shown beside a photooxidized cell (right) for comparison. (d) High-magnification TEM of a photooxidized dividing Δ PBP5 cell showing labeled extracellular PG and stained intracellular precursors (IP); IP staining is depicted as a continuous contour on the cytoplasmic side of the plasma membrane (PM; low-contrast region between bands).

of the photooxidation region. In cells that partially occupied both illuminated and non-illuminated areas, we detected membrane staining only in light-exposed areas (**Supplementary Fig. 8**), thus verifying that the observed contrast was due to light-deposited DAB precipitates.

Click-EM imaging of peptidoglycan synthesis

Finally, we applied click-EM to image bacterial peptidoglycan, an essential cell component of most bacterial species that is also the target of many antibiotics. As previously reported in the context of fluorescence imaging, we used D-propargylglycine (AlkDAla)³⁴ (Fig. 5a and Supplementary Fig. 9) to label peptidoglycan (PG) in L. monocytogenes. There are two primary routes through which D-amino acids are incorporated into L. monocytogenes PG: (i) periplasmic (extracellular) addition to crosslinked PG peptides, and (ii) cytosolic (intracellular) incorporation into PG precursors^{35,36} (Fig. 5b). Previous analyses using MS and light microscopy suggest that AlkDAla is incorporated via the cytosolic pathway in L. monocytogenes³⁴; however, it is difficult to selectively distinguish cytosolic AlkDAla-containing intermediates with conventional fluorescence imaging. In the cytosolic addition mechanism, D-amino acids are incorporated into a precursor that transfers a phospho-N-acetylmuramyl-pentapeptide moiety to undecaprenol, forming the membrane-anchored intermediate lipid I. Subsequent attachment of N-acetylglucosamine to lipid I results in the formation of lipid II, the final cytosolic intermediate in PG biosynthesis. During cell growth, lipid II is transported to the extracellular face of the bacterial membrane for subsequent incorporation into the PG mesh via transglycosylation.

Direct imaging of labeled intermediates along the cytoplasmic leaflet of the plasma membrane would provide definitive proof of a cytosolic route of AlkDAla incorporation. However, distinguishing intracellular precursors from extracellular PG by light microscopy is challenging because they are separated only by the thickness of the cell membrane (~7 nm in *L. monocytogenes*)³⁷. Because EM provides exquisite resolution, we anticipated that click-EM would



Figure 6 | Click-EM imaging of wild-type and ramoplanin-treated L. *monocytogenes.* (a) Schematic of expected staining pattern (top) and click-EM imaging (bottom) of AlkDAla-labeled wild type *L. monocytogenes* cells. AlkDAla is removed along the cell length by the endogenous PBP5, resulting in polar staining of extracellular PG (black arrowheads); labeled intracellular precursors are observed as a continuous contour on the cytoplasmic face of the cell membrane (red arrows). (b) Schematic of the expected staining pattern (top) and click-EM imaging (bottom) of ramoplanin-treated wild-type *L. monocytogenes* cells. Labeling of extracellular PG is not detected on drug-treated cells (white arrowheads), but labeled intracellular precursors remained visible (red arrows).

enable us to unambiguously distinguish labeled extracellular PG and its cytoplasmic intermediates. In an initial analysis, we labeled *L. monocytogenes* cells lacking PBP5 (Δ PBP5)³⁸, an extracellular D,D-carboxypeptidase that removes D-amino acids from PG along the length of the organism, but not at polar regions³⁴. Elimination of PBP5 was used to ensure maximal incorporation of the analog into PG and its biosynthetic intermediates. After overnight labeling with AlkDAla, cells were fixed, subjected to CuAAC ligation with DBF-azide and subsequently used for DAB photooxidation.

EM imaging of photooxidized cells revealed electron-dense staining along the cell perimeter (Fig. 5c). High-magnification imaging showed two distinct bands of staining: a thick band of extracellular PG and a thin intracellular band, separated by a region of reduced contrast representing the plasma membrane (Fig. 5d and Supplementary Fig. 10a). To confirm the accuracy of these assignments, we labeled wild-type L. monocytogenes cells using a short pulse with AlkDAla (40 min). Previous fluorescence imaging of wild-type cells labeled under similar conditions revealed a predominantly septal and polar localization of AlkDAla, presumably due to removal of the analog along the cell length by endogenous PBP5 (ref. 34). EM imaging revealed thick segments of staining at the poles of labeled wild-type cells, thus confirming our ability to identify extracellular PG (Fig. 6a and Supplementary Fig. 10b). We additionally detected a thin and continuous contour of staining in wild-type cells that was separated from extracellular PG by the plasma membrane, as in Δ PBP5 cells.

