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Two Hox cofactors, the Meis/Hth homolog UNC-62 and the Pbx/Exd homolog CEH-20, function together during *C. elegans* postembryonic mesodermal development

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Abstract

The TALE homeodomain-containing PBC and MEIS proteins play multiple roles during metazoan development. Mutations in these proteins can cause various disorders, including cancer. In this study, we examined the roles of MEIS proteins in mesoderm development in *C. elegans* using the postembryonic mesodermal M lineage as a model system. We found that the MEIS protein UNC-62 plays essential roles in regulating cell fate specification and differentiation in the M lineage. Furthermore, UNC-62 appears to function together with the PBC protein CEH-20 in regulating these processes. Both *unc-62* and *ceh-20* have overlapping expression patterns within and outside of the M lineage, and they share physical and regulatory interactions. In particular, we found that *ceh-20* is genetically required for the promoter activity of *unc-62*, providing evidence for another layer of regulatory interactions between MEIS and PBC proteins.

Keywords

Meis; Hth; *unc-62*; Pbx; Exd; *ceh-20*; Hox; *C. elegans*; mesoderm

Introduction

Hox genes encode homeodomain (HD)-containing proteins that establish body plan and specify cell fates during embryogenesis and organogenesis (McGinnis and Krumlauf, 1992; Krumlauf, 1994; Pearson et al., 2005). As monomers, Hox proteins bind to DNA with poor selectivity and affinity *in vitro*. However, interactions with other DNA-binding proteins can increase the DNA-binding specificity and affinity of Hox proteins *in vitro* and therefore facilitate Hox proteins to bind distinct *in vivo* biological targets. Several members of the TALE (Three-Amino-Acid Loop Extension) class of homeodomain proteins can function as Hox partners to modulate Hox-dependent transactivation in invertebrates and vertebrates (Mann and Affolter, 1998; Moens and Selleri, 2006). These Hox cofactors are further grouped into PBC and MEIS classes based on the additional motifs located N-terminal to the homeodomain (Bürglin, 1997, 1998; Mukherjee and Bürglin, 2007).

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The IPBC class of homeodomain proteins is represented by the vertebrate Pbx proteins and the *Drosophila* Extradenticle (Exd) protein (Bürglin and Ruvkun, 1992; Rauskolb et al., 1993; Bürglin, 1997, 1998; Mukherjee and Bürglin, 2007). In addition to the homeodomain, this class of proteins contains two other conserved domains, PBC-A and PBC-B, and the PBC-A domain mediates the interaction of PBC proteins with the MEIS subfamily of proteins (Chang et al., 1997; Knoepfler et al., 1997). The MEIS class of homeodomain proteins includes the vertebrate Meis and Prep1 proteins and the *Drosophila* Homothorax (Hth) protein (Bürglin, 1997, 1998; Mukherjee and Bürglin, 2007). Recent studies found that both Hth and Meis1 encode two types of isoforms, one HD-containing and one HD-less (Noro et al., 2006). All MEIS proteins share two domains at their N-termini: HM1 and HM2 (also referred to as MEIS A and MEIS B), which are essential for mediating interactions between MEIS and PBC proteins (Rieckhof et al., 1997; Ryoo et al., 1999, Bürglin, 1997, 1998; Mukherjee and Bürglin, 2007).

PBC and MEIS proteins primarily modulate Hox functions by forming heterodimeric or trimeric complexes with Hox proteins on DNA (For examples, see Maconochie et al., 1997; Kroon et al., 1998; Swift et al., 1998; Ryoo et al., 1999; Shanmugam et al., 1999; Shen et al., 1999; Ferretti et al., 2000; Ebner et al., 2005). In these complexes, the PBC or MEIS proteins either directly bind DNA or serve as non-DNA binding partners to stabilize the complex. PBC and MEIS can also form stable heterodimers with each other in the presence or absence of DNA (Chang et al., 1997; Rieckhof et al., 1997; Abu-Shaar et al., 1999; Berthelsen et al., 1999; Jaw et al., 2000). In addition to their physical interaction, PBC and MEIS proteins also display mutual regulatory interactions. A number of studies have shown that MEIS proteins are required for the nuclear localization and stability of PBC proteins (Pai et al., 1998; Abu-Shaar et al., 1999; Affolter et al., 1999; Berthelsen et al., 1999; Kurant et al., 1998; Stevens and Mann, 2007). Work in *Drosophila* showed that in *exd* mutants, HTH protein levels are undetectable in the developing brain and epidermis and greatly reduced in multiple other embryonic tissues, suggesting that the maintenance of HTH protein level is dependent on EXD activity (Kurant et al., 1998; Nagao et al., 2000).

C. elegans contains both PBC and MEIS proteins. CEH-20 and two divergent members, CEH-40 and CEH-60, comprise the PBC class (Bürglin, 1997, 1998; Van Auken et al., 2002; Mukherjee and Bürglin, 2007). UNC-62 is the HD-containing MEIS1/hth ortholog, while PSA-3 is a HD-less variant of the PREP family that belongs to the ancient MEIS class (Bürglin, 1997; Van Auken et al., 2002, Arata et al., 2006; Mukherjee and Bürglin, 2007). A number of studies have shown that Hox and CEH-20 function in a complex to directly regulate downstream gene expression during mesodermal and vulval development, and in regulating apoptosis (Liu and Fire, 2000; Koh et al., 2002; Shemer and Podbilewicz, 2002; Liu et al., 2006; Takacs-Vellai et al., 2007; Potts et al., 2009). However, the roles of MEIS proteins and their functional interactions with Hox and PBC proteins in *C. elegans* are not well characterized. *unc-62* has been shown to function during embryogenesis, Q neuroblast migration, vulval formation and VC motor neuron survival (Van Auken et al., 2002; Yang et al., 2005; Potts et al., 2009). While *unc-62* interacts genetically with *ceh-20* and *ceh-40* during embryogenesis, these genes do not appear to function together with any of the Hox genes (Van Auken et al., 2002). Similarly, loss of *unc-62* or *ceh-20* activity causes Q cell migration and vulval formation defects different than those seen upon loss of Hox activity, suggesting that UNC-62 and CEH-20 may function independently of Hox proteins in regulating these processes (Yang et al., 2005). Finally, the HD-less protein PSA-3 appears to function as a non-DNA binding partner of CEH-20 and the Hox protein NOB-1 in regulating T cell asymmetric cell division (Arata et al., 2006).

We use the *C. elegans* postembryonic mesodermal lineage, the M lineage, as a model system to investigate the roles of MEIS proteins and their relationships with PBC and Hox proteins during mesodermal development. The *C. elegans* M lineage originates from a single embryonic

pluripotent precursor, the M mesoblast (Sulston and Horvitz, 1977). At the first larval stage (L1) in hermaphrodites, the M lineage undergoes four to five rounds of cell division and produces 18 descendants, including two non-muscle coelomocytes (CCs), 14 striated bodywall muscles (BWMs) and two sex myoblasts (SMs) (Fig. 1A). The two SMs migrate from the posterior to the presumptive vulval region and during the L3 stage divide three times to generate 16 precursor cells that eventually differentiate into four classes of non-striated sex muscles required for egg-laying: type I and II vulval muscles (VM1 and VM2) and type I and II uterine muscles (UM1 and UM2). Vulval muscles form tubular muscles and attach to the vulva, while uterine muscles ingress, form thin sheets and attach to the uterus (Sulston and Horvitz, 1977).

Previous studies have shown that two medial group Hox genes, *mab-5* and *lin-39*, share redundant roles in the M lineage, and that they function together with the Hox cofactor *ceh-20* in regulating the specification and diversification of the M lineage (Liu and Fire, 2000). One of the direct target genes of the Hox/CEH-20 complex is the *hlh-8* gene, which encodes the *C. elegans* Twist ortholog and is expressed in all undifferentiated cells in the M lineage (Harfe et al., 1998; Liu and Fire, 2000). The homeobox gene *mls-2* is likely another direct target of CEH-20, although the activation of *mls-2* by CEH-20 appears to be Hox-independent (Jiang et al., 2005; Jiang et al., 2008).

In this study, we set out to determine the functions of MEIS proteins in M lineage development. We showed that while *psa-3* does not have an apparent role in the M lineage, *unc-62* has multiple functions throughout M lineage development. Furthermore, *unc-62* genetically interacts with *ceh-20* and *mab-5* to regulate the expression of downstream target genes such as *hlh-8*. Finally, we provide evidence that *unc-62* and *ceh-20* can bind to each other in the yeast two-hybrid system and that they exhibit mutual regulatory interactions: while *unc-62* is required for the nuclear accumulation of CEH-20, *ceh-20* is required for the promoter activity of *unc-62*.

