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Mono-unsaturated fatty acids link H3K4me3 modifiers to *C. elegans* lifespan

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Chromatin and metabolic states both influence lifespan, but how they interact in lifespan regulation is largely unknown. The COMPASS chromatin complex, which trimethylates lysine 4 on histone H3 (H3K4me3), regulates lifespan in *Caenorhabditis elegans*. However, the mechanism by which H3K4me3 modifiers affect longevity, and whether this mechanism involves metabolic changes, remain unclear. Here we show that a deficiency in H3K4me3 methyltransferase, which extends lifespan, promotes fat accumulation in worms with a specific enrichment of mono-unsaturated fatty acids (MUFAs). This fat metabolism switch in H3K4me3 methyltransferase-deficient worms is mediated at least in part by the downregulation of germline targets, including S6 kinase, and by the activation of an intestinal transcriptional network that upregulates delta-9 fatty acid desaturases. Notably, the accumulation of MUFAs is necessary for the lifespan extension of H3K4me3 methyltransferase-deficient worms, and dietary MUFAs are sufficient to extend lifespan. Given the conservation of lipid metabolism, dietary or endogenous MUFAs could extend lifespan and healthspan in other species, including mammals.

Chromatin state, which encompasses post-translational modification of histone proteins, integrates environmental signals to influence gene expression and downstream cellular processes. Enzymes that deposit or remove histone modifications regulate lifespan in several species^{1,2}. For example, methyltransferase or demethylase complexes that modify H3K4me3, H3K27me3 and H3K36me3 modulate lifespan in yeast, worms and flies^{1,2}. Several chromatin-modifying enzymes require intracellular metabolites as co-factors (for example, NAD+, α -ketoglutarate or S-adenosyl methionine), rendering their activity highly sensitive to metabolic states^{1,2}. A subset of these chromatin-modifying enzymes, such as deacetylases, in turn influence metabolism^{1,2}. However, the impact of methyltransferases and demethylases on metabolism remains unclear.

Fat metabolism plays an important role in many physiological and pathological processes. It influences long-term energy storage, intercellular and intracellular signalling and membrane homeostasis. In humans, excessive fat storage in the form of triglycerides (TAGs) is associated with diseases such as atherosclerosis and type 2 diabetes³. On the other hand, studies in both invertebrates and mammals have suggested that specific alterations in fat profiles, and even elevated fat storage, can be associated with longevity⁴. Whether fat metabolism links chromatin modifiers to lifespan extension is unknown.

H3K4me3 modifiers alter fat metabolism

The COMPASS chromatin complex, which catalyses trimethylation of lysine 4 on histone H3 (H3K4me3)⁵, regulates lifespan in *C. elegans*⁶. The H3K4me3 global landscape is remodelled during ageing and in cellular senescence in mammals^{1,2}. However, the mechanism by which H3K4me3 modifiers influence longevity remains unclear. To determine whether the COMPASS H3K4me3 methyltransferase complex affects fat metabolism, we used Oil-Red-O (ORO) staining. In fixed tissues, ORO stains neutral lipids such as TAGs⁷. Worms with a deficiency in the COMPASS H3K4me3 methyltransferase SET-2 (homologue of mammalian SET1), which are long-lived⁶ (Supplementary Table 1), exhibited increased ORO staining (Fig. 1a). Conversely, worms deficient for the H3K4me3 demethylase RBR-2 (homologue of mammalian JARID1), which are slightly short-lived⁶ (Supplementary Table 1), displayed decreased ORO staining (Fig. 1a). RNA interference (RNAi)-mediated knockdown of *ash-2* (homologue of mammalian *ASH2L*) and *wdr-5.1* (homologue of mammalian *WDR5*), which are shared components between the COMPASS and Trithorax-related H3K4me3 complexes⁵, and of *set-2*, a specific component of the COMPASS H3K4me3 complex⁵, also extended lifespan⁶ (Supplementary Table 1) and increased ORO staining (Fig. 1b and Extended Data Fig. 1a). By contrast, knockdown of *set-16* (homologue of mammalian *MLL*) and *utx-1* (homologue of mammalian *UTX*, also known as *KDM6A*), specific components of the Trithorax-related H3K4me3 complex⁵, did not significantly affect ORO levels (Fig. 1b). Thus, the COMPASS H3K4me3 methyltransferase complex specifically influences fat levels.

Fat accumulated most prominently in the intestine in *ash-2* knockdown worms (Fig. 1c), and this was observed at the initiation of reproduction, during reproduction and after reproduction (Fig. 1c and Extended Data Fig. 1b, c). The intestinal fat accumulation in *ash-2* knockdown worms was also observed using Nile red, which stains neutral lipids in fixed tissues⁸ (Extended Data Fig. 1d–f). Furthermore, gas chromatography coupled to mass spectrometry (GC–MS) showed that *ash-2* knockdown worms had significantly higher levels of TAGs than control worms, but similar levels of phospholipids (Fig. 1d). Fat accumulation in *ash-2* knockdown worms is unlikely to be caused by a shift in fat resources resulting from reduced fertility, because *ash-2* knockdown worms had a similar number of progeny to wild-type worms⁶ (Extended Data Fig. 1g). Thus, H3K4me3 methyltransferase-deficient worms, which are long-lived and fertile, accumulate intestinal fat in the form of TAGs.

H3K4me3 modifiers act mostly in the germline to extend lifespan⁶, but fat accumulation occurs in the intestine (Fig. 1c). To determine whether H3K4me3 modifiers act in the germline or intestine to regulate fat accumulation, we performed tissue-selective RNAi knockdown and transgenic rescue experiments. Knockdown of *ash-2* in the intestine

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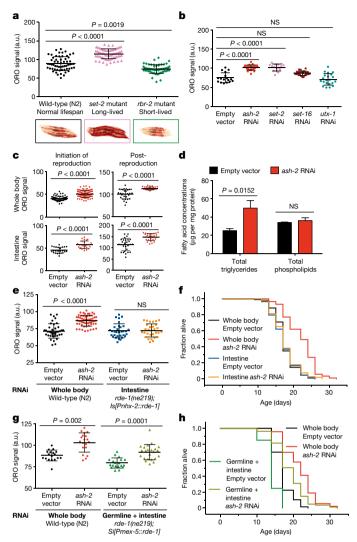


Figure 1 | H3K4me3 methyltransferase deficiency in the germline leads to fat accumulation in the intestine. a, ORO staining images and quantification. Mean \pm s.d., $n \ge 55$ worms per condition. b, As in a, $n \ge 16$ worms per condition. c, As in a, $n \ge 29$ worms (upper panels) or $n \ge 25$ intestinal cells (lower panels) per condition. d, GC–MS quantification. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. e, As in a, $n \ge 26$ worms per condition. f, *ash-2* RNAi extends lifespan in wild-type (P < 0.0001, log-rank) but not in *rde-1(ne219)*; *Is*[*Pnhx-2::rde-1*] worms. g, As in a, $n \ge 18$ worms per condition. h, *ash-2* RNAi extends lifespan in both wild-type and *rde-1(ne219)*; *Si*[*Pmex-5::rde-1*] worms (P < 0.0001, log-rank). b, c, e–h, Representative of two experiments. *P* values: a, b, e, g, Kruskal-Wallis with Dunn's correction; c, d, two-tailed Mann-Whitney.

(*rde-1(ne219*);*Is*[*Pnhx-2::rde-1*]⁹) did not promote fat accumulation or lifespan extension (Fig. 1e, f and Extended Data Fig. 1h, i). Likewise, knockdown of *set-2* in the intestine did not increase fat accumulation (Extended Data Fig. 1j), and overexpression of *set-2* in the intestine of *set-2(ok952)* mutants did not abrogate the elevated fat levels found in *set-2(ok952)* mutants (Extended Data Fig. 1k–m). These results suggest that H3K4me3 modifiers act mostly outside of the intestine to influence fat accumulation. As strains that allow germline-specific RNAi knockdown do not currently exist in *C. elegans*, we used the *rde-1(ne219);Si[Pmex-5::rde-1]* strain, which restricts RNAi to the germline and intestine ¹⁰. Interestingly, *ash-2* knockdown in the germline and extended Data Fig. 1i). Similarly, *set-2* knockdown in the germline and intestine and intestine also increased fat levels

(Extended Data Fig. 1j). *ash-2* knockdown in *rrf-1(pk1417)* mutants, in which RNAi is restricted to the germline, intestine and some hypodermal seam cells¹¹, also increased fat levels and extended lifespan (Extended Data Fig. 1n and Supplementary Table 1). Thus, the COMPASS H3K4me3 methyltransferase complex acts mostly in the germline to regulate intestinal fat accumulation and lifespan, implying that there is communication between the germline and intestine.

