Proximity-dependent protein labeling provides a powerful in vivo strategy to characterize the interactomes of specific proteins. We previously optimized a proximity labeling protocol for Caenorhabditis elegans using the highly active biotin ligase TurboID. A significant constraint on the sensitivity of TurboID is the presence of abundant endogenously biotinylated proteins that take up bandwidth in the mass spectrometer, notably carboxylases that use biotin as a cofactor. In C. elegans, these comprise POD-2/acyl-CoA carboxylase alpha, PCCA-1/propionyl-CoA carboxylase alpha, PYC-1/pyruvate carboxylase, and MCCC-1/methylcrotonyl-CoA carboxylase alpha. Here, we developed ways to remove these carboxylases prior to streptavidin purification and mass spectrometry by engineering their corresponding genes to add a C-terminal His$_{10}$ tag. This allows us to deplete them from C. elegans lysates using immobilized metal affinity chromatography. To demonstrate the method’s efficacy, we use it to expand the interactome map of the presynaptic active zone protein ELKS-1. We identify many known active zone proteins, including UNC-10/RIM, SYD-2/liprin-alpha, SAD-1/BRSK1, CLA-1/CLArinet, C16E9.2/Sentryn, as well as previously uncharacterized potentially synaptic proteins such as the ortholog of human angiomotin, F59C12.3 and the uncharacterized protein R148.3. Our approach provides a quick and inexpensive solution to a common contaminant problem in biotin-dependent proximity labeling. The approach may be applicable to other model organisms and will enable deeper and more complete analysis of interactors for proteins of interest.

Protein–protein interactions underlie most biological processes. Methods that highlight such interactions can provide key insights into basic molecular mechanisms and help target strategies for therapeutic interventions (1). One such method is proximity-dependent protein labeling, which can both characterize protein–protein interaction networks and define the proteomes of subcellular structures (1, 2). The proximity labeling enzymes most widely used in living cells and organisms are currently engineered variants of the Escherichia coli biotin ligase BirA, such as TurboID, the ascorbate peroxidase APEX and its derivatives, and horseradish peroxidase (HRP) (1–3). All these enzymes release activated biotin, which reacts with amine groups of exposed lysine residues in the vicinity (4–6).

Biotin is a critical cofactor of carboxylation reactions. Biotin-dependent carboxylases, notably acetyl-CoA carboxylase, propionyl-CoA carboxylase (PCC), 3-methylcrotonyl-CoA carboxylase (MCC), and pyruvate carboxylase (PC), are widespread in nature and important for the metabolism of fatty acids, carbohydrates, and amino acids. In mammals, which do not synthesize or store biotin, biotin deficiency causes severe syndromes, including ataxia and neurological dysfunction (7, 8). Biotin is covalently linked to the carboxylase apoproteins by holocarboxylase synthetase (7). The nematode Caenorhabditis elegans has one ortholog of holocarboxylase synthetase, BPL-1 (biotin protein ligase 1), which biotinylates the carboxylases MCC-1/MCCC (methylcrotonoyl coenzyme A carboxylase), PCCA-1/PCCA (propionyl coenzyme A carboxylase alpha subunit), POD-2/ACACA (acyetyl coenzyme A carboxylase), and PYC-1/PC (9). These four endogenously biotinylated carboxylases, from here on referred to collectively as MP3, and their orthologs, dominate the biotinylated proteome of many organisms, including worms, flies, mice, and humans. MP3 proteins provide by far the most abundantly detected peptides when streptavidin affinity-purified samples are analyzed by mass spectrometry (MS) (10–12).

High abundance proteins (HAPs) in biological samples limit detection of low abundance proteins (LAPs) by MS (13). Significant effort has been spent developing ways to deplete HAPs from different samples to increase detection of LAPs by MS. For example, collagenase depletion of collagen significantly increases the number and variety of proteins detected from extracellular matrix–derived protein extracts (14). Albumins, immunoglobulins, fibrinogen, transferrin, and haptoglobin comprise more than 90% of the plasma proteome and therefore mask the detection of LAPs. Depleting albumins and immunoglobulins by affinity chromatography prior to MS analysis makes it possible to identify many LAPs in blood plasma (15). EDTA-functionalized nanoparticles effectively double the number of proteins detected in urine samples (16).

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Extraction protocols that favor recovery of LAPs over HAPs (beta-conglycinin and glycinin) from soybean seeds allow more LAPs to be identified (17).

