Gluconeogenesis and PEPCK are critical components of healthy aging and dietary restriction life extension

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Abstract

High glucose diets are unhealthy, although the mechanisms by which elevated glucose is harmful to whole animal physiology are not well understood. In Caenorhabditis elegans, high glucose shortens lifespan, while chemically inflicted glucose restriction promotes longevity. We investigated the impact of glucose metabolism on aging quality (maintained locomotory capacity and median lifespan) and found that, in addition to shortening lifespan, excess glucose negatively impacts locomotory healthspan. Conversely, disrupting glucose utilization by knockdown of glycolysis-specific genes results in large mid-age physical improvements via a mechanism that requires the FOXO transcription factor DAF-16. Adult locomotory capacity is extended by glycolysis disruption, but maximum lifespan is not, indicating that limiting glycolysis can increase the proportion of life spent in mobility health. We also considered the largely ignored role of glucose biosynthesis (gluconeogenesis) in adult health. Directed perturbations of gluconeogenic genes that specify single direction enzymatic reactions for glucose synthesis decrease locomotory healthspan, suggesting that gluconeogenesis is needed for healthy aging. Consistent with this idea, overexpression of the central gluconeogenic gene pck-2 (encoding PEPCK) increases health measures via a mechanism that requires DAF-16 to promote pck-2 expression in specific intestinal cells. Dietary restriction also features DAF-16-dependent pck-2 expression in the intestine, and the healthspan benefits conferred by dietary restriction require pck-2. Together, our results describe a new paradigm in which nutritional signals engage gluconeogenesis to influence aging quality via DAF-16. These data underscore the idea that promotion of gluconeogenesis might be an unappreciated goal for healthy aging and could constitute a novel target for pharmacological interventions that counter high glucose consequences, including diabetes.

Author summary

It is known that high levels of dietary sugar can negatively impact human health, but the mechanisms underlying this remain unclear. Here we use the facile Caenorhabditis elegans genetic model to extend understanding of the impact of glucose and glucose metabolism...
on health and aging. We show that the two opposing glucose metabolism pathways–glycolysis and gluconeogenesis–have dramatically opposite effects on health: glycolytic activity responsible for sugar catabolism is detrimental, but driving gluconeogenesis promotes healthy aging. The powerful longevity regulator DAF-16 is required for the healthspan effects of gluconeogenesis. Our data highlight the intriguing possibility that driving the biosynthetic gluconeogenesis pathway could be a novel strategy for healthspan promotion. Indeed, we find that increasing levels of the core gluconeogenic enzyme PEPCK (PCK-2) in just a few intestinal cells can increase overall health in a DAF-16-dependent manner. Dietary restriction, which can promote health and longevity across species, increases PCK-2 levels in the intestine via DAF-16, and PCK-2 is required for the health benefits seen when calories are limited. Our results define gluconeogenic metabolism as a key component of healthy aging, and suggest that interventions that promote gluconeogenesis may help combat the onset of age-related diseases, including diabetes.

Introduction

How to live at peak health over adult life is a challenge of our time. Successful aging, defined by some as the prolonged maintenance of physical and cognitive function coupled with avoidance of debilitating disease, can most certainly be influenced by activity and diet [1]. At odds with the goal of healthy aging, the “American diet” has been criticized for high sugar and high fat content that promotes muscular, cardiac, immune, and neuronal decline [2–8]. Diets rich in carbohydrates, which readily break down to glucose, can exacerbate diabetes health risks [9] and lead to sarcopenic muscle mass loss [10–12]. Although epidemiological studies support that high sugar diets impair health, tightly controlled studies of metabolic influences on aging quality are difficult to conduct in the human population, which is heterogeneous in genetics, activity, and food consumption patterns.

Animal models have been highly instructive regarding conserved metabolic states that promote or defer age-associated decline. In the facile aging model Caenorhabditis elegans, high glucose in growth medium shortens lifespan independently of associated osmotic change or glucose metabolism by bacterial food, in part via insulin-like signaling [13–20]. The insulin/IGF-1 receptor ortholog DAF-2 activates the AGE-1 PI3 kinase to signal downstream to inhibit the activity of longevity-promoting transcription factors including DAF-16/FOXO. Glucose stimulates insulin signaling to decrease lifespan in part by inhibiting DAF-16 [16, 18, 19]. Conversely, low insulin signaling activity activates DAF-16-mediated transcription to promote longevity [21–28].

Restricting glucose use can confer health benefits. For example, limiting C. elegans glucose catabolism by administering non-metabolizable 2-deoxyglucose results in longevity [19, 29]. More generally, dietary restriction (DR), a physiological state induced by reduced caloric intake, increases lifespan across species. In mammals, DR reduces blood glucose and insulin levels, and decreases glycolytic gene expression [30–32]. In C. elegans, complex pathways can induce DR benefits of longevity and prolonged vigor [33–43]. Still, several methods of food limitation extend lifespan via a DAF-16-dependent mechanism, suggesting that DR increases longevity by engaging a mechanism linked to the insulin signaling pathway under at least some conditions [34, 38, 40, 43].

How glucose impacts tissue-specific features of age-associated decline, such as sarcopenic loss of muscle mass and strength, is a central issue in healthy maintenance that remains poorly understood. Likewise, metabolic circuits linking glucose, DR, and insulin signaling remain to
be fully elaborated [17]. To begin to address these relationships, we examined the impact of genetically perturbing glycolytic and gluconeogenic flux on maintained *C. elegans* muscle function and on survival during adult life.

Here we report on how glucose metabolism intersects with longevity pathways to influence aging quality. We show that high glucose levels and high glycolytic activity negatively impact adult locomotory and general health, while conversely, gluconeogenic activity promotes maintenance of healthy muscle function and mid-life survival. Conserved transcription factor DAF-16 is crucial for these effects: glycolysis inhibits DAF-16 to compromise healthspan, while DAF-16 signaling directly promotes gluconeogenic gene expression to extend healthspan. DAF-16-dependent expression of gluconeogenesis-specific enzyme PEPCK PCK-2 in specific intestinal cells is critical for this regulation, suggesting that metabolic regulation in specific cells drives animal-wide health. Multiple DR pathways elevate gluconeogenic gene expression via DAF-16, and we show this gluconeogenic gene expression is required for extended healthspan under DR. Our results demonstrate that interventions that promote gluconeogenic metabolism can improve overall health, possibly by inducing a DR-like physiological state. We suggest that interventions that promote gluconeogenesis constitute a novel strategy for combating age-associated diseases and accelerated aging-related glucose homeostasis dysregulation.

**Results**

**High glucose accelerates sarcopenic decline of muscle-controlled *C. elegans* behaviors**

Elevated glucose systemically impairs human health with consequences particularly potent in middle to older age. The mechanisms by which high glucose compromises individual tissue functions remain poorly understood at the molecular level. In particular, how high glucose diets relate to conserved degenerative processes such as sarcopenia—the progressive loss of muscle mass and muscle strength over time that lowers quality of life and increases frailty—is unclear.

*C. elegans* locomotion rates decline with age in a manner that is roughly correlated with the degree of sarcopenic muscle deterioration [44, 45] but that also involves neuronal health [46, 47]. Cellular phenotypes associated with human muscle skeletal decline, including disorganization and loss of sarcomeres as well as fat infiltration of muscle, are observed in bodywall muscles of aging nematodes that progressively slow in locomotory capacity. Likewise, *C. elegans* cardiac-like muscle that makes up the pharynx undergoes significant structural and functional decline with age [45, 48, 49].

To examine the impact of high glucose on locomotory ability, we exposed wild-type *C. elegans* to increasing levels of glucose and recorded swim vigor in older age. (Swim behavior is directed by worm skeletal muscle-like bodywall muscle [50, 51]; here and throughout the text we use a swim vigor measure (number of head bends/30 sec.) as a measure of locomotion capacity). We found that 4% glucose media impairs swimming ability of young adults (5 days old, Supplemental S1A Fig); middle-aged adults (Fig 1A, left-hand graph, 8 days old), and animals of more advanced age (13 days old, Supplemental Fig 1B); 2% glucose, which has been shown to decrease lifespan [16, 20, 52], did not impact swimming in our hands (Supplemental S1C Fig). In addition, we found that exposure to high glucose reduces older age pharyngeal pumping mediated by cardiac-like pharyngeal muscle (Fig 1A, right-hand graph; 5 day old measure scored because pharyngeal muscle function declines faster than bodywall muscle). Together, these outcomes reveal a progeric influence of high glucose on physical activity and muscle function reminiscent of that suggested for glucose impact on muscle-associated decline.
Fig 1. Disrupting Glycolysis-promoting Genes Extends C. elegans Healthspan, Whereas High Glucose and Gluconeogenic Gene Disruptions Shorten Healthspan. (A) Left-hand graph: swimming rates for WT animals raised on control media or media containing 4% glucose. We recorded the number of head bends (i.e., head “thresholds”) / 30 seconds for individual animals placed in liquid media on day 8 of life. 4% glucose significantly decreases WT locomotory ability (**P = 0.0005, unpaired t test; averages from three trials, n = 30 animals per condition; error bars represent SEM). Trials with 5 and 13 day old animals also showed impaired older age locomotion in 4% glucose (Supplemental S1A and S1B Fig). Right-hand graph: We recorded pharyngeal pumping of individual wild-type 5 day old animals raised from the egg stage on control media or media containing 4% glucose; 30 second records. 4% glucose significantly impairs pumping rate (**P = 0.0008, unpaired t test, n = 35 animals per condition, error bars represent SEM; averages from three trials per condition). (B) C. elegans glycolytic and gluconeogenic pathways. Although most enzymes of glycolysis catalyze both forward (green, glycolysis) and reverse (red, gluconeogenesis) reactions, PFK/FBP and PYK/PCK are enzyme pairs that catalyze one-way, opposing reactions: glycolytic enzymes PFK (phosphofructokinase) and PYK (pyruvate kinase) and gluconeogenic enzymes FBP (fructose-1,6-bisphosphatase) and PCK (phosphoenolpyruvate carboxykinase). Glu-6-P = glucose-6-phosphate; Fru-6-P = fructose-6-phosphate; Fru-1,6-BP = fructose-1,6-bisphosphate; DHAP = dihydropyruvate phosphate; G3P = glyceraldehyde-3-phosphate; PEP = phosphoenolpyruvate. * Gene disruptions studied most extensively in this work (C. elegans ORF designations indicated). We note that, in our hands, disruption of C50F4.2 (pfk-1.2), F25H5.3 (pyk-1), and W05G11.6 (pkc-1) did not impact lifespan. Throughout the paper, data on genes that normally promote glycolysis are represented in green; data on genes that normally promote gluconeogenesis are presented in red. (C) Swimming rate profiles for WT animals treated with glycolysis-limiting pfk(RNAi) and pyk-2(RNAi) or gluconeogenesis-limiting fbp-1(RNAi) and pck-2(RNAi). age-1(RNAi) is used as a positive control known to extend old age locomotory ability. Animals treated with glycolysis-limiting pfk(RNAi) and pyk-2(RNAi) swim significantly better than controls on days 10 and 15 of life; (pfk(RNAi) confers a 42% increase in swimming ability on day 10 and a 59% increase on day 15; pyk-2(RNAi) confers a 63% increase on day 10 and an 83% increase on day 15). It may be worth noting that these increases we recorded in this experiment are higher than those seen for the age-1 positive control RNAi (for age-1(RNAi), a 26% and a 46% increase in swimming ability on days 10 and 15, respectively, in this study) so locomotory impact appears relatively substantial, assuming age-1(RNAi) efficacy. On day 15, fbp-1(RNAi) decreases swimming rates by 44%, and pck-2(RNAi) animals moved too slowly to measure (or were dead) (marked as N/A). Data shown are pooled from two similar independent trials, n = 40 animals per condition per trial. Error bars represent SEM. ***P < 0.0001, ****P < 0.0001; n.s. = not significant; all statistical analyses by one-way ANOVA. (D) Survival curves of WT animals treated with RNAi targeting glycolytic genes pfk-1(RNAi) and pyk-2(RNAi). Log-rank test), maximal lifespan is not significantly affected (Supplemental S1D Fig). Conversely, animals treated with RNAi directed against gluconeogenic genes fbp-1 and pck-2 have lowered median survival (a 10.94% decrease for fbp-1 and 21.88% decrease for pck-2 disruptions, Log-rank test), and exhibit a 10.94% decrease for fbp-1 disruptions and a 21.88% decrease for pck-2 disruptions) and exhibit lower survival relative to controls (P < 0.0001 for both pfk and pyk-2 disruptions, Log-rank test). Animals treated with the age-1(RNAi) positive control show a 46.88% increase in median survival and significantly increased survival (P < 0.0001, Log-rank; refer to S1 Table for survival data with age-1(RNAi)). Data shown are pooled from 6 independent trials, n = 60 animals per condition per trial.

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in human elderly [11, 12, 53] and consistent with recent studies of glucose toxicity on old-age mobility [13, 20].

Glucose catabolism is associated with accelerated aging traits

Aging induces glycolytic gene expression in mammals [30, 32], and, in C. elegans under standard feeding conditions, inhibiting glucose catabolism with the glucose analog 2-deoxyglucose or with genetic inhibition of at least one glycolytic enzyme (pyruvate kinase, or PYK) promotes longevity [19, 52]. In addition, treatment with the glycolysis intermediate DHAP has a large detrimental impact on C. elegans lifespan [52]. These observations suggest that the process, or consequences, of glucose utilization could be linked to age-associated decline in locomotion.

