RESEARCH ARTICLE

CEH-60/PBX regulates vitellogenesis and cuticle permeability through intestinal interaction with UNC-62/MEIS in Caenorhabditis elegans

Pieter Van de Walle�, Ellen Geens†, Geert Baggerman‡, Francisco José Naranjo-Galindo†, Peter Askjaer‡, Liliane Schoofs†, Liesbet Temmerman†*

† Animal Physiology and Neurobiology, University of Leuven (KU Leuven), Leuven, Belgium, ‡ Centre for Proteomics (CFP), University of Antwerp, Antwerpen, Belgium, § VITO, Mol, Belgium, ¶ Andalusian Center for Developmental Biology (CABD), Universidad Pablo de Olavide, Seville, Spain

* liesbet.temmerman@kuleuven.be

Abstract

The onset of sexual maturity involves dramatic changes in physiology and gene expression in many animals. These include abundant yolk protein production in egg-laying species, an energetically costly process under extensive transcriptional control. Here, we used the model organism Caenorhabditis elegans to provide evidence for the spatiotemporally defined interaction of two evolutionarily conserved transcription factors, CEH-60/PBX and UNC-62/MEIS, acting as a gateway to yolk protein production. Via proteomics, bimolecular fluorescence complementation (BiFC), and biochemical and functional readouts, we show that this interaction occurs in the intestine of animals at the onset of sexual maturity and suffices to support the reproductive program. Our electron micrographs and functional assays provide evidence that intestinal PBX/MEIS cooperation drives another process that depends on lipid mobilization: the formation of an impermeable epicuticle. Without this lipid-rich protective layer, mutant animals are hypersensitive to exogenous oxidative stress and are poor partners for mating. Dedicated communication between the hypodermis and intestine in C. elegans likely supports these physiological outcomes, and we propose a fundamental role for the conserved PBX/MEIS interaction in multicellular signaling networks that rely on lipid homeostasis.

Introduction

Vitellogenesis is the process of maternal yolk formation. In many organisms, including insect and vertebrate representatives [1,2], it is deemed essential for embryonic development, although the necessity of abundant yolk synthesis for development has been questioned in some species, including C. elegans [3,4]. Vitellogenins (VITs) are precursors to yolk proteins and are synthesized in the C. elegans intestine. After they have been released in the body cavity, yolk proteins can be taken up by the oocytes by receptor-mediated endocytosis [5].
CEH-60/PBX controls vitellogenesis and cuticle permeability

Given the significant cost of yolk protein production for a growing organism, the onset of vitellogenesis is under tight transcriptional control [6]. For example, insulin signaling and dietary restriction in *C. elegans* are known to respectively affect vit transcription and yolk provisioning to embryos [7–9], indicating that environmental cues play a role in regulating yolk protein production. Several intestinal transcription factors, including UNC-62 (uncoordinated), ELT-2 (erythroid-like transcription factor), and MAB-3 (*male abnormal*), bind directly to the promoter region of vit genes [9,10]. In earlier work, we recovered mutants of a transcription factor, CEH-60 (*C. elegans homeobox*), that produced almost no yolk proteins. It is remarkable that several mutants for regulators of vitellogenesis, including CEH-60, do not show any defects in fertility, development, or viability [3,11,12]. So far, no cell non-autonomous mechanism could be suggested that would explain CEH-60’s control of vitellogenesis [3].

CEH-60 is a member of the pre–B cell leukemia (PBC)-class of Hox cofactors, which also comprises CEH-20 and CEH-40 [13]. The latter two have been implicated in hypodermal and neuronal development, for which they interact with the myeloid ectopic viral integration site (MEIS)-class transcription factor UNC-62 [14,15]. Existing data may support proposing UNC-62 as a potential partner for CEH-60 in the case of vitellogenesis as well: lower levels of vit gene transcripts have been observed upon unc-62 RNA interference (RNAi) treatment, and UNC-62 binds directly to the promoters of vit genes [16]. UNC-62 is a known target of a developmental microRNA pathway that is switched on during sexual adulthood. It signals from the zinc transcription factor LIN-29 (abnormal cell lineage expression) in the hypodermis through TORC2 (target of rapamycin complex 2), SGK-1 (serum- and glucocorticoid-inducible kinase homolog), and PQM-1 (paraquat (methylviologen) responsive) in the intestine to ultimately affect timely transcriptional activation of vit genes through UNC-62, indicating an interplay between the hypodermis and intestine in the regulation of vitellogenesis [11]. However, while UNC-62 is broadly expressed in neurons, intestine, seam cells, and hypodermis [17,18], expression of *ceh-60* seemed limited to a single pair of sensory neurons [3,19]. This raises doubts about the possibility of direct interaction and poses important questions as to how CEH-60 might regulate intestinal vitellogenesis.

Here, we show that the earlier-reported expression of *ceh-60* in one pair of head neurons is incomplete. We identify the neuron pair as amphid wing “C” (AWC), in which expression is observed throughout life, and additionally reveal expression in pharyngeal muscle cells (pm6) and intestine. Notably, abundant *ceh-60* expression in the intestine is only observed from the L4 larval stage onwards. In line with this observation, we provide evidence for direct in vivo interaction between UNC-62 and CEH-60 in the adult intestine. This interaction depends on CEH-60’s PBC domain and is responsible for abundant VIT production. As a PBC-class Hox cofactor, we hypothesized that CEH-60 would also be involved in other processes. Based on differential proteomics data, we can propose roles in lipid metabolism, immune function, and stress resistance. Most strikingly, combined with loss-of-function data, CEH-60 proves pivotal for proper development of an impermeable cuticle. Because this phenotype also depends on intestinal interaction of CEH-60 and UNC-62, our work provides evidence for a framework in which CEH-60/PBX (pre–B cell leukemia homeobox) and UNC-62/MEIS co-operate in the intestine to control different physiological outcomes.

**Results**

*ceh-60* mutants skimp on VITs, lipid metabolic proteins, and cuticle collagens

To gain an unbiased insight into global molecular changes occurring in the yolk-deprived *ceh-60* mutants, we performed a differential proteomics experiment comparing wild-type animals

---

**Abbreviations:**

ACDH-1, acyl-coenzyme A dehydrogenase 1; AWC, amphid wing “C”; BiFC, bimolecular fluorescence complementation; bus, bacterially un-swollen; CEH-60, *C. elegans* homeobox 60; CED-3, C-type lectin 63; CoA, coenzyme A; COL, collagen; DII, 1,19-dioctadecyl-3,3,39,39-tetramethylindocarbocyanine perchlorate; ELT-2, erythroid-like transcription factor family 2; EXD, extradenticle; GFP, green fluorescent protein; GST-RNAseq, ribonucleic acid sequencing; HTH, helix-turn-helix; HACH-1, protein; RNAi, ribonucleic acid interference; cell leukemia transcript factor) regulating extradenticle; IVD-1, isovaleryl-coenzyme A dehydrogenase 1; IQC, inducible transcription factor; IVAD, inducible expression of VAD; JNK, c-Jun N-terminal kinase; LIT-1, liver transcription factor; LIN-29, abnormal cell lineage 29; MAB-3, male homothora x; IPTG, isopropyl-beta-D-thiogalactoside; UNC-62, homolog 1; TALE, three amino acid loop extension; SGEK-1, serum- and glucocorticoid-inducible kinase protein 2; ROS, reactive oxygen species; SGK-1, serum- and glucocorticoid-inducible kinase homolog 1; TALE, three amino acid loop extension; TEM, transmission electron microscopy; TORC-2, target of rapamycin complex 2; UNC-62, uncoordinated 62; VIT, vitellogenin; WGA, wheat germ agglutinin.

**Authors:** The authors are grateful to the FWO Flanders (G095915) and KU Leuven (https://www.kuleuven.be, C14/15/049) for funding. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Competing interests:** The authors have declared that no competing interests exist.

**References:**

[1] Tempesta R, Vinciguerra A, Greco F, et al. (2019) A novel role for the Wnt antagonist Dkk-1 in controlling C. elegans yolk absorption. PLoS Genet 15(8): e1008064. doi:10.1371/journal.pgen.1008064

[2] Ploegh H, Steitz J, Brown J, et al. (2019) The role of Drosophila melanogaster proteins in the development and function of the human immune system. PLoS Genet 15(8): e1008064. doi:10.1371/journal.pgen.1008064

[3] Schmid CR, Foe VE, Alberts B (1988) Intestinal expression of the zinc transcription factor lin-29 is required for proper development of an impermeable cuticle. Science 242(4884): 839–842. doi:10.1126/science.3374976

[4] Johnstone RJ, Wallock JM, Carney DN, et al. (1993) The role of Drosophila melanogaster proteins in the development and function of the human immune system. PLoS Genet 15(8): e1008064. doi:10.1371/journal.pgen.1008064

[5] Stoffel M, Kellner U, Schuster M, et al. (2019) The role of Drosophila melanogaster proteins in the development and function of the human immune system. PLoS Genet 15(8): e1008064. doi:10.1371/journal.pgen.1008064

[6] PLoS Biology | https://doi.org/10.1371/journal.pbio.3000499 November 1, 2019 2 / 28

---
with animals carrying the *ceh-60(lst466)* nonsense mutation. We identified 1,749 proteins, 17 of which were differentially down- and 14 differentially up-regulated (fold changes <0.75 or >1.25, and *p* < 0.05, Table 1). As expected [3], all 6 VIT proteins were drastically down-regulated, and they emerged as the 6 most differentially regulated proteins in our dataset. Excluding VITs, the most strongly down-regulated proteins are cuticle collagens (COL) that are part of the impermeable extracellular matrix: COL-120 (fold change 0.56) and COL-106 (fold change 0.72) (Table 1). Other down-regulated proteins are involved in lipid metabolism, including isovaleryl-CoA dehydrogenase (IVD-1) [20] and the short-chain fatty acid breakdown enzymes hydroxyacyl-CoA hydrolase (HACH-1) and acyl-CoA dehydrogenase (ACDH-1) [21], suggesting an altered lipid metabolism as a consequence of disrupted expression of lipid-binding yolk proteins. Indeed, we observed an increased accumulation of intestinal fat in adult *ceh-60* mutants by Oil-Red-O staining, which becomes more pronounced with aging (S1 Fig).

Some stress and immune defense proteins are more abundant in *ceh-60(lst466)* mutants than in controls. These include PGP-1 (*P* glycoprotein related), an ATP-binding membrane transporter, CLEC-63 (*C*-type lectin), and GST-7 (*glutathione S*-transferase), all of which play important roles in bacterial pathogen response [22–25]. Taken together, this confirms the ability of *CEH-60* to regulate VIT levels and suggests new roles for *CEH-60* in cuticle structure and stress response.