Proximity labeling has been applied extensively in cultured cells but its use in C. elegans is in its infancy. Overexpressing a patrocinin:TurboID fusion protein in the worm gut successfully highlighted the interactome of this microtubule-binding protein (18). Fusing TurboID to an anti-GFP nanobody allowed tissue-specific analysis of the interactomes of the GFP-tagged centrosomal proteins PLK-1 and SPD-5 (19). We identified interactors of the presynaptic active zone protein ELKS-1 by knocking in a TurboID:mNeongreen cassette at the elks-1 locus (10). We also found that knock-ins that express TurboID fusions at endogenous levels enhance the number of interacting partners identified compared with transgenically over-expressing the corresponding fusion. However, for proteins of interest (POIs) expressed at low levels, or in a small number of cells, detecting biotinylated proximal interactors by MS becomes difficult because of the abundant copurifying MP3 proteins. Here, we devise a way to deplete MP3 proteins prior to streptavidin-based affinity purification, thus increasing the signal/noise ratio for TurboID-based proximity labeling in C. elegans. Using ELKS-1 as our POI, we show that depleting MP3 enables many more known or putatively synaptic proteins to be identified compared with analysis of undepleted controls.

Results

To increase the signal/noise ratio in biotin-dependent proximity labeling, we first sought to prevent endogenous MP3 biotinylation. To achieve this, we tried to deplete BPL-1, the enzyme that biotinylates MP3. BPL-1 is required for efficient de novo fatty acid biosynthesis, and deleting bpl-1 results in maternal-effect embryonic lethality (9). To conditionally disrupt bpl-1, we knocked in an auxin-inducible degron (AID) just upstream of the bpl-1 termination codon. Incubating bpl-1::AID worms with auxin from the egg, L1, or L2 stages arrested their growth. By contrast, worms transferred to auxin plates from the mid-L3 stage and onward grew to adulthood. However, when we harvested these animals as young adults and probed Western blots of their extracts with streptavidin, we did not observe reduced biotinylation of MP3 (Fig. S1A). Incubating bpl-1::AID worms on auxin plates for longer periods, for 48 or 64 h post-YA stage, depleted biotinylated POD-2 but not the other three proteins (Fig. S1B).

Since knocking down BPL-1 only had marginal effects on MP3 biotinylation, we sought to deplete each carboxylase individually. Using CRISPR/Cas9-mediated genome editing, we tagged each protein with an AID at its C terminus. mccc-1 and pcca-1 only express one isoform of 73.7 and 79.7 kDa, respectively, whereas pod-2 expresses three isoforms of 91.4, 230.6, and 242.6 kDa and pyc-1 two isoforms of 67.8 and 129.3 kDa. We chose to tag the C termini because they are shared by all isoforms. We grew the mccc-1::3XFLAG-AID; pcca-1::3XFLAG-AID; pyc-1::3XFLAG-AID expressing animals (MP2-deg worms) on auxin plates for several generations but only achieved mild depletion of these three mitochondrial matrix proteins (Fig. S1C). Growing animals that expressed pod-2::3XFLAG-AID as well as MP2::AID (henceforth referred to as MP3::AID) on auxin plates caused embryonic or L1 lethality. Transferring MP3::AID worms from no auxin control plates to auxin plates depletes POD-2::AID within 5 h and caused strong phenotypes but did not alter levels of the other carboxylases (Fig. S1C). Presumably, a 5-h incubation on auxin is insufficient to deplete MCCC-1, PCCA-1, and PYC-1 because they are localized in the mitochondrial matrix. Ubiquitination does occur in different mitochondrial compartments, and mitochondrial proteins can be degraded in a ubiquitin–proteasome system–dependent manner (20). We therefore asked if targeting TIR1 to mitochondria can promote efficient degradation of MP3::AID proteins. We generated an eft-3p::hsp-6 MTS::TIR1::tagBFP (MTS [mitochondrial targeting sequence]) plasmid and used it to make transgenic animals that express mitochondria-localized TIR1::tagBFP. We found, however, that F1 animals expressing TIR-1::tagBFP in mitochondria arrested at the L1 stage, and we abandoned this approach after several unsuccessful attempts to recover edited worms.

Since we were unable to deplete all four carboxylases and their various isoforms satisfactorily using the AID system, we sought a different way to remove biotinylated carboxylases from worm extracts. Immobilized metal affinity chromatography (IMAC), using for example nickel–nitrilotriacetic acid (Ni–NTA) resins, allows efficient purification of proteins bearing a polyhistidine tag on either terminus. IMAC separation offers several advantages: the interaction between the polyhistidine tag and Ni2+ is strong and stable in harsh environments; Ni2+–NTA resins have high protein-binding capacity and can be regenerated; and the resins are relatively cheap (21). We decided to take an IMAC-based approach to deplete MP3 from C. elegans lysates (Fig. 1A). We engineered each of the genes encoding MP3 carboxylases to add His10 tags to their C termini (MP3-His). We then introduced transgenes that express either mNeongreen alone (rab-3p::mNG) or a TurboID:mNeongreen fusion throughout the nervous system (rab-3p::TurboID::mNG) into the MP3-His10 genetic background to optimize an MP3 depletion protocol. Incubating lysates with Ni–NTA resin efficiently depleted MP3 from the samples with minimal loss of other biotinylated proteins (Fig. 1B). Subjecting the samples to two rounds of Ni–NTA resin–based depletion was not necessary to capture most of the His10-tagged proteins (Fig. 1B); a second round of depletion did not significantly impact the MS analysis (data not shown). As expected, the Ni–NTA resin did not capture biotinylated proteins lacking a His10 tag, confirming the specificity of the resin (Fig. S2A).

In previous work, we had engineered the endogenous elks-1 gene, which encodes a synaptic active zone protein, so as to tag ELKS-1 C-terminally with TurboID–mNeongreen. Affinity-purifying extracts from these animals with streptavidin enabled us to identify bona fide synaptic proteins using MS (10). To benchmark our MP3 depletion strategy, we probed the ELKS-1 interactome further. Western blot data showed efficient
depletion of His\textsubscript{10}-tagged MP3 from the samples that we submitted for MS analysis (Fig. 1C). HPLC-UV analysis of peptides obtained from on-bead digestion following streptavidin purification revealed an approximately 10-fold decrease in total peptide concentration in samples depleted of MP3-His\textsubscript{10} compared with undepleted controls (Fig. S2C). To match the amount of proteins analyzed across samples, we injected 5% of the undepleted samples and 50% of the depleted samples into the reverse-phase HPLC column upstream of the MS. We also loaded 5% of the depleted samples to the column for comparison. LC–MS/MS analysis cumulatively identified 2430 proteins present in wildtype, depleted, or undepleted ELKS-1 samples (Table S1).

Using the amica proteomics data analysis platform (22), we found that 180 proteins were enriched and 36 reduced (log\textsubscript{2} ≥1.5-fold change threshold, adjusted \( p \leq 0.05 \)) in undepleted ELKS-1::TbID samples compared with wildtype (Fig. S2C). To match the amount of proteins analyzed across samples, we injected 5% of the undepleted samples and 50% of the depleted samples into the reverse-phase HPLC column upstream of the MS. We also loaded 5% of the depleted samples to the column for comparison. LC–MS/MS analysis cumulatively identified 2430 proteins present in wildtype, depleted, or undepleted ELKS-1 samples (Table S1).

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Further enriched for 214 and 133 proteins and reduced for 19 and 81 proteins, respectively, compared with undepleted ELKS-1::TbID samples (Fig. 2, B–D). Although many previously characterized synaptic proteins were enriched in both depleted and undepleted lists, enrichment was significantly greater in the depleted list (Figs. 2E and S3, A and B). Most, although not all, synaptic proteins already enriched in the undepleted list were further enriched in the depleted list, although, to a lesser extent than proteins that we could not identify without depletion (Fig. 2E).

As well as enrichment of many known synaptic proteins, we identified uncharacterized proteins enriched in the 5% and 50% ELKS-1::TbID depleted samples that we had not identified in the undepleted samples (Table S1 and Fig. 3, A–C). Among these proteins, R148.3 and F59C12.3 stood out because of the low number of spectral counts in the three replicates of undepleted ELKS-1–TbID samples (Table S1).
and Fig. 3, A–C). We expressed mNeongreen translational fusion transgenes for these proteins under the control of the DA9 neuronal promoter *itr-1* and showed that they colocalized with ELKS-1::mScarlet at presynaptic active zones (23) (Fig. 3D). Altogether, our findings suggest that depleting MP3 in *C. elegans* lysates significantly improves the sensitivity of proximity-labeling using TurboID. The biochemical depletion approach we take may be applicable to other model organisms to enhance characterization of protein interaction networks.

Figure 2. Interactome analysis of the presynaptic active zone protein ELKS-1 by comparing carboxylase depleted versus undepleted samples. A, significantly enriched proteins in ELKS-1 undepleted samples compared with wildtype (log₂ ≥1.5-fold change threshold, adjusted *p* ≤ 0.05). As expected, many synaptic proteins are highly enriched. B and C, significantly enriched proteins when data obtained from analysis of 50% (B) or 5% (C) of depleted samples are compared with 5% of undepleted ELKS-1 samples (log₂ ≥1.5-fold change threshold, adjusted *p* ≤ 0.05). D, Venn diagram highlighting number of significantly enriched proteins between samples. See also Fig. S3A. E, table showing log₂ fold enrichment and adjusted *p* value of synaptically annotated proteins. ED: elks-1::TbID::mNG; mp3-His10 carboxylase depleted; WT: mp3-His10; E: elks-1::TbID::mNG carboxylase undepleted.

