Migrations of the *Caenorhabditis elegans* HSNs are regulated by *egl-43*, a gene encoding two zinc finger proteins

Gian Garriga,1-3 Catherine Guenther,1 and H. Robert Horvitz2

1Department of Molecular and Cell Biology, University of California, Berkeley, California 94720 USA; 2Howard Hughes Medical Institute, Department of Biology, Massachusetts Institute of Technology, Cambridge, Massachusetts 02139 USA

During embryonic development, the two *Caenorhabditis elegans* HSN motor neurons migrate from their birthplace in the tail to positions near the middle of the embryo. Here, we demonstrate that of all cells that undergo long-range migrations, only the HSNs are affected in animals that lack function of the *egl-43* gene. We also show that *egl-43* function is required for normal development of phasmid neurons, which are sensory neurons located in the tail. The *egl-43* gene encodes two proteins containing zinc finger motifs that are similar to the zinc fingers of the murine Evi-1 proto-oncoprotein. Our genetic and molecular results suggest that *egl-43* encodes two transcription factors and acts to control HSN migration and phasmid neuron development, presumably by regulating other genes that function directly in these processes.

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During metazoan development, many cells migrate before they differentiate. Although cell migration is important for many aspects of development, the extent of the cell migration that occurs during the development of nervous systems is particularly striking. For example, populations of migrating cells generate much of the vertebrate peripheral nervous system (LeDouarin 1982). Neuronal precursors that migrate from the neural crest ventrally along different routes to various positions produce sensory, autonomic, and enteric neurons as well as glia, pigment cells, mesenchymal cells, and chromaffin cells of the adrenal medulla. Extensive cell migration also occurs during the development of the vertebrate central nervous system. Neuroblasts migrate radially from the ventricular zone, where they are born, past older neurons to more superficial layers of the cortex (Rakic 1974). Moreover, after reaching the appropriate cortical layer, many of these cells undergo extensive secondary migrations. For example, clonally related cells of the chick optic tectum migrate along three separate pathways, adopting distinct fates (Gray and Sanes 1991).

Although many studies have established that cell migrations play a major role during metazoan development, the mechanisms that regulate these migrations remain largely undefined. To identify the components involved in cell migration, investigators have taken two general approaches. Biochemical methods have been used to identify and characterize molecules that mediate cell migration and have been particularly successful in defining cell surface and extracellular matrix molecules (for review, see Hatten 1990; Reichardt and Tomaselli 1991; Hynes and Lander 1992). An alternative strategy is to use genetic methods, which, in principle, should identify not only genes that encode cell-surface and extracellular matrix molecules but also genes that regulate and respond to these molecules to control the timing, direction, and extent of cell migration. Genