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Narayan, Vikram; Ly, Tony; Pourkarimi, Ehsan; Brenes Murillo, Alejandro; Gartner, Anton; Lamond, Angus; Kenyon, Cynthia

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Supplemental Information

Deep Proteome Analysis Identifies Age-Related Processes in *C. elegans*

Vikram Narayan, Tony Ly, Ehsan Pourkarimi, Alejandro Brenes Murillo, Anton Gartner, Angus I. Lamond, and Cynthia Kenyon
Figure S1

Panel A: Graph showing the percentage of worms alive over time for different RNAi treatments.

Panel B: Graph showing the number of eggs produced over time for different RNAi treatments.

Panel C: Graph showing the percentage of worms in different classes over time.

Panel D: Graph showing the pumping rate of animals over time, with a 50% drop indicated.

Panel E: Images showing the effect of FUdR concentration on egg production.

Panel F: Graph showing the percentage of worms alive over time for different FUdR concentrations.

Table:

| Background | Experiment | Treatment  | #Total worms | #Censored | #Scored | Mean lifespan (days of adulthood) |
|------------|------------|------------|--------------|-----------|---------|----------------------------------|
| N2         | 1          | 0 uM FUdR  | 100          | 21        | 79      | 16.4                             |
| N2         | 1          | 50 uM FUdR | 100          | 6         | 94      | 16.2                             |
| N2         | 2          | 0 uM FUdR  | 100          | 25        | 75      | 17.2                             |
| N2         | 2          | 50 uM FUdR | 100          | 24        | 76      | 16      |
Figure S2

A

B

C

D

E

Figure S2

A

B

C

D

E
Protein-coding genes with at least 1 GO Cellular Component annotation

Number of GO Cellular Component annotations per gene

![Graph F](image)

Protein-coding genes
- 20,359
- 7,848
- 4,954

Protein-coding genes with at least 1 manually curated GO Cellular Component annotation

Method of curation
- Automated
- Inferred
- Experimental
- Computational
- Manual
- Curator Statements
- Author Statements

Number of GO Cellular Component annotations

![Graph G](image)

![Graph H](image)
MaxQuant Evidence File

Data pooled from 3 biological replicates

1,706,820 entries

1,487,547 entries

Quality Control: Step I

Remove:
- Contaminants
- Reverse hits
- E. coli proteins

30,779 protein groups

Assemble Protein Groups

Aggregate SILAC ratios (median), intensity (sum) & peptide count (sum) by 'Proteins', 'Gene.Names' & 'Experiment'

Identification

Quality Control: Step II

Select:
Protein groups identified in 3 out of 3 biological replicates with 1 or more Razor+Unique peptides

9,398 protein groups identified

30,779 protein groups

Data pooled from 3 biological replicates

Identification

Quantification

Quality Control: Step III

Select:
Protein groups identified in 3 out of 3 biological replicates with 2 or more Razor+Unique peptides

7,380 protein groups quantified

9,398 protein groups identified

Select:
Protein groups identified in 3 out of 3 biological replicates with 1 or more Razor+Unique peptides

637 Low

5,882 Medium

861 High

Sort Data

Distribute:
Protein groups into 3 abundance bins:
- log10(Intensity) ≤ 8.5
- >8.5 & <10.5
- ≥10.5

Significant age-dependent changes in protein abundance

627 protein groups

2σ Significant Changes

Apply:
95% Confidence Intervals
Remove:
Age-invariant proteins with ratios within mean ± 2 standard deviations for each SILAC pair

5,665 protein groups

p < 0.05 Significant Changes

Apply:
ANOVA
- Randomly select 3 values from a set of 25 unchanging controls (values closest to 0) for each ratio in each abundance bin and calculate ANOVA for each protein group in that bin relative to these
- Loop this randomization & ANOVA estimation 100 times (FDR)
- Calculate mean p-value for each protein group
Remove:
Protein groups with FDR-corrected p-value ≥ 0.05 i.e. less than 95% significance

5,665 protein groups

95% Confidence Intervals
Remove:
Age-invariant proteins with ratios within mean ± 2 standard deviations for each SILAC pair

627 protein groups

Downstream Analysis

GO-term enrichment

Clustering

Network Analysis

C. elegans proteins

Human homologs

Downstream Analysis

GO-term enrichment

Clustering

Network Analysis

C. elegans proteins

Human homologs
Figure S4

A

Day 10 versus Day 1

Day 10 versus Day 5

Day 5 versus Day 1

B

Protein groups

Proteins (with unique peptides)

