Longevity is dictated by a combination of environmental and genetic factors. One of the key mechanisms to regulate life-span extension is the induction of protein chaperones for protein homeostasis. Ectopic activation of the unfolded protein response of the endoplasmic reticulum (UPRER), specifically in neurons is sufficient to enhance organismal stress resistance and extend life span. Here, we find that this activation not only promotes chaperones but also facilitates ER restructuring and ER function. This restructuring is concomitant with lipid depletion through lipophagy. Activation of lipophagy is distinct from chaperone induction and is required for the life-span extension found in this paradigm. Last, we find that overexpression of the lipophagy component, ehpb-1, is sufficient to deplete lipids, remodel ER, and promote life span.Therefore, UPR induction in neurons triggers two distinct programs in the periphery: the proteostasis arm through protein chaperones and metabolic changes through lipid depletion mediated by EH domain binding protein 1 (EHBPP-1).

INTRODUCTION
The homeostatic regulation of protein folding (proteostasis), which is monitored in specific subcellular compartments, is an integral player in stress resistance and longevity. The endoplasmic reticulum (ER), in particular, is a central regulator of stress monitoring as it controls nearly a third of the cell’s proteins, provides an internal medium for lipid homeostasis and cell signaling, and communicates directly with all other organelles to maintain cellular secretion. Thus, cells have evolved numerous quality control machineries dedicated to protecting the ER both under basal and stressed conditions.

Notably, the ER has evolved three primary branches of its unfolded protein response (UPRER) to maintain proper secretion, protein folding, and lipid homeostasis. The most studied of the three UPRER branches involves the membrane localized/endoribonuclease inositol-requiring protein-1 (IRE-1), which is activated via oligomerization upon unfolded protein stress in the ER lumen or lipid disequilibrium in the ER membrane. This results in splicing of a specific intron from the mRNA of the transcription factor, X-box binding protein-1 (XBP-1), to create the spliced version of xbp-1 (xbp-1s), which then up-regulates the expression of protein degradation, protein folding, and lipid metabolism gene targets. Another branch of the UPRER is the membrane-localized protein kinase, R (PKR)-like ER kinase (PERK), which, upon ER stress, globally reduces protein translation via phosphorylation of the translation initiation factor, eukaryotic initiation factor 2 (eIF2α). Transcripts that promote protein folding and apoptosis, however, are preferentially translated. The last of the three UPRER branches is the ER membrane-localized activating transcription factor-6 (ATF6), which, upon luminal stress, moves to the Golgi to become a mature transcription factor capable of up-regulating canonical proteostasis genes (1, 2).

While these pathways have been intensively studied for the past two decades, much less is known about the adaptive responses of the ER under long-lived conditions. The small nematode, Caenorhabditis elegans, is an excellent model to simultaneously study the genetic underpinnings of cell-cell communication under adaptive conditions while also enabling the monitoring of its cell biology and organismal health (e.g., life span and stress resistance). Work with C. elegans has shown that its cells become less capable in protein folding and also less able to induce stress responses to proteotoxicity with advanced age (3–9). Overexpression of xbp-1s, specifically in neurons (heretofore referred to as neuronal xbp-1s), extends organismal life span and increases ER stress tolerance in a cell nonautonomous manner (9). While the precise, small ER stress signal was not identified, small clear vesicles are required for this beneficial effect, which could be host to numerous neurotransmitters. The intestine, the putative target tissue in this paradigm, induces chaperones in an XBP-1–dependent manner. However, it is well understood that UPRER induction not only alters protein homeostasis through chaperones but also is important mechanism for metabolic change and handling of lipid stress (10).

We hypothesized that induction of the UPRER in neurons, which reverses the age-dependent loss of ER proteostasis, also enacts a marked restructuring of ER morphology, which, in turn, imparts a beneficial metabolic change and promotes longevity. Although whole-organismal metabolic restructuring has been a topic of intense study in the aging field, much less is known about the adaptive responses of organelles in long-lived conditions. Here, we find that neuronal xbp-1s animals have notable ER restructuring and lipid depletion. These changes are distinct from chaperone induction and are dependent on the conserved lipophagy complex, RAB-10/EHBP-1/RME-1 (ras-related GTP-binding protein/EH domain binding protein/receptor mediated endocytosis). Thus, we argue that the beneficial effects of nonautonomous UPRER are dependent on two independent, yet equally important, arms of UPRER: the protein...
homeostasis arm, including chaperone induction and the metabolic arm, which induces ER remodeling and lipophagy.

**RESULTS**

Ectopic UPR\textsuperscript{ER} induction results in ER remodeling

To determine whether preserving ER protein homeostasis would alter age-associated changes in the ER structure, we interrogated ER morphology and organization in the peripheral tissue (intestine) of neuronal \textit{xbp-1s} animals (Fig. 1A). We visualized the ER during various stages of adulthood in \textit{C. elegans} using an mCherry::HDEL fusion protein that localizes to the ER lumen through a SEL-1 (Suppressor/Enhancer of Lin-12) signal sequence. At day 1 of adulthood, the ER fills the intestinal lumen and can be visualized as a sponge-like, interconnected, and uniform structure. As early as day 4, intestinal ER is less uniform and begins to collapse, forming disorganized, aggregated...
structures with increased severity by day 7. Neuronal xbp-1s animals display no difference in intestine ER structures at day 1 compared with wild-type animals. Unexpectedly, at day 4, when wild-type animals begin to exhibit collapse of the ER, neuronal xbp-1s animals have a spherical ER morphology that is not seen at any age in wild-type animals. These structures are visible as early as day 2; by day 7, these structures disappear, and the ER network collapses in a manner indistinguishable from wild-type animals. Therefore, constitutive UPRER in neurons is not sufficient to protect ER quality and structure during aging and only show significant differences in early adulthood.

To better understand the nature of ER remodeling in neuronal xbp-1s animals, we performed high-magnification transmission electron micrograph imaging and identified denser patches of rough ER compared with controls and an increase in the number of ER whors (Fig. 1, B and C). ER whors have been previously observed upon up-regulation of UPRER/ER-phagy (11–13). To ensure that the changes in ER structure and organization were not due to artifacts of the imaging methods, we interrogated ER morphology in neuronal xbp-1s animals using two additional ER markers: mRuby::HDEL localized to the ER through an HSP-4 (heat shock protein) signal sequence (Fig. 1D) and a green fluorescent protein (GFP) fusion to an ER transmembrane protein, SP12 (fig. S1A). Both ER markers showed ER remodeling in neuronal xbp-1s animals similar to mCherry::HDEL and electron microscopy. Moreover, the formation of these structures is partially dependent on xbp-1s in the periphery, as RNA interference (RNAi) knockdown of xbp-1 partially resolved these structures (Fig. 1D and fig. S1A). Strong overexpression of xbp-1s directly in the intestine is sufficient to induce ER remodeling and is dependent on xbp-1 (fig. S1B). However, systemic activation of the UPRER through ER stress by treatment with the chemical agent tunicamycin results in collapsed ER structure distinct from ER re-structuring in neuronal and intestinal xbp-1s (fig. S1C). These data suggest that ectopic xbp-1s expression induces a unique phenomenon distinct from direct activation of UPRER under conditions of proteotoxic stress.

As a critical organelle in secretory vesicle formation, we hypothesized that the major ER remodeling found in neuronal xbp-1s animals could result in increased intestinal secretion. To measure secretion, we compared VIT-2::GFP deposition in neuronal xbp-1s animals compared with control animals. VIT-2::GFP is a fluorescently tagged version of vitellogenin, which is a common reporter used to measure protein secretion as a proxy for ER function in the intestine of C. elegans (14, 15). This protein is integral to maternal yolk formation and, under normal conditions, is secreted solely by the intestine and endocytosed by developing eggs. We find that neuronal xbp-1s animals have increased VIT-2::GFP deposition in their eggs. Moreover, this increase is dependent on xbp-1, suggesting that neuronal xbp-1s animals have a significant increase in ER secretory function in the intestine (Fig. 1, E and F).

**ER remodeling upon UPRER induction is concomitant with lipid depletion**

A massive increase in secretion would be expected to result in lipid depletion due to the expansion of ER membranes. We tested whether intestinal ER remodeling of neuronal xbp-1s animals occurred concurrently with lipid depletion. We measured intestinal lipids using a combination of whole-organism and tissue-specific quantification. Our lab had previously developed a methodology and software, called LAMPro, to exploit underused positional information inherent to BioSorter data, which takes fluorescent traces of many animals at once and defines key regions of interest to various cellular phenomena (16). This enabled us to overcome the practical limits of microscopy (e.g., low n) to reliably measure the distribution of total lipids across animals. Using LAMPro technology, we measured whole-body neutral lipid content (via BODIPY 409/503 staining) of several hundred animals per condition. Briefly, animals were staged when ER remodeling was prominent, fluorescent integrated intensity was measured across the length of BODIPY-stained worms, and data were represented as a heat map of median profiles of relative neutral lipid distribution. Notably, we found that neuronal xbp-1s animals had a marked depletion of lipids compared with wild-type animals (Fig. 2A).

Next, we tested whether the decline in lipid content is due to the loss of lipid droplets (LDs), the primary organelle that stores intestinal lipids in C. elegans (17). We visualized LDs with the intestine-specific, fluorescently labeled, short-chain dehydrogenase DHS-3::GFP fusion protein, which is abundant on the surface of C. elegans intestinal LDs (18, 19). To ensure that LAMPro technology could reliably measure changes in lipid content using DHS-3::GFP, we used sbp-1 RNAi as a positive control for decreased lipid content. sbp-1 is the C. elegans ortholog of SREBP1 (sterol regulatory element–binding protein 1) and is essential for fat accumulation (20), and our LAMPro methods can reliably detect a decrease in lipid levels as expected in sbp-1 knockdown animals (fig. S2). Using this marker, we found that LD content was significantly lower in neuronal xbp-1s animals compared with wild-type animals (Fig. 2, B and C). Last, we find that lipid depletion in neuronal xbp-1s animals was partially dependent on xbp-1, as RNAi knockdown of xbp-1 partially suppressed this phenotype, similar to ER remodeling (Fig. 2, D and E).

**A lipophagy complex driven by EBHP-1 promotes lipid depletion and is required for life-span extension of neuronal xbp-1s animals**

Another study has also presented that neuronal xbp-1s animals have significant lipid depletion, which is dependent on lysosomal expansion and increased activity of lysosomal lipases in the intestine (21), yet independent of the core autophagy machinery. Similarly, we find an expansion of lysosomes in neuronal xbp-1s animals and that these lysosomes colocalize with distinct ER membranes that occur during ER remodeling (fig. S3, A and B). Under conditions of UPRER activation and ER expansion, induction of ER phagy and lipophagy can aid in the redistribution of lipids (12, 22). While Imanikia et al. (21) investigated the role of lysosomal lipases in lipid depletion, here we questioned whether the lipid depletion in neuronal xbp-1s animals was dependent on the elements of the RAB-10/EBHP-1/RME-1 lipophagy complex. We find that RNAi knockdown of ehp-1 fully suppressed the lipid depletion found in neuronal xbp-1s animals, even more than RNAi knockdown of xbp-1 (Fig. 2, D and E). While xbp-1 and ehp-1 RNAi alone both increase lipids in wild-type animals, neuronal xbp-1 RNAi fails to deplete lipids under xbp-1 or ehp-1 knockdown, providing evidence that both xbp-1 and ehp-1 are essential for the lipid depletion phenotype found in this animal (fig. S4). RNAi knockdown of rme-1 only partially suppressed the lipid depletion, and knockdown of rab-10 had no effect. The failure of rme-1 and rab-10 knockdown could be due in part to the fact that they have been associated with induction of autophagy (23). Moreover, knockdown of rab-10 and rme-1 causes phenotypes associated with calorie restriction, including egg laying and growth defects, and an increased life span, which could confound a lipid rescue. Together, these...
data suggest that EHBP-1, potentially through the RAB-10/RME-1/EHBP-1 complex, is involved in the ER remodeling and lipid depletion found in neuronal xbp-1s animals.

Changes in metabolism, specifically lipid depletion, are directly downstream of UPRER induction by xbp-1s. Therefore, we sought to determine whether RAB-10, RME-1, and EHBP-1 were linked to canonical chaperone induction mediated by xbp-1s. To our surprise, RNAi knockdown of any of these lipophagy components did not affect induction of chaperones, as measured by the reporter hsp-4p::GFP (fig. S5, A to D). However, and most unexpectedly, knockdown of...
**DISCUSSION**

Our lab and others have developed cell nonautonomous signaling models (i.e., neuron to intestine) to provide new paradigms in the study of longevity and organismal adaptation to stress. Previously, we developed a model in which neuronal *xbp-1s* overexpression (Fig. 3, A and B). Intestinal *ehbp-1* overexpression also resulted in changes in ER structure, reminiscent of *xbp-1s* overexpression, and displays life-span extension comparably to neuronal *xbp-1s* overexpression (Fig. 3, C and D). However, unlike *xbp-1s* overexpression, overexpression of *ehbp-1* is not sufficient to induce chaperone induction, although there is a mild increase in stress-induced UPR\(^{\text{ER}}\) (Fig. 3, E and F). Therefore, increased lipophagy via EHBP-1 does not promote chaperone induction. These data suggest that overexpression of *ehbp-1* is sufficient to phenocopy the metabolic signaling downstream of *xbp-1s*, providing further evidence that the lipid depletion and extension of life span found in metabolic remodeling through *xbp-1s* signaling are driven by EHBP-1–mediated lipid depletion distinct from canonical chaperone signaling.

When we characterized the intestinal ER morphology of neuronal *xbp-1s* adult animals, we found that the changes in ER structure closely resembled endocytic structures, similar to ER whorls identified in previous studies (24). The major structural change in the ER can be due to two potential reasons: increased secretion from the ER or increased autophagic flux of the ER in these animals. ER whorls...
neuronal another study has identified that lysosomal expansion found in animals colocalized with lysosomal markers and
et al 6 ,

have been identified during conditions of hypersecretion (25, 26) and during stress for ER turnover (24). Here, we found that neuronal xbp-1s animals exhibited increased secretion as measured by vitellogenin secretion, and thus, it is possible that changes in ER morphology occur during early to mid-adulthood to keep up with the secretory demands of egg formation.

A second possibility in the changes to intestinal ER morphology is through increased autophagy. Autophagy has been observed to be up-regulated in response to ER stress (27–29). Specifically, induction of ER stress, either through phenobarbital treatment, or Tm treatment, or induction of unstable dimers of membrane proteins in the ER, is associated with a concomitant ER expansion and induction of autophagy (11–13, 26, 30). We find that the unique ER structures observed in neuronal xbp-1s animals colocalized with lysosomal markers and that these animals exhibited an expansion of lysosomes. Similarly, another study has identified that lysosomal expansion found in neuronal xbp-1s animals is critical for the life-span extension of this

paradigm (21). Therefore, it is possible that there is also an increase in autophagic flux of the ER, which may also contribute to the ER remodeling and lysosomal expansion.

To accommodate for the ER expansion and remodeling, the adaptive UPR ER stress response may also massively up-regulate gene expression of membrane lipid enzymes. For example, ectopic expression of xbp-1s increases lysosomal lipases and δ9 desaturases to drive lipid depletion (21). Here, we find that lipid depletion in neuronal xbp-1s animals is dependent on the conserved lipophagy machinery, RAB-10/EHBP-1/RME-1. Whether this complex works in parallel or in the same molecular pathway as lysosomal lipases has yet to be determined. However, since we find that hyperactivation of lipophagy via overexpression of ehp-b-1 is sufficient to drive all the phenotypes described here—lipid depletion, ER remodeling, and increased life span—independent of chaperone induction, it is likely that the phenomenon described here is independent of the previously described mechanism involving lysosomal lipases and δ9 desaturases. Last, because knockdown of rab-10 and rme-1 causes phenotypes associated with calorie restriction, it is difficult to determine their roles in lipid metabolism. Since overexpression of ehp-b-1 is sufficient to drive lipid depletion while its knockdown is sufficient to increase lipids, it is possible that EHBP-1 can regulate lipid levels outside of its canonical role in lipophagy through the RAB-10/EHBP-1/RME-1 complex.

Last, although this phenotype was described in C. elegans, the phenomena of cell nonautonomous UPR ER transmission—from the brain to the liver, for instance—is a conserved and well-described process (1, 31–33). It is envisioned, due to the conservation of the RAB-10/ EHBP-1/RME-1 complex, that these findings might also apply to understanding the health-promoting effects seen in the liver responding to signals from the nervous system under conditions of activated UPR ER.

METHODS

Strains and maintenance

All C. elegans strains used are derivatives of N2 from the Caenorhabditis Genetics Center (CGC) and are listed in table S1. All worms were maintained at 15°C on nematode growth media (NGM) agar plates and fed the OP50 Escherichia coli B strain of bacteria. To avoid genetic drift, all worms were maintained for a maximum of 4 months (16 to 20 generations) before a fresh thaw was performed. For experimentation, worms were bleached to synchronize by degrading carcasses with bleach solution (1.8% sodium hypochlorite and 0.375 M KOH), then washing four times intact eggs with M9 solution (22 mM KH2PO4, 42.3 mM Na2HPO4, 85.6 mM NaCl, and 1 mM MgSO4), and plating eggs onto RNAi bacteria on NGM agar plates containing 1 μM isopropyl-β-D-thiogalactopyranoside, carbenicillin (100 μg/ml), and tetracycline (10 μg/ml) until the desired stages of adulthood. RNAi bacteria are derived from the HT115 E. coli K12 strain of bacteria containing pL4400 empty vector control or expressing double-stranded RNA containing the sequence of the target gene. RNAi strains were all isolated from Vidal or Ahringer libraries and sequence verified before use. All RNAi used in this study have been knockdown verified previously (7, 34–36).

rgf-1p::xbp-1s worms were synthesized by injecting N2 worms with pRTS (for rgf-1p::xbp-1s) at 25 ng/μl, pEK2 (myo-2p::tdtomato) at 2.5 ng/μl, and pD64 (100 ng/μl) vehicle as filler DNA. Worms positive for myo-2p::tdtomato were selected to identify for stable arrays. Integration was performed by gamma irradiation. Briefly, L4 worms were irradiated with 4000 to 4400 rem (roentgen-equivalent-man)
of radiation, and integrants were identified by selecting animals that maintained myo-2p::tdTomato at 100% frequency in the F3 generation. Two independent lines were isolated and backcrossed to N2 animals eight times to eliminate mutations, and animals with the most similar phenotypes to the array animals were used for experimentation. vha-6p::ehbp-1 worms were synthesized by injecting N2 worms with pSM1 (for vha-6p::xbp-1s) at 25 ng/μl, pEK1 (myo-2p::GFP) at 2.5 ng/μl, and pD64 vehicle (100 ng/μl) as filler DNA. Worms positive for myo-2p::GFP were selected to identify for stable arrays. All experiments were performed by positively selecting for array animals.

Staging of animals for all ER and lipid imaging and quantification was performed at late day 2/early day 3 of adulthood, as this is the earliest time point when changes to ER morphology were detected. This is the stage of adulthood where egg laying is at its maximum, approximately 90 to 100 hours after hatch at 20°C in standard growth conditions.