To determine whether AlkDAla is incorporated solely by the cytoplasmic mechanism, we labeled wild-type cells with AlkDAla in the presence of ramoplanin, a glycolipodepsipeptide antibiotic that inhibits the transglycosylation step of PG synthesis³⁹. Ramoplanin prevents the transfer of disaccharide-pentapeptide monomers into growing PG strands by binding lipid II extracellularly, thereby halting addition of D-amino acid to PG via the cytosolic route. Thus, the absence of extracellular labeling on ramoplanin-treated cells would support a solely cytosolic route of AlkDAla incorporation, whereas extracellular detection of AlkDAla would suggest dual modes of addition. Because ramoplanin acts extracellularly, we anticipated that the drug would have no effect on cytoplasmic synthesis of PG intermediates and thus expected cells to retain intracellular staining.

Consistent with this prediction, we observed intracellular staining in ramoplanin-treated cells, demonstrating that ramoplanin does not prevent the incorporation of AlkDAla into cytoplasmic intermediates (**Fig. 6b**). Extracellular staining was not detected on ramoplanin-treated cells, thus suggesting that AlkDAla is added to *L. monocytogenes* PG exclusively through the cytosolic route and that extracellular (periplasmic) addition of the analog to crosslinked PG peptides does not occur.

DISCUSSION

As demonstrated here, the click-EM photooxidation technique provides a simple and direct means for determining the detailed cellular distribution of metabolically tagged (nonprotein) biomolecules. Click-EM is advantageous over conventional immuno-EM techniques because the biomolecules of interest are labeled before fixation and the auxiliary reagents required for generating contrast (O_2 , DAB and OsO_4) are small molecules that readily diffuse into fixed cells. Click-EM permits high-quality preservation of EM-visible landmarks, the integrity of which is essential to assigning the precise location of labeled biomolecules. Application of the DAB photooxidation technique allows electron density to be deposited at labeled sites throughout entire cellular volumes, thus enabling 3D biomolecular localization via stereoscopic analysis, electron tomography and SBEM.

Metabolic tagging with radioisotopic labels has been used to image biomolecules via autoradiographic EM detection⁴⁰, but these methods are tedious, rarely used and have much lower resolution than alternative techniques^{41,42}. Previously reported techniques relying on immunogold detection of incorporated metabolic labels are burdened by the limitations and challenges associated with the membrane impermeability of antibodies and the random detection of a small fraction of total epitopes. For instance, EM imaging of propargyl-choline by attachment of biotin followed by detection using anti-biotin and gold-nanoparticle-conjugated protein A results in sparse labeling of intracellular membranes¹¹. In contrast, click-EM imaging of AzCho reveals continuous staining along membrane contours (Fig. 3c) that not only aligns with fluorescence imaging of choline analogs but also more accurately depicts the distribution and packing density of choline-modified phospholipids in cell membranes.

We exploited the advantages of click-EM for nanoscale interrogations of metabolically tagged DNA, RNA and lipids in cultured mammalian cells and neurons. Additionally, we used click-EM to track the route of AlkDAla metabolism from its initial incorporation into cytosolic lipid I to its eventual integration into *L. monocytogenes* PG (**Figs. 5** and **6**). The high resolution of click-EM enabled us to distinguish extracellular PG from its intracellular intermediates (which are separated by only 7 nm) and was essential for our analysis. Because gold nanoparticles used in immuno-EM are bulky (typically 5–25 nm in diameter) and attached to large macromolecules (an immunoglobulin is ~9 nm, and protein A is ~5 nm in length), individual labels can be situated many nanometers away from a labeled site. Thus, use of immunogold staining may not have provided the resolution necessary to distinguish various AlkDAla-labeled species in the bacterium.