| Protein  | E-5% vs WT-5% | adj. p-value | ED-50% vs E-5% | adj. p-value | ED-5% vs E-5% | adj. p-value |
|----------|----------------|--------------|----------------|--------------|----------------|--------------|
| TOM-1    | 0.12           | 0.89         | 3.77           | 0.00         | 1.11           | 0.21         |
| UNC-41   | 0.93           | 0.21         | 3.48           | 0.00         | 2.17           | 0.06         |
| UNC-31   | 1.40           | 0.19         | 3.45           | 0.01         | 2.78           | 0.02         |
| SAD-1    | 1.30           | 0.39         | 3.24           | 0.02         | 3.11           | 0.03         |
| RIC-4    | 2.51           | 0.12         | 2.90           | 0.04         | 2.24           | 0.10         |
| AIPR-1   | -0.20          | 0.84         | 2.87           | 0.02         | 2.87           | 0.02         |
| GIT-1    | 0.90           | 0.20         | 2.36           | 0.02         | 1.89           | 0.04         |
| CPX-1    | 4.05           | 0.00         | 2.33           | 0.01         | 1.79           | 0.01         |
| C16E9.2  | 1.49           | 0.15         | 2.05           | 0.04         | 1.32           | 0.32         |
| CLA-1    | 8.72           | 0.00         | 2.04           | 0.01         | 0.56           | 0.32         |
| SYD-2    | 5.33           | 0.00         | 1.85           | 0.03         | 1.67           | 0.07         |
| SIPA-1   | 4.30           | 0.00         | 1.63           | 0.01         | 1.74           | 0.01         |
Discussion

In *C. elegans* and other species, four endogenously biotinylated carboxylases, MCCC-1/MCCC, PCCA-1/PCCA, POD-2/ACACA, PYC-1/PC (MP3), dominate the biotinylated proteome. Peptides from these large and abundant carboxylases obscure peptides from less abundant biotinylated proteins when samples affinity-purified on streptavidin beads are analyzed by MS. Using CRISPR/Cas9 genome editing, we tagged the MP3 proteins with a His10 tag, allowing us to deplete them using affinity purification and thereby substantially increasing the sensitivity of MS analysis. We expect depletion will be especially useful when the POI tagged with TurboID is expressed at low levels or only in a subset of cells.

Depleting the carboxylases not only reduces noise by removing confounding peptides but also increases signal as it allows more of the sought after peptides to be loaded onto the mass spectrometer without saturating the run.

Adding a depletion step to TurboID experiments using the active zone protein ELKS-1 highlights many proteins known to be synaptically enriched that are not highlighted when we omit depletion. These proteins include SEL-5/AAK1, an ortholog of human AP2-associated kinase 1 (24); UNC-41/stonin, which functions with the AP2 complex to recycle synaptic vesicle membranes (25); RIC-8, which localizes to cholinergic synapses along the ventral nerve cord (26); the aryl hydrocarbon receptor–interacting protein (AIP)–related protein AIPR-1,
which physically interacts with ryanodine receptor at worm synapses (27); UNC-31/CAPS; PAK-1/PAK1; SAD-1/Sad kinase; synaptobrevin SNB-1; SYD-2/ripin-alpha; NAB-1/PPP1R9A; GIT-1/GIT1; RIC-4/SNAP23; and TOM-1/STXB5. MP3 depletion also highlights ELKS-1 interactors whose enrichment at synapses is less well documented. We confirmed the synaptic localization of two such proteins, R148.3 and F59C12.3. R148.3 is predicted to be a trans membrane protein with both extracellular and intracellular domains. When we express R148.3::mNG in DA9 neurons, we observe strong synaptic colocalization with ELKS-1::mScarlet (Fig. 5D). A previous study reports that R148.3 is expressed in neurons, pharynx, body wall muscles, and vulva and is an ortholog of human FKBP15 (28). Knockdown of R148.3 decreases fat accumulation and shortens lifespan (28); however, the function of R148.3 remains to be elucidated. F59C12.3 is predicted to be a 548-residue protein orthologous to human AMOT (angiomotin). The human Angiomotin family is a group of scaffolding proteins associated with tight junctions (29). AMOT-p130 accumulates at postsynaptic densities in rat hippocampal neurons and interacts with actin, the multiple PDZ domain proteins MUPP1 and PSD-95 (30). Our colocalization data indicate that AMOT also localizes presynaptically.

The high binding capacity, simplicity, and low price of IMAC resins are significant advantages to their use for MP3 depletion. However, there are also some limitations that should be considered before employing this approach. IMAC resins are often incompatible with buffers containing SDS, EDTA, or DTT, all of which were present in the buffers of our original TurboID protocol (this study; 31). SDS is an anionic detergent used to solubilize and denature proteins and is particularly effective for solubilizing membrane proteins, for example, active zone proteins. We tested several detergents including CHAPS, Tween-20, and Triton X-100 to avoid using SDS. Although capture of MP3 was effective in the absence of SDS, protein extraction was not nearly as effective as when we used the original SDS buffer (Fig. S2B). To overcome this obstacle, we started the protein purification protocol with 1% SDS and then reduced this high concentration using desalting columns and a dilution step in Ni–NTA binding buffer. Schlager et al. (31) removed SDS from protein solutions through precipitation by cooling combined with low concentrations of sarcosyl (32). We tested this protocol but did not achieve an effective level of MP3 depletion (data not shown). In addition to SDS, the reducing agent DTT and chelating agent EDTA also substantially decrease binding of His-tagged proteins to divalent metal-based (mostly nickel or cobalt) resins. DTT reduces and EDTA sequesters the metal ions of an IMAC resin, making them unsuitable for IMAC applications. We therefore excluded EDTA and DTT from our buffers. Affinity purification using short peptide tags, such as FLAG, might provide an alternative to using His tags to deplete MP3 and may not require the removal of SDS, EDTA, or DTT from extraction buffers. As highlighted previously (12), urea-containing buffers should be prepared fresh—MP3 depletion was significantly impaired when we used old 8 M urea Ni–NTA buffer (data not shown). Including low levels of imidazole is usually advised to minimize nonspecific binding to IMAC-based columns or resins (33). Although His-tagged protein purity was not our primary concern, we compared binding of His-tagged proteins and other proteins to Ni–NTA resin in the presence or the absence of 5 mM imidazole in Ni–NTA binding buffer. We did not observe a noticeable change in the bound proteins as judged by Western blot analysis and Coomassie staining and therefore omitted imidazole from our buffers (data not shown).