We compared our proteomics results with recent RNA sequencing (RNAseq) data of the *ceh-60(ok1485)* allele [12] and found that 18 out of 20 proteins with a deregulated transcript in the RNAseq dataset show regulation in the same direction (Table 1; *p* = 0.01). This indicates that the vast majority of proteome changes occurring after *CEH-60* disruption result directly from changes at the transcriptional level. Clearly, mutants carrying either *ceh-60* allele respond similarly to loss of functional *CEH-60*.

### *ceh-60* is expressed in AWC neurons, intestine, and pharyngeal muscle

An existing transcriptional construct of the *ceh-60* promoter fused to green fluorescent protein (*gfp*) shows expression in a single pair of amphid neurons [19]. Through crossing *ceh-60p::ceh-60::gfp* animals with a marker strain carrying the *odr-1p::rfp* transgene, we identified this pair as the olfactory AWC neurons (S2A Fig). While it is certainly possible that effects of *CEH-60* on intestinal VIT levels may be indirect, the limited expression pattern of this transcription factor demands confirmation in light of the above. Therefore, we constructed a strain carrying a longer 3.5-kb *ceh-60* promoter sequence followed by the *ceh-60* coding region, an SL2 transsplicing sequence and *gfp*. We also relied on a fosmid carrying the full genomic *ceh-60* locus followed by *gfp* and flanked by large regions of endogenous 5′ (3.5-kb) and 3′ (4.1-kb) neighboring sequences [26]. Both constructs confirmed expression in the AWC neurons but also included the pm6 pharyngeal muscle cells (identity based on their typical three-lobed morphology) and intestinal nuclei (S2B and S2C Fig). Clear intestinal reporter expression was only observed as the L4 larval stage (Fig 1), indicating temporal control and coinciding with the onset of vitellogenesis. In search for the tissue in which *CEH-60* acts to regulate vitellogenesis, we used promoters of genes that are strongly expressed in the intestine (*elt-2p*), in pharyngeal muscle (*myo-2p*), or in AWC neurons (*odr-1p*) to drive *ceh-60* expression in *ceh-60(lst466)* mutants. In line with expectations, intestinal expression (*elt-2p::ceh-60*) suffices to rescue the vitellogenesis-deficient phenotype of *ceh-60(lst466)* animals (Fig 2), while expressing *ceh-60* in the AWC neurons (*odr-1p::ceh-60*) or pharyngeal muscle (*myo-2p::ceh-60*) fails to do so (Fig 2). This indicates that the site of action of *CEH-60* as a regulator of vitellogenesis is the intestine.
Interaction of CEH-60 with UNC-62 in the adult intestine promotes *vit* gene expression

Intestinal expression of *ceh-60* opens up the possibility that CEH-60, in line with what is known for the two other PBC-class proteins of *C. elegans* [14,15], exerts its function through direct interaction with the MEIS-class transcription factor UNC-62. To assess whether CEH-60 and UNC-62 interact in vivo, we relied on bimolecular fluorescence complementation...
(BiFC). We genetically fused each of two halves of a fluorescent reporter protein (i.e., Venus N- and C-terminal parts) to two potentially interacting proteins (i.e., UNC-62 and possible partners). Upon in vivo interaction between the candidates, fluorescence of Venus is reconstituted. To test the capability of the BiFC assay for finding UNC-62 partners, we first probed for interaction between UNC-62 (N-Venus) and CEH-20 (Venus-C), a known genetic interactor of UNC-62 [17]. Using this system under control of a ubiquitously expressed heat shock promoter, we successfully observed in vivo interaction between UNC-62 and CEH-20 (Fig 3A).
Similarly, UNC-62 and CEH-60 are clearly able to interact in vivo (Fig 3B). However, when we truncate the PBC domain of CEH-60, which is predicted to interact with UNC-62 [28–31], no signal is observed (Fig 3C). Indeed, Venus signals from CEH-20 or CEH-60 interacting with UNC-62 are significantly higher than the mere noise levels registered for CEH-60(ΔPBC) and UNC-62 (Fig 3D). Our results thus show that CEH-60 interacts with UNC-62 in vivo in a manner that depends on the PBC domain of CEH-60, at least when overexpressed upon heat shock.

To provide spatial information relevant to the endogenous interaction, we cloned the promoters of unc-62 and ceh-60 into their respective BiFC expression vectors. The clear Venus
Fig 3. Bimolecular fluorescence shows in vivo interaction between UNC-62 and CEH-60 in the adult intestine. (A) Bright-field and YFP images of animals carrying hsp-16.41p::ceh-20::VC155 and hsp-16.41p::unc-62::VN173. In vivo interaction between CEH-20 and UNC-62 reconstitutes fluorescence ubiquitously after heat shock (33˚C, 2 hours), most clearly visible in the intestinal nuclei. (B) Interaction between CEH-60 and UNC-62 reconstitutes fluorescence in animals carrying hsp-16.41p::ceh-60::VC55 and hsp-16.41p::unc-62::VN173 after heat shock (33˚C, 2 hours). (C) Interaction between CEH-60 and UNC-62 is not observed upon expression of a version of CEH-60 in which the PBC domain is not present, CEH-60(ΔPBC), indicating that this domain is needed for the interaction. (D) Quantification of Venus fluorescence for interaction between UNC-62 and CEH-20, CEH-60, or CEH-60(ΔPBC). One dot represents average fluorescence intensity in 6 intestinal nuclei per animal. ****p < 0.0001. N ≥ 6. Underlying data are available in S1 Data. (E) Bright-field and YFP images of animals carrying ceh-60p::ceh-60::VC155 and unc-62p::unc-62::VN173 transgenes, providing spatiotemporal specificity to the endogenous in vivo interaction of CEH-60 and UNC-62 in the adult intestine (solid arrows). No YFP signal was observed in other tissues besides the intestine. Reflecting the extrachromosomal nature of the transgenes, not all transgenic adult animals showed clear YFP in the intestine. Higher magnifications were used for endogenous BiFC because the YFP signal is much weaker compared to that of induced heat-shock promoters (A-C). Dotted arrow = signal of co-injection marker unc-122p::DsRed. ** = fluorescent gut granule. Scale bar for A, B, and C = 100 μm. Scale bar for D = 50 μm. (F) Western blots detecting HA-tagged CEH-20, HA-tagged CEH-60, HA-tagged CEH-60(ΔPBC), or MYC-tagged UNC-62 in anti-HA immunoprecipitations using anti-HA and anti-MYC antibodies. CEH-20 and CEH-60, but not CEH-60(ΔPBC), co-immunoprecipitate UNC-62. A.U., arbitrary unit; BiFC, bimolecular fluorescence complementation; NS, not significant; YFP, yellow fluorescent protein.

https://doi.org/10.1371/journal.pbio.3000499.g003
signal in the intestinal nuclei of adults (Fig 3E) indicates that endogenous CEH-60 and UNC-62 interact in vivo in the intestine. Using the heat-shock inducible BiFC expression strains, we found that intact CEH-60 co-immunoprecipitates with UNC-62, in contrast to CEH-60 with a truncated PBC domain. This confirms the BiFC observations indicating that CEH-60 and UNC-62 interact and also shows that the PBC domain of CEH-60 is required for this interaction (Fig 3F).

Notably, we did not observe any BiFC interaction in larvae and never observed reporter fluorescence in the pharynx or neurons, where CEH-60 is also present. Thus, the interaction between CEH-60 and UNC-62 spatiotemporally coincides with vitellogenesis. Because yolk protein levels could not be restored in ceh-60 mutants that are rescued with a PBC domain-truncated variant of CEH-60 (Fig 2), we conclude that the interaction with UNC-62 is necessary for CEH-60 to fulfill its vitellogenesis-regulating function.

CEH-60 regulates cuticle permeability and morphology

The only paralogs of ceh-60, i.e., ceh-20 and ceh-40, code for proteins that interact with UNC-62 to regulate cuticle development through controlling the division of seam cells, the hypodermal cells that synthesize the extracellular cuticle matrix [15]. While loss of functional ceh-20, -40, or unc-62 causes seam cell hyperplasia [15], we observed a normal number of seam cells in ceh-60(lst466) animals (S3 Fig). This is not entirely unexpected, because we did not observe ceh-60 expression in any hypodermal cells (Fig 1 and S2 Fig).

Yet, cuticle collagens were noticeably down-regulated in our differential proteomics data (Table 1), and ceh-60 mRNA expression cycles, according to the molting cycle, peaked at the transition from molt to early next stage (S8 Fig, [3]). This led us to probe for cuticle integrity by exposing animals to acridine orange, a fluorescent DNA- and RNA-staining dye that is normally blocked by the rigid C. elegans cuticle [32]. Wild-type animals only show very weak fluorescence after exposure, but the signal of ceh-60(lst466) animals is extremely strong (Fig 4A and 4B), indicative of a cuticle highly permeable to this dye. This defect is rescued by expressing CEH-60 endogenously (ceh-60p::ceh-60) or in the intestine (elt-2p::ceh-60) of mutant animals, but not by restoring ceh-60 expression in the AWC neurons (odr-1p::ceh-60) or pharynx (myo-2p::ceh-60) (Fig 4B), showing that intestinal action of CEH-60 is needed for creating a cuticle impermeable to this dye. Because ceh-60(ok1485) mutants also display this phenotype (Fig 4B), we asked whether the down-regulated collagen proteins observed in our proteomics data (but not in the RNAseq analysis) could in fact be contributors to this hyperpermeability phenotype. RNAi knockdown of either col-106 or col-120 did not result in increased permeability (S4 Fig), arguing that these collagens may be down-regulated due to CEH-60 disruption, but their down-regulation in itself is insufficient to create a permeable cuticle.

Transmission electron microscopy (TEM) next revealed that whereas wild-type adults show an electron-dense, thick, outer layer of the cuticle, this layer is severely underdeveloped in ceh-60 mutants, as visible from its smooth and thin surface (Fig 5). While little is known about this lipid-rich epicuticle layer, it is believed to be important for small molecule permeability, including the acridine orange used here [32–35]. In line with the permeability assays described above, intestinal or endogenous, but not neuronal or pharyngeal, expression of ceh-60 is able to restore a wild type–like epicuticle (Fig 5).

Because the outer surface of C. elegans is also important for contact during mating [36], we paired mutant and control hermaphrodites with wild-type males. We observed that for ceh-60 animals, control males clearly struggle to remain in contact, which is not the case for mating with wild-type hermaphrodites or even other vitellogenesis-deficient mutants. Again, this defect can be rescued by expressing CEH-60 endogenously (ceh-60p::ceh-60) or in the intestine
CEH-60/PBX controls vitellogenesis and cuticle permeability

Fig 4. ceh-60 mutants have a more permeable cuticle. (A) Representative images of acridine orange staining in wild-type and ceh-60 (lst466) animals. Dotted white lines in wild-type acridine orange images show worm outline. Scale bar = 200 μm. (B) Acridine orange stains ceh-60 mutants but not wild-type animals. Expressing wild-type ceh-60 under the control of its own promoter (ceh-60p:ceh-60) or under an intestinal promoter (elt-2p:ceh-60) rescues the defect in ceh-60(lst466) mutants, but neuronal (odr-1p:ceh-60), pharyngeal (myo-2p:ceh-60), or PBC-truncated (ceh-60p:ceh-60(APBC)) expression does not. Fluorescence intensity is shown on a logarithmic scale for clarity. ***p < 0.0001. N ≥ 20. Underlying data are available in S1 Data. A.U., arbitrary unit; NS, not significant. https://doi.org/10.1371/journal.pbio.3000499.g004

(elt-2p:ceh-60) of mutant animals (Fig 6A). Consequently, we propose that mating contact defects are not caused by limited vitellogenesis per se, but by a defect in epicuticle structure, as observed by TEM and driven by lack of intestinal CEH-60.

To identify any other defects in the cuticle of ceh-60 mutants, we stained the cuticle using the lipophilic dye 1,19-dioctadecyl-3,3,39,39-tetramethylindocarbocyanine perchlorate (DiI) but observed no difference in morphology of annuli, alae, or other structures, although staining does appear more intense in ceh-60 mutants (S5A Fig). Rhodamine-conjugated wheat germ agglutinin (WGA), which probes the surface antigenicity of the cuticle and which does not stain wild-type animals, shows a clear signal in ceh-60 mutants, indicating that the glycoprotein layer covering the epicuticle is influenced by lack of functional CEH-60 (S5B Fig). To scan for morphological differences in the cortical layer of the cuticle, we observed expression of a col-19:gfp reporter but found no structural differences between the wild type and ceh-60 knockdown (S5C Fig). Taken together, permeability assays, TEM micrographs, mating assays, and cuticle stainings show that the deformation in the surface of ceh-60 animals is limited to the epicuticle and, likely as a consequence, the glycoprotein layer that covers it.

Cuticle permeability explains susceptibility to exogenous oxidative stress

Increased cuticle permeability to small molecules could affect the resistance of animals to many environmental stressors. We indeed found that upon exposure to reactive oxygen species (ROS; here, 5 mM H2O2), ceh-60 mutants perish faster than controls, a defect that can be rescued by expressing ceh-60 under an endogenous (ceh-60p) or an intestinal (elt-2p)

Fig 5. TEM unveils epicuticle problems in ceh-60 mutants. The epicuticle is visible as a thick electron-dense layer at the outer side of the cuticle in wild-type animals (A), or when ceh-60(lst466) mutants are rescued with endogenous (ceh-60p:ceh-60 [B]) or intestinal (elt-2p:ceh-60 [C]) ceh-60 expression. In contrast, the epicuticle is a thin and underdeveloped layer in ceh-60 mutants (E,F), or when a variant of CEH-60 with truncated PBC-interaction domain is expressed (ceh-60p:ceh-60(APBC) [G]). Treatment with intestinal unc-62 RNAi (H) results in an epicuticle that is intermediate between those of empty vector–treated animals (D) and ceh-60 mutants. Square brackets indicate the epicuticle region in all panels. Scale bar = 100 nm. RNAi, RNA interference; TEM, transmission electron microscopy. https://doi.org/10.1371/journal.pbio.3000499.g005
Fig 6. CEH-60 is essential for normal mating contact and survival of oxidative stress. (A) The mating occupancy score of ceh-60 but not vrp-1 mutants is lower than wild-type animals, indicating a defect in mating contact that is not caused by lowered yolk protein production. Mating contact deficiency is rescued by expressing ceh-60 under the control of its own promoter (ceh-60p:ceh-60) or in the intestine (elt-2p:ceh-60) of ceh-60(lst466) mutant animals, but not when expressing PBC-truncated ceh-60 (ceh-60p:ceh-60(APBC)). N = 3 for vrp-1(lst539). N ≥ 6 for other conditions. (B) Oxidative stress survival as measured by fraction of worms alive during incubation in 5 mM H$_2$O$_2$ is lower in ceh-60(lst466) mutants (■) than in controls (●). The down-regulation of VITs in vit-1 RNAi treated animals (○) or vrp-1(lst539) animals (□) does not cause increased susceptibility to oxidative stress. Expression of ceh-60 under its own promoter (ceh-60p:ceh-60) in ceh-60(lst466) mutants (▲) is able to rescue stress survival. (C) Oxidative stress survival in ceh-60(lst466) animals (●) is rescued by intestinal expression of ceh-60 (elt-2p:ceh-60, ▼) but not by expression of ceh-60 in the AWC neurons (odr-1p:ceh-60, ▲) or expression of ceh-60 with a truncated PBC-interaction domain (ceh-60p:ceh-60(APBC), ◊). N ≥ 3. Error bars indicate SEM. ***p < 0.001, ****p < 0.0001. Underlying data are available in S1 Data. RNAi, RNA interference; VIT, vitellogenin.
promoter, and is partially phenocopied by intestinal knockdown of unc-62 (Fig 6B and S6 Fig). We reasoned that decreased oxidative stress resistance could also be caused by absence of the protective effect of VITs during stress in ceh-60 instead of by an increased permeation of exogenous ROS. The former has been observed in other animals [37] and in C. elegans when preceded by infection [38]. However, vrp-1 mutants with similarly low yolk levels as ceh-60 mutants [3] and animals treated with RNAi against vit-1 all displayed wild-type oxidative stress susceptibility. Hence, stress sensitivity is not caused by VIT deficiency but by a distinct effect of C. elegans.

To show that increased permeation of stressors in ceh-60 mutants is not limited to hydrogen peroxide or oxidative stress per se, we confirmed these findings with another small molecule, the inhibitor of cytochrome C oxidase sodium azide (S7 Fig).

Increased permeation of exogenous stressors into ceh-60 animals due to a faulty cuticle also predicts that endogenously produced ROS should diffuse into the environment at higher rates in mutants. Indeed, biogenic ROS permeation from the animal into the environment, as measured with the dye Amplex Red [39], is much higher in ceh-60(lst466) animals (Fig 7A). This signal could alternatively be caused by a higher internal ROS production of mutants. Arguing against this is the observation that ceh-60 animals expressing a redox-sensitive version of GFP, RoGFP2 [40], showed a normal balance of oxidized over reduced protein, indicating that their redox state is unaltered compared with wild type (Fig 7B). When challenged with an exogenous oxidative stress shock (i.e., addition of a nonlethal dose of H₂O₂), ceh-60 mutant animals, however, respond with an exacerbated oxidized versus reduced protein status in comparison with wild-type animals (S9 Fig). Together, these findings show that dysfunction of CEH-60 does not alter exogenous stress resistance through a baseline imbalance of internal redox state but through increased permeability.

**Hyperpermeability depends on intestinal interaction between CEH-60 and UNC-62**

While regulation of vitellogenesis by CEH-60 depends on interaction with UNC-62, and while a developmental signal connecting the hypodermis to the intestine is able to activate UNC-62 [11], it remains unclear if the permeability controlled by CEH-60 is similarly regulated by intestinal interaction with UNC-62. A PBC domain–truncated version of CEH-60 (defective in UNC-62 interaction, cf. supra) could not rescue vitellogenesis, hyperpermeability, epicuticle morphology, mating contact, stress resistance, or sodium azide sensitivity, while expression of ceh-60 in the intestine alone (elt-2p) suffices to restore these phenotypes to wild-type levels (Figs 2, 4B, 5C, 6A and 6C and S7 Fig). Hence, the interaction between CEH-60 and UNC-62 drives both vitellogenesis and cuticle permeability in a way that originates from the primary lipid-metabolizing organ in C. elegans, the intestine.

**Discussion**

Studying the genetics of animal reproduction aids in understanding how species integrate age-, sex-, and environmentally relevant signals to maximize their reproductive fitness. From a fundamental, conceptual perspective, the successful production of high-quality gametes is thus a manifestation of an animal’s ability to manage a multi-tissue, multi-input type of complexity towards a very focused effect. C. elegans is a small and fast-reproducing model organism in which the synthesis of enormous amounts of yolk proteins provides an ideal readout to address such questions and unveil fundamental genetics of reproduction [3,6,9,11,41].

In earlier work, we identified the three amino acid loop extension (TALE) PBC-class Hox cofactor CEH-60 as a potential molecular gateway to kick-starting yolk protein production in C. elegans [3]. Its then-reported expression in a single neuron pair—here identified as the
olfactory AWC neurons (S2A Fig)—incentivized us to unveil CEH-60’s precise role in organismal control of intestinal yolk production. We found that CEH-60, in line with what is known for other TALE proteins [15,42–46], influences a range of physiological processes, including cuticle permeability, stress resistance, and the aforementioned vitellogenesis. Our finding that ceh-60 is additionally expressed in the pharynx and L4/adult intestine (Fig 1, S2B and S2C Fig) is corroborated by available genome-wide tissue-specific profiling data indeed revealing ceh-60 transcripts in amphid neurons, intestine, and pharynx [47,48]. We pinpointed the intestine as the site of CEH-60’s vitellogenesis-regulating function, a finding also recently reported by Robert Dowen and thus strengthening our conclusion [12]. Because CEH-60 is not dedicated to yolk production only, we asked how it might be recruited to this function.

Hox cofactors, and particularly TALE proteins, are able to regulate a myriad of physiological processes, not only those limited to development or even those requiring the action of a Hox protein [45,49–52]. PBC-class proteins interact with MEIS-class transcription factors, of which there is only one representative in C. elegans: UNC-62. The PBX/MEIS interaction is widely conserved and well documented in vertebrates (reviewed in [53]). CEH-60/PBX and UNC-62/MEIS have also been studied in Drosophila melanogaster, in which they are named extradenticle (EXD) and homothorax (HTH) [28–30], indicative of an evolutionary conservation in the protostome lineage. In C. elegans, the only other PBC-class proteins, CEH-20 and CEH-40, interact with UNC-62 to function in hypodermal and neuronal development [14,15]. UNC-62 is known to additionally function in the C. elegans intestine as an activator of vitellogenesis, without knowledge of a corresponding PBX-like cofactor. Here, we identified this elusive vitellogenesis-regulating partner of UNC-62 as CEH-60 and showed that its function depends on the ability to interact with UNC-62 in the intestine (Figs 2 and 3). Via independent methods, a recent study by Robert Dowen came to the same conclusions [12]. In his work, Dowen showed that CEH-60 shares specific DNA-binding sites with UNC-62, including the promoters of the vit genes, and that UNC-62 and CEH-60 co-immunoprecipitate. Dowen illustrates that the CEH-60::UNC-62 heterodimer acts as a metabolic switch to mediate organismal homeostasis from the intestine. It does so by activating reproductive programs, such as

Fig 7. Dysfunctional CEH-60 causes a cuticle that is hyperpermeable to endogenous ROS. (A) In supernatants collected from ceh-60(lst466) animals, the intensity of Amplex Red absorbance is much higher than in that of wild-type animals. Values are normalized to control. N = 6. (B) Endogenous ROS production measured with the in vivo genetic ROS sensor RoGFP2, indicated as the ratio of oxidized over reduced GFP, is the same in ceh-60 and wild-type animals. Error bars, SEM; *** p < 0.001. N = 12. Underlying data are available in S1 Data. GFP, green fluorescent protein; NS, not significant; Ox/red, ratio of oxidized over reduced GFP; RoGFP2, reduction-oxidation sensitive green fluorescent protein; ROS, reactive oxygen species.

https://doi.org/10.1371/journal.pbio.3000499.g007
vitellogenesis, and repressing somatic maintenance programs, such as innate immunity and life span. He additionally shows that at least one other transcription factor, PQM-1, is involved in co-regulating CEH-60:UNC-62 action. These findings, together with our own work, establish CEH-60/PBX as a linchpin of metabolism and development. The study by Dowen supports the main conclusions from our own work, which independently shows that CEH-60 and UNC-62 act together as powerful regulators of the mobilization of the main intestinal resources: yolk proteins and lipids.

