A terminal selector prevents a Hox transcriptional switch to safeguard motor neuron identity throughout life

Weidong Feng1,2, Yinan Li1,3, Pauline Dao1, Jihad Aburas1, Priota Islam4,5, Benayahu Elbaz6, Anna Kolarzyk6, André EX Brown4,5, Paschalis Kratsios1,2,3,7*

1Department of Neurobiology, University of Chicago, Chicago, United States; 2Committee on Development, Regeneration and Stem Cell Biology, University of Chicago, Chicago, United States; 3Committee on Neurobiology, University of Chicago, Chicago, United States; 4MRC London Institute of Medical Sciences, London, United Kingdom; 5Institute of Clinical Sciences, Imperial College London, London, United Kingdom; 6Department of Neurology, Center for Peripheral Neuropathy, University of Chicago, Chicago, United States; 7The Grossman Institute for Neuroscience, Quantitative Biology, and Human Behavior, University of Chicago, Chicago, United States

Abstract To become and remain functional, individual neuron types must select during development and maintain throughout life their distinct terminal identity features, such as expression of specific neurotransmitter receptors, ion channels and neuropeptides. Here, we report a molecular mechanism that enables cholinergic motor neurons (MNs) in the C. elegans ventral nerve cord to select and maintain their unique terminal identity. This mechanism relies on the dual function of the conserved terminal selector UNC-3 (Collier/Ebf). UNC-3 synergizes with LIN-39 (Scr/Dfd/Hox4-5) to directly co-activate multiple terminal identity traits specific to cholinergic MNs, but also antagonizes LIN-39’s ability to activate terminal features of alternative neuronal identities. Loss of unc-3 causes a switch in the transcriptional targets of LIN-39, thereby alternative, not cholinergic MN-specific, terminal features become activated and locomotion defects occur. The strategy of a terminal selector preventing a transcriptional switch may constitute a general principle for safeguarding neuronal identity throughout life.

Introduction

Every nervous system is equipped with distinct neuron types essential for different behaviors. Fundamental to nervous system function is the precise establishment and maintenance of neuron type-specific gene expression programs. Integral components of such programs are effector genes that encode proteins critical for neuronal function (e.g., neurotransmitter [NT] biosynthesis components, ion channels, NT receptors, neuropeptides) (Deneris and Hobert, 2014; Hobert, 2008; Hobert, 2011; Hobert, 2016). These effector genes, referred to as terminal identity genes herein, are expressed continuously, from development throughout life, in post-mitotic neurons in a combinatorial fashion (Hobert, 2008). Hence, it is the unique overlap of many effector gene products in a specific neuron type that determines its distinct terminal identity, and thereby function. However, the molecular mechanisms that select, in individual neuron types, which terminal identity genes should be expressed and which ones should be repressed are poorly defined. Understanding how neuron type-specific batteries of terminal identity genes are established during development and, perhaps most importantly, maintained throughout life represents one key step towards
understanding how individual neuron types become and remain functional. Providing molecular insights into this fundamental problem may also have important biomedical implications, as defects in terminal identity gene expression are associated with a variety of neurodevelopmental and neurodegenerative disorders (Deneris and Hobert, 2014; Shibuya et al., 2011; Imbrici et al., 2013; Sgadò et al., 2011).

Seminal genetic studies in multiple model systems revealed a widely employed molecular principle: neuron type-specific transcription factors (TFs) often coordinate the expression of ‘desired’ terminal identity genes with the exclusion of ‘unwanted’ terminal identity genes (Morey et al., 2008; Sagasti et al., 1999; Britanova et al., 2008; Cheng et al., 2004; Kala et al., 2009; Lopes et al., 2012; Mears et al., 2001; Nakatani et al., 2007). These TFs exert a dual role: they are not only required to induce a specific set of terminal identity features critical for the function of a given neuron type, but also to simultaneously prevent expression of molecular features normally reserved for other neuron types. Consequently, neurons lacking these TFs fail to acquire their unique terminal identity, and concomitantly gain features indicative of alternative identities. For example, mouse striatal cholinergic interneurons lacking Lhx7 lose their terminal identity and acquire molecular features indicative of GABAergic interneuron identity (Lopes et al., 2012). In midbrain neurons, removal of Gata2 results in loss of GABAergic identity and simultaneous gain of terminal identity features specific to glutamatergic neurons (Kala et al., 2009). However, the molecular mechanisms underlying the dual function of most neuron type-specific TFs remain poorly defined. How can the same TF, within the same cell, promote a specific identity and simultaneously prevent molecular features of alternative neuronal identities? In principle, the same TF can simultaneously operate as direct activator of neuron type-specific terminal identity genes and direct repressor of alternative identity genes (Lodato et al., 2014; Wyler et al., 2016). Another possibility is indirect regulation. For example, a neuron type-specific TF can prevent adoption of alternative identity features by repressing expression of an intermediary TF that normally promotes such features (Cheng et al., 2004). Other mechanisms involving TF competition for cell type-specific enhancers or cell type-specific TF-TF interactions have also been described (see Discussion) (Andzelm et al., 2015; Gordon and Hobert, 2015; Rhee et al., 2016; Thaler et al., 2002). It remains unclear, however, whether these mechanisms of action of neuron type-specific TFs are broadly applicable in the nervous system.

Although the aforementioned studies begin to explain how neurons select their terminal identity features during development (Morey et al., 2008; Sagasti et al., 1999; Britanova et al., 2008; Cheng et al., 2004; Kala et al., 2009; Lopes et al., 2012; Mears et al., 2001; Nakatani et al., 2007), the function of neuron type-specific TFs is rarely assessed during post-embryonic stages. Hence, the molecular mechanisms that maintain neuronal terminal identity features, and thereby neuronal function, are largely unknown. Is the same neuron type-specific TF continuously required, from development through adulthood, to induce a specific set of terminal identity genes and simultaneously prevent ‘unwanted’ features? Alternatively, a given neuron type could employ different mechanisms for selection (during development) and maintenance (through adulthood) of its function-defining terminal features. Addressing this fundamental problem has been challenging in the vertebrate nervous system, in part due to its inherent complexity and difficulty to track individual neuron types with single-cell resolution from embryo to adult.

To study how neurons select and maintain their terminal identity features, we use as a model the well-defined motor neuron (MN) subtypes of the Caenorhabditis elegans ventral nerve cord (equivalent to vertebrate spinal cord). Five cholinergic (DA, DB, VA, VB, AS) and two GABAergic (DD, VD) MN subtypes are located along the nerve cord and control locomotion (Figure 1A) (Van Stetten et al., 2006; White et al., 1986). Because they are present in both C. elegans sexes (males and hermaphrodites), we will refer to them as ‘sex-shared’ MNs. In addition, there are two subtypes of ‘sex-specific’ cholinergic MNs: the hermaphrodite-specific VC neurons control egg laying (Portman, 2017; Schafer, 2005), and the male-specific CA neurons are required for mating (Schindelman et al., 2006) (Figure 1A). In addition to distinct morphology and connectivity, each subtype can be molecularly defined by the combinatorial expression of known terminal identity genes, such as ion channels, NT receptors, and neuropeptides (Figure 1B). An extensive collection of transgenic reporter C. elegans animals for MN subtype-specific terminal identity genes is available, thereby providing a unique opportunity to investigate, at single-cell resolution, the effects of TF gene removal on developing and adult MNs.
Figure 1. An extensive collection of terminal identity markers for distinct motor neuron subtypes of the *C. elegans* ventral nerve cord. (A) Schematic showing distinct morphology for each motor neuron subtype in the *C. elegans* hermaphrodite. Below, colored dots represent the invariant cell body position of all MNs of the ventral nerve cord (VNC). Red: 39 sex-shared cholinergic MNs (DA2–7 = 6 neurons, DB3–7 = 5, VA2–11 = 10, VB3–11 = 9, Figure 1 continued on next page

| Terminal identity genes | Cholinergic MNs | GABAergic MNs |
|--------------------------|----------------|--------------|
| | **both sexes** | **both sexes** |
| TF: unc-3/ COE | DA DB VA VB AS VC CA | DD VO |
| unc-17/ VAChT | | |
| cha-1/ ChAT | | |
| cho-1/ ChT | | |
| aor-2/ AChR | | |
| aor-5/ AChR | | |
| aor-16/ AChR | | |
| unc-53/ Nav2 | | |
| lgc-36/ GABAR | | |
| del-1/ ENaC | | |
| unc-8/ ENaC | | |
| bpi-1/ TRP | | |
| unc-77/ VGCl | | |
| slo-2/ K channel | | |
| unc-129/ TGFb | | |
| dbi-1/ TGFb-like | | |
| inx-12/INX | | |
| ida-1/ IA-2 | | |
| gh-5/ GluR | | |
| scr-16/ GPCR* | | |
| TF: unc-30/ Pitx | | |
| unc-25/ GAD | | |
| unc-46/ LAMP | | |
| unc-47/ VGAT | | |
| br-39 | | |
| F21D12.3 | | |
| Neuro- | | |
| peptides, ion | | |
| channels, signaling | | |
| molecules, etc. | | |
| | | |
UNC-3, the sole C. elegans ortholog of the Collier/Olf/Ebf (COE) family of TFs, is selectively expressed in all sex-shared cholinergic MNs of the nerve cord (Figure 1B) (Kratsios et al., 2017; Kratsios et al., 2012; Pereira et al., 2015; Prasad et al., 2008; Prasad et al., 1998). Animals lacking unc-3 display striking locomotion defects (Brenner, 1974). UNC-3 is known to directly activate a large battery of terminal identity genes expressed either in all sex-shared cholinergic MNs (e.g., the NT identity genes unc-17/VAChT and cha-1/ChAT), or in certain subtypes (e.g., ion channels, NT receptors, signaling molecules) (Kratsios et al., 2012) (Figure 1B–C). Based on its ability to broadly co-regulate many distinct terminal identity features, unc-3 has been classified as a terminal selector gene (Hobert, 2008). Besides its well-established function as activator of terminal identity genes in cholinergic MNs, whether and how UNC-3 can prevent expression of terminal features of alternative neuronal identities remains unclear.

Here, we describe a dual role for UNC-3 that enables sex-shared cholinergic MNs to select during development and maintain throughout life their terminal identity features. We find that UNC-3 is continuously required - from development through adulthood - not only to activate cholinergic MN identity genes, but also to prevent expression of terminal features normally reserved for other MN subtypes of the nerve cord, namely the sex-shared GABAergic VD neurons (DD2/C05 = 4); Blue: nine sex-shared GABAergic DD neurons (DD2–5 = 9); With the exception of VC, all other subtypes have 1–3 extra neurons located at the flanking ganglia (retrovesicular and pre-anal) of the VNC (not shown). Individual neurons of each subtype intermingle along the VNC. (B) Table summarizing expression of terminal identity markers for VNC MNs. The sex-shared GABAergic MNs (DD, VD) and the sex-specific MNs (VC, CA) do not express UNC-3. Conversely, the sex-shared cholinergic MNs (DA, DB, VA, VB, AS) and the sex-specific MNs (VC, CA) do not express UNC-30/Pitx. For the genes indicated with an asterisk (*), a detailed expression pattern is provided in Figure 1—figure supplement 1. Of note, the male-specific MNs of the CP subtype are also not shown. (C) Schematic that summarizes the known function of UNC-3 (activator of cholinergic MN identity genes) and the question under investigation: does UNC-3 prevent expression of terminal identity features reserved for other MN subtypes?

The online version of this article includes the following figure supplement(s) for figure 1:

**Figure supplement 1.** Detailed characterization of the expression pattern of VC and VD terminal identity markers.

To uncover the molecular mechanism underlying the dual role of UNC-3, we conducted an unbiased genetic screen, which led to the identification of the Hox protein LIN-39 (Scr/Dfd/Hox4-5) as the intermediary factor necessary for expression of alternative neuronal identity features (e.g., VD, VC) in unc-3-depleted MNs. Unlike previously described cases of TFs that act indirectly to prevent alternative neuronal identities by repressing intermediary factors (discussed earlier), UNC-3 does not repress lin-39 and both factors are co-expressed in cholinergic MNs. However, UNC-3 antagonizes the ability of LIN-39 to induce terminal features of alternative identities. Intriguingly, UNC-3 also synergizes with LIN-39 to co-activate multiple terminal identity features specific to cholinergic MNs. Consequently, loss of unc-3 causes a switch in the transcriptional targets of LIN-39, thereby alternative, not cholinergic MN-specific, terminal identity features become activated and locomotion defects occur. Given that terminal selectors and Hox proteins are expressed in a multitude of neuron types across species (Deneris and Hobert, 2014; Hobert and Kratsios, 2019; Philippidou and Dasen, 2013; Estacio-Gómez and Díaz-Benjumea, 2014), the strategy of a terminal selector preventing a Hox transcriptional switch may constitute a general principle for safeguarding neuronal identity throughout life.
**Results**

**UNC-3 has a dual role in distinct populations of ventral nerve cord (VNC) motor neurons**

Neuron type-specific TFs often promote a specific identity and simultaneously suppress features reserved for other, functionally related neuronal types (Arlotta and Hobert, 2015). To test this notion for UNC-3, it was essential to identify a set of terminal identity markers for all unc-3-negative MN subtypes of the VNC, namely the GABAergic (VD, DD) and sex-specific (VC, CA) MNs (Figure 1B). We undertook a candidate gene approach and examined the precise expression pattern of terminal identity genes (e.g., NT receptors, signaling proteins, ion channels, neuropeptides) reported to be expressed in unc-3-negative MNs (www.wormbase.org). In total, we carefully characterized at single-cell resolution the expression of 15 genes in wild-type animals of both C. elegans sexes at the fourth larval stage (L4) (see Materials and methods and Figure 1—figure supplement 1). This analysis provided nine terminal identity markers highly specific to unc-3-negative MNs that fall into four categories (Figure 1B): (a) two VD-specific markers (ser-2-serotonin receptor [ortholog of HTR1D]; oig-1-one Ig domain protein), (b) one DD-specific marker (flp-13/FMRF like neuropeptide), (c) three markers for sex-specific (VC in hermaphrodites, CA in males) MNs (glr-5/glutamate receptor [ortholog of GRID/GRIK]; srb-16/serpentine GPCR receptor; ida-1/ortholog of protein tyrosine phosphatase PTPRN), and (d) three markers expressed in both GABAergic subtypes (DD, VD) and sex-specific MNs (flp-11/FMRF like neuropeptide, twk-46/potassium channel [ortholog of KCNK1], ils-4/invertebrate type lysozyme).

These nine markers enabled us to test whether unc-3-depleted MNs gain expression of terminal features normally reserved for other MN subtypes. By using animals carrying a strong loss-of-function (null) allele for unc-3 (n3435) (Prasad et al., 2008), we first assessed any putative effects on terminal markers for the sex-shared GABAergic MNs (DD, VD). Although the DD-specific marker flp-13 is unaffected (Figure 2—figure supplement 1, panel A), ectopic expression of the VD-specific markers (ser-2, oig-1) was observed in unc-3-depleted MNs (Figure 2A–B). Interestingly, this ectopic expression was region-specific, observed in cholinergic MNs of the mid-body region of the VNC with 100% penetrance (Figure 2A–B). Importantly, 12.1 ± 2.6 (mean ± STDV) out of the 39 unc-3-depleted MNs in the VNC were ectopically expressing these VD markers, suggesting that not all unc-3-depleted MNs acquire VD terminal identity features. Given that GABAergic and cholinergic MNs are generated in normal numbers in unc-3 animals (Kratsios et al., 2012), the increase in the number of neurons expressing the VD markers cannot be attributed to early developmental defects affecting MN numbers. We next asked whether these ~12 MNs adopt additional VD terminal identity features, such as expression of genes involved in GABA biosynthesis (unc-25/GAD and unc-47/VGAT), or selectively expressed in GABAergic MNs (trr-39, klp-4). However, this does not appear to be the case, arguing against a complete cell fate switch (Figure 2—figure supplement 1, panel A). We conclude that, in the absence of unc-3, cholinergic MNs not only lose their original terminal identity, but a third of them (~12 out of 39) in the mid-body VNC region also gain some terminal identity features normally reserved for the sex-shared VD neurons (Figure 1). We will refer to these unc-3-depleted MNs as ‘VD-like’ (Figure 2G). We also uncovered the identity of these cells across multiple unc-3 mutant animals and conclude that it is the same 12 neurons that become VD-like across animals (Figure 2—figure supplement 1, panel B).