To address how glucose catabolism influences functional aging under standard feeding conditions, we disrupted the specific C. elegans enzymes that catalyze unidirectional, irreversible steps in glycolysis. Although most reactions of glycolysis are readily reversible, distinct enzymes execute the forward and reverse reactions for the fructose-6-phosphate/fructose-1,6 bisphosphate interconversion (PFK/FBP) and the P-enolphosphate/pyruvate interconversion (PYK/PCK) (see Fig 1B). We anticipated that RNAI-mediated disruption of the reactions that specifically promote glycolysis (i.e., enzymes PFK and PYK) should have a predominant metabolic effect of limiting glycolytic flux, as appears to be the case for 2-deoxyglucose [19, 29]. Indeed, we found that knockdown of the glycolysis-promoting phosphofructokinase (pfk-1.1, hereafter referred to as pfk) and pyruvate kinase (pyk-2) genes results in significant increases in swim ability in mid- and late life (Fig 1C). We conclude that limiting glycolysis can robustly protect against age-associated mobility decline, consistent with a previous report [20]. Furthermore, we find that disruptions of glycolytic genes pfk and pyk-2 also result in increases in median survival (Fig 1D and S1Table), supporting that limiting glycolysis promotes overall health in aging adult populations.
It is interesting to note that the gains in swimming ability and median survival we observe with unidirectional glycolysis gene disruptions are not accompanied by comparably large increases in maximal lifespan (Supplemental S1D Fig). This observation is important in light of the study of Bansal et al. [54] that suggests that some long-lived C. elegans mutants spend extended periods of time in a frail state at the end of life, in proportion to their increased maximal lifespan. The glycolysis interventions we highlight appear to maintain youthful physiology in mid-life without extending late-life frailty. The inability of glycolysis interventions to extend maximal lifespan may indicate midlife-specific metabolic consequences, or alternatively might be attributed to confounding failures in older animals not improved by glycolytic inhibition. Blocking glycolysis may result in the build-up of glucose, which could reach toxic levels later in life. In sum, knockdown of enzymes that specifically catalyze one-way steps critical for glucose breakdown can improve multiple measures of healthy aging, supporting that glycolytic flux confers an overall negative impact on physiology that promotes functional aging.

Inhibiting unidirectional glycolysis-promoting reactions increases mid-life health via DAF-16

DAF-16/FOXO is a critical downstream transcription factor in longevity-promoting pathways [22], and inhibition of DAF-16 activity has been previously implicated in the glucose toxicity mechanism [16, 19]. We therefore tested whether DAF-16/FOXO might act to promote healthy aging when glycolysis is genetically impaired. We first sought evidence of DAF-16 activation under conditions of pfk and pyk-2 RNAi using a well-characterized reporter of DAF-16 activity, SOD-3::GFP, a direct transcriptional target of C. elegans DAF-16 [55, 56]. We found that both pfk(RNAi) and pyk-2(RNAi) result in significant increases in SOD-3::GFP levels (Fig 2A), suggesting that glycolytic gene disruptions may generally increase DAF-16 transcriptional activity. Indeed, daf-16 is critical for the strong healthspan effects of glycolytic gene disruptions we document in Fig 1: the large increases in swimming ability and the median survival extension of wild-type animals treated with RNAi against glycolytic genes pyk-2 and pfk are largely eliminated in daf-16 null mutants (Fig 2B and 2C and S1 Table, overall survival analyses; we note that, although median survival does not increase with pfk(RNAi) in the daf-16 mutant background, the daf-16 lifespan curve is significantly right-shifted late in life with pfk(RNAi) treatment, suggesting that glycolytic interventions may impact longevity independently of DAF-16 in older animals).

If DAF-16 were activated by knockdown of glycolytic pfk and pyk-2 via an insulin signaling pathway, we might expect that pfk and pyk-2 disruption in an insulin pathway activation background would not further enhance health and lifespan phenotypes. We tested this possibility by performing pfk and pyk-2 RNAi in the age-1 insulin signaling pathway mutant, well characterized for activated DAF-16 and associated longevity [26]. We found that disrupting glycolytic pfk or pyk-2, which extend locomotory healthspan and mid-life survival in wild type (Fig 1C and 1D and S1 Table), does not enhance locomotory ability in the age-1 mutant background (S2A Fig) and pfk(RNAi) does not significantly improve age-1 lifespan curves (S1 Table). The lack of benefits of pfk and pyk-2 knockdown in age-1 is consistent with a model in which pfk and pyk-2 disruption can activate DAF-16, possibly via a common insulin signaling pathway: enhanced DAF-16 activity can bypass deleterious consequences of glycolysis. We note that pyk-2(RNAi) can extend age-1 lifespan (S1 Table), suggesting that the glycolytic pathway may also influence lifespan via pathways other than the insulin pathway and DAF-16 (as supported by our results with pfk(RNAi) in the daf-16 background, described above).

Together these data on daf-16 demonstrate that inhibiting unidirectional glycolytic enzymes increases locomotory healthspan and median lifespan via a DAF-16-dependent...
Fig 2. DAF-16/FOXO is Required for Locomotory Healthspan and Lifespan Benefits of Genetic Glycolysis Disruption. (A) Fluorescence intensity levels of wild-type animals expressing a GFP transcriptional reporter for sod-3, which expresses superoxide dismutase and is a direct transcriptional target of DAF-16. We measured in vivo fluorescence using a spectrophotometer on day 5 of life. As expected from previous work [74], decreased insulin signaling with age-1(RNAi) results in increased SOD-3::GFP expression. Similarly, knockdown of glycolytic genes via pfk(RNAi) and pyk-2(RNAi) significantly increases SOD-3::GFP fluorescence levels. Disrupting the fbp-1 and pck-2 gluconeogenic genes, on the other hand, does not significantly change GFP fluorescence levels. Data are pooled...
Glycolysis and gluconeogenesis differentially impact *C. elegans* healthspan

Average from 5 independent trials, n = 50 animals per condition per trial. Error bars represent SEM. *** P < 0.001; **** P < 0.0001; n.s. = not significant; all by one way ANOVA with Dunnett’s multiple comparisons test. (B) Swimming rates of *daf-16*(mgDf50) mutants treated with glycolytic and gluconeogenic RNAi. Disrupting glycolysis-promoting genes *pfk* or *pyk-2* in the *daf-16* null background eliminates old-age benefits that occur with *pfk*(RNAi) or *pyk-2*(RNAi) (refer to Fig 1C), indicating a requirement for *daf-16* in the benefits of glycolysis impairment. In contrast, disruption of gluconeogenic genes *fbp-1* or *pck-2* results in significant decreases in swimming rates in *daf-16* animals, similar to wild-type. Data shown are pooled from two independent trials, n = 40 animals per condition per trial. Measurements were taken on day 10 of life. Error bars represent SEM. *** P < 0.001; **** P < 0.0001; n.s. = not significant; all by one way ANOVA with Dunnett’s multiple comparisons test. *daf-16* mutants are documented to have lowered locomotory ability vs. WT later in life [91, 92]; we note that, although comparison of scores across experiments is not possible due to variation in baseline values, the swimming rate of *daf-16*(mgDf50) control animals was slightly lower than WT controls on the same day of life (in Fig 1C; mean of 46.17 head thrashes/30 sec. for *daf-16* vs. 52.17 head thrashes/30 sec. for WT on day 10). (C) Survival curves of *daf-16*(mgDf50) null mutants treated with RNAi against glycolytic genes *pfk* and *pyk-2*. The beneficial effects of glycolytic gene disruptions seen in wild-type animals (refer to Fig 1D) are mostly eliminated in the *daf-16* null background: median survival is not increased in *daf-16* animals treated with either RNAi, and the survival curve of *daf-16* mutants raised on *pyk-2*(RNAi) is not significantly (n.s.) different than controls (Log-rank test). *pfk*(RNAi) slightly increases the survival of *daf-16* mutants later in life (as indicated by a right-shifted survival curve in late-life; Log rank calculated *P* = 0.0128), suggesting possible *daf-16*-independent mechanisms for lifespan extension in older animals with glycolytic inhibitions. Data are pooled from 5 independent trials, n = 60 animals per condition per trial, details in S1 Table.

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Mechanism. Data also support that modulation of insulin signaling is associated with glycolytic flux changes, consistent with previous studies on *C. elegans* glucose toxicity [13, 16–20].

**Reversing glycolytic flux (i.e., promoting gluconeogenesis) promotes healthy aging**

The existence of enzymes that specifically catalyze the reverse reactions of PFK and PYK-2 enabled us to test whether disruption of gluconeogenesis promotes effects opposite to the positive outcomes of RNAi-mediated glycolysis inhibition. We found that animals treated with RNAi against genes that encode the unidirectional gluconeogenic enzymes fructose-1,6-bisphosphatase (FBP-1), which catalyzes the gluconeogenic reaction reverse to PFK, and phosphoenolpyruvate carboxykinase (PCK-2), which catalyzes the gluconeogenic reaction reverse to PYK-2 (Fig 1B), exhibit major impairments in locomotory ability in late life (disrupting *pck-2* knockdown has such strong effects that animals were moving too slowly (or too few animals remained alive) to measure on day 15) (Fig 1C). Disrupting gluconeogenic genes *fbp-1* and *pck-2* also significantly shortens median lifespan (Fig 1D and S1 Table).

To rule out that *pck-2*(RNAi) might merely confer general sickness, we tested *pck-2* inhibition in the *age-1* background. We find that *pck-2*(RNAi) does not reduce locomotory rates or lifespan extension in the *age-1* background (S2A and S2B Fig and S1 Table), supporting that impairing gluconeogenic activity does not universally compromise health. We conclude that *fbp-1* and *pck-2* dependent reactions that promote gluconeogenesis are needed to maintain normal adult healthspan. Our findings indicate that gluconeogenesis pathways are critical for late life health under standard growth conditions, and constitute the first focused demonstration that *fbp-1* and *pck-2* deficiency can be progeric for *C. elegans*.

In sum, inhibiting the enzymes that specifically promote glycolysis confers beneficial effects on both locomotory healthspan and median survival, whereas inhibiting enzymes that specifically promote gluconeogenesis negatively impacts these two indicators of adult health. Data suggest that glycolysis is deleterious, and gluconeogenesis is beneficial for healthy aging, especially as judged by locomotory capacity.
Overexpression of gluconeogenic PEPCK-C gene \( pck-2 \) extends locomotory healthspan and lifespan

Phosphoenolpyruvate carboxykinase (PEPCK) catalyzes the rate-controlling step of gluconeogenesis, and is thus a central player in glucose homeostasis. In mammals, there are two forms of the enzyme: cytosolic PEPCK-C and mitochondrial PEPCK-M, with PEPCK-C thought to play the larger role in metabolic regulation [57]. Given the striking requirement for gluconeogenic \( pck-2 \) in maintained locomotory health and normal lifespan (Fig 1C and 1D and S1 Table) and previous indications that PEPCK can drive metabolic states [58, 59], we focused on mechanisms of \( pck-2 \) contributions to adult health (\( pck-2 \) does not include a detectable mitochondrial targeting sequence and PCK-2 expressed from a rescuing transgene does not localize to mitochondria (S3A Fig), supporting \( pck-2 \) encodes a PEPCK-C; in our hands, sequence-confirmed RNAi knockdown of \( pck-1 \), which encodes another \( C.\ elegans \) PEPCK ortholog [59] did not impact aging, S1 Table).

Since disruption of \( pck-2 \) is deleterious to health, we first asked whether elevated \( C.\ elegans \) PEPCK expression might suffice to promote healthy association with enhanced late adult maintenance. We examined a strain carrying multiple integrated copies of \( P_{pck-2}::gfp \) for potential benefits of \( pck-2 \) over-expression in healthy aging. We found that the Is\( [P_{pck-2}::gfp] \) over-expression line exhibits increases in midlife locomotory ability and median survival (but not maximal lifespan) as compared to controls that express only GFP from the \( pck-2 \) promoter (S3B and S3C Fig and S1 Table; the promoter-only Is\( [P_{gfp}] \) control construct had no effect on wild-type locomotory ability or lifespan, S3C, S3D and S3E Fig and S1 Table). We conclude that over-expression of \( pck-2 \), which encodes the unidirectional, rate-limiting enzyme of gluconeogenesis, can extend locomotory healthspan and median lifespan.

\( pck-2 \) overexpression lifespan extension requires gluconeogenesis

In mammals, PEPCK is widely expressed, and, in addition to its role in gluconeogenic tissues, PEPCK plays an anaplerotic role in non-gluconeogenic cells (i.e., in glyceroneogenesis; [57]). To ask whether \( pck-2 \) overexpression acts via its gluconeogenesis role, we tested the requirement for the glucose-6-phosphatase complex in the improved mid-life survival seen in our \( pck-2 \) overexpressing strain. In mammals, glucose-6-phosphatase is specifically expressed in gluconeogenic tissues where it catalyzes the final step of gluconeogenesis [60]. Here we targeted the ortholog of glucose-6-phosphate translocase, a component of the glucose-6-phosphatase complex in vertebrates ([61]; see Supplemental S4 Fig for homology alignment; note that the \( C.\ elegans \) glucose-6-phosphatase has not been experimentally identified). We found that the lifespan benefits of \( pck-2 \) overexpression are absent when glucose-6-phosphate translocase is disrupted (Fig 3A and S1 Table). Thus \( pck-2 \) overexpression acts via a gluconeogenesis pathway to improve adult health.