Click-EM may be further improved through development of strategies for localizing labeled biomolecules in the context of specific proteins or genomic loci. Dual labeling of this kind may be achieved through combining click-EM with correlative superresolution imaging of fluorescent proteins that survive osmium staining⁴³ or selective targeting of fluorescent dyes that can be imaged in-resin⁴⁴. Though we did not attempt to determine the detection sensitivity of the method, we predict that click-EM can be used to image low-abundance molecules through the catalytic amplification afforded by DAB photooxidation. Nevertheless, the sensitivity of the method may be improved through identification of dyes with increased ¹O₂ yields that can be specifically attached to metabolically labeled structures or that are 'switched on' after ligation^{45,46}. Alternatively, increasing dye-ligation yields by subjecting cells to multiple rounds of CuAAC can augment the detection of labeled biomolecules without compromising selectivity⁹.

Click-EM should find widespread applications in the ultrastructural localization of many diverse classes of biomolecules. In principle, the method is applicable to any biomolecule that can be metabolically tagged with a bioorthogonal functional group. Throughout this work, we compared photooxidized and nonphotooxidized cells to identify DAB-specific signals on EM micrographs. In several cases, image interpretation was further aided by imaging of drug-treated cells, mutant cells or cells at different stages of the cell cycle. Future click-EM users should employ similar controls to ensure that DAB-derived signals can be distinguished from ultrastructural contrast. Light-absorbing compounds (such as certain pharmacological agents or fluorescent counterstains) should be tested for potential ¹O₂-photogenerating activity before use. Although the applications reported here were focused on nonprotein biomolecules, click-EM should also be useful for imaging discrete subpopulations of proteins, such as those that are posttranslationally modified⁸, or proteins with a specific enzymatic activity⁴⁷. Also, click-EM could be coupled with molecule-specific, proximity-enhanced approaches such as those recently described for labeling glycans and individual proteins^{48,49}. Global incorporation of azide- or alkyne-containing noncanonical amino acids^{12,50} in combination with click-EM may be an advantageous strategy for imaging localized neuronal protein synthesis in dendrites, a neuronal subregion that is difficult to resolve by light microscopy. Additional applications for click-EM include imaging of labeled biomolecules in other cellular subregions and structures that are difficult to fully resolve using light microscopy, such as axon terminals, mitochondria, synaptic vesicles and autophagosomes.

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METHODS

Methods and any associated references are available in the online version of the paper.

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Author contributions

J.T.N., S.R.A., T.J.D., D.B., F.R.-R. and S.F.P. performed experiments. J.T.N., S.R.A., T.J.D., D.B., F.R.-R., S.F.P., C.R.B., M.H.E. and R.Y.T. analyzed data and prepared the manuscript.

Competing financial interests

The authors declare no competing financial interests.

Additional information

Any supplementary information, chemical compound information and source data are available in the online version of the paper. Reprints and permissions information is available online at http://www.nature.com/reprints/index.html. Correspondence and requests for materials should be addressed to R.Y.T.

ONLINE METHODS

Metabolic labeling reagents. EdU and EU were purchased from Life Technologies. AzCho (azido-choline) was synthesized according to previously reported procedures³¹ and verified by mass spectrometry before use. AlkDAla (D-propargylglycine) was purchased from Acros.

Reactive dyes. Fluorescein-azide was purchased from Lumiprobe. Alexa Fluor 488–azide and Alexa Fluor 647–azide were purchased from Life Technologies. TAMRA-azide and SR101-azide were purchased from Click Chemistry Tools. Cy5.5-azide was purchased from AAT Bioquest.

Synthesis of reactive dyes. N-hydroxysuccinimide (NHS) esters of Alexa Fluor 633, methylene blue and IRDye700DX were purchased from Life Technologies, ATTO-TEC GmbH, and LI-COR, respectively. Eosin isothiocyanate (ITC) was purchased from Sigma.

Synthesis of DBF. To a stirred suspension of 5(6)-carboxyfluorescein (376 mg, 1 mmol) in 80% aqueous acetic acid (5 ml) at room temperature, bromine (102 μ L, 2 mmol) was added dropwise. The solid dissolved and after stirring overnight, the crude product was collected by filtration, washed with water, and dried *in vacuo* over P₂O₅ (yield 0.5 g, 94%). LC-MS revealed the desired product (56% by absorbance at 254 nm) with unreacted starting material (1%) and mono- (5.5%), tri- (15.6%) and tetrabrominated (21.7%) products, and was used without further purification. ES-MS (*m*/*z*): [M]⁺ for C₂₁H₁₀Br₂O₇, 534.9; found, 534.8.

