Genome-wide RNAi Screen for Fat Regulatory Genes in C. elegans Identifies a Proteostasis-AMPK Axis Critical for Starvation Survival

Highlights
- Opposing proteostasis pathways regulate C. elegans fat and starvation survival
- Extended starvation survival with ARS RNAi requires elevated proteasome activity
- Proteasome RNAi shortens starvation survival and blocks AMPK activation
- AMPK is the critical determinant of starvation survival with altered proteostasis

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In Brief
Using a genome-wide screen for fat regulatory genes in C. elegans, Webster et al. define a proteostasis-AMPK signaling axis that is central to organismal starvation defenses. The results suggest that enhanced survival under elevated proteasome activity is due to AMPK activation and not to increased fuel availability.
SUMMARY

Organisms must execute metabolic defenses to survive nutrient deprivation. We performed a genome-wide RNAi screen in Caenorhabditis elegans to identify fat regulatory genes indispensable for starvation resistance. Here, we show that opposing proteostasis pathways are principal determinants of starvation survival. Reduced function of cytoplasmic aminoacyl tRNA synthetases (ARS genes) increases fat mass and extends starvation survival, whereas reduced proteasomal function reduces fat and starvation survival. These opposing pathways converge on AMP-activated protein kinase (AMPK) as the critical effector of starvation defenses. Extended starvation survival in ARS deficiency is dependent upon increased proteasome-mediated activation of AMPK. When the proteasome is inhibited, neither starvation nor ARS deficiency can fully activate AMPK, leading to greatly diminished starvation survival. Thus, activity of the proteasome and AMPK are mechanistically linked and highly correlated with starvation resistance. Conversely, aberrant activation of the proteostasis-AMPK axis during nutritional excess may have implications for obesity and cardiometabolic diseases.

INTRODUCTION

Starvation is among the most ancient of selection pressures. In response to repeated cycles of nutrient deprivation, organisms have evolved elaborate defenses that permit survival through periods of starvation. A major starvation defense is the ability to store long-term energy stores as triglyceride (Hardie, 2012; Viscarra and Ortiz, 2013). In humans, the majority (85%) of available energy is stored as adipose triglyceride, even in normal weight individuals (Cahill et al., 1968; Saudek and Felig, 1976). Perhaps the best evidence that adipose triglyceride stores represent a starvation defense is a report that obese individuals can survive for nearly 1 year under total starvation without complications (Thomson et al., 1966).

Total body protein also undergoes cyclic changes with periods of fasting and feeding (Millward et al., 1996), but a far more modest amount of available calories are present in body protein. Further, liberating these calories can be achieved, but with the potential for great harm (Cahill et al., 1968; Saudek and Felig, 1976). Indeed, death from starvation occurs in humans when approximately half of body protein stores have been depleted (Garrow et al., 1965).

These observations indicate that it is not only lipid metabolism that is critical for starvation defense, rather, orchestration of both protein and lipid anabolism and catabolism. In multicellular organisms, this balance is achieved in part by changes in endocrine signals that signal nutrient adequacy. One major signal indicating nutrient adequacy across phylogeny is the insulin/insulin-like growth factor (IGF) signaling pathway. Whether more ancient systems exist to orchestrate starvation defenses remains an open question.

The highly conserved, cellular energy sensing serine-threonine protein kinase, AMPK, activates catabolic pathways when nutrients are scarce (Hardie, 2011; Viollet et al., 2003). In response to signals indicating nutrient sufficiency, AMPK is inhibited, restraining lipid oxidation and protein salvage pathways (Cuervo et al., 2005; Kang et al., 2007; Singh et al., 2009). Alternatively, when the cellular AMP to ATP ratio rises, AMPK is activated, curtailing energy intensive processes such as lipogenesis and activating lipid catabolism, proteolysis, and starvation defenses (Viollet et al., 2003, 2009; Cunningham et al., 2014; Narbonne and Roy, 2009).
Figure 1. Genome-wide RNAi Screen Identifies Enrichment of Proteostasis Genes

(A) Schematic of genome-wide RNAi screen completed in *eri-1* animals and summary of results. See also Table S1.
(B) GSEA using KEGG identifiers indicates a significant enrichment of ARS genes among high-fat-inducing RNAi clones. See also Table S2.

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In spite of advances in our understanding of signaling pathways that modulate anabolism and catabolism, comprehensive understanding of the genetic landscape regulating starvation resistance in multicellular organisms is lacking. In the present study, we identify a starvation defense system that links proteostasis to lipid metabolism, through a validated genomewide RNAi screening methodology for studying fat mass in *C. elegans* (Pino et al., 2013). Prior screens for fat regulators in the worm (Ashari et al., 2003), while illuminating, suffered from methodological difficulties whereby fluorescent dyes fed to living worms were primarily localized to the acidified lysosome related organelle rather than to lipid droplets, thus complicating the discovery of true lipid regulators (Brooks et al., 2009; O’Rourke et al., 2009; Yen et al., 2010). Here, we describe an elaborate interplay between proteostasis and lipid metabolism that enables starvation resistance through efficient storage and utilization of energy rich triglyceride. Inactivation of any of the entire complement of cytoplasmic aminoacyl tRNA synthetases (ARS genes) leads to a massive 200%–400% increase in triacylglycerol stores, increased lipid synthesis, decreased lipid oxidation, and a doubling to tripling of starvation survival. Conversely, we identify the 26S proteasomal machinery, one of two major proteolytic mechanisms in starvation (Finn and Dice, 2006), as being indispensable for normal accrual of fat mass and defense against starvation. We find that increased starvation survival upon ARS gene RNAi is strictly dependent upon increased proteasomal activity, and further, completely abolished upon inhibition of AMPK. Strikingly, altered proteasomal activity governs starvation survival not through direct provision of essential energy and nutrients by protein turnover, but through acting as a critical, upstream regulator of AMPK. These results highlight the complexity of starvation defenses and identify cross talk between ancient genetic systems involved in proteostasis, long-term triglyceride energy stores, and starvation defenses activated by the master energy sensor AMPK.