Peptides

53.8% overlap

31.6% overlap

21.6% overlap

C

Relative protein abundance

2 cluster model

Days of adulthood

D

Decreased abundance

Increased abundance

Cluster 1

Cluster 2

- Nucleosome assembly
- Response to heat
- Negative regulation of mitotic cell cycle
- Mitotic cell cycle process
- Nuclear division
- Determination of adult lifespan
- Multi-organism process
- Single organism reproductive process
- Developmental process involved in reproduction
- Embryo development ending in birth or egg hatching

- Threonine catabolic process
- Purine nucleobase biosynthetic process
- de novo IMP biosynthetic process
- Unsaturated fatty acid biosynthetic process
- Flavanoid biosynthetic process
- Alpha-amino acid biosynthetic process
- Sulfur compound metabolic process
- Oxidation-reduction process

Cluster 1 (334)

Cluster 2 (293)
Figure S6

A

Distal end of Gut

Gut cells

Day 1 Day 5 Day 10

GFP

B

Control: No Heat Shock

Day 1 Day 5 Day 10

Bright Field

GFP

Day 1 Day 5 Day of adulthood

Heat Shock

Day 10

Day 1 Day 5

Day of adulthood

C

Whole worm GFP intensity

Mean Intensity

( Arbitrary Units)

Day of adulthood

D

Number of peroxisomes

(sum across 81 z-stacks per worm)

GFP-SKL Foci

(indicative of peroxisomes)

Phsp-16.2::gfp-SKL worms

Pges-1::gfp-SKL worms

E

Number of Eggs

( sum across 81 z-stacks per worm)

Number of Eggs

Control RNAi prx-5 RNAi

p = 0.00019
Microscopy data (n = 20 individual worms except for SQST-1 Day 10 males, where n = 5 individual worms)
Mass spectrometry data (n = 3 biological replicates)
Graphs show sample mean ± standard deviation
SUPPLEMENTAL FIGURE LEGENDS

Figure S1. FUdR treatment does not affect adult C. elegans lifespan [Related to Figure 1]

(A) Wild-type N2 (Bristol) animals were grown on HT115 E. coli at 20°C from hatching and transferred to the indicated RNAi bacteria at mid-late L4. Animals were scored as alive or dead on each day of adulthood until the entire population was dead; missing, bagged or exploded animals were censored. n = 89 (L4440 empty vector RNAi control), n = 94 (orn-1 RNAi) and n = 78 (daf-16 RNAi); n does not include censored animals; n = 100 per condition prior to censoring. The graph on the right was plotted without depicting any data points, for clarity, and shows the mean lifespan for wild-type and orn-1(RNAi) animals. (B) Graph of number of eggs laid in wild-type N2 (Bristol) worms at 20°C between days 1 to 10 of adulthood. Day 1 was considered to be 3 days after hatching (24 h after L4). 20 adult animals were scored; mean eggs laid ± standard error of mean is indicated on the graph. The time points used in this study (days 1, 5 and 10 of adulthood) are highlighted. (C) The lifespan of wild-type N2 (Bristol) worms (n = 100 per experiment) at 20°C was measured; missing, bagged or exploded animals were censored. On each day of adulthood, the number of Class A, B and C animals, designated based on motility as described in Herndon et al, 2002, was scored. Shown is the mean ± standard deviation from two independent experiments. (D) Graph of the rate of pharyngeal pumping per minute plotted against the age of the animals. Pharyngeal pumping decreases markedly early in adulthood and is virtually undetectable by day 13. Animals (n = 20; see figure for details) were scored for 3 x 30 s intervals every 24 h and the mean pumping rate per minute for each animal calculated. The graph shows the average of the mean pumping rate per minute per worm ± standard error of mean. Pumping could not be detected in some animals after day 7. (E) N2 (Bristol) animals were treated with the indicated concentrations of FUdR beginning from the mid-late L4 larval stage. Shown are images taken on day 2 of adulthood (left) and total number of eggs laid (right; n = 20; graph shows mean ± standard deviation) up to day 10 of adulthood. All FUdR doses tested completely prevented eggs from hatching, as is evident from the images. The total number of eggs laid was significantly lower at doses higher than 50 µM (* = p-value from t-test compared to 0 µM control < 0.05; ** = p-value < 0.01). (F) The lifespan of N2 (Bristol) animals on 0 or 50 µM FUdR (treatment was initiated at mid-late L4) at 20°C was scored. Missing, bagged or exploded animals were censored. Shown is the mean ± standard error of mean from 2 independent experiments.