Sample preparation and microscopy

Worm fixing protocol and sample imaging for confocal microscopy

Staged worms were rinsed off plates with M9 and added to 15-ml conical tubes. Animals were washed three times with M9 to remove bacteria (do not exceed ~1000 g; 1000 relative centrifugal force (RCF) conical tubes. Animals were pelleted, excess M9 was aspirated to recommended for transferring of worms since they will adhere to plastic surfaces). Animals were pelleted, excess M9 was aspirated to 100 μl, and 100 μl of buffered paraformaldehyde (PFA) was added directly to worms [4% PFA, 340 μM CaCl2, 1.2% glucose, 40 mM NaH2PO4, and 60 mM Na2HPO4 (pH 7.4)]. Tubes were rocked at room temperature for 20 min, ~100 μl of liquid was removed, and 1.3 ml of M9 was added. Animals were then left at 4°C overnight. Animals were then spun down, solution was aspirated down to 100 μl, and 200 μl of 70% glycerol buffered in phosphate-buffered saline was mixed in. Animals were stored at −20°C until being imaged. For imaging, samples were mounted directly on glass slides, and a coverslip was placed directly on the specimen and sealed with nail polish. Mounted slides were kept at 4°C for at least 2 days before imaging to allow for settling of specimen and rehydration with glycerol. Fixed animals were viewed on a ZEISS LSM700 inverted confocal microscope, with constant acquisition settings when comparing specimens within a given experiment. Unless otherwise noted, images are maximum intensity projections (using the “Processing” tab in Zen software) of stacks spanning the width of various tissues within a single transgenic animal. Neutral lipid (BODIPY 409/503) staining was performed as previously described in Klapper et al. (37).

Live-cell imaging using wide-field/compound microscopy

When preparing samples for live-cell microscopy, individual worms were picked off plates and resuspended in M9 containing 100 nM sodium azide. A coverslip was placed directly on the sample, sealed with nail polish, and imaged immediately. Specimens were viewed on a ZEISS AxioObserver.Z1 microscope equipped with a lumencor SOLA light engine and a ZEISS Axiocam 506 camera, driven by ZEISS ZEN Blue software using a 63×/1.4 Plan Apochromat objective and standard GFP filter (ZEISS filter set 46 HE), and a standard dsRed filter was used (ZEISS filter set 43).

BioSorter analysis

Staged worms were washed off plates and resuspended in M9 into 15-ml conical tubes. We used a Union Biometrica COPAS (complex object parameter analysis sorter) BioSorter (product no. 350-5000-000) using both 561- and 488-nm light sources. BioSorter calibration, cleaning, and sample running were performed as described in (16). For vha-6p::ehbp-1 worms, myo-2p::GFP was used as a coinjection marker. Therefore, all BioSorter analysis was performed in the lower 70% of the animal, cutting off signal from the top 30% (i.e., the contribution from the pharynx GFP).

Software specifications and data analysis

Worm profile data were collected using the Biosort 5401.1 software provided for use with the BioSorter machine. Before running an orientation algorithm ["LAMPro" (16)], all profiles were converted from one of multiple .dat file(s) into a single .txt file using the “Export as Text” function in the Profile Reader 16.1 software (also sold by Union Biometrica) and then saved in a folder alongside the .lmd and .dat files that were saved by the BioSorter at the time of data acquisition. All orientation algorithms were written in Perl5 and can be interpreted in Cygwin (a Linux application programming interface that can run on Windows) through additional Perl5 packages. Cygwin can be downloaded at the following address: Cygwin.com/install.html. During Cygwin installation, users will be given the option to install the additional Perl5 packages. Data analysis, significance testing, and data plotting were run using MATLAB scripts (version R2015a), which have been integrated into a single graphical user interface (“LAMPro Suite 2.0”). To quantify the number and/or size of fluorescently labeled cell markers (e.g., MLS::GFP, LMP-1::GFP, and GFP::RAB-10 punctae), stacks were normalized to the same threshold on a dark background using ImageJ. A smooth function eliminated scattered pixels. The area containing the intestine was highlighted, and the “Analyze Particles” function was used to give a count. These counts were normalized to “control” images (e.g., rab-3p::xbp-1s signal divided by the count from animals without rab-3p::xbp-1s). Enrichment relative to controls was then calculated for at least three biological replicates, and a t test was calculated to measure if this was above or below 1. Colocalization was calculated using the “JACoP” (Just another Colocalization Plugin) for experiments that contained at least three biological replicates, and a t test was performed to see if the variance of the set was above or below matched controls. All violin plots were made in R.