In summary, His10 tags can be added to endogenously biotinylated carboxylases without disrupting their essential function. The tagged enzymes can then be depleted efficiently using IMAC, providing a reliable way to improve detection and enrichment of biotinylated proteins in proximity labeling experiments. Here, we used depletion with TurboID labeling, but the method is applicable to proximity labeling experiments that use other enzymes to generate reactive biotin, for example, APEX2 or HRP. We have successfully employed this protocol to different POIs, ranging from transcription factors to structural proteins, localized in different subcellular compartments, for example, endoplasmic reticulum, cytoplasm, and nucleus (data not shown). Carboxylase depletion will improve the reach of proximity labeling to single cells and subcellular structures and is likely applicable to many model organisms because of the high evolutionary conservation of these carboxylases.

Experimental procedures

Strains

Worms were grown at room temperature (22 °C) on nematode growth medium (NGM) plates seeded with the biotin auxotroph E. coli MG1655. C. elegans husbandry otherwise followed standard laboratory culture conditions (34). C. elegans strains used in this study include:

N2, the wildtype Bristol strain

AX7884 pod-2(syb1772[pod-2::His10]) II; mccc-1(syb1666[ccc-1::His10]) IV; pyc-1(syb1680[pyc-1::His10]) V; pcca-1(syb1626[pcca-1::His10]) X quadruple His10-tag knock in strain

AX7885 pod-2(syb1772[pod-2::His10]) II; elks-1(syb1710[els-1::CeTurboID:mNeongreen::3xFLAG]) mccc-1(syb1666[ccc-1::His10]) IV; pyc-1(syb1680[pyc-1::His10]) V; pcca-1(syb1626[pcca-1::His10]) X

AX7897 dbls37[rab-3p::mNeongreen::3xFLAG]; pod-2(syb1772[pod-2::His10]) II; mccc-1(syb1666[ccc-1::His10]) IV; pyc-1(syb1680[pyc-1::His10]) V; pcca-1(syb1626[pcca-1::His10]) X

AX7898 dbls24[rab-3p::CeTurboID::mNeongreen::3xFLAG]; pod-2(syb1772[pod-2::His10]) II; mccc-1(syb1666[ccc-1::His10]) IV; pyc-1(syb1680[pyc-1::His10]) V; pcca-1(syb1626[pcca-1::His10]) X

AX8273 bpl-1(db1372[bpl-1::degron(AID)]) iei57[eft-3p::TIR1::mRuby::unc-54 3’UTR+Cbr unc-119(+)] II
AX8489 mcc-1(db1433/mcc-1::degron(AID)::3xFLAG)) IV; pyc-1(db1434/pyc-1::degron(AID)::3xFLAG)) V; pca-1(db1437 [pca-1::degron(AID)::3xFLAG]) X; ieSiS77lefT-3p::TIR1:mRuby::unc-54 3’UTR+Cbr unc-119(+)) II
AX8490 pod-2(db1461/pod-2::degron(AID)::3xFLAG)) II; mcc-1(db1433/mcc-1::degron(AID)::3xFLAG)) IV; pyc-1(db1434/pyc-1::degron(AID)::3xFLAG)) V; pca-1(db1437 [pca-1::degron(AID)::3xFLAG]) X; ieSiS77lefT-3p::TIR1:mRuby::unc-54 3’UTR+Cbr unc-119(+)) II
AX8466 dbEx1339[itr-1p::F59C12.3:mNeongreen; itr-1p::elks-1 cDNA::mScarlet, myo-2p::RFP]
AX8541 dbEx13345[itr-1p::R148.3::mNeongreen; itr-1p::elks-1 cDNA::mScarlet, myo-2p::RFP]

Molecular biology
Cloning uncharacterized ELKS-1 interactors

To generate the pPD95.75-mNG and pPD95.75-mSc plasmids, the pPD95.75 plasmid was digested with KpnI and BsmI restriction enzymes (NEB), and the linearized plasmid gel-extracted and assembled with PCR-amplified codon-optimized mNeongreen or mScarlet using In-fusion cloning (Takara). To generate pPD95.75-itr-1p-mNG or pPD95.75-itr-1p-mSc plasmids, pPD95.75-mNG plasmid was digested with HindIII and PstI, pPD95.75-mSc with PstI and BamHI restriction enzymes (NEB), the linearized plasmids were gel-extracted and assembled with PCR-amplified itr-1 promoter using In-fusion cloning.