Materials and Methods

C. elegans strains

Strains were manipulated under standard conditions as described by Brenner (1974). Analyses were performed at 20°C, unless otherwise noted. The wild-type reference strain was LW0081 [*ccIs4438 (intrinsic CC::gfp) III; ayIs2 (egl-15::gfp) IV; ayIs6 (hlh-8::gfp) X*] (Jiang et al., 2005). Other M lineage-specific reporters, including *Nde-box::gfp*, *arg-1::gfp*, *rgs-2::gfp* and *mls-2::lacZ*, were as described in Kostas and Fire (2002). The mutant alleles used in this work are: LGIII, *ceh-20(ay9, ay42, n2513, ok541, os39, os114)*, *unc-36(e251)* and *unc-119(ed3)*; LGV, *unc-62(e917, e644, ku234)*, *dpy-11(e224)* and *him-5(e1467)*; LGX, *psa-3(os8)*.

RNA interference

Plasmids to make dsRNA are: p15.121A (Wang et al., 1993) for *lin-39*, pJKL422.1 (Liu and Fire, 2000) for *ceh-20*, and *yk414a1* (gift from Yuji Kohara, National Institute of Genetics, Japan) for *unc-62*. For *mab-5* RNAi, primers JKL-601 and JKL-602 were used to amplify the insert in the *mab-5* RNAi clone III-4I09 from the Ahringer library (Kamath et al., 2003). dsRNA synthesis was following the protocol of Fire and colleagues (Fire et al., 1998). For *ceh-20*, *mab-5* or *lin-39* RNAi, dsRNA was injected into gravid adults of different genotypes. Progeny of injected animals were scored for M lineage and vulval defects. Water-injected animals were used as controls. RNAi soaking was used for *unc-62(RNAi-soaking)* by modifying the soaking protocol described in Ahringer (2006). Embryos and the subsequent newly hatched L1s were soaked in dsRNA against *unc-62* or in soaking buffer alone for 24-48 hours at 20°C. The soaked L1 worms were transferred to NGM plates with OP50 and followed through development for M lineage phenotypes or *gfp* expression.

Transgenic animals and *mls-1::lacZ* expression

2.9 kb of the *unc-62* promoter sequence (-2883 to -1 exon 1a) were amplified using iProof DNA polymerase (Bio-Rad Laboratories) using genomic DNA as template and YJ-178 and YJ-179 as primers. The PCR products were used to generate a reporter construct, pYJ167 (-2.9 kb *unc-62p::gfp::let-858* 3' UTR). pJKL790 and pJKL791 are *unc-62* rescuing constructs that contain the *unc-62* promoter driving *unc-62* 1a-7b and *unc-62* 1b-7b cDNAs respectively.

pHK110 (a gift from Thomas Burglin, Karolinska Institutet, Sweden) and pJKL775 were used for analyzing *ceh-20* expression pattern. pHK110 contains 2.4 kb of the *ceh-20* promoter sequence (-2385 to -1), the entire *ceh-20* coding region fused with *gfp* sequence at the C-terminus, and the *unc-54* 3' UTR. pJKL775 is a transcriptional reporter of *ceh-20* with *nls::gfp* under the control of the *ceh-20* promoter (-2385 to -1).

Reporters were microinjected with the plasmid pRF4 (Mello et al., 1991) or the *pha-1* rescuing plasmid pC1 (Granato et al., 1994) as markers using standard techniques. LW1175 was a spontaneous integrant of pHK110. pSAK534 (*mls-1::lacZ*) (Kostas and Fire, 2002) was microinjected with pRF4 and pJKL449.1 (*myo-2::gfp*) into N2 worms using standard techniques (Mello et al., 1991). Animals transgenic for *jjEx[pSAK534.1(mls-1::lacZ)+rol-6(su1006)+pJKL449.1(myo-2::gfp)]* were crossed to animals of *unc-62(ku234)V*; *ayIs6(hlh-8::gfp)* X to generate *unc-62(ku234)V*; *ayIs6(hlh-8::gfp)* X; *jjEx[pSAK534.1(mls-1::lacZ)+rol-6(su1006)+pJKL449.1(myo-2::gfp)]* animals. L3 larvae were fixed following the protocol of Harfe and colleagues (Harfe et al., 1998) and used for immunostaining. The following antibodies were used: goat anti-GFP antibodies (Rockland Immunochemicals) (1:5000), mouse anti- β -galactosidase antibodies (Promega) (1:50), Cy3-conjugated donkey anti-mouse antibodies (Jackson ImmunoResearch Laboratories) (1:100) and FITC-conjugated donkey anti-goat antibodies (Jackson ImmunoResearch Laboratories) (1:100).

Yeast two-hybrid assay

Two-hybrid analysis was performed using the protocol described by James and colleagues (James et al., 1996). The following plasmids were used in the two-hybrid assay:

pYJ188: *ceh-20* aa1-196 in pGAD-C2

pJKL841: *ceh-20* aa1-196 in pGBD-C2

pJKL842: *unc-62* isoform 1a7b aa1-287 in pGBD-C2

pJKL843: *unc-62* isoform 1a7b aa1-287 in pGAD-C2

pJKL844: *unc-62* isoform 1b7b aa1-250 in pGAD-C1

pJKL845: *unc-62* isoform 1b7b aa1-250 in pGBD-C1

Plasmids were transformed into the yeast strain PJ69-4A and selected for on -Trp (pYJ188, pJKL843 and pJKL844) or -Leu (pJKL841, pJKL842 and pJKL845) plates. Transformed strains were tested for auto-activation (growth) on -Ade plates. Transformed yeast colonies were mated in liquid culture and plated in serial dilutions on control (-Leu -Trp) and selection (-Leu -Trp -Ade) plates and grown at 30°C for 2-3 days before being scored for growth.

Results

The Meis protein UNC-62 plays multiple roles in the postembryonic M lineage

C. elegans contains two Meis-related genes, *unc-62* and *psa-3* (Van Auken et al., 2002 and Arata et al., 2006). We first examined the role of *psa-3* in the M lineage. Neither *psa-3* (*RNAi*) animals nor mutant animals homozygous for a loss-of-function *psa-3(os8)* allele (Arata

et al., 2006) exhibited any M lineage defects, suggesting that *psa-3* is not likely to play a role in M lineage development. In contrast, animals with reduced levels of *unc-62* displayed multiple M lineage defects.

There are multiple *unc-62* alleles available, and the majority of them cause either zygotic or maternal-effect embryonic lethality (Van Auken et al., 2002). However, two weak zygotic alleles, *ku234* and *e644*, can produce viable progeny. *ku234* has a point mutation specifically affecting the start codon of the 1b isoforms, changing the first amino acid from Met to Ile, while *e644* has an early stop codon in the homeodomain region encoded by exon 7b (Van Auken et al., 2002). We introduced the M lineage specific GFP markers into *ku234* and *e644* mutants and examined their M lineage phenotypes.

The early M lineage appeared completely normal in *ku234* mutants, producing two SMs, two CCs and fourteen BWMs (Table 1, Figure 1E). Just like in wild-type animals, the two SMs in *ku234* mutants migrated to the presumptive vulval region and underwent three rounds of division to produce 16 *hlh-8::gfp*-positive precursor cells. However, unlike wild-type animals in which the 16 sex muscle precursors differentiate into 8 vulval muscles (4 VM1s and 4 VM2s) and 8 uterine muscles (4 UM1s and 4 UM2s), in 100% (n=200) of *ku234* mutants examined, the majority (ranging between 9 and 14) of the 16 sex muscle precursor cells differentiated into VM1-like cells that expressed *egl-15::gfp* (Fig. 1E, compare Figure 2A-B). These extra VM1-like cells appeared to be a result of fate transformation from UMs and VM2s to VM1s based on the following observations. First, in wild-type animals, *Nde-box::gfp* labels all 16 differentiated sex muscles while highlighting the distinct cell morphologies of VMs and UMs (Harfe et al., 1998; Fig. 2C). *ku234* mutants had 16 *Nde-box::gfp*-expressing cells; however, there was an increase of VM-like cells and a concomitant decrease of UM-like cells (Fig. 2D). Second, *rgs-2::gfp*, which preferentially labels UMs but also faintly labels VMs in wild-type animals (Kostas and Fire, 2002; Fig. 2E), showed an increase of the faint VM-like signal and a decrease of the bright UM-like signal in *ku234* mutants (Fig. 2F). Finally, all the eight vulval muscles (both VM1s and VM2s) in wild-type animals express *arg-1::gfp* (Kostas and Fire, 2002; Fig. 2G). *ku234* mutants had 10 to 14 *arg-1::gfp*-positive cells that all exhibited the morphology of VM1-like cells (Fig. 3H). Thus, most of the UMs and VM2s are transformed to VM1s in *ku234* mutants.

Previous work by Kostas and Fire (2002) showed that the T-box gene *mls-1* is a cell fate determinant of uterine muscles (UMs): *mls-1* is specifically expressed in the UM and VM2 precursor cells (Fig. 2I) and loss of *mls-1* activity results in the doubling of the numbers of both VM1s and VM2s due to a fate transformation of UMs to VMs. The *ku234* mutant phenotype is similar, but not identical, to that of *mls-1* mutants. We therefore tested if *mls-1* expression is affected in *unc-62(ku234)* mutants. We detected normal expression of a functional *mls-1::lacZ* reporter in the SM lineage of *ku234* mutants (Fig. 2J), suggesting that the UM to VM fate transformation in *ku234* mutants is not due to the loss of or altered *mls-1* expression.

Unlike *ku234* mutants, *e644* mutants exhibited variable M lineage defects. We used *hlh-8::gfp* to follow the M lineage in *e644* mutants as *hlh-8::gfp* expression appeared unaffected in *e644* mutants. Among 200 *e644* worms examined, 22% of them had fewer (0-1) M-derived CCs and extra (3-4) SMs (Table 1). Direct observation M lineage cell divisions in three *e644* animals showed that while M lineage patterning and proliferation were normal, both M-derived CCs and 1 or 2 M-derived BWMs were transformed to SMs (data not shown). The remaining 78% of *e644* animals contained normal numbers of SMs; however, 69% of these mutant animals with 2 SMs (n=69) gave rise to extra VM1s, ranging from 5 to 10 (Fig. 1D, Table 1). Using the cell type specific GFP markers that were used for analyzing the *ku234* mutants, we found that these extra VM1s were produced at the expense of UMs and VM2s (data not shown),

similar to the phenotypes observed in the *ku234* mutants. The multiple M lineage defects in *e644* mutants suggest that *unc-62* is not only involved in UM and VM decisions in the SM lineage, but also involved in CC, SM and BWM cell fate decisions in the early M lineage.

Because both *ku234* and *e644* are partial loss-of-function alleles of *unc-62* and *unc-62* is essential for embryogenesis (Van Auken et al., 2002), we also soaked synchronized L1 wild-type worms with dsRNA against *unc-62* (referred to as *unc-62(RNAi-soaking)*, see Materials and Methods) and examined the M lineage phenotypes in the soaked animals. Soaking buffer alone was used as a negative control. Unlike the control animals, *unc-62(RNAi-soaking)* worms exhibited defects in the vulva and the somatic gonad (data not shown), as well as in the M lineage. The M lineage phenotypes can be grouped into two categories. The majority of *unc-62(RNAi-soaking)* worms (87%, n=137) had no differentiated M-derived CCs, VM1s, or BWMs (Fig. 1F), or occasionally no M-derived CCs and BWMs, but with 1-4 elongated *egl-15::gfp*-positive cells (data not shown). These phenotypes resemble phenotypes observed in *ceh-20(RNAi)* or *lin-39(0) mab-5(0)* mutants (Liu and Fire, 2000). Unlike *ceh-20(RNAi)* or *lin-39(0) mab-5(0)* mutants in which *hlh-8::gfp* expression is undetectable throughout postembryonic development (Liu and Fire, 2000), *unc-62(RNAi-soaking)* worms showed normal *hlh-8::gfp* expression in the early M lineage during the L1 stage when CC, SM and BWM fates are specified. The early expression of *hlh-8::gfp* in the *unc-62(RNAi-soaking)* worms is likely due to the timing of soaking being too late to affect the early M lineage. The remaining 13% of *unc-62(RNAi-soaking)* animals had *hlh-8::gfp* expression throughout the M lineage, but exhibited fate transformation from one or both M-derived CCs to SMs (Fig. 1G) and randomized division orientation in the SM lineage (data not shown).

The range of M lineage defects observed in *unc-62(RNAi-soaking)* animals and in *ku234* and *e644* mutants suggest that *unc-62* plays multiple roles at multiple stages in the M lineage, including regulating CC/SM/BWM fate specification and differentiation, and VM/UM fate specification.

Analysis of a weak *ceh-20* allele, *ay9*, revealed multiple roles of the PBC protein CEH-20 in the M lineage

We have previously shown that CEH-20, but not CEH-40, is the PBC protein that functions in the M lineage (Liu and Fire, 2000; Jiang et al., 2008). Furthermore, CEH-20 is required for the diversification of the M lineage (Liu and Fire, 2000). These previous studies are based on analyses of *ceh-20(RNAi)* and a strong loss-of-function allele *n2513*. We extended our analysis to additional alleles of *ceh-20*, including *os114* and *os39* (Arata et al., 2006), and *ay42* and *ay9* (Takacs-Vellai et al., 2007). Among these alleles, *os114* and *os39* appear to be the strongest loss-of-function, and possibly null, alleles of *ceh-20*, as 99% of *os114* and *os39* mutant animals (n=87 for *os114* and n=56 for *os39*) failed to generate any M lineage descendants, consistent with previous studies of *ceh-20(RNAi)* or *n2513* (Liu and Fire, 2000). *ay42* is likely a partial loss-of-function allele as 62% of *ay42* worms (n=224) failed to produce any M lineage products (data not shown).

ay9 contains a missense mutation, changing a highly conserved Met78 residue to Ile (Takacs-Vellai K. et al., 2006), and exhibits multiple M lineage phenotypes. In *ay9* mutant animals, *hlh-8::gfp* expression was detectable throughout the M lineage; however, compared to wild-type worms, the intensity of *hlh-8::gfp* signal was moderately decreased throughout the M lineage in 20% of *ay9* mutants (n=200) (Fig. 3A-D). All *ay9* mutants exhibited normal M lineage patterning and proliferation; however, 9% of *ay9* animals examined (n=200) exhibited earlier defects in CC/SM/BWM fate decisions, resulting in variable numbers of M-derived CCs, SMs, and BWMs in *ay9* mutants (Table 1). In addition to these earlier M lineage defects, *ay9* mutants also exhibited defects in the SM lineage. 56% of the *ay9* animals (n=75) that contained the normal number of SMs exhibited fate transformations from UMs and VM2s to

VM1s, resulting in an increase of the number of *egl-15::gfp*-positive VM1s (5-10 VM1s) in *ay9* mutants (Table 1, Fig. 3F). VM1s often did not display proper orientation and attachment (Fig. 3E), likely due to the vulval defects in *ay9* mutants.

The broad range of M lineage defects seen in the various *ceh-20* mutant animals suggest that like *unc-62*, *ceh-20* plays multiple roles at multiple stages in the M lineage, including regulating CC/SM/BWM fate specification, VM/UM fate specification and *hlh-8* expression.

***unc-62* genetically interacts with *ceh-20* and *mab-5* to regulate *hlh-8*/CeTwist expression and cell fate specification in the M lineage**

The similar M lineage defects caused by mutations or RNAi of *unc-62* and *ceh-20* suggest that *unc-62* and *ceh-20* may function together to regulate proper M lineage development. To test this hypothesis, we generated double mutants between *ceh-20(ay9)* and *unc-62(e644)* or *unc-62(ku234)*.

Compared to *unc-62(e644)* single mutants, *ceh-20(ay9)/+; unc-62(e644)/unc-62(e644)* animals showed a roughly 20% increase (from 22% to 45%) in penetrance of fate transformations from M-derived CCs to SMs (Table 1). *ceh-20(ay9); unc-62(e644)* double homozygous animals exhibited significantly more penetrant and severe M lineage defects compared to *ceh-20(ay9)* and *unc-62(e644)* single mutants (Table 1). First, unlike in each single mutant, *hlh-8::gfp* was undetectable (97%) or very faint (3%, n=200) throughout the M lineage in *ceh-20(ay9); unc-62(e644)* double mutants (Fig. 3G-H). Second, 100% of *ceh-20(ay9); unc-62(e644)* double mutants (n=83) had no M-derived CCs, compared to 9% (n=200) of *ceh-20(ay9)* animals and 22% (n=200) of *unc-62(e644)* animals that had a CC to SM fate transformation (Table 1, Fig. 3K). Thus there appeared to be a dose-dependent effect of *ceh-20* on *unc-62(e644)* mutants regarding *hlh-8* expression and early M lineage fate specification. Similar synergistic effect on M lineage development was also observed between *ceh-20(ay9)* and *unc-62(ku234)* (Table 1, Fig. 3I, J, L). These observations are consistent with *ceh-20* and *unc-62* functioning together in regulating fate specification and *hlh-8::gfp* expression during M lineage development. In addition to the M lineage defects, we also observed enhanced penetrance of embryonic lethality and vulval defects in *ceh-20; unc-62* double mutants (data not shown), suggesting that these two genes function together in multiple tissues at multiple developmental stages.

In addition to the genetic interactions between *unc-62* and *ceh-20*, we also observed genetic interactions between *unc-62* and the Hox gene *mab-5*. *mab-5(0)* or *mab-5(RNAi)* mutants lack *hlh-8::gfp* expression in the early M lineage, but show normal *hlh-8::gfp* expression in the SM lineage. Furthermore, the M-derived CCs and 1-3 M-derived BWMs are transformed to SMs in *mab-5(0)* or *mab-5(RNAi)* mutants (Harfe et al., 1998; Liu and Fire, 2000; Amin et al., 2007, Table 1). 41% (n=85) of *mab-5(RNAi); unc-62(e644)* mutant animals showed no *hlh-8::gfp* expression throughout the M lineage, lacked all M-derived CCs, and had zero or 1-5 elongated *egl-15::gfp* positive cells. These phenotypes resemble those observed in *lin-39(0) mab-5(0)* and *ceh-20(RNAi)* animals (Liu and Fire, 2000). These observations suggest that in addition to *ceh-20*, *unc-62* also functions together with the Hox gene *mab-5* to regulate *hlh-8::gfp* expression and M lineage fate specification. Similar synergistic interactions were also observed in *mab-5(RNAi) ceh-20(ay9)* animals (Table 1), consistent with previous findings on the cooperative actions of *mab-5* and *ceh-20* in regulating *hlh-8* expression and M lineage fate specification.

***unc-62* and *ceh-20* are co-expressed throughout the M lineage and in other cell types**

Because *ceh-20* and *unc-62* appear to function together to regulate multiple processes of M lineage development at multiple developmental stages, we next determined when and where these two genes are expressed in the M lineage.

unc-62 uses two alternative start codons, exon 1a or 1b (Van Auken et al., 2002). To determine the temporal and spatial expression pattern of *unc-62*, we generated a transcriptional *gfp* reporter (*unc-62p::gfp*) using 2.9kb sequences upstream of the start codon of exon 1a (see Materials and Methods). We first found that *unc-62* cDNAs under the control of this 2.9kb promoter element can rescue the M lineage defects of *unc-62(e644)* and *unc-62(ku234)* mutants. We then examined the expression pattern of *unc-62p::gfp* in transgenic animals carrying the reporter construct. Similar GFP expression patterns were observed in three independent transgenic lines. *unc-62p::gfp* expression was first detectable during mid-embryogenesis in a subset of cells that include AB lineage derivatives (data not shown). In newly hatched L1 larvae, *unc-62p::gfp* was observed in multiple cell types including the M mesoblast, as well as a few neurons in the head and tail, and dorsal and ventral hypodermis (Fig. 4A-B). In the L1 larva, *unc-62p::gfp* expression continued throughout the M lineage in all dividing M descendants (data not shown). The GFP signal was transiently detectable in M-derived CCs and BWMs before they terminally differentiate, but remained detectable in the SMs (Fig. 4C-D) and throughout the SM lineage (data not shown), including the differentiated vulval muscles (Fig. 4E-F). Outside of the M lineage, the neuronal expression in the head and tail was retained throughout post-embryonic development, while hypodermal expression appeared more variable. *unc-62p::gfp* expression was also observed in a subset of ventral nerve cord (VNC) cells after the L2 stage (data not shown) and in the P lineage with initial faint GFP expression in P3.p, P4.p and P5.p in L1 animals, which expanded to P6.p, P7.p and P8.p cells at the L2 stage (Fig. 4C-D). Upon vulval induction, L3 and L4 larvae displayed strong GFP expression in P5.p, P6.p and P7.p and their descendants, but expression in P3.p, P4.p and P8.p and their descendants became weaker and eventually undetectable by mid-L4 stage. *unc-62p::gfp* expression persisted in all differentiated vulval cells after vulval morphogenesis in adults (data not shown).

ceh-20 shared an overlapping, but not identical, expression pattern with *unc-62* both within and outside of the M lineage. Using a functional *ceh-20::gfp* construct (see Materials and Methods), we detected nuclear GFP signal in a wide array of cells outside of the M lineage, including a subset of embryonic cells, hypodermal cells, gut cells, bodywall muscles, VNC neurons and VPCs (data not shown), consistent with previous reports on the expression pattern of *ceh-20* (Yang et al., 2005; Liu et al., 2006; Takacs-Vellai et al., 2006). Within the M lineage, CEH-20::GFP was detected in the M cell (Fig. 4G-H) and throughout the early M lineage. It persisted in the differentiated BWMs as well as in the SMs and all the SM descendants (data not shown). In addition to the translational *gfp* reporter, we also generated a transcriptional *ceh-20p::gfp* reporter (see Materials and Methods). Transgenic animals carrying *ceh-20p::gfp* showed similar expression pattern to the translational *ceh-20::gfp* reporter outside of the M lineage, but failed to show expression in the M lineage (Fig. 4I-J). Because the only difference between the translational *ceh-20::gfp* and the transcriptional *ceh-20p::gfp* constructs is the presence or absence of the introns, the above observations suggest that the introns of *ceh-20* may contain enhancer element(s) required for *ceh-20* expression in the M lineage.

The similar expression patterns of *unc-62* and *ceh-20* throughout the M lineage are consistent with their multiple and cooperative roles in the M lineage, as indicated by the genetic interactions described earlier.

***unc-62* and *ceh-20* display physical and regulatory interactions**

Because both *unc-62* and *ceh-20* share similar expression patterns in the M lineage, the synergistic interactions between them could be due to direct physical interactions between these two factors, or regulatory interactions, or both. To test these possibilities, we first used the yeast two-hybrid system and tested whether UNC-62 and CEH-20 can interact with each other. When fused with the GAL4 DNA binding domain, full length UNC-62 and CEH-20 each alone activated reporter expression (data not shown). We thus made truncations of each protein and found that the region containing the PBC domain of CEH-20 (aa1-196) interacts with the N-terminal 287aa (isoform 1a) or 250aa (isoform 1b) of UNC-62, which contains the HM domain (Fig. 5A). These results suggest that CEH-20 and UNC-62 can physically interact with each other.

Previous studies have shown that *unc-62* is required for the nuclear localization of CEH-20 (Van Auken et al., 2002; Potts et al., 2009). Consistent with these findings, we found that *unc-62(RNAi)* led to the reduction of the translational CEH-20::GFP signal (Fig. 5B-E), but not the transcriptional *ceh-20p::gfp* signal (Fig. 5F-I). To determine whether CEH-20 plays a role in regulating the expression of *unc-62*, we assayed the expression of the *unc-62p::gfp::let-858 3'UTR* reporter in the strong loss-of-function, and possibly null, alleles of *ceh-20*, *os39*. We found that 96 of 97 *ceh-20(os39)* mutant animals lacked *unc-62p::gfp* signal in the M mesoblast and the P lineages, but did express *unc-62p::gfp* in neuronal or hypodermal cells (Fig. 5J-L). Similar observations were made using the strong loss-of-function *ceh-20(n2513)* mutants (data not shown). Since the M mesoblast is still present in the *os39* mutant animals and the *unc-62p::gfp* reporter uses the well characterized *let-858 3'UTR* (Kelly et al., 1997), the above results suggest that *ceh-20* is required to positively regulate the promoter activity of *unc-62* in the M lineage and P lineage, but not in neuronal or hypodermal cells.

Discussion

Meis/Hth/*unc-62* functions in mesodermal development

MEIS family proteins have been previously shown to play important roles in multiple aspects of development in both vertebrates and invertebrates (for example, see Abu-Shaar and Mann, 1998; Henderson and Andrew, 2000; Choe et al., 2002; Choe and Sagerstrom, 2004; Mercader et al., 2005). Similarly, the *C. elegans* MEIS-related proteins *psa-3* and *unc-62* have been shown to function in regulating T cell asymmetric cell division (*psa-3*, Arata et al., 2006), regulate embryogenesis, Q neuroblast migration, vulval formation and VC motor neuron survival (*unc-62*, Van Auken et al., 2002; Yang et al., 2005; Potts et al., 2009). In this study, we have found that a loss-of-function mutation *psa-3(os8)* or *psa-3(RNAi)* did not affect the postembryonic mesodermal lineage, the M lineage. However, reducing *unc-62* activity led to pleiotropic M lineage phenotypes including defects in cell fate specification and differentiation, indicating that *unc-62* is a crucial regulator of mesodermal development in *C. elegans*.

In *Drosophila*, the MEIS protein Hth functions in the visceral mesoderm to regulate *dpp* expression (Stultz et al., 2006). Murine *meis-2* exhibits a dynamic expression pattern in differentiating somitic mesodermal cells (Ceconi et al., 1997). Similar temporal expression of *meis-2* has also been observed in the lateral mesoderm in zebrafish (Biemar et al., 2001; Zerucha and Prince, 2001). Thus MEIS family proteins may share a conserved role in mesodermal development.

UNC-62 functionally cooperates with CEH-20 to regulate mesoderm development

Previous work has shown that *ceh-20* is required for the specification and diversification of the M lineage, as well as the proper specification of CC and BWM fates (Liu and Fire, 2000). Our analysis of the phenotypes of a weak *ceh-20(ay9)* allele revealed another role of *ceh-20*

in the proper specification of uterine muscles (UMs) and vulval muscles (VMs). *ceh-20* and *unc-62* appear to function together in this process, as two partial loss-of-function *unc-62* alleles, *e644* and *ku234*, both caused fate transformations from UMs and type II vulval muscles (VM2s) to type I vulval muscles (VM1s). In particular, the *ku234* mutants exhibited 100% penetrance of the UM and VM2 to VM1 fate transformation without affecting the early M lineage (Table 1, Fig. 2). Because *ku234* is a point mutation located in the first intron of the 1a isoform, but mutating the translation initiation codon of the 1b isoform, either the *ku234* mutation disrupts a VM/UM precursor-specific enhancer element located in the first intron, or the 1b isoform of *unc-62* is specifically expressed in the VM/UM precursor cells. *unc-62* is also required for proper specification and differentiation of M-derived CCs and BWMs. In addition to the similar M lineage phenotypes shared by *ceh-20* and *unc-62* mutants, we observed synergistic genetic interactions between *ceh-20* and *unc-62*. As shown in Table 1, both *ceh-20(ay9); unc-62(e644)* and *ceh-20(ay9); unc-62(ku234)* double mutants exhibited more severe M lineage defects compared to each of the corresponding single mutants. For example, even though *unc-62(ku234)* single mutants did not exhibit any early M lineage defects, the *ceh-20(ay9); unc-62(ku234)* double mutant animals showed severe early M lineage defects in CC/BWM/SM fate specification.

The genetic interactions between *unc-62* and *ceh-20* could be due to 1) a physical interaction, 2) regulatory interactions, or 3) both physical and regulatory interactions between the two factors. We have found that *unc-62* and *ceh-20* share similar expression patterns within and outside of the M lineage, and that they can physically interact in the yeast two-hybrid system. We further showed that the two genes also share regulatory interactions: while *unc-62* is required for the nuclear localization of the CEH-20 protein, *ceh-20* is required for the expression of *unc-62* in the M lineage. Because the M cell is present in *ceh-20* mutants and the *unc-62p::gfp* reporter uses the well characterized *let-858* 3'UTR (Kelly et al., 1997), the lack of expression of the *unc-62p::gfp* reporter in the M and P lineages in *ceh-20* mutants suggests that *ceh-20* may positively regulate the transcription of *unc-62*. However, at this point, we do not know whether this regulation is direct or indirect. The regulation of the nuclear localization of PBC proteins by MEIS proteins have been previously reported in multiple systems (Pai et al., 1998; Abu-Shaar et al., 1999; Affolter et al., 1999; Berthelsen et al., 1999; Jaw et al., 2000; Rieckhof et al., 1997; Potts et al., 2009). However, transcriptional regulation of MEIS by PBC proteins has not been previously reported. In *Drosophila*, *hth* transcription appears not to be affected by the loss of *exd* (Abu-Shaar and Mann, 1998) even though Hth protein level is greatly reduced in embryos lacking both maternal and zygotic EXD (Kurant et al., 1998) or in *exd* mutant clones (Abu-Shaar and Mann, 1998). Thus our findings add one more level to the complex interactions between MEIS and PBC family of proteins: possible transcriptional regulation of *meis* genes by PBC proteins.

The MEIS and PBC proteins have been implicated as cofactors of Hox proteins that cooperatively bind to target DNA sequences with Hox proteins (Maconochie et al., 1997; Kroon et al., 1998; Swift et al., 1998; Ryoo et al., 1999; Shanmugam et al., 1999; Shen et al., 1999; Ferretti et al., 2000; Ebner et al., 2005). The early M lineage defects of *unc-62(e644)*, *unc-62(RNAi-soaking)* and *ceh-20(ay9)* mutants regarding CC/BWM/SM fate specification and differentiation are very similar to the previously described phenotypes of *mab-5(0)* mutants (Harfe et al., 1998). Furthermore, the *ceh-20(ay9); unc-62(e644)*, *ceh-20(ay9); unc-62(ku234)*, *mab-5(RNAi); unc-62(e644)* and *mab-5(RNAi) ceh-20(ay9)* double mutants exhibited more severe M lineage phenotypes that resemble the *lin-39(0) mab-5(0)* double mutant phenotypes. These observations suggest that UNC-62 may interact with CEH-20 and Hox to activate downstream targets required for the diversification and proper BWM/CC/SM fate specification of the M lineage. At this point, it is not clear whether the cooperative role of UNC-62 and CEH-20 in UM/VM fate specification is dependent on the Hox factors. *lin-39(0)* mutants have no defects in the M lineage, while *lin-39(0) mab-5(0)* mutants and *mab-5*

(0) mutants both have earlier defects in the M lineage, which could potentially mask the role of *mab-5* and *lin-39* in the late SM lineage.

It has been shown that LIN-39 or MAB-5 and CEH-20 can form heterodimers and bind to the corresponding Hox-PBC binding sites in the promoter region of *hll-8* to regulate its expression (Liu and Fire, 2000). Since *hll-8::gfp* expression is absent or barely detectable in *ceh-20(ay9); unc-62(e644)*, *ceh-20(ay9); unc-62(ku234)*, *mab-5(RNAi); unc-62(e644)* and *mab-5(RNAi) ceh-20(ay9)* mutants, it is possible that UNC-62 may participate in the DNA-bound Hox-CEH-20 complex in regulating *hll-8* expression. However, we did not find any potential MEIS binding site (TGACAG, Chang et al., 1997; Knoepfler et al., 1997; Shen et al., 1997; Berthelsen et al., 1998) adjacent to the Hox-CEH-20 consensus sequence in the characterized *hll-8* promoter region (Liu and Fire, 2000). Furthermore, attempts using chromatin immunoprecipitation (ChIP) on the *hll-8* promoter failed, possibly due to the short time window of early M lineage development and the small number of M-derived cells (1-18) relative to the whole animal (~900 somatic cells). It is possible that UNC-62 binds to some novel sequences in the *hll-8* promoter that are different from the canonical MEIS binding site, or UNC-62 acts as a non-DNA-binding partner in the Hox-CEH-20-UNC-62 trimeric complex. The physical interaction that we observed between CEH-20 and UNC-62 is consistent with the second hypothesis.

In addition to regulating *hll-8* expression, *unc-62* and *ceh-20* may have additional targets in the M lineage. We have previously found that the HMX factor MLS-2 is likely a direct target of CEH-20 in the M lineage (Jiang et al., 2008). We observed normal *mls-2* expression in the early M lineage of *unc-62(RNAi-soaking)* animals (data not shown). However, since *unc-62(RNAi-soaking)* may not completely knock out *unc-62* activity in the early M lineage, we cannot rule out the possibility that *mls-2* is regulated by *unc-62*.

In the SM lineage, a T-box transcription factor MLS-1 is both necessary and sufficient for UM fate specification (Kostas and Fire, 2002). *mls-1* is expressed in both UM and VM2 precursor cells and is regulated by *hll-8* (Kostas and Fire, 2002). While *unc-62(ku234)* mutants exhibited fate transformations from UMs and VM2s to VM1s (Fig. 2A-H), *mls-1* expression appeared normal (Fig. 2I-J). Thus, *unc-62* must be involved in regulating *mls-1* activity through post-transcriptional processes. For instance, *unc-62* could be required to activate MLS-1 cofactor(s) in the SM lineage, which then function(s) together with MLS-1 to promote UM fate.

In summary, UNC-62/MEIS/HTH, CEH-20/PBX/EXD and Hox factors (MAB-5 and LIN-39) play multiple roles in the *C. elegans* postembryonic mesoderm. Previous work has found roles of CEH-20 and Hox factors MAB-5 and LIN-39 in cell proliferation in the M lineage as well as in the specification and differentiation of M lineage-derived striated body-wall muscles (BWMs) and non-muscle coelomocytes (CCs) (Harfe et al., 1998; Liu and Fire, 2000; Jiang et al. (2008). We showed in this work that UNC-62 also participates in these functions. Furthermore, we demonstrated that UNC-62 and CEH-20 are also required for the proper specification of non-striated vulval and uterine muscles (VM and UMs). While all three factors are required for the M lineage expression of CeTwist, HLH-8 (Liu and Fire, 2000; this work), CEH-20 appears to function in a Hox-independent manner in regulating the HMX gene *mls-2* (Jiang et al., 2005; 2008). The mode in which these three factors function together or independently will be elucidated upon the identification and characterization of additional M lineage specific targets of each.

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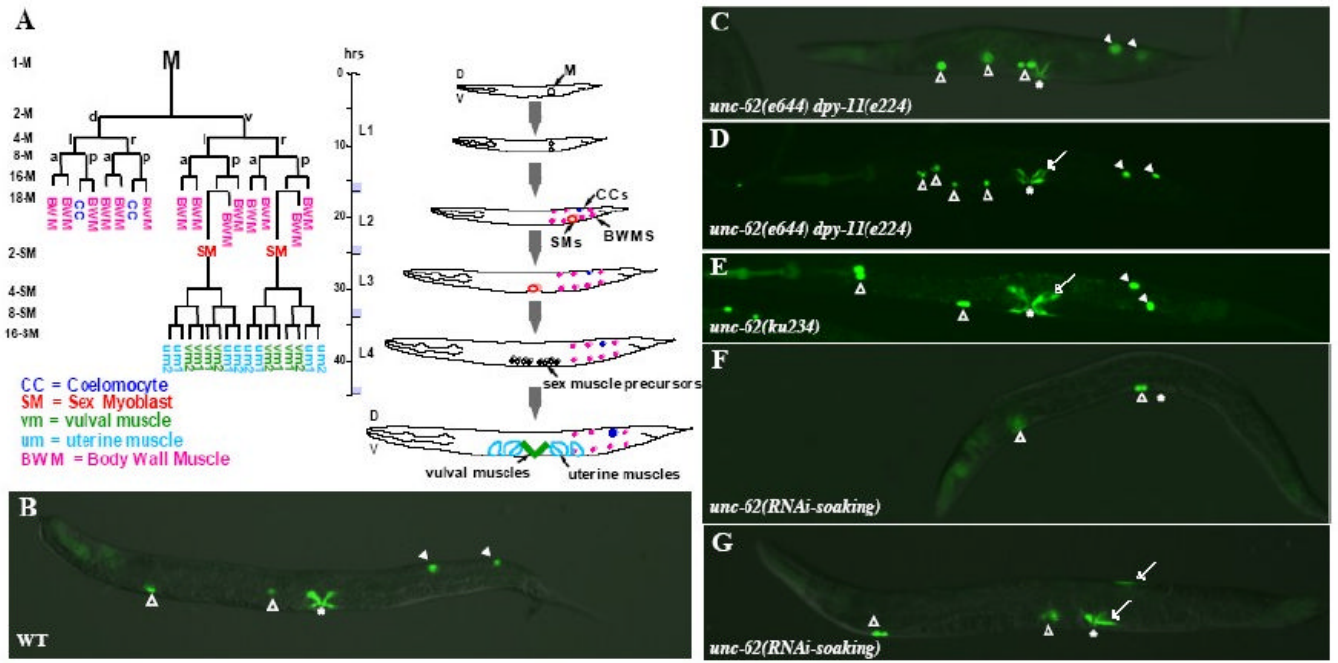