\( pck-2 \) is expressed in specific intestinal cells via a \( daf-16 \)-dependent mechanism

To better understand how gluconeogenic PCK-2 improves adult health, we probed the expression pattern of \( pck-2 \). We used the Is\( [P_{pck-2} pck-2::gfp] \) reporter to determine the tissues/cells in which PCK-2 is likely needed for adult health impact. In younger animals under standard growth conditions, the Is\( [P_{pck-2} pck-2::gfp] \) translational fusion is expressed in the pharynx, in the intestine, in bodywall muscle, and in hypodermis. In adults, Is\( [P_{pck-2} pck-2::gfp] \) expression becomes strikingly restricted to the very most anterior and posterior intestinal cells (Fig 3B).
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A

Fraction survival

0.0 0.2 0.4 0.6 0.8 1.0

Days

Pck-2gfp + EV RNAi
pck-2 OE + EV RNAi
Pck-2gfp + G6PTase(RNAi)
pck-2 OE + G6PTase(RNAi)

****

B

ls[Pck-2pck-2::gfp] + empty vector control

ls[Pck-2pck-2::gfp] + daf-16(RNAi)

Pck-2 Predicted DAF-16 BS

Pck-2pck-2::gfp
5’-tttcccacttttcattataat
cttatcaatatttcgatttt-3’

Ex[Pck-2mutpck-2::gfp] + empty vector control

C

No. head thresses / 30 sec

0 50 100 150

Pck-2gfp + EV RNAi
pck-2 OE + EV RNAi
Pck-2gfp + daf-16(RNAi)
pck-2 OE + daf-16(RNAi)

**

D

Fraction survival

0.0 0.2 0.4 0.6 0.8 1.0

Days

Pck-2gfp + EV RNAi
pck-2 OE + EV RNAi
Pck-2gfp + daf-16(RNAi)
pck-2 OE + daf-16(RNAi)

****
is required for DAF-16 expression healthspan effects. Indeed, the pck-2 over-expressor strain and treated with empty vector control (EV RNAi) or F47B8.10(RNAi).

We find that disrupt the DAF-16 binding site in the pck-2 promoter is shown on the bottom left in red; green highlighting indicates altered base pairs in the binding site mutant. Disruption of the DAF-16 binding site 1086 base pairs upstream of the intestinal GFP pattern on day 7) as well as in the pharynx (arrowhead; all animals displayed pharyngeal GFP on day 7).

daf-16(RNAi) treatment specifically eliminates the intestinal GFP signal (top right-hand panels; no intestinal GFP was detected in any of the 60 observed animals on day 7). Bar, 200 μm. (C) Swimming rates of wild-type animals expressing pck-2::gfp signal (bottom right; no intestinal GFP was detected in any of the 60 observed animals on day 7). Bar, 200 μm. (D) Survival curves of wild-type animals expressing a transcriptional reporter for pck-2 (P\textsubscript{pck-2}::gfp, used as the control strain) or the intact pck-2 gene expressed from its own promoter (P\textsubscript{pck-2}::pck-2::gfp, here indicated as “pck-2 OE”, used as the pck-2 over-expressor strain) treated with an empty vector control RNAi (EV RNAi) or daf-16(RNAi). pck-2 overexpression (OE) significantly increases swimming rates vs. wild-type controls (P < 0.01, one way ANOVA). Swimming fitness is disrupted with daf-16(RNAi) in both WT controls and pck-2 OE animals (P < 0.01, one way ANOVA). Swimming rates of control and pck-2 OE animals treated with daf-16(RNAi) are not significantly different from one another, one way ANOVA. Data shown are from averages from three trials, n = 40 animals per condition. Measurements were taken on day 9 of life. Error bars represent SEM. ** P < 0.01. All one way ANOVA analyses were performed with Dunnett’s multiple comparisons test. (D) Survival curves of wild-type animals expressing a transcriptional reporter for pck-2 (P\textsubscript{pck-2}::gfp, used as the control strain) or the intact pck-2 gene expressed from its own promoter (P\textsubscript{pck-2}::pck-2::gfp, “pck-2 OE”, used as the pck-2 over-expressor strain) treated with an empty vector control RNAi (EV RNAi) or daf-16(RNAi). pck-2 overexpression (OE) significantly increases survival (P = 0.0017, Log-rank test). The pck-2 OE survival benefit is abolished with daf-16(RNAi) (control and pck-2 OE animals treated with daf-16(RNAi) have shortened lifespans that are not significantly (n.s.) different from one another, Log-rank test). Data shown are pooled from 2 independent trials, n = 60 animals per condition per trial. **** P < 0.0001. Details in S1 Table.

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We also note that, consistent with its assignment as a PEPCCK-C ortholog, PCK-2::GFP appears localized to the cytoplasm in the cells in which it is expressed (S3A Fig).

Since we had determined that daf-16 is important for the benefits associated with glycolysis downregulation (Fig 2), we wondered whether daf-16 might be important for the pck-2 overexpression healthspan effects. Indeed, the pck-2 promoter includes a consensus DAF-16 binding site, and has been identified as a direct target of DAF-16 [62, 63]. To test whether DAF-16 is required for pck-2 expression, we performed daf-16(RNAi) in the Is[P\textsubscript{pck-2} pck-2::gfp] strain. We find that daf-16(RNAi) eliminates the GFP signal specifically in adult intestinal cells but not in other tissues in younger animals (Fig 3B), establishing that the intestinal expression of pck-2 is DAF-16-dependent. To confirm direct targeting of pck-2 by DAF-16, we altered the putative DAF-16 binding site in the pck-2 promoter (Fig 3B) and examined Ex[P\textsubscript{pck-2 mut} pck-2::gfp] expression. We find that disruption of the DAF-16 consensus binding site eliminates intestinal expression, supporting that DAF-16 directly regulates pck-2 expression in specific intestinal cells via consensus DAF-16 target sites (Fig 3B). Furthermore, the locomotory and median lifespan benefits associated with pck-2 over-expression are completely abolished by daf-16(RNAi): the mid-age health phenotypes of pck-2 over-expressing animals under daf-16 RNAi knockdown conditions are similar to that of control animals treated with daf-16(RNAi) (Fig 3C, D).

Although DAF-16 is required for increased lifespan with both reduced insulin signaling [21–28] and pck-2 overexpression, pck-2 is not required for the long lifespan of age-1 insulin pathway mutants (S2B Fig), suggesting that gluconeogenic activity may lie upstream of or parallel to the insulin pathway to impact lifespan and healthspan.
Overall, our results support that DAF-16 positively regulates pck-2, which is required to extend *C. elegans* healthspan, and suggest that specific cells in the anterior and posterior intestine are central to this regulation. DAF-16 thus plays a critical role in promoting gluconeogenesis and healthy aging at least in part by transcriptional regulation of key biosynthetic enzyme cytoplasmic PCK-2.

**Expression of transcriptional fusion** $P_{pck-2}\text{-gfp}$ **is induced under food limitation via a daf-16-dependent mechanism**

For our studies of pck-2 expression we also constructed a pck-2 transcriptional reporter (Is[$P_{pck-2}\text{-gfp}$]) in which the pck-2 promoter is fused to GFP and all pck-2 coding and intron sequences are missing. We noted a striking difference in GFP expression between the functional PCK-2::GFP translational fusion and the $P_{pck-2}$ transcriptional reporter: while the PCK-2::GFP translational fusion is constitutively expressed, the pck-2 promoter-only transcriptional reporter is expressed solely in the intestine and only under food limitation conditions (Fig 4A and 4B). We tested multiple food limitation regimens published to induce dietary restriction-like metabolism (absence of food, food dilution, metformin administration, and the eat-2 feeding-impaired mutant) to document that animals carrying the $P_{pck-2}\text{-gfp}$ transcriptional reporter exhibit strong GFP fluorescence in the anterior and posterior intestine cells (Fig 4A and 4B) (similar to the constitutive Is[$P_{pck-2}\text{-pck-2}\text{-gfp}$] intestinal cell expression pattern (Fig 3B); expression in other cells that express $P_{pck-2}\text{-gfp}$ is not evident for Is[$P_{pck-2}\text{-gfp}$]). Strikingly, however, under *ad lib* conditions little, if any, $P_{pck-2}\text{-gfp}$ expression is apparent in anterior or posterior intestinal cells. These data raise the possibility of enhanced pck-2 expression under food limitation conditions. Increased intestinal GFP produced from the $P_{pck-2}\text{-gfp}$ transcriptional reporter in the DR-mimetic eat-2 mutant requires daf-16 (Fig 4B). Thus, food limitation is associated with daf-16-dependent transcriptional expression of Is[$P_{pck-2}\text{-gfp}$] in specific intestinal cells.

We confirmed an increase in native pck-2 transcripts in the eat-2 DR-mimetic mutant relative to WT later in life by qPCR analysis (Fig 4C). While pck-2 expression decreases in wild-type animals as they age (0.44 and 0.39 fold change as compared to the WT day 3 level for days 7 and 10, respectively), pck-2 levels remain remarkably level throughout the eat-2 lifespan (0.72, 0.70 and 0.80 fold change as compared to WT day 3 level for day 3, 7, and 10, respectively) (Fig 4C). Notably, pck-2 transcript levels under DR are elevated in mid-to-late life when compared to *ad lib* fed animals, just as we see with the $P_{pck-2}\text{-gfp}$ reporter (Fig 4B). Levels of the gluconeogenic gene encoding the *C. elegans* glucose-6-phosphate translocase ortholog are also increased under eat-2 DR (Fig 4C), suggesting that DR metabolism may generally feature elevated transcripts of the enzymes of gluconeogenesis, with consequent increased gluconeogenic capacity.

The difference in expression profiles between full length translational fusion (constitutive gut expression) and the transcriptional fusion (food limitation-induced gut expression) might be attributed to different sequence inclusions in the transcription and translational constructs, although an alternative possibility can be considered, grounded in the fact that the translational fusion constitutes a PEPCK over-expression situation. Because in WT, qPCR shows that pck-2 transcript levels decline with age, but in the eat-2 DR model pck-2 transcript levels remain elevated (Fig 4C), it is tempting to speculate that the elevation of PCK-2 protein associated with the over-expressed functional translational fusion might be the consequence of increased PCK-2 activity that acts in a positive feedback loop to maintain a gluconeogenic metabolic state (see Discussion).