A switch to MUFA metabolism

Like mammals, worms have different categories of fatty acids: saturated fatty acids (SFAs), mono-unsaturated fatty acids (MUFAs) and polyunsaturated fatty acids (PUFAs) (Fig. 2a). To identify specific lipid species associated with longevity in H3K4me3 methyltransferase-deficient worms, we profiled long-chain fatty acids by GC–MS. At all ages tested, *ash-2* knockdown increased MUFAs, while SFAs and PUFAs remained mostly unchanged (Fig. 2b). The MUFAs palmitoleic acid and *cis*vaccenic acid were significantly elevated at all ages, whereas the MUFA oleic acid was raised after reproduction in *ash-2* knockdown worms and in worms deficient for *set-2* (Fig. 2b, c). By contrast, MUFAs remained largely unaffected in worms deficient for *set-16* and *utx-1* (Extended Data Fig. 2a). Thus, the COMPASS H3K4me3 methyltransferase complex specifically influences MUFA metabolism.

To determine whether this switch to MUFA accumulation arises from changes in fat synthesis, storage or breakdown, we examined the expression of the specific enzymes that control these steps (Fig. 2a). Knockdown of ash-2 strongly increased the expression of the delta-9 fatty acid desaturase genes fat-5 and fat-7 (Fig. 2d). FAT-5, a palmitoyl-coenzyme A (CoA) desaturase, preferentially converts palmitic acid into the MUFA palmitoleic acid, whereas FAT-7 and FAT-6, two stearoyl-CoA desaturases, convert stearic acid into the MUFA oleic acid¹² (Fig. 2a). The expression of fat-5 and fat-7 was also upregulated in set-2 mutant or knockdown worms (Extended Data Fig. 2b). By contrast, the expression of fat-2, which catalyses the first step in converting MUFAs to PUFAs¹³, and the expression of other fat metabolism enzymes (for example, lipases), was not consistently affected by ash-2 knockdown (Fig. 2d). The expression of fat-5 and fat-7 was elevated in the intestine but not in the germline of ash-2 knockdown worms (Fig. 2e), and *ash-2* acts mostly in the germline to promote this intestinal fat-5 and fat-7 elevation (Fig. 2e). Transgenic worms expressing green fluorescent protein (GFP) translational reporters for FAT-5 or FAT-7 also showed increased GFP fluorescence in the intestine upon ash-2 knockdown (Fig. 2f). Thus, a deficiency in germline H3K4me3 methyltransferase leads to the upregulation of delta-9 desaturases in the intestine, suggesting that MUFA accumulation occurs at the level of synthesis.

We next examined the role of delta-9 desaturases in the accumulation of MUFA in H3K4me3 methyltransferase-deficient worms. Knockdown of *fat-7* in *ash-2*-deficient worms specifically decreased oleic acid levels at the initiation of reproduction and reduced all three MUFAs (oleic, palmitoleic and *cis*-vaccenic acid) after reproduction (Fig. 2g, h and Extended Data Fig. 2c). By contrast, *fat-5* knockdown in *ash-2*-deficient worms increased palmitoleic and *cis*-vaccenic acid (Extended Data Fig. 2d), probably owing to compensatory upregulation of *fat-7* (Extended Data Fig. 2e). These results suggest that the MUFA switch in H3K4me3 methyltransferase-deficient worms is driven by FAT-7 upregulation in the intestine.

Germline targets of H3K4me3 modifiers

The H3K4me3 methyltransferase complex acts mostly in the germline to trigger intestinal upregulation of delta-9 desaturases. To understand the germline-to-intestine communication better, we identified germline targets of the H3K4me3 methyltransferase complex. We generated an RNA sequencing (RNA-seq) dataset using microdissected germlines and intestines from *ash-2* knockdown worms and analysed this dataset (Extended Data Fig. 3a, b and Supplementary

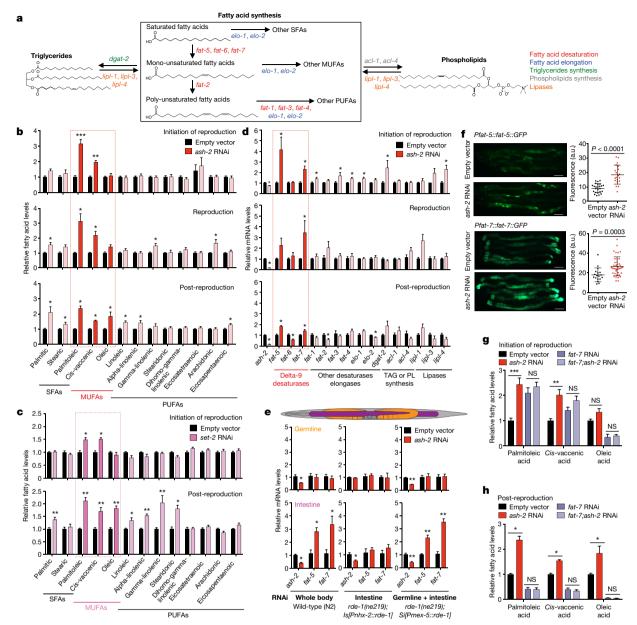


Figure 2 | H3K4me3 methyltransferase deficiency results in elevated MUFAs and upregulation of delta-9 desaturases. a, Fat metabolism pathways with genes used in RT–qPCR. b, c, GC–MS quantification of fatty acids. Mean \pm s.e.m. of two or three independent experiments, each with three biological replicates. d, RT–qPCR. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. e, RT–qPCR. Mean \pm s.e.m. of two independent experiments, each with

two or three biological replicates. **f**, Images and quantification of GFP fluorescence. Mean \pm s.d., $n \ge 18$ worms per condition. Representative of two (FAT-5) or three (FAT-7) experiments. Scale bars, $100 \,\mu$ m. **g**, **h**, GC–MS quantification of MUFAs. Mean \pm s.e.m. of three (**g**) or two (**h**) independent experiments, each with two or three biological replicates. *P* values: two-tailed Mann–Whitney (with Benjamini–Hochberg correction for >2 conditions). **P* < 0.05, ***P* < 0.01, ****P* < 0.001.

Table 2–5), as well as other large-scale datasets^{6,14,15}. Because H3K4me3 methyltransferase complexes generally activate gene expression⁵, we selected targets that were downregulated in H3K4me3 methyltransferase-deficient worms (Extended Data Table 1). Using quantitative PCR with reverse transcription (RT–qPCR), we confirmed that *ash-2* knockdown led to the significant downregulation of four candidate genes in the germline (*rsks-1, tbh-1, nhr-10* and *unc-132*), one candidate in both the germline and intestine (*F21A3.11*) and two candidates in the intestine (*asm-2* and *dod-23*) (Fig. 3a and Extended Data Fig. 3c).

We next assessed the functional involvement of these candidates in fat metabolism. Knockdown of *rsks-1*, *tbh-1*, *F21A3.11* or *unc-132* led to increased ORO staining (Fig. 3b and Extended Data Fig. 3d–g). Among these four targets, *rsks-1* was the only one that mostly functioned in

the germline to regulate fat accumulation (Extended Data Fig. 3d–g). RSKS-1 (homologue of mammalian S6 kinase) is a key conserved substrate of mechanistic target of rapamycin (mTOR) complex 1¹⁶. A deficiency in RSKS-1 extends lifespan in *C. elegans* and mice¹⁷ and promotes fat accumulation in *C. elegans*¹⁸. We investigated whether *rsks-1* could mediate the fat metabolism switch of H3K4me3 methyltransferase-deficient worms. Knockdown or mutation of *rsks-1* led to increased fat accumulation, MUFA levels and *fat-7* expression (Fig. 3b–d and Extended Data 3h–j). Notably, *ash-2* knockdown could not further increase fat accumulation and *fat-7* expression in *rsks-1*-deficient worms (Fig. 3d and Extended Data 3i, j). Likewise, knockdown of the RSKS-1 upstream regulator *let-363* (homologue of mammalian *mTOR*) also led to increased fat accumulation, MUFA levels and *fat-7* expression (Extended Data Fig. 3k–n), and *ash-2* knockdown

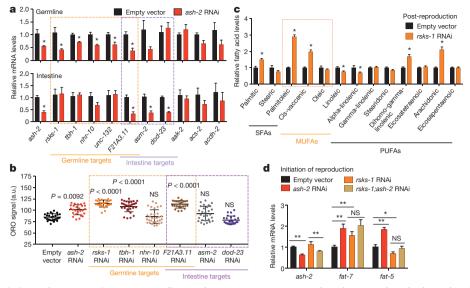


Figure 3 | H3K4me3 methyltransferase germline targets influence fat metabolism. a, RT-qPCR. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. b, ORO quantification. Mean \pm s.d., $n \ge 23$ worms per condition. Representative of two experiments. c, GC-MS profiles of fatty acids. Mean \pm s.e.m. of two independent

did not further increase *fat-7* expression in these worms (Extended Data Fig. 3k). Deficiencies in RSKS-1 or its upstream regulators (*let-363* or *daf-15*, homologue of mammalian *Raptor*) resulted in lifespan extension, and the ability of *ash-2* or *set-2* deficiency to extend lifespan was diminished in the context of *rsks-1* deficiency (Extended Data Fig. 30, p). Collectively, these data suggest that RSKS-1 is a germline target of the H3K4me3 methyltransferase complex that mediates at least part of the intestinal upregulation of delta-9 desaturases leading to MUFA accumulation.