Figure S2. Differential buffer extraction reveals putative sub-cellular localization profiles for ~6,000 C. elegans proteins [Related to Figure 1 and Table S1]

(A) Venn diagram (not area-proportionate; generated using Interactivenn) showing proteins identified in all 3 biological replicates using the S-buffer differential extraction method employed in this study that were unique to each buffer, as well as the overlap between buffers. Buffer 1 is a 4% SDS-based buffer (see Supplemental Experimental Procedures) and buffers 2-5 are from the Qiagen QProteome Cell Compartment Kit (CE1, CE2, CE3 and CE4 respectively). CE1 = cytoplasmic fraction, CE2 = membrane fraction, CE3 = nuclear fraction, CE4 = cytoskeletal fraction and insoluble materials. (B) Bar plot showing the number of proteins identified using each of the 5 buffers in (A) with 1 or more, 2 or more and 4 or more peptides. Also included is the aggregate data wherein peptides identified using each buffer were pooled and compiled into a single dataset (“Combined”). (C) Heat map showing the distribution of 6,283 proteins across the indicated sub-cellular compartments, isolated using the QProteome Cell Compartment Kit (Qiagen), in day 1 adult C. elegans (also see Table S1, sheet 2). (D) 2.5 µg of each sub-cellular fraction or 4% SDS-extracted whole-cell lysate were run out on a 4-12% NuPAGE gel (Thermo Fisher) and analyzed by immunoblotting with the following antibodies: anti-GAPDH (Sigma; 1:5000), anti-Histone H3 (Abcam; 1:4000) and anti-ATP synthase subunit α (Mitosciences; 1:2000). (E) GO-term enrichment (Cellular Component) of the proteins present in each sub-cellular compartment was performed using PANTHER and plotted using REVIGO, in order to validate the efficacy of the extraction methodology in nematodes. The size of the bubbles is indicative of the number of proteins annotated with that GO-term (range: 9-309 for cytoplasm; 7-168 for membrane; 5-14 for nucleus; 4-60 for cytoskeleton); bubbles are color-coded according to significance (Bonferroni-corrected p-value). To reduce GO-term redundancy, the ‘PANTHER GO-Slim Cellular Component’ annotation data set was used unless the output was predominantly ‘unclassified’, in which case ‘GO Cellular Component complete’ was used. Only those proteins whose levels in a single fraction were found to be 75% or more of their total abundance across all fractions were included in the analysis [i.e. (intensity in a single fraction / sum of intensities across all 4 fractions) x 100 >= 75]. This subset included 2,297 cytoplasmic proteins, 354 membrane proteins, 78 nuclear proteins and 321 cytoskeletal proteins (see Table S1, sheets 3-6). (F) Area-proportionate Venn diagram (generated using BioVenn) showing that only ~39% of the theoretical C. elegans protein-coding genes (20,359 according to
WormBase Release WS250, 31 July 2015) have at least one Cellular Component GO-term annotation (7,848; GO_CC), and of these, only 4,954 have a manually curated GO_CC annotation. (G) Histogram showing the number of GO_CC annotations assigned to each of the 7,848 C. elegans protein-coding genes with a GO_CC annotation. (H) Graph showing the breakdown of GO_CC assignments to the 7,848 C. elegans protein-coding genes with at least one GO_CC annotation. As stated in (F), 4,954 of these genes (63%) have a manually curated GO_CC annotation. However, only a subset of the manually curated GO_CC annotations (~10,000 annotations, corresponding to 2,121 protein-coding genes) is based on experimental evidence. Thus, only about 10% (2,121 out of 20,359) of all C. elegans protein-coding genes have a GO_CC assignment based on experimental data. (F-H) Raw data was obtained from WormBase, and mined/queried using WormMine.

Figure S3. Data analysis workflow [Related to Figures 1 and 2, and Tables S2-S4] Schematic representation of the data analysis pipeline used in this study.