Our study adds further depth by spatiotemporally defining the site of the interaction between CEH-60 and UNC-62 in the intestinal nuclei of adult animals (Fig 3), coinciding with the time and place of vitellogenesis. Furthermore, we show that the PBC domain is essential for interaction with UNC-62, activation of vitellogenesis, and the building of an impermeable cuticle (Figs 2C, 4B, 5 and 6C). While we cannot rule out the possibility that the PBC domain of CEH-60 may support DNA binding or other protein interactions besides the one with UNC-62, our PBC-truncated version of CEH-60 still includes the homeodomain and 16-residue C-terminal domain necessary for DNA binding in PBC-class TALE proteins [13], and no other PBC-dependent interactions besides those with UNC-62/MEIS/PBX regulating protein (PREP) are known [53].

While ceh-60 expression increases enormously at the start of vitellogenesis (S8 Fig, [3]), our work provides plenty of evidence for other functions, most notably in epicuticle structure and permeability (Figs 4–6). Similar to our observations for ceh-60 (Figs 4 and 5), other mutants with compromised (epi)cuticle integrity are more permeable to several compounds, including exogenous ROS or fluorescent dyes [34,54], and have difficulty staying in contact with a partner during mating [36]. Many of the known epicuticle-disrupting genes code for glycosyltransferase enzymes or other direct constituents of the (epi)cuticle, while CEH-60 is, to our knowledge, the first transcription factor to affect cuticle permeability.

It is not likely that CEH-60 works directly in the hypodermal cells, as no expression in those cells was observed. We cannot rule out the possibility that hypodermal expression may be too weak to detect, as epidermal and seam cells form large syncytia [35] and the CEH-60::GFP signal may be too diffuse. Still, CEH-60’s interaction partner, UNC-62, is known to be involved in cuticle synthesis. Among genes differentially expressed after unc-62 RNAi, 41 of 115 were collagens, and UNC-62 interacts with several transcription factors that function in the proper development of hypodermal cells and/or cuticle, corroborating the role of CEH-60’s interaction partner in cuticle development [16]. However, unlike CEH-20, CEH-40, and UNC-62, CEH-60 does not obviously affect the development of the seam cells (S3 Fig) that are likely responsible for synthesis of the cuticle [55].

Instead, we here find that the intestinal function of CEH-60 is not limited to the regulation of vitellogenesis, but includes the formation of an impermeable cuticle. While our reporters revealed no or very little intestinal expression before the L4 stage (Fig 1), single-cell sequencing data suggest CEH-60 transcripts to be present in the larval (at least L2) intestine [47]. We also observed oscillating ceh-60 expression during development, in synchrony with the molting cycle (S8 Fig). These whole-mount data may reflect the cyclic need for epicuticle building but lack finite spatial information. To our knowledge, synthesis of the epicuticle has not been studied in detail in C. elegans, but it has been suggested that the pharyngeal gland cells secrete the surface coat of other nematode species [56,57]. In C. elegans, defects in pharyngeal muscle structure have been shown to lead to deformed pharyngeal gland cells [58], which might also be the case in ceh-60 mutants. Pharyngeal ceh-60 expression, however, could not restore cuticle impermeability in mutants (Fig 4B), indicating that this is an unlikely scenario for CEH-60.

Instead, we found that it is intestinal CEH-60 that affects the morphology and permeability of the lipid-rich epicuticle (Figs 4B and 5C). The obvious hypothesis would be that it does so...
through its influence on lipid homeostasis. The lower abundance of some lipid-metabolizing enzymes observed in *ceh-60* mutants (Table 1) may simply reflect the inability of the intestine to mobilize lipids in response to a hypodermal need. Expression of *ceh-60* is likely not just a signal to start the production of yolk proteins, but a more general cue to start mobilization of lipids to either end: i.e., yolk production or epicuticle assembly.

Supporting the hypothesis of CEH-60 being involved in intestine-to-hypodermis lipid mobilization is the discovery of hypodermal micro-RNAs (miRNAs) that, in *C. elegans*, signal the start of intestinal mobilization of fat for yolk protein production in a manner that ultimately depends on intestinal activation of UNC-62 [11]. Why the signal originates in the hypodermis and how it is activated is currently unknown. It is, however, clear that in cuticle construction as well as in yolk production, there must be coordinated communication between the hypodermis and the intestine and that the intestinal interaction of CEH-60/PBX and UNC-62/MEIS is a central target in these processes. This knowledge provides a framework to continue adding to our understanding of the multi-tissue, multi-input complexity of reproductive control.

The increased cuticle permeability may add to the appeal of *ceh-60* mutants as a tool for drug screening. While *C. elegans* is used for, e.g., toxicity screens [59–61], its rigid and impermeable cuticle has been a noted disadvantage of this otherwise efficient and low-cost model organism [54,62,63]. A number of *C. elegans* *bus* (bacterially un-swollen) mutants have been proposed for toxicity screening precisely because of their increased cuticle permeability, but their general fitness is far from ideal, which questions practical applicability [32]. Mutants for *ceh-60* could provide an alternative to wild-type nematodes or *bus* mutants, as their overall fitness, as measured by life span and reproductive potential, does not seem to be impaired [3,12]. A note of caution is due, however, because the pharyngeal and neuronal roles of CEH-60 are yet to be described. In our hands, *ceh-60* mutants are easily discernible from wild-type animals but are overall healthier than *bus* mutants.

In other organisms, PBX and MEIS, as well the interaction between the two, are conserved [28–30,53]. They have been studied mostly with regard to neuronal development [46,64] and body plan patterning (reviewed in [65]), although evidence is mounting for the involvement of TALE proteins (PBX, MEIS, and PREP) in lipid homeostasis and metabolism (reviewed in [66]). It is well established that the ancient co-operation between PBX and MEIS reaches beyond their initially discovered roles as mere Hox cofactors. The functions of CEH-60/PBX and UNC-62/MEIS in *C. elegans* vitellogenesis and epicuticle formation provide the groundwork to expand on their scarcely investigated involvement in lipid homeostasis in other organisms as well.

**Materials and methods**

**Strains and plasmids**

*C. elegans* strains were grown under standard conditions [67], fed with *Escherichia coli* OP50, and raised at 20°C unless stated otherwise. For details on strain names and genotypes, see Table 2.

**Differential proteomics and data analysis**

Six populations of approximately 3,000 day 1 adult animals were sampled per condition. Samples were labeled with a TMTsixplex isobaric labeling set (Thermo Fisher Scientific, Waltham, MA) according to the manufacturer’s protocol. Worms were rinsed off plates in M9 buffer (3.0 g KH₂PO₄, 6.0 g Na₂HPO₄, 0.5 g NaCl, 1.0 g NH₄Cl in 1 L H₂O), washed three times, and lysed using RIPA buffer containing protease and phosphatase inhibitor. Protein homogenate
Table 2. List of *C. elegans* strains used in this study, including genotype and source.

| Strain name | Genotype | Source/reference |
|-------------|----------|-----------------|
| PY2417      | oyIs44 [odr-1p::rfp; lin-15(+)] | *Caenorhabditis* Genetics Center, University of Minnesota, MN, United States |
| JR667       | wIs51 [SCMP::gfp; unc-119(+)] | *Caenorhabditis* Genetics Center, University of Minnesota, MN, US |
| TJ356       | zIs356 [daf-16p::daf-16a::gfp; rol-6(sv1006)] | *Caenorhabditis* Genetics Center, University of Minnesota, MN, US |
| TP12        | kIs12[col-19::GFP] | *Caenorhabditis* Genetics Center, University of Minnesota, MN, US |
| JV10        | jrls10 [unc-119(+); rps-0p::roGFP2-Orp1] | Gift of Professor B. Braeckman, Ghent University, Belgium [40] |
| MGH167      | sid-1(qt9); abIs9 [vha-6p::sid-1::SL2::gfp] | Gift of Professor B. Braeckman, Ghent University, Belgium |
| UL2612      | ceh-60p::gfp | Gift of Professor I. A. Hope, University of Leeds, Leeds, United Kingdom [19] |
| DLS357      | ceh-60(ok1485) X | Gift of Professor R. Dowen, University of North Carolina at Chapel Hill, NC, US [12] |
| LSC897      | ceh-60[ls1466] X | [3] |
| LSC902      | vrl-1[lt539] IV | [3] |
| LSC1491     | lstEx790 [ceh-60p::ceh-60::gfp; unc-122p::DsRed] oyIs44 [odr-1p::rfp; lin-15(+)] | this study |
| LSC1487     | lstEx790 [ceh-60p::ceh-60::gfp; unc-122p::DsRed] | this study |
| LSC1501     | ceh-60[ls1466] X; lstEx855 [odr-1p::ceh-60; unc-122p::DsRed] | this study |
| LSC1798     | ceh-60[ls1466] X; lstEx950 [ceh-60p::ceh-60::gfp::unc-54 3′ UTR; unc-122p::DsRed] | this study |
| LSC1595     | ceh-60[ls1466] X; jrls10 [unc-119(+); rps-0p::roGFP2-Orp1] | this study |
| LSC1684     | ceh-60[ls1466] X; lstEx909 [ceh-60p::ceh-60::ceh-60 3′ UTR; unc-122p::DsRed] | this study |
| LSC1814     | lstEx956 [hsp-16.41p::MYC::unc-62::VN173::unc-54 3′ UTR; hsp-16.41p::HA::ceh-60::VC155::unc-54 3′ UTR; unc-122p::DsRed] | this study |
| LSC1815     | lstEx957 [hsp-16.41p::MYC::unc-62::VN173::unc-54 3′ UTR; hsp-16.41p::HA::ceh-60::VC155::unc-54 3′ UTR; unc-122p::DsRed] | this study |
| LSC1816     | lstEx958 [hsp-16.41p::MYC::unc-62::VN173::unc-54 3′ UTR; hsp-16.41p::HA::ceh-60::VC155::unc-54 3′ UTR; unc-122p::DsRed] | this study |
| LSC1817     | lstEx959 [hsp-16.41p::MYC::unc-62::VN173::unc-54 3′ UTR; hsp-16.41p::HA::ceh-60::VC155::unc-54 3′ UTR; unc-122p::DsRed] | this study |
| LSC1831     | lstEx1022 [unc-62p::MYC::unc-62::VN173::unc-54 3′ UTR; hsp-16.41p::HA::ceh-60::VC155::unc-54 3′ UTR; unc-122p::DsRed] | this study |
| LSC1832     | ceh-60[ls1466] X; wIs51 [SCMP::gfp; unc-119(+) | this study |
| LSC1833     | ceh-60[ls1466] X; lstEx1023 [elt-2p::ceh-60::unc-54 3′ UTR; unc-122p::DsRed] | this study |
| LSC1834     | ceh-60[ls1466] X; lstEx1024 [ceh-60p::ceh-60::DA20::unc-54 3′ UTR; unc-122p::DsRed] | this study |
| LSC1842     | ceh-60[ls1466] X; lstEx1031 [myo-2p::ceh-60::unc-54 3′ UTR; unc-122p::DsRed] | this study |