**RESULTS**

**Genome-wide RNAi Screen Identifies Protein Translation and Proteasome Genes as Reciprocal Regulators of Whole-Body Fat Mass**

We conducted a quantitative screen using a validated, lipid staining technique for gene knockdowns that impact body fat mass with RNAi libraries which target greater than 94% of the *C. elegans* genome (Kamath et al., 2003; Kim et al., 2005; Pino et al., 2013). With a tiered screening strategy, we identified 514 RNAi (targeting 475 unique genes) that cause significantly increased or decreased fat accumulation (Figure 1A). 386 RNAi clones increased fat mass, whereas 128 RNAi clones decreased fat mass (Figure 1A; Table S1). All RNAi clones were sequence verified. The screening strategy was validated by recovery of low-fat RNAi corresponding to genes known to positively regulate lipogenesis such as *sbo-1* (sterol regulatory element-binding protein-1), *pod-2* (acetyl-CoA carboxylase), and *fasn-1* (fatty acid synthase) (Table S1) (McKay et al., 2003). Using a Kyoto Encyclopedia of Genes and Genomes (KEGG)-based classification system for gene set enrichment analysis (GSEA; Falk et al., 2008; Huang et al., 2009), we identified RNAi to ARS genes as the most overrepresented pathway within the dataset causing elevated fat mass (Figures 1B and 1C; Table S2). Conversely, RNAi to core proteasome components were the most overrepresented class among low-fat gene inactivations (Figures 1D and 1E; Table S3). Other overrepresented classes included purine and pyrimidine synthesis genes among high-fat-inducing RNAi and oxidative phosphorylation genes among low-fat-inducing RNAi (Tables S2 and S3).

Strikingly, 14 of 21 cytoplasmic ARS genes, whose canonical cellular role is to charge the tRNA molecule with its cognate amino acid (Ibba and Soll, 2000), emerged as high-fat-inducing RNAi (Figure 1C). We confirmed an increase in triacylglycerol stores using quantitative biochemical analysis of lipids in animals treated with trytophanyl-tRNA synthetase (war-1) RNAi and animals bearing a temperature-sensitive mutation in valyl-tRNA synthetase (vars-2) (Figure S1A, vars-2(bn2ts) also known as *glp-4*) (Rastogi et al., 2015). We similarly witnessed an increase in fat when worms were treated with the translation elongation inhibitor cycloheximide (Figure S1B) (Schneider-Poetsch et al., 2010).

Proteasomal gene inactivations were among the most statistically enriched functional classes contributing to low-fat mass. The ubiquitin proteasome system (UPS) serves as a key cellular degradation pathway required to maintain proteostasis (Lee et al., 2013). RNAi of members of five different classes of the UPS, belonging to the β-type subunits, β-type subunits, the 19S non-ATP regulatory subunit, ATPases associated with diverse cellular activities (AAA) ATPase subunits, along with a ubiquitin activating enzyme, leads to significant reduction in lipid accumulation (Figure 1E). Exposure to the proteasomal inhibitor bortezomib also leads to significant reductions in fat mass (Figure S1E). These data suggest that protein synthesis and protein breakdown are opposing proteostasis pathways that antagonistically regulate normal triglyceride stores.

**Inactivation of ARS Genes Protects against Fat Mass Loss during Starvation and Exercise**

To determine which of our fat regulatory genes played a role in mobilizing triacylglycerol stores in response to nutrient deprivation, we inactivated our 475 fat regulatory genes by RNAi from early larval development to young adulthood in wild-type *C. elegans*, thereafter subjecting animals to starvation and...
exercise in minimal media or pure starvation (by including the chemical paralytic 2,3-butanedione monoxime (BDM) which prevents muscle contraction) (Figure 2A). Starvation or starvation plus exercise for 16 hr is sufficient to reduce fat mass in control animals by 23% and 42%, respectively (Figure S2A).

As we conducted this analysis in wild-type animals, this also provided the opportunity to analyze concordance of RNAi fat phenotypes between the initial screen in eri-1 animals and wild-type animals in the fed state. Overall, 78% of fat regulatory gene RNAi show identical directionality between eri-1 and N2 wild-type animals. Linear regression analysis indicates a highly significant positive correlation between values obtained in N2 wild-type versus eri-1 fat levels ($R^2 = 0.53$, $p < 0.0001$). Effects of any given RNAi on fat mass are less pronounced in wild-type than eri-1 animals (clusters 1 and 7; Figures 2B and S2B; Table S4). These clusters contained the entire membership of cytoplasmic ARS genes previously identified to be high-fat inducing in our primary screen (Figures 2B, 2C, and S2C; Table S4). We confirmed that increased fat mass fails to decrement with starvation following ARS gene RNAi by lipid staining and quantitative lipid biochemistry (Figures S2D and S2E). This response was not a general feature of all high-fat states, as animals deficient in insulin receptor ortholog daf-2 have elevated triacylglycerol mass at baseline, but showed the predicted decrement in fat mass upon starvation (Figure 2C).