Figure S4. Assessment of proteomics data quality [Related to Figures 1 and 2] (A) Scatter plots depicting log2-transformed SILAC ratios (for each pair-wise comparison of the 3 time-points used in this study) between the 3 biological replicates. Reproducibility of protein quantification across the replicates was high, as indicated by the Pearson’s r values shown on the upper left corner of each scatter plot. (B) Left: Area-proportionate Venn diagram (generated using BioVenn) showing the overlap between proteins identified in this study (adults only; “Experimental”) versus the theoretical C. elegans proteome comprising all developmental stages, as defined by WormBase (Release WS243, 28 March 2014 containing 20,480 protein-coding genes; “Theoretical”). Center: Area-proportionate Venn diagram of the theoretical C. elegans proteome (as above) and the number of proteins reproducibly identified with sequence-unique peptides in this study (“Experimental”). The “Experimental” dataset here does not, therefore, include peptides that were assigned to protein groups rather than individual proteins. Right: A theoretical tryptic digest of the C. elegans proteome was generated and a list of putative MS-flyable peptides compiled (conditions imposed: peptides must have at least one Lys or Arg, and must be 6-35 amino acids in length; “Theoretical”). This list was compared to the sequence-unique peptides identified in this study (“Experimental”) and is represented as an area-proportionate Venn diagram. (C) Cartoons showing representative profiles of the 627 age-variant proteins (see main Figure 2) grouped by hierarchical clustering into either 2 clusters (top) or 6 clusters (bottom). (D) Hierarchical clustering of the 627 proteins using Ward’s minimum variance method into 2 groups based on trend profiles. To minimize redundancy in clustering due to large differences in the magnitude of the abundance changes, the SILAC ratios were scaled to range between -1 (maximum decrease) and +1 (maximum increase) with ratios near 0 indicating no change in abundance during the 3 time points considered in this study. Results from GO-term enrichment analysis (multiple hypothesis [Bonferroni corrected p < 0.05] of each cluster using PANTHER are also depicted.

Figure S5. Gene Set Enrichment Analysis of proteomics data reveals pathways altered during nematode aging [Related to Figures 3 and 4] (A) Protein quantification data for day-10 versus day-5 of adulthood, as measured in this study, were analyzed using gene set enrichment analysis to identify KEGG pathways that change during aging. Shown is the GSEA output using the algorithm from the Broad Institute. (B) Density plots depicting the abundance of a gene set comprising KEGG ‘peroxisome’ genes relative to the entire dataset. The curve for peroxisome genes includes those only those genes that were detected both in a study by Youngman et al. 2011 measuring mRNA levels and in this study, measuring protein levels. Top: density plot for peroxisome mRNA transcripts relative to the entire Youngman et al. 2011 mRNA dataset; Bottom: density plot for peroxisome protein levels relative to the entire protein dataset measured in this study. (C) Scatter plot showing mRNA levels measured in a study by Youngman et al. 2011 plotted against the levels of their corresponding protein products as measured in this study. The mRNA and protein data show poor correlation (r = 0.25 [Pearson]). A linear model, calculated using the function lm in R, is depicted by the red line. 95% confidence intervals for the linear model are shaded light red.

Figure S6. Control experiments for GFP-SKL C. elegans strains [Related to Figure 5] (A) In order to ensure that the GFP-SKL foci observed in young Phsp-16.2::gfp-SKL worms after induction of GFP-SKL expression by heat shock were not merely artifacts or stress granules, Phsp-16.2::gfp animals, lacking the peroxisome-targeting -SKL signal were used (strain CL2070). No foci were detected in these animals after heat shock. Similar to data from GFP-SKL animals (see below), hsp-16.2 driven GFP expression was markedly decreased by day 10 of adulthood. (B & C) Whole-body hsp-16.2-driven GFP-SKL intensity was measured and found to increase significantly between days 1
and 5 of adulthood, and then drop markedly by day 10 (p-value from unpaired t-test = 1.02 x 10^{-4} for day 1 versus day 5; 1.70 x 10^{-5} for day 5 versus 10; and 0.067 for day 1 versus day 10, n = 10 in all cases). Shown are bright field and GFP images taken with (right) and without (left) heat shock (B) as well as a graph showing quantification of GFP intensity from the images (C). (D) The number of foci resolved ("peroxisomes") at day 1 and day 5 (based on data presented in main Figures 5C and 5D) was summed across the entire z-stack (81 sections per worm) for GFP-SKL driven by the hsp-16.2 promoter (left) and the ges-1 promoter (right). The number of dots resolved was found to increase significantly by day 5 (p-value from unpaired t-test = 0.0048 (left) and 3.07 x 10^{-9} (right)). (E) N2 (Bristol) worms were treated with control or prx-5 RNAi from late L4/young adult stage, and the number of eggs laid on each day of adulthood measured (left). Also shown is the total number of eggs laid during days 1 to 10 of adulthood (right). Graphs show mean ± standard deviation (n = 20 for control RNAi animals and n = 26 for prx-5(RNAi) animals). The total number of eggs laid in prx-5 RNAi animals was found to be significantly less than control animals using t-test (p-value = 0.00019).