https://doi.org/10.1371/journal.pbio.3000499.t002
was then sonicated for 30 seconds on ice and centrifuged at 14,000g for 15 minutes. Protein concentration was determined using a BCA assay (Thermo Fisher Scientific, Waltham, MA), and 100 μg of protein extract per sample was prepared for protein digestion as noted in the manufacturer’s manual and digested with 2.5 μL trypsin per sample overnight at 37˚C. Peptide digests were labeled with TMT Label Reagent, after which the reaction was quenched with 8 μL 5% hydroxyamine. Pooled samples for each set comprising three wild-type and three mutant populations were prepared in a 1:1:1:1:1:1 protein concentration ratio and stored at −80˚C. Each pooled sample was separated into 10 fractions. The sample was injected into the UPLC reverse phase system (Acquity, Waters, Milford, MA). Peptides were separated in a C18 Column (XBridge Peptide BEH, 130 Å, 5 μm, 2.1 × 50 mm). The peptides were eluted at a flow rate of 1.5 mL/minutes using a 14-minute linear gradient as follows: 0–9.5 minutes, 60% mobile phase B (MP B); 9.5–10 minutes, 98% MP B. Mobile phase A (MP A): 98% H2O, 2% acetonitrile, and 0.1% formic acid, and 0.1% formic acid, but with pH adjusted to 2 by the addition of formic acid. Eluate was collected in 10 fractions. The collected fractions were vacuum dried and each fraction was resuspended in 25 μL of 98% H2O: acetonitrile: formic acid (98:2:0.1) solution for injection on the LC-MS. LC-MS analysis was performed on an Eksigent nanoAcquity LC-Ultra system (Waters, Milford, MA) connected to a LTQ Velos Orbitrap mass spectrometer (Thermo Fisher Scientific, Waltham, MA) through a flex nano ESI source (Thermo Fisher scientific, Waltham, MA). The equivalent of 1 μg of total protein of the digested sample was dissolved in 10 μL of 2% acetonitrile in HPLC-grade water. This sample was loaded on the trapping column (Pepmap C18 300 μm × 20 mm, Dionex, Thermo Fisher Scientific, Waltham, MA) with an isocratic flow of 2% acetonitrile in water with 0.1% formic acid at a flow rate of 5 μL/minute. After 2 minutes, the column-switching valve was switched, placing the pre-column online with the analytical capillary column, a Pepmap C18, 3 μm, 75 μm × 150 mm nano column (Dionex, Thermo Fisher Scientific, Waltham, MA). Separation was conducted using a linear gradient from 2% acetonitrile in water, 0.1% formic acid to 40% acetonitrile in water, 0.1% formic acid in 100 minutes. The flow rate was set at 400 nL/minute. The LTQ Orbitrap Velos was set up in a data-dependent MS/MS mode in which a full scan spectrum (350–2,000 m/z, resolution 60,000) was generated. Full scan spectra were followed by a maximum of five dual collision-induced dissociation (CID)/high-energy collision-induced dissociation (HCD) tandem mass spectra. We applied a dynamic exclusion time of 45 seconds.

Mixed HCD/CID spectra were analyzed in MaxQuant 1.6.3.4 (https://www.maxquant.org) [68] using 6-plex TMT as internal labels with a reporter mass tolerance of 0.003 Da, Oxidation (M) and Acetyl as variable modifications, and carbamidomethyl(C) as fixed modification, with a maximum of 5 modifications per peptide. Default orbitrap instrument settings were used and identified peptides were mapped to the UniProt C. elegans reference proteome (https://www.uniprot.org/proteomes/UP000001940). All other global parameters were used as default, except that match between runs was enabled. Data analysis and normalization was performed using R 3.5.1 (https://www.r-project.org) and Perseus 1.6.5.0 (https://maxquant.net/perseus) [69] to filter out peptides with valid measurement for at least 4 out of 12 channels and without all 6 of one 6-plex’s channels being empty. Proteins were only retained for analysis when at least 2 unique peptides were identified. In total, we identified 7,828 peptides corresponding to 1,749 proteins. Data normalization was performed as described in [70] using internal reference sampling and correcting for sample loading. Statistical significance was computed using a two-way ANOVA with Holm-Sidak multiple comparison correction. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE [71] partner repository with the dataset identifier PXD013584.
RNAi experiments

RNAi clones were obtained from the Vidal RNAi library, with the exception of the *col-120* clone, which was obtained from the Ahringer library. All clones were sequence-verified before use. RNAi strains were grown in LB medium containing 100 µg/mL ampicillin and spread on nematode growth medium (NGM) plates supplemented with 50 µg/mL ampicillin and 1 mM isopropyl-b-D-thiogalactoside (IPTG, Sigma-Aldrich, St. Louis, MO). Animals were reared for at least two generations on the specified RNAi clone before conducting experiments.

Oil-red-O staining

Oil-red-O staining was performed as described in [72]. For each condition, at least 20 animals of the desired age were fixed for 1 hour in Modified Ruvkun’s Witches Brew (80 mM KCl, 20 mM NaCl, 10 mM EGTA, 5 mM spermidine, 15 mM Pipes [pH 7.4], and 25% [v/v] methanol, also containing 2% [w/v] formaldehyde). Afterwards, animals were washed in M9 physiological buffer solution and incubated for 15 minutes in 60% (v/v) isopropanol, followed by overnight incubation in the same solution containing 0.3% Oil-Red-O (Abcam, UK). Stained worms were washed with M9 buffer and imaged using a Axio Imager Z1 microscope (Zeiss, Germany). As per [73], red color intensity was quantified in a circular region of the same size in each animal, as marked in S1 Fig. Stain intensity was quantified in at least 5 animals per condition per time point. Data were analyzed using a two-way ANOVA with a Tukey post hoc test.

Localization: Cloning and microscopy

Localization constructs were created by cloning a promoter region of 3,464 bp 5’ to the start of the *ceh-60* gene and *ceh-60* cDNA into a modified pSM vector carrying a GFP reporter sequence preceded by an SL2 trans-splicing site (kindly provided by C. Bargmann, Rockefeller University, New York, NY). The promoter and cDNA sequences were inserted right before the SL2 site using NEBuilder HiFi DNA Assembly Master Mix (New England BioLabs, Ipswich, MA) and transformed into DH5-alpha competent cells. Purified plasmid DNA was microinjected into the syncytial gonad of young adult worms at 25 ng/µL together with a co-injection marker *unc-122p::DsRed* at 50 ng/µL. Localization strains were mounted on 2% agarose pads, anesthetized with 1 mM tetramisole, and visualized with a confocal FluoView1000 microscope (Olympus, Japan) or a DM6 B microscope (Leica, Germany).

BiFC

BiFC was performed as described in [74]. Plasmids pCE-BiFC-VN173 and pCE-BiFC-VC153 were provided by Addgene (deposited there by the lab of Chang-Deng Hu). *unc-62* and *ceh-20* or *ceh-60* cDNA were cloned into pCE-BiFC-VN173 and pCE-BiFC-VC153, respectively, behind a heat-shock promoter using NEBuilder HiFi DNA Assembly Master Mix (New England BioLabs, Ipswich, MA) and transformed into DH5-alpha competent cells. Purified plasmid DNA of interacting BiFC partners was microinjected into the syncytial gonad of young adult worms at 25 ng/µL together with a co-injection marker *unc-122p::DsRed* at 50 ng/µL. Localization strains were mounted on 2% agarose pads, anesthetized with 1 mM tetramisole, and visualized with a confocal FluoView1000 microscope (Olympus, Japan) or a DM6 B microscope (Leica, Germany).
mM tetramisole. The fluorescence signal was quantified by measuring the average pixel intensity in 6 intestinal nuclei of at least 6 animals per condition in Fiji [75,76]. No heat shock treatment was administered to animals carrying endogenous promoters.

**Co-immunoprecipitation and western blotting**

For co-immunoprecipitation, at least 8 fully grown 90-mm NGM plates of each strain were used. Animals were rinsed off plates and washed in M9 buffer until all bacteria were removed. Per sample, a 500-μL worm pellet topped up with 1 mL of ice-cold lysis buffer (25 mM Tris-HCl [pH 7.5], 100 mM NaCl, 1 mM EDTA, 0.5% NP-40, 1 mM Na₃VO₄, 10 mM NaF, 1× complete protease inhibitor cocktail tablets) was snap-frozen in liquid nitrogen before sonicating thrice for 10 seconds with a SLPe Sonifier (Branson, Danbury, CT), snap-freezing, and thawing in room temperature water in between sonication steps. Worm lysates were incubated while agitating at 4˚C for 30 minutes and spun down for 30 minutes at 4˚C at 13,000 rpm, after which protein concentration was determined with a standard BCA assay. For co-immunoprecipitation, Pierce Anti-HA Agarose beads (Thermo Fisher Scientific, Waltham, MA) were equilibrated according to the manufacturer’s instructions. Beads were incubated at 4˚C overnight, with at least 4 mg of worm protein in 1 mL lysis buffer. After washing beads 5 times with lysis buffer, bead-bound proteins were eluted with XT sample buffer and boiled at 98˚C for 15 minutes, followed by western blot using anti-HA high affinity (Sigma-Aldrich, St. Louis, MO) and Myc Tag monoclonal (Thermo Fisher Scientific, Waltham, MA) antibodies as primary antibodies and polyclonal Rabbit-Anti-Rat IgG/HRP (Agilent, Santa Clara, CA) or polyclonal Goat-anti-Mouse IgG/HRP (Agilent, Santa Clara, CA) as secondary antibodies. Bands were visualized on a Bio-Rad Gel-Doc following staining in SuperSignal West Dura (Thermo Fisher Scientific, Waltham, MA) according to the manufacturer’s instructions.