Synthesis of DBF-NHS. Crude 5(6)-carboxy-4',5'-dibromofluorescein (200 mg, 0.37 mmol) was dissolved in anhydrous THF (10 ml) under N₂, and *N*-hydroxysuccinimide (52 mg, 0.45 mmol) was added followed by DIPC (59 μ l, 0.38 mmol). After 4 h stirring at room temperature, LC-MS indicated an incomplete reaction, so an additional 25 μ L of DIPC was added. After overnight stirring, the reaction mix was evaporated, dissolved in dry DMSO (1 mL), filtered and purified by preparative HPLC eluting with a gradient of 10–90% acetonitrile-water-0.05% TFA in 20 min. The collected fractions were pooled and immediately frozen and lyophilized to give a red solid (yield 98 mg, 42%; ES-MS (*m*/*z*), [M]⁺ for C₂₅H₁₃Br₂NO₉, 632.9; found, 632.9).

Synthesis of DBF-azide. DBF-NHS (5.1 mg, mixture of 5'- and 6'-isomers, 73% pure by HPLC with 21% hydrolyzed to DBF-carboxylic acid, 5.9 µmol) dissolved in dry DMSO (100 µl) was reacted with 3-azidopropylamine (Click Chemistry Tools; 80 µmol, 8 mg) at room temperature for 1 h. LC-MS of the reaction mixture confirmed complete reaction; the mixture was acidified with glacial acetic acid (175 µmol, 10 µl) and the desired product separated by RP-HPLC (Luna2 C18 columns, Phenomenex) with a water-acetonitrile-0.05% TFA gradient. Fractions containing the product were lyophilized, dissolved in DMSO, and quantified by absorbance in 0.1 N NaOH using an extinction coefficient of 95,000 M-1 cm-1 (ref. 51). Yield, 67%; 1H NMR (500 MHz, DMSO-*d*₆) δ p.p.m. 1.71 (t, *J* = 6.85 Hz, 4 H); 1.82 (quin, *J* = 6.72 Hz, 4 H); 3.45 (t, *J* = 6.72 Hz, 4 H); 6.65 (d, *J* = 7.83 Hz, 1 H); 6.66 (d, *J* = 7.58 Hz, 1 H); 6.77 (d, *J* = 6.85 Hz, 1 H); 6.79 (d, *J* = 6.85 Hz, 1 H); 7.49 (dd, *J* = 7.95, 0.61 Hz, 1 H); 7.74 (t, *J* = 0.98 Hz, 1 H); 8.10 (dd, *J* = 8.07, 0.73 Hz, 1 H); 8.18 (dd, *J* = 8.07, 1.47 Hz, 1 H); 8.26 (dd, J = 8.07, 1.71 Hz, 1 H); 8.48 (dd, J = 1.47, 0.73 Hz, 1 H); 8.65 (t, J = 5.62 Hz, 1 H); 8.87 (t, J = 5.62 Hz, 1 H); 10.98 (s, 1 H); 10.99 (s, 1 H). ¹³C NMR (126 MHz, DMSO-*d*₆) δ p.p.m. 28.14, 28.24, 36.81, 36.84, 48.44, 48.47, 82.66, 82.76, 97.48, 97.53, 110.28, 110.34, 112.69, 112.80, 126.19, 127.49, 127.66, 127.76, 136.36, 140.80, 148.49, 148.57, 152.37, 154.12,156.79, 164.44, 164.53, 167.79, 167.89. HRMS (*m/z*), [M]⁺ for C₂₄H₁₆Br₂N₄O₆, 613.9437; found, 613.9436.