MUFAs mediate longevity by H3K4me3 modifiers

To examine the role of MUFAs in the modulation of longevity by H3K4me3 modifiers, we first investigated the molecular mechanisms that underlie MUFA accumulation in worms with H3K4me3 methyltransferase deficiency. Delta-9 desaturases are known targets of a highly conserved intestinal transcriptional network that involves the transcription factor SBP-1 (homologue of mammalian SREBP)¹⁹ and the Mediator complex subunit MDT-15 (homologue of mammalian MED15)²⁰, as well as other transcription factors, such as NHR-49 (functional homologue of mammalian PPAR α)²¹ and NHR-80 (homologue of mammalian HNF4)²² (Fig. 4a). Knockdown of ash-2 and set-2, as well as mutation of set-2, increased nuclear accumulation of SBP-1 in intestinal cells (Fig. 4b and Extended Data Fig. 4a), and this accumulation was reduced in rsks-1-deficient worms (Extended Data Fig. 4b-d). We next tested whether SBP-1 and MDT-15 are necessary for upregulation of delta-9 desaturases and accumulation of MUFAs in H3K4me3 methyltransferase-deficient worms. Knockdown of sbp-1 or mdt-15 decreased the basal levels of fat-5 and fat-7 mRNA and reduced their upregulation by ash-2 knockdown (Fig. 4c and Extended Data Fig. 4e). Knockdown of sbp-1 or mdt-15 also blocked the elevation of MUFAs (palmitoleic, cis-vaccenic and oleic acid) in ash-2-deficient worms after reproduction (Fig. 4d and Extended Data Fig. 4f). Thus, SBP-1 and MDT-15 are critical for delta-9 desaturase upregulation and MUFA accumulation in H3K4me3 methyltransferase-deficient worms.

We next used this molecular knowledge to assess whether the MUFA metabolic switch causes the lifespan extension of H3K4me3 methyltransferase-deficient worms. Interestingly, knockdown or mutation of *sbp-1* or *mdt-15* abolished fat accumulation and lifespan extension in *ash-2* knockdown worms (Fig. 4e, f, Extended Data

experiments, each with two or three biological replicates. **d**, RT–qPCR. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. *P* values: **b**, Kruskal–Wallis with Dunn's correction; **a**, **c**, **d**, two-tailed Mann–Whitney with Benjamini–Hochberg correction. **P* < 0.05, ***P* < 0.01.

Fig. 4g, h and Supplementary Table 1). Likewise, sbp-1 knockdown abrogated the longevity of set-2 mutant worms (Supplementary Table 1). By contrast, nhr-49 mutation only partially diminished the ability of ash-2 deficiency to promote fat accumulation and lifespan extension (Extended Data Fig. 4i, j) and nhr-80 mutation had no effect (Extended Data Fig. 4k, l). We also tested whether delta-9 desaturases are required for the longevity of H3K4me3 methyltransferase-deficient worms. Knockdown of fat-7 or double mutations in fat-6 and fat-7, which produce a strong deficiency in oleic acid synthesis²³, blocked the fat accumulation and longevity of ash-2-deficient worms (Fig. 4g, h and Extended Data Fig. 5a, b). By contrast, single hypomorphic mutations in fat-7 or fat-6 (or double mutations in fat-5 and fat-7 or fat-5 and fat-6), which produce only mild deficiencies in MUFA synthesis^{22,23}, did not abolish the longevity of ash-2-knockdown worms (Extended Data Fig. 5c-f). Thus, a conserved pathway involving SBP-1, MDT-15 and the downstream delta-9 desaturase FAT-7 is critical for the longevity of H3K4me3 methyltransferase-deficient worms.

To assess the specific role of MUFAs in H3K4me3 methyltransferasedeficient worms, we supplemented oleic acid in the diet (Extended Data Fig. 5g, h). Dietary oleic acid largely restored the high-fat phenotype and lifespan extension in *ash-2* and *fat-7* double knockdown worms (Fig. 4g, h and Extended Data Fig. 5i, j). Similarly, dietary oleic acid restored high fat levels in *ash-2* and *sbp-1* double knockdown worms (Extended Data Fig. 5k). Furthermore, knockdown of *fat-2*, which normally converts oleic acid into downstream PUFA¹³, increased fat accumulation and lifespan extension, and *ash-2* deficiency did not further increase fat levels or extend lifespan in *fat-2* knockdown worms (Extended Data Fig. 5l–o). Thus, the MUFA oleic acid plays a key role in the longevity of H3K4me3 methyltransferase-deficient worms.

Dietary MUFAs extend lifespan

The role of oleic acid in lifespan regulation in the context of H3K4me3 methyltransferase deficiency led us to examine the importance of MUFAs and their downstream PUFAs in the regulation of lifespan under physiological conditions (Fig. 5a). Dietary supplementation of individual MUFAs (oleic, palmitoleic or *cis*-vaccenic acid; Fig. 5b, d and Extended Data Fig. 6a) was sufficient to extend lifespan (Fig. 5c, e, f and Extended Data Fig. 6b). By contrast, dietary supplementation of the PUFAs linoleic and alpha-linolenic acid (Extended Data Fig. 6c, e) did not significantly affect lifespan (Extended Data

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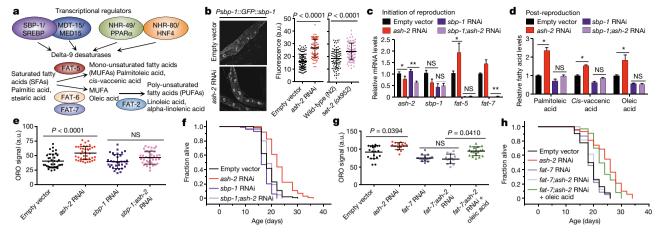
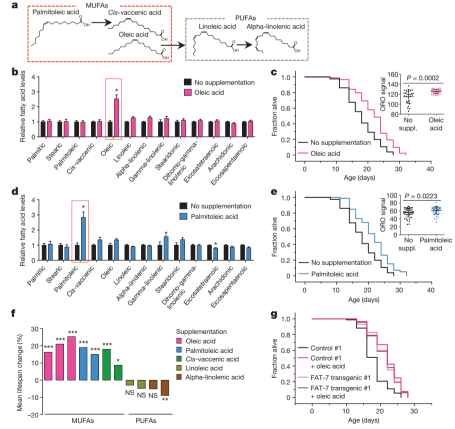


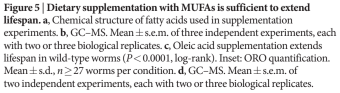
Figure 4 | Accumulation of MUFAs is critical for the longevity of H3K4me3 methyltransferase-deficient worms. a, Conserved transcriptional network regulating delta-9 desaturase genes. b, Images and quantification of SBP-1 nuclear accumulation. Mean \pm s.d. of two independent experiments, each with n = 4-6 nuclei per worm and ≥ 8 worms per condition. c, RT–qPCR. Mean \pm s.e.m. of three independent experiments, each with three biological replicates. d, GC–MS. Mean \pm s.e.m. of two independent experiments, each with two or three biological replicates. e, ORO quantification. Mean \pm s.d.,

Fig. 6d, f and Fig. 5f). All MUFAs and PUFAs tested increased total fat accumulation, except for the PUFA alpha-linolenic acid (Fig. 5c, e and Extended Data Fig. 6b, d, f). Furthermore, overexpression

 $n \ge 32$ worms per condition. **f**, Lifespan extension by *ash-2* RNAi is reduced in *sbp-1* RNAi (9.78%) worms compared to control worms (21.90%) (P = 0.0003, two-way ANOVA). **g**, As in **e**, $n \ge 13$ worms per condition. **h**, Dietary oleic acid extends lifespan in the context of *fat-7* and *ash-2* double RNAi (P < 0.0001, log-rank). **g**, **h**, Representative of two experiments. *P* values: **b**, two-tailed Mann-Whitney; **e**, **g**, Kruskal-Wallis with Dunn's correction; **c**, **d**, two-tailed Mann-Whitney with Benjamini–Hochberg correction. *P < 0.05, **P < 0.01.

of the oleic acid-synthesizing enzyme FAT-7 in the intestine increased fat accumulation and extended lifespan (Fig. 5g and Extended Data Fig. 6g–i). Oleic acid supplementation did not further extend the long





e, Palmitoleic acid supplementation extends lifespan in wild-type worms (P < 0.0001, log-rank). Inset: as in **c**, $n \ge 45$ worms per condition. **f**, Mean lifespan changes in independent supplementation experiments (P values: log-rank). **g**, FAT-7 overexpression extends lifespan (P < 0.0001, log-rank), and this lifespan is not extended further by dietary oleic acid. P values: two-tailed Mann-Whitney (Benjamini–Hochberg correction for more than one comparison). *P < 0.05, **P < 0.01, ***P < 0.001.

lifespan of FAT-7-overexpressing worms (Fig. 5g and Extended Data Fig. 6i). Collectively, these results suggest that dietary supplementation and endogenous accumulation of MUFAs, but not downstream PUFAs, are beneficial for lifespan.