We next hypothesized that fat regulatory gene inactivations would be enriched for those that impact starvation survival. Accordingly, following 3 days of RNAi from the synchronous first larval stage (L1), wild-type animals were washed free from food, subjected to starvation in minimal media, and examined at the onset of and at two discreet time points of starvation, one at day 5 (when virtually no vector RNAi-treated animals are dead) and another at day 19 of starvation (when more than 99% of vector RNAi-treated animals are dead) and another at day 19 of starvation (when more than 99% of vector RNAi-treated animals are dead). Over 40% of genes showed some alteration in starvation survival versus vector control at these two screening time points (Figure 2B; Table S4). Many RNAi exhibiting extended starvation survival grouped in clusters also exhibited more judicious rationing of fat mass (Figure 2B, “Starvation Resistant” clusters). Alternatively, more than 70% of genes in low-fat clusters, which includes UPS genes, exhibited diminished starvation survival (Figures 2B and 2D, “Starvation Sensitive” clusters).

**Inactivation of ARS Genes Prolongs Starvation Survival**
Longitudinal starvation survival analysis indicated that starvation survival is increased up to nearly triple the mean of control animals upon RNAi of wars-1, vars-2, and lysyl-tRNA synthetase (kars-1) (Figure 3A; Table S5). Genetic inactivation of vars-2 similarly increased starvation survival (Figure 3B; Table S5). None of the ARS RNAi tested led to prolongation of lifespan under fed conditions except for a 12.9% increase for wars-1 RNAi (Figure S3A; Table S6), that paled in comparison to the ~150% increase in starvation survival, indicating a starvation survival advantage specific to starvation. Following inactivation of wars-1 by RNAi, we observed a 40% decrease in oxygen consumption (Figure S3B) and a concomitant increase in fatty acid synthase (fasn-1) mRNA, consistent with increased lipid synthesis (Figure S3C). These changes occurred without an increase in food consumption rate, as indicated by the pharyngeal pumping rate (Figure S3D). A small, but significant, decrease in pharyngeal pumping is seen in ARS-deficient animals, which could indicate negative feedback in the face of high body fat mass stores.

As translation consumes approximately 25%–35% of cellular energy stores (Rolf and Brown, 1997; Siems et al., 1992), we surmised that the excess fat mass of ARS-deficient animals would be dependent upon excess cellular reducing power and energy. To address this, we treated control and ARS-deficient animals with the mitochondrial uncoupling agent dinitrophenol (DNP). DNP-treated control and ARS-deficient animals both demonstrate a proportionally equivalent, dose-dependent decrease in fat mass, suggesting that excess proton-motive force at the level of the mitochondrion is partially, but not entirely, responsible for high-fat mass in ARS deficiency (Figure S3E). Finally, day 1 starved adults treated with vector RNAi show a marked extension in starvation survival that is exaggerated further upon inactivation of kars-1 when compared to identically treated animals starved at the L4 developmental stage (Figure S3F). Thus, while developmental stage at the onset of starvation is important in determining starvation survival, ARS deficiency extends survival irrespective of the paradigm used.

**Inactivation of Ubiquitin-Proteasome Genes Shortens Starvation Survival**
The very low-fat mass resulting from UPS gene inactivation (Figures 1D and 1E) screen level data indicating shortened starvation survival (Figure 2B, clusters 10–11; Table S4) led us to more closely study the role of the proteasome in starvation resistance. Longitudinal starvation analysis of animals treated with RNAi to regulatory proteasome subunit rpm-6 indicated nearly 80% shortening of mean starvation survival versus vector control (Figure 3C; Table S5). We reasoned that bortezomib should similarly reduce starvation resistance and acute treatment would allow us to examine starvation survival without potential negative pleiotropic effects on development. Indeed, acute inhibition of bortezomib at adulthood, 6 hr prior to initiation of starvation, led to ~60% shortening of mean starvation survival (Figure 3D; Table S5).

**Metabolic Shifts in ARS-Deficient Animals Are Distinct from Canonical Nutrient and Stress Sensing Pathways**
Ablation of germline stem cells in C. elegans leads to high-fat mass and increased longevity in a manner dependent upon the FoxO ortholog daf-16 (Hsin and Kenyon, 1999; Lapierre et al., 2011). A temperature-sensitive mutation in the vars-2 gene also leads to relative deficiency in germline stem cells (Beanan and Strome, 1992). To determine whether the increase in fat mass and starvation defense is similarly dependent upon loss of germline stem cells in ARS-inactivated animals,
### Figure 2. Inactivation of ARS and UPS Genes Has Opposing Effects on Fat Mass

(A) Schematic workflow of secondary RNAi screen in wild-type animals for starvation-based fat mobilization.

(B) k-means clustering of 475 fat regulatory genes into 11 groups based on fixative Nile red fat levels in conditions of fed, starved (with paralytic), or starved and exercised (no paralytic) (left, within dashed line). The alternating black and white bar to the right of the heatmap indicates cluster divisions. Immediately adjacent are the results from longitudinal starvation survival analysis, with red indicating shortened starvation survival and green indicating extended starvation survival (right, within shaded area). Starvation-resistant and starvation-sensitive gene clusters are shown. See also Figure S2 and Table S4.