General synthesis of reactive dyes from dye-NHS or dye-ITC. The dye-NHS or dye-ITC (1–5 mg) dissolved in dry DMF or DMSO was reacted with a 10to 20-fold excess of 3-azidopropylamine at room temperature for 1 h. After neutralization with glacial acetic acid, the desired product was purified by RP-HPLC (Luna2 C18 columns, Phenomenex) with a water-acetonitrile-0.05% TFA gradient except for Alexa Fluor 633–azide and IRDye700DX-azide, which used a 0.1 M triethylammonium acetate (pH 7.0)-acetonitrile gradient. Fractions containing the product were lyophilized, dissolved in DMSO, and quantified by absorbance using published or manufacturer's extinction coefficients (eosin-azide: HRMS (m/z), [M]⁻ for C₄₀H₁₃F₂N₆O₁₁S₆, 1,006.0734, found, 1,006.0728; methylene blue–azide: HRMS (m/z), [M]⁺ for $\begin{array}{l} C_{21}H_{26}N_7OS^+, \ 590.1277, \ found, \ 590.1265; \ IRDye700DX-azide: \ HRMS \ (m/z), \\ [M]^- \ for \ C_{73}H_{103}N_{15}O_{24}S_6Si_3, \ 10 \ 1,849.4932, \ found, \ 1,849.4942. \end{array}$

Synthesis of monobromo-TAMRA-azide. Monobromo-TAMRA-azide was prepared from TAMRA carboxylic acid (Novabiochem) following the method described for dibromination of rhodamine 123 (ref. 52). LC-MS analysis of the reaction mixture indicated 2.3% monobromo-, 85.5% dibromo- and 12.2% starting material. The evaporated reaction mixture was subsequently treated with 1.1 equivalents of TSTU and excess triethylamine in DMSO for 4 h. Subsequent addition of an excess of 3-azidopropylamine gave the desired product, which was isolated by RP-HPLC (HRMS (m/z): $[M]^+$ for $C_{28}H_{28}BrN_6O_4$, 590.1277, found, 590.1265).

Synthesis of monobromo-Cy5-azide. 1-(5'-carboxypentynyl)-2,3,3trimethylindoleninium-5-sulfonate (70 mg, 0.2 mmol)53 bromomalonaldehyde dianil bromide (38.2 mg 0.1 mmol)⁵⁴ and anhydrous potassium acetate (40 mg, 0.4 mmol) were refluxed in ethanol (0.5 ml) for 30 min. The reaction mix was cooled and left to stand overnight, and crude product collected by filtration as an iridescent blue solid (62 mg). Further purification of 30 mg by RP-HPLC gave 9.5 mg pure monobromo-Cy5-bis-carboxylic acid (yield, 24%; ES-MS (m/z): [M]⁺ for C₃₇H₄₆BrN₂O₁₀S₂⁺, 821.2, found, 821.2). The mono-NHS and azide were made by reaction of this product (5.9 mg, 7.1 µmol) with TSTU (2.2 mg, 7.1 µmol) in dry DMF (200 µl) and pyridine (20 µl) overnight. LC-MS revealed a mixture of mono- (32%) and di-NHS (4%) esters with unreacted starting material (64%). Excess 3-azidopropylamine (10 µl) was added, and after 30 min, the reaction was quenched with acetic acid (10 µl) to give the desired product (1.2 mg, 19%) following purification by RP-HPLC (HRMS (*m*/*z*): [M]⁺ for C₄₀H₅₂BrN₆O₉S₂⁺, 902.2342, found, 902.2338).

Synthesis of IRDye700DX-DIBAC. The product was prepared from the dye-NHS by reaction with excess DBCO-amine (Click Chemistry Tools) in DMF followed by purification by RP-HPLC using a 0.1 M triethylammonium acetate pH 7.0-acetonitrile gradient (HRMS (m/z): [M]⁻ for C₈₈H₁₁₁N₁₃O₂₅S₆Si₃, 2,025.5446; found, 2,025.5441).