To test whether UNC-3 also prevents expression of terminal identity features of sex-specific cholinergic MNs, we examined three VC-specific terminal markers (glr-5, srb-16, ida-1 in Figure 1B) in hermaphrodite nematodes lacking unc-3. Again, we observed region-specific effects with 100% penetrance in the same cells across multiple animals (Figure 2C–D, Figure 2—figure supplement 1, panel B). All three markers were ectopically expressed in 10.5 ± 3.7 (mean ± STDV) of the 39 unc-3-depleted MNs located in the mid-body region of the VNC (Figure 2C–D). These results are in agreement with a previous study reporting ectopic ida-1 expression in unc-3-depleted MNs (Prasad et al., 2008). If these ~11 MNs fully adopt the VC terminal identity, then they should also express genes necessary for acetylcholine biosynthesis since VC neurons are cholinergic. However, this is not the case as expression of unc-17/VACHT and cho-1/ChT is dramatically affected in unc-3-depleted MNs (Kratsios et al., 2012). These data suggest that ~11 of the 39 unc-3-depleted MNs in the mid-body VNC region adopt some, but not all, VC terminal identity features. We will therefore refer to these unc-3-depleted MNs as ‘VC-like’ (Figure 2G).
**Figure 2.** UNC-3 has a dual role in cholinergic ventral cord motor neurons. (A) Terminal identity markers of VD neurons (ser-2, oig-1) are ectopically expressed in unc-3-depleted MNs. Representative images of larval stage 4 (L4) hermaphrodites are shown. Similar results were obtained in adult animals. Arrowheads point to MN cell bodies with gfp marker expression. Green fluorescence signal is shown in white for better contrast. Dotted black box indicates imaged area. (B) Quantification of VD markers (ser-2, oig-1) in WT and unc-3 (n3435) at L4. N > 15. ***p<0.001. For details on box plots, Figure 2 continued on next page
Are the VD-like and VC-like neurons in unc-3 hermaphrodites distinct populations? To test this, we generated unc-3 hermaphrodites that carry a green fluorescent reporter for VC terminal identity (ida-1::gfp) and a red reporter for VD identity (ser-2::rfp). We found no overlap of the two reporters, indicating that the VD-like and VC-like neurons represent two distinct populations (Figure 2E–F). We further corroborated this result by taking advantage of the invariant lineage and cell body position of all MNs along the C. elegans nerve cord (Figure 2—figure supplement 1, panel B). Of note, the VC-like population appears to be lineally related to VC neurons, whereas the VD-like population is not lineally related to VD neurons (Figure 2—figure supplement 1, panels B-C). Lastly, terminal identity markers normally expressed in both VD and VC neurons (flp-11, ilys-4, twk-46) display an additive effect in unc-3 mutants, as they are ectopically expressed in both VD-like and VC-like populations, further suggesting the presence of distinct unc-3 MN populations (Figure 2—figure supplement 1, panels D-E).

To summarize, there are 39 unc-3-expressing MNs along the wild-type nerve cord in hermaphrodites. While loss of unc-3 uniformly leads to loss of cholinergic identity in all these MNs (Kratsios et al., 2012), one population (~12 MNs) acquires VD-like molecular features, while a second population (~11 MNs) acquires VC-like molecular features, uncovering a dual role of UNC-3 in these populations (Figure 2G). Of note, the remaining MNs (~16) in the VNC of unc-3 mutants [39 - (12 VD-like + 11 VC-like)=16] do not gain either VD or VC terminal identity features.

**The dual role of UNC-3 in cholinergic MNs extends to both C. elegans sexes**

To test whether the dual function of UNC-3 applies to both sexes, we extended our analysis to C. elegans males. First, we showed that loss of unc-3 in males resulted in loss of several cholinergic MN terminal identity features (Figure 2—figure supplement 2). Second, we observed ectopic expression of VD-specific terminal identity markers (oig-1, ser-2) in 11.9 ± 3.9 (mean ± STDV) out of the 39 unc-3-depleted MNs, indicating the presence of ‘VD-like’ neurons in the male nerve cord (Figure 2—figure supplement 2). Lastly, we asked whether unc-3 loss leads to ectopic expression of terminal identity markers (ida-1, srb-16, glr-5) for male-specific CA neurons. Indeed, we found this to be the case (Figure 2—figure supplement 2), suggesting the adoption of ‘CA-like’ features by a population of unc-3-depleted MNs. Similar to hermaphrodites, these VD-like and CA-like cells were observed in the mid-body region of the male nerve cord with 100% penetrance (Figure 2—figure supplement 2).

Taken together, our findings uncover a dual role for UNC-3 in sex-shared cholinergic MNs. UNC-3 is not only required to activate cholinergic MN identity genes (Kratsios et al., 2012), but also to prevent expression of molecular features normally reserved for three other, functionally related neuronal subtypes of the nerve cord (VD, VC, CA). In both sexes, UNC-3 prevents expression of select terminal features of VD neurons in a specific population of cholinergic MNs. In a second population, UNC-3 prevents expression of terminal features normally reserved for sex-specific MNs, that is VC features in hermaphrodites and CA features in males. In the ensuing sections, we focus our analysis on C. elegans hermaphrodites to dissect the molecular mechanism underlying the dual role of UNC-3.
UNC-3 is continuously required to prevent expression of VD and VC terminal identity features

Neuron type-specific TFs that promote a specific identity and simultaneously prevent alternative features have been previously described (see Introduction). However, whether this dual role is required transiently (during development), or continuously (throughout life) remains unclear. The UNC-3 case provides an opportunity to distinguish between these two possibilities because ectopic expression of VC and VD features is observed at both larval and adult stages in unc-3 null animals (Figure 2, Figure 2—figure supplement 1). To this end, we employed the auxin-inducible degron (AID) system that enables depletion of UNC-3 in a temporally controlled manner (Zhang et al., 2015). This system requires tagging the UNC-3 protein with the AID degron fused to a fluorescent reporter gene (mNeonGreen, mNG). When UNC-3::mNG::AID and the plant-specific F-box protein TIR1 are co-expressed in MNs (by crossing animals carrying the unc-3::mNG::AID allele with eft-3::TIR1 transgenic animals), application of the plant hormone auxin on these double transgenic animals induces degradation of UNC-3::mNG::AID (Figure 3A–C). Auxin administration at the L4 stage (last larval stage before adulthood) on unc-3::mNG::AID; eft-3::TIR1 animals resulted in a dramatic depletion of UNC-3 at day one adult animals (24 hr after auxin). UNC-3 depletion was accompanied by ectopic expression of VD and VC terminal identity features in nerve cord MNs, demonstrating a post-embryonic requirement for UNC-3 (Figure 3D–E). Similar results were obtained when auxin was applied at different time points (Figure 3D, legend). These findings suggest that UNC-3 is continuously required to prevent expression of VD and VC terminal identity features.

UNC-3 acts indirectly to prevent expression of VD and VC terminal identity genes

How does UNC-3 activate cholinergic MN identity genes and simultaneously prevent terminal features of alternative MN identities (e.g., VD, VC) (Figure 2G)? Based on previous reports, the same TF, within the same neuron, can act as a direct activator for a set of genes and a direct repressor for another set of genes (Lodato et al., 2014; Wyler et al., 2016; Borromeo et al., 2014). While it is known that UNC-3 acts directly – through its cognate binding site (COE motif) – to activate expression of a large battery of cholinergic MN identity genes, we did not find any COE motifs in the cis-regulatory region of VD or VC terminal identity genes (Supplementary file 1). This contrasts the previously described function of UNC-3 as direct repressor (through the COE motif) of terminal identity genes in the chemosensory ASI neurons of C. elegans (Kim et al., 2005).

To test the possibility of indirect repression via an intermediary factor, we focused on VD neurons because, unlike VC neurons, a known activator of VD terminal features has been reported (Cinar et al., 2005; Eastman et al., 1999; Jin et al., 1994). In wild-type animals, the TF UNC-30, ortholog of human PITX1-3, is required to induce VD terminal identity genes. Since UNC-30 is not expressed in cholinergic MNs (Jin et al., 1994), we hypothesized that UNC-3 prevents expression of UNC-30/PITX, leading to inactivation of VD terminal identity genes. However, this is not the case because: (1) ectopic unc-30 expression is not observed in unc-3-depleted MNs, and (2) the ectopic expression of the VD marker (ser-2) in unc-3 mutants was not abolished in unc-3; unc-30 double mutants (Figure 4—figure supplement 1). These observations suggest that UNC-3 may act indirectly to prevent expression of VD and VC terminal identity genes through as yet unknown intermediary factors.

The mid-body Hox protein LIN-39 (Scr/Dfd/Hox4-5) is the intermediary factor necessary for ectopic expression of VD and VC features in unc-3 mutants

If the hypothesis of indirect repression is correct, mutation of the intermediary factor(s) in the unc-3 mutant background would selectively eliminate ectopic expression of VD and/or VC terminal identity genes in unc-3-depleted MNs. To identify such factor(s), we embarked on an unbiased genetic screen. For the screen, we chose a transgenic gfp reporter strain for flp-11, an FMRF-like neuropeptide-encoding gene expressed in both VD and VC neurons (Figure 1B, Figure 1—figure supplement 1), which is markedly affected by UNC-3 (Figure 4A–B, Figure 2—figure supplement 1, panels D–E). We mutagenized unc-3 (n3435); flp-11::gfp animals with ethyl methanesulfonate (EMS) and visually screened ~4200 haploid genomes for mutants in which ectopic flp-11::gfp expression in
Figure 3. UNC-3 is continuously required to prevent expression of VD and VC terminal identity features. (A) Schematic showing time window of auxin administration. (B) Animals of the unc-3::mNG::AID; eft-3::TIR1 genotype were either administered ethanol (EtOH) or auxin at the L4 stage. Twenty four hours later, expression of endogenous unc-3 reporter (unc-3::mNG::AID) is severely reduced in the nuclei of VNC MNs (arrowheads) at the young adult stage (day 1). The same exact region was imaged in EtOH- and auxin-treated worms. mNG green fluorescent signal is shown in white for better contrast. White dotted line indicates the boundary of intestinal tissue (gut), which tends to be autofluorescent in the green channel. (C) Quantification of number of MNs expressing the unc-3::mNG::AID reporter after EtOH (control) and auxin treatment. N > 12. ***p<0.001. (D) Auxin or ethanol (control) were administered at larval stage 3 (L3) on unc-3::mNG::AID; eft-3::TIR1 animals carrying the VD marker ser-2::gfp. Images were taken at the young adult stage (day 1.5). A significant increase in the number of MNs expressing the VD marker was evident in the auxin-treated animals compared to EtOH-treated controls. For comparison, quantification is provided for ser-2::gfp expressing MNs of wild-type animals and unc-3(n3435) mutants. Similar results were obtained when auxin was applied at L4 or day 1 adult animals. N > 20. ***p<0.001. (E) Auxin or ethanol (control) were administered at larval stage 4 (L4) on unc-3::mNG::AID; eft-3::TIR1 animals carrying the VC marker glr-5::gfp. Images were taken at the young adult stage (day 2). A significant increase in the number of MNs expressing the VC marker was evident in the auxin-treated animals compared to EtOH-treated controls. N > 11. *p<0.05; ***p<0.001.
Figure 4. A genetic screen identifies the mid-body Hox protein LIN-39 (Scr/Dfd/Hox4-5) as necessary for ectopic expression of VD and VC terminal features. (A) Representative images of L4-stage WT, unc-3(n3435), unc-3(n3435); kas1, and unc-3(n3435); lin-39(n1760) animals carrying flp-11::gfp (VD/VC marker). Arrowheads point to MN cell bodies with gfp marker expression. (B) Quantification graph summarizing results from panel A. The two rightmost bars show quantification of two independent transgenic lines driving lin-39 RNAi specifically in cholinergic MNs (Punc-3 >lin-39 RNAi) of unc-3.

Figure 4 continued on next page
unc-3-depleted MNs is suppressed. We isolated one mutant allele (kas1) (Figure 4A–B). The phenotype was 100% penetrant as all unc-3 (n3435); flp-11::gfp animals carrying kas1 in homozygosity consistently displayed a dramatic reduction in ectopic flp-11 expression.

Gross morphological examination of unc-3 (n3435); kas1; flp-11::gfp hermaphrodites revealed that, unlike unc-3 (n3435); flp-11::gfp animals, the introduction of kas1 is accompanied by a lack of the vulva organ (vulvaless phenotype). Upon a literature survey for TF mutants that are vulvaless, we stumbled across the mid-body Hox gene lin-39 (ortholog of Dfd/Scr in flies and Hox4-5 in vertebrates) (Aboobaker and Blaxter, 2003; Clark et al., 1993), and hypothesized that the molecular lesion of kas1 may lie in the lin-39 locus. Indeed, Sanger sequencing uncovered a point mutation on the splice acceptor site (WT: A → G > A) in the second intron of lin-39 (Figure 4C). Similar to unc-3 (n3435); kas1 animals, unc-3 (n3435) mutants carrying a previously published strong loss-of-function (premature STOP) allele of lin-39 (n1760) (Clark et al., 1993) displayed the same loss of ectopic flp-11 expression (Figure 4A–C), suggesting that kas1 is a loss-of-function mutation of lin-39. The ectopic expression of flp-11 in unc-3(n3435); kas1 animals can be, at least partially, rescued by (1) selective expression of lin-39 cDNA in cholinergic MNs, and (2) introduction of the lin-39 wild-type locus in the context of a ~ 30 kb genomic clone (fosmid) (Figure 4—figure supplement 1), corroborating that the kas1 lesion in the lin-39 locus is the phenotype-causing mutation.

Because flp-11 is expressed in both VD and VC neurons, we next tested whether lin-39 is required for ectopic expression of VD-specific (ser-2, oig-1) and VC-specific (ida-1) terminal identity genes in unc-3-depleted MNs. We found this to be the case by either generating unc-3 (n3435); lin-39 (n1760) double mutants (for VD markers) or by performing cholinergic MN-specific RNAi for lin-39 in unc-3 (n3435) animals (for VC marker) (Figure 4D). RNAi was necessary because VC neurons do not survive in lin-39 (n1760) animals (Potts et al., 2009), and the use of the n1760 allele could confound our VC marker quantifications. Of note, all other nerve cord MN subtypes are normally generated in lin-39 (n1760) single and unc-3 (n3435); lin-39 (n1760) double mutants (Stefanakis et al., 2015), indicating that suppression of the unc-3 phenotype, that is, loss of ectopic VD gene expression in the double mutants is not due to MN elimination. Taken together, our genetic screen identified the mid-body Hox gene lin-39 to be necessary for ectopic expression of both VD and VC terminal features in unc-3-depleted MNs (Figure 4F).

Interestingly, this finding contradicts our initial hypothesis of UNC-3 repressing an intermediary TF in order to prevent expression of VD and VC features because lin-39 is co-expressed with (not repressed by) unc-3 in wild-type cholinergic MNs at the mid-body region of the VNC (Figure 4D), as evident by our single-cell analysis of unc-3 and lin-39 reporters (Figure 4—figure supplement 2). Of note, 28 cholinergic MNs co-express unc-3 and lin-39, which is in close agreement with the total number of VD-like (12.1 ± 2.6) and VC-like (10.5 ± 3.7) cells observed in unc-3 mutants (Figure 4E–F). Moreover, we found that lin-39 acts cell-autonomously as cholinergic MN-specific RNAi against lin-39 in unc-3 (n3435) animals resulted in a significant reduction of ectopic terminal identity marker (flp-11, ida-1) expression (Figure 4B–D). In the following Results sections, we describe the molecular mechanism through which UNC-3 and LIN-39/Hox select and maintain throughout life key terminal features of cholinergic MNs (Figure 4F).
UNC-3 prevents a switch in the transcriptional targets of LIN-39 in cholinergic motor neurons

What is the function of LIN-39 in wild-type cholinergic MNs of the VNC? Our previous findings suggested that LIN-39 and UNC-3, together with another mid-body Hox protein, MAB-5 (Antp/Hox6-8) (Salser et al., 1993), act synergistically to control expression of two cholinergic MN terminal identity genes (unc-129,ortholog of human BMP; del-1/Degenerin like sodium channel [ortholog of human SCNN1G]) (Kratsios et al., 2017). To test the extent of this synergy, we examined in lin-39 and mab-5 null animals the expression of 4 additional cholinergic MN terminal identity genes known to be controlled by UNC-3 (acr-2/nicotinic acetylcholine receptor; db-1/DPP/BMP-like; unc-77/sodium channel [ortholog of human NALCN], slo-2/potassium sodium-activated channel [ortholog of human KCNT1]) (Kratsios et al., 2012). In all four cases, we found a statistically significant decrease in lin-39 mutants, and this effect was exacerbated in lin-39; mab-5 double mutants (Figure 5A–B), indicating that the synergy of LIN-39 with MAB-5 (and UNC-3) extends to multiple terminal identity genes in cholinergic MNs (WT panel in Figure 5D). The observed effects were 100% penetrant and consistent with the previously described region-specific expression pattern of lin-39 and mab-5 in VNC MNs (Figure 5A) (Kratsios et al., 2017). Of note, while MAB-5 collaborates with LIN-39 to activate cholinergic MN identity genes (Figure 5B), it does not affect the ectopic expression of VD or VC genes observed in unc-3 mutants (Figure 5—figure supplement 1).