(C) Fat levels by fixative Nile red fat staining of wild-type animals treated with RNAi against empty vector, the insulin-like receptor *daf-2*, tryptophanyl-tRNA synthetase *wars-1*, or valyl-tRNA synthetase *vars-2*, under fed, starved, or starved and exercised conditions. *p < 0.05 and ****p < 0.01 by two-way ANOVA with Bonferroni correction. Bars represent mean ± SEM (n = 6 biological replicates).

(D) Fat levels by fixative Nile red fat staining of wild-type animals treated with RNAi to empty vector, the indicated proteasome regulatory genes (rpn and rpt), or ubiquitin E1 ligase *uba-1*, under fed, starved, or starved and exercised conditions. ****p < 0.0001 by two-way ANOVA with Bonferroni correction. The bars represent mean ± SEM, (n = 3 biological replicates).

<table>
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<th>RNAi to</th>
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<th>Orbital Shaking</th>
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<td>Fixation Nile Red</td>
<td>Altered Starvation Metabolic Response</td>
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Starvation Gene RNAi Screening in Triplicate

- **Fed Control Worms**
- **Fed Control Worms (± Paralytic)**
- **Fed Fat RNAi Worms (± Paralytic)**
- **Starved Control Worms (± Paralytic)**
- **Starved and Exercised Control Worms (± Paralytic)**
- **Starved Fat RNAi Worms (± Paralytic)**
- **Starved Fat RNAi Worms**

Normal Response: Fat Mass Reduced

Starvation Resistant Response: No Fat Mass Reduction

Starvation Survival

- **Fed**
- **Starved**
- **Starved and Exercised**

Relative Fat Level

- **Empty Vector**
- **daf-2**
- **wars-1**
- **vars-2**

- **rpt-5**
- **rpn-11**
- **rpt-2**
- **uba-1**

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we performed RNAi against ARS genes in daf-16 null mutant animals. In contrast to daf-2/insulin receptor-deficient animals, which require daf-16/FoxO for elevated fat mass, fat mass remains elevated in ARS-deficient animals lacking daf-16 (Figures 4A and S4A). Additionally, while fat mass in germline stem cell-deficient glp-1 mutants (Priess et al., 1987) is daf-16 dependent, the elevated lipid mass of daf-16;vars-2 double mutant animals remains as high as vars-2 single mutants (Figure 4B). RNAi knockdown of canonical metabolic mediators and genes involved in insulin/phosphatidylinositol 3-kinase (PI3K) signaling, hypoxia signaling, and stress responses did not suppress the high triacylglycerol phenotype of ARS-deficient animals (Figure S4B; Table S7). In aggregate, these data suggest the metabolic shift evident in ARS-deficient animals is not dependent upon these metabolic regulatory pathways or the germline-mediated response.

Opposing ARS and UPS Pathways Exert Metabolic Effects via Action in Somatic Tissues

A compelling possibility is that the ARS and UPS genes exert opposing metabolic effects via action in the same tissues. To test this possibility and to gain insight as to the site of action of ARS and UPS genes in metabolic regulation, we made use of a mutant in the rrf-1 gene, which encodes an RNA-directed RNA polymerase required for the somatic, but not germ-line, actions of RNAi (Sijen et al., 2001). Fat levels of animals with ARS genes inactivated were decreased in rrf-1 mutants, suggesting that reduced levels of ARS gene function were necessary in somatic tissues to trigger this starvation defense pathway (Figure 4C). This change occurred in rrf-1 mutant animals despite preservation of moderate somatic and normal germline RNAi competency (Kumsta and Hansen, 2012). We also tested a ppw-1 mutant, which failed to produce a

![Graphs and figures](https://example.com/graphs.png)

**Figure 3. Inactivation of Proteostasis Machinery Results in Altered Starvation Survival**

(A) Starvation survival in minimal media of wild-type animals treated with empty vector, wars-1, vars-2, or kars-1 RNAi and fed ad libitum to the L4 larval stage prior to starvation onset.

(B) Starvation survival in minimal media of wild-type animals or vars-2(bn2ts) mutants treated with empty vector RNAi and fed ad libitum to the L4 stage.

(C) Starvation survival in minimal media of wild-type animals treated with empty vector RNAi or RNAi to proteasome regulatory subunit rpn-6.1 and fed ad libitum to the L4 stage.

(D) Starvation survival in minimal media of wild-type animals fed ad libitum with E. coli OP50 until the young adult stage, then washed into minimal media containing 5 μg/mL bortezomib or vehicle control. For all graphs, the bars represent mean ± SEM, n > 30 worms/time point measurement, and the points represent mean of 2–5 biological replicates. All ARS, UPS, and bortezomib curves are statistically different from control by linear regression analysis. See also Figure S3 and Table S5 experiments 1–5 for tabular data, replicates, and statistical analysis.
PIWI, AGO, and Zwille (PAZ)/P-element induced Wimpy testis (PIWI)-domain containing protein required for efficient germline RNAi (Tijsterman et al., 2002). The lipid levels of ARS-deficient animals remained elevated in ppw-1 mutants, confirming the essential lipid-regulatory role of ARS in the soma (Figure 4D).

We next performed UPS gene inactivation in the nrf-1 and ppw-1 mutant genetic backgrounds to identify the site of action of proteasomal genes. In an identical fashion to ARS gene inactivation, RNAi of UPS genes pas-5 and rpn-6.1 no longer led to significant decreases in fat mass when somatic RNAi efficiency is blunted, as in the nrf-1 mutant (Figure 4E). Confirming that these effects are not via action in germline stem cells, UPS gene RNAi reduced fat mass in a ppw-1 mutant in a fully penetrant manner (Figure 4F).