**Figure S7. Comparison of mass spectrometric predictions with in vivo measurements of age-dependent protein changes by microscopy [Related to Figure 6]**

(A-F) Graphs showing mass spectrometry-determined changes in protein abundance during aging (red) compared to in vivo fold-change measurements calculated by imaging GFP::protein fusion worm strains (black). For microscopy, 20 individual worms were analyzed (see (G) and main Figure 6A for snapshots); quantification was performed using Cell Profiler (Broad Institute). Shown is the mean ± standard deviation. Note that the individual variation in protein expression for all GFP::protein fusion strains increased markedly by day 10. SQST-1::GFP worms (strain HZ859) were in a him-5(e1490) background; thus, both hermaphrodites (E and main Figure 6A) and males (F, G) were examined. (H) HIS-24::GFP (left), HIS-71::GFP (center) and F55B11.4::GFP-expressing (right) worms were grown on the indicated RNAi (initiated at L1 stage) and GFP-fluorescence was quantified on days 1, 5 and 10 of adulthood. Graphs show mean ± standard deviation (n = 20 worms). Significance was calculated using an unpaired t-test (p-value < 0.05 = *, < 0.01 = **, < 0.001 = ***). Thedaf-2 and hsf-1 RNAi are in different vector backbones (see Supplemental Experimental Procedures); thus, the corresponding empty vector control for each was used.

**Figure S8. Comparison between this study and recent proteomics studies (Liang et al. 2014; Walther et al. 2015; Zimmerman et al. 2015) [Related to Figure 1 and Discussion]**

(A) Area-proportionate Venn diagram showing that 97.4% of the proteins for which quantification data from the Liang et al. 2014 study was publicly available, were also detected in our study. (B) Significant fold-changes as determined in the Liang et al. 2014 study based on undefined thresholds, were projected onto our thresholds for significance (95% confidence interval calculated from the mean ± 1.96 standard deviations based on quantification of 7,380 proteins from 3 biological replicates). The majority of the changes deemed significant in the Liang et al. 2014 study were not scored as significant in our study based on our stricter thresholds. The Liang et al. 2014 data (upper panel) is compared to our quantification data (lower panel) for the same proteins. Also indicated are the mismatches (6 proteins) between the two datasets. (C) Area-proportionate Venn diagram showing that 92% of the proteins detected in the Walther et al. 2015 study were also detected in our study. (D) Scatter plot for proteins detected in both the Walther et al. 2015 study and this study, showing reproducibility of protein quantification between the studies. A modest correlation was observed (r = 0.54 [Pearson]); this is likely adversely affected by the fact that different quantification strategies were used in the two studies (spike-in SILAC versus triple-SILAC). Additionally, the measured time-points were not identical (day 21 / day 1 data from the Walther et al. 2015 study were compared to day 10 / day 1 data from our study). A linear model, calculated using the function lm in R, is depicted by the red line. 95% confidence intervals for the linear model are shaded light red. Although the magnitude of the fold-change for each protein in the datasets was only moderately similar (reflected in the modest r = 0.54), 74% of the proteins showed similar trends (i.e. increase or decrease) in both datasets. (E) Area-proportionate Venn diagram showing that 82% of the proteins detected in the Zimmerman et al. 2015 study were also detected in our study. (F) Scatter plot for proteins detected in both the Zimmerman et al. 2015 study and this study, showing reproducibility of protein quantification between the studies. A modest correlation was observed (r = 0.47 [Pearson]); this is likely adversely affected by the fact that different quantification strategies were used in the two studies (dimethyl labeling versus SILAC). Additionally, the measured time-points were not identical. A linear model, calculated using the function lm in R, is depicted by the red line. 95% confidence intervals for the linear model are shaded light red.
SUPPLEMENTAL EXPERIMENTAL PROCEDURES

Reagents
EZQ protein assay, Triscarboxyethylphosphine (TCEP), CBQCA assay, Pepmap C18 (2 cm x 75 μm) trap columns and EasySpray C18 columns (2 μm particles, 50 cm x 75 μm) were from Thermo Fisher Scientific (Waltham, MA). MS Grade Trypsin was from Promega. Sep Pak C18 μ-elution 96-well plates were purchased from Waters. Purified arginine and lysine for SILAC experiments were from Cambridge Isotopes Laboratories (Tewksbury, MA) via CK Gases and were as follows: Light (Arg0, ULM-8347; Lys0, ULM-8766); Medium (Arg6, CLM-2265; Lys4, DLM-2640); and Heavy (Arg10, CNLM-539; Lys8, CNLM-291). All other materials were obtained from Sigma, unless otherwise indicated.