Since UNC-3 controls directly, via its cognate binding site, cholinergic MN terminal identity genes (Kratsios et al., 2012), we then asked whether this is the case for LIN-39. We analyzed available ChIP-Seq data for LIN-39 from the modENCODE project (Boyle et al., 2014) and found evidence for direct LIN-39 binding in the cis-regulatory of all six cholinergic MN terminal identity genes (unc-129, del-1, acr-2, db-1, unc-77, slo-2) (Figure 5C, Figure 5—figure supplement 1). Moreover, we identified multiple consensus LIN-39 binding sites (previously defined as GATTGATG) (Boyle et al., 2014) located within the LIN-39 ChIP-Seq peaks in the cis-regulatory region of the aforementioned genes (Supplementary file 2).

This analysis strongly suggests that LIN-39, similar to UNC-3, regulates directly the expression of multiple terminal identity genes in cholinergic MNs (Figure 5D). However, in the absence of UNC-3, the function of LIN-39 in cholinergic MNs is modified. Instead of activating cholinergic MN identity genes, LIN-39 activates VD or VC terminal identity genes in unc-3-depleted MNs (Figure 4). Taken together, our data suggest that UNC-3 antagonizes the ability of LIN-39 to activate alternative identity genes, thereby preventing a switch in the transcriptional targets of LIN-39 (model schematized in Figure 5D). If this hypothesis is correct, one would expect decreased LIN-39 binding in the cis-regulatory region of cholinergic MN terminal identity genes in unc-3 mutants. By performing ChIP-Seq for LIN-39 in unc-3 mutant animals, we indeed observed decreased LIN-39 binding in the cis-regulatory region of the aforementioned genes (Figure 5C, Figure 5—figure supplement 1). As a positive control, LIN-39 binding in unc-3 mutant animals was observed in other loci, including the lin-39 locus itself (Figure 5—figure supplement 1), consistent with the known role of LIN-39 in regulating its own expression (Niu et al., 2011). Similar results were obtained by ChIP-qPCR for LIN-39 targets in unc-3 mutants animals (Figure 5—figure supplement 1). We conclude that, in the absence of UNC-3, LIN-39 is released from cholinergic MN terminal identity gene promoters, presumably leading to increased availability of LIN-39 and thereby activation of alternative identity genes.

LIN-39 is continuously required to control expression of terminal identity genes in cholinergic MNs

The neuronal function of Hox proteins at post-developmental stages is largely unknown (Hutlet et al., 2016). The continuous expression of mid-body Hox lin-39 in both developing and adult cholinergic MNs led us to investigate whether lin-39 is required to maintain expression of terminal identity genes in these neurons. To test this idea, we employed clustered regularly interspaced short palindromic repeats (CRISPR)/Cas9-based genome engineering and generated an auxin-inducible lin-39 allele (lin-39::mNG::3xFLAG::AID) that also serves as an endogenous lin-39 reporter (mNG). Animals carrying lin-39::mNG::3xFLAG::AID display no developmental phenotypes and show nuclear mNG expression in MNs located at the mid-body region of the VNC during development and adult stages (Figure 6A), corroborating previous observations with a LIN-39 antibody (Maloof and Kenyon, 1998). Upon crossing the lin-39::mNG::3xFLAG::AID animals with the ef-3::
Figure 5. UNC-3 prevents a switch in the transcriptional targets of LIN-39 in cholinergic motor neurons. (A) Schematic summarizing the expression pattern of lin-39 and mab-5 in VNC cholinergic MNs. Below, representative images are shown of unc-77::GFP in WT, lin-39 (n1760), mab-5 (1239) and lin-39 (n1760); mab-5 (1239) animals at L4 stage. Arrowheads point to MN cell bodies with gfp marker expression. Green fluorescence signal is shown in white for better contrast. Dotted black box indicates imaged area. (B) Quantification of cholinergic MN terminal identity markers (unc-77, dbl-1, acr-2, ser-2, glr-5, ida-1, srb-16, flp-11).}

**Figure 5 continued on next page**
Figure 5 continued

slo-2 in WT, lin-39 (n1760), mab-5 (1239) and lin-39 (n1760); mab-5 (1239) animals at L4. N > 15. **p<0.01; ***p<0.001. (C) ChIP-Seq tracks are shown for LIN-39 on four cholinergic MN terminal identity genes (acr-2, unc-129, db1-1, del-1). The WT data come from the modENCODE project (Boyle et al., 2014), whereas the unc-3 (-) data were obtained by performing ChIP-Seq for LIN-39 on unc-3 (n3435); lin-39 (kas9 [lin-39::mNG::3xFLAG::AID] animals. (D) Schematic showing the transcriptional switch in LIN-39 targets. In WT animals, UNC-3, MAB-5 and LIN-39 co-activate subtype-specific genes in cholinergic MNs (e.g., unc-77, db1-1, unc-129, acr-2). In unc-3 mutants, LIN-39 is no longer able to activate these genes, and instead switches to VD- or VC-specific terminal identity genes. Black font: gene expressed. Gray font: gene not expressed. Gray arrows indicate inactive genetic interactions. COE motif taken from Kratsios et al. (2012) and LIN-39 site taken from Weirauch et al. (2014) are represented with black rectangles and dots, respectively. The online version of this article includes the following figure supplement(s) for figure 5:

**Figure supplement 1.** MAB-5 is not required for ectopic VD or VC marker expression and LIN-39 binding on cholinergic MN genes is affected in unc-3 mutants.

TIR1 line, we observed hypomorphic effects in the expression of two cholinergic MN identity genes (acr-2, unc-77) (Figure 4—figure supplement 2, panel C). Although LIN-39 protein is present in the nuclei of cholinergic MNs of lin-39::mNG::3xFLAG::AID; eft-3::TIR1 animals (Figure 6A), these effects are likely due to a mild reduction in LIN-39 levels triggered by TIR1. However, post-embryonic auxin administration on these animals resulted in efficient LIN-39 protein depletion and significantly enhanced these effects (Figure 6A–C, Figure 4—figure supplement 2, panel C). We therefore conclude that LIN-39 is continuously required to maintain terminal identity features in cholinergic MNs.

Next, we sought to determine whether LIN-39 is continuously required for the ectopic activation of VD and VC terminal features observed in unc-3 null animals. Indeed, auxin administration at L4 stage on unc-3(n3435); lin-39::mNG::3xFLAG::AID animals carrying either a VD (ser-2), VC (glr-5), or VD/VC (flip-11) marker resulted in a statistically significant suppression of the unc-3 phenotype when compared to control (treated with ethanol) (Figure 6D–F).

To sum up, our findings with the auxin-inducible (Figure 4—figure supplement 2, panel C) and null lin-39 alleles (Figures 4F and 5A–B) indicate that, in the presence of UNC-3, LIN-39 is required to induce and maintain expression of cholinergic MN terminal identity genes (Figure 5D). In the absence of UNC-3 (Figure 6), LIN-39 is also continuously required - from development and possibly throughout life - for ectopic activation of VD and VC terminal identity genes (Figure 5D).

**LIN-39 is an activator of VD and VC terminal identity genes**

The observation that lin-39 is required for ectopic activation of both VD and VC terminal identity genes in unc-3-depleted MNs prompted us to examine the role of lin-39 in VD and VC neurons of wild-type animals. Does LIN-39 control the same VD- and VC-specific terminal identity genes that become ectopically expressed in unc-3 mutants?

To this end, we leveraged our endogenous lin-39 reporter (lin-39::mNG::3xFLAG::AID) to assess expression in wild-type VD neurons at the mid-body region of the VNC, and found this to be the case (Figure 7A, Figure 4—figure supplement 2). Next, we found that LIN-39 is required to induce expression of VD terminal identity genes (ser-2, oig-1) (Figure 7B). To gain further mechanistic insights, we then asked whether lin-39 acts together with UNC-30, the known activator of GABAergic MN identity genes (Eastman et al., 1999; Jin et al., 1994). Apart from confirming previous observations of UNC-30 controlling the VD-specific oig-1 gene (Cinar et al., 2005; Howell et al., 2015), we also found that ser-2 (Figure 7B) and flip-11 (Figure 4—figure supplement 2, panel E) constitute novel UNC-30 targets in VD neurons. To test for synergistic effects, we focused on ser-2 and flip-11, two VD-expressed terminal identity genes mildly affected in lin-39 or unc-30 single mutants. We generated lin-39; unc-30 double mutants and observed stronger effects than either single mutant (Figure 7B, Figure 4—figure supplement 2, panel E). Such additive effects indicate that lin-39 and unc-30 act in parallel to activate VD terminal identity genes. Importantly, expression of other UNC-30 targets in GABAergic MNs, such as flip-13 (DD-specific terminal identity marker) (Cinar et al., 2005; Shan et al., 2005; Yu et al., 2017) and genes expressed in both DD and VD neurons (unc-25/GAD, unc-47/VGAT), is unaffected in lin-39 mutants (Figure 7B, Figure 4—figure supplement 2, panel F). Unlike UNC-30 that broadly controls multiple terminal features (NT identity and VD-specific terminal features) in VD neurons, we conclude that lin-39 is selectively required for activation of VD-specific terminal identity genes (Figure 7F). To test for a maintenance role in VD
neurons, we administered auxin at various post-developmental stages (L3, L4, day one adult) on animals carrying the lin-39::mNG::3xFLAG::AID allele. We found that LIN-39 is continuously required to maintain expression of the VD terminal identity gene ser-2 (Figure 7C).

The above genetic analysis indicates that LIN-39 and UNC-30/PITX activate expression of VD-specific genes (left panel in Figure 7F). Similarly, LIN-39 and UNC-3 directly co-activate terminal identity genes in cholinergic MNs (left panel in Figure 5D). Since the absence of UNC-3 leads to ectopic activation of VD-specific genes (Figure 5D), we next considered the converse possibility: Does the absence of UNC-30/PITX lead to ectopic activation of cholinergic MN terminal identity genes in GABAergic VD neurons? However, this appears not to be the case as expression of 4 cholinergic
Figure 7. LIN-39 is an activator of VD terminal identity genes. (A) Schematic summarizing unc-30 and lin-39 expression in VD and DD neurons populating the VNC. In addition, 4 VD and 2 DD neurons are located in ganglia flanking the VNC (not shown because they were excluded from our analysis). Raw data on lin-39 expression described in Figure 4—figure supplement 2. (B) Quantification of two VD (ser-2::gfp, oig-1::gfp) and one DD (flp-13::gfp) markers in WT and lin-39 (n1760) animals at L4. Both VD markers were also tested in unc-30 (e191) mutants. Double lin-39 (n1760); unc-30
(e191) mutants showed a more severe reduction in expression of the VD marker ser-2::gfp compared to each single mutant. N > 15. ***p<0.001. N. S: not significant. (C) Auxin or ethanol (control) were administered at larval stage 3 (L3) on lin-39::mNG::3xFLAG::AID; eft-2::TIR1 animals carrying the VD marker ser-2::gfp. Images were taken at the young adult stage (day 1.5). A significant decrease in the number of MNs expressing the VD marker was evident in the auxin-treated animals compared to EtOH-treated controls. Similar results were obtained when auxin administration occurred at L4 or day one adult animals. For comparison, quantification of marker expression is also provided in WT and lin-39 (n1760) animals. N > 15. **p<0.01, ***p<0.001. N. S: not significant. (D) Several terminal identity markers of cholinergic neurons (acr-2, slo-2, unc-129, del-1) are not ectopically expressed in unc-30-depleted GABAergic MNs. A strong loss-of-function allele e191 for unc-30 was used (Brenner, 1974; Eastman et al., 1999). Arrowheads point to MN cell bodies with gfp marker expression. Green fluorescence signal is shown in white for better contrast. (E) Quantification of data presented in panel D. N. S: not significant. (F) Schematic summarizing the function of LIN-39 and UNC-30 in GABAergic VD neurons. LIN-39 site is taken from Weirauch et al. (2014). UNC-30 site is taken from Yu et al. (2017).

LIN-39 acts through distinct cis-regulatory elements to control oig-1 expression in VD and VD-like motor neurons

Does LIN-39 act directly or indirectly to activate VD and VC terminal identity genes? We used the auxin-inducible lin-39::mNG::3xFLAG::AID allele to address this question because, unlike all other nerve cord MNs, the VC neurons do not survive in lin-39 (n1760) null animals (Potts et al., 2009). We applied auxin at a late larval stage (L3-L4) to knockdown LIN-39 and observed that VC neurons do not die, providing an opportunity to test for putative effects on VC terminal identity gene expression. Indeed, we found a statistically significant reduction in the number of VC neurons expressing srb-16 (compare auxin and ethanol in Figure 4—figure supplement 2, panel D).

Taken together, lin-39 is required for expression of VD- and VC-specific terminal identity genes. In VD neurons, LIN-39 acts together with UNC-30/PITX to activate expression of VD-specific genes (left panel in Figure 7F). Collectively, these findings on VD and VC neurons together with observations on cholinergic MNs (Figure 5D) show that, in different MN subtypes, the mid-body Hox gene lin-39 controls expression of distinct terminal identity genes, likely due to collaboration with distinct TFs (i.e., UNC-3 and MAB-5 in cholinergic MNs versus UNC-30 in VD neurons [compare Figure 5D and Figure 7F]).

MN markers (acr-2, slo-1, unc-129, del-1), normally co-activated by UNC-3 and LIN-39 (Figure 5A–B), is unaffected in unc-30 mutants (Figure 7D–F).

Similar to its role in sex-shared VD neurons, does lin-39 control expression of terminal identity genes in sex-specific VC neurons? We used the auxin-inducible lin-39::mNG::3xFLAG::AID allele to address this question because, unlike all other nerve cord MNs, the VC neurons do not survive in lin-39 (n1760) null animals (Potts et al., 2009). We applied auxin at a late larval stage (L3-L4) to knockdown LIN-39 and observed that VC neurons do not die, providing an opportunity to test for putative effects on VC terminal identity gene expression. Indeed, we found a statistically significant reduction in the number of VC neurons expressing srb-16 (compare auxin and ethanol in Figure 4—figure supplement 2, panel D).

Collectively, these findings on VD and VC neurons together with observations on cholinergic MNs (Figure 5D) show that, in different MN subtypes, the mid-body Hox gene lin-39 controls expression of distinct terminal identity genes, likely due to collaboration with distinct TFs (i.e., UNC-3 and MAB-5 in cholinergic MNs versus UNC-30 in VD neurons [compare Figure 5D and Figure 7F]).

LIN-39 acts through distinct cis-regulatory elements to control oig-1 expression in VD and VD-like motor neurons

Does LIN-39 act directly or indirectly to activate VD and VC terminal identity genes? Analysis of available ChIP-Seq data (modENCODE project) indicates direct regulation of these genes by LIN-39 (Figure 8A, Figure 8—figure supplement 1). However, the low resolution of ChIP-Seq data does not allow the identification of the exact DNA sequence recognized by LIN-39. Therefore, we interrogated the cis-regulatory region of two VD terminal identity genes oig-1 and ser-2 for the presence of the consensus LIN-39 binding site GATTGATG (Boyle et al., 2014) and found several copies located within the boundaries of the LIN-39 ChIP-Seq peaks in oig-1 and ser-2 (Supplementary file 2). To test the functionality of these putative LIN-39 binding sites, we honed in on oig-1 and performed a systematic cis-regulatory analysis in the context of transgenic reporter animals. A previous study identified a minimal 125 bp cis-regulatory element (contained within the LIN-39 peak boundaries) upstream of oig-1 as sufficient to drive reporter gene expression in VD neurons (Howell et al., 2015) (Figure 8A). We independently confirmed this observation, and further found that the 125 bp element contains a single LIN-39 site. Mutation of this site in the context of transgenic oig-1 reporter animals (oig-1 125bp LIN-39 site MUT::tagRFP) leads to a significant reduction of tagRFP expression in VD neurons (Figure 8A–B), phenocopying the effect observed in lin-39 (n1760) null mutants (Figure 7B).