**26S Ubiquitin Proteasome Activity Is Central to Starvation Defense**

Given the common site of action of ARS and UPS genes in somatic tissues, we next examined mechanistic connections between the two pathways. Since low proteasomal activity leads to low-fat mass and is detrimental to starvation survival (Figures 2D and 3C), it logically follows that increased activity might provide greater resistance to starvation. We tested this by asking first whether gene knock downs of the high-fat ARS genes lead to an increase in proteasomal catalytic activity. Extracts of animals after short-term gene inactivation of vars-2 and kars-1 led to a significant increase in chymotrypsin-like and caspase-like proteasomal activities (Figures 5A and 5B), while trypsin-like activity did not show a significant increase (Figure 5C). All demonstrable proteasomal activity was suppressible by addition of the proteasomal inhibitor MG132 (data not shown).

These data suggest a genetic relationship between the ARS and UPS systems, moreover that the UPS might be epistatic to beneficial effects seen in ARS RNAi animals. To determine the nature of this relationship, vars-2 (bn2ts) mutants were treated with RNAi to genes encoding proteasomal regulatory and catalytic subunits. The elevated fat mass of vars-2 mutants was suppressed by knocking down the genes encoding UPS subunits (Figure 5D). These data confirm that the UPS is epistatic to high-fat mass seen upon inactivation of ARS genes.

To determine whether the epistasis extended to starvation survival, we performed a starvation survival analysis on ARS RNAi animals treated with bortezomib. As we previously found, RNAI knock down of kars-1, vars-2, or vars-1 leads to significant increases in starvation longevity as compared to control (Figures 5E–5G; Table S5). Inhibition of the proteasome with bortezomib significantly, and in some cases completely, reversed the extended starvation survival witnessed with ARS gene knockdown (Figures 5E–5G; Table S5). The most potent ARS gene knockdown, kars-1, nearly triples starvation survival and is unable to do so when animals are treated with bortezomib (Figure 5E). These data place the proteasome downstream of the...
ARS genes in a genetic pathway critical to organismal starvation defenses.

Proteostasis Pathways Modulate Starvation Survival through Regulation of AMPK

Our observation that the uncoupling agent DNP was unable to fully reverse metabolic shifts in ARS knockdown to a greater extent than it does in wild-type animals (Figure S3E), suggests that the proteasome activates starvation defenses through additional mechanisms, rather than scavenging proteins to be used as cellular fuels. AMPK is a master sensor of cellular energy levels that is well poised to activate cellular starvation defenses.

In order to determine whether the proteasome is mechanistically coupled to AMPK activation and subsequent starvation defenses, we exposed a population of animals to 24 hr of starvation with and without treatment with bortezomib. This period of starvation is sufficient to increase phosphorylation of the alpha subunit of AMPK on the homologous residue to mammalian AMPK.

Figure 5. The 26S Proteasome Operates Functionally Downstream of the ARSs for Starvation and Fat Mass

(A–C) Proteasomal chymotrypsin-like (A), caspase-like (B), and trypsin-like (C) activity measurements from lysates of L3 larval stage animals treated for 18 hr with empty vector, vars-2, and kars-1 RNAi and fed ad libitum (n = 3 biological replicates). *p < 0.05 and **p < 0.01 by one-way ANOVA.

(D) Fat levels as indicated by fixative Nile red staining in vars-2(bn2ts) mutant animals treated with empty vector, rpn-2, rpn-6.1, rpt-2, rpt-3, and pas-6 RNAi from L1 stage (n = 3 biological replicates). ****p < 0.0001 by one-way ANOVA.

(E–G) Starvation survival in minimal media of wild-type animals treated with empty vector and kars-1 (E), vars-2 (F), or wars-1 (G) RNAi with and without the proteasomal inhibitor bortezomib (5 μg/mL). Animals were treated with the indicated RNAi and fed ad libitum to the L4 stage prior to starvation onset. See also Table S5 experiments 3–5 for tabular data, replicates, and statistical analyses. All bars represent mean ± SEM.
threonine 172 (p-AMPK\(^{172}\)) corresponding to Thr243 in AAK-2 isoform CE31222, a required A-loop activating site phosphorylated in mammals by the upstream kinase liver kinase B1 (Lkb1) (Figures 6A and 6C) (Shaw et al., 2004; Woods et al., 2003). Treatment with bortezomib completely blunts the starvation-mediated phosphorylation of p-AMPK\(^{172}\), indicating that proteasomal activity acts as an essential upstream activator of AMPK in the response to starvation (Figures 6A and 6C).

As proteasomal activity is increased in ARS-deficient animals, we anticipated increased activation of AMPK following ARS gene knockdown. A profound and significant increase in p-AMPK\(^{172}\) was observed in animals treated with RNAi to the kars-1 gene, which was further augmented by starvation (Figures 6B and 6C). As expected, proteasomal inhibition with bortezomib prevented the ARS- and starvation-mediated increases in p-AMPK\(^{172}\).

To address our hypothesis that AMPK sits genetically downstream of the ARS genes, we analyzed starvation survival in AMPK (aak-1 and aak-2) mutant animals with gene inactivation of vars-2, wars-1, and kars-1 (Figures 6D–6I; Table S5). While knockdown of each ARS gene provides parallel starvation survival advantage to both control and aak-1 mutant animals, loss of aak-2 leads to severe deficiency in starvation survival and prevents any increase in starvation survival upon RNAi of ARS genes. Thus, similar to proteasomal inhibition, loss of function of aak-2 leads to profound reductions in starvation survival. As expected, double mutants lacking aak-1 and aak-2 show further shortened starvation survival, and RNAi to kars-1 and other ARS genes is unable to extend starvation survival in these mutants (Figure 6J; Table S5).