Discussion

Our study shows that histone methylation modifiers influence MUFA metabolism. We propose a model in which H3K4me3 methyltransferase deficiency causes the downregulation of specific targets (for example, RSKS-1) in the germline, which in turn triggers a germline-to-intestine signal leading to activation of SBP-1, expression of delta-9 desaturases and accumulation of MUFAs in the intestine (Extended Data Fig. 6j). The nature of this germline-to-intestine signal is unknown, but it could be a diffusible lipid signal ('lipokine'), a signal from the maturing eggs or an alteration in levels of lipid-binding proteins. Deficiency in the H3K4me3 methyltransferase complex might also affect fat metabolism by altering the methyl pool available for other enzymatic reactions or by methylating non-histone substrates that have not yet been identified.

Our work also shows that dietary MUFAs, but not their downstream PUFAs, positively affect lifespan. Previous studies have tested supplementation of individual fatty acids on lifespan^{24–28} and implicated other PUFAs in lifespan regulation^{24,26}. Dietary oleic acid did not extend the lifespan of wild-type worms in previous studies^{25,27,28}, although these studies used different oleic acid supplementation protocols. MUFAs could contribute to longevity by promoting membrane fluidity, minimizing oxidative stress, enhancing energy storage or activating signal-ling pathways. The beneficial effects of MUFAs are exciting because of their dietary availability (for example, oleic acid in olives) and microbiotic sources (for example, palmitoleic acid in *E. coli*)²⁹. In humans, diets rich in MUFAs have been associated with a decreased risk of cardiovascular disease and diabetes³⁰. The general conservation of lipid metabolism between worms and mammals suggests that the benefits of MUFAs on healthspan and lifespan could be conserved in other species.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Supplementary Information is available in the online version of the paper.

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Author Contributions S.H. conceived the study under the guidance of A.B. S.H. performed all the experiments except those specified below. E.A.S. planned and performed Nile Red and FAT-5/FAT-7 reporter imaging, time-course RT–qPCR and one oleic acid supplementation lifespan experiment. C.G.S.-G. planned and performed SBP-1 localization experiments and generated *set-2* transgenic lines under the guidance of W.B.M. K.H. analysed the RNA-seq data. S.H. wrote the paper with the help of A.B. and E.A.S. C.G.S.-G., K.H. and W.B.M. provided feedback.

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METHODS

No statistical methods were used to predetermine sample size. The experiments were not randomized and the investigators were not blinded to allocation during experiments and outcome assessment except where indicated.

Worm strains. All strains were maintained on standard nematode growth medium (NGM) plates with Escherichia coli (OP50-1, streptomycin-resistant). Unless otherwise noted, worms were grown on the RNAi strain HT115 (DE3) for all experiments. The wild-type N2 strain (ABR) was provided by M.-W. Tan. The NL2098 rrf-1(pk1417) strain, which restricts RNAi to the germline, intestine and partially the hypodermal seam cells^{11,31}, was provided by A. Fire. The germline and intestine RNAi strain AMJ345 rde-1(ne219);jamSi2[Pmex-5::rde-1(+)] and corresponding wild-type N2 (AMJ) were provided by A. Jose¹⁰. The intestinespecific RNAi strain VP303 rde-1(ne219);kbIs7[nhx-2p::rde-1 + rol-6(su1006)] was obtained from the Caenorhabditis Genetics Center (CGC). Other strains obtained from the CGC were as listed: RB1025 set-2(ok952), ZR1 rbr-2(tm1231), DA1116 eat-2(ad1116), XA7702 mdt-15(tm2182), STE68 nhr-49(nr2041), BX165 nhr-80(tm1011), BX153 fat-7(wa36), BX106 fat-6(tm331), BX160 fat-7(wa36); fat-5(tm420), BX110 fat-6(tm331);fat-5(tm420), BX156 fat-6(tm331);fat-7(wa36), RB1206 rsks-1(ok1255), and fluorescence translational fusion reporters BX150 lin-15B&lin-15A(n765);waEx18[fat-5::GFP + lin15(+)], BX113 lin-15B&lin-15A(n765);waEx15[fat-7::GFP + lin-15(+)] and CE548 sbp-1(ep79);epEx141 [sbp-1::GFP::SBP-1 + rol-6(su1006)]. Mutants set-2(ok952), rbr-2(tm1213), eat-2(ad1116), and transgenic strain CE548 were outcrossed to our laboratory N2 (ABR) at least six times. During backcrossing, the sbp-1(ep79) mutation in CE548 was removed to generate a line expressing a translational fusion between GFP and SBP-1 in a wild-type background. See Supplementary Table 6 for a list of strains. All experiments were performed using hermaphrodites.

RNA interference. RNAi was performed as described⁶. HT115 (DE3) bacteria were transformed with vectors expressing double-stranded (ds)RNA targeting the following genes of interest: *ash-2, set-2, wdr-5.1, set-16, utx-1, jmjd-3.1, fat-2, fat-5, fat-7, mdt-15, sbp-1, tbh-1, nhr-10, F21A3.11, dod-23 (Ahringer Library,* A. Fire), *let-363* (X. Long), *daf-15* (A. Hsu), *rsks-1, asm-2* and *unc-132* (Vidal Library, Dharmacon). All vectors were confirmed by sequencing. Empty vector bacteria, containing the RNAi plasmid without an RNAi insert, served as a control for all RNAi experiments. RNAi culture was concentrated 30-fold, and stored at 4°C for no more than 2 weeks. For double RNAi treatments, two cultures were mixed in a 1:1 ratio by volume. To obtain a synchronized worm population, egglay was performed on appropriate RNAi plates for 1–4h. All RNAi treatments began at hatching, except for ORO and lifespan experiments using *sbp-1, mdt-15, fat-2, let-363* or *daf-15* RNAi, which began at the initiation of reproduction (day 2.5 of life) to minimize the effects on development.

Lifespan assays. Worm lifespan assays were performed at 20 °C as described³². Kaplan–Meir lifespan curves were generated using GraphPad Prism, and statistical results were obtained by log-rank analysis using JMP software. Two-way ANOVA was used to assess lifespan interactions between conditions. Lifespan assays that were blinded or repeated by an independent investigator are indicated in Supplementary Table 1. Representative Kaplan–Meir lifespan curves are shown in the figures and all statistics are included in Supplementary Table 1.

Oil Red O and Nile Red staining and quantification. ORO staining of fixed worms was conducted as described⁷, except that 2% paraformaldehyde in PBS was used to fix worm tissues. ORO staining of dissected intestines was performed for 1 h. Nile Red staining of fixed worms was performed as described³³, except that staining was conducted in 1.5-ml Eppendorf tubes. ORO- or Nile Red-stained worms were mounted onto 2% agar pads, and imaged at $10 \times$ magnification using a Zeiss AxioSkop 2 Plus. Images were acquired using an AxioCam MRc camera with AxioVision 4.7 software and saved as TIF files. The same exposure settings were used across all conditions within each experiment. Using FIJI image processing software, raw images were subtracted from background, converted to greyscale, inverted and thresholded to outline the worm body. The same threshold values were used across all conditions within each experiment. 'Analyze Particles' was used to measure mean ORO intensity per worm. To quantify anterior intestinal cells of whole worms or extruded intestines of dissected worms, specific regions in the raw images were selected using the oval brush tools (intestinal cells) or the segmented line tool (extruded intestines), saved to the Region of Interest (ROI) manager, processed as described above, and quantified for mean ORO intensity per selected region. Mean intensity values in arbitrary units (a.u.) were graphed using Graphpad Prism, and statistical significance was determined using a twotailed Mann-Whitney test (two conditions) or the Kruskal-Wallis test with Dunn's correction for multiple comparisons (more than two conditions). For figures containing straightened whole worms or intestines, unmodified images were selected and straightened, and representative worms or intestines from each condition were concatenated into montages.

GFP quantification. For quantification of FAT-5 and FAT-7 expression, transgenic lines expressing a translational fusion between FAT-5 and GFP or FAT-7 and GFP were used. Worms at active reproduction (day 3 of life) were anaesthetized in M9 (22 mM KH2PO4, 42 mM Na2HPO4, 86 mM NaCl and 1mM MgSO4, dissolved in water) containing 100µM levamisole and mounted on 2% agar pads. All images were acquired on Zeiss AxioSkop 2 Plus using an AxioCam MRc camera with AxioVision 4.7 software with identical exposure settings for the same experiment. Mean fluorescence intensity in arbitrary units (a.u.) of the whole worm was quantified in FIJI as described above. These experiments were conducted in a blinded manner. For visualization of intestinal nuclear SBP-1 expression, a transgenic line expressing a translational fusion between GFP and SBP-1 was used. Worms at active reproduction (day 3 of life) were mounted on 2% agarose pads and anaesthetized with 20 mM tetramisole in M9. Images were acquired using a Zeiss Axio Imager.M2 microscope equipped with an ApoTome.2 system and an AxioCam MRc camera using identical exposure settings for the same experiment. Images showing the maximum number of defined intestine nuclei were selected in each animal and saved as TIF files. The mean fluorescence intensity in arbitrary units (a.u.) of each nucleus was quantified using the oval brush tool in FIJI as described above.