Life-span analysis

Life-span measurements were performed on solid NGM agar plates spotted with RNAi bacteria (HT115 E. coli strain K12). Worms were synchronized by bleaching, plated onto RNAi from hatch, and grown to adulthood at 20°C. Adult worms were moved away from progeny by moving worms onto fresh RNAi plates every day until days 7 to 10 when progeny were no longer visible. Animals were then scored every 1 to 2 days for death until all animals were scored. Animals with bagging, vulval explosion, or other age-unrelated deaths were censored and removed from quantification. Prism 5 software was used for statistical analysis. Log-rank (Mantel-Cox) method was used to determine significance.

Epifluorescence microscopy

Staged worms were picked at random (under white light) from a population and immobilized in 100 nM sodium azide. Immobilized worms were aligned on a solid NGM agar plate, and images were captures using a Leica M250FA automated fluorescent stereo microscope equipped with a Hamamatsu ORCA-ER camera.
Induction of ER stress
Protocol was adapted from Tian et al. (36). Briefly, synchronized L4 stage animals were treated with solution (25 ng/µl) of tunicamycin in M9 buffer for 4 hours. Control animals were treated in an equivalent solution of dimethyl sulfoxide (DMSO) (1%) in M9 buffer. Animals were rotated in DMSO or tunicamycin in 15-ml conical tubes. Drugs were washed three times with M9, and animals were plated on OP50 to recover overnight at 20°C and then imaged at day 1.

Electron microscopy
Whole-worm samples were processed for electron microscopy as previously reported (38). Briefly, staged worms were subjected to high-pressure freezing (BAL-TEC HPM 010) and then freeze substituted with an acetone/resin series (25% resin, then 50, 75, and 100%). Worms were cured in pure resin, then sectioned (70-nm sections), and then imaged (FEI Tecnai 12 Transmission Electron Microscope) on formvar-coated mesh grids.

SUPPLEMENTARY MATERIALS
Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/6/1/eaaz1441/DC1

REFERENCES AND NOTES
Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/6/1/eaaz1441/DC1
R.H.-S., the National Science Foundation Graduate Research Fellowship to J.W.P., 1F31AG060660-01 through NIA to M.G.M., and 4R01AG042679-04 through the NIA and the Howard Hughes Medical Institute to A.D. We thank the CGC for availability of strains. **Author contributions:** J.R.D. developed the story, performed life spans, performed confocal and BioSorter imaging and data analysis, and helped in preparing figures and writing the manuscript. R.H.-S. developed the story, performed life spans, completed wide-field, confocal, and BioSorter imaging and data analysis, prepared the figures, and wrote the manuscript. J.D. and S.M. helped in developing the story and performed life spans, confocal imaging, BioSorter imaging, and strain construction. V.R. helped in the UPRER activation experiments and strain construction. M.S. performed all sample preparation and imaging for electron microscopy. S.U.T., N.K., M.G.M., and P.A.F. helped in life spans and imaging. G.G. performed strain construction for ER imaging. C.B. and M.Z. helped in the data analysis. D.J.E. made essential additions to the LAMPro software, which enabled data analysis in this paper to be performed. L.J. and A.D. helped in developing the story and assisted in manuscript preparation. **Competing interests:** The authors declare that they have no competing interests. **Data and materials availability:** All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from authors. All strains synthesized in this manuscript are derivatives of N2 or other strains from CGC and are either made available on CGC or available upon request.

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UPRER promotes lipophagy independent of chaperones to extend life span

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