Surprisingly, neither acute proteasomal inhibition with bortezomib upon initiation of starvation or genetic deficiency in AMPK affected the rate of fat mass loss during starvation (Figure S3). These data suggest, in contrast to prior studies of aak-2 mutant animals (Narbonne and Roy, 2009), that loss of UPS activity and AMPK activity do not affect starvation survival by strictly affecting the rate of fat mass loss.

These data suggest that a principal mechanism by which proteasomal inhibition with bortezomib shortens starvation survival is through preventing activation of AMPK. If this is the case, then genetic activation of AMPK should prevent bortezomib from reducing starvation survival. Indeed, a C. elegans transgenic strain carrying a gain-of-function aak-2 isoform (where the equivalent A-loop phosphorylation site to T172 is mutated to an phospho-mimic aspartate residue) was resistant to reductions in starvation survival upon bortezomib treatment (Figure 6K; Table S5). Taken together, the above data place AMPK genetically downstream of the proteasome and of proteostasis genes in general in regulating starvation survival (Figure 7).

**DISCUSSION**

In this study, we find that hundreds of genes contribute to regulation of body energy stores and survival under nutrient deprivation. Beginning with an unbiased study for genes that activate triglyceride accumulation, we identified 475 gene inactivations that both increase and decrease fat mass. Among those are genes belonging to the aminacyl tRNA synthetase family, where knockdown by RNAi leads to massive lipid accumulation and prolonged starvation survival. We find that proteasomal activity is increased in starvation-resistant ARS-deficient animals, and that proper function of the proteasome is required for starvation survival in ARS-deficient and wild-type animals alike. In support of the importance of the proteasome in starvation defenses, the primary screen also identified a large group of UPS encoding genes among low-fat and starvation-susceptible gene inactivations. Remarkably, these opposing proteostasis pathways do not govern starvation survival strictly based upon provision of cellular energy, rather, both converge on AMPK, and AMPK is the major determinant of starvation survival. We find that decreased proteasomal activity prohibits AMPK from responding to low cellular energy stores. Thus, enhanced AMPK activity mechanistically explains the enhanced survival of ARS deficiency, whereas reduced AMPK activity is responsible for shortened survival of proteasome deficiency. We suggest that cellular sensing and integration of proteostasis is communicated to metabolic shifts and survival in times of nutrient deprivation through AMPK.

**Cellular Proteostasis Pathways Are Major Determinants of Starvation Resistance**

In general, for animals to survive prolonged periods of starvation, metabolic adaptation is necessary in the form of increased nutrient storage and greater efficiency of energy utilization (Hardie, 2012). Previous studies indicate that inhibition of protein translation and elongation can defend against nutrient deprivation at the cellular level (Lepivier et al., 2013). Here, we show that disabling ARS encoding genes activates starvation defenses at the organismal level, including a massive shift toward increased triglyceride storage and decreased utilization. These data are consistent with observations in Drosophila, where genetic activation of protein synthesis reduces fat storage and starvation survival (Telemans et al., 2005). Conversely, reduced function of UPS genes decreases lipid accumulation and survival under conditions of nutrient deprivation. Mutations in the gene encoding immunoproteasome subunit proteasome subunit meta 8 (PSMB8) lead to a rare form of human lipodystrophy, whereby adipocytes cannot accumulate normal triglyceride stores (Agarwal et al., 2010; Arima et al., 2011; Arimochi et al., 2016; Kitamura et al., 2011). We suggest that regulation of long-term energy stores is an essential and conserved role of the proteasome across phylogeny.

We find low-fat mass evident with life-long proteasomal inhibition is not critical for shortened starvation survival. Indeed, our data indicate that AMPK activation is the essential output of proteasomal initiation and is necessary for starvation survival rather than fat mass per se. AMPK is necessary for survival under nutrient deprivation in arrested L1 and dauer larvae (Baugh, 2013; Baugh and Sternberg, 2006; Fukuyama et al., 2012; Narbonne and Roy, 2009). Further, in spite of massive lipid stores, we found that inhibition of the proteasome or AMPK completely abrogates the survival benefit evident with ARS gene deficiency. We also did not find altered rates of fat mass loss with acute inhibition of the proteasome or with genetic aak-2 deficiency,
indicating that inactivation of this UPS-AMPK axis affects starvation survival in a manner independent of the macroscopically apparent rate of overall lipid utilization. This contrasts with one previous study indicating accelerated fat mass loss with starvation in C. elegans aak-2 dauer larvae (Narbonne and Roy, 2009), but is in agreement with findings of Cunningham et al. (2014). Nonetheless, our results suggest that fat mass utilization can only extend starvation survival if the proteasome and AMPK are active, and that additional metabolic processes other than gross lipolysis and fatty acid oxidation are key targets of proteasome and AMPK activity during starvation. In aggregate, these data suggest that dysregulation of nutrient utilization during starvation rather than accumulated fat stores are the critical determinant of starvation survival. However, these data are not inconsistent with the idea that
accumulated triglyceride facilitates starvation survival. We suggest that ARS-deficient animals are extremely starvation resistant due to increased triglyceride stores and mechanisms mediated by activated AMPK.