**Gluconeogenic PCK-2 is required for the health benefits of DR**

Our observations raised the question as to whether PCK-2, induced under DR, is required for the healthspan benefits associated with the DR state. To address this question, we measured
Fig 4. Dietary Restriction Induces Gluconeogenic Gene pck-2 Expression via DAF-16, and pck-2 Expression is Needed for DR-associated Healthspan Benefits. (A) Representative wild-type animals expressing a GFP transcriptional reporter for pck-2. (B) % Animals with GFP Signal. (C) pck-2 and GAPDh expression fold change. (D) No. head thresses/30 sec. (E) Survival curves for eat-2(ad1116) with or without pck-2 RNAi. (F) Survival curves for WT with or without pck-2 RNAi. (G) Excitation at peak fluorescence intensity (n.m). (H) Relative change in PCK-2-GFP. (I) Schematic diagram illustrating the role of DAF-16 in regulating glucose and fatty acid metabolism in C. elegans.
promoter GFP fusion, lacking pck-2 coding sequences) under ad libitum (AL) or dietary restricted (DR) conditions, day 4 of life. DR animals were raised on diluted (7 × 10^9 cfu/ml) OP50; AL is undiluted (7 × 10^10 cfu/ml) bacteria. A = anterior, P = posterior, intestinal GFP fluorescence indicated by arrows. Quantitation of GFP observations is described in (B). Bar, 200 μm. (B) pck-2 transcriptional reporter GFP expression under three distinct food limitation conditions. Left: as in (A), wild-type animals expressing the P_pck-2::gfp reporter were raised on plates seeded with undiluted (7 × 10^10 cfu/ml) or diluted (7 × 10^6 cfu/ml) bacteria and the number of animals displaying GFP fluorescence was scored on day 4 of life. Ampicillin was added to plates to inhibit bacterial growth. The number of animals with GFP fluorescence increases with bacterial dilution (P < 0.009, unpaired t test). Data are pooled averages from 3 independent trials, n = 10 animals per condition per trial. Error bar represents SEM. * P < 0.05. Center: metformin is a drug used to treat type-2 diabetes that induces features of dietary restriction-like metabolism in C. elegans [41]. Animals expressing P_pck-2::gfp were raised from the egg stage on plates containing either no metformin or 50 mM metformin, and GFP expression was observed on day 4 of life. The number of animals with GFP expression increases on 50 mM metformin plates (P = 0.04, unpaired t test). Data are pooled averages from 3 independent trials, n = 60 animals per condition per trial. Error bar represents SEM. * P < 0.05. Right: GFP fluorescence in eat-2(ad1116) mutants expressing P_pck-2::gfp and treated with either an empty vector control RNAi or daf-16(RNAi). Under DR, daf-16 (RNAi) reduces the frequency of observing GFP expression on all tested days (data shown are from day 8 of life, P = 0.05, unpaired t test). Wild-type animals carrying injectable P_pck-2::gfp showed no GFP signal on day 8. Data are pooled averages from 3 independent trials, n = 60 animals per condition per trial. Error bar represents SEM. * P < 0.05. (C) Left-hand graph: qRT/PCR-evaluated G6P translocase (G6PTase). Data are averages from 3 independent trials, n = 10 animals per condition per trial. Error bar represents SEM. ** P < 0.001; *** P < 0.0001. All one way ANOVA analyses performed with Dunnett’s multiple comparisons test. (E) Survival curves of long-lived eat-2(ad1116) mutants treated with RNAi against pck-2. Pck-2 RNAi results in small but significant increases in swimming rates early in life (day 5; P < 0.01, one way ANOVA, a 12% increase) and no later change; glycolytic gene pck-2(RNAi) did not significantly affect swimming ability on any tested day. Disrupting glycolytic gene pck-2 significantly reduces swimming rates on day 15 (P < 0.0001 by one way ANOVA, a 58% reduction). Data are pooled from 3 independent trials, n = 40 animals per condition per trial. Error bars represent SEM. * P < 0.001; *** P < 0.0001. All one way ANOVA analyses performed with Dunnett’s multiple comparisons test. (F) Survival curves of wild-type animals raised under ad libitum (AL) or dietary restriction (DR) conditions, with and without pck-2(RNAi). Median survival of animals expressing the empty vector control was higher under DR conditions as compared to well-fed controls (an 18.62% increase, see S1 Table for details) (survival differences P = 0.0039, Log-rank test). Disrupting glycolytic gene pck-2 abolished these beneficial DR effects: animals raised on diluted gluconeogenic RNAi bacteria had survival curves that were not significantly different from those of well-fed animals (n.s., Log-rank test). Data are pooled from 3 independent trials, n = 60 animals per condition per trial. Error bars represent SEM. * P < 0.001; n.s. = not significant. (G) Excitation wavelengths corresponding to peak age pigment fluorescence intensities in wild-type and eat-2(ad1116) dietary restricted mutants raised on empty vector control RNAi, and in wild-type animals treated with RNAi against glycolytic gene pfk. Dietary restriction lowers age pigment peak excitation wavelengths, and this shift to lower excitation wavelengths (the ExMax shift) can be used as a biomarker for DR [64]. Like eat-2 mutants, pfk(RNAi) animals show ExMax shifts to lower excitation wavelengths on day 5 of life (the average ExMax value for wild-type animals with pfk disruptions is 334.0 nm vs. 344.0 nm for wild-type controls; ExMax values for eat-2(ad1116) animals is 330.0 nm; P < 0.0001 for eat-2(ad1116) + vector control and P < 0.001 for WT + pfk(RNAi), one way ANOVA). Data are averages from 2 independent trials, n = 30 animals per condition per trial. Error bars represent SEM. * P < 0.001; *** P < 0.0001. All one way ANOVA analyses performed with Dunnett’s multiple comparisons test. (H) Wild-type animals expressing P_pck-2::gfp and treated with a vector control or with RNAi against glycolytic gene pfk on day 7 of life. Pfk(RNAi) significantly increases GFP fluorescence in P_pck-2::gfp-expressing animals, as quantitated in a spectrophotometer (P = 0.0383, unpaired t test). Data are averages from 3 independent trials, n = 30 animals per condition per trial. Error bar represents SEM. * P < 0.05. (I) Summary model for interactions of glycolysis, gluconeogenesis, and healthy aging. Under ad libitum (AL) conditions, insulin-like signaling promotes glucose transport and processing via glycolysis. Glycolytic activity, in turn, suppresses the healthspan-promoting transcription factor DAF-16 to limit healthy aging. Under dietary restriction (DR) or RNAi disruption of pfk or pck-2, glucose levels and glycolytic activity decrease, while DAF-16 is activated to increase transcription of pro-healthspan targets including gluconeogenic gene pck-2. Elevated gluconeogenic activity (dependent on DAF-16-promoted PCK-2 expression) acts in a feed-forward capacity to further enhance the gluconeogenic pathway and may inhibit glycolysis/insulin signaling to promote/maintain DAF-16 signaling and healthy metabolism (inhibition represented by a dashed line).
DAF-16 positively regulates pck-2 expression under DR or glucose limitation conditions, in a metabolic shift critical for maintaining healthspan. We find that RNAi-mediated knockdown of fbp-1, the other unidirectional enzyme that promotes gluconeogenesis, also eliminates longevity phenotypes of eat-2 and of WT reared under food limitation (Supplemental S5A and S5B Fig). Together, our data support that gluconeogenic flux may be a critical element in maintaining healthy aging in older animals (Fig 1C and 1D) and in healthy DR metabolism.

**Glycolytic disruptions extend healthspan by promoting gluconeogenic activity and DR-like metabolism**

Limiting glycolysis promotes DAF-16 activity (Fig 2A), and DAF-16 is needed for healthspan benefits of glycolysis inhibition (Fig 2B and 2C), gluconeogenic gene pck-2 overexpression (Fig 3C and 3D), and some forms of dietary restriction [34]. Our findings raise an intriguing question on metabolic flux: do glycolytic disruptions trigger gluconeogenic / DR metabolism to promote healthy aging?

To address this possibility, we first asked whether the locomotory healthspan effects of glycolytic gene disruption might act in the same pathway as locomotory healthspan effects of feeding limitation eat-2 DR by addressing whether or not an additional (possibly additive) improvement occurs when both perturbations are operative. We find that glycolytic disruption in the eat-2 background does not increase older age swimming above that recorded for eat-2 (Fig 4D), consistent with a model in which glycolytic gene disruptions and DR increase locomotory healthspan via the same pathway (although we cannot be certain that ceiling effects might exist in this particular situation).

We next asked whether glycolytic disruption induces additional features of dietary restriction. We previously showed that under multiple DR conditions, the excitation wavelength corresponding to the peak fluorescence of age pigments/lipofuscin shifts to a wavelength lower than any other longevity pathway mutants or any animal grown in abundant food [64]. Indeed, we found that for glycolytic gene disruption pfk(RNAi), the peak age pigment fluorescence excitation wavelength shifts downward, similar to that found in dietary-restricted eat-2 mutants (Fig 4G) and in other food limitation conditions, suggesting that pfk(RNAi) induces a switch to DR-like metabolism. Interestingly, DR prevents the upregulation of glycolytic PFK with age in mammals [30] and inhibits pfk expression in *C. elegans* [65], supporting that decreased glycolysis-promoting phosphofructokinase activity may be a key and conserved feature of DR metabolism.

Finally, we found that glycolysis disruptions via pfk(RNAi) significantly increase gluconeogenic gene pck-2 expression, as measured by increased PCK-2::GFP (expressed from Is[P_pck-2::gfp]) levels (Fig 4H). Thus, genetically limiting glycolysis flux increases gluconeogenic enzyme expression, likely via daf-16-mediated loops that direct a yin/yang metabolic balance between glucose utilization and biosynthesis. Our data support a model in which enhanced gluconeogenic metabolism is key to health benefits under dietary restriction and is needed for normal healthy aging, a new concept in healthy metabolism.

**Discussion**

We examined the relationship between *C. elegans* glucose metabolism pathways and healthy aging, with an emphasis on the maintenance of physical function and overall mid-life survival health outcomes. A high glucose diet accelerates age-associated decline in mobility and survival robustness. Glycolysis exerts a particularly strong negative influence on healthy aging: limiting glycolysis by disrupting one-way glucose degradation steps can result in large old-age locomotory improvements and increases in median lifespan. A key point in our study is
focused on the opposing gluconeogenic pathway: gluconeogenic activity is essential for normal healthy aging, as disrupting dedicated gluconeogenic genes induces progeric decline of locomotory activity and compromises survival. Gluconeogenic pathway integrity is also needed for the beneficial effects of dietary restriction. Importantly, interventions that disrupt glycolysis or increase gluconeogenic activity promote healthy physiology in mid-life without markedly extending maximal lifespan, suggesting that these metabolic interventions improve the quality of aging without extending the period of late-life frailty. Data also indicate that expression of the gluconeogenic PEPCK PCK-2 in just a few intestinal cells can modulate animal-wide metabolic state. Overall, our data highlight a novel paradigm in which glycolysis and gluconeogenesis work opposite to one another to influence the maintenance of adult health. We speculate that this metabolic yin/yang underlies major differences in the quality of aging, which if conserved, validates warnings against high sugar diets.

High glucose accelerates muscle decline across species

Type II diabetes mellitus is associated with accelerated muscle loss during human aging [10–12] and loss of mouse glycolytic muscle is associated with metabolic dysfunction, a consequence that can further compromise glucose homeostasis [66]. We emphasize that a high glucose diet also accelerates *C. elegans* locomotory decline, likely reflective of muscle dysfunction. Previous reports on mobility parameters are consistent with a negative impact of glucose on adult mobility maintenance [14, 20, 67]. Moreover, high glucose, which can compromise mammalian heart health [68], also promotes functional decline of *C. elegans* cardiac-like pharyngeal muscle. Although the swimming assay we focus on measures functionality rather than muscle integrity per se, our data support *C. elegans* is a plausible model for development of protective strategies that maintain muscle functionality against the stress of high glucose/high glycolysis metabolism associated with high sugar diets or diabetes [69].

Glucose processing negatively impacts healthy aging

Lee et al. show that excessive glucose promotes fatty acid synthesis to shorten lifespan, and suggest that glycolytic and fatty acid synthesis pathway activity may result in the accumulation of toxic intermediate metabolites [52]. Consistent with this suggestion, treatment with glycolytic intermediate DHAP reduces lifespan in glucose-fed nematodes, and inhibition of glycolytic enzyme pyruvate kinase increases lifespan both in high-glucose and control conditions [52]. We show that inhibiting glycolysis increases median lifespan and promotes locomotory ability in older animals. A paradox is that glycolytic interventions also would be anticipated to elevate glucose levels, which can be toxic. That inhibiting glycolysis can be beneficial raises the possibility that, in early- and mid-life, the buildup of toxic metabolites from glucose processing (i.e. DHAP) rather than the overall glucose levels has a significant impact on aging quality. Since maximal lifespan does not appear to be changed by glycolytic inhibition, it is possible that glucose build-up could reach toxic levels in older animals. In future studies it will be of interest to measure glucose levels over adult life and to address whether promoting gluconeogenic activity, like inhibiting glycolysis, can mitigate the harmful effects of excess glucose exposure.

DAF-16/FOXO promotes gluconeogenic metabolism via a mechanism critical for healthy aging

How exactly does glucose metabolism impact aging quality? When glycolysis is active, daf-16/FOXO expression is suppressed, with deleterious consequences for locomotory and whole-animal aging [16]. We document that when glycolysis is inhibited, DAF-16 activity increases, enabling direct positive regulation of key gluconeogenic gene *pck-2*. Without *pck-2* increase,
aging is accelerated. Thus, our data suggest that one mechanism by which DAF-16 promotes healthspan is by engaging gluconeogenic pathways via increasing PCK-2 production. This model is supported by chromatin profiling studies showing that pck-2 is a direct target of DAF-16 [70] and our demonstration that DAF-16 consensus binding sequences are required for in vivo effects of pck-2 expression. Since mammalian DAF-16 ortholog FOXO1 targets PEPCK to regulate hepatic gluconeogenesis [71, 72], our data support that DAF-16/FOXO regulation of pck-2 may be a facet of a conserved mechanism that controls glucose levels.

**Different roles of PEPCK isoforms as drivers of healthy maintenance: PCK-2 in intestine as a key metabolic regulatory point for healthy aging**

The overall impact of gluconeogenic pathway flux in *C. elegans* is likely carried out by two distinct PEPCKs, with critical points of action in different tissues. There are three PEPCK genes encoded by the *C. elegans* genome. *pck-1* encodes a cytoplasmic PEPCK-C that contributes 85% of the measurable PEPCK activity in young *C. elegans* (*pck-2* appears to contribute the rest of measurable PEPCK activity at this time) [59]. PCK-1 is expressed in bodywall muscle, pharyngeal muscle and intestine, and has been suggested to be the workhorse energy/metabolic modulator PEPCK in *C. elegans* [59, 73]. Overexpression of *pck-1* from its native promoter [59] or only in bodywall muscle is sufficient to increase overall lifespan (expression of *pck-1* in intestine is not effective; [73]). Disruption of *pck-1* can shorten lifespan [73], although we (S1 Table) do not find this outcome for RNAi interventions (technical differences likely explain the outcomes).

Interestingly, *pck-2*, which we show is another PEPCK-C homolog, seems to serve a different role from *pck-1* in promoting adult health. The restriction of *pck-2* expression to the very anterior and posterior gut in adults suggests a non-autonomous instructive role for PCK-2 in locomotory aging, and aging of the whole animal. More specifically, although a *pck-2* translational reporter (native promoter + whole protein GFP tagged) is expressed in multiple tissue types including the muscles and intestine, treatment with *daf-16*(RNAi) or altering the DAF-16 binding site in the *pck-2* promoter abolishes PCK-2 expression only in particular cells of the intestine, yet this deficit is sufficient to disrupt health benefits. Although GFP-based transgene reporters are not always accurate in reflecting native expression, our finding that adult PCK-2::GFP expression in intestine can drive whole-animal metabolic coordination and hence benefits in locomotion (nerve and muscle) and mid-life survival (system-wide health), implies that metabolic changes in certain gut cells can set the animal-wide maintenance tone. Future studies that express *pck-2* specifically in the intestine in a *pck-2* null background should further test this model.