The ORF for F59C12.3 (~5.7 kb) and R148.3 (~7.6 kb) was amplified from C. elegans genomic DNA by PCR and cloned into pPD95.75-itr-1p-mNG vector digested with PstI and KpnI by In-fusion cloning. elks-1 complementary DNA (cDNA) (~2.5 kb) was PCR amplified from a C. elegans cDNA library and cloned into pPD95.75-itr-1p-mSc vector digested with BamHI and KpnI by In-fusion reaction. The resulting expression vectors were injected into the gonad of day 1 adult N2 worms at a concentration of 25 ng/μl.

CRISPR/Cas9-mediated genome editing

The AX7884 strain was obtained by crossing strains PHX1772 pod-2(syb1772[pod-2::His10]) II, PHX1666 mcc-1(syb1666/mcc-1::His10]) IV, PHX1680 pyc-1(syb1680[pyc-1::His10]) V, and PHX1626 pce-1(syb1626[pce-1::His10]) X to obtain the quadruple His10-tag KI strain. PHX1772, PHX1666, PHX1680, and PHX1626 were generated by SunyBiotech upon our request. AX8273 was generated by engineering bpl-1 so as to add an AID degron to the C terminus of BPL-1. AX8490 was generated by knocking in sequences that encode an AID degron::3xFLAG just before the stop codon of each of the four carboxylase-encoding genes and using genetic crosses to assemble the four edited genes into a single strain. AX8273 and AX8490 strains were generated according to previously published CRISPR editing protocol (35).

Primers used for cloning

F59C12.3 genomic DNA (~5.7 kb):

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Forward: F59C12.3-GGTTGTTGGAAGGACACACG
TGATACCAGGAGAGACACATTTTAG
Reverse: F59C12.3-CTCCCTCTGACACACCATGGCAGAAA
ATTGGTATTCCGGAAGATTTGAC
R148.3 genomic DNA (~7.6 kb):
Forward: R148.3-GGTGGTGAAGACACACG
CCGCCCCAAATTTGAC
Reverse: R148.3-CTCCCCCTCACGTGGTTGCC
AAGATGTGAGCAAGATCTG
elks-1 cDNA (~2.5 kb):
Forward: elks-1-cDNA-CTGCCCTTTCCTTTGCAACCATTAC
TTCCAGAGTTGGTTCCGAGAC
Reverse: elks-1-cDNA-CTGCCCTTTCCTTTGCAACCATTAC
GGCGGCCCCAATTCGTCAGCATCG
itr-1 promoter for pPD95.75-mScarlet plasmid:
Forward: itr-1p-GAAATAAGCTTGCATGCTTGAC
ATTCCAGAGTTGGTTCCGAGAC
Reverse: itr-1p-CCCTTGCCAAATCCGGGATACCG
TGGTTCCTCACACCATAGAC
itr-1 promoter for pPD95.75-mNeongreen plasmid:
Forward: itr-1p-CAACTTGGAAATGAAATAAGCTT
ATTCCAGAGTTGGTTCCGAGAC
Reverse: itr-1p-GATCCTCTAGAGCTCAGCTACG
GTGCCTTCACACCATAGAC
Codon-optimized mScarlet for pPD95.75 plasmid:
Forward: mScarlet-GGAGGACCCCTGGAGGGATCC
GGTTAGCAACGTCCCAAGGCAG
Reverse: mScarlet-CAGTGGAAATTCTAGGATGTTT
ACTTTGAAAGCTCATCCATCATT
Codon-optimized mNeongreen for pPD95.75 plasmid:
Forward: mNG-GGGAGCCCTTGGAGGGATCC
GTGTCGAAGGGAGAGGCAG
Reverse: mNG-CAGTGGAAATTCTAGGATGTTT
ACCCCTGGAGGGATCC

Light microscopy
Confocal microscopy images of transgenic C. elegans expressing fluorescent proteins were acquired using a Leica SP8 inverted laser scanning confocal microscope with a 63×1.2 numerical aperture oil-immersion objective, using the LAS X software platform (Leica). The Z-project function in ImageJ (Rasband W.S., US National Institutes of Health; http://rsweb.nih.gov/ij/) was used to obtain the figures used in the panels. Animals were mounted on 2% agarose pads and immobilized with 100 μM of sodium azide.

AID
AID assays were conducted as described previously (36). Briefly, age-synchronized animals were grown on biotin auxotroph E. coli strain MG1655-seeded NGM plates in the presence or the absence of 1 mM auxin (indole-3-acetic acid; Sigma).