Metabolic labeling of mammalian cells and neurons. Mammalian cells and neurons were maintained at 37 °C in a humidified cell culture incubator with 5% CO2. Authenticated HEK293 and HeLa cell lines (both American Type Culture Collection) cells were purchased directly from the vendor, and the lines were morphologically correct. Cells were not tested for mycoplasma contamination. HeLa and HEK293 cells were cultured in imaging dishes containing poly-Dlysine-coated coverslip glass bottoms (P35GC-0-14-C, MatTek Corporation) in DMEM supplemented with 10% FBS, 20 U/ml penicillin and 50 µg/ml streptomycin. Cortical neurons were dissociated by papain from postnatal day 2 (P2) Sprague-Dawley rats (male and female were used) and cultured in imaging dishes containing poly-D-lysine-coated coverslip glass bottoms (P35GC-0-14-C, MatTek Corporation) in Neurobasal A medium supplemented with 1× B27 supplements (both from Life Technologies), 2 mM GlutaMAX (Life Technologies), 20 U/ml penicillin and 50 µg/ml streptomycin. All animal procedures were approved by the Institutional Animal Care and Use Committee of UC San Diego. Experiments were not randomized, and no statistical method was used to predetermine sample size.

EdU labeling. HeLa and HEK293 cells were labeled with EdU overnight by addition to of the analog to culture medium at a final concentration of $10 \,\mu$ M.

EU labeling. HeLa cells were labeled with EU for 2.5 h by addition of the analog to culture medium at a final concentration of 2 mM. To inhibit RNAP I activity, actinomycin D was added to culture medium (at a final concentration of 100 nM) 30 min before the addition of EU. To inhibit RNAP II and RNAP III, α -amanitin was added to culture medium (at a final concentration of 25 µg/ml) 30 min before the addition of EU.

AzCho labeling. HeLa cells were labeled with AzCho overnight by addition of the analog to culture medium at a final concentration of 75 μ M. Cortical neurons were isolated from rats at 15–20 days *in vitro* (DIV) and labeled with AzCho under the same conditions.

Metabolic labeling of *L. monocytogenes* cells. Wild-type *L. monocytogenes* EGD-e cells were grown in Brain Heart Infusion (BHI) medium at 37 °C with shaking and labeled for 40 min or overnight. Δ PBP5 *L. monocytogenes* cells

(*lmo2754::tn*) were labeled at a final concentration of 5 mM AlkDAla under the same growth conditions. For ramoplanin inhibition studies, cells were grown in the presence of both 1 ug/ml ramoplanin and 5 mM AlkDAla for 40 min. Following labeling experiments, liquid cultures were washed twice with BHI medium to remove excess AlkDAla and prepared as glycerol stocks to be stored at -80 °C until processed for electron microscopy. Cells were thawed, rinsed once in PBS and resuspended in fresh PBS before adsorption onto poly-D-lysine-coated glass-bottom imaging dishes by pipetting cultures directly on the glass surface. After 30 min at room temperature, excess liquid culture was removed and the dishes were rinsed once with fresh PBS before fixation and further processing as described below.

Cell fixation. Cells were rinsed once with PBS and subsequently fixed using 2.5% (w/v) glutaraldehyde (Electron Microscopy Sciences) in a solution of 0.1 M sodium cacodylate buffer, pH 7.4 (Ted Pella Incorporated) containing 1 mM CaCl₂. Cell fixation was allowed to proceed for 5 min at 37 °C and then on ice for 1 h. The fixative was then removed and cells were rinsed with ice-cold cacodylate buffer (3 × 3 min) followed rinsing with an ice-cold solution of PBS containing 2% BSA (w/v) (2 × 3 min). Cells were kept on ice in cold PBS-BSA solution while dye reaction mixtures (described below) were prepared at room temperature.

CuAAC ligation reactions. All dyes were stored as concentrated stocks in DMSO and diluted into reaction mixtures before use. For conjugation of azide-functionalized dyes to alkyne-labeled biomolecules (EdU, EU, and AlkDAla) we used the Click-iT Cell Reaction Buffer Kit (Life Technologies) according to the manufacturer's protocol with the following modifications: (i) Cells were fixed with 2.5% (w/v) glutaraldehyde as described in the preceding section; (ii) cells were not subjected to permeabilization with Triton X-100 or another detergent, (iii) the azide-functionalized dyes described above were used at concentrations of $5-10 \,\mu$ M, and 4) the Click-iT cell buffer additive (Component C) was replaced by a 20 mg/ml solution of sodium ascorbate in ddH₂O (prepared fresh each time immediately before use). The sodium ascorbate solution was added as the last component of the reaction cocktails by dilution (tenfold, v/v) into reaction mixtures.