We conclude that, in wild-type animals, LIN-39 acts directly, by recognizing its cognate site, to activate expression of the VD-specific gene oig-1. Interestingly, a functional binding site for UNC-30/PITX also exists in this 125 bp element (Howell et al., 2015; Yu et al., 2017), and is spaced 11 base pairs apart from the LIN-39 site (Figure 7D), indicating that LIN-39 and UNC-30 control oig-1 by recognizing distinct and in close proximity cis-regulatory motifs. Moreover, available UNC-30 ChIP-Seq data further support this possibility as UNC-30 and LIN-39 ChIP-Seq peaks largely overlap at this 125 bp element (Figure 8A). Lastly, deletion of the region where LIN-39 and UNC-30 peaks
Figure 8. LIN-39 acts through distinct cis-regulatory elements to activate oig-1 expression in VD and VD-like neurons. (A) ChIP-Seq tracks are shown for UNC-30 (the top two) and LIN-39 (the bottom two) on VD gene oig-1 locus. The UNC-30 data were obtained from Yu et al. (2017). The LIN-39 data come from the modENCODE project (Boyle et al., 2014). Four LIN-39 peaks are annotated with peak#3 largely overlapping with UNC-30 peak. The results of cis-regulatory analysis in both WT and unc-3 mutants are shown in the lower panel (aligned to the ChIP-seq tracks). Expression patterns of at

Figure 8 continued on next page
least two transgenic lines were analyzed for each construct. ‘+’ indicates consistent and bright expression in ventral nerve cord (VNC) MNs (either VD or cholinergic). ‘+/−’ indicates consistent and bright expression in noticeable less number of VNC MNs. ‘−’ indicates no or extremely dim expression in VNC MNs. ‘N.D.’ Not determined. In the schematic of the transgenes, a known UNC-30 site is shown as a blue box and a bioinformatically predicted LIN-39 site is represented as a black circle (filled circle indicates the presence of the site while unfilled one indicates deletion of the site). MUT indicates deletion of the LIN-39 site and DEL indicates deletion of the respective LIN-39 peak region. 

VNC MNs. ‘N.D.’: Not determined. In the schematic of the transgenes, a known UNC-30 site is shown as a blue box and a bioinformatically predicted expression in cholinergic MNs of selected constructs in the Figure supplement 1.

The online version of this article includes the following figure supplement(s) for figure 8:

**Figure supplement 1.** LIN-39 binds directly to the cis-regulatory region of VD and VC terminal identity genes.

overlap in the context of a 2.6 kb oig-1 reporter (oig-1 2.6kb LIN-39 peak #3 DEL) abolish reporter expression in VD neurons (Figure 8A–B).

We next asked whether LIN-39 acts through the same or distinct cis-regulatory elements to drive oig-1 expression in VD versus VD-like neurons of unc-3 mutants. While oig-1 reporters in the context of a large (32.5 kb) genomic clone (fosmid) or a 2.6 kb intergenic region do show expression in both VD and VD-like neurons of unc-3 mutants, reporter animals carrying 1 kb of cis-regulatory sequence (that contains the 125 bp element) immediately upstream of ATG showed expression only in VD neurons (Figure 8A). Conversely, a distal 1.6 kb element displayed expression in VD-like cells, but no expression in VD neurons of either WT or unc-3 animals, suggesting the VD and VD-like elements are physically separated on the genome. Within the 1.6 kb element, there is a LIN-39 binding peak (peak #4) based on available ChIP-Seq data on WT animals. Deletion of this peak in the context of a 2.6 kb oig-1 reporter (oig-1 2.6kb LIN-39 peak #4 DEL) resulted in loss of expression in VD-like cells of unc-3 mutants, whereas reporter expression was maintained in VD neurons (Figure 8A–B). This analysis strongly suggests that LIN-39 acts through distinct cis-regulatory elements to activate oig-1 expression in VD versus VD-like cells.

**The LIN-39-mediated transcriptional switch depends on UNC-3 and LIN-39 levels**

How does the absence of UNC-3 lead to ectopic and lin-39-dependent activation of VD terminal identity genes in cholinergic MNs (Figure 9D)? In principle, UNC-3 and LIN-39 could physically interact in order to co-activate expression of cholinergic MN terminal identity genes. In the absence of unc-3, this interaction would be disrupted and LIN-39 becomes available, in cholinergic MNs, to assume its VD function, that is to activate VD-specific terminal identity genes (Figure 9D). Although our co-immunoprecipitation (co-IP) experiments on UNC-3 and LIN-39 in a heterologous system (HEK cells) did not provide evidence for physical interaction (Figure 9—figure supplement 1), the heterologous context of this experiment still leaves open the possibility that, in cholinergic MNs in vivo, UNC-3 directly (or indirectly) recruits LIN-39 on terminal identity gene promoters. This scenario is supported by the observed decrease of LIN-39 binding on cholinergic MN gene loci in unc-3 mutants (Figure 5C). Lastly, the gene dosage experiments presented below firmly suggest there is a close stoichiometric relationship between UNC-3 and LIN-39, reminiscent of LIM homeodomain TF stoichiometries described in vertebrate MNs (Song et al., 2009).

Because the decrease of LIN-39 binding is accompanied by ectopic activation of VD terminal identity genes in unc-3-depleted MNs, we hypothesized that LIN-39 is the rate-limiting factor present in limited amount in cholinergic MNs. That is, in the presence of UNC-3, LIN-39 activates cholinergic MN identity genes, but in its absence LIN-39 becomes available to activate alternative identity (e.g., VD) genes. Quantification of the endogenous expression levels of both proteins indeed showed lower levels of LIN-39 expression compared to UNC-3 (Figure 9A). Supporting the aforementioned hypothesis, we found a gene dosage relationship between unc-3 and lin-39. Loss of one unc-3 copy (unc-3 (n3435)/+) caused slight ectopic expression of VD genes (ser-2 in Figure 9B and flp-11 in Figure 9—figure supplement 2, panel A), but that ectopic expression is decreased by loss of one lin-39 copy (unc-3 (n3435)/+; lin-39 (n1760)/+) (Figure 9B, Figure 9—figure supplement 2, panel A). Accordingly, loss of one lin-39 copy in unc-3 null animals (unc-3 (n3435); lin-39 (n1760)/+...
Figure 9. Gene dosage experiments suggest that LIN-39 is the rate-limiting factor. (A) The endogenous expression levels of UNC-3 are higher than LIN-39. The DIC and mNG channels of two worms next to each other on the same slide with the genotype of lin-39::mNG::3xFLAG::AID (top, anterior right, ventral up) and unc-3::mNG::3xFLAG::AID (bottom, anterior left, ventral down) respectively are shown on the left. VNC regions indicated by dashed frame (a, b) are zoomed in at the middle panel with dashed circles around MN nuclei. The identities of these MNs are shown (e.g., VA8, VB9). Figure 9 continued on next page
also reduced, but did not eliminate, ectopic expression of VD genes (Figure 9A, Figure 9—figure supplement 2, panel A). Moreover, knock-down of unc-3 with RNAi specifically in cholinergic MNs also led to ectopic expression of VD genes (Figure 9C, Figure 9—figure supplement 2, panels B-C), whereas ectopic expression of UNC-3 in VD neurons resulted in repression of VD gene expression, presumably by recruiting LIN-39 away from VD promoters (Figure 9C, Figure 9—figure supplement 2, panels B-C). Lastly, we asked whether LIN-39 is sufficient to induce expression of VD terminal identity genes in cholinergic MNs. Indeed, we found this to be the case (Figure 9C, Figure 9—figure supplement 2, panels D-E). In conclusion, we propose that the LIN-39-mediated transcriptional switch observed in unc-3 mutants critically depends on UNC-3 and LIN-39 levels, with the latter being the rate-limiting factor (Figure 9D).

Ectopic expression of VD terminal identity genes in cholinergic motor neurons is associated with locomotion defects

The dual role of UNC-3 revealed by our molecular analysis (Figure 2) led us to posit that the severe locomotion defects observed in unc-3 animals may represent a composite phenotype (Brenner, 1974; Yemini et al., 2013). In other words, these defects are not only due to loss of expression of cholinergic MN terminal identity determinants (e.g., unc-17/VACHT, cha-1/ChAT, del-1/Degen-erin-like sodium channel, acr-2/acetetylcholine receptor [ortholog of CHRNE]), but also due to the ectopic expression of VD and VC terminal features (e.g., ser-2/serotonin receptor [ortholog of HTR1D], flp-11/FRMR-like neuropeptide, glr-5/Glutamate receptor [ortholog of GRID], srb-16/GPCR) in unc-3-depleted MNs. To genetically separate these distinct molecular events, we generated unc-3 (n3435); lin-39 (n1760) double mutants, which do display loss of cholinergic MN terminal identity genes, but the ectopic expression of VD and VC terminal features is suppressed (Figure 4F). We predicted that if ectopic expression of VD and VC genes contributes to locomotion defects, then unc-3 (n3435) mutants would display more severe locomotion defects than unc-3 (n3435); lin-39 (n1760) double mutants. To test this, we performed high-resolution behavioral analysis of freely moving adult (day 1) C. elegans animals using automated multi-worm tracking technology (Javer et al., 2018b; Yemini et al., 2013). This analysis can quantify multiple features related to C. elegans locomotion (e.g., speed, crawling amplitude, curvature, pause, forward and backward locomotion) and, most importantly, each feature can be localized to a specific part of the nematode’s body (e.g., head, mid-body, tail). Since unc-3 and lin-39 expression uniquely overlaps in mid-body nerve cord MNs that innervate mid-body muscles, we hypothesized that loss of unc-3 and/or lin-39 genes would have effects on mid-body posture and motion, and thereby focused our analysis on mid-body curvature features. Of the 49 mid-body features examined, 29 were significantly different in unc-3 single mutants when compared to wild-type (N2 strain) animals (see Supplementary file 3 for all 49 features). Intriguingly, 12 of these 29 features (41.37%) were significantly suppressed in unc-3; lin-39 double mutants (Figure 10A, Figure 10—figure supplement 1, panel A, Supplementary file 3), suggesting that suppression of these behavioral defects could be attributed to suppression of the ectopically expressed VD and VC terminal identity genes in these double mutants. We found no evidence for suppression of the remaining 17 features in unc-3; lin-39 double mutants, likely due to the fact that UNC-3 also controls other terminal identity genes, such as NT pathway genes (Figure 9D), independently of LIN-39.
Figure 10. Ectopic expression of VD terminal identity genes in cholinergic motor neurons is associated with locomotion defects. (A) Examples of three mid-body locomotion features that are significantly affected in unc-3 (n3435) animals, but markedly improved in unc-3 (n3435); lin-39 (n1760) double mutant animals. Each black dot represents a single adult animal. The unit for the first two graphs is 1/microns. The unit for the graph on the right is 1/(microns*seconds). N = 12. Additional mid-body features affected in unc-3 (n3435) animals, but improved in unc-3 (n3435); lin-39 (n1760) mutants are Figure 10 continued on next page
Next, we asked whether ectopic expression of VD terminal identity genes in otherwise wild-type animals can lead to locomotion defects. To test this, we took advantage of our transgenic animals that selectively over-express LIN-39 in cholinergic MNs (Pcho-1 >LIN-39, Punc-3 >LIN-39) (Figure 9C, Figure 10—figure supplement 1, panel D). First, we confirmed that in these animals expression of cholinergic MN terminal identity genes is unaffected (Figure 10—figure supplement 1, panel D). Second, we found that LIN-39 overexpression led to ectopic activation of VD, but not VC, terminal identity genes in cholinergic MNs (Figure 10—figure supplement 1, panel D), providing an opportunity to specifically assess the consequences of ectopic VD gene expression on animal locomotion. We found that 9 of the 29 (31.03%) mid-body features affected in unc-3 (n3435) animals were also altered in animals over-expressing lin-39 in cholinergic MNs (Figure 10B, Figure 10—figure supplement 1, panels B-C, Supplementary file 3).

In conclusion, our behavioral analysis is in agreement with our molecular findings. At the molecular level, we found that lin-39/Hox is necessary for the ectopic expression of VD terminal identity genes in unc-3 mutants. At the behavioral level, this lin-39-dependent, ectopic expression of terminal identity genes is accompanied by locomotion defects.

**Discussion**

During development, individual neuron types must select their unique terminal identity features, such as expression of NT receptors, ion channels and neuropeptides. Continuous expression of these features - from development through adulthood - is essential for safeguarding neuronal terminal identity, and thereby ensuring neuronal function (Deneris and Hobert, 2014; Hobert, 2011; Hobert, 2016). Here, we provide critical insights into the mechanisms underlying selection and maintenance of neuron type-specific terminal identity features by using the well-defined MN populations of the C. elegans nerve cord as a model. First, we report that, in cholinergic MNs, the terminal selector-type TF UNC-3 has a dual role; UNC-3 is not only required to promote cholinergic MN identity features (Kratsios et al., 2012), but also to prevent expression of multiple terminal features normally reserved for three other ventral cord neuron types (VD, VC, CA). Second, we provide evidence that cholinergic MNs can secure their terminal identity throughout life by continuously relying on UNC-3’s dual function. Third, we propose an unusual mechanism underlying this dual function, as we find UNC-3 necessary to prevent a switch in the transcriptional targets of the mid-body Hox protein LIN-39 (Scr/Dfd/Hox4-5) (Figure 9D). Lastly, our findings shed light upon the poorly explored, post-embryonic role of Hox proteins in the nervous system by uncovering that LIN-39 is continuously required to maintain expression of multiple terminal identity genes in MNs.

**UNC-3 determines the function of the rate-limiting factor LIN-39/Hox in cholinergic motor neurons**

Numerous cases of neuron type-specific TFs with a dual role have been previously described in both vertebrate and invertebrate models systems (Britanova et al., 2008; Cheng et al., 2004; Kala et al., 2009; Lopes et al., 2012; Mears et al., 2001; Morey et al., 2008; Nakatani et al., 2007; Sagasti et al., 1999). Although the underlying mechanisms often remain unclear, recent studies proposed two modes of action. First, such TFs can act directly to activate ‘desired’ terminal identity features and repress (also directly) alternative identity features (Lodato et al., 2014; Wyler et al., 2016). Second, neuron type-specific TFs can act indirectly by controlling intermediary factors. For example, in the mouse spinal cord, a complex of three TFs (Isl1, Lhx3, NLI) specifies MN...
identity by recognizing specific DNA elements in the cis-regulatory region of MN-specific genes and the homeodomain TF Hb9, and activates their expression. Hb9 functions as a transcriptional repressor of alternative (V2a interneuron) identity genes, thereby consolidating MN identity (Lee et al., 2008; Song et al., 2009; Thaler et al., 2002). An analogous mechanism operates in V2a interneurons and involves Chx10, a homeodomain protein that represses alternative neuronal identity programs (Clovis et al., 2016; Lee et al., 2008). Hence, mouse mutants for Hb9 or Chx10 result in ectopic expression of alternative identity genes (Arber et al., 1999; Clovis et al., 2016; Thaler et al., 1999). Several TFs (unc-4/UnCx, mab-9/Tbx20, unc-55/COP, bnc-1/BNC) with repressor activity are known to control aspects of cholinergic MN development in C. elegans (Kerk et al., 2017; Pflugrad et al., 1997; Pocock et al., 2008; Von Stetina et al., 2007; Winnier et al., 1999). However, their genetic removal did not result in ectopic expression of alternative (VD, VC) identity features in cholinergic MNs (data not shown). Although we cannot exclude the involvement of yet-to-be identified transcriptional repressors acting downstream of UNC-3, our genetic and biochemical analyses led us to propose the following mechanism underlying UNC-3’s dual role.