Immunoblotting
Synchronized populations of C. elegans grown on E. coli MG1655 were harvested at L4 or young adult stage, washed...
three times in M9 buffer, and flash-frozen after adding 4× Bolt LDS sample buffer supplemented with fresh DTT. The samples were then thawed, boiled for 10 min at 90 °C, vortexed mildly for 10 min, centrifuged for 30 min at 15,000 rpm at 4 °C, and the supernatant was collected. Proteins were transferred to a polyvinylidene difluoride membrane (Thermo Fisher Scientific) following electrophoresis using Bolt 4 to 12% Bis–Tris Plus gels (Thermo Fisher Scientific). Membranes were blocked for 1 h at room temperature with 1% casein blocking buffer and incubated for 1 h at room temperature with fluorescently labeled streptavidin or with HRP-conjugated antibodies. Membranes were then washed three times with Tris-buffered saline with Tween-20 (TBS-T). The following antibodies or protein–HRP conjugates were used for this study: IRDye 800CW Streptavidin (1:10,000 dilution in casein) (LI-COR Biosciences), anti-FLAG M2-peroxidase (1:5000 dilution in 1% casein buffer) (catalog no.: A8592; Sigma), anti–alpha tubulin–HRP (1:10,000 dilution in 1% casein buffer) (DM1A; Abcam; catalog no.: ab40742). Membranes were imaged using ChemiDoc Imaging System (Model MP; Bio-Rad).

**TurboID-based enzymatic protein labeling and extraction of biotinylated proteins from C. elegans**

Gravid adult *C. elegans* were bleached, and the eggs were transferred to NGM plates seeded with *E. coli* MG1655 to obtain synchronized populations of worms. The animals were harvested at L4 or young adult stage, washed three times in M9 buffer, incubated at room temperature (22 °C) in M9 buffer supplemented with 1 mM biotin and *E. coli* MG1655 for 2 h unless stated otherwise. Two hours later, the worms were washed three times in M9 buffer and allowed to settle on ice after the last wash. After completely aspirating the M9 buffer, one volume of 2× TBS buffer supplemented with 1 mM PMSF and COmplete EDTA-free protease inhibitor cocktail (Roche Applied Science) was added to one volume of packed worms and then added dropwise to liquid nitrogen to obtain frozen worm “popcorn.” A Spex 6875D cryogenic mill was used to grind frozen *C. elegans* to a fine powder, which was then stored at −80 °C. Worm powder was thawed in a 50 ml falcon tube while rolling it on a tube roller at room temperature. After the sample was completely thawed, it was centrifuged (1000 rpm, 1 min) to collect the sample at the bottom of the tube. SDS was added to the sample to a final concentration of 1% from a stock solution of 20% SDS. The tubes were gently inverted a few times and immediately incubated at 90 °C for 5 min. After heat treatment, the samples were sonicated continuously for 1 min twice, with brief cooling between the two sonication steps. Sonication used a probe sonicator microtip (QSonica 700; microtip 4417, 1.6 mm and an amplitude setting of 50/maximum). The samples were cooled to room temperature following sonication and adjusted to 2 M urea using a stock solution (8 M urea, 1% SDS, and TBS buffer). The samples were then centrifuged at 60,000 RPM for 30 min at 22 °C using a benchtop ultracentrifuge Optima MP and MLA-80 rotor (Beckman), and the clear supernatant between the pellet and surface lipid layer transferred to a new tube.

Zeba spin desalting columns (7 K molecular weight cutoff) (Thermo Fisher Scientific) were equilibrated three times with 5 ml TBS buffer containing 2 M urea, freshly supplemented with protease inhibitors (Roche cOmplete EDTA-free protease inhibitor cocktail one tablet/25 ml; PMSF 1 mM) by centrifugation at 1000g for 5 min (or until the buffer was completely eluted from resin). Around 4 ml of clarified sample was then loaded onto the equilibrated spin column and desalted by centrifugation at 1000g for 5 min (or until the sample was completely eluted from resin) to remove free biotin. The desalting step was repeated once more using freshly equilibrated columns. Protein concentration in the samples was measured using Pierce 660 nm protein assay reagent supplemented with Ionic Detergent Compatibility Reagent (Thermo Fisher Scientific).

**IMAC-mediated depletion of His-tagged carboxylases**

One milliliter of PureCube Ni–NTA agarose resin slurry (Cube Biotech) was transferred to a 15 ml Falcon tube, centrifuged for 1 min at 2000 rpm, and the supernatant was removed. The Ni–NTA agarose resin slurry was equilibrated twice for 10 min in 10 ml of Ni–NTA binding buffer (100 mM NaH2PO4, 10 mM Tris–HCl, 8 M urea, pH 8.0) before mixing it with the protein lysate. One volume of total protein lysate (containing 12.5 mg of proteins) was mixed with three volumes of Ni–NTA binding buffer in a Falcon tube and mixed with equilibrated Ni–NTA resin. The mix was incubated for 2 h at room temperature using a tube roller. The tube was centrifuged at 2000 rpm for 1 min, and the supernatant was transferred to a new Falcon tube. Note: extra care must be paid not to transfer Ni–NTA resin along with the supernatant. To confirm efficient carboxylase depletion, the Ni–NTA resin was processed for His-tagged protein purification and Western blot analysis.