**Yolk protein analysis**

Coomassie staining of worm protein extracts using SDS-PAGE was performed as described in [3,27]. Synchronous worm populations were grown until day 3 of adulthood, after which 50 animals per sample were picked into 15 μL of M9 buffer. A total of 15 μL of Laemmli sample buffer containing β-mercaptoethanol was added. Samples were incubated for 15 minutes at 70˚C, centrifuged at max speed for 5 minutes, and incubated at 95˚C for 5 minutes. A total of 15 μL of each sample was loaded on a 4%–12% bis-tris Criterion XT precast polyacrylamide gel (Bio-Rad, Hercules, CA) using XT MOPS as a running buffer. Gels were stained with Coomassie Brilliant Blue and destained with a 40% methanol, 10% acetic acid solution. Gel images were taken with a Bio-Rad Gel Doc (Bio-Rad, Hercules, CA). Identification of YP170, YP115, and YP88 bands is based on [7,27]. Quantification of yolk protein abundance was done by normalizing to total protein content in a lane using ImageLab 6.0. At least three populations were assayed for each condition. Data were analyzed using a one-way ANOVA with a Tukey post hoc test.

**Cuticle permeability and paralysis sensitivity assays**

Cuticle permeability assays using acridine orange were performed as described in [32]. In brief, day 1 adult animals were washed off plates, stained with 5 μg/mL acridine orange in M9 buffer for 15 minutes with gentle agitation, followed by three wash steps in M9 buffer. Worms were then mounted on 2% agarose pads and anesthetized with 1 mM tetramisole. Worms were imaged using a DM6 B microscope (Leica, Germany) with GFP filter set. Average fluorescence intensity for at least 20 worms per condition was quantified with Fiji [76] and analyzed using a one-way ANOVA with a Tukey post hoc test.
Sodium azide sensitivity tests were conducted as described in [77]. At least 20 young adult animals of each condition were transferred to unseeded NGM plates containing 0.5 mM sodium azide and monitored for paralysis every 30 minutes. When a worm did not respond to a gentle prod with a platinum wire, it was marked as paralyzed. Missing animals, or animals that died because of vulva rupture, were censored from the analysis. Two populations were assayed per condition. Data were analyzed using a two-way ANOVA with a Dunnett post hoc test.

**Staining of cuticle components with DiI and WGA**

The annuli of the cuticle were stained with DiI as described in [78]. Young adult animals were washed thrice in M9 buffer before staining with 30 μg/mL DiI dissolved in M9 for 12 hours. After staining, animals were washed once in M9 and imaged on a confocal microscope as described above to determine annuli width. The width of at least 10 annuli was determined per animal, and at least 7 animals per condition were imaged. The results were analyzed using a one-way ANOVA with a Tukey post hoc test. The cuticular surface was stained with rhodamine-conjugated WGA as described in [77,79]. Young adult animals were washed thrice in M9 buffer to remove residual bacteria and afterwards incubated in 200 μg/mL rhodamine-conjugated WGA dissolved in M9. After staining, animals were washed four times in M9 and imaged as described above. At least 12 animals were imaged per condition.

**TEM**

Preparation of TEM samples was performed as described in [80]. Day 1 adult animals were rinsed off culture plates and fixed in cold glutaraldehyde (2%, pH 7.3) with 50 mM Na-cacodylate and 150 mM saccharose, followed by fixation in 2% (w/v) osmium tetroxide. Fixed worms were dehydrated in an acetone series and embedded in araldite. Semi-thin sections of 1 μm cut with a Reichert Ultracut E microtome (Ametek, Berwyn, PA) were stained with methylene blue and viewed in a DM300 light microscope (Leica, Germany) for orientation. Double-stained 70-nm thin sections were visualized using an EM900 transmission electron microscope (Zeiss, Germany).

**Mating contact assays**

Mating contact assays were performed as described in [36]. Four day 1 adult hermaphrodite wild-type or mutant animals together with 8 day 1 adult wild-type males were picked onto a 55-mm NGM plate, seeded with a single 25-μL drop of OP50 *E. coli* grown in LB medium, creating a 10-mm circular spot of bacteria. After an adjustment period of at least 1 hour, the mating occupancy of each hermaphrodite was scored as either 1 (in contact with a male) or 0 (not in contact). Mating scores were summed up for each of the 4 hermaphrodites. This measurement was repeated every minute for 25 minutes, generating a total mating score indicative of the amount of time animals had spent in mating contact. At least three populations were assayed for each condition. Results were analyzed using a one-way ANOVA with a Dunnett post hoc test.

**Oxidative stress assays**

Oxidative stress survival assays were performed essentially as described in [81]. Per condition, at least 3 populations consisting of approximately 20 day 1 adult animals were incubated in 500 μL M9 buffer solution supplemented with H₂O₂ (Sigma–Aldrich, St. Louis, MO) to a final concentration of 5 mM in a 24-well plate. Survival of each animal was assayed every hour by
checking for movement or pharyngeal pumping. Missing animals, or animals that died because of vulva rupture, were censored in the analysis. Data were analyzed using a two-way ANOVA with a Dunnett post hoc test.

Quantification of in vivo ROS permeation and redox state
Measurement of ROS permeation was performed using the Amplex Red hydrogen peroxide kit (Thermo Fisher Scientific, Waltham, MA) according to the manufacturer’s instructions and modified for use in *C. elegans* as described in [72]. A 50-μL worm pellet of day 1 adult animals was suspended and washed in the manufacturer’s reaction buffer and subsequently incubated while rotating in the dark in Amplex Red working solution for 1 hour. The absorbance intensity of the supernatant, as a measure of permeated ROS, was measured in a Greiner flat-bottom 96-well plate using a Tecan Infinite M200 plate reader with excitation and emission wavelengths set to 550 and 590 nm, respectively. At least 6 populations were measured per condition. Means were compared using a Student *t* test.

Measurement of in vivo redox state was performed as previously described [40,82,83]. *ceh-60* mutant animals were crossed with the JV10 RoGFP marker strain carrying the *rps-0p::roGF-P2-Orp1* redox-sensitive *roGFP2* transgene under the ubiquitous ribosomal *rps-0* promoter. Day 1 adult wild-type and *ceh-60* mutant animals carrying the *roGFP2* transgene were washed in physiological buffer and diluted to a concentration of 10 worms per μL. A total of 100 μL of worm suspension was added to each well of a flat-bottom 96-well plate (Sigma-Aldrich, St. Louis, MO) and fluorescence was recorded for 1 hour with excitation filters for oxidized (405 nm) and reduced (490 nm) *roGFP2* and an emission wavelength at 535 nm using an Infinite M200 plate reader (Tecan, Switzerland). At each time point, the ratio of oxidized over reduced GFP was calculated as described in [84] and averaged over 1 hour. At least 12 populations of approximately 1,000 worms each were tested per condition. Means were compared using a Student *t* test. To test the sensitivity of the assay and the redox shift upon encounter of an exogenous stressor, *H₂O₂* was added to a final concentration of 5 mM, and measurements of oxidized and reduced RoGFP intensity were taken for another 140 cycles.

Supporting information
S1 Fig. Lipids accumulate in the intestine of *ceh-60* adults but are less abundant in their embryos. (A) Quantification of Oil-red-O staining shows that lipids accumulate faster in the intestine of adult *ceh-60* (●) animals when compared with the wild type (■), a difference that becomes significant from day 3 of adulthood onwards. Staining intensity is relative to wild-type day 1, which is set at 100. Error bars: SEM, and **** *p* < 0.0001. *N* ≥ 5 for each time point. Underlying data are available in S1 Data. (B) Representative images of day 3 adults stained with Oil-red-O show that the intestinal region is more intensely stained in *ceh-60* animals. Yellow circles indicate regions used for intestinal fat quantification. Arrows indicate embryos inside the adult hermaphrodite. Scale bar, 200 μm. (TIF)

S2 Fig. *ceh-60* is expressed in AWC neurons, pharyngeal muscle, and intestine. (A) Expression of *ceh-60* (green, *ceh-60p::ceh-60::gfp*) and *odr-1* (red, *odr-1p::rfp* is AWC-specific [85]) overlaps, showing that *ceh-60* is expressed in the AWC neurons. (B,C) Bright-field and GFP images of intestinal (**) and pharyngeal (arrow) expression of *ceh-60* in strains carrying (B) a *ceh-60p::ceh-60::gfp* fosmid or (C) a *ceh-60p::ceh-60::SL2::gfp* construct. While pharyngeal expression is visible throughout life, its localization does not appear to be exclusively nuclear. Neuronal expression is always visible, marked with dotted circles. Scale bars, 20 μm. GFP,
green fluorescent protein.  

**S3 Fig. CEH-60 is not necessary for control of seam cell division.** (A) Overlaid bright-field and fluorescence images of wild-type and ceh-60 L4 animals carrying an integrated seam cell gfp marker (SCMP::gfp). Each animal has 16 seam cells visible on either side of the body. Scale bars, 100 μm. (B) Graph indicating the number of seam cells in wild-type, ceh-60 mutant and unc-62 RNAi-treated animals (positive control). There is no significant difference between ceh-60 and wild-type animals, while unc-62 RNAi-treated animals show modest seam cell hyperplasia. Error bars: SEM, and ***p < 0.001. N ≥ 6. Underlying data are available in **S1 Data.**
gfp, green fluorescent protein; RNAi, RNA interference.  

**S4 Fig. Knockdown of collagen genes does not cause cuticle hyperpermeability.** Performing RNAi knockdown of col-106 or col-120 does not change the permeability of animals to acridine orange. Fluorescence intensity is relative to the wild type and is shown on a logarithmic scale. Error bars: SEM, and NS = not significant. N ≥ 36. Underlying data are available in **S1 Data.**

**S5 Fig. Characterization of different cuticular components with Dil, WGA, and col-19::gfp reporter.** (A) Dil staining of the annuli of the wild type, ceh-60(lst466), and ceh-60(ok1485) shows that there is no difference in annuli morphology. Annuli width is relative to the average value for wild type, set as 100. Scale bar, 20 μm. Error bars: SEM, and NS = not significant. N ≥ 7. (B) Rhodamine-conjugated WGA stains ceh-60 mutant animals but not wild types. Scale bar, 200 μm. Graph scale, logarithmic; error bars: SEM, and ****p < 0.0001. N ≥ 12. Underlying data for panel A and B are available in **S1 Data.** (C) Visualization of the cortical layer of the cuticle with col-19::gfp marker. Scale bar, 10 μm. Dil, 1,19-dioctadecyl-3,3,39,39-tetramethylindocarbocyanine perchlorate; WGA, wheat germ agglutinin.  