In cholinergic MNs, unc-3 and lin-39 are co-expressed, albeit the latter in lower levels (Figure 9A), suggesting that LIN-39/Hox is a rate-limiting factor whose function is determined by UNC-3. In wild-type animals, UNC-3 and LIN-39 occupy cis-regulatory elements of cholinergic MN terminal identity genes, resulting in their activation (Figure 5C) (Kratsios et al., 2012). In the absence of UNC-3, LIN-39 is released from these elements and becomes available to activate alternative identity genes, such as VD-specific terminal identity genes. Several lines of evidence support this conclusion. First, ChIP-seq data show that LIN-39 binding is decreased in cholinergic MN gene loci in unc-3 mutants (Figure 5C). Second, our gene dosage experiments show that either lowering unc-3 levels or increasing lin-39 levels in cholinergic MNs results in ectopic activation of VD identity genes (Figure 9B–C). Lastly, we performed an extensive cis-regulatory analysis of one VD-specific gene (oig-1) and identified the element through which LIN-39 acts to induce oig-1 expression in VD-like cells of unc-3 mutants (Figure 8). Together, these data suggest that the role of UNC-3 in cholinergic MNs is not simply to activate gene expression with LIN-39, but also to ‘recruit’ LIN-39 away from promoters of alternative identity genes, thereby antagonizing its ability to activate those genes. Supporting this scenario, ectopic UNC-3 expression in VD neurons results in decreased expression of VD-specific genes (Figure 9C). Given that the mouse ortholog of unc-3, Ebf2, is co-expressed with Hox genes in cholinergic MNs of the spinal cord (Catela et al., 2019; Kratsios et al., 2017), the molecular mechanism described here may be conserved across species. Interestingly, a seminal study recently described a conceptually similar mechanism in the mouse retina, where CRX recruits MEF2D to retina-specific enhancers, resulting in selective activation of photoreceptor genes (Andzelm et al., 2015).

Insights into how neurons maintain their terminal identity features throughout life

Is there a need for mechanisms that continuously prevent expression of alternative identity features in a post-mitotic neuron? Or, do such mechanisms become superfluous once neurons have restricted their developmental potential by committing to a specific terminal identity? This fundamental question is poorly explored, in part due to the fact that most neuron type-specific TFs have been studied during embryonic stages. For example, it is not known whether CRX is continuously required to activate retina-specific enhancers and simultaneously prevent expression of alternative identity genes (Andzelm et al., 2015). Our temporally controlled protein depletion experiments uncovered a continuous requirement for the dual role of UNC-3. Post-embryonic depletion of UNC-3 not only results in failure to maintain cholinergic MN terminal features (Kratsios et al., 2012), but is also accompanied by ectopic expression of alternative identity features (e.g., VD, VC). These findings reveal a simple and economical mechanism that can enable individual neuron types to select and maintain their distinct terminal identity features. That is, the same TF is continuously required - from development throughout life - to not only activate neuron type-specific identity genes, but also prevent expression of alternative identity features.
Maintenance of terminal identity features: A new function of hox proteins in the nervous system

Across model systems, a large body of work on motor neurons and other neuron types has established that, during early development, Hox proteins are required for neuronal diversity, cell survival, axonal path finding and circuit assembly (Baek et al., 2013; Catela et al., 2016; Estacio-Gómez and Díaz-Benjumea, 2014; Estacio-Gómez et al., 2013; Karlsson et al., 2010; Mendelsohn et al., 2017; Miguel-Aliaga and Thor, 2004; Moris-Sanz et al., 2015; Philippidou and Dasen, 2013). However, the function and downstream targets of Hox proteins during post-embryonic stages are largely unknown. Our contributions towards this knowledge gap are twofold. First, we found that the mid-body Hox protein LIN-39 is continuously required, from development through adulthood, to control expression of MN terminal identity genes, thereby revealing a novel role for Hox proteins in maintaining neuronal identity. Second, we uncovered multiple terminal identity genes as downstream targets of LIN-39 in different MN subtypes (cholinergic MNs: acr-2, dbl-1, unc-77, slo-2; VD neurons: oig-1, ser-2, flp-11; VC neurons: srb-16). Since continuous expression of these genes is essential for MN function, these findings may provide a molecular explanation for the uncoordinated locomotion defects observed in lin-39 mutants (Figure 10—figure supplement 1). Given the maintained expression of Hox genes in the adult nervous system of flies, mice and humans (Baek et al., 2013; Takahashi et al., 2004; Hutlet et al., 2016), our findings may be broadly transferable.

Impact on the concept of terminal selector genes

TFs able to broadly activate many distinct terminal identity features of a specific neuron type (e.g., NT biosynthesis components, NT receptors, ion channels, neuropeptides) have been termed ‘terminal selectors’ (Hobert, 2008). Several dozens of terminal selectors have been described thus far in multiple model systems including worms, flies and mice (Hobert, 2011; Hobert, 2016; Hobert and Kratsios, 2019). However, it is unclear whether terminal selectors are also required to prevent expression of alternative identity features. Our findings suggest this to be the case by revealing a dual role for UNC-3, the terminal selector of cholinergic MN identity in C. elegans. In the future, it will be interesting to see whether other terminal selectors also exert a dual role in order to safeguard neuronal terminal identity. Supporting this possibility, Pet-1, the terminal selector of mouse serotonergic neurons has been recently shown to repress several terminal identity genes (Wyler et al., 2016).

Limitations and lessons learned about the control of neuronal terminal identity

The examination of multiple MN terminal identity markers at single-cell resolution enabled us to make an interesting observation. Although all unc-3-depleted nerve cord MNs uniformly lose their cholinergic identity, one subpopulation acquires VD terminal features (‘VD-like’ neurons) and another subpopulation acquires VC terminal features (‘VC-like’ neurons). This intriguing observation may be analogous to findings described in the mammalian neocortex, where genetic removal of the TF Satb2 leads to loss of pyramidal neuron identity (UL1 subtype), and concomitant gain of molecular features specific to two other pyramidal neuron subtypes (DL, UL2) (Britanova et al., 2008). Together, the cases of UNC-3 and Satb2 support the notion that neuron type-specific TFs often suppress features of functionally related neuronal subtypes (Arlotta and Hobert, 2015).

Although our study employs an extensive repertoire of terminal identity markers for distinct MN subtypes, the extent of alternative identity features (e.g., VD, VC) being ectopically expressed in unc-3-depleted MNs remains unknown. Future unbiased transcriptional profiling of unc-3-depleted MNs could help address this issue. In addition, the strong axonal defects in MNs of unc-3 mutants preclude any further attempts to assess whether the observed VD-like and VC-like cells, as defined by molecular markers, also acquire morphological features of VD and VC neurons, respectively (Prasad et al., 1998). However, the VD-like neurons of unc-3 mutants do not acquire GABAergic identity like wild-type VD neurons (Figure 2—figure supplement 1, panel A), arguing against a complete cell fate transformation.
**Evolutionary implications of this study**

Our findings highlight the employment of economical solutions to evolve novel cell types in the nervous system. The same Hox protein (LIN-39) collaborates with distinct terminal selectors in different MNs, and this collaboration determines the specificity of LIN-39/Hox function. In GABAergic (VD) neurons, LIN-39 works together with UNC-30/PITX to control expression of VD terminal identity genes, whereas in cholinergic MNs LIN-39 synergizes with UNC-3 to control cholinergic MN identity genes ([Figure 7F, 9D](#)). We speculate that the unc-3 mutant ‘state’ may constitute the ‘ground state’. That is, the ‘VD-like’ neurons, for example, in unc-3 mutants that express LIN-39 may represent an ancient cell type that was altered to become a new cell type through the recruitment of distinct terminal selectors (conceptual model in [Figure 10C](#)). Hence, the amount of genetic information required for evolution of new cell types is kept to minimum. The recruitment of UNC-30/PITX enabled ‘VD-like’ cells to fully adopt GABAergic VD neuron terminal identity, as evident by the ability of UNC-30/PITX to control expression of GABA synthesis proteins ([Eastman et al., 1999; Jin et al., 1994](#)) ([Figure 7F](#)). Similarly, recruitment of UNC-3 enabled ‘VD-like’ cells to become cholinergic MNs. In this ‘new’ cholinergic cell type, UNC-3 exerts a dual role: it antagonizes the ability of LIN-39 to activate VD-specific genes, and also synergizes with LIN-39 to co-activate cholinergic MN terminal identity genes ([Figure 9D](#)). We hope the strategy described here of a terminal selector preventing a Hox transcriptional switch may provide a conceptual framework for future studies on terminal identity and evolution of neuronal cell types.

**Materials and methods**

**Key resources table**

| Reagent type (species) or resource | Designation | Source or reference | Identifiers | Additional information |
|-----------------------------------|-------------|---------------------|-------------|------------------------|
| Gene (Caenorhabditis elegans)     | unc-3       | Wormbase            | WBGene00006743 |
| Gene (Caenorhabditis elegans)     | unc-30      | Wormbase            | WBGene00006766 |
| Gene (Caenorhabditis elegans)     | lin-39      | Wormbase            | WBGene00003024 |
| Gene (Caenorhabditis elegans)     | mab-5       | Wormbase            | WBGene00003102 |
| Strain, strain background (Caenorhabditis elegans) | unc-3 (n3435) | Bob Horvitz (MIT, Cambridge MA) | MT10785 | Null Allele: deletion |
| Strain, strain background (Caenorhabditis elegans) | unc-30 (e191) | Caenorhabditis Genetics Center | CB845 | Allele: substitution |
| Strain, strain background (Caenorhabditis elegans) | lin-39(n1760)/dpy-17(e164) unc-32(e189) III. | Caenorhabditis Genetics Center | MT4009 | Null Allele: substitution |
| Strain, strain background (Caenorhabditis elegans) | mab-5 (n1239) III; him-5 (e1490) V | Caenorhabditis Genetics Center | CB3531 | Allele: substitution |
| Strain, strain background (Caenorhabditis elegans) | him-8 (e1489) IV | Caenorhabditis Genetics Center | CB1489 | Allele: substitution |

*Continued on next page*
| Reagent type (species) or resource | Designation | Source or reference | Identifiers | Additional information |
|-----------------------------------|-------------|--------------------|------------|------------------------|
| Strain, strain background (Caenorhabditis elegans) | iesi57 II; unc-3 (ot837 [unc-3::mNG::AID]) | Caenorhabditis Genetics Center | OH13988 | CRISPR-generated allele |
| Strain, strain background (Caenorhabditis elegans) | lin-39 [kas9 [lin-39::mNG::AID]] | This paper | KRA110 | See Materials and methods, Section Targeted genome editing |
| Strain, strain background (Caenorhabditis elegans) | iesi57 [left-3prom::tir1] | Caenorhabditis Genetics Center | CA1200 | Genotype: iesi57 II; unc-119(ed3) III. |
| Strain, strain background (Caenorhabditis elegans) | ser-2::gfp | Caenorhabditis Genetics Center | OH2246 | Genotype: otls107 |
| Strain, strain background (Caenorhabditis elegans) | org-1::gfp | Caenorhabditis Genetics Center | OH3955 | Genotype: pha-1(e2123) III; otEx193 |
| Strain, strain background (Caenorhabditis elegans) | ida-1::gfp | Caenorhabditis Genetics Center | BL5717 | Genotype: inls179 II; him-8(e1489) IV |
| Strain, strain background (Caenorhabditis elegans) | glr-5::gfp | Aixa Alfonso (University of Illinois, Chicago IL) | AL270 | Genotype: icls270 X |
| Strain, strain background (Caenorhabditis elegans) | srb-16::gfp | Caenorhabditis Genetics Center | BC14820 | Genotype: dpy-5(e907) I; sEx14820 |
| Strain, strain background (Caenorhabditis elegans) | flp-11::gfp | Caenorhabditis Genetics Center | NY2040 | Genotype: ynlS40 V |
| Strain, strain background (Caenorhabditis elegans) | twk-46::gfp | Caenorhabditis Genetics Center | BC13337 | Genotype: dpy-5(e907) I; sIs12928 V |
| Strain, strain background (Caenorhabditis elegans) | ilyS-4::targfp | This paper | KRA22 | Genotype: pha-1(e2123) III; kasEx22 |
| Strain, strain background (Caenorhabditis elegans) | flp-13::gfp | Caenorhabditis Genetics Center | NY2037 | Genotype: ynlS37 III |
| Strain, strain background (Caenorhabditis elegans) | lin-11::mCherry | Oliver Hobert (Columbia University, New York NY) | OH11954 | Genotype: lin-11::mCherry + myo-2::GFP V |
| Strain, strain background (Caenorhabditis elegans) | klp-4::gfp | Caenorhabditis Genetics Center | BC11799 | Genotype: dpy-5(e907) I; sEx11799 |
| Strain, strain background (Caenorhabditis elegans) | alr-1::egfp | Caenorhabditis Genetics Center | OP200 | Genotype: unc-119(ed3) III; wgls200 X |

Continued on next page
| Reagent type (species) or resource | Designation | Source or reference | Identifiers | Additional information |
|-----------------------------------|-------------|---------------------|-------------|------------------------|
| Strain, strain background (Caenorhabditis elegans) | *irx-1::egfp* | Caenorhabditis Genetics Center | OP536 | Genotype: *unc-119*(tm4063) III; *wgls536* I |
| Strain, strain background (Caenorhabditis elegans) | *del-1::gfp* | Caenorhabditis Genetics Center | NC138 | |
| Strain, strain background (Caenorhabditis elegans) | *acr-2::gfp* | Caenorhabditis Genetics Center | CZ631 | Genotype: *juls14* IV |
| Strain, strain background (Caenorhabditis elegans) | *unc-129::gfp* | Caenorhabditis Genetics Center | evls82b | Genotype: evls82b IV |
| Strain, strain background (Caenorhabditis elegans) | *dbl-1::gfp* | Caenorhabditis Genetics Center | BW1935 | Genotype: *unc-119*(ed3) III; *ctIs43 him-5*(e1490) V |
| Strain, strain background (Caenorhabditis elegans) | *nca-1::gfp* | Caenorhabditis Genetics Center | BC15028 | Genotype: dpy-5(e907) I; sEx15028 |
| Strain, strain background (Caenorhabditis elegans) | *slo-2::gfp* | Caenorhabditis Genetics Center | BC10749 | Genotype: dpy-5(e907) I; sEx10749 |
| Strain, strain background (Caenorhabditis elegans) | *ttr-39::mCherry* | Caenorhabditis Genetics Center | CZ8332 | Genotype: *juls223* IV |
| Strain, strain background (Caenorhabditis elegans) | *cho-1::rfp* | Caenorhabditis Genetics Center | OH13646 | Genotype: *pha-1*(e2123) III; *him-5*(e1490) *otIs544* V |
| Strain, strain background (Caenorhabditis elegans) | *unc-17::gfp* | Caenorhabditis Genetics Center | LX929 | Genotype: *vsIs48* X |
| Strain, strain background (Caenorhabditis elegans) | *unc-25::gfp* | Caenorhabditis Genetics Center | CZ13799 | Genotype: *juls76* II |
| Strain, strain background (Caenorhabditis elegans) | *unc-47::mChOpti* | Caenorhabditis Genetics Center | OH13105 | Genotype: *him-5*(e1490) *otIs564* V |
| Strain, strain background (Caenorhabditis elegans) | *unc-30::gfp* | Caenorhabditis Genetics Center | OP395 | Genotype: *unc-119*(tm4063) III; *wgls395 |
| Strain, strain background (Caenorhabditis elegans) | *ser-2::rfp* | Mark Alkema (University of Massachusetts, Worcester MA) | AL270 | Genotype: *zfIs8* IV |
| Strain, strain background (Caenorhabditis elegans) | *oig-1(fosmid)::GFP* | Caenorhabditis Genetics Center | OH11809 | Genotype: *otls450* |