Worm tissue micro-dissection. Synchronized worms grown on appropriate RNAi bacteria were collected at specified stages of life. At least 20 worms from each condition were washed three times with M9 buffer and deposited onto a glass slide in drops of M9 (one worm per drop) with a glass Pasteur pipette. Under the dissection light microscope, individual worms were decapitated using 22.5° knives (Surgistar) to extrude their anterior intestines and full germlines (including developing oocytes). For ORO or Nile Red staining, reproductive worms (day 5 of life) with extruded intestines were used for staining. For RT-qPCR or RNA-seq, fully detached intestines and germlines from worms during active reproduction (day 3 of life) or at initiation of reproduction (day 2.5 of life) were harvested. To isolate the full germline containing maturing oocytes, an incision was made before the spermatheca to minimize sperm contamination. Dissected germlines were immediately collected in 500 µl Trizol LS reagent (Invitrogen) using a glass Pasteur pipette whose tip was pulled under the flame to thin out the opening. To isolate the anterior intestine from the same worms, an incision was made near the 10 pairs of anterior intestinal cells. The anterior intestinal samples were collected in Trizol in separate tubes. All samples were stored at -80 °C until RNA extraction. Quantitative RT-PCR. For experiments using whole worms, at least 100 agesynchronized worms in biological triplicate for each condition were harvested, washed three times with M9 and resuspended in 500 µl Trizol. For experiments using dissected tissues, 20-40 germlines or intestines in biological triplicate for each condition were harvested. To extract total RNA, worm or tissue pellets in Trizol underwent six freeze-thaw cycles in a dry ice-ethanol bath. RNA was extracted according to the Trizol procedure, and resuspended in RNase- and DNase-free water. RNA from whole worms was quantified using the Qubit 2.0 fluorometer (Invitrogen), and at least 500 ng total RNA per condition was used for cDNA synthesis. RNA from dissected tissues was assessed for quality and quantified using the Agilent Bioanalyzer (RNA 6000 Pico), and at least 2 ng total RNA per condition was used for cDNA synthesis.

Total RNA was treated with DNase I Amp Grade (Invitrogen), then reversetranscribed using Oligo-dT primers (Invitrogen) and Superscript III reverse transcriptase (Invitrogen), based on the manufacturer's instructions. RT–qPCR was performed using diluted cDNA on an ABI7900HT or a Bio-Rad C1000 thermal cycler using iTaq master mix (Bio-Rad) containing a passive reference dye (ROX). Primers used in RT–qPCR were designed to span exon–exon junctions towards the 3' end of the gene and were used at a final concentration of 250 nM each. Primers are listed in Supplementary Table 7. Melt curves were examined to ensure primer specificity. Results were analysed using the standard $\Delta\Delta C_{\rm T}$ method. For each biological replicate, the mean $C_{\rm T}$ value of 3–4 technical replicates was analysed. *act-1* served as the reference gene in all analyses, and changes in mRNA levels relative to *act-1* were confirmed using *cdc-42, ama-1* or *pmp-3* as alternate reference genes in independent experiments (data not shown).

RNA-seq on micro-dissected tissues. For each of the five replicates, full germlines and anterior intestines were collected from 20 worms treated with *ash-2* RNAi or empty vector control. At the initiation of reproduction (day 2.5 of life), worms from each pair of *ash-2* RNAi and empty vector plates were dissected within the same hour. Tissue dissection and RNA extraction procedures were the same as described above and 20 full germlines or approximately 200 intestinal cells were used as the initial input for individual RNA-seq libraries. For cDNA synthesis, 150 pg (intestine) or 750 pg (germline) of extracted RNA was treated with RQ1 RNase-free DNase (Promega), which was later removed, and reverse transcribed using an oligo(dT)-primed, Clontech SMART-Seq v4 Ultra Low Input RNA Kit (634889) according to the manufacturer's instructions. Poly-A selected RNA

seq libraries were generated using Illumina Nextera XT DNA Library Prep Kit (FC-131-1024). All 20 libraries were processed together to minimize batch effects. The libraries were uniquely barcoded and multiplexed onto one lane, and 150 base pair paired-end reads were sequenced on NextSeq 500 (Illumina).

RNA-seq analysis. RNA-seq reads were aligned to the WBcel235 genome and gene read counts were calculated using STAR v2.5.1b³⁴. Low-coverage genes were filtered out: only genes with at least one read count per million mapped reads, in at least five samples, were included in the analysis, resulting in a dataset of 10,442 genes and 20 samples. More genes were detected in the germline samples (185 genes with a median count number of 0) than in the intestinal samples (682 genes with a median count number of 0), with 76% of the genes having a higher median read count in germline samples than in intestinal samples. Because of these tissue-specific differences, we did not perform normalization across samples (which assumes that most genes are not differentially expressed) and instead used FPKM values to compare germline and intestinal samples. The read counts were transformed to log₂(FPKM+1) values, which were used for principal component analysis (PCA) on the samples from both tissues together. A gene was determined to be germline-enriched if the lowest expression value $(log_2(FPKM+1))$ in the germline empty-vector samples was at least twofold higher than the highest expression value in the intestine empty-vector samples (5,494 germlineenriched genes). A gene was determined to be intestine-enriched if the lowest expression value (log₂(FPKM+1)) in the intestine empty-vector samples was at least twofold higher than the highest expression value in the germline empty-vector samples (1,418 intestine-enriched genes).

We then compared empty-vector and *ash-2* RNAi samples for each tissue separately. Low-coverage genes were filtered out: only genes with at least one read count per million mapped reads in at least five samples were included in the analysis. This resulted in a germline dataset of 9,073 genes and 10 samples, and an intestine dataset of 7,288 genes and 10 samples. Using DESeq2 v1.10.1³⁵, the read counts were normalized using the variance stabilizing transformation (VST) within each tissue and PCA was performed on the germline and intestine datasets. For each tissue, differentially expressed genes between empty-vector and *ash-2* RNAi were examined, accounting for a plate effect (GE ~ condition plus plate). Genewise *P* values were corrected for multiple hypothesis testing using Benjamini–Hochberg and FDR = 0.05 was used as a significance threshold.

Selection of ASH-2 candidate targets. ASH-2 candidate targets were selected using the following datasets: 1) our RNA-seq dataset, 2) microarrays on *ash-2* knockdown larval worms⁶ and *ash-2* mutant adult germlines¹⁴, and 3) ASH-2 chromatin immunoprecipitation followed by DNA microarray (ChIP–chip) analysis on worm embryos¹⁵. Predicted candidate expression in the germline or intestine for datasets 2 and 3 was based on published gene expression datasets^{36–38} (A. Rechtsteiner and S. Strome, personal communication) and pre-publication RNA-seq datasets (R. Waterston, personal communication). The following criteria were used to select potential targets: i) downregulation upon *ash-2* deficiency, as the H3K4me3 methyltransferase complex is generally associated with gene activation; ii) expression in the germline or intestine, as these are the two key tissues involved in this fat metabolic switch; and/or iii) previously identified role in lifespan or fat metabolism.

 fragment was generated by PCR amplification from pAS2.1 (a gift from F. Palladino) using the following primers (pSH9_*set-2.a*_F: 5'-ATCTATTACATATC TTATCTTTGAATTCAGATGTCCACACATGATATGAACCATCATCCT-3' and pSH9_*set-2.a*_R: 5'-AACTAGGTGAAAGTAGGATGAGACAGCCCCTCA ATTAAGATATCCACGACACGTCTTCGC-3'), and was corrected for a short missing sequence. The pIM26 fragment was amplified by PCR using the following primers (pSH9_pIM26_F: 5'-GCGAAGACGTGTCGGGAT ATCTAATTGAGGGGCTGTCCTCATCCTACTTTCACCTAGTT-3' and pSH9_ pIM26_R: 5'-ACATTTTTCGTCATCATAGTTCGGGAGTTTGGGGATCCTCT AGAGTCGACCTGCAGGCAT-3'). Three fragments were assembled via Gibson assembly³⁹.

All fragments generated via PCR amplification were sequence-verified in the final plasmid to ensure no mutations were introduced.