**His-tagged protein purification**

After completely removing protein supernatant in Ni–NTA buffer, the Ni–NTA resin was washed twice by adding 10 ml of Ni–NTA buffer and rotating for 10 min on a tube rotator. After the washes, the Falcon tube was centrifuged at 2000 rpm for 1 min and the buffer was aspirated. The resin was transferred to an Eppendorf tube by mixing with 1 M imidazole solution (1:1 ratio V:V) for elution of His-tagged proteins from the resin (a small aliquot can be processed to determine carboxylase depletion).

The tube was incubated at room temperature for 30 min using a rotator.

The Eppendorf tube was centrifuged at 2000 rpm for 1 min, and the supernatant was transferred into a new tube for Western blot analysis.

**Streptavidin magnetic bead acetylation**

Pierce Streptavidin magnetic beads (Thermo Fisher Scientific; 100 μl bead slurry per 12.5 mg of *C. elegans* total protein...
lysate) were washed three times with 1 ml of buffer 1 (50 mM Hepes–NaOH, pH 7.8, and 0.2% Tween-20), briefly incubating them in the buffer and using magnetic separation to retain beads while discarding buffer. Washed beads were resuspended in a mix of 190 μl of buffer 1 and 10 μl of 100 mM Pierce Sulfo-NHS-Acetate (Thermo Fisher Scientific; dissolved in dimethyl sulfoxide), and incubated for 1 h at room temperature to acetylate free amines (37). Beads were washed three times with 1 ml of buffer 2 (50 mM ammonium bicarbonate and 0.2% Tween-20) to stop the reaction.

**Biotinylated-protein pulldown and elution**

Carboxylase-depleted supernatant was mixed with acetylated magnetic beads in a 15 ml Falcon tube and incubated overnight on a tube roller at room temperature. To collect beads, a neodymium magnet was taped to the side of the Falcon tube and incubated on a rocking platform for 1 h or until all the magnetic beads bound to the magnet. Unbound lysate was aspirated, and the beads were transferred to a 2 ml LoBind protein tube (Eppendorf) using 2% SDS wash buffer

MS

Beads were resuspended in 50 μl 1 M urea and 50 mM ammonium bicarbonate. Disulfide bonds were reduced with 2 μl of 250 mM DTT for 30 min at room temperature before adding 2 μl of 500 mM iodoacetamide and incubating for 30 min at room temperature in the dark. Remaining iodoaceticamide was quenched with 1 μl of 250 mM DTT for 10 min. Proteins were digested on the beads with 150 ng LysC (Wako) adding 2 μl of 250 mM DTT for 30 min at room temperature in the dark. Remaining iodoacetamide was quenched with 1 μl of 500 mM iodoacetamide and incubating for 1 h at room temperature to acetylate free amines (37). Beads were washed three times with 1 ml of buffer 2 (50 mM ammonium bicarbonate and 0.2% Tween-20) to stop the reaction.

**Data analysis using R scripts**

MaxQuant output tables were further processed in R (R Core Team, 2018) (40). Reverse database identifications, contaminant proteins, protein groups identified only by a modified peptide, protein groups with less than three quantitative values in one experimental group, and protein groups with less than two razor peptides were removed for further analysis. Missing values were replaced by randomly drawing data points from a normal distribution modeled on the whole dataset (data mean shifted by −1.8 standard deviations, width of distribution of 0.3 standard deviations). Differences between groups were statistically evaluated using the LIMMA package (41) at 5% false discovery rate (Benjamini–Hochberg).

**Data availability**

Proteomics data deposition: the MS proteomics data have been deposited at the ProteomeXchange Consortium via the PRIDE partner repository (42) with the dataset identifier PXD034639.

**Ethics**

The work used the free-living nematode *C. elegans*, for which there is no requirement for review and approval from an institutional animal care and use committee. Transgenic
experiments were carried out following Institute of Science and Technology Austria guidelines for such work.

Supporting information—This article contains supporting information.

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Abbreviations—The abbreviations used are: ACACA, acetyl coenzyme A carboxylase; AID, auxin-inducible degron; BPL-1, biotin protein ligase 1; cDNA, complementary DNA; HAP, high abundance protein; HRP, horseradish peroxidase; IMAC, immobilized metal affinity chromatography; LAP, low abundance protein; MCC, 3-methylcrotonyl-CoA carboxylase; MCCC, methylcrotonyl coenzyme A carboxylase; MS, mass spectrometry; NGM, nematode growth medium; Ni–NTA, nickel–nitritolriatic acid; PC, pyruvate carboxylase; PCC, propionyl-CoA carboxylase; PCCA, propionyl coenzyme A carboxylase alpha subunit; POL, protein of interest; TBS-T, Tris-buffered saline with Tween-20.

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