The opposing proteostasis pathways identified highlight two groups of genes representing not only opposing lipid and starvation phenotypes when knocked down, but also of opposing molecular machines. The ARS genes are intimately involved in an anabolic process as protein translation, and the proteasome is involved in protein catabolism. We find that these two physiologically and molecularly distinct pathways are mechanistically linked. ARS gene inactivation leads to activation of starvation defenses in a manner completely dependent upon elevated proteasomal activity. Conversely, while it is possible that proteasome inhibition leads to sickness rather than true starvation sensitivity, the fact that ARS gene inactivations lead to elevated proteasome activity and that elevated activity contributes to starvation defenses argues against this possibility.

A large body of literature indicates reciprocal regulation of protein translation and protein turnover during starvation (Amherdt et al., 1974; Cahill et al., 1968; Finn and Dice, 2006; Millward et al., 1996; Saudek and Felig, 1976). Under conditions where insulin and growth factors are scant and nutrients are deficient, protein translation is curtailed and protein turnover is activated (the latter by either the UPS or autophagy pathway) (Cahill et al., 1968; Finn and Dice, 2006; Kim and Lee, 2014; Millward et al., 1996; Saudek and Felig, 1976). Our findings, taken together with the historical starvation and protein homeostasis literature, suggest that the proteasome is activated as a consequence of translational inhibition. We hypothesize the existence of an as yet unidentified cellular signal, present under fed conditions when protein translation is activated, serves to inhibit the UPS. When animals are starved, this signal is no longer available, triggering UPS activity.

**AMPK Integrates Starvation Responses Received by Altered Proteostasis**

We observe significant increases in zAMPK<sup>T172</sup> phosphorylation upon ARS gene inhibition. Normally, activation of AMPK in the face of low ATP levels is associated with lipid catabolism to supply the cell with energy (Hardie, 2003, 2011; Hardie et al., 2012; Viollet et al., 2009). In the case of ARS gene inactivation, triglyceride levels are increased in spite of the observed increase in AMPK activation. Consistent with these findings, activation of AMPK in white adipose tissue suppresses lipolysis while AMPK-deficient mice have small adipocytes (Daval et al., 2005). Our results indicate that AMPK activation in starvation, which is exaggerated in ARS deficiency, prolongs starvation survival in a manner independent of baseline fat storage or the rate of fat mass loss in starvation. These results contrast with prior studies of the role of AMPK in fat storage and starvation survival (Cunningham et al., 2014; Narbonne and Roy, 2009). We conclude, at the level of the whole organism, that reduced ARS gene function leads to elevated UPS activity, which is necessary for starvation-induced activation of AMPK.

In cultured mammalian cells, UPS inhibition has been reported to activate AMPK, possibly through relieving proteasomal degradation of AMPK subunits (Jiang et al., 2015; Zungu et al., 2011). Two likely possibilities exist to explain why this phenomenon is not orthologous between cells in culture and *C. elegans*: (1) that cells in culture are not representative of in vivo signaling and physiology and/or (2) that UPS inhibition of AMPK in *C. elegans* involves a multisite signaling relay. AMPK has been shown to act in the nervous system and also in peripheral tissues in order to modulate aging (Burkewitz et al., 2015; Cunningham et al., 2014). It remains to be seen whether distinct sites of action of AMPK are required for efficient starvation defense at the organismal level versus cell autonomous effects.
Altered Proteostasis Activates an Ancient Starvation Defense Pathway

We suggest that starvation resistance is due to specific metabolic adaptation to decreased protein translation, which serves to activate AMPK in a proteasome-dependent manner following ARS gene knockdown. Coupled to increased fat levels, elevated UPS-AMPK activity in ARS-deficient animals permits extended starvation survival. More generally, it is easy to speculate that evolution favored mechanistic links between catabolism of protein and lipid in the face of starvation. Our data suggest that one such link could be activation of AMPK in a manner dependent upon the proteasome. In circumstances when protein turnover through the UPS is activated, activation of AMPK would serve to upregulate starvation defenses. The adaptive nature of such a shift is readily evident. While in higher eukaryotes, endocrine signaling pathways have evolved that balance protein and lipid anabolic and catabolic pathways, it is tempting to think that the protein degradation-AMPK link we report here represents an ancient adaptation to repeated cycles of feast and famine.

EXPERIMENTAL PROCEDURES

Strains and Maintenance

C. elegans were routinely grown and maintained on nematode growth media (NGM) seeded with Escherichia coli OP50. The wild-type strain was N2 Bristol. The following C. elegans strains were used: MGH55 daf-16(mgD174) 3x, CB4037 glp-1(e1491), MG1627 aak-2(pk524) 4x, MG1627 aak-1(tm1944) 4x, MG16276 aak-1(tm1944)/aak-2(pk524), NL2086 ror-1(pk1417) 2x, NL3511 pcp-1(pk1425) 2x, GR1373 eri-1(mg366) X, AGD487 uhr-1(e490), and SS104 vars-2(glp-4(e1271)) 2x (vars-2(glp-4(e1271)) encodes missense mutation Gly296Asp) [Rastogi et al., 2015].

Genome-wide Fat RNAi Screen

RNAi analysis for fat regulatory genes was conducted as previously described (Pino et al., 2013) in the C. elegans. Phenotypes from 18,179 RNAi clones were analyzed in duplicate for fixative Nile red fat levels and compared to animals treated with empty RNAi vector. From this, 1,364 clones were selected as having an altered fat phenotype based upon one replicate resulting in fat levels ≥ 2 SD away or both replicates ≥ 1 SD away from the empty vector mean. In the secondary screen, the 1,364 clone sublibrary was assayed for significant fat phenotypes in the eir-1 strain for an additional six biological replicates, identifying 475 fat regulatory genes (represented by 514 independent, sequence verified RNAi clones) in the Starvation and Exercise Screen, the 1,364 clone sublibrary was assayed for significant fat phenotypes away or both replicates.