The specific expression for PCK-2 suggests that metabolic regulation/change in particular sets of cells might constitute a driver mechanism that directs whole-animal changes. PCK-2 might serve as a *daf-16*-dependent first responder that signals for metabolic shift in other cells and tissues. General metabolic director cells would be of interest to identify as their specific manipulation might be targeted for enhanced health maintenance during aging.

**The intestine as a driver for whole animal health**

*C. elegans* intestine is thought to/appears to perform multiple endodermal activities for the worm, functioning as digestive organ, liver and pancreas. Increasing *daf-16* in intestine upregulates DAF-16 activity in other tissues [74], with FOXO-to-FOXO signaling downregulating intestinal insulin *ins-7* [75]. In flies, overexpressing dFOXO specifically in the fat body reduces insulin-like peptide gene *dilp-2* expression in neurons and reduces insulin/IGF-1 signaling in peripheral tissues [76, 77]. Brief FOXO induction in young adult fly intestine is sufficient to
extend lifespan [15, 78]. Details of how PEPCK relates to DAF-16 signaling in aging remain to be worked out in worms, flies and possibly mammals, but the possibility of conservation of mechanism has been raised [79].

Indeed, the gut has previously been shown to differentially express food-dependent master regulators of DR [42, 80]. As pck-2 is essential for healthspan promotion under DR, the anterior and posterior intestinal cells may monitor food/energy levels and signal non-autonomously to adjust metabolism across the entire body when food becomes limiting. The nature of this signal is not yet known, although it is appealing to speculate that it might parallel mammalian glucagon signaling.

Regarding the question of how PEPCK acts in healthy aging, it might be worth noting that L1 starvation has been demonstrated to use a daf-16 dependent mechanism to restructure carbohydrate metabolism to drive carbon flux through glyoxylate and gluconeogenesis towards trehalose synthesis [81]. In C. elegans, excess glucose can be stored as glycogen or in the form of the two-carbon sugar trehalose [20, 82]. Although we have not focused on the energy storage outcome of promoting/preventing gluconeogenesis, high glucose has been shown to increase stored glycogen [20, 82] and genetic disruption of glycogen synthesis induces a daf-16-dependent shift toward trehalose elevation associated with longevity and enhanced healthspan [20]. Promotion of gluconeogenesis might be anticipated to enhance trehalose production as has been documented for the daf-16-dependent L1 starvation response [81]. In the future, it will be of interest to define how energy stores are impacted by pck-2.

**Dietary restriction-mediated longevity requires pck-2**

Our data show that at least two modes of dietary restriction, genetically induced eat-2 and food limitation, require pck-2 expression for healthspan benefits, and that eat-2 mutants exhibit elevated levels of pck-2 expression late into life. Our findings interface well with published findings documenting that DR and longevity in general involve a metabolic shift away from glycolysis and toward fatty acid metabolism [59, 61, 83, 84]. A role for the PCK-1 PEPCK-C in DR has been independently documented [73]. We define gluconeogenic metabolism as a critical requirement for DR-associated extensions of healthspan. We also emphasize that, as pck-2 and fbp-1 are required to maintain health in older wild-type animals under typical feeding conditions, gluconeogenic activity is not only required for the extended healthspan under DR, but for healthy aging in general.

Together, our results support that promoting gluconeogenic metabolism via PEPCK activity engages a conserved metabolic reorganization that results in significant healthspan benefits. How enhanced gluconeogenesis impacts whole-animal/tissue-specific glucose levels remains to be determined, but a switch to gluconeogenic metabolism may result in a shift to health-promoting storage or utilization of energy stores. For example, gluconeogenesis requires amino acid fuel provided by autophagic proteolysis [85]. Increasing gluconeogenic flux may further promote autophagy, which is required for most longevity pathways (including DR; [86]), and may engage healthy metabolism. That gluconeogenesis can promote healthy aging is supported by studies in other models linking enhanced gluconeogenesis to increased lifespan [87–89], suggesting a conserved pro-longevity mechanism. Interestingly, pck-2 expression decreases as C. elegans age [90], but gluconeogenic genes, including pck-2, are up-regulated in long-lived dauer and daf-2 insulin signaling mutants [61, 83, 84]. We also show that pck-2 expression remains elevated during adulthood in long-lived eat-2 DR mutants. We suggest that gluconeogenic capacity, possibly in key cells, is a common component of healthy aging, and compromising this capacity later in life undermines successful maintenance.
A positive feedback loop for metabolic conditions that promote adult health?

Disrupting glycolytic gene pfk can both induce a fluorescent biomarker for the DR state and increase PCK-2 expression, supporting a model in which glycolytic disruptions increase DAF-16 signaling to trigger gluconeogenic/DR metabolism and extend healthspan (Fig 4I). This model is supported by our data showing that DAF-16 is required for the healthspan benefits seen with glycolytic gene disruptions.

Interestingly, although a transcriptional reporter for PCK-2 is expressed in a food-dependent manner, the functional PCK-2 translational reporter is not sensitive to food status. Given published datasets that suggest that elevated PCK-2 transcription levels track with increased longevity/healthspan [61, 83, 84], it seems possible that the high constitutive levels we find when intact pck-2 is over-expressed reflect a gain-of-activity consequence of high PCK-2. We speculate that high PCK-2 induces a metabolic state in which DAF-16 activity increases, resulting in a positive regulatory loop that stimulates DAF-16 activity and reinforces a healthy metabolic state. As reduced insulin signaling does not require PCK-2 to promote healthy aging via DAF-16, PCK-2 may signal parallel to the insulin pathway to intersect with DAF-16. Alternatively, PCK-2 may increase DAF-16 activity by downregulating insulin signaling, which may lie downstream of PCK-2 in regulating lifespan and healthspan (Fig 4I).

Overall, our data support a model in which nutritional signals engage glucose metabolism to influence the quality of aging via DAF-16 (Fig 4I). Under DR or glucose limitation, increased DAF-16 activity promotes gluconeogenic gene expression, resulting in lifespan and healthspan benefits. When food is plentiful, glycolytic activity effectively suppresses DAF-16, which compromises both locomotory healthspan and lifespan. As high levels of glucose inhibit DAF-16 [16], buildup of toxic metabolites [52] may result from inhibition of gluconeogenic metabolism consequent to elevated glycolytic activity. Our data mechanistically tie high glucose intake with suppression of an anti-aging, pro-healthspan gluconeogenic pathway, underscoring a rationale for limiting dietary glucose. The importance of the gluconeogenesis flux in adult health suggests gluconeogenesis process as a novel target for anti-aging interventions as well as for anti-glucose toxicity approaches.

Materials and methods

C. elegans cultures

The following strains used in this study were provided by the Caenorhabditis Genetics Center (CGC, University of Minnesota), which is funded by NIH Office of Research Infrastructure Programs (P40 OD010440): wild-type Bristol N2, TJ1052 age-1(hx546), GR1307 daf-16 (mgDf50), DA1116 cat-2(ad1116), RB1966 R11A5.4(ok2586), CF1553 N2;muIS84[P_sod-3:gf]. The following are transgenic lines generated for these studies: ZB4912 bzIs191[P_pck-2:gf P_mec-4:cherry], ZB4913 bzIs192[P_pck-2:gf P_mec-4:cherry], ZB4914 bzEx283[P_pck-2:gf P_mec-4:cherry]. All strains were grown under standard conditions at 20˚C [93] on Escherichia coli strain OP50-1 or HT115 for RNAi. All experiments were performed at 20˚C. Metformin (Sigma-Aldrich, Catalog # D15,095–9) was added directly to the NGM agar media to a final concentration 50 mM from a 1 M aqueous stock. For high-glucose plates, D-(-)-Glucose (Sigma-Aldrich, Catalog # D9434) was added to the liquid media to a final concentration of 2% or 4% from a 24% aqueous stock.
Generation of the \( fbp-1 \) (K07A3.1) RNAi clone

As the RNAi feeding vector for \( fbp-1 \) targeting was not available in the Ahringer library [94, 95], we constructed our own vector using the following primers for cloning into the pL4440 plasmid using NheI cut sites:

\[
5'-CGCGCGCTAGCATACGG AATCGCTG-3' \\
5'-GCGCGGGCTAGCATCGATTTTTTTTAA-3'
\]

Generation of N2;Is\([P_{pck-2}\text{-gfp}] \), N2;Is\([P_{pck-2}\text{-pck-2::gfp}] \), and N2;Ex\([P_{pck-2\text{mut}}\text{-pck-2::gfp}] \) lines

The \( P_{pck-2}\text{-gfp} \) and \( P_{pck-2}\text{-pck-2::gfp} \) vectors were constructed using the In-Fusion HD cloning system (Clontech). For \( P_{pck-2}\text{-gfp} \), we amplified 2.9 kb of sequence upstream of the \( pck-2 \) start codon using the primers

\[
5'-CGACTCTAGAGGATCCGACTGATTGAATGAATGACTGGAG TGTATT GG-3' \\
5'-CCAATCCCGGGGATCCGATTCTCTACACCGACTGTGCCGAAAC TTT-3'
\]

and inserted the product into the pPD95.77 vector (Addgene) linearized with BamHI. For \( P_{pck-2}\text{-pck-2::gfp} \), we amplified the \( pck-2 \) coding region and 3'-UTR using the primers

\[
5'-GGAGGAC CCTTGAGGTACCAGATCGTTGATCCAAACCTTCTTACTCC -3' \\
5'-TCATTTT TTCTACCGGTACCGG CAATGTCTGGACTCTCTTCTCTTGAGC -3'
\]

and inserted the product into the \( P_{pck-2}\text{-gfp} \) vector linearized with KpnI. The \( P_{pck-2}\text{-mut} \) vector containing the mutated DAF-16 binding site in the \( pck-2 \) promoter region was generated using the QuikChange II site-directed mutagenesis kit (Agilent) with the \( P_{pck-2}\text{-pck-2::gfp} \) vector and the following primers:

\[
5' - GAACTGGAGAAAAAAACAAACTTTCCACTTTTCTATAATAT GTTCTGCA TTCTAAGCTCTTCCCACTTTTCTATAATAT GTCCTGCA TTTCCAAAAATTTTTTTTCTTAAAGAACATTAACTTTTAAAT-3' \\
5' - ATTTAA AGTTAACTGGAGAAAAAAACAAACTTTCCACTTTTCTATAATAT GTCCTGCA TTTCCAAAAATTTTTTTTCTTAAAGAACATTAACTTTTAAAT-3'
\]

All constructs were injected at 50 ng/ul into wild-type animals along with a \( P_{mec-4}\text{-mCherry} \) co-injection marker [96] (100 ng/ul). \( P_{pck-2}\text{-gfp} \) and \( P_{pck-2}\text{-pck-2::gfp} \) transgenics were γ-irradiated to identify stably transformed lines as described [97].

Lifespan assays

Lifespan analyses were performed in the same manner for all strains. For each experiment, except where otherwise noted, the lifespan of 60 animals was measured in each trial. RNAi assays were performed using a feeding library as described [94, 95], with some modifications. For each RNAi clone tested, we placed 15 L4-stage larvae on 3 RNAi plates containing 4 mM IPTG with 5 animals per plate and allowed these to develop to adulthood and then lay eggs over 24 hours. These parental animals were then removed from the plates. 48 hours later, 60 (day 2) L4 larvae were transferred to fresh plates. These animals were transferred to fresh plates every day during the progeny production period, and then every other day thereafter. Animals that did not move when gently prodded were scored as dead. Animals that crawled off the plate or died from vulva bursting or internal hatching were not included in lifespan counts.

\( pck-2 \) overexpression studies

Studies with the \( pck-2 \) over-expressor (\( P_{pck-2}\text{-pck-2::gfp} \)) used GFP expressed from the \( pck-2 \) promoter (\( P_{pck-2}\text{-gfp} \)) as a control. We used \( P_{pck-2}\text{-gfp} \) (lacking the PKC-2 coding sequence) to guard against the possibility that elevating the promoter alone might have biological consequences (such as titrating out transcription factors like DAF-16). Potential effects of having high copy numbers of the \( pck-2 \) promoter would be not be apparent in non-transgenic wild-type animals. In Supplemental S3C Fig, nontransgenic wild-type animals generated during the outcrossing of the transgenic lines were used as controls in lifespan studies with \( P_{pck-2}\text{-gfp} \) and \( P_{pck-2}\text{-pck-2::gfp} \) animals.
Locomotion assays

Animals were raised from eggs similar to the lifespan assays. 40 individuals were measured for body bend rate in liquid. Briefly, 4 animals were placed in 20 μl M9 buffer on a glass slide and filmed for 30 seconds using a Qimaging Rotera-XR digital camera attached to a dissecting microscope and Streampix imaging software (ver. 3.17.2, NorPix). Swimming rates were calculated using CeleST [92]. Note that although assays in some different panels were sampled at different timepoints in mid-late adulthood due to convenience, locomotory decline progresses over time and comparison was always done to control on the same day. Absolute scores can vary between days/experiments, but relative decline is reproducible. Because baseline scores can vary, our conclusions are only drawn from data from trials within single experiments, not across different experiments.