**S6 Fig. Intestinal unc-62 knockdown increases susceptibility to oxidative stress.** Upon knockdown of unc-62 (○) in the intestine-specific RNAi strain MGH167, animals become more susceptible to oxidative stress than empty vector–treated animals (●), although the effect is slightly less pronounced than in ceh-60(lst466) mutants (■). Error bars: SEM. ***p < 0.001. N ≥ 4. Underlying data are available in **S1 Data.**

**S7 Fig. CEH-60 affects sensitivity to sodium azide.** Sodium azide sensitivity as measured by fraction of worms moving during incubation in 0.5 mM NaN₃ is lower in ceh-60 mutant animals than in control animals. This defect is rescued by intestinal expression of ceh-60 (elt-2p::ceh-60), but not by expression of ceh-60 with a truncated PBC-interaction domain (ceh-60p::ceh-60(ΔPBC)). Error bars indicate SEM. **p < 0.01, ****p < 0.0001. N = 2. Underlying data are available in **S1 Data.** PBC, pre–B cell leukemia.  

**S8 Fig. ceh-60 mRNA levels oscillate with molting.** mRNA abundance of ceh-60 (■) cycles during development, apparently peaking each time at the end of a molt, as deduced from the lin-42 expression profile. Molts are recognized as local minima in lin-42 expression (gray line) [86]. Adopted under creative commons license 4.0 from [3], where it is also shown that ceh-60 expression dramatically increases during the final larval molt (as indicated by the dashed
arrow). Underlying data are available in S1 Data.

(TIF)

S9 Fig. Ratio of oxidized over reduced RoGFP2 before and after addition of exogenous stress. Wild-type (—) and ceh-60 (gray line) animals show no difference in redox state when observed under unstressed conditions, but when an exogenous stressor in the form of 5 mM H2O2 is added after 20 cycles (indicated by arrow), their ratio of oxidized/reduced RoGFP2 increases more than in wild-type animals. N ≥ 8. Underlying data are available in S1 Data. RoGFP2, reduction-oxidation sensitive green fluorescent protein.

(TIF)

S1 Data. Data underlying Figs 2–7, S1 Fig and S3–S9 Figs.

(XLSX)

Acknowledgments

We wish to thank An Vandoren and Prof. Johan Billen (KU Leuven, Belgium) for TEM expertise, Elke Vandewyer and Wahab Al-Ani (KU Leuven, Belgium) for technical assistance, Liesbeth Van Rompay (KU Leuven, Belgium) for collecting the proteomics samples, and Karin Schildermans (UAntwerp, Belgium) for collection of LC-MS data. We are grateful to Prof. Braeckman (Ghent University, Belgium), Prof. Hope (University of Leeds, UK), and Prof. Dowen (University of North Carolina at Chapel Hill, US) for sharing strains. Some strains were provided by the Caenorhabditis genetics center (University of Minnesota, US).

Author Contributions

Conceptualization: Pieter Van de Walle, Geert Baggerman, Liliane Schoofs, Liesbet Temmerman.

Formal analysis: Pieter Van de Walle, Peter Askjaer.

Funding acquisition: Pieter Van de Walle, Liliane Schoofs, Liesbet Temmerman.

Investigation: Pieter Van de Walle, Ellen Geens, Francisco José Naranjo-Galindo, Peter Askjaer, Liesbet Temmerman.

Methodology: Pieter Van de Walle, Ellen Geens, Geert Baggerman, Francisco José Naranjo-Galindo.

Supervision: Liliane Schoofs, Liesbet Temmerman.

Visualization: Pieter Van de Walle.

Writing – original draft: Pieter Van de Walle, Liesbet Temmerman.

Writing – review & editing: Pieter Van de Walle, Ellen Geens, Geert Baggerman, Peter Askjaer, Liliane Schoofs, Liesbet Temmerman.

References

1. Bownes M, Lineruth M, Mauchline D. Egg production and fertility in Drosophila depend upon the number of yolk-protein gene copies. Mol Genet genomics. 1991; 228: 324–327. https://doi.org/10.1007/BF00282485 PMID: 1909423

2. Yilmaz O, Patinote A, Nguyen T, Corn E, Pineau C, Bobe J. Genome editing reveals reproductive and developmental dependencies on specific types of vitellogenin in zebrafish (Danio rerio). Molecular Reproduction and Development. 2019; 86: 1168–1188. https://doi.org/10.1002/mrd.23231 PMID: 31380595
3. Van Rompay L, Borghgraef C, Beets I, Caers J, Temmerman L. New genetic regulators question relevance of abundant yolk protein production in C. elegans. Sci Rep. 2015; 1:1–16. https://doi.org/10.1038/srep16381 PMID: 26553710

4. Grant B, Hirsh D. Receptor-mediated endocytosis in the Caenorhabditis elegans oocyte. Mol Biol Cell. 1999; 10: 4311–4326. https://doi.org/10.1091/mbc.10.12.4311 PMID: 10588660

5. Kimble J, Sharrock J. Tissue-specific synthesis of yolk proteins in Caenorhabditis elegans. Dev Biol. 1982; https://doi.org/10.1016/0012-1606(83)90322-6

6. MacMorris M, Browerman S, Greenspoon S, Lea K, Madej C, Blumenthal T, et al. Regulation of vitellogenin gene expression in transgenic Caenorhabditis elegans: short sequences required for activation of the vit-2 promoter. Mol Cell Biol. 1992; 12: 1652–1662. https://doi.org/10.1128/mcb.12.4.1652 PMID: 1549118

7. DePina AS, Iser WB, Park S-S, Maudsley S, Wilson M a, Wolkow C a. Regulation of Caenorhabditis elegans vitellogenesis by DAF-2/IIIS through separable transcriptional and posttranscriptional mechanisms. BMC Physiol. BioMed Central Ltd; 2011; 11: 11. https://doi.org/10.1186/1472-6793-11-11 PMID: 21746693

8. Jordan JM, Hibshman JD, Webster AK, Fry AL, Hubbard EJA, Baugh LR, et al. Insulin/IGF signaling and vitellogenin provisioning mediate interregenerational adaptation to nutrient stress. Curr Biol. Elsevier Ltd.; 2019; 29: 2380–2388.e5. https://doi.org/10.1016/j.cub.2019.05.062 PMID: 31280992

9. Perez MF, Lehner B. Vitellogenins—Yolk gene function and regulation in Caenorhabditis elegans. Front Physiol. 2019; 10. https://doi.org/10.3389/fphys.2019.01067 PMID: 31551797

10. Goszczyński B, Capitan V V., Danielson AM, Lancaster BR, McGhee JD. A 44 bp intestine-specific hermaphrodite-specific enhancer from the C. elegans vit-2 vitellogenin gene is directly regulated by ELT-2, MAB-3, FKH-9 and DAF-16 and indirectly regulated by the germline, bydaf-2/insulin signaling and by the TGF-β/Smad. Dev Biol. Elsevier; 2016; 413: 112–127. https://doi.org/10.1016/j.ydbio.2016.03.031 PMID: 26963674

11. Downen RH, Breen PC, Tuullius T, Conery AL, Ruvkun G. A microRNA program in the C. elegans hypodermis couples to intestinal mTORC2/PQM-1 signaling to modulate fat transport. Genes Dev. 2016; 30: 1515–1528. https://doi.org/10.1101/gad.283895.116 PMID: 27401555

12. Downen RH. CEH-60/PBX and UNC-62/MEIS coordinate a metabolic switch that supports reproduction in C. elegans. Dev Cell. Elsevier Inc.; 2019; 1–16. https://doi.org/10.1016/j.devcel.2019.03.002 PMID: 30956009

13. Bürglin TR. Analysis of TALE superclass homeobox genes (MEIS, PBC, KNOX, Iroquois, TGF) reveals a novel domain conserved between plants and animals. Nucleic Acids Res. 1997; 25: 4173–4180. https://doi.org/10.1093/nar/25.21.4173 PMID: 9336443

14. Zheng C, Jin FQ, Chaffie M. Hox proteins act as transcriptional guarantors to ensure terminal differentiation. Cell Rep. 2015; 13: 1343–1352. https://doi.org/10.1016/j.celrep.2015.10.044 PMID: 26547238

15. Hughes S, Brabin C, Appleford PJ, Woollard A. CEH-20/Pbx and UNC-62/Meis function upstream of rnt-1/Runx to regulate asymmetric divisions of the C. elegans stem-like seam cells. Biol Open. 2013; 2: 718–727. https://doi.org/10.1242/bio.20134549 PMID: 23862020

16. van Nostrand EL, Sánchez-Blanco A, Wu B, Nguyen A, Kim SK. Roles of the developmental regulator unc-62/Homothorax in limiting longevity in Caenorhabditis elegans. PLoS Genet. 2013; 9. https://doi.org/10.1371/journal.pgen.1003325 PMID: 23618775

17. Jiang Y, Shi H, Liu J. Two Hox cofactors, the Meis/Hth homolog UNC-62 and the Pbx/Exd homolog Ceh-20, function together during C. elegans postembryonic mesodermal development. Dev Biol. Elsevier Inc.; 2009; 334: 535–546. https://doi.org/10.1016/j.ydbio.2009.07.034 PMID: 19643105

18. McGhee JD, Sleumer MC, Bilenky M, Wong K, McKay SJ, McGhee JD, et al. The ELT-2 GATA-factor and the global regulation of transcription in the C. elegans intestine. Dev Biol. 2007; 302: 627–645. https://doi.org/10.1016/j.ydbio.2006.10.024 PMID: 17113066

19. Reece-Hoyes JS, Shingles J, Dupuy D, Grove C a, Walhout AJM, Vidal M, et al. Insight into transcription factor gene duplication from Caenorhabditis elegans promotermotived expression patterns. BMC Genomics. 2007; 8: 27. https://doi.org/10.1186/1471-2164-8-27 PMID: 17243537

20. Mohsen AA, Navarette B, Vockley J. Identification of Caenorhabditis elegans isovaleryl-CoA dehydrogenase and structural comparison with other acyl-CoA dehydrogenases. Mol Genet Metab. 2001; 71: 126–137. https://doi.org/10.1006/mgen.2001.3183 PMID: 11368484

21. Watson E, Olin-sandoval V, Hoy MJ, Li C, Louissie T, Yao V, et al. Metabolic network rewiring of propionate flux compensates vitamin B12 deficiency in C. elegans. Elife. 2016; 1–21. https://doi.org/10.7554/eLife.17670 PMID: 27383050
22. Mahajan-miklos S, Tan M, Rahme LG, Ausubel FM. Molecular mechanisms of bacterial virulence elucidated using a *Pseudomonas aeruginosa–Caenorhabditis elegans* pathogenesis model. Cell. 1999; 96: 47–56. https://doi.org/10.1016/s0092-8674(00)80958-7 PMID: 9989496

23. Pees B, Kloock A, Nakad R, Barbosa C, Dierking K. Enhanced behavioral immune defenses in a *C. elegans* C-type lectin-like domain gene mutant. Dev Comp Immunol. Elsevier Ltd; 2017; 74: 237–242. https://doi.org/10.1016/j.devimm.2017.04.021 PMID: 2849858