Figure 1. *unc-62* is required for proper development of the M lineage

(A) The *C. elegans* hermaphrodite postembryonic M lineage. Times are indicated post-hatching at 25°C. The M lineage showing all differentiated cell types is shown on the left (modified from Sulston and Horvitz, 1977), while a schematic lateral view of the M lineage through larval development is shown on the right. The different stages of the M lineage are also depicted as 1-M through 16-SM. D, dorsal; V, ventral; L, left; R, right; A, anterior; P, posterior. (B-H) Wild-type (B), *unc-62(e644) dpy-11(e224)* (C-D), *unc-62(ku234)*, and *unc-62(RNAi-soaking)* (F-G) adult animals visualized using an *intrinsic CC::gfp* and *egl-15::gfp*. Open arrowheads point to embryonic CCs, while solid arrowheads point to M-derived CCs. Arrows point to either extra type I vulval muscles (VM1s) in the vulval region (the vulva is denoted by an asterisk) or ectopic *egl-15::gfp*-positive cells in the posterior. (C) A *unc-62(e644) dpy-11(e224)* animal with normal numbers of CCs and VM1s. (D) A *unc-62(e644) dpy-11(e224)* animal with normal numbers of CCs but extra VM1s. (E) A *unc-62(ku234)* animal with normal numbers of CCs but extra VM1s. (F) A *unc-62(RNAi-soaking)* animal with no M-derived CCs and VM1s. (G) A *unc-62(RNAi-soaking)* animal with no M-derived CCs and extra VM1-like cells located both in the vulval region and on the dorsal side of the animal.

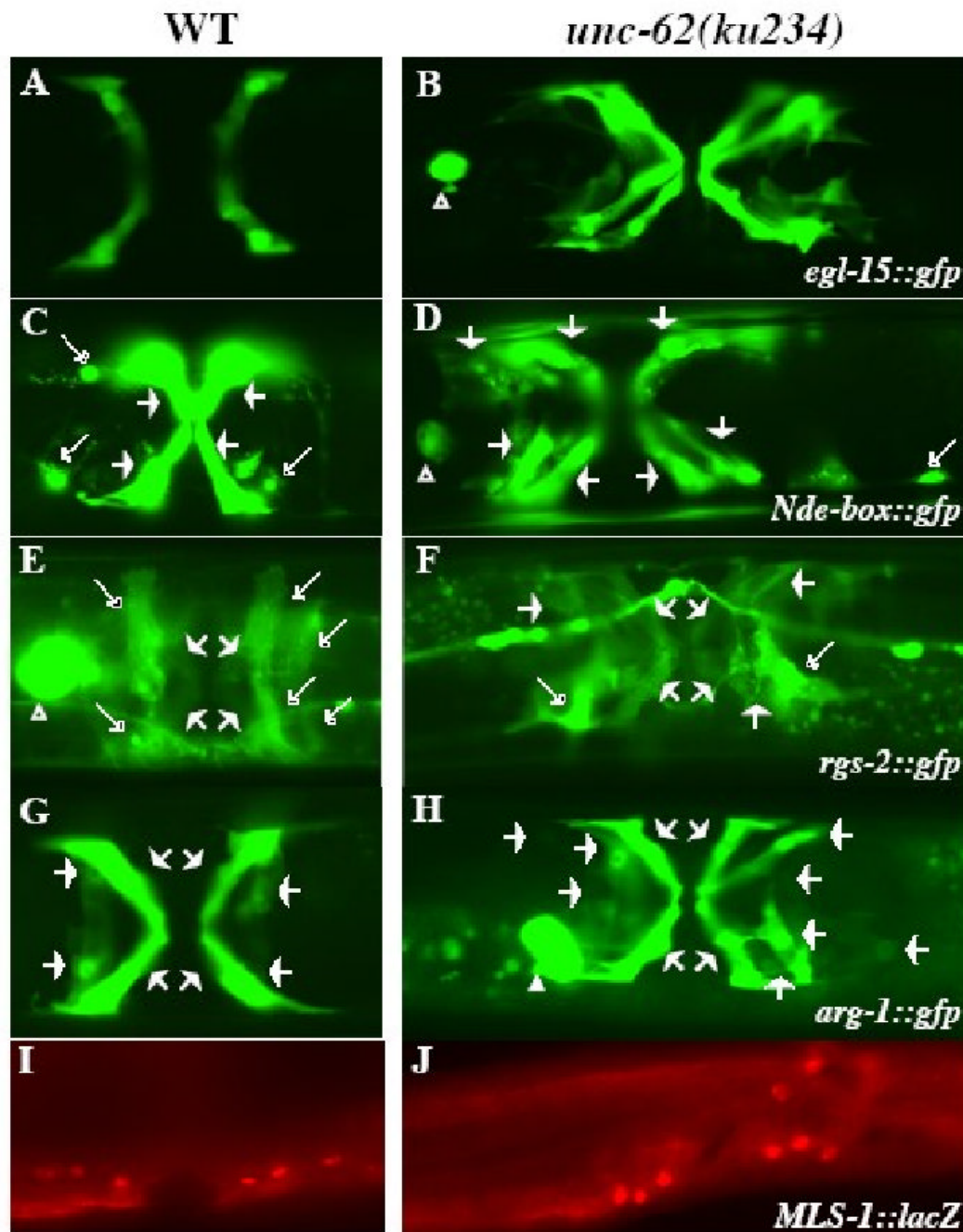


Figure 2. Uterine muscles (UMs) and type II vulval muscles (VM2s) are transformed to type I vulval muscles (VM1s) in *unc-62(ku234)* mutants

Ventral views of wild-type (A, C, E, G) and *unc-62(ku234)* (B, D, F, H) animals showing the differentiated sex muscles. Four VM1s in wild-type (A) and 12 VM1s in *unc-62(ku234)* (B) animals visualized by *egl-15::gfp*. UMs and VMs visualized by *Nde-box::gfp* (C, D) and *rgs-2::gfp* (E, F) in wild-type (C, E) and *unc-62(ku234)* (D, F) animals. The morphology of UMs (marked by longer open arrows) and VMs (marked by short solid arrows) are distinct. *unc-62(ku234)* animals (D, F) have reduced numbers of cells with the UM morphology and extra numbers of cells with the VM morphology. *arg-1::gfp* in wild-type (G) and *unc-62(ku234)* (H) animals showing both VM1s and VM2s. The *unc-62(ku234)* animal in panel H

has 12 VM1s. (I, J) *mls-1::LacZ* expression pattern in wild-type (I) and *unc-62(ku234)* (J) animals. Six out of the 12 (I) and 8 out of the 12 (J) *mls-1::lacZ*-expressing cells are shown in panels I and J, respectively. The other *mls-1::lacZ*-expressing cells are on a different focal plane. Both wild-type and *unc-62(ku234)* animals show similar *mls-1::lacZ* expression patterns.