RNAi experiments
RNA interference experiments were performed using the RNAi via feeding strategy [Ahringer library; (Kamath et al. 2001)]. The exception was the daf-2 RNAi construct used (pAD48-daf-2-RNAi; Addgene plasmid #34834 and corresponding empty vector control pAD12; Addgene plasmid #34832), which was made in-house, and has been characterized previously (Dillin et al. 2002). RNAi of orn-1 for SILAC experiments was performed as described previously (Larance et al. 2015; Larance et al. 2011). prx-5 RNAi was initiated during ‘day 0 of adulthood’, just prior to the appearance of eggs in the hermaphrodite animal, as described in Zhou et al. 2012. All other RNAi experiments were initiated at the L1 stage and the resulting adults were imaged as described in the main Experimental Procedures.

Lifespan assays
Lifespan assays were performed at 20°C as described previously, except that days of adulthood (day 1 = 24 h post L4) were reported in our study (Hsin & Kenyon 1999). Additionally, for orn-1 and daf-16 RNAi experiments, RNAi treatment was initiated at L4 and FUdR was not used; the animals were instead transferred to fresh RNAi plates every other day. For FUdR lifespan assays (dose response), synchronized wild-type L4 animals were transferred onto to NGM/OP50 plates containing the required concentration of FUdR and assessed for survival on each day of adulthood. Animals that exploded, bagged or were missing were censored in all cases.

Egg-laying assays
Egg-laying assays were performed at 20°C. In brief, N2 animals were synchronized as L1 larvae by filtering [11 μm filter; (Larance et al. 2011)] and at L4, 30 animals were transferred onto fresh 6 cm NGM/OP50 plates at 3 animals per plate. 24 h post L4 (day 1), the adults were transferred onto fresh NGM/OP50 plates and this was repeated every 24 h until no more eggs were laid. On each day, the plates containing laid eggs and/or L1 larvae (from which the adults were transferred onto new plates) were incubated at 25°C overnight, followed by a 30 min incubation at 4°C to temporarily immobilize the animals. Dead eggs and larvae (viable eggs) were scored. For FUdR egg-laying assays, FUdR treatment was initiated at L4 and adults were transferred onto fresh NGM/OP50 plates containing FUdR on each day of adulthood. prx-5 RNAi treatment was initiated at day 0 of adulthood, as described previously (Zhou et al. 2012) and adults were transferred onto fresh RNAi plates on each day of adulthood.

Determination of class A, B, C animals
Class A, B, C animals were identified as described previously (Herndon et al. 2002). Measurements were made at 20°C using N2 (Bristol) animals. Experiments were performed in a similar manner to lifespan assays (see above) although no FUdR was used, and animals were scored as being class A, B or C on every other day of adulthood. Exploded, bagged or missing animals were censored. In brief, animals that moved well were scored as Class A, animals showing uncoordinated movements were scored as Class B, and animals that showed little or no movement were scored as Class C.

Pumping assays
Synchronized single N2 (Bristol) animals maintained at 20°C were observed. On each day of adulthood, the number of contractions in the terminal bulb of the pharynx was counted for 3 x 30 s intervals (n = 20 or as indicated in the figure legends).