To initiate the CuACC reaction, the PBS–BSA solution was removed from the imaging dishes and immediately replaced with complete (ascorbatecontaining) dye-reaction mixture. Dishes were then gently and briefly swirled before allowing reactions to proceed without further agitation for 30 min at room temperature, protected from light. For some EdU-labeled cells (**Supplementary Figs. 4** and **6**), and all EU-labeled cells (**Fig. 3**), a total of two rounds of CuAAC reactions (using fresh reaction components) were applied in succession. Following the reaction, cells were placed on ice and rinsed with ice-cold PBS–BSA solution (3×3 min), followed by rinsing with ice-cold cacodylate buffer (2×3 min) before proceeding to the blocking step.

Copper-free click chemistry reactions. Prior to exposure of cells to IRDye700DX-DIBAC, free thiols were alkylated by treatment of fixed cells with a solution of *N*-ethylmaleimide (NEM, 100 mM in PBS) for 45 min at room temperature. Cells were rinsed once in PBS–BSA solution before adding a 4.4 μ M solution of IRDye700DX-DIBAC in PBS (dye was diluted from a 220 uM stock in DMSO). Reactions were allowed to proceed protected from light for 30 m at room temperature. Following the reaction, cells were rinsed with ice-cold PBS–BSA solution (3 × 3 min), followed by rinsing with ice-cold cacodylate buffer (2 × 3 min) before proceeding to the blocking step.

Blocking step. To prevent background precipitation of DAB by endogenous cellular components, cells were treated for 30 min on ice in a solution of 0.1 M sodium cacodylate buffer (pH 7.4) containing 50 mM glycine, 10 mM KCN, 5 mM aminotriazole, and 1 μ l of 30% H₂O₂ per 30 mL blocking solution.

Confocal fluorescence imaging and photooxidation. Cells were imageg and photooxidized using an inverted confocal microscope outfitted with a stage chilled to 4 °C. Confocal fluorescence and transmitted light images were taken with minimum exposure to identify transfected cells for correlative light microscopic imaging. For photooxidation, DAB (Sigma-Aldrich) was dissolved

in 0.1 N HCl at a concentration of 5.4 mg/ml and subsequently diluted tenfold (v/v) into sodium cacodylate buffer, pH 7.4 (with a final buffer concentration of 0.1 M), mixed, and passed through a 0.22- μ m syringe filter. DAB solutions were freshly prepared on the day of photooxidation and placed on ice and protected from light before being added to the cells. Regions of interest were identified by fluorescence, and images were recorded with minimal excitation of the sample to avoid photobleaching. For photooxidation, samples were illuminated through an appropriate bandpass filter (as indicated for each dye in **Supplementary Fig. 2b**) with intense light from a 150 W xenon lamp. In our initial screen of potential click-EM dyes we used a fixed illumination period of 5 min. Samples processed for EM analysis were illuminated for 5–10 min, depending on the initial fluorescence intensity, the brightness of the illumination, and the optics used.

Staining of cells for TEM and imaging. Multiple areas on a single dish were photooxidized as described in the preceding section. Subsequently, cells were placed on a bed of ice and washed using ice-cold cacodylate buffer ($5 \times 2 \min$) to remove unpolymerized DAB. After washing, cell were post-fixed with 1% osmium tetroxide (Electron Microscopy Sciences) in 0.1 M sodium cacodylate buffer for 30 min on ice, then washed with ice-cold cacodylate buffer (3 × 2 min) and rinsed once in ice-cold distilled water. The samples were then dehydrated with an ice-cold graded ethanol series (20% (v/v), 50% (v/v), 70% (v/v), 90% (v/v), 100% (anhydrous), and 100% (anhydrous) for 3 min each) and washed once with room-temperature 100% (anhydrous) ethanol for 3 min. Samples were then infiltrated with Durcupan ACM resin (Electron Microscopy Sciences, prepared according to the manufacturer's protocol) using a 1:1 solution of anhydrous ethanol/resin for 30 min on a platform with gentle rocking, then with 100% resin overnight with rocking. The next day, the resin was removed from dishes (by decanting and gentle scraping with care to avoid touching cells), replaced with freshly prepared 100% resin (3 \times 30 min with rocking), and polymerized in a vacuum oven at 60 °C for 48 h. Subsequently, photooxidized areas of interest were identified using a dissecting microscope, sawed out using a jeweler's saw, and mounted on dummy acrylic blocks with cyanoacrylic adhesive. The coverslip was carefully removed and ultrathin sections (80-nm thick) were cut using an ultramicrotome. Electron micrographs were recorded using a JEOL 1200 EX transmission electron microscope operating at 60 kV.