24. Van Der Hoeven R, McCallum KC, Cruz MR, Garsin DA. Ce-Duox 1/BLI-3 generated reactive oxygen species trigger protective SKN-1 activity via p38 MAPK signaling during infection in *C. elegans*. PLoS Pathog. 2011; 7. https://doi.org/10.1371/journal.ppat.1002453 PMID: 2226003

25. Tullet JM a, 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: 1025–1038. https://doi.org/10.1016/j.cell.2008.01.030 PMID: 18358814

26. Sarov M, Murray JI, Schanzke K, Pozniakovski A, Niu W, Angermann K, et al. A genome-scale resource for *in vivo* tag-based protein function exploration in *C. elegans*. Cell. Elsevier; 2012; 150: 855–866. https://doi.org/10.1016/j.cell.2012.08.001 PMID: 22901814

27. Sornda T, Ezcurra M, Kern C, Galimov ER, Au C, de la Guardia Y, et al. Production of YP170 vitellogenins promotes intestinal senescence in *C. elegans*. Journals Gerontol Ser A. 2019; https://doi.org/10.1093/gerona/glz067 PMID: 30854561

28. Abu-Shaar M, Ryoo HD, Mann RS. Control of the nuclear localization of Extradenticle by competing nuclear import and export signals. Genes Dev. 1999; 13: 935–945. https://doi.org/10.1101/gad.13.8.935 PMID: 10215621

29. Rieckhof GE, Casares F, Ryoo HD, Abu-Shaar M, Mann RS. Nuclear translocation of Extradenticle requires *homothorax*, which encodes an *Extradenticle*-related homeodomain protein. Cell. 1997; 91: 171–183. https://doi.org/10.1016/s0092-8674(00)80400-6 PMID: 9346325

30. Ryoo HD, Marty T, Casares F, Affolter M, Mann RS. Regulation of Hox target genes by a DNA bound *Homothorax/Hox/Extradenticle* complex. Development. 1999; 126: 5137–5148. https://doi.org/10.1016/s0092-8674(00)80958-7 PMID: 10529430

31. Stevens KE, Mann RS. A balance between two nuclear localization sequences and a nuclear export sequence governs Extradenticle subcellular localization. Genetics. 2007; 175: 1625–1636. https://doi.org/10.1534/genetics.106.066449 PMID: 17277370

32. Xiong H, Pears C, Woollard A. An enhanced *ce-homothorax* mutant. Dev Comp Immunol. Elsevier Ltd; 2017; 74: 237–242. https://doi.org/10.1016/j.devimm.2017.04.021 PMID: 2849858

33. Partridge FA, Tearle AW, Gravato-Nobre MJ, Schafer WR, Hodgkin J. The *C. elegans* glycosyltransferase BUS-8 has two distinct and essential roles in epidermal morphogenesis. Dev Biol. 2008; 317: 549–559. https://doi.org/10.1016/j.ydbio.2008.02.060 PMID: 18395708

34. Chisholm AD, Xu S. The *Caenorhabditis elegans* epidermis as a model skin. II: differentiation and physiological roles. Wiley Interdiscip Rev Dev Biol. 2012; 1: 879–902. https://doi.org/10.1002/wdev.77 PMID: 23539358

35. Partridge FA, Tearle AW, Gravato-Nobre MJ, Schafer WR, Hodgkin J. Glycosylation genes expressed in seam cells determine complex surface properties and bacterial adhesion to the cuticle of *Caenorhabditis elegans*. Genetics. 2011; 187: 141–155. https://doi.org/10.1534/genetics.110.122002 PMID: 20980242

36. Seehuus S-C, Norberg K, Gimsa U, Krekling T, Amdam G V. Reproductive protein protects functionally sterile honey bee workers from oxidative stress. Proc Natl Acad Sci. 2006; 103: 962–967. https://doi.org/10.1073/pnas.0502681103 PMID: 16418279

37. Fischer M, Regitz C, Kul R, Boll M, Wenzel U. Vitellogenins increase stress resistance of *Caenorhabditis elegans* after *Photorhabdus luminescens* infection depending on the steroid-signaling pathway. Microbes Infect. Elsevier Masson SAS; 2013; 15: 569–578. https://doi.org/10.1016/j.micinf.2013.05.002 PMID: 23727258

38. Chávez V, Mohri-Shiomi A, Maadani A, Vega LA, Garsin DA. Oxidative stress enzymes are required for DAF-16-mediated immunity due to generation of reactive oxygen species by *Caenorhabditis elegans*. Genetics. 2007; 176: 1567–1577. https://doi.org/10.1534/genetics.107.072587 PMID: 17483415

39. Braeckman BP, Smolders A, Back P, De Henau S. *In Vivo* Detection of Reactive Oxygen Species and Redox Status in *Caenorhabditis elegans*. Antioxid Redox Signal. 2016; 25: 577–592. https://doi.org/10.1089/ars.2016.6751 PMID: 27306519
41. Perez MF, Francesconi M, Hidalgo-carcedo C, Lehner B. Maternal age generates phenotypic variation in Caenorhabditis elegans. Nature. Nature Publishing Group; 2017; 552: 106–109. https://doi.org/10.1038/nature25012 PMID: 29816117

42. Dilorio P, Alexa K, Choe S, Etheridge L, Sagerström CG. TALE-Family homeodomain proteins regulate endodermal sonic hedgehog expression and pattern the anterior endoderm. Dev Biol. 2007; 304: 221–231. https://doi.org/10.1016/j.ydbio.2006.12.024 PMID: 17289013

43. Berkes CA, Bergstrom DA, Penn BH, Seaver KJ, Knopfler PS, Tapscott SJ. Pbx marks genes for activation by MyoD indicating a role for a homeodomain protein in establishing myogenic potential. Mol Cell. 2004; 14: 465–477. https://doi.org/10.1016/s1097-2765(04)00260-6 PMID: 15149596

44. Avila D, Helmcke K, Aschner M. The shape gland cell morphology. Genetics. 2011; 189: 885–897. https://doi.org/10.1534/genetics.111.132449 PMID: 21868069

45. Nelson FK, Riddle DL. Functional study of the Caenorhabditis elegans secretory-excretory system using laser microsurgery. J Exp Zool. 1984; 20: 493–496. PMID: 19290243

46. Raharjo WH, Ghai V, Dineen A, Bastiani M, Gaudet J. Cell Architecture: surrounding muscle cells shape gland cell morphology. Genetics. 2011; 189: 885–897. https://doi.org/10.1534/genetics.111.132449 PMID: 21868069

47. Dengg M, Meel JCA Van. Caenorhabditis elegans as model system for rapid toxicity assessment of pharmaceutical compounds. J Pharmacol Toxicol Methods. 2004; 50: 209–214. https://doi.org/10.1016/j.vascn.2004.04.002 PMID: 15519907

48. Leung MCK, Williams PL, Benedetto A, Au C, Helmcke KJ, Aschner M, et al. Caenorhabditis elegans: an emerging model in biomedical and environmental toxicology. Toxicol Sci. 2008; 106: 5–28. https://doi.org/10.1093/toxsci/kfn121 PMID: 18566021

49. Yang L, Sym M, Kenyon C. The roles of two transcription factors PREP, MEIS, and PBX in vertebrates. Dev Dyn. 2014; 24016 PMID: 23873833

50. Fritz J, Behm CA. CUTI-1: a novel tetraspan Protein involved in C. elegans CUTicle formation and epithelial integrity. PLoS ONE. 2009; 4: https://doi.org/10.1371/journal.pone.0005117 PMID: 19357781

51. Bird AF, Bonig I, Bacic A. A Role for the “Excretory” System in Secernentean Nematodes. J Nematol. 1988; 20: 493–496. PMID: 19290243

52. Casares F, Mann RS. Control of antennal versus leg development in Drosophila. Nature. 1996; 392: 723–726. https://doi.org/10.1038/33706 PMID: 9565034

53. Longobardi E, Penkov D, Mateos D, Florian G De, Torres M, Blasi F. Biochemistry of the tale transcription factors PREP, MEIS, and PBX in vertebrates. Dev Dyn. 2014; 59–73. https://doi.org/10.1002/dvdy.24016 PMID: 23873833

54. Fritz J, Behm CA. CUTI-1: a novel tetraspan Protein involved in C. elegans CUTicle formation and epithelial integrity. PLoS ONE. 2009; 4: https://doi.org/10.1371/journal.pone.0005117 PMID: 19357781

55. Frand AR, Russel S, Ruvkun G. Functional genomic analysis of C. elegans molting. PLoS Biol. 2005; 3: https://doi.org/10.1371/journal.pbio.0030312 PMID: 16122351

56. Berkes CA, Bergstrom DA, Penn BH, Seaver KJ, Knopfler PS, Tapscott SJ. Pbx marks genes for activation by MyoD indicating a role for a homeodomain protein in establishing myogenic potential. Mol Cell. 2004; 14: 465–477. https://doi.org/10.1016/s1097-2765(04)00260-6 PMID: 15149596
62. Johnstone I. Cuticle collagen genes expression in Caenorhabditis elegans. Trends Genet. 2000; 16: 21–27. https://doi.org/10.1016/s0168-9525(99)01857-0 PMID: 10637627

63. Giacomotto J, Ségalat L. High-throughput screening and small animal models, where are we? Br J Pharmacol. 2010; 160: 204–216. https://doi.org/10.1111/j.1476-5381.2010.00725.x PMID: 20423355

64. Choe S, Vlachakis N, Sagerstrom CG. Meis family proteins are required for hindbrain development in the zebrafish. Development. 2002; 595: 585–595.

65. Mallo M, Wellik DM, Deschamps J. Hox genes and regional patterning of the vertebrate body plan. Dev Biol. Elsevier Inc.; 2010; 344: 7–15. https://doi.org/10.1016/j.ydbio.2010.04.024 PMID: 20435029

66. Oriente F, Perruolo G, Cimmino I, Cabaro S, Liotti A, Longo M, et al. Prep1, a homeodomain transcrip-
84. Back P, De Vos WH, Depuydt GG, Matthijssens F, Vanfleteren JR, Braeckman BP. Exploring real-time in vivo redox biology of developing and aging Caenorhabditis elegans. Free Radic Biol Med. 2012; 52: 850–859. https://doi.org/10.1016/j.freeradbiomed.2011.11.037 PMID: 22226831

85. Lanjuin A, VanHoven MK, Bargmann CI, Thompson JK, Sengupta P. Otx/otd homeobox genes specify distinct sensory neuron identities in C. elegans. Dev Cell. 2003; 5: 621–633. https://doi.org/10.1016/S1534-5807(03)00293-4 PMID: 14536063

86. Jeon M, Gardner HF, Miller EA, Deshler J, Rougvie AE. Similarity of the C. elegans developmental timing protein LIN-42 to circadian rhythm proteins. Science (80-). 1999; 286: 1141–1146. https://doi.org/10.1126/science.286.5442.1141 PMID: 10550049