Microscopy and imaging
Induction of GFP-SKL expression from the hsp-16.2 promoter was carried out as follows: day-1 adults (or as indicated in the figure legends) grown at 20°C were shifted for 2 h to 34°C, followed by 30 min at 20°C, then 1 h at 34°C, after which the worms were returned to 20°C for 12 h. Animals were
anesthetized on 2% agarose pads in 10 mM levamisole for no more than 10 min immediately prior to viewing. Imaging of GFP-SKL strains (intestinal cells) was performed using an UltraVIEW Vox CSU-X1 confocal imaging system (Perkin Elmer), incorporating an Olympus IX70 microscope, a 488 nm solid-state laser, and an Olympus 60X/1.3 NA oil-immersion objective. Images were processed using Velocity 15 software (Perkin Elmer) and analyzed using Cell Profiler (Lamprecht et al. 2007) as follows: z-sections were taken over 40 µm at 0.5 µm intervals. The 81 resulting images per worm were quantified individually for peroxisomes (using a modification of the speckle counting tool in Cell Profiler) and cytosol (total GFP intensity minus peroxisomes). Mean intensities from each section were then summed to get total intensity, peroxisomal intensity and cytosolic intensity per worm. These were then graphed using R. Additionally, the number of peroxisomes was counted and graphed from the sum of the 81 z-stacks as well as from the single z-section that had the maximum number of peroxisomes. All other microscopy was performed using a Leica MZ16FA microscope (Leica; Bannockburn, IL) and images were analyzed using Cell Profiler. Here, fluorescence intensity in the whole animal was measured and subtracted from background fluorescence (i.e. mean fluorescence) outside the worm and plotted using R.

### SILAC labeling, sample preparation, fractionation and clean-up

For isotopic labeling, N2 (Bristol) worms were grown on NGM-N plates at 20°C as described previously (Larance et al. 2011) with the following adaptations: 25 mid-late L4 larvae were picked on to NMG-N plates supplemented with 1 mM IPTG and 1 µg/ml carbenicillin, and seeded with *E. coli* SLE1 grown in light, medium or heavy label as described previously (Larance et al. 2011). Labeling was for 2 generations at 20°C. L1 animals from the F2 generation were collected by filtering through an 11 µm filter (Millipore) and 40,000 L1 larvae per condition (light, medium or heavy) were transferred to 9 cm NGM-N plates seeded with labeled SLE1 at 4,000 worms per plate. Approximately 2 days later, when the worms were mid-late L4 animals, they were washed off the plates with M9 buffer and transferred to fresh NGM-N plates (seeded with the required labeled bacteria) containing 50 µM FUdR. Care was taken to ensure the animals did not starve during the experiment – if required, additional concentrated, labeled SLE1 was added to the plates. The animals were then collected as required (Heavy – Day 1 adults, Medium – Day 5 and Light – Day 10) by washing off the plate with M9 buffer, followed by a further 3-5 washes in M9 to remove bacteria and eggs. Worms were collected by light centrifugation or by gravity at room temperature (5-10 min in a falcon tube on the bench). Worm pellets were snap-frozen in liquid N2 and upon thawing, were mixed at a 1:1:1 ratio by volume of heavy:medium:light. 5% of the mixture was directly in SDS buffer (4% SDS, 150 mM NaCl, 10 mM phosphate buffer pH 7, 50 mM N-ethylmaleimide, 25 mM TCEP) as follows: 2x volume of buffer and 1x volume of 0.7 mM Zirconia beads (BioSpec) were added and the sample was lysed by bead beating (Mini Beadbeater 8, BioSpec) at max speed for 3 x 20 s with 20 s intervals. The sample was then heated to 65°C for 10 min, centrifuged at 16,100 g for 15 min at room temperature and the supernatant collected. The remainder of the worm mixture (95%) was fractionated using the Q-Proteome Cell Compartment Kit (Qiagen) as described previously (Pourkarimi et al. 2012), but with the following modifications: (1) worm pellets were aliquotted into 100 µl aliquots and each aliquot was processed separately; (2) 100 µl 0.7 mM Zirconia beads were used per aliquot, together with 250 µl buffer CE1; subsequent buffers were scaled accordingly, maintaining the ratios recommended by the manufacturer; (3) homogenizing using a Qiashredder (Qiagen) was adjusted as follows: the sample was added to a shredder and centrifuged for 10 s at 100 g, after which the eluate was transferred to a fresh shredder and the process repeated. SDS, TCEP and NEM were added to each fraction, to final concentrations of 2%, 25 mM and 50 mM respectively, following which the samples were heated at 65°C for 10 min. Equal amounts of protein (500 µg, measured by EZQ Assay) from the SDS-lysate and sub-cellular fractions were concentrated by chloroform-methanol precipitation and resuspended in 8 M urea in 100 mM triethylammonium bicarbonate (TEAB). Samples were diluted 8-fold using 100 mM TEAB and supplemented with CaCl2 to a final concentration of 1 mM. Trypsin was added at a 1:50 ratio of enzyme to protein and incubated for 16 h at 37°C. Sample desalting and offline SAX chromatography was performed as described previously (Larance et al. 2015; Ly et al. 2014).