Continued on next page
| Reagent type (species) or resource | Designation | Source or reference | Identifiers | Additional information |
|----------------------------------|-------------|---------------------|-------------|------------------------|
| Strain, strain background (Caenorhabditis elegans) | *lin-39::gfp* | Caenorhabditis Genetics Center | OP18 | Genotype: unc-119(ed3) III; wgls18 |
| Genetic reagent (Caenorhabditis elegans) | Poig-1_1 kb::gfp | Oliver Hobert (Columbia University, New York NY) | otEx5993; otEx5995 | |
| Genetic reagent (Caenorhabditis elegans) | Poig-1_1.6 kb::gfp | This paper | kasEx147; kasEx148 | See Materials and methods |
| Genetic reagent (Caenorhabditis elegans) | Poig-1_2.6 kb_LIN-39 site #3 DEL::gfp | This paper | kasEx149; kasEx150 | See Materials and methods |
| Genetic reagent (Caenorhabditis elegans) | Poig-1_2.6 kb_LIN-39 site #4 DEL::gfp | This paper | kasEx151; kasEx152 | See Materials and methods |
| Genetic reagent (Caenorhabditis elegans) | Poig-1_125 bp_:tagrfp | This paper | kasEx80; kasEx81; kasEx82 | See Materials and methods |
| Genetic reagent (Caenorhabditis elegans) | Poig-1_ LIN-39 site mut 125 bp_:tagrfp | This paper | kasEx91; kasEx92; kasEx93 | See Materials and methods |
| Genetic reagent (Caenorhabditis elegans) | Punc-3_558bp > *lin-39* RNAi + myo-2::gfp | This paper | kasEx68; kasEx69; kasEx70; kasEx71; kasEx72 | See Materials and methods |
| Genetic reagent (Caenorhabditis elegans) | Punc-3_558bp > *unc-3* RNAi + myo-2::gfp | This paper | kasEx73; kasEx74; kasEx78; kasEx79 | See Materials and methods |
| Genetic reagent (Caenorhabditis elegans) | Punc-3_558bp > *lin-39* cDNA OE + myo-2::gfp | This paper | kasEx35; kasEx36; kasEx37; kasEx76; kasEx77 | See Materials and methods |
| Genetic reagent (Caenorhabditis elegans) | Punc-47 > *unc-3* cDNA + myo-2::gfp | This paper | kasEx75 | See Materials and methods |
| Genetic reagent (Caenorhabditis elegans) | Pcho-1_280bp > *lin-39* cDNA OE + myo-2::gfp | This paper | kasEx38; kasEx39; kasEx41 | See Materials and methods |
| Genetic reagent (Caenorhabditis elegans) | *lin-39* fosmid | WRM0616E11 + myo-2::gfp | kasEx33; kasEx34 | See Materials and methods |
| Antibody | anti-Myc (Rabbit polyclonal) | Abcam | #ab9106; RRID: AB_307014 | 1:1000 dilution |
| Antibody | anti-Flag (Mouse monoclonal) | Sigma | #F3165; RRID: AB_259529 | 1:1000 dilution |
| Antibody | anti-Flag (Rabbit polyclonal) | Sigma | #SA4301135; RRID: AB_2811010 | 1:1000 dilution |
| Antibody | Clean-Blot IP Detection Reagent (Mouse monoclonal) | Thermo Fisher | #21230; RRID: AB_2576514 | See Materials and methods |
| Antibody | Flag antibody coated beads (Mouse monoclonal) | Sigma | #A2220; RRID: AB_10063035 | See Materials and methods |

Continued on next page
Continued

| Reagent type (species or resource) | Designation | Source or reference | Identifiers | Additional information |
|-----------------------------------|-------------|---------------------|-------------|------------------------|
| Antibody                         | anti-FLAG M2 magnetic beads (Mouse monoclonal) | Sigma-Aldrich | M8823; RRID: AB_2637089 | See Materials and methods |
| Recombinant DNA reagent          | pcDNA 3.1(+)-C-Flag (Plasmid) | Genscript | pcDNA 3.1(+) | C-terminus Flag-tagged UNC-3 |
| Recombinant DNA reagent          | pcDNA 3.1(+)-N-Myc (Plasmid) | Genscript | pcDNA 3.1(+) | N-terminus Myc-tagged LIN-39 |
| Recombinant DNA reagent          | Fosmid clone WRM0616aE11 | Source BioScience | WRM0616aE11 | lin-39::GFP fosmid clone |
| Commercial assay or kit          | Gibson Assembly Cloning Kit | NEB | #55105 | |
| Commercial assay or kit          | QiAquick PCR Purification Kit | QIAGEN | #28104 | |
| Commercial assay or kit          | Ampure XP beads | Beckman Coulter Life Sciences | A63881 | |
| Commercial assay or kit          | TOPO XL-2 Complete PCR Cloning Kit | Thermo Fisher | K8050 | |
| Chemical compound, drug          | Auxin (indole-3-acetic acid) | Alfa Aesar | #10196875 | |
| Software, algorithm              | ZEN | ZEISS | Version 2.3.69.1000, Blue edition, RRID: SCR_013672 | |
| Software, algorithm              | Image J | Image J | Version 1.52i, RRID: SCR_003070 | |
| Software, algorithm              | RStudio | RStudio | Version 1.2.5001 | |
| Software, algorithm              | Adobe Photoshop CS6 | Adobe | Version 13.0 x 64 | |
| Software, algorithm              | Adobe Illustrator CS6 | Adobe | Version 16.0.0 x 64 | |

**C. elegans strains**

Worms were grown at 15 °C, 20 °C or 25 °C on nematode growth media (NGM) plates seeded with bacteria (E.coli OP50) as food source (Brenner, 1974).

**Forward genetic screen**

EMS mutagenesis was performed on unc-3 (n3435); yns40 [fpl-11::GFP] animals using standard procedures (Kutscher and Shaham, 2014). Mutagenized L4 animals were visually screened at a dissecting fluorescence microscope for changes in fpl-11::GFP expression in VNC MNs. One mutant (kas1) was retrieved.

**Generation of transgenic reporter animals**

Reporter gene fusions for cis-regulatory analysis of terminal identity genes were made using either PCR fusion (Hobert, 2002) or Gibson Assembly Cloning Kit (NEB #55105). Targeted DNA fragments were fused (ligated) to tagrfp coding sequence, which was followed by unc-54 3’ UTR. The TOPO XL PCR cloning kit was used to introduce the PCR fusion fragments into the pCR-XL-TOPO vector (Invitrogen). Mutations on LIN-39 motifs were introduced via mutagenesis PCR. The product DNA fragments were either injected into young adult pha-1(e2123) hermaphrodites at 50 ng/µl using pha-1 (pBX plasmid) as co-injection marker (50 ng/µl) and further selected for survival, or injected into young adult N2 hermaphrodites at 50 ng/µl (plus 50 ng/µl pBX plasmid) using myo-2::gfp as co-injection marker (3 ng/µl) and further selected for GFP signal.

The fosmid clone WRM0616aE11 (genomic region: III:7519128..7554793) (Source BioScience) that contains the entire lin-39 locus was linearized by restriction enzyme digestion, mixed with sonicated
bacterial genomic DNA (12 ng/μl) and injected into young adult N2 hermaphrodites at 15 ng/μl using myo-2::gfp as co-injection marker (3 ng/μl).

**Generation of transgenic animals for RNAi or over-expression**

The cDNA (for over-expression) or the exon-rich genomic region (for RNAi) of unc-3 and lin-39 were amplified by PCR and then ligated to cholinergic (cho-1, unc-3) or GABAergic (unc-47) MN promoters using Gibson Assembly Cloning Kit (NEB #5510S). For unc-3 RNAi, we targeted exons 2-5 with the following primers: FRW: GTCTGTAAAGATGAGAACCACCGG, RVS: CTGTCATAAAC TGGACGCTTG. For lin-39 RNAi, we targeted exons 3-5 with the following primers: FRW: gtggcattccggaacttaatg, RVS: gaagggcggaaatgtgtgtaaac. For over-expression constructs, DNA products were purified using a PCR purification protocol (QIAGEN), and then injected into young adult WT hermaphrodites at 50 ng/μl together with 50 ng/μl pBS plasmid (filler DNA) and 3 ng/μl of myo-2::gfp (co-injection marker). For RNAi constructs, complementary sense and anti-sense exon-rich genomic regions of unc-3 and lin-39 were PCR purified and injected into young adult WT or unc-3 (n3435) hermaphrodites each at 100 ng/μl with myo-2::gfp as co-injection marker (3 ng/μl) following previously established procedures (Esposito et al., 2007).

**Targeted genome engineering**

To generated the lin-39 (kas9 [lin-39::mNG::AID]) allele, CRISPR/Cas9 genome editing was employed to introduce the mNG::3xFLAG::AID cassette into the lin-39 gene locus before the stop codon. Micro-injection, selection and strain establishment were performed as previously described (Dickinson et al., 2015).

**Temporally-controlled protein degradation**

In the presence of TIR1, AID-tagged proteins are conditionally degraded when exposed to auxin in the presence of TIR1 (Zhang et al., 2015). Animals carrying auxin-inducible alleles of lin-39 (kas9 [lin-39::mNG::AID]) or unc-3 (ot837 [unc-3::mNG::AID]) (Kerk et al., 2017) were crossed with ieSi57 [eft-3prom::tir1] animals that express TIR1 ubiquitously. Auxin (indole-3-acetic acid [IAA]) was dissolved in ethanol (EtOH) to prepare 400 mM stock solutions which were stored at 4˚C for up to one month. NGM agar plates with fully grown OP50 bacteria were coated with auxin solution to a final concentration of 4 mM, and allowed to dry overnight at room temperature. To induce protein degradation, worms of the experimental strains were transferred onto auxin-coated plates and kept at 20˚C. As control, worms were transferred onto EtOH-coated plates instead. Auxin solutions, auxin-coated plates, and experimental plates were shielded from light.

**Microscopy**

Worms were anesthetized using 100 mM of sodium azide (NaN₃) and mounted on a 4% agarose pad on glass slides. Images were taken using an automated fluorescence microscope (Zeiss, Axio Imager. Z2). Acquisition of several z-stack images (each ~1 μm thick) was taken with Zeiss Axioval 503 mono using the ZEN software (Version 2.3.69.1000, Blue edition, RRID:SCR_013672). Representative images are shown following max-projection of 1–8 μm Z-stacks using the maximum intensity projection type. Image reconstruction was performed using Image J software (RRID:SCR_003070; Schindelin et al., 2012).

**Chromatin immunoprecipitation (ChIP)**

ChIP assay was performed as previously described (Yu et al., 2017; Zhong et al., 2010) with the following modifications. Synchronized unc-3 (n3435); lin-39 (kas9 [lin-39::mNG::3xFLAG::AID]) worms at L1 stage were cultured on 10 cm plates seeded with OP50 at 20˚C overnight. Early L3 worms were cross-linked and resuspended in FA buffer supplemented with protease inhibitors (150 mM NaCl, 10 μl 0.1 M PMSF, 100 μl 10% SDS, 500 μl 20% N-Lavroyl sarsosine sodium, 2 tablets of cOmplete ULTRA Protease Inhibitor Cocktail [Roche Cat.# 05892970001] in 10 ml FA buffer). The sample was then sonicated using a Covaris S220 at the following settings: 200 W Peak Incident Power, 20% Duty Factor, 200 Cycles per Burst for 60 s. Samples were transferred to centrifuge tubes and spun at the highest speed for 15 min. The supernatant was transferred to a new tube, and 5% of the material was saved as input and stored at −20˚C. Twenty (20) μl of equilibrated anti-FLAG M2 magnetic
beads (Sigma-Aldrich M8823) were added to the remainder. The lin-39 (kas9 [lin-39::mNG::3xFLAG::AID]) CRISPR-generated allele was used in order to precipitate the immunocomplex comprising the endogenous LIN-39 protein and the bound DNA. The immunocomplex was incubated and rotated overnight at 4˚C. On the next day, the beads were washed at 4˚C twice with 150 mM NaCl FA buffer (5 min each), once with 1M NaCl FA buffer (5 min). The beads were transferred to a new centrifuge tube and washed twice with 500 mM NaCl FA buffer (10 min each), once with TEL buffer (0.25 M LiCl, 1% NP-40, 1% sodium deoxycholate, 1 mM EDTA, 10 mM Tris-HCl, pH 8.0) for 10 min, twice with TE buffer (5 min each). The immunocomplex was then eluted in 200 µl elution buffer (1% SDS in TE with 250 mM NaCl) by incubating at 65˚C for 20 min. The saved input samples were thawed and treated with the ChIP samples as follows. One (1) µl of 20 mg/ml proteinase K was added to each sample and the samples were incubated at 55˚C for 2 hr and then at 65˚C overnight (12–20 hr) to reverse cross-link. The immunoprecipitated DNA was purified with Ampure XP beads (A63881) according to manufacturer’s instructions. Library preparation and Illumina sequencing was performed at the Genomics Core facility of the University of Chicago. The LIN-39 ChIP-Seq data on wild-type animals were generated by the modENCODE project (RRID:SCR_006206).

**Real-time quantitative PCR (qPCR) analysis of ChIP DNA**

ChIP was performed on unc-3 (n3435); wgIs18 (lin-39 fosmid::GFP) animals as described above. qPCR analysis of ChIP DNA was performed to probe enrichment of predicted LIN-39 binding sites at four target genes (acr-2, dbl-1, unc-129, lin-39). Three biological replicates were included. The primers used are provided in 5’–3’ orientation: acr-2 LIN-39 site (FRW: acattcgcaccaacaaagcg; RVS: aaag-gagggacaccagcag), acr-2 3’ UTR (FRW: ttccagccacagttggttg; RVS: atgtcaggctgtgcagagag), dbl-1 LIN-39 site (FRW: gcacaatccctcgggatcaa; RVS: TAAGTTTTGCGCTGCTGCTG), dbl-1 3’ UTR (FRW: ataccgctttctgtgc; RVS: cctgacacatggcagcaaa), unc-129 LIN-39 site (FRW: atctgtgtcttgagcaggg), unc-129 3’ UTR (FRW: tttgcctgctctttccct; RVS: tttgccaagaaacaaagagagcag), lin-39 LIN-39 site (FRW: gacgtctccctctttctcct; RVS: tttgccaagaaacaaagagagcag). The amplification was conducted in a QuantStudio three using the Power SYBR Green PCR Master Mix (ThermoFisher Cat.# 4367659), with the following program: Step 1: 95˚C for 10 min; Step 2: 95˚C for 15 s; Step 3: 60˚C for 1 min. Repeat steps 2–3 for 40 times.

**Motor neuron identification**

Motor neuron (MN) subtypes were identified based on combinations of the following factors: (a) co-localization with fluorescent markers with known expression pattern, (b) invariant cell body position along the ventral nerve cord, or relative to other MN subtypes, (c) MN birth order, and (d) number of MNs that belong to each subtype.

**Bioinformatic analysis**

To predict the UNC-3 binding site (COE motif) in the cis-regulatory region of unc-129, del-1, acr-2, unc-77 and slo-2, we used the MatInspector program from Genomatix (Cartharius et al., 2005) (RRID:SCR_008036). The Position Weight Matrix (PWM) for the LIN-39 binding site is catalogued in the CIS-BP (Catalog of Inferred Sequence Binding Preferences database) (Weirauch et al., 2014). To identify putative LIN-39 sites on the cis-regulatory regions of unc-129, del-1, acr-2, unc-77, slo-2, oig-1, and ser-2, we used FIMO (Find Individual Motif Occurrences) (Grant et al., 2011), which is one of the motif-based sequence analysis tools of the MEME (Multiple Expectation maximization for Motif Elicitation) bioinformatics suite (http://meme-suite.org/). To predict the binding site for the transcription factor UNC-30, we performed FIMO analysis using the UNC-30 binding motif (WNTAATCHH) described in Cinar et al. (2005). The p-value threshold for the analysis was set at p<0.005.

**Automated worm tracking**

Worms were maintained as mixed stage populations by chunking on NGM plates with E. coli OP50 as the food source. The day before tracking, 30–40 L4 larvae were transferred to a seeded NGM plate and incubated at 20˚C for approximately 24 hr. Five adults are picked from the incubated plates to each of the imaging plates (see below) and allowed to habituate for 30 min before
recording for 15 min. Imaging plates are 35 mm plates with 3.5 mL of low-peptone (0.013% Difco Bacto) NGM agar (2% Bio/Agar, BioGene) to limit bacteria growth. Plates are stored at 4°C for at least two days before use. Imaging plates are seeded with 50 µL of a 1:10 dilution of OP50 in M9 the day before tracking and left to dry overnight with the lid on at room temperature.

**Behavioral feature extraction and analysis**

All videos were analyzed using Tierpsy Tracker (Javer et al., 2018a) to extract each worm’s position and posture over time. These postural data were then converted into a set of behavioral features as previously described (Javer et al., 2018b). From the total set of features, we only considered 48 that are related to midbody posture and motion, as well as the midbody width (see Supplementary file 3 for feature descriptions and their average values for each strain). For each strain comparison, we performed unpaired two-sample t-tests independently for each feature. The false discovery rate was controlled at 5% across all strain and feature comparisons using the Benjamini Yekutieli procedure (Kim and van de Wiel, 2008). The p-value threshold to control the false discovery rate at 0.05 is 0.0032.