Generation of transgenic worm strains. The strain carrying extra-chromosomal arrays of [*Pges-1::fat-7:myo-3::mCherry*] was generated by injecting wild-type N2 (ABR) worms with pSH1 and a co-injection plasmid pCFJ104, with a muscle-specific *myo-3* promoter driving the mCherry gene (*Pmyo-3::mCherry*, made by E. Jorgensen, Addgene plasmid number 19328). The strain carrying extra-chromosomal arrays of [*Pges-1::set-2a cDNA;myo-3::mCherry*] was generated by injecting *set-2(ok952)* (ABR10) worms with pSH9 and *Pmyo-3::mCherry*. The injection mixture of both plasmids had a final concentration of 50 ng µl⁻¹. Control strains were generated by injecting wild-type N2 (ABR) worms with the co-injection marker (*Pmyo-3::mCherry*) at a final concentration of 50 ng µl⁻¹.

Worms containing extra-chromosomal arrays were selected on the basis of mCherry fluorescence in the body wall muscles. In transgenic worms also carrying a GFP reporter, GFP fluorescence was used as a primary marker for strain maintenance. Overexpression of *fat-7* in wild-type transgenic worms was confirmed by RT–qPCR with primers designed to detect *fat-7* mRNA. Overexpression of *set-2.a* in *set-2(ok952)* mutant transgenic worms was confirmed by RT–qPCR with primers designed to detect transgenic (wild-type) but not mutant *set-2.a* mRNA. Functional validation of *set-2* intestine rescue transgenic lines was conducted by RT–qPCR analysis of two intestinal tawrgets of COMPASS H3K4me3 complex, *dod-23* and *asm-2*, which are normally downregulated in *set-2(ok952)* mutants. Primer sequences are listed in Supplementary Table 7.

GC-MS quantification of fatty acids. Quantification of total triglycerides and total phospholipids levels, and profiling of long-chain fatty acids, were performed as described³³ with the following modifications. For each condition, 500-1,000 age-synchronized animals were used. For GC-MS, an Agilent 7890A gas chromatograph equipped with an HP-5MS column ($30 \text{ m} \times 0.25 \text{ mm}$ $I.D. \times 0.25\,\mu\text{m},$ Agilent) using helium as a carrier gas was used, and $1\,\mu\text{l}$ of each sample was injected at a 10:1 split ratio. The initial oven temperature was set to 150 °C, increased to 220 °C at 20 °C per min, increased to 230 °C at 2 °C per min, increased to 320 °C at 50 °C per min, and held for 2 min at this temperature. Fatty acid abundance was expressed as a ratio (µg per mg) of fatty acid concentration (µg per ml) to protein concentration (mg per ml) for each sample. In all GC-MS figure panels involving comparisons to the empty vector control (unless otherwise mentioned), raw fatty acid concentration (µg per mg) of each condition was normalized to the raw fatty acid concentration (µg per mg) of the empty vector control. The final ratio is expressed as relative fatty acid levels. Raw GC-MS data for all figure panels are provided in Supplementary Table 8.

Fertility assessment. The fertility of N2 worms treated with *ash-2* RNAi or the empty vector control was assayed at 20 °C. After a 1-h egglay, eggs were maintained at 20 °C until worms reached the initiation of reproduction (2.5 days of life). Each of the 30 reproductive worms from each condition was transferred onto an individual 3.5-cm plate seeded with corresponding bacteria. Individual worms were transferred onto a new plate every day, and the number of fertilized eggs (eggs with visible shell) and unfertilized oocytes (eggs without visible shell) were assessed on the old plate. The old plate with eggs was incubated at 20 °C for an additional day, and the number of hatched progeny (live brood size) was counted. Fertility was assessed daily until no more hatched worms, fertilized eggs or unfertilized ocytes were seen on each plate. This experiment was conducted in a blinded manner.

Fatty acid supplementation. The fatty acid supplementation protocol was adapted from a previous study⁴⁰ with several modifications. To facilitate fatty acid dissolution, the detergent Tergitol (type NP-40, Sigma-Aldrich) was added to a final concentration of 0.001% in liquid NGM agar medium for both non-supplemented and supplemented plates before autoclaving. NGM plates for RNAi experiments were prepared as described⁶, and HT115 bacteria were used for all supplementation experiments. A final concentration ranging from 0.1 mM to 4 mM of oleic acid was used during the initial experimental optimization, and a final concentration of 0.8 mM was used for all fatty acids in subsequent supplementation experiments. HT115 bacteria expressing the empty vector or the appropriate RNAi were seeded onto plates containing the respective fatty acids at room temperature

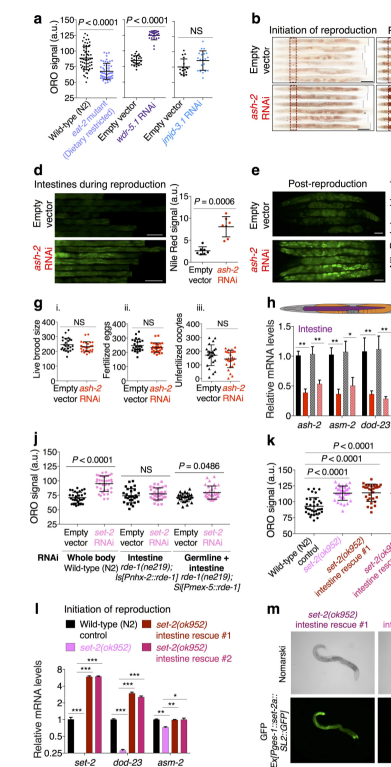
for 24–48 h before the addition of worms to ensure incorporation of fatty acids into feeding bacteria. Successful incorporation of supplemented fatty acids into the HT115 bacteria and into the worms was confirmed by GC–MS.

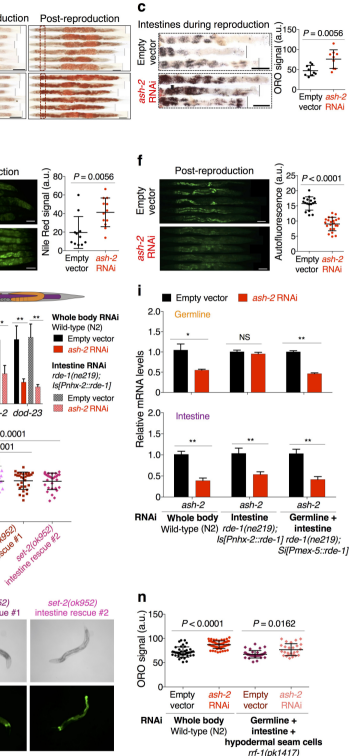
Data availability. RNA-seq data are available at Sequence Read Archive (SRA) under accession number PRJNA343151. The code for RNA-seq analysis is available on the Brunet Laboratory Github: https://github.com/brunetlab.

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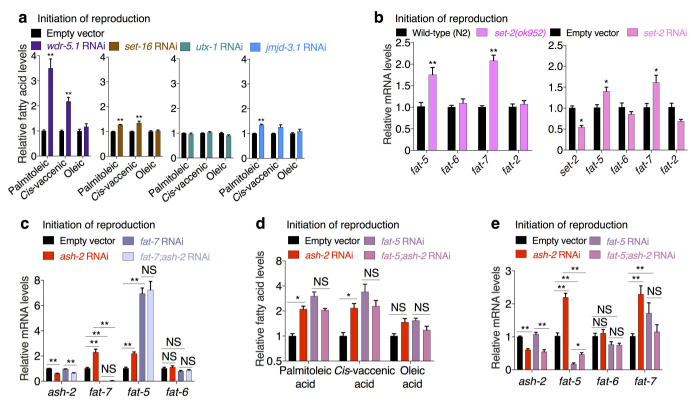




Extended Data Figure 1 | Deficiency in H3K4me3 modifiers leads to fat accumulation in the intestine without altering fertility. a, ORO quantification. Mean \pm s.d., $n \ge 17$ worms per condition. b, ORO images. Scale bars, 100 µm. c, ORO quantification. Mean \pm s.d., $n \ge 9$ dissected intestines per condition. d, e, Nile Red staining and quantification. Mean \pm s.d., $n \ge 7$ dissected intestines (d) or $n \ge 11$ worms (e) per condition. Scale bars, 100 µm. f, Autofluorescence and quantification. Mean \pm s.d., $n \ge 17$ worms per condition. Scale bars, 100 µm. g, Fertility quantification of live brood size (i), fertilized eggs (ii), and unfertilized occytes (iii) laid per worm. Mean \pm s.d., $n \ge 25$ worms per condition. h, RT–qPCR. Mean \pm s.e.m. of two independent experiments, each with two or three biological replicates. i, RT–qPCR. Mean \pm s.e.m. from two

independent experiments, each with two or three biological replicates. **j**, ORO quantification. Mean \pm s.d., $n \ge 26$ worms per condition. **k**, ORO quantification. Mean \pm s.d., $n \ge 29$ worms per condition. **l**, RT–qPCR. Mean \pm s.e.m. of three biological replicates. **m**, Differential interference contrast (DIC) (Nomarski) and GFP fluorescence images. **n**, ORO quantification. Mean \pm s.d., $n \ge 26$ worms per condition. **a**–**f**, **k** (rescue line number 1), **n**, Representative of two experiments. *P* values: **a**, **c**–**g**, two-tailed Mann–Whitney; **h**, **i**, two-tailed Mann– Whitney with Benjamini–Hochberg correction; **j**, **k**, **n**, Kruskal–Wallis with Dunn's correction; **l**, two-tailed *t*-test with Benjamini–Hochberg correction. **P* < 0.05, ***P* < 0.01.