Starvation and Exercise Screen

N2 animals fed individually on 475 fat regulatory gene RNAi library were either Starvation and Exercise Screen, the 1,364 clone sublibrary was assayed for significant fat phenotypes away or both replicates. Starvation and Exercise Screen, the 1,364 clone sublibrary was assayed for significant fat phenotypes away or both replicates.

Starvation Survival Analysis

For the starvation analysis screen, we prepared 96-well plates with the 475 fat regulatory gene RNAi library either harvested at L4 stage (fed), or washed twice and transferred to fresh 150 μL M9 minimal liquid media in 96-well U-bottom plates (falcon). Animals were suspended in either M9 (starved and exercised) or M9 containing 100 mM BDM (Sigma) paralytic agent (starved). Starvation was carried out at 20°C for 16-20 hr with 90 revolutions per minute (RPM) orbital shaking to prevent sedimentation of worms (n = 4 biological replicates). Clustering was performed using the R/Bioconductor k-means function, using a least-squares analysis to determine the number of clusters.

Body fat mass was measured as previously described (Pino et al., 2013). All analyses were performed on identically staged adult worms (unless otherwise stated). Body fat mass was scored by analysis of Nile red-stained C. elegans lipid droplets using MATLAB scripts, which use bright field image processing to detect worms and the GFP/ITC channel for quantitation.

Biochemical Triacylglycerol to Phospholipid Measurement

Triacylglycerol to phospholipid (TG/PL) was determined as previously described on 5,000 identically staged adult worms (unless otherwise stated) by solid phase extraction (SPE), fatty acid methyl ester (FAME) derivitization, and gas chromatography-mass spectrometry (GC/MS) analysis (Pino et al., 2013). Normalized total abundance of each lipid species was used for TG/PL calculations.

Proteasome Activity Measurement

Proteasome activity was measured as previously described (Vilchez et al., 2012). Briefly, 5,000 synchronized L2/L3 C. elegans larvae were collected and washed three times in M9 buffer, resuspended in 600 μL of lysis buffer (250 mM sucrose, 50 mM Tris pH 7.4, 5 mM MgCl₂, 0.5 mM EDTA, 2 mM ATP, and 1 mM DTT), and subjected to two rounds of 20 s high intensity pulse sonication on ice. Lyases were centrifuged at 10,000 g for 20 min at 4°C. The soluble fraction with exception of the surface layer was isolated and added to AMG substrates (Enzo Life Sciences) to monitor the chymotrypsin-like, caspase-like, and trypsin-like enzymatic activities. The reactions were carried out in 100 μL of reaction buffer (50 mM Tris pH 7.5, 1 mM MgCl₂, 25 mM KCl, 10 mM NaCl and 75 μM AMCA substrate, and 50–150 μg of total protein), and three observations were made per sample over the course of 40 min on a SpectraMax M3 ( Molecular Devices) microplate reader at 20°C using excitation 360 nm and emission 460 nm. Proteasome inhibitor MG132 was supplied at 50 μM to determine background rates of non-proteasomal substrate hydrolysis.

Pharyngeal Pumping Rates

Pharyngeal pumping rates were determined for 10–12 synchronized, well-fed day late L4 stage animals per strain/RNAi grown on the same food source using a Canon Vixia HF S30 HD video camera attached to a Nikon SMZ1500 microscope. Pharyngeal contractions in 20 s time periods were counted using iMovie (Apple) with 50 frame-per-second capture rate slowed down to 1/4 speed, and pumping rates per minute were calculated.

Oxygen Consumption

Synchronized, late L4 stage worms were washed three times in M9 medium. Oxygen consumption of 2,500 worms was measured immediately in 1.5 mL of M9 medium using a Strathkelvin 928 6-Channel Clark electrode respirometer (n = 6 per strain/RNAi, normalized to total dry weight).

Longevity Assay

Longevity assays were performed as previously described (Soukas et al., 2009). Briefly, synchronous L1 animals were placed on RNAi plates seeded with the appropriate RNAi clone. As day 1 adults, 100 animals per genotype were transferred to fresh RNAi plates with the corresponding RNAi supplemented with 50 μM FUDR solution. Live, dead, and censored worms were calculated every 2–3 days in the worm populations by scoring movement with gentle prodding when necessary. Data were analyzed and statistics performed through OASIS (http://sbi.postech.ac.kr/oasis/surv).

Altered Proteostasis Activates an Ancient Starvation Defense Pathway

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Statistical Analysis

Sample sizes were chosen to detect a difference of 50% assuming a SD of 20% with a p value of 0.05 and a power of 80%. This equated to a minimum of biological triplicates, and often experiments were repeated in 4x to 6x biological replicates, as indicated in the figure legends. Statistical tests performed to compare groups are indicated in each figure legend. No samples were excluded from the analysis. All data were blinded and only unblinded following completion of the analyses. No strict randomization was conducted, as identical populations of synchronous C. elegans were used for each experiment. GraphPad Prism software was used for all statistical analyses unless otherwise indicated.

SUPPLEMENTAL INFORMATION

Supplemental Information includes five figures and seven tables and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2017.06.068.

AUTHOR CONTRIBUTIONS


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