**Cloning, western blot, and immunoprecipitation**

UNC-3, and LIN-39 cDNAs were cloned into the mammalian expression vectors pcDNA 3.1(+)C-Flag plasmid and the pcDNA 3.1(+)N-Myc plasmid by GeneScript, to generate C-terminus Flag-tagged UNC-3 and N-terminus Myc-tagged LIN-39. The constructs were verified by sequencing at the sequencing core facility of University of Chicago. The tagged proteins were expressed in HEK293 cells. Protein expression was detected by standard western blot. Expression of Myc tagged LIN-39 was detected using anti-Myc (Abcam, #ab9106, RRID:AB_307014), expression of Flag-tagged UNC-3 in the total cell lysate was detected using mouse anti-Flag (Sigma, #F3165, RRID:AB_259529), expression of Flag-tagged UNC-3 in the IP was detected using rabbit anti-Flag (Sigma, #SAB4301135, RRID:AB_2811010). Immunoprecipitation of Flag-tagged UNC-3 was performed using Flag antibody coated beads (Sigma, #A2220). For the IP, the ‘Clean-Blot IP Detection Reagent’ (Thermo Fisher, #21230) was used as secondary antibody.

**Quantification of fluorescence intensity**

Images of worms carrying the lin-39::mNG::3xFLAG::AID or unc-3::mNG::3xFLAG::AID alleles were taken on the same slide with the same camera settings at the same development stage. Acquisition of four z-stack images (each 0.53 µm thick) covering the middle portion of targeted MN cell bodies was taken with Zeiss Axiocam 503 mono using the ZEN software (Version 2.3.69.1000, Blue edition, RRID:SCR_013672). Image reconstruction was performed using Image J software following average-projection the Z-stacks using the average intensity projection protocol. The chosen cells for quantification of mNG fluorescence intensity for both genotypes are the same 10 cholinergic MNs: AS2, DB3, DA2, VA3, VB4, AS3, DA3, VA4, VB5 and DB4. Targeted cell areas were manually selected with minimum background as region of interest (ROI) and the total fluorescence intensity was measured, calculated, and then represented by Image J as Integrated Density – IntDen (ROI). Background was additionally selected and IntDen (Background) was calculated. The net fluorescence intensity increase is represented as NetIncr = IntDen(ROI)/Area(ROI) – IntDen (Background)/Area (Background). 12 NetIncrs of both genotypes were calculated and data were normalized by dividing the Median of NetIncr (lin-39::mNG::3xFLAG::AID) for better contrast and presented as arbitrary units (a.u).

**Statistical analysis**

For data quantification, graphs show values expressed as mean ± standard deviation (STDEV). The statistical analyses were performed using the unpaired t-test (two-tailed). Calculations were performed using the Evan’s Awesome A/B Tools online software (https://www.evanmiller.org/ab-testing/t-test.html). Differences with p<0.05 were considered significant. Quantifications are provided in the form of box-and-whisker plots (Tukey boxplot) with individual data point dot-plotted. In all boxplots, middle horizontal line represents the median value (equals to Q2). The box illustrates the interquartile range (IQR), that is from Q1 to Q3. The upper limit indicates either the maximum
value if maximum < $Q_3 + 1.5 \times IQR$, or the value that is not higher than $Q_3 + 1.5 \times IQR$. Similarly, the lower limit indicates either the minimum value or the value that is not lower than $Q_1 - 1.5 \times IQR$.

Acknowledgements
We thank the Caenorhabditis Genetics Center (CGC), which is funded by NIH Office of Research Infrastructure Programs (P40 OD010440), for providing strains. We thank Anthony Osuma, Melanie Le Gouez, and Minhkhoi Nguyen for generating lin-39 and oig-1 reporter strains. We are grateful to Oliver Hobert, Elizabeth Hecksher, Robert Carillo, Catarina Catela, and Daniele Canzio for comments on this manuscript. This work was funded by an NINDS grant (R00NS084988) and a Whitehall Foundation grant to PK.

Additional information

**Funding**

| Funder                                           | Grant reference number               | Author               |
|--------------------------------------------------|--------------------------------------|----------------------|
| National Institute of Neurological Disorders and Stroke | K99/R00: Pathway to Independence Award | Paschalis Kratsios   |
| Whitehall Foundation                             | 2017-12-50                           | Paschalis Kratsios   |

The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

**Author contributions**

Weidong Feng, Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Writing—review and editing; Yinan Li, Pauline Dao, Formal analysis, Validation, Investigation, Visualization, Writing—review and editing; Jihad Aburas, Priota Islam, Benayahu Elbaz, Anna Kolarzyk, Investigation; André EX Brown, Project administration, Writing—review and editing; Paschalis Kratsios, Conceptualization, Supervision, Funding acquisition, Investigation, Writing—original draft, Project administration, Writing—review and editing

**Author ORCIDs**

Paschalis Kratsios [https://orcid.org/0000-0002-1363-9271](https://orcid.org/0000-0002-1363-9271)

**Decision letter and Author response**

**Decision letter** [https://doi.org/10.7554/eLife.50065.sa1](https://doi.org/10.7554/eLife.50065.sa1)
**Author response** [https://doi.org/10.7554/eLife.50065.sa2](https://doi.org/10.7554/eLife.50065.sa2)

**Additional files**

**Supplementary files**

- Supplementary file 1. UNC-3 binding sites (COE motifs) are not found in the cis-regulatory region of VD- and VC-expressed terminal identity genes.
- Supplementary file 2. LIN-39/Hox targets in cholinergic and GABAergic (VD) motor neurons. Asterisk (*) highlights novel LIN-39 targets; N. D: Not Determined. The selected cis-regulatory regions are LIN-39 ChIP-seq peaks that fall within the DNA sequence used for our reporter gene constructs (except for del-1). The UNC-3 binding sites (COE motifs 23 bp) have been previously described in Kratsios et al. (2012). The LIN-39 binding sites were predicted by a FIMO search (p<0.005). The UNC-30 binding site on ser-2 locus was predicted by a FIMO search. The UNC-30 site on oig-1 was experimentally validated in Howell et al. (2015).
- Supplementary file 3. Locomotion features assessed by automated worm tracking analysis.
- Transparent reporting form
References

Aboobaker AA, Blaxter ML. 2003. Hox gene loss during dynamic evolution of the nematode cluster. Current Biology 13:37–40. DOI: https://doi.org/10.1016/S0960-9822(02)01399-4, PMID: 12526742

Andzelm MM, Cherry TJ, Harmin DA, Boeke AC, Lee C, Hemberg M, Pawlyk B, Malik AN, Flavell SW, Sandberg MA, Raviole E, Greenberg ME. 2015. MEF2D drives photo receptor development through a genome-wide competition for tissue-specific enhancers. Neuron 86:247–263. DOI: https://doi.org/10.1016/j.neuron.2015.02.038, PMID: 25801704

Arber S, Han B, Mendelsohn S, Smith M, Jessell TM, Sockanathan S. 1999. Requirement for the homeobox gene Hb9 in the consolidation of motor neuron identity. Neuron 23:659–674. DOI: https://doi.org/10.1016/S0896-6273(01)80206-X, PMID: 10482234

Arlotta P, Hobert O. 2015. Homeotic Transformations of Neuronal Cell Identities. Trends in Neurosciences 38: 751–762. DOI: https://doi.org/10.1016/j.tins.2015.10.005

Baek M, Enriquez J, Mann RS. 2013. Dual role for Hox genes and Hox co-factors in conferring leg motoneuron survival and identity in Drosophila. Development 140:2027–2038. DOI: https://doi.org/10.1242/dev.090902

Borromeo MD, Meredith DM, Castro DS, Chang JC, Tung K-C, Guillemot F, Johnson JE. 2014. A transcription factor network specifying inhibitory versus excitatory neurons in the dorsal spinal cord. Development 141: 2803–2812. DOI: https://doi.org/10.1242/dev.108666

Boyle AP, Araya CL, Brdlik C, Cayting P, Cheng C, Cheng Y, Gardner K, Hillier LW, Janette J, Jiang L, Kasper D, Kawli T, Kheradpour P, Kundaje A, Li JJ, Ma L, Niu W, Rehm EJ, Rozowsky J, Slattery M, et al. 2014. Comparative analysis of regulatory information and circuits across distant species. Nature 512:453–456. DOI: https://doi.org/10.1038/nature13668, PMID: 25164757

Brenner S. 1974. The genetics of Caenorhabditis elegans. Genetics 77:71–94. PMID: 4366476

Britanova O, de Juan Romero C, Cheung A, Kwan KY, Schwark M, Gyorgy A, Vogel T, Akopov S, Mitkovski M, Agoston D, Sestan N, Molnár Z, Tarabykin V. 2008. Satb2 is a postmitotic determinant for upper-layer neuron specification in the neocortex. Neuron 57:378–392. DOI: https://doi.org/10.1016/j.neuron.2007.12.028, PMID: 18255031

Cartharius K, Frech K, Grote K, Klocke B, Haltmeier M, Klinkenhoff A, Frisch M, Bayerlein M, Werner T. 2005. MatInspector and beyond: promoter analysis based on transcription factor binding sites. Bioinformatics 21: 2933–2942. DOI: https://doi.org/10.1093/bioinformatics/bti473, PMID: 15860560

Catela C, Shin MM, Lee DH, Liu JP, Dassen JS. 2016. Hox proteins coordinate motor neuron differentiation and connectivity programs through ret/Gfrc genes. Cell Reports 14:1901–1915. DOI: https://doi.org/10.1016/j.celrep.2016.01.067, PMID: 26904995

Catela C, Correa E, Wen K, Aburas J, Croli L, Consagale GG, Kratsios P. 2019. An ancient role for Collier/Olf/Ebf (COE)-type transcription factors in axial motor neuron development. Neural Development 14:2. DOI: https://doi.org/10.1186/s13064-018-0125-6

Cheng L, Arata A, Mizuguchi R, Qian Y, Karunaratne A, Gray PA, Arata S, Shirasawa S, Bouchard M, Luo P, Chen CL, Busslinger M, Onimaru H, Ma Q. 2004. Tlx3 and Tlx1 are post-mitotic selector genes determining glutamatergic over GABAergic cell fates. Nature Neuroscience 7:510–517. DOI: https://doi.org/10.1038/nn1221, PMID: 15064766

Cinari H, Kees S, Jin Y. 2005. Expression profiling of GABAergic motor neurons in Caenorhabditis elegans. Current Biology 15:340–346. DOI: https://doi.org/10.1016/j.cub.2005.02.025, PMID: 15723795

Clark SG, Chisholm AD, Horvitz HR. 1993. Control of cell fates in the central body region of C. elegans by the homeobox gene lin-39. Cell 74:43–55. DOI: https://doi.org/10.1016/0092-8674(93)90293-Y, PMID: 8101475

Clovis YM, Seo SY, Kwon JS, Rhee JC, Yeo S, Lee JW, Lee S, Lee SK. 2016. Chx10 consolidates V2a interneuron identity through two distinct gene repression modes. Cell Reports 16:1642–1652. DOI: https://doi.org/10.1016/j.celrep.2016.06.100, PMID: 27477290

Deneris ES, Hobert O. 2014. Maintenance of postmitotic neuronal cell identity. Nature Neuroscience 17:899–907. DOI: https://doi.org/10.1038/nn.3731, PMID: 24926660

Dickinson DJ, Pani AM, Heppert JK, Higgins CD, Goldstein B. 2015. Streamlined genome engineering with a Self-Excising drug selection cassette. Genetics 200:1035–1049. DOI: https://doi.org/10.1534/genetics.115.178335, PMID: 26044593

Eastman C, Horvitz HR, Jin Y. 1999. Coordinated transcriptional regulation of the unc-25 glutamic acid decarboxylase and the unc-47 GABA vesicular transporter by the Caenorhabditis elegans UNC-30 homeodomain protein. The Journal of Neuroscience 19:6225–6234. DOI: https://doi.org/10.1523/JNEUROSCI.19-15-06225.1999, PMID: 10414952

Esposito G, Di Schiavi E, Bergamasco C, Bazzicalupo P. 2007. Efficient and cell specific knock-down of gene function in targeted C. elegans neurons. Gene 395:170–176. DOI: https://doi.org/10.1016/j.gene.2007.03.002, PMID: 17459615

Estacio-Gómez A, Moris-Sanz M, Schäfer AK, Pereira D, Herrero P, Díaz-Benjumea FJ. 2013. Bithorax-complex genes sculpt the pattern of leucokinergic neurons in the Drosophila central nervous system. Development 140: 2139–2148. DOI: https://doi.org/10.1242/dev.090423, PMID: 23633511

Feng et al. eLife 2020;9:e50065. DOI: https://doi.org/10.7554/eLife.50065

Data availability
All data generated or analysed during this study are included in the manuscript and supporting files.
Estacio-Gómez A, Diaz-Benjumeda FJ. 2014. Roles of hox genes in the patterning of the central nervous system of Drosophila. Fly 8:26–32. DOI: https://doi.org/10.14616/fly.27424, PMID: 24406332

Faumont S, Rondeau G, Thiele TR, Lawton KJ, McCormick KE, Sottile M, Griesbeck O, Heckscher ES, Roberts WM, Doe CQ, Lockery SR. 2011. An image-free opto-mechanical system for creating virtual environments and imaging neuronal activity in freely moving Caenorhabditis elegans. PLOS ONE 6:e24666. DOI: https://doi.org/10.1371/journal.pone.0024666, PMID: 21959859

Gordon PM, Hobert O. 2015. A competition mechanism for a homeotic neuron identity transformation in C. elegans. Developmental Cell 34:206–219. DOI: https://doi.org/10.1016/j.devcel.2015.04.023, PMID: 26096732

Grant CE, Bailey TL, Noble WS. 2011. FIMO: scanning for occurrences of a given motif. Bioinformatics 27:1017–1018. DOI: https://doi.org/10.1093/bioinformatics/btr664, PMID: 21330290

Hobert O. 2002. PCR fusion-based approach to create reporter gene constructs for expression analysis in transgenic C. elegans. BioTechniques 32:728–730. DOI: https://doi.org/10.1093/biotechniques/34.4.728, PMID: 11962590

Hobert O. 2008. Regulatory logic of neuronal diversity: terminal selector genes and selector motifs. PNAS 105: 20067–20071. DOI: https://doi.org/10.1073/pnas.0806070105, PMID: 19104055

Hobert O. 2011. Regulation of terminal differentiation programs in the nervous system. Annual Review of Cell and Developmental Biology 27:681–696. DOI: https://doi.org/10.1146/annurev-cellbio-092910-154226, PMID: 21985672

Hobert O. 2016. Terminal selectors of neuronal identity. Current Topics in Developmental Biology 116:455–475. DOI: https://doi.org/10.1016/bs.ctdb.2015.12.007, PMID: 26970634

Hobert O, Kratsios P. 2019. Neuronal identity control by terminal selectors in worms, flies, and chordates. Current Opinion in Neurobiology 56:97–105. DOI: https://doi.org/10.1016/j.conb.2018.12.006, PMID: 30665084

Howell K, White JG, Hobert O. 2015. Spatiotemporal control of a novel synaptic organizer molecule. Nature 523:83–87. DOI: https://doi.org/10.1038/nature14545, PMID: 26083757

Hutlet B, Theys N, Coste C, Abn MT, Doshi-Shagolli K, Lizen B, Gofflot F. 2016. Systematic expression analysis of hox genes at adulthood reveals novel patterns in the central nervous system. Brain Structure and Function 221:1223–1243. DOI: https://doi.org/10.1007/s00429-014-0965-8, PMID: 25527350

Imbrici P, Camerino DC, Tricario D. 2013. Major channels involved in neuropsychiatric disorders and therapeutic perspectives. Frontiers in Genetics 4:76. DOI: https://doi.org/10.3389/fgene.2013.00076, PMID: 23675382

Javer A, Currie M, Lee CW, Hokanson J, Li K, Martineau CN, Yemini E, Grundy LJ, Li C, Ch’ng Q, Schafer WR, Nollen EAA, Kerr R, Brown AEX. 2018a. An open-source platform for analyzing and sharing worm-behavior data. Nature Methods 15:645–646. DOI: https://doi.org/10.1038/s41592-018-0112-1, PMID: 30171234