Age pigment fluorescence spectroscopy

Age pigment fluorescence intensity was measured as described [64]. Wild-type animals were raised from eggs on plates as in the lifespan assays until day 5 of life. On day 5, 50 animals were transferred to 50 μl 10 mM NaN₃ solution in a single well of a 96-well white FluoroNunc plate (Nalge Nunc Int’l). The animals were scanned using an in vivo spectrofluorimeter (Fluorolog-3, Jobin Yvon Inc., Edison NJ). Peak age pigment fluorescence intensity was determined by scanning through a range of excitation wavelengths from 280–410 nm and an emission wavelength of 430 nm. DataMax data acquisition software (v. 2.20, Jobin Yvon Inc.) and Grams/32 data manipulation software (v. 4.14, Galactic Industries Corp.) were used to process the emission data. Scores are the average age pigment fluorescence intensity levels of three independent trials.

Pharyngeal pumping assays

Wild-type animals were raised from eggs on NGM control or 2% or 4% glucose plates similar to the lifespan assays. On day 5 of life, the pharyngeal pumping rates of 35 individuals were measured in real time by scoring pharyngeal pumping by eye under a dissecting microscope for 30 seconds.

DR assays on solid media

For the experiment described in Fig 4A, 4B and 4F and S5 Fig, bleached eggs were placed on plates layered with either undiluted (AL) OP50-1 (for Fig 4A) or HT115 (for Fig 4F) overnight culture or 1:10 (DR) diluted OP50-1 (for Fig 4A) or HT115 (for Fig 4F) culture (the HT115 overnight cultures were grown in the presence of 4 mM IPTG to induce RNAi expression). OP50-1/HT115 were killed immediately after drying on AL and DR plates using 2 x 10⁶ uJ at 254 nm using a Stratagene UV Stratalinker 1800 instrument. For the experiment described in Fig 4B, live OP50-1 cultures were placed on plates containing 100 μg/ml ampicillin to arrest growth at 7 x 10⁶ cfu (AL) and 7 x 10⁹ cfu (DR) concentrations. N2;Is[Pₚₛₒ₅₋₃::gfp] animals were placed on these plates at the L4 stage (day 2 of life) and assays were performed on day 4 of life.

Quantitation of Pₛₒ₅₋₃::GFP and Pₚck₋₂::gfp expression

N2; mu184[Pₛₒ₅₋₃::gfp] or N2; Is[Pₚck₋₂::gfp] animals were raised from eggs as in the lifespan assays. On day 5 of life for N2; mu184[Pₛₒ₅₋₃::gfp] and day 7 for N2; Is[Pₚck₋₂::gfp] animals, we measured GFP fluorescence in 50 animals per condition using a spectrofluorimeter as in the age pigment quantitation assays described above.
Fluorescence microscopy

N2;Is[Pck-2::gfp], N2;Is[Pck-2::gfp] and N2;Ex[Pck-2::gfp] animals were raised from eggs on RNAi plates similar to the lifespan assays. On day 7 of life, animals were placed in 10 mM NaN₃ or M9 liquid media and observed under a 40x objective lens using a Zeiss Axioplan 2 microscope equipped with an X-cite Series 120 (EXPO Photonic Solution, Inc.) fluorescence illuminator. Micrographs were obtained using an Optronics digital microscope camera and Magnafire processing software.

cDNA synthesis and qPCR

C. elegans samples (approximately 10,000 animals per sample) were collected by washing using M9 buffer with 0.01% Tween20. To remove residual OP50 E. coli culture, the worm pellet was applied onto M9 buffer with 10% sucrose solution and spun in a clinical centrifuge at full speed for 1 min. The resulting pellet was transferred into a pre-chilled (with liquid N2) mortar and ground with mortar and pestle. The resulting sample grind was stored at -80°C. Total RNA was extracted from the frozen grind of the worms sample using TRIzol extraction (Life Technology, 15596–026). The resulting sample was treated with DNase I (NEB M0303L) and followed by phenol:chloroform (1,1) purification. The resulting total RNA was re-suspended with dH₂O and stored at -80°C. First strand cDNA synthesis was done using SuperScript III RT kit (Life Technology, 18080–044) and using 1ug of the total RNA and oligo(dT)20 for mRNA enrichment. The resulting cDNA samples were used for qPCR. All qPCR reactions were done in triplicate, using KAPA SYBR FAST kit (KAPA Biosystems). Each mRNA level was quantified with reference to the mRNA level of tba-1 [98]. The primers used for pck-2 were: CGATATCACCACATGGCTTG and GCTTTCCCAGTCTGGATGAA; for F47B8.10: GCTTCAACAAGCTGATTCTC and CGAAGACGTACACGGATGA.

Mitochondria staining

N2;Is[Pck-2::gfp] transgenic animals were raised from eggs on plates similar to the lifespan assays. On day 7 of life, 20 animals were washed and resuspended in 1 μg/ml MitoTracker Red CMXRos (Molecular Probes, M7512) in M9 for 6 hours at 20°C in the dark. Animals were washed four times in M9, then placed on seeded NGM plates for one hour at 20°C in the dark. Mitochondrial staining was imaged as described under Fluorescence Microscopy above.

Statistical analyses

Log-rank (Mantel-Cox) tests, Gehan-Breslow-Wilcoxon tests, ANOVA, and unpaired t tests were performed using GraphPad Prism version 5.00 for Windows (GraphPad Software, San Diego California).

Supporting information

S1 Table. Lifespan data. These are data described in the main body of the text or for the Supplemental Figures. “N total” = the total number of animals monitored in all trials for a particular condition; “N dead” = the number of animals that were counted as dead due to “old age”, i.e. not lost from the assay or due to any developmental or physical defect; “N censored” = number of animals removed from the assay due to causes other than “old age”; these include internal hatching, vulval bursting, or crawling off the plate. AL = ad libitum; DR = dietary restriction. P values are from Log-rank tests comparing lifespan curves. The RNAi clones used in this study were sequence confirmed to be specific for their target genes.

(TIF)
S1 Fig. Excess glucose availability negatively impacts healthspan of young and old animals; conversely, the benefits of disrupting glycolysis do not include extended maximal lifespan.

(A) Swimming rates of wild-type animals raised on control plates or plates containing 4% glucose. Exposure to excess glucose results in significant decreases in swimming rates in young animals (day 5 from hatching; \( P = 0.0004 \), unpaired t test). \( n = 40 \) animals per condition from a single trial; error bars represent SEM; \( *** P < 0.001 \). (B) Swimming rates of wild-type animals raised on control plates or plates containing 4% glucose. Exposure to excess glucose results in significant decreases in swimming rates late in life (day 13 from hatching; \( P = 0.0062 \), unpaired t test). \( n = 40 \) animals per condition from a single trial; error bars represent SEM; \( ^* P < 0.01 \). (C) Swimming rates of wild-type animals raised on control plates or plates containing 2% glucose. Unlike 4% glucose (Figs 1A and S1A and S1B), exposure to 2% glucose does not impact swimming rates on day 8 of life (n.s., not significant, unpaired t test). \( n = 40 \) animals per condition from a single trial; error bars represent SEM. (D) Maximal lifespan values of WT animals treated with pfk(RNAi), pyk-2(RNAi), and age-1(RNAi). While glycolytic gene disruptions significantly increase median survival (see Fig 1D and S1 Table), maximal lifespans of animals treated with pfk(RNAi) or pyk-2(RNAi) are not significantly different than those of empty vector (EV) controls (not significant, n.s., by one way ANOVA), suggesting that inhibiting glycolysis and decreasing insulin signaling might increase healthspan via a common pathway. Also unlike in wild-type, inhibiting gluconeogenesis does not significantly decrease locomotory healthspan in the age-1 background (fbp-1(RNAi) and pck-2(RNAi) vs. empty vector control; n.s., not significant by one way ANOVA), demonstrating that reduced insulin signaling can rescue the detrimental healthspan effects seen with gluconeogenic gene disruptions. Data are pooled from 6 independent trials, \( n = 60 \) animals per condition per trial (see Fig 1D and S1 Table). \( **** P < 0.0001 \); one way ANOVA analyses performed with Dunnett’s multiple comparisons test. Maximal lifespan is defined as the mean age at death of the longest-lived 10% of a given population [100].

(TIF)

S2 Fig. The healthspan effects of glycolytic and gluconeogenic disruptions are absent in the age-1 insulin signaling mutant background.

(A) Swimming rates of long-lived age-1 insulin pathway mutants treated with RNAi against glycolytic genes pfk and pyk-2 and gluconeogenic genes fbp-1 and pck-2. Unlike in wild-type (Fig 1C), disrupting glycolysis does not further increase healthspan in the age-1 background (pfk(RNAi) and pyk-2(RNAi) vs. empty vector control; n.s., not significant by one way ANOVA), suggesting that inhibiting glycolysis specifically promotes mid-life vigor without extending end-of-life frailty. age-1 (RNAi), which extends maximal lifespan by inhibiting insulin signaling [99], was used as a positive control. Maximal lifespan data values are averages from 6 independent trials, \( n = 60 \) animals per condition per trial (see Fig 1D and S1 Table). \( **** P < 0.0001 \); one way ANOVA analyses performed with Dunnett’s multiple comparisons test. (B) Survival curves of age-1 mutants treated with RNAi against gluconeogenic gene pck-2. The harmful effects of pck-2 RNAi (Fig 1D) are absent in age-1 mutants; the survival curve is not altered with pck-2 RNAi treatment (n.s., not significant, Log-rank), and median survival is actually increased by 5.13%.

(TIF)

S3 Fig. Expression of a pck-2 translational reporter increases healthspan measures, and expression of a pck-2 promoter -only transcriptional reporter does not affect healthspan measures.

(A) Wild-type animals expressing the pck-2 translational reporter on day 7 of life. In the panel on the left, PCK-2::GFP is observed to be in the cytoplasm in posterior cells of the
intestine; the panel on the right shows mitochondria in the same intestinal cells stained with the red-fluorescent dye MitoTracker. 24 out of 60 animals displayed this PCK-2::GFP localization pattern on day 7. Bar, 50 μm. (B) Swimming rate profiles of wild-type animals carrying integrated forms of either a transcriptional reporter for *pck-2* ([*Is*][P*\_pck-2::gfp*]) or *pck-2* expressed from its own promoter from an integrated transgene array ([*Is*][P*\_pck-2::gfp*], "*pck-2 OE"), used as the over-expressor strain) on day 9 of life. *pck-2* overexpression (OE) significantly increases locomotory ability (*P* < 0.0001, unpaired t test). Data are from a single trial, n = 30 animals per condition. Error bars represent SEM. *** *P* < 0.0005, unpaired t test. (C) Survival curves of wild-type animals expressing a transcriptional reporter for *pck-2* ([*Is*][P*\_pck-2::gfp*], "*pck-2 OE"), nontransgenic wild-type siblings generated during the outcrossing of the [*Is*][P*\_pck-2::gfp*] line ("WT Cntl for *pck-2* OE Outcross"), *pck-2* expressed from its own promoter ([*Is*][P*\_pck-2::gfp*], "*pck-2 OE"), and wild-type animals generated during the outcrossing of the [*Is*][P*\_pck-2::gfp*] line ("WT Cntl for *pck-2* OE Outcross"), all raised on *E. coli* strain OP50. While the survival curve of the [*P*\_pck-2::gfp*] animals is not significantly different from the curve of nontransgenic wild-type controls ("WT Cntl for *pck-2* OE Outcross"; *P* = 0.6283, Log-rank), the curve of the [*P*\_pck-2::gfp*] animals is significantly right-shifted as compared to both nontransgenic wild-type controls ("WT Cntl for *pck-2* OE Outcross"), and [*P*\_pck-2::gfp*] animals (*P* < 0.0001 for both, Log-rank). Data are pooled from two independent trials, n = 100 animals per line per trial. n.s. = not significant; **** *P* < 0.0001. See S1 Table for details. (D) Swimming rates of wild-type (WT) animals and wild-type animals expressing a *pck-2* transcriptional reporter (WT; [*Is*][P*\_pck-2::gfp*]) on day 11 of life. Swimming rates of the two groups are not significantly different in mid-life (n.s., unpaired t test). Data are from a single trial, n = 30 animals per condition. Error bars represent SEM. n.s. = not significant. (E) Survival curves of wild-type (WT) animals and wild-type animals expressing a *pck-2* transcriptional reporter ([*Is*][P*\_pck-2::gfp*], "*pck-2 OE"), used as the over-expressor strain, to WT (*P* < 0.0001, Log-rank) and to WT (*P* = 0.0009, Log-rank). Also as in S3C Fig, *pck-2* over-expressors have a survival curve that is significantly right-shifted as compared to [*P*\_pck-2::gfp*] animals (*P* = 0.0092, Log-rank) and to WT (*P* = 0.00118, Log-rank). Data are pooled from 7 independent trials for WT controls; 4 independent trials for *pck-2* mutants; 2 independent trials for [*P*\_pck-2::gfp*] animals; 6 independent trials for *pck-2* over-expressors; approximately 60 animals per trial per strain. See S1 Table for details.