### LC-MS/MS and analysis of spectra

~1 µg of each desalted SAX fraction containing peptides in 5% (vol/vol) formic acid (~10 µl) was injected onto a 0.3 mm id × 5 mm PepMap-C18 pre-column and chromatographed on a 75 µm × 15 cm PepMap-C18 column. Using the following mobile phases: 2% ACN incorporating 0.1% FA (Solvent A) and 80% ACN incorporating 0.1% FA (Solvent B), peptides were resolved using a linear gradient from 5% B to 35% B over 4 hours with a constant flow of 200 nL min⁻¹ as described previously (Ly et al. 2014). Peptides were ionized by electrospray ionization at +2.0 kV. Tandem mass spectrometry analysis was carried out on a Q Exactive mass spectrometer (Thermo Fisher Scientific) using HCD.
fragmentation. The data-dependent acquisition method used acquired MS/MS spectra on the top 10 most abundant ions at any one point during the gradient. The raw data produced by the mass spectrometer were analysed using the quantitative proteomics software MaxQuant including the Andromeda search engine (http://www.maxquant.org, version 1.4.1.2) (Cox & Mann 2008). Peptide and protein level identification were both set to a false discovery rate of 1% using a target-decoy based strategy. The databases supplied to the search engine for peptide identifications were both the C. elegans (December 2013) and E. coli Uniprot databases (July 2012) containing 22,998 and 4,431 entries respectively. The mass tolerance was set to 7ppm for precursor ions and MS/MS mass tolerance was set at 20 ppm. Enzyme was set to trypsin/p with up to 2 missed cleavages. Deamidation of Asn and Gln, oxidation of Met, pyro-Glu (with N-term Gln), and acetylation of the protein N-terminus were set as variable modifications. N-ethylmaleimide on Cys was searched as a fixed modification. The output from MaxQuant provided peptide level data as well as protein level data, grouped by protein isoforms. Protein groups were defined as a set of unique peptides that are either shared between multiple protein isoforms, or belong to a single protein isoform (Figure S3). While this may lead to higher redundancy it enables us to quantify changes in individual protein isoforms. All raw proteomics data from this study have been deposited to the ProteomeXchange Consortium (http://proteomecentral.proteomexchange.org) via the MassIVE partner repository with the dataset identifiers PXD002777 (ProteomeXchange) and MSV000079263 (MassIVE). The processed data are additionally provided in a searchable, online database (https://www.peptracker.com/epd/).

**Data analysis, statistics, homology mapping and graphing**

Most of the subsequent data analysis was done using R version 3.1.3 (R Core Team n.d.) employing Rstudio 0.98.1091 (RStudio Team n.d.) and the ggplot2 package for generating graphs (Wickham 2009). The MaxQuant output file “evidence.txt” was used to compile protein groups as depicted in Figure S3; the MaxQuant “proteinGroups.txt” output file was used for analysis of the sub-cellular fractionation data presented in Figure S2 (see Table S1). Only those proteins identified in all 3 biological replicates with at least 2 razor+unique peptides were included in our “quantified proteins” dataset (Table S3). As measurement error correlates inversely with intensity, prior to p-value calculations by ANOVA, the dataset was grouped into three bins based on protein abundance (i.e. measured intensity). Control proteins for each ANOVA calculation were randomly sampled from a set of age-invariant proteins (with SILAC pair ratios or log2 fold changes ≈0) in the same abundance bin as the protein in consideration. The random sampling and ANOVA calculation for each protein was looped 100 times and used to estimate the false discovery rate (FDR). GO-term enrichment was performed using DAVID (Huang, Sherman & Lempicki 2009a; Huang, Sherman & Lempicki 2009b) unless otherwise indicated, with the full C. elegans proteome supplied by DAVID used as the background list, and plotted to reduce redundancy using ReviGo (Supek et al. 2011); Venn diagrams were generated using BioVenn (Hulsen et al. 2008). PCA analysis was done using Partek Genomics Suite (Partek Inc, St. Louis, MO). GSEA analysis was done using the algorithm from the Broad Institute (Subramanian et al. 2005) using either a user-defined gene set or the KEGG pathway database (Kanehisa et al. 2014; Kanehisa & Goto 2000). Human homologs of C. elegans genes were obtained using information from the WormBase FTP site (ftp://ftp.wormbase.org/pub/wormbase/); files containing both best BLAST hits and orthologs were used and merged into a single file using R. Microscopy images were analyzed and quantified using Cell Profiler (Lamprecht et al. 2007).
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