Javer A, Ripoll-Sánchez L, Brown AEX. 2018b. Powerful and interpretable behavioural features for quantitative phenotyping of Caenorhabditis elegans. Philosophical Transactions of the Royal Society B: Biological Sciences 373:20170375. DOI: https://doi.org/10.1098/rstb.2017.0375

Jin Y, Hoskins R, Horvitz HR. 1994. Control of type-D GABAergic neuron differentiation by C. elegans UNC-30 homeodomain protein. Nature 372:780–783. DOI: https://doi.org/10.1038/372780a0

Kala K, Haugas M, Lillevall K, Guimera J, Wurst W, Salmenlin M, Partanen J. 2009. Gata2 is a tissue-specific post-mitotic selector gene for midbrain GABAergic neurons. Development 136:253–262. DOI: https://doi.org/10.1242/dev.029900

Karlsson D, Baumgardt M, Thor S. 2010. Segment-specific neuronal subtype specification by the integration of anteroposterior and temporal cues. PLOS Biology 8:e1000368. DOI: https://doi.org/10.1371/journal.pbio.1000368, PMID: 20485487

Kerk SY, Kratsios P, Hart M, Mourao R, Hobert O. 2017. Diversification of C. elegans Motor Neuron Identity via Selective Effector Gene Repression. Neuron 93:80–98. DOI: https://doi.org/10.1016/j.neuron.2016.11.036, PMID: 28056346

Kim K, Colosimo ME, Yeung H, Sengupta P. 2005. The UNC-3 olf/EBF protein represses alternate neuronal programs to specify chemosensory neuron identity. Developmental Biology 286:136–148. DOI: https://doi.org/10.1016/j.ydbio.2005.07.024, PMID: 16143223

Kim KL, van de Wiel MA. 2008. Effects of dependence in high-dimensional multiple testing problems. BMC Bioinformatics 9:114. DOI: https://doi.org/10.1186/1471-2105-9-114, PMID: 18298808

Kratsios P, Stolfi A, Levine M, Hobert O. 2012. Coordinated regulation of cholinergic motor neuron traits through a conserved terminal selector gene. Nature Neuroscience 15:205–214. DOI: https://doi.org/10.1038/nn.2989

Kratsios P, Kerk SY, Catela C, Liang J, Vidal B, Bayer EA, Feng W, De La Cruz ED, Croci L, Consalez GG, Gifford DK, Arlotta P. 2014. Gene co-regulation by Fezf2 selects neurotransmitter identity and connectivity of corticospinal neurons. Nature Neuroscience 17:1046–1054. DOI: https://doi.org/10.1038/nn.3757, PMID: 24997765
Lopes R, Verhey van Wijk N, Neves G, Pachnis V. 2012. Transcription factor LIM homeobox 7 (Lhx7) maintains subtype identity of cholinergic interneurons in the mammalian striatum. PNAS 109:3119–3124. DOI: https://doi.org/10.1073/pnas.1109251109, PMID: 22315402

Maloof JN, Kenyon C. 1998. The Hox gene lin-39 is required during C. elegans vulval induction to select the outcome of Ras signaling. Development 125:181–190. PMID: 9486792

Mears AJ, Kondo M, Swain PK, Takada Y, Bush RA, Saunders TL, Sieving PA, Swaroop A. 2001. Nrl is required for rod photoreceptor development. Nature Genetics 29:447–452. DOI: https://doi.org/10.1038/ng774, PMID: 11694879

Mendelsohn Al, Dasen JS, Jessell TM. 2017. Divergent hox coding and evasion of retinoid signaling specifies motor neurons innervating digit muscles. Neuron 93:792–805. DOI: https://doi.org/10.1016/j.neuron.2017.01.017, PMID: 28190640

Miguel-Aliaga I, Thor S. 2004. Segment-specific prevention of pioneer neuron apoptosis by cell-autonomous, postmitotic Hox gene activity. Development 131:6093–6105. DOI: https://doi.org/10.1242/dev.01521

Morey M, Yee SK, Herman T, Nern A, Blanco E, Zipursky SL. 2008. Coordinate control of synaptic-layer specificity and rhodopsins in photoreceptor neurons. Nature 456:795–799. DOI: https://doi.org/10.1038/nature07419

Moris-Sanz M, Estacio-Gomez A, Sanchez-Herrero E, Díaz-Benjumea FJ. 2015. The study of the Bithorax-complex genes in patterning CCAP neurons reveals a temporal control of neuronal differentiation by Abd-B. Biology Open 4:1132–1142. DOI: https://doi.org/10.1242/bio.012872

Nakatani T, Minaki Y, Kumai M, Ono Y. 2007. Helt determines GABAergic over glutamatergic neuronal fate by repressing Ngn genes in the developing mesencephalon. Development 134:2783–2793. DOI: https://doi.org/10.1242/dev.02870

Niu W, Lu ZJ, Zhong M, Sarov M, Murray JI, Brdlik CM, Janette J, Chen C, Alves P, Preston E, Slightham C, Jiang L, Hyman AA, Kim SK, Waterston RH, Gerstein M, Snyder M, Reinke V. 2011. Diverse transcription factor binding features revealed by genome-wide ChIP-seq in C. elegans. Genome Research 21:245–254. DOI: https://doi.org/10.1101/gr.114587.110, PMID: 21177963

Pereira L, Kratsios P, Serrano-Saiz E, Sheftel H, Mayo AE, Hall DH, White JG, LeBoeuf B, Garcia LR, Alon U, Hobert O. 2015. A cellular and regulatory map of the cholinergic nervous system of C. elegans. eLife 4:e12432. DOI: https://doi.org/10.7554/eLife.12432, PMID: 26705699

Pflugrad A, Meir JY, Barnes TM, Miller DM. 1997. The Groucho-like transcription factor UNC-37 functions with the neurospecificity gene unc-4 to govern motor neuron identity in C. elegans. Development 124:1699–1709. PMID: 9165118

Philippidou P, Dasen JS. 2013. Hox genes: choreographers in neural development, architects of circuit organization. Neuron 80:12–34. DOI: https://doi.org/10.1016/j.neuron.2013.09.020, PMID: 24094100

Pocock R, Mione M, Hussain S, Maxwell S, Pontecorvi M, Aslam S, Gerrelli D, Sowden JC, Woollard A. 2008. Neuronal function of Tbx20 conserved from nematodes to vertebrates. Developmental Biology 317:671–685. DOI: https://doi.org/10.1016/j.ydbio.2008.02.015, PMID: 18358469

Portman DS. 2017. Sexual modulation of sex-shared neurons and circuits in Caenorhabditis elegans. Journal of Neuroscience Research 95:527–538. DOI: https://doi.org/10.1002/jnr.23912, PMID: 27870393

Potts MB, Wang DP, Cameron S. 2009. Trithorax, hox, and TALE-class homeodomain proteins ensure cell survival through repression of the BH3-only gene egl-1. Developmental Biology 329:374–385. DOI: https://doi.org/10.1016/j.ydbio.2009.02.022, PMID: 19254707

Prasad BC, Ye B, Zackhary R, Schrader K, Seydoux G, Reed RR. 1998. unc-3, a gene required for axonal guidance in Caenorhabditis elegans, encodes a member of the O/E family of transcription factors. Development 125:1561–1568. PMID: 9502737

Prasad B, Karakuzu O, Reed RR, Cameron S. 2008. unc-3-dependent repression of specific motor neuron fates in Caenorhabditis elegans. Developmental Biology 323:207–215. DOI: https://doi.org/10.1016/j.ydbio.2008.08.029, PMID: 18517768

Rhee HS, Closser M, Guo Y, Bashkirova EV, Tan GC, Gifford DK, Wichterle H. 2016. Expression of terminal effector genes in mammalian neurons is maintained by a dynamic relay of transient enhancers. Neuron 92:1252–1265. DOI: https://doi.org/10.1016/j.neuron.2016.11.037, PMID: 27939581

Sagasti A, Hobert O, Troemel ER, Ruvkun G, Bargmann CI. 1999. Alternative olfactory neuron fates are specified by sequential activation of directly acting Hox genes. Development 125:1714–1724. DOI: https://doi.org/10.1016/S0012-1606(99)00027-9

Salser SJ, Loer CM, Kenyon C. 1993. Multiple HOM-C gene interactions specify cell fates in the nematode central nervous system. Genes & Development 7:1714–1724. DOI: https://doi.org/10.1101/gad.7.9.1714, PMID: 8103754

Schafer WR. 2005. Egg-laying. WormBook. DOI: https://doi.org/10.1895/wormbook.1.38.1, PMID: 18050396

Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Longair M, Pietzsch T, Preibisch S, Rueden C, Saalfeld S, Schmid B, Tinevez JY, White DJ, Hartenstein V, Eliceiri K, Tomancak P, Cardona A. 2012. Fiji: an open-source platform for biological-image analysis. Nature Methods 9:676–682. DOI: https://doi.org/10.1038/nmeth.f.1862, PMID: 22743772

Schindelman G, Whittaker AJ, Thum JY, Gharib S, Sternberg PW. 2006. Initiation of male sperm-transfer behavior in Caenorhabditis elegans requires input from the ventral nerve cord. BMC Biology 4:26. DOI: https://doi.org/10.1186/1741-7007-4-26, PMID: 16911797
Sgadò P, Dunleavy M, Genovesi S, Provenzano G, Bozzi Y. 2011. The role of GABAergic system in neurodevelopmental disorders: a focus on autism and epilepsy. *International Journal of Physiology, Pathophysiology and Pharmacology* 3:223–235. PMID: 21941613

Shan G, Kim K, Li C, Walthall WW. 2005. Convergent genetic programs regulate similarities and differences between related motor neuron classes in Caenorhabditis elegans. *Developmental Biology* 280:494–503. DOI: https://doi.org/10.1016/j.ydbio.2005.01.032, PMID: 15882588

Shibuya K, Misawa S, Ariki K, Nakata M, Kanai K, Yoshiyama Y, Ito K, Isose N, Noto Y, Nasu S, Sekiguchi Y, Fujimaki Y, Ohmori S, Kitamura H, Sato Y, Kuwabara S. 2011. Markedly reduced axonal potassium channel expression in human sporadic amyotrophic lateral sclerosis: an immunohistochemical study. *Experimental Neurology* 232:149–153. DOI: https://doi.org/10.1016/j.expneurol.2011.08.015, PMID: 21906595

Siponen MI, Wisniewska M, Lehtio L, Johansson I, Svensson L, Raszewski G, Nilsson L, Sigvardsson M, Berglund H. 2010. Structural determination of functional domains in early B-cell factor (EBF) family of transcription factors reveals similarities to rel DNA-binding proteins and a novel dimerization motif. *Journal of Biological Chemistry* 285:25875–25879. DOI: https://doi.org/10.1074/jbc.C110.150482, PMID: 20592035

Song M-R, Sun Y, Bryson A, Gill GN, Evans SM, Pfaff SL. 2009. Islet-to-LOM stoichiometries control the function of transcription complexes that specify motor neuron and V2a interneuron identity. Development *136*:2923–2932. DOI: https://doi.org/10.1242/dev.037986

Stefanakis N, Carrera I, Hobert O. 2015. Regulatory Logic of Pan-Neuronal Gene Expression in C. elegans. *Neuron* 87:733–750. DOI: https://doi.org/10.1016/j.neuron.2015.07.031

Takahashi Y, Hamada J, Murakawa T, Takada M, Tada M, Nogami I, Hayashi N, Nakamori S, Monden M, Miyamoto M, Katoh H, Morii T. 2004. Expression profile of 39 HOX genes in normal human adult organs and anaplastic thyroid Cancer cell lines by quantitative real-time RT-PCR system. *Experimental Cell Research* 293:144–153. DOI: https://doi.org/10.1016/j.yexcr.2003.09.024, PMID: 14729064

Thaler J, Harrison K, Sharma K, Lettieri K, Kehrl J, Pfaff SL. 1999. Active suppression of interneuron programs during motor neuron development revealed by analysis of homeodomain factor HB9. *Neuron* 23:657–687. DOI: https://doi.org/10.1016/S0896-6273(01)80027-1, PMID: 10482235

Thaler JP, Lee SK, Jurata LW, Gill GN, Pfaff SL. 2002. LIM factor Lhx3 contributes to the specification of motor neuron and interneuron identity through cell-type-specific protein-protein interactions. *Cell* 110:237–249. DOI: https://doi.org/10.1016/S0092-8674(02)00823-1, PMID: 12150931

Thorvaldsdóttir H, Robinson JT, Mesirov JP. 2013. Integrative genomics viewer (IGV): high-performance genomics data visualization and exploration. *Briefings in Bioinformatics* 14:178–192. DOI: https://doi.org/10.1093/bib/bbs017, PMID: 22517427

Von Stetina SE, Treinin M, Miller DM. 2006. The motor circuit. *International Review of Neurobiology* 69:125–167. DOI: https://doi.org/10.1016/S0077-7742(05)69005-8, PMID: 16492464

Von Stetina SE, Fox RM, Watkins KL, Starich TA, Shaw JE, Miller DM. 2007. UNC-4 represses CEH-12/HB9 to specify synaptic inputs to VA motor neurons in C. elegans. *Genes & Development* 21:332–346. DOI: https://doi.org/10.1101/gad.1502107, PMID: 17289921

Weirauch MT, Yang A, Albu M, Cote AG, Montenegro-Montero A, Drewe P, Najafabadi HS, Lambert SA, Mann I, Cook K, Zheng H, Goity A, Van Bakel H, Lozano JC, Galli M, Lewsey MG, Huang E, Mukherjee T, Chen X, Reece-Hoyes JS, et al. 2014. Determination and inference of eukaryotic transcription factor sequence specificity. *Cell* 158:1431–1443. DOI: https://doi.org/10.1016/j.cell.2014.08.009, PMID: 25215497

White JG, Southgate E, Thomson NJ, Brenner S. 1986. The structure of the nervous system of the nematode Caenorhabditis elegans. *Philosophical Transactions of the Royal Society B: Biological Sciences* 314:1–340. DOI: https://doi.org/10.1098/rstb.1986.0056

Winemaker AR, Meir JY-J, Ross JM, Tavernarakis N, Driscoll M, Ishihara T, Katsura I, Miller DM. 1999. UNC-4/UNC-37-dependent repression of motor neuron-specific genes controls synaptic choice in Caenorhabditis elegans. *Genes & Development* 13:2774–2786. DOI: https://doi.org/10.1101/gad.13.21.2774

Wyler SC, Spencer WC, Green NH, Rood BD, Crawford L, Craige C, Gresch P, McMahon DG, Beck SG, Deneris E. 2016. Pet-1 switches transcriptional targets postnatally to regulate maturation of serotonin neuron excitability. *The Journal of Neuroscience* 36:1758–1774. DOI: https://doi.org/10.1523/JNEUROSCI.3798-15.2016, PMID: 26843655

Yemini E, Jucikas T, Grundy LJ, Brown AE, Schaffer WR. 2013. A database of Caenorhabditis elegans behavioral phenotypes. *Nature Methods* 10:877–879. DOI: https://doi.org/10.1038/nmeth.2560, PMID: 23852451

Yu B, Wang X, Wei S, Fu T, Dzakah EE, Waqas A, Walthall WW, Shan G. 2017. Convergent translational programs regulate cAMP levels in C. elegans GABAergic Motor Neurons. *Developmental Cell* 43:212–226. DOI: https://doi.org/10.1016/j.devcel.2017.09.013

Zheng Z, Ward JD, Cheng Z, Dernburg AF. 2015. The auxin-inducible degradation (AID) system enables versatile conditional protein depletion in C. elegans. *Development* 142:4374–4384. DOI: https://doi.org/10.1242/dev.129635, PMID: 26552885

Zhong M, Niu W, Lu ZJ, Sarov M, Murray JI, Janette J, Raha D, Sheaffer KL, Lam HY, Preston E, Slightham C, Hillier LW, Brock T, Agarwal A, Auerbach R, Hyman AA, Gerstein M, Mango SE, Kim SK, Waterston RH, et al. 2010. Genome-wide identification of binding sites defines distinct functions for Caenorhabditis elegans PHA-4/FOXA in development and environmental response. *PLOS Genetics* 6:e1000848. DOI: https://doi.org/10.1371/journal.pgen.1000848, PMID: 20174564