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**S4 Fig.** Alignment between *C. elegans* glucose-6-Phosphate translocase ortholog and human, mouse, zebrafish, and fly glucose-6-Phosphate exchanger. Alignment of the *C. elegans* glucose-6-phosphate translocase ortholog (F47B8.10) with the human glucose-6-phosphate exchanger isoform 1 (E value = 3e-31; 26.56% identity; NCBI reference sequence NP_001157749), mouse glucose-6-phosphate exchanger SLC37A4 isoform b (E value = 5e-30; 25.30% identity; NCBI reference sequence NP_001280559), zebrafish glucose-6-phosphate translocase isoform X1 (E value = 3e-24; 26.48% identity; NCBI reference sequence XP_009289762.1), and Mediterranean fruit fly Ceratitis capitata glucose-6-phosphate exchanger SLC37A2 isoform X2 (E value = 3e-04; 23.00% identity; NCBI reference sequence XP_004522689.1). No ortholog has been documented in *Drosophila melanogaster*. "Cooler" blue colors indicate residues not conserved, while "hotter" red colors indicate increased sequence conservation. Final alignment performed with the PRALINE multiple sequence alignment tool.
alignment website (http://www.ibi.vu.nl/programs/pralinewww/). A domain/motif search using Prosite (http://prosite.expasy.org/) reveals a single consensus domain (MFS, or major facilitator superfamily domain, a transmembrane substrate transporter domain), which spans the orthologous sequences.

S5 Fig. Gluconeogenic gene fbp-1 expression is required for healthspan extension under dietary restriction. (A) Survival curves of long-lived, dietary-restricted cat-2(ad1116) mutants treated with RNAi against gluconeogenic gene fbp-1. fbp-1(RNAi) decreases median survival by 10% and shifts the survival curve to the left ($P < 0.0001$, Log-rank). Data are pooled from 5 independent trials, n = 60 animals per condition per trial. (B) Survival curves of wild-type animals raised under ad libitum (AL) or dietary restriction (DR) conditions, with and without fbp-1(RNAi). Median survival of animals expressing the vector control was higher under DR conditions as compared to well-fed controls (an 18.62% increase, see S1 Table for details), and survival was increased ($P = 0.0039$, Log-rank test). Disrupting gluconeogenic gene fbp-1 completely abolished these beneficial DR effects: animals raised on diluted gluconeogenic RNAi bacteria had survival curves that were not significantly (n.s., Log-rank) different from those of well-fed animals. Data are pooled from 2 independent trials, n = 60 animals per condition per trial.

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References

1. de Cabo R, Carmona-Gutierrez D, Berri M, Hall MN, Madeo F. The search for antiaging interventions: from elixirs to fasting regimens. Cell. 2014; 157(7):1515–26. Epub 2014/06/21. https://doi.org/10.1016/j.cell.2014.05.031 PMID: 24949965; PubMed Central PMCID: PMC4254402.

2. Adler AI, Boyko EJ, Ahroni JH, Stensel V, Forsberg RC, Smith DG. Risk factors for diabetic peripheral sensory neuropathy. Results of the Seattle Prospective Diabetic Foot Study. Diabetes care. 1997; 20(7):1162–7. Epub 1997/07/01. https://doi.org/10.2337/diabetes.20.7.1162 PMID: 9203456.

3. Ford ES, Giles WH, Dietz WH. Prevalence of the metabolic syndrome among US adults: findings from the third National Health and Nutrition Examination Survey. JAMA: the journal of the American Medical Association. 2002; 287(3):356–9. Epub 2002/01/16. https://doi.org/10.1001/jama.287.3.356 PMID: 11790215.

4. Hanefeld M, Fischer S, Julius U, Schulze J, Schwanebeck U, Schmechel H, et al. Risk factors for myocardial infarction and death in newly detected NIDDM: the Diabetes Intervention Study, 11-year follow-up. Diabetologia. 1996; 39(12):1577–83. Epub 1996/12/01. https://doi.org/10.1007/s001250050617 PMID: 8960845.

5. Klein R. Hyperglycemia and microvascular and macrovascular disease in diabetes. Diabetes care. 1995; 18(2):258–68. Epub 1995/02/01. https://doi.org/10.2337/diacare.18.2.258 PMID: 7729308.

6. Standl E, Balletshofer B, Dahl B, Weichenhain B, Stiegler H, Hornmann A, et al. Predictors of 10-year macrovascular and overall mortality in patients with NIDDM: the Munich General Practitioner Project. Diabetologia. 1996; 39(12):1540–5. Epub 1996/12/01. https://doi.org/10.1007/s001250050612 PMID: 8960840.

7. Uusitupa MI, Niskanen LK, Siitonen O, Voutilainen E, Pyorala K. Ten-year cardiovascular mortality in relation to risk factors and abnormalities in lipoprotein composition in type 2 (non-insulin-dependent) diabetic and non-diabetic subjects. Diabetologia. 1993; 36(11):1175–84. Epub 1993/11/01. https://doi.org/10.1007/BF00401063 PMID: 8270133.

8. Wei M, Gaskill SP, Haffner SM, Stern MP. Effects of diabetes and level of glycemia on all-cause and cardiovascular mortality. The San Antonio Heart Study. Diabetes care. 1998; 21(7):1167–72. Epub 1998/07/08. https://doi.org/10.2337/diacare.21.7.1167 PMID: 9653614.

9. American Diabetes A, Bantle JP, Wylie-Rosett J, Albright AL, Apovian CM, Clark NG, et al. Nutrition recommendations and interventions for diabetes: a position statement of the American Diabetes Association. Diabetes care. 2008; 31 Suppl 1:S61–78. Epub 2008/01/10. https://doi.org/10.2337/dc08-S061 PMID: 18165339.

10. Ostler JE, Maurya SK, Dials J, Roof SR, Devor ST, Ziolo MT, et al. Effects of insulin resistance on skeletal muscle growth and exercise capacity in type 2 diabetic mouse models. Am J Physiol Endocrinol Metab. 2014; 306(6):E592–605. Epub 2014/01/16. https://doi.org/10.1152/ajpendo.00277.2013 PMID: 24425761; PubMed Central PMCID: PMC3948983.

11. Park SW, Goodpaster BH, Lee JS, Kuller LH, Boudreau R, de Rekeneire N, et al. Excessive loss of skeletal muscle mass in older adults with type 2 diabetes. Diabetes care. 2009; 32(11):1993–7. Epub 2009/06/25. https://doi.org/10.2337/dc09-0264 PMID: 19549734; PubMed Central PMCID: PMC2768193.

12. Park SW, Goodpaster BH, Strotmeyer ES, Kuller LH, Broudeau R, Kammere C, et al. Accelerated loss of skeletal muscle strength in older adults with type 2 diabetes: the health, aging, and body composition study. Diabetes care. 2007; 30(6):1507–12. Epub 2007/03/17. https://doi.org/10.2337/dc06-2537 PMID: 17383749.

13. Alcantar-Fernandez J, Navarro RE, Salazar-Martinez AM, Perez-Andrade ME, Miranda-Rios J. Caenorhabditis elegans respond to high-glucose diets through a network of stress-responsive transcription factors. PLoS One. 2018; 13(7):e0199888. Epub 2018/07/11. https://doi.org/10.1371/journal.pone.0199888 PMID: 29990370; PubMed Central PMCID: PMC5903904.

14. Choi SS. High glucose diets shorten lifespan of Caenorhabditis elegans via ectopic apoptosis induction. Nutr Res Pract. 2011; 5(3):214–8. Epub 2011/07/23. https://doi.org/10.4162/nrp.2011.5.3.214 PMID: 21779524; PubMed Central PMCID: PMC3133753.

15. Dobson AJ, Eczurra M, Flanagan CE, Summerfield AC, Piper MDW, Gens D, et al. Nutritional Programming of Lifespan by FOXO Inhibition on Sugar-Rich Diets. Cell Rep. 2017; 18(2):299–306. Epub 2017/01/12. https://doi.org/10.1016/j.celrep.2016.12.029 PMID: 28076775; PubMed Central PMCID: PMC5263231.

16. Lee SJ, Murphy CT, Kenyon C. Glucose shortens the lifespan of C. elegans by downregulating DAF-16/FOXO activity and aquaporin gene expression. Cell Metab. 2009; 10(5):379–91. Epub 2009/11/04. https://doi.org/10.1016/j.cmet.2009.10.003 PMID: 19883616; PubMed Central PMCID: PMC2887095.
17. Marcellino BK, Ekasumara N, Mobbs CV. Dietary Restriction and Glycolytic Inhibition Reduce Proteotoxicity and Extend Lifespan via NHR-49. Curr Neurobiol. 2018; 9(1):1–7. Epub 2019/03/02. PMID: 30820135; PubMed Central PMCID: PMC6390974.

18. Schlotterer A, Kukudov G, Bozorgmehr F, Hutter H, Du X, Oikonomou D, et al. C. elegans as model for the study of high glucose-mediated life span reduction. Diabetes. 2009; 58(11):2450–6. Epub 2009/08/14. https://doi.org/10.2337/db09-0567 PMID: 19675139; PubMed Central PMCID: PMC2768179.

19. Schulz TJ, Zarse K, Voigt A, Urban N, Birringer M, Ristow M. Glucose restriction extends Caenorhabditis elegans life span by inducing mitochondrial respiration and increasing oxidative stress. Cell Metab. 2007; 6(4):280–93. Epub 2007/10/03. https://doi.org/10.1016/j.cmet.2007.08.011 PMID: 17906557.

20. Seo Y, Kingsley S, Walker G, Mondoux MA, Tissenbaum HA. Metabolic shift from glycolgen to trehalose promotes lifespan and healthspan in Caenorhabditis elegans. Proc Natl Acad Sci U S A. 2018; 115(12):E2791–E800. Epub 2018/03/08. https://doi.org/10.1073/pnas.1714781115 PMID: 29511104; PubMed Central PMCID: PMC5866546.

21. Friedman DB, Johnson TE. A mutation in the age-1 gene in Caenorhabditis elegans lengthens life and reduces hermaphrodite fertility. Genetics. 1988; 118(1):75–86. Epub 1988/01/01. PMID: 6808934; PubMed Central PMCID: PMC1203268.

22. Hsu AL, Murphy CT, Kenyon C. Regulation of aging and age-related disease by DAF-16 and heat-shock factor. Science. 2003; 300(5622):1142–5. Epub 2003/05/17. https://doi.org/10.1126/science.1083701 PMID: 12750321.

23. Kenyon C, Chang J, Gensch E, Rudner A, Tabtiang R. A C. elegans mutant that lives twice as long as wild type. Nature. 1993; 366(6540):461–4. Epub 1993/12/02. https://doi.org/10.1038/366461aa PMID: 8247153.

24. Kimura KD, Tissenbaum HA, Liu Y, Ruvkun G. daf-2, an insulin receptor-like gene that regulates longevity and diapause in Caenorhabditis elegans. Science. 1997; 277(5328):942–6. Epub 1997/08/15. https://doi.org/10.1126/science.277.5328.942 PMID: 9252323.

25. Lin K, Dorman JB, Walker G, Mondoux MA, Tissenbaum HA. Metabolic shift from glycogen to trehalose promotes lifespan and healthspan in Caenorhabditis elegans. Proc Natl Acad Sci U S A. 2018; 115(12):E2791–E800. Epub 2018/03/08. https://doi.org/10.1073/pnas.1714781115 PMID: 29511104; PubMed Central PMCID: PMC5866546.

26. Friedman DB, Johnson TE. A mutation in the age-1 gene in Caenorhabditis elegans lengthens life and reduces hermaphrodite fertility. Genetics. 1988; 118(1):75–86. Epub 1988/01/01. PMID: 6808934; PubMed Central PMCID: PMC1203268.

27. Tullet JM, Hertweck M, An JH, Baker J, Hwang JY, Liu S, et al. Direct inhibition of the longevity-promoting factor SKN-1 by insulin-like signaling in C. elegans. Cell. 2008; 132(6):1025–38. Epub 2008/03/25. https://doi.org/10.1016/j.cell.2008.03.010 PMID: 18358814; PubMed Central PMCID: PMC2367249.

28. Vowels JJ, Thomas JH. Genetic analysis of chemosensory control of dauer formation in Caenorhabditis elegans. Genetics. 1992; 130(1):105–118. Epub 1992/01/01. PMID: 13711061; PubMed Central PMCID: PMC137532.

29. Priebe S, Menzel U, Zarse K, Groth M, Platzer M, Ristow M, et al. Extension of life span by impaired glucose metabolism in Caenorhabditis elegans is accompanied by structural rearrangements of the transcriptomic network. PLoS One. 2013; 8(10):e77776. Epub 2013/11/10. https://doi.org/10.1371/journal.pone.0077776 PMID: 24204961; PubMed Central PMCID: PMC3813781.

30. Lee CK, Allison DB, Brand J, Weindruch R, Prolla TA. Transcriptional profiles associated with aging and middle age-onset caloric restriction in mouse hearts. Proc Natl Acad Sci U S A. 2002; 99(23):14988–93. Epub 2002/11/07. https://doi.org/10.1073/pnas.232308992 PMID: 12419651; PubMed Central PMCID: PMC1204785.

31. Masoro EJ, McCarter RJ, Katz MS, McMahan CA. Dietary restriction alters characteristics of glucose fuel use. J Gerontol. 1992; 47(6):B202–8. Epub 1992/11/01. https://doi.org/10.1093/geronj/47.6.b202 PMID: 1430849.

32. Mobbs CV, Mastaitis JW, Zhang M, Isoda F, Cheng H, Yen K. Secrets of the lac operon. Glucose hysteresis as a mechanism in dietary restriction, aging and disease. Interdiscip Top Gerontol. 2007; 35:39–68. Epub 2006/10/26. https://doi.org/10.1159/000096555 PMID: 17063032; PubMed Central PMCID: PMC2755922.

33. Bishop NA, Guarante L. Two neurons mediate diet-restriction-induced longevity in C. elegans. Nature. 2007; 447(7144):545–9. Epub 2007/06/01. https://doi.org/10.1038/nature05904 PMID: 17538612.