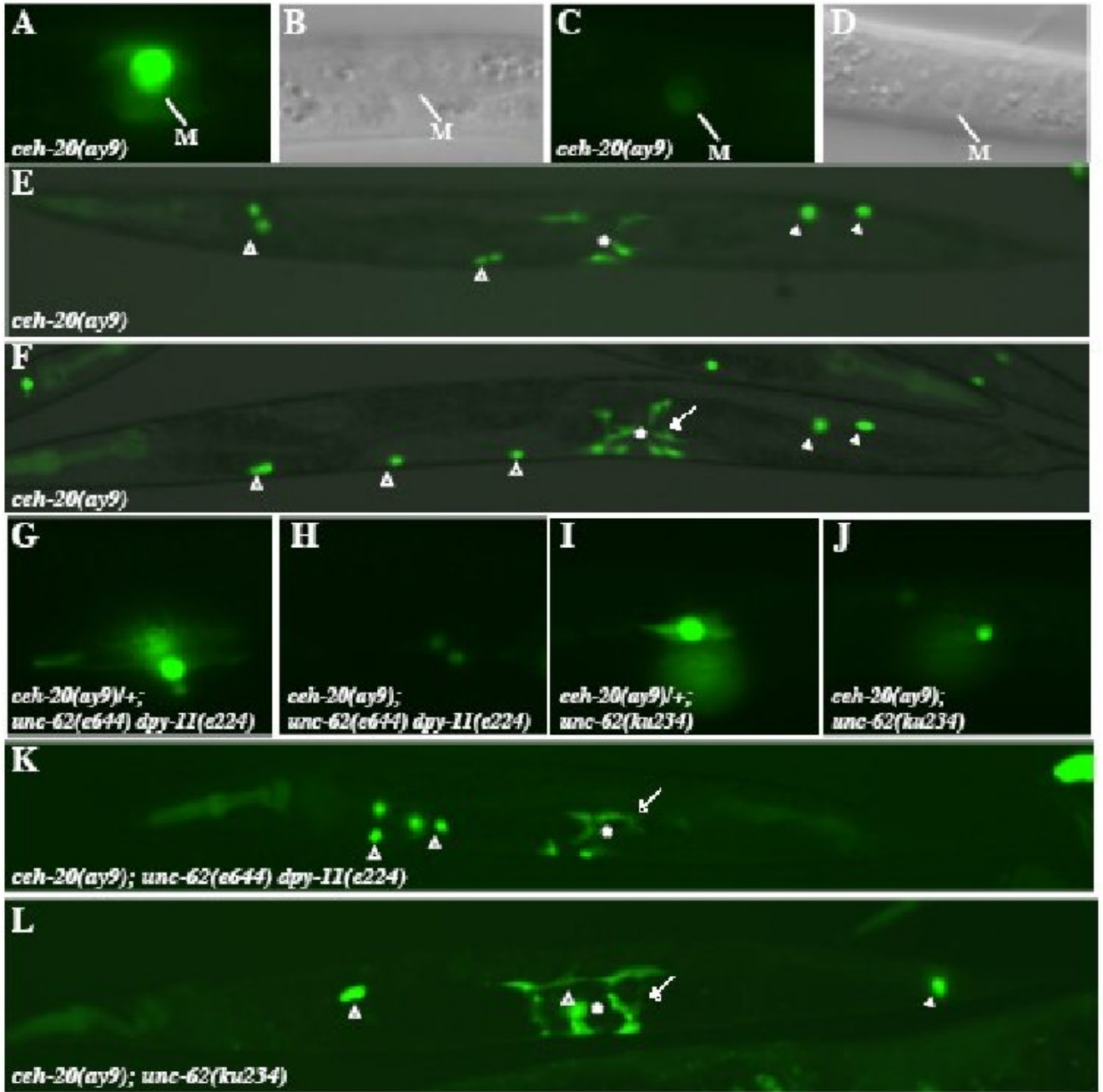


Figure 3. *ceh-20* genetically interacts with *unc-62* to regulate M lineage development
 (A-F) various M lineage phenotypes of *ceh-20(ay9)* animals. (A-D) *ceh-20(ay9)* animals at the 1-M stage with *hhl-8::gfp* (A, C) and the corresponding DIC images (B, D). 80% (n=200) of *ceh-20(ay9)* animals exhibit strong *hhl-8::gfp* expression (A), while 20% showing much reduced *hhl-8::gfp* expression (C). Panels A and C are under the same exposure. (E-F) Adult *ceh-20(ay9)* animals visualized using an *intrinsic CC::gfp* and *egl-15::gfp*. (E) An *ay9* animal with the normal number of M-derived CCs and VM1s that are not properly attached. (F) An *ay9* animal with the normal number of M-derived CCs but a total number of 10 VM1s. (G-J) *hhl-8::gfp* expression in the 2 SMs of *ceh-20(ay9)/+; unc-62(e644) dpy-11(e224)* (G), *ceh-20(ay9); unc-62(e644) dpy-11(e224)* (H), *ceh-20(ay9)/+; unc-62(ku234)* (I) and *ceh-20(ay9); unc-62(ku234)* (J). (K-L) *hhl-8::gfp* expression in the 2 SMs of *ceh-20(ay9); unc-62(e644) dpy-11(e224)* (K) and *ceh-20(ay9); unc-62(ku234) dpy-11(e224)* (L).

unc-62(ku234) (J) animals. Notice the reduced level of *hlh-8::gfp* expression in *ceh-20(ay9); unc-62(e644) dpy-11(e224)* (H) and *ceh-20(ay9); unc-62(ku234)* (J) animals compared to that in *ceh-20(ay9)/+; unc-62(e644) dpy-11(e224)* (G) and *ceh-20(ay9)/+; unc-62(ku234)* (I) animals. (K-L) Adult *ceh-20(ay9); unc-62(e644) dpy-11(e224)* (K) and *ceh-20(ay9); unc-62(ku234)* (L) animals visualized using an *intrinsic CC::gfp* and *egl-15::gfp*. (K) A *ceh-20(ay9); unc-62(e644) dpy-11(e224)* animal with no M-derived CCs and extra VM1s. (L) A *ceh-20(ay9); unc-62(ku234)* animal with one M-derived CCs and extra VM1s. Open arrowheads: embryonically-derived CCs. Solid arrowheads: M-derived CCs. Arrows: ectopic *egl-15::gfp*-expressing VM1-like cells. *: location of the vulva.

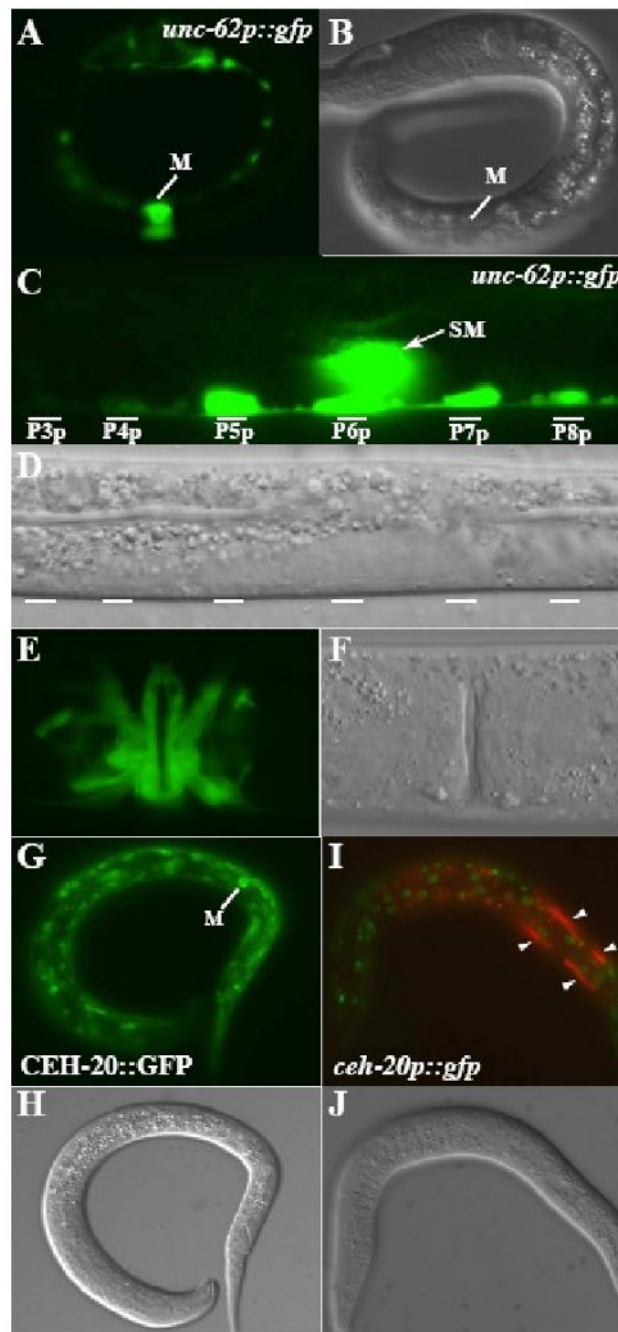


Figure 4. Both *unc-62* and *ceh-20* are expressed in the M lineage

(A-F) *unc-62p::gfp* (A, C, E) and corresponding DIC images (B, D, F) of 1-M stage L1 (A-B), 2-SM stage L2 (C-D) and adult stage animals. *unc-62p::gfp* is expressed throughout the M lineage, including mature vulval muscles (E) and in the P lineages (C). (G-J) GFP (G, I) and corresponding DIC images showing the expression of a functional translational CEH-20::GFP in the M mesoblast (G-H) and the lack of M lineage expression of the transcriptional *ceh-20p::gfp* in the M lineage (I-J). Green in I: *ceh-20p::gfp*. Red in I: *hllh-8p::rfp*. Notice that *ceh-20p::gfp* is expressed in the other cell types that express CEH-20::GFP, but not in cells of the M lineage (marked by arrowheads).