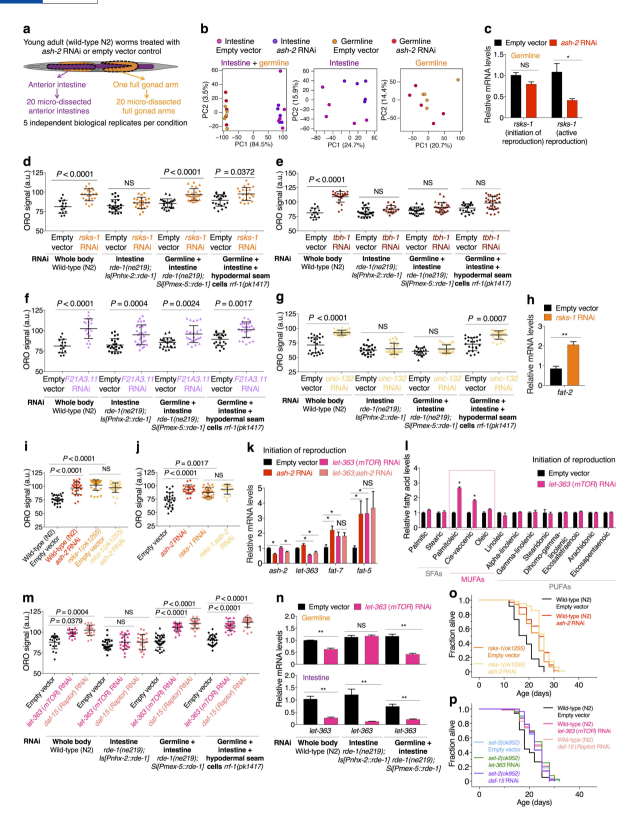
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Extended Data Figure 2 | The delta-9 desaturases FAT-5 and FAT-7 support MUFA accumulation in H3K4me3 methyltransferase-deficient worms. a, GC–MS quantification of MUFAs. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. b, c, RT–qPCR. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. d, GC–MS quantification of MUFAs.

Mean \pm s.e.m. of two independent experiments, each with two or three biological replicates. **e**, RT–qPCR. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. *P* values: **a**–**e**, two-tailed Mann–Whitney with Benjamini–Hochberg correction. **P* < 0.05, ***P* < 0.01.

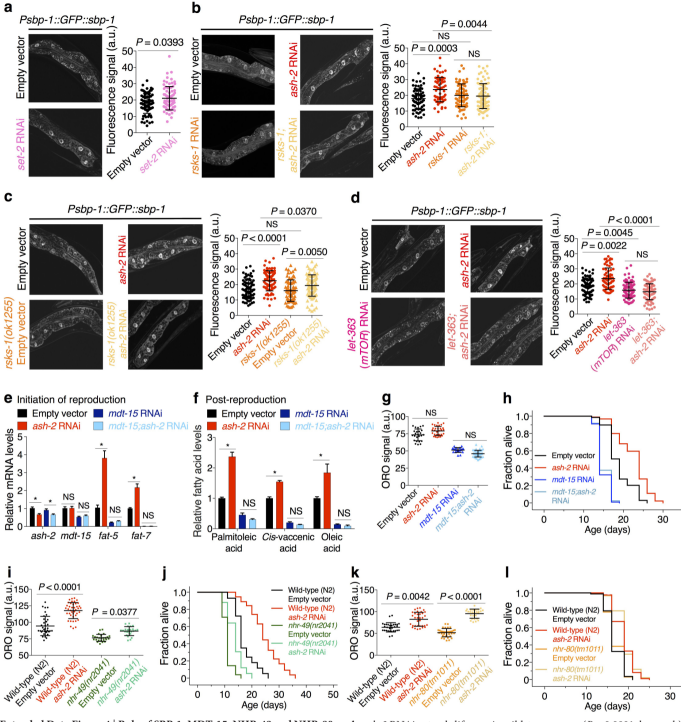




Extended Data Figure 3 | RNA-seq on micro-dissected germlines and intestines and functional validation of ASH-2 targets. a, RNA-seq tissue sample collection pipeline. b, Principal component analysis (PCA) with both intestinal and germline samples (left), only intestinal samples (middle) or only germline samples (right). c, RT-qPCR. Mean \pm s.e.m. of two independent experiments, each with two or three biological replicates. d-g, ORO quantification. Mean \pm s.d., $n \ge 15$ worms per condition. h, RT-qPCR. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. i, j, ORO quantification. Mean \pm s.d., $n \ge 21$ (i) and $n \ge 27$ (j) worms per condition. k, RT-qPCR. Mean \pm s.e.m. of two independent experiments, each with three biological replicates.

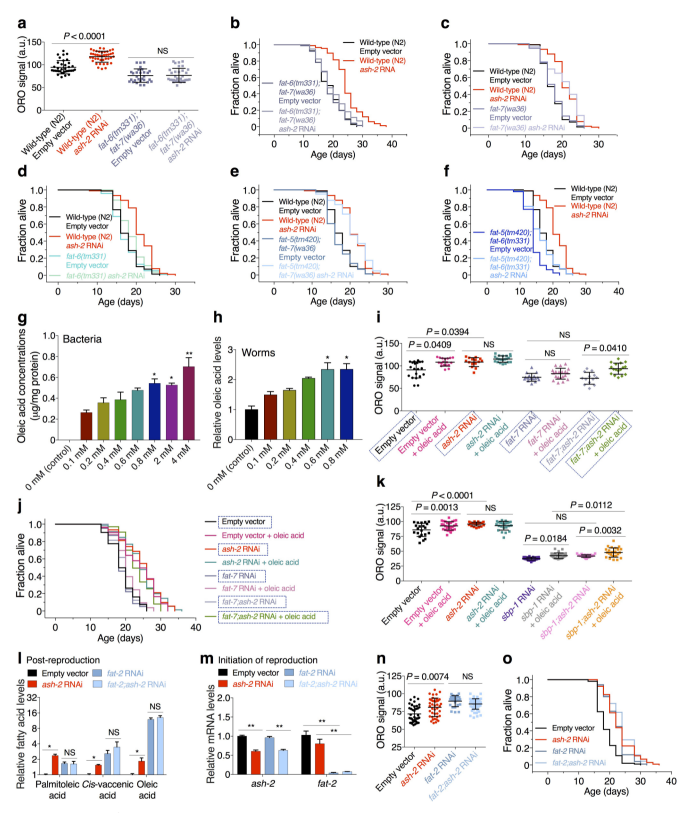
l, GC–MS. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. **m**, ORO quantification. Mean \pm s.d., $n \ge 19$ worms per condition. **n**, RT–qPCR. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. **o**, Lifespan extension by *ash-2* RNAi is reduced in *rsks-1* mutants (13.12%) compared to wild-type worms (29.20%) (P < 0.0001, two-way ANOVA). **p**, *let-363* RNAi and *daf-15* RNAi extend lifespan in wild-type worms (P < 0.0001, log-rank), but not in *set-2* mutants. **d** (except *rrf-1* data), **i**, **j**, Representative of two experiments. *P* values: **c**, **k**, **l**, **n**, two-tailed Mann–Whitney with Benjamini–Hochberg correction; **h**, two-tailed Mann–Whitney; **d–g**, **i**, **j**, **m**, Kruskal–Wallis with Dunn's correction. *P < 0.05, **P < 0.01.