Preparation of staining solutions for SBEM analysis. The following solutions were prepared on day 1 of the staining procedure immediately before use.

Solution A. A solution containing 3% (w/v) potassium ferrocyanide and 6mM CaCl₂ in 0.2 M cacodylate buffer was freshly prepared and mixed (1:1, v/v) with a 4% (w/v) aqueous solution of osmium tetroxide (Electron Microscopy Sciences). This solution was chilled on ice before use.

Solution B. A fresh solution of thiocarbohydrazide was prepared (while cells were incubating Solution A, see details below) by adding 100 mg thiocarbohydrazide (Ted Pella) to 10 ml ddH₂O. The solution was then heated in a 60 °C oven for 30 min with gentle agitation and swirling every 10 min to ensure dissolution of the material.

Solution C. 2% (w/v) osmium tetroxide was prepared by mixing equal volumes of ddH_2O and a 4% (w/v) aqueous solution of osmium tetroxide.

Solution D. 2% uranyl acetate (w/v) was prepared by dissolution in ddH_2O . The following solutions were prepared on day 2 of the staining procedure immediately before use.

Solution E. 0.998 g L-aspartic acid (Sigma-Aldrich) was dissolved in 250 ml ddH₂O.

Solution F. 0.066 g lead nitrate was dissolved in 10 ml Solution E, and the solution pH was adjusted to 5.5 by addition of 1 N KOH dropwise. The solution was then heated in a 60 °C oven for 30 min until completely dissolved.

Staining of cells for SBEM and imaging. Multiple areas on a single dish were photooxidized as described in the preceding section. Subsequently, unpolymerized DAB was removed from cells by rinsing cells with ice-cold cacodylate buffer ($5 \times 2 \min$) on a bed of ice. Cells were then incubated in Solution A on ice for 30 m while Solution B was prepared. At the end of the 30-min incubation, Solution A was removed and cells were rinsed ($5 \times 2 \min$) in ddH₂O at room

temperature and subsequently incubated in Solution B at room temperature. After 10 min, Solution B was removed and cells were rinsed (5 × 2 min) with ddH₂O at room temperature and subsequently incubated in Solution C. After 10 min, Solution C was removed and cells were rinsed (5 × 2 min) with ddH₂O at room temperature and subsequently incubated with Solution D overnight in a refrigerator kept at ~4 °C.

The following day cells were rinsed (5 × 2 min) in ddH₂O at room temperature before *en bloc* Walton's lead aspartate staining was performed by incubating cells in warm Solution F for in a 60 °C oven. After 10 min, cells were removed from the oven and rinsed (5 × 2 min) in ddH₂O at room temperature. The cells were then subjected to dehydration using a series of ice-cold ethanol solution in water for 3 min each on a bed of ice in the following order: 20% (v/v), 50% (v/v), 70% (v/v), 90% (v/v). Subsequently, the cells were further dehydrated using two washes in ice-cold 100% (anhydrous) ethanol for 3 min each, then removed from the bed of ice and washed once in room temperature 100% (anhydrous) ethanol for 3 min. Samples were then infiltrated with Durcupan ACM resin (Electron Microscopy Sciences, prepared according to the manufacturer's protocol) using a 1:1 solution of anhydrous ethanol/resin for 30 min on a platform with gentle rocking, then with 100% resin overnight with rocking. The next day, the resin was removed from dishes (by decanting

and gentle scraping with care to avoid touching cells), and replaced with freshly prepared 100% resin (3×30 min with rocking). Cells were then submerged using a small amount of resin and covered with a square piece of Aclar film to create a flat surface. The resin was allowed to polymerize in a vacuum oven at 60 °C for 48 h. Finally, photooxidized areas of interest were identified, removed with a jeweler's saw, and mounted for SBEM imaging with a Gatan 3View system (Gatan Inc.) incorporated into a Zeiss Merlin Compact Scanning Electron Microscope (Zeiss), and images were recorded at 60-nm cutting intervals.

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