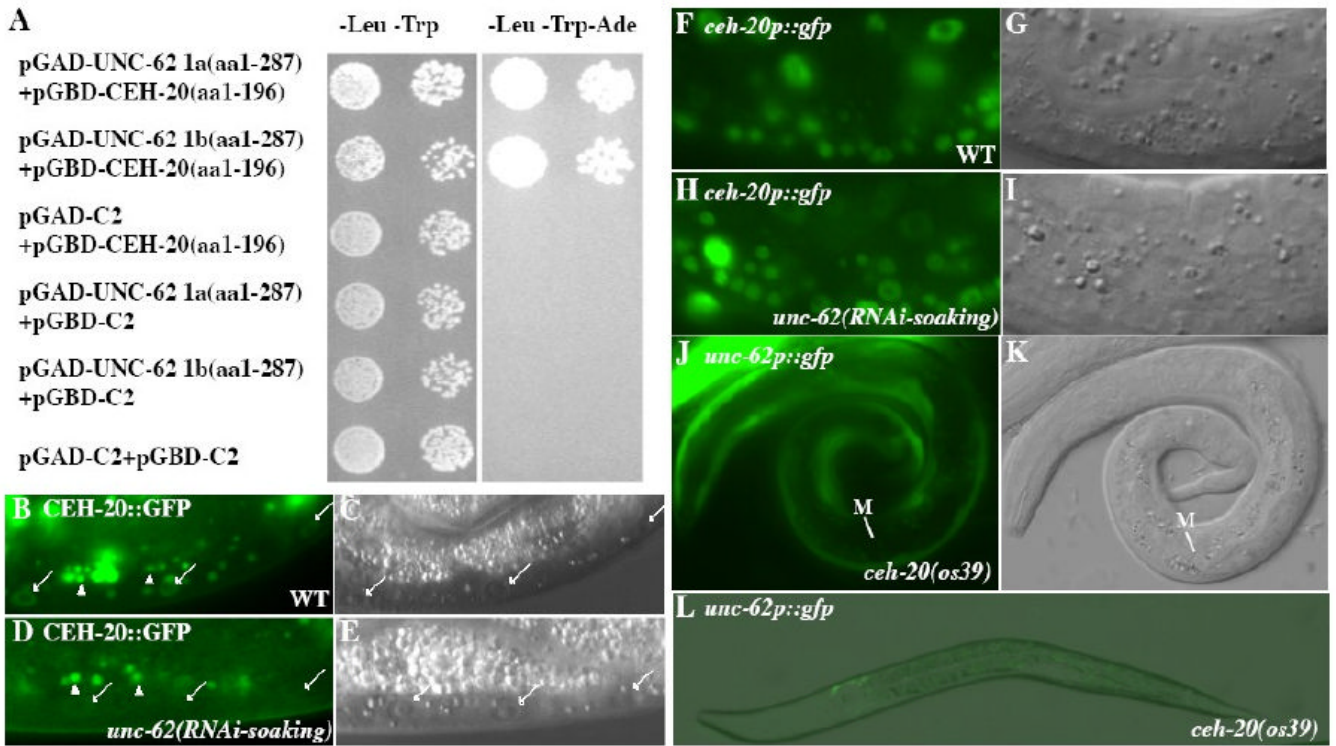


Figure 5. UNC-62 and CEH-20 exhibit physical and regulatory interactions

(A) The N-termini of both the 1a and 1b splicing isoforms of *unc-62* that contain the HM domain can bind to the PBC domain (aa 1-196) of CEH-20 in the yeast two-hybrid assay. Interaction was detected in one direction using the PBC domain (aa 1-196) of CEH-20 fused to the GAL4 DNA binding domain (GBD). The GBD-UNC-62 fusions autoactivate and were not included in the tests. (B-E) CEH-20::GFP (B, D) and the corresponding DIC images (C, E) in wild-type (B-C) and *unc-62(RNAi-soaking)* (D-E) animals. CEH-20::GFP is present in the nucleus, but not the nucleolus, of multiple cell types in wild-type animals (B-C). Notice the significant reduction or loss of nuclear CEH-20::GFP in *unc-62(RNAi-soaking)* animals (arrows in panel D) compared to wild-type animals (arrows in panel B). Additional bright gut autofluorescence signals in panels B and D are marked by arrowheads. (F-I) *ceh-20p::gfp* (F, H) and the corresponding DIC images (G, I) in wild-type (F-G) and *unc-62(RNAi-soaking)* (H-I) animals. The expression of the transcriptional *ceh-20p::nls::gfp* is not affected by *unc-62(RNAi-soaking)*. (J-L) *unc-62p::gfp* in *ceh-20(os39)* animals. Expression of *unc-62p::gfp* is lost in the M cell in an L1 animal (J-K) and most of the cells in an L2 animal (L).

Table 1
***unc-62* genetically interacts with *ceh-20* and *mab-5* to regulate M lineage development**

Genotype ¹	<i>hlh-8::gfp</i> expression	Early M lineage phenotypes ²	SM lineage phenotypes ³
Wild-type	bright (100%) n=200	2 CCs, 2 SMs, 14 BWMs (100%) n=200	4 VM1s (100%) n=200
<i>ceh-20(ay9)</i>	bright (80%) faint ⁴ (20%) n=200	2 CCs, 2 SMs, 14 BWMs (91%) 1-3 CCs, 1 SM & 13-14 BWMs (9%) n=200 ⁵	4 VM1s (44%) >4 VM1s (56%) n=75
<i>unc-62(e644) dpy-11(e224)</i>	bright (100%) n=200	2 CCs, 2 SMs, 14 BWMs (78%) 0-1 CCs, 3-4 SMs, 12-14 BWMs (22%) n=200 ⁶	4 VM1s (44%) >4 VM1s (56%) n=69
<i>unc-62(ku234)</i>	bright (100%) n=200	2 CCs, 2 SMs, 14 BWMs (100%) n=200 ⁷	>4 VM1s (100%) n=200
<i>mab-5(RNAi)</i>	no early expression bright SM lineage expression (100%) n=156	2 CCs, 2 SMs, 14 BWMs (6%) 0 CCs, 3-5 SMs, 13-14 BWMs (94%) n=156	4 VM1s (6%) >4 VM1s (94%)
<i>ceh-20(ay9)/+; unc-62(e644) dpy-11(e224)</i>	bright (100%) n=123	2 CCs, 2 SMs, 14 BWMs (55%) 0-1 CCs, 3-4 SMs (45%) n=60	4 VM1s (7%) >4 VM1s (93%) n=123
<i>ceh-20(ay9); unc-62(e644) dpy-11(e224)</i>	faint (3%) no expression (97%) N=200	0-1 CCs ⁸ (100%) n=83	>4 VM1s (100%) n=45
<i>ceh-20(ay9)/+; unc-62(ku234)/+</i>	bright (100%) n=23	2 CCs, 2 SMs, 14 BWMs (100%) n=23	4 VM1s (100%) n=23
<i>ceh-20(ay9)/+; unc-62(ku234)</i>	bright (100%) n=163	2 CCs, 2 SMs, 14 BWMs (72%) 0-1 CCs, 2 SMs (28%) n=163	>4 VM1s (100%) n=163
<i>ceh-20(ay9); unc-62(ku234)</i>	faint (7%) no expression (93%) N=200	2 CCs, 2 SMs, 14 BWMs (13%) 0-1 CCs, 9-14 BWMs ⁹ (87%) n=200	>4 VM1s (100%) n=200
<i>mab-5(RNAi); unc-62(e644) dpy-11(e224)</i>	no expression (41%) bright in SM lineage ¹⁰ (59%) n=200	2 CCs (1%) 0-1 CCs ⁸ (99%) n=200	0 VM1s ¹¹ (58%) 4 VM1s (1%) >4 VM1s (41%) n=200
<i>mab-5(RNAi) ceh-20(ay9)</i>	no expression (52%) bright in SM lineage ¹⁰ (48%) n=200	2 CCs (9%) 0-1 CCs ⁸ (91%) n=200	0 VM1s ¹¹ (60%) 4 VM1s (9%) >4 VM1s (31%) n=200

¹ All animals examined carried the following GFP markers for the M lineage: *intrinsic CC::gfp(ccIs4438)* III, *ayIs2(egl-15::gfp)* IV; *hlh-8::gfp(ayIs6)* X.

² Phenotypes focus on specification of the 18 M lineage descendants (14 BWMs, 2 CCs and 2 SMs) produced at the L1 stage.

³ The SM lineage phenotypes shown here focused on the number of *egl-15::gfp* expressing VM1-like cells.

⁴ *hlh-8::gfp* signal was faint in both the early M lineage and the SM lineage.

⁵ The number of BWMs was obtained via examining 6 animals.

⁶ The number of BWMs was obtained via examining 5 animals.

⁷ The number of BWMs was obtained via examining 3 animals.

⁸ The number of BWMs and SMs was not determined in this mutant strain.

⁹ The number of BWMs was obtained via examining 5 animals.

¹⁰ There was no *hlh-8::gfp* expression in the early M lineage.

¹¹ Some of the animals had 1-3 elongated *egl-15::gfp* positive cells randomly located in the posterior of the worm.