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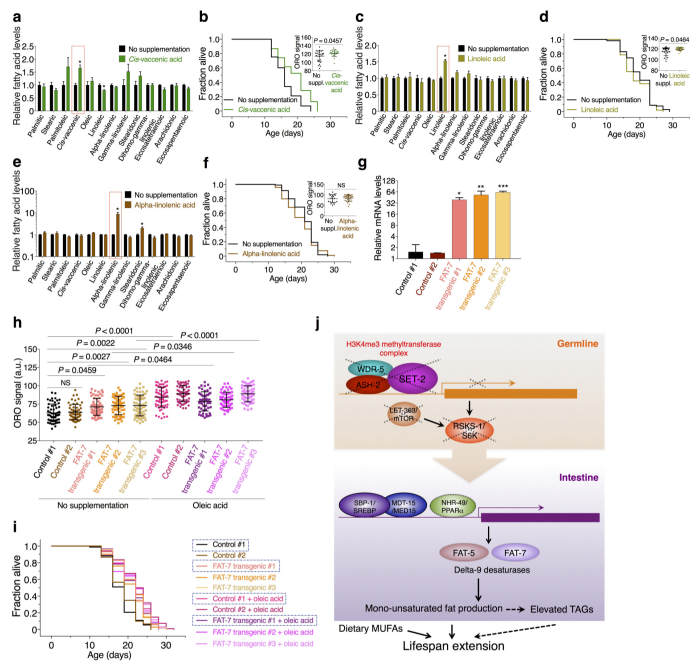
Extended Data Figure 4 | Role of SBP-1, MDT-15, NHR-49 and NHR-80 in the fat accumulation and longevity of H3K4me3 methyltransferasedeficient worms. a–d, Images and quantification of SBP-1 nuclear accumulation. Mean \pm s.d. of two independent experiments, each with 4–6 nuclei per worm of \geq 8 worms per condition. e, RT–qPCR. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. f, GC–MS quantification of MUFAs. Mean \pm s.e.m. of two independent experiments, each with two or three biological replicates. g, ORO quantification. Mean \pm s.d., $n \geq 21$ worms per condition.

h, *ash-2* RNAi extends lifespan in wild-type worms (P < 0.0001, log-rank), but not in *mdt-15* RNAi worms. **i**, ORO quantification, Mean \pm s.d., $n \ge 24$ worms per condition. **j**, *ash-2* RNAi extends lifespan in both wild-type worms and *nhr-49* mutants (P < 0.0001, log-rank). **k**, ORO quantification. Mean \pm s.d., $n \ge 27$ worms per condition. **l**, *ash-2* RNAi extends lifespan in both wild-type worms and *nhr-80* mutants (P < 0.0001, log-rank). **i**–**k**, Representative of two experiments. *P* values: **a**, two-tailed Mann–Whitney; **b–d**, **g**, **i**, **k**, Kruskal–Wallis with Dunn's correction; **e**, **f**, two-tailed Mann–Whitney with Benjamini–Hochberg correction. *P < 0.05.



Extended Data Figure 5 | See next page for caption.

Extended Data Figure 5 | Delta-9 desaturases FAT-6 and FAT-7 and MUFA oleic acid mediate the longevity of H3K4me3 methyltransferasedeficient worms. a, ORO quantification. Mean \pm s.d., $n \ge 29$ worms per condition. b, *ash-2* RNAi leads to lifespan extension in control (P < 0.0001, log-rank), but not in *fat-6* and *fat-7* double mutants. c, *ash-2* RNAi extends lifespan in both wild-type worms and *fat-7* single mutants (P < 0.0001, log-rank). d, *ash-2* RNAi extends lifespan in *fat-6* single mutants (P < 0.0001, log-rank). d, *ash-2* RNAi extends lifespan in *fat-6* single mutants (P < 0.0001, log-rank), but lifespan extension by *ash-2* RNAi is reduced in *fat-6* mutants (9.26%) compared to wild-type worms (20.46%) (P = 0.0072, two-way ANOVA). e, *ash-2* RNAi extends lifespan in *fat-5* and *fat-7* double mutants (P < 0.0001, log-rank), f. *ash-2* RNAi extends lifespan in *fat-5* and *fat-6* double mutants (P < 0.0001, log-rank), f. *ash-2* RNAi extends lifespan extension by *ash-2* RNAi is reduced in *fat-5* and *fat-6* double mutants (P < 0.0001, log-rank). f. *ash-2* RNAi extends lifespan extension by *ash-2* RNAi is reduced in *fat-5* and *fat-6* double mutants (P < 0.0001, log-rank), for lifespan extension by *ash-2* RNAi is reduced in *fat-5* and *fat-6* double mutants (P < 0.0002, log-rank), but lifespan extension by *ash-2* RNAi is reduced in *fat-5* and *fat-6* double mutants (P < 0.0002, log-rank), but lifespan extension by *ash-2* RNAi is reduced in *fat-5* and *fat-6* double mutants (P < 0.0002, log-rank), but lifespan extension by *ash-2* RNAi is reduced in *fat-5* and *fat-6* double mutants (P < 0.0358, two-way ANOVA). g, h, GC-MS quantification of oleic acid. Mean \pm s.e.m. of three biological replicates. **i**, ORO quantification. Mean \pm s.d., $n \ge 13$ worms per condition. Boxed conditions are identical to Fig. 4g. **j**, Oleic acid supplementation extends lifespan (P < 0.0001, log-rank), which is not further extended by *ash-2* RNAi. Oleic acid supplementation extends lifespan in *ash-2* and *fat-7* double RNAi worms (P < 0.0001, log-rank). Boxed conditions are identical to Fig. 4h. **k**, ORO quantification. Mean \pm s.d., $n \ge 21$ worms per condition. **I**, GC–MS quantification of MUFAs. Mean \pm s.e.m. of two independent experiments, each with two or three biological replicates. **m**, RT–qPCR. Mean \pm s.e.m. of two independent experiments, each with two or three biological replicates. **m**, $n \ge 25$ worms per condition. **o**, *ash-2* RNAi extends lifespan in control (P < 0.0001, log-rank) but not *fat-2* RNAi worms. **c**, **i**, **j**, **n**, **o**, Representative of two experiments. *P* values: **a**, **g–i**, **k**, **n**, Kruskal–Wallis with Dunn's correction; **l**, **m**, two-tailed Mann–Whitney with Benjamini–Hochberg correction. *P < 0.05, **P < 0.01.



Extended Data Figure 6 | Dietary supplementation with MUFAs, but not PUFAs, extends lifespan in wild-type worms. a, GC–MS. Mean \pm s.e.m. of two independent experiments, each with two or three biological replicates. b, *Cis*-vaccenic acid supplementation extends lifespan in wild-type worms (P < 0.0001, log-rank). Inset: ORO quantification. Mean \pm s.d., $n \ge 20$ worms per condition. c, GC–MS. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. d, Linoleic acid supplementation does not extend lifespan in wild-type worms. Inset: ORO quantification. Mean \pm s.d., $n \ge 23$ worms per condition. e, GC–MS. Mean \pm s.e.m. of two independent experiments, each with three biological replicates. f, Alpha-linolenic acid supplementation does not increase the lifespan of wild-type worms. Inset: ORO quantification. Mean \pm s.d.,

 $n \ge 23$ worms per condition. **g**, RT–qPCR. Mean \pm s.e.m. of three biological replicates. **h**, ORO quantification. Mean \pm s.d., $n \ge 54$ worms per condition. **i**, Overexpression of FAT-7 extends lifespan (P < 0.0001, log-rank), but this lifespan is not extended further by dietary oleic acid. Boxed regions are identical to Fig. 5g. **j**, Proposed model by which *ash*-2 deficiency in the germline could lead to the fat metabolic switch in the intestine. **b**, **d**, **f**, Representative of two experiments. *P* values: **a**, **c**, **e**, two-tailed Mann–Whitney; **g**, Kruskal–Wallis test with Dunn's correction (non-significant, probably owing to small sample size). One-way ANOVA with Bonferroni's correction. **h**, Kruskal–Wallis with Dunn's correction; *P < 0.05, **P < 0.01, ***P < 0.001.

Extended Data Table 1 | List of ASH-2 candidate targets

Candidates	Datasets	Functions	Expression
tbh-1	RNA-seq of ash-2 knockdown worms (adult, micro-dissected tissues)	Tyramine beta-hydroxylase, Required for biosynthesis of the octopamine neurotransmitter, Longevity	Germline
nhr-10	RNA-seq of ash-2 knockdown worms (adult, micro-dissected tissues)	Nuclear hormone receptor, Fat metabolism	Intestine, Germline
unc-132	RNA-seq of ash-2 knockdown worms (adult, micro-dissected tissues)	Unknown roles in fat metabolism and longevity	Germline
F21A3.11	RNA-seq of ash-2 knockdown worms (adult, micro-dissected tissues)	Predicted to have acid phosphatase activity and metal ion binding activity, Longevity	Intestine
asm-2	RNA-seq of <i>ash-2</i> knockdown worms (adult, micro-dissected tissues) Microarray of <i>ash-2</i> knockdown worms (larval, whole worms)	Acid sphingomyelinase	Intestine
lod-23	RNA-seq of <i>ash-2</i> knockdown worms (adult, micro-dissected tissues) Microarray of <i>ash-2</i> knockdown worms (larval, whole worms)	Determinant of adult lifespan	Intestine
acs-2	Microarray of ash-2 knockdown worms (larval, whole worms)	Fatty acid beta-oxidation	Intestine (predicted)
acdh-2	Microarray of ash-2 knockdown worms (larval, whole worms)	Fatty acid beta-oxidation	Intestine (predicted)
rsks-1	ASH-2 ChIP-chip (whole embryos)	S6 kinase (S6K), Fat metabolism, Longevity	Germline (predicted)
aak-2	ASH-2 ChIP-chip (whole embryos)	AMPK subunit, Nutrient sensing, Longevity	Intestine, Germline (predicte

List of candidate targets of ASH-2 based on our gene expression dataset as well as published datasets.