34. Greer EL, Brunet A. Different dietary restriction regimens extend lifespan by both independent and overlapping genetic pathways in C. elegans. Aging Cell. 2009; 8(2):113–27. Epub 2009/02/26. https://doi.org/10.1111/j.1474-9726.2009.00459.x PMID: 19239417; PubMed Central PMCID: PMC2680339.

35. Greer EL, Dowlatshahi D, Banko MR, Vilen J, Hoang K, Blanchard D, et al. An AMPK-FOXO pathway mediates longevity induced by a novel method of dietary restriction in C. elegans. Curr Biol. 2007; 17
Laranjeiro R, Harinath G, Hewitt JE, Hartman JH, Meyer JN, et al. Swim exercise in Caenorhabditis elegans.

Laranjeiro R, Harinath G, Burke D, Braeckman BP, Driscoll M. Single swim sessions in C. elegans.

Houthoofd K, Braeckman BP, Johnson TE, Vanfleteren JR. Life extension via dietary restriction is independent of the Ins/IGF-1 signalling pathway in Caenorhabditis elegans. Exp Gerontol. 2003; 38 (9):947–54. Epub 2003/09/05. https://doi.org/10.1016/s0531-5555(03)00161-x PMID: 12954481.

Kaeberlein TL, Smith ED, Tsuchiya M, Welton KL, Thomas JH, Fields S, et al. Lifespan extension in Caenorhabditis elegans by complete removal of food. Aging Cell. 2006; 5(6):487–94. Epub 2006/11/04. https://doi.org/10.1111/j.1477-9476.2006.00238.x PMID: 17081160.

Lee D, Jeong DE, Son HG, Yamaoka Y, Kim H, Seo K, et al. SREBP and MDT-15 protect C. elegans.

Lee GD, Wilson MA, Zhu M, Wolfkow CA, de Cabo R, Ingram DK, et al. Dietary deprivation extends lifespan in Caenorhabditis elegans. Aging Cell. 2006; 5(6):515–24. Epub 2006/11/14. https://doi.org/10.1111/j.1477-9476.2006.00241.x PMID: 17096674; PubMed Central PMID: PMC2546582.

Lee GD, Wilson MA, Zhu M, Wolfkow CA, de Cabo R, Ingram DK, et al. Dietary deprivation extends lifespan in Caenorhabditis elegans. Aging Cell. 2006; 5(6):515–24. Epub 2006/11/14. https://doi.org/10.1111/j.1477-9476.2006.00241.x PMID: 17096674; PubMed Central PMID: PMC2546582.

Masoro EJ. Overview of caloric restriction and ageing. Mech Ageing Dev. 2005; 126(9):913–22. Epub 2005/05/12. https://doi.org/10.1016/j.mad.2005.03.012 PMID: 15885742.

Onken B, Driscoll M. Metformin induces a dietary restriction-like state and the oxidative stress response to extend C. elegans Healthspan via AMPK, LKB1, and SKN-1. PLoS One. 2010; 5(1): e8758. Epub 2010/01/22. https://doi.org/10.1371/journal.pone.0008758 PMID: 20090112; PubMed Central PMID: PMC2807458.

Panowski SH, Wolff S, Aguilian H, Durieux J, Dillin A. PHA-4/Foxa mediates diet-restriction-induced longevity of C. elegans. Nature. 2007; 447(7144):550–5. Epub 2007/05/04. https://doi.org/10.1038/nature05837 PMID: 17476212.

Walker G, Houthoofd K, Vanfleteren JR, Gems D. Dietary restriction in C. elegans: from rate-of-living effects to nutrient sensing pathways. Mech Ageing Dev. 2005; 126(9):929–37. Epub 2005/05/18. https://doi.org/10.1016/j.mad.2005.03.014 PMID: 15896824.

Glenn CF, Chow DK, David L, Cooke CA, Gami MS, Iser WB, et al. Behavioral deficits during early stages of aging in Caenorhabditis elegans result from locomotory deficits possibly linked to muscle frailty. The journals of gerontology Series A, Biological sciences and medical sciences. 2004; 59 (12):1251–60. Epub 2005/02/09. https://doi.org/10.1093/gerona/59.12.1251 PMID: 15699524; PubMed Central PMCID: PMC1458366.

Herrdorn LA, Schmeissner PJ, Dudaronek JM, Brown PA, Listner KM, Sakano Y, et al. Stochastic and genetic factors influence tissue-specific decline in ageing C. elegans. Nature. 2002; 419(6909):808–14. Epub 2002/10/25. https://doi.org/10.1038/nature01135 PMID: 12397350.

Liu J, Zhang B, Lei H, Feng Z, Liu J, Hsu AL, et al. Functioning in the nervous system contributes to age-dependent motor activity decline in C. elegans. Cell Metab. 2013; 18(3):392–402. Epub 2013/09/10. https://doi.org/10.1016/j.cmet.2013.08.007 PMID: 24011074; PubMed Central PMCID: PMC3811915.

Evasion K, Huang C, Yamben I, Covey DF, Kornfeld K. Anticonvulsant medications extend worm life-span. Science. 2005; 307(5707):258–62. Epub 2005/01/18. https://doi.org/10.1126/science.1105299 PMID: 15653505.

Laranjeiro R, Harinath G, Burke D, Braeckman BP, Driscoll M. Single swim sessions in C. elegans induce key features of mammalian exercise. BMC biology. 2017; 15(1):30. Epub 2017/04/12. https://doi.org/10.1186/s12915-017-0369-4 PMID: 28395669; PubMed Central PMCID: PMC5385602.

Laranjeiro R, Harinath G, Hewitt JE, Hartman JH, Royal MA, Meyer JN, et al. Swim exercise in Caenorhabditis elegans extends neuromuscular and gut healthspan, enhances learning ability, and protects against neurodegeneration. Proc Natl Acad Sci U S A. 2019; 116(47):23829–39. Epub 2019/11/07. https://doi.org/10.1073/pnas.1909211116 PMID: 31685639; PubMed Central PMCID: PMC6876156.

Lee D, Jeong DE, Son HG, Yamaoka Y, Kim H, Seo K, et al. SREBP and MDT-15 protect C. elegans from glucose-induced accelerated aging by preventing accumulation of saturated fat. Genes Dev. 2015; 29(23):2490–503. Epub 2015/12/08. https://doi.org/10.1101/gad.266304.115 PMID: 26637528; PubMed Central PMCID: PMC4691952.
Glycolysis and gluconeogenesis differentially impact C. elegans healthspan
71. Oh KJ, Han HS, Kim MJ, Koo SH. CREB and FoxO1: two transcription factors for the regulation of hepatic glucose homeostasis. BMB Rep. 2013; 46(12):567–74. Epub 2013/11/19. https://doi.org/10.5483/bmbrep.2013.46.12.248 PMID: 24238363; PubMed Central PMCID: PMC4133859.

72. Salih DA, Brunet A. FoxO transcription factors in the maintenance of cellular homeostasis during aging. Curr Opin Cell Biol. 2008; 20(2):126–36. Epub 2008/04/09. https://doi.org/10.1016/j.ceb.2008.02.005 PMID: 18394876; PubMed Central PMCID: PMC2387118.

73. Yuan Y, Hakimi P, Kao C, Kao A, Liu R, Janocha A, et al. Reciprocal Changes in Phosphoenolpyruvate Carboxykinase and Pyruvate Kinase with Age Are a Determinant of Aging in Caenorhabditis elegans. J Biol Chem. 2016; 291(3):1307–19. Epub 2015/12/04. https://doi.org/10.1074/jbc.M115.691766 PMID: 26631730; PubMed Central PMCID: PMC4714217.

74. Libina N, Berman JR, Kenyon C. Tissue-specific activities of C. elegans DAF-16 in the regulation of lifespan. Cell. 2003; 115(4):489–502. Epub 2003/11/19. https://doi.org/10.1016/s0092-8674(03)00869-4 PMID: 14622602.

75. Murphy CT, Lee SJ, Kenyon C. Tissue entrainment by feedback regulation of insulin gene expression. Nature. 2004; 429(6991):562–6. Epub 2004/06/04. https://doi.org/10.1038/nature02549 PMID: 15175753.

76. Giannakou ME, Goss M, Junger MA, Hafen E, Leevers SJ, Partridge L. Dynamics of the action of dFOXO on adult mortality in Drosophila. Aging Cell. 2007; 6(4):429–38. Epub 2007/05/01. https://doi.org/10.1111/j.1474-9726.2007.00290.x PMID: 17465980.

77. Yuan Y, Hakimi P, Kao C, Kao A, Liu R, Janocha A, et al. Reciprocal Changes in Phosphoenolpyruvate Carboxykinase and Pyruvate Kinase with Age Are a Determinant of Aging in Caenorhabditis elegans. J Biol Chem. 2016; 291(3):1307–19. Epub 2015/12/04. https://doi.org/10.1074/jbc.M115.691766 PMID: 26631730; PubMed Central PMCID: PMC4714217.

78. Orlandi I, Staicu A, Vai M. Altered Expression of Mitochondrial NAD(+) Carriers Influences Yeast Chronological Lifespan by Modulating Cytosolic and Mitochondrial Metabolism. Front Genet. 2018; 9:676. Epub 2019/01/09. https://doi.org/10.3389/fgene.2018.00676 PMID: 30619489; PubMed Central PMCID: PMC6305841.
89. Vall-Llaurà N, Mir N, Garrido L, Vived C, Cabisco E. Redox control of yeast Sir2 activity is involved in acetic acid resistance and longevity. Redox Biol. 2019; 24:101229. Epub 2019/06/04. https://doi.org/10.1016/j.redox.2019.101229 PMID: 31153044; PubMed Central PMCID: PMC64719053.

90. Budovskaya YV, Wu K, Southworth LK, Jiang M, Tedesco P, Johnson TE, et al. An elt-3/elt-5/elt-6 GATA transcription circuit guides aging in C. elegans. Cell. 2008; 134(2):291–303. Epub 2008/07/30. https://doi.org/10.1016/j.cell.2008.05.044 PMID: 18662544; PubMed Central PMCID: PMC4719053.

91. Kashyap L, Perera S, Fisher AL. Identification of novel genes involved in sarcopenia through RNAi screening in Caenorhabditis elegans. The journals of gerontology Series A, Biological sciences and medical sciences. 2012; 67(1):56–65. Epub 2011/05/20. https://doi.org/10.1093/gerona/grl072 PMID: 21593014; PubMed Central PMCID: PMC3260486.

92. Restif C, Ibanez-Ventoso C, Vora MM, Guo S, Metaxas D, Driscoll M. CeleST: computer vision software for quantitative analysis of C. elegans swim behavior reveals novel features of locomotion. PLoS Comput Biol. 2014; 10(7):e1003702. Epub 2014/07/18. https://doi.org/10.1371/journal.pcbi.1003702 PMID: 25033081; PubMed Central PMCID: PMC4102393.

93. Brenner S. The genetics of Caenorhabditis elegans. Genetics. 1974; 77(1):71–94. Epub 1974/05/01. PMID: 4366476; PubMed Central PMCID: PMC1213120.

94. Kamath RS, Ahringer J. Genome-wide RNAi screening in Caenorhabditis elegans. Methods. 2003; 30 (4):313–21. Epub 2003/06/28. https://doi.org/10.1016/s1046-2023(03)00050-1 PMID: 12828945.

95. Kamath RS, Fraser AG, Dong Y, Poulin G, Durbin R, Gotta M, et al. Systematic functional analysis of the Caenorhabditis elegans genome using RNAi. Nature. 2003; 421(6920):231–7. Epub 2003/01/17. https://doi.org/10.1038/nature01278 PMID: 12529635.

96. Pinan-Lucarre B, Gabel CV, Reina CP, Hulme SE, Shevkoplyas SS, Slone RD, et al. The core apoptotic executor proteins CED-3 and CED-4 promote initiation of neuronal regeneration in Caenorhabditis elegans. PLoS Biol. 2012; 10(5):e1001331. Epub 2012/05/26. https://doi.org/10.1371/journal.pbio.1001331 PMID: 22629231; PubMed Central PMCID: PMC3358320.

97. Rosenbluth RE, Cuddeford C, Baillie DL. Mutagenesis in Caenorhabditis elegans. II. A spectrum of mutational events induced with 1500 r of gamma-radiation. Genetics. 1985; 109(3):493–511. Epub 1985/03/01. PMID: 3978912; PubMed Central PMCID: PMC1216284.

98. Zhang Y, Chen D, Smith MA, Zhang B, Pan X. Selection of reliable reference genes in Caenorhabditis elegans for analysis of nanotoxicity. PLoS One. 2012; 7(3):e31849. Epub 2012/03/23. https://doi.org/10.1371/journal.pone.0031849 PMID: 22438870; PubMed Central PMCID: PMC3305280.

99. Friedman DB, Johnson TE. Three mutants that extend both mean and maximum life span of the nematode, Caenorhabditis elegans, define the age-1 gene. J Gerontol. 1988; 43(4):B102–9. Epub 1988/07/01. https://doi.org/10.1093/gerong/43.4.b102 PMID: 3385139.

100. Sulphin GL, Kaieberlein M. Dietary restriction by bacterial deprivation increases life span in wild-derived nematodes. Exp Gerontol. 2008; 43(3):130–5. Epub 2007/12/18. https://doi.org/10.1016/j.exger.2007.10.019 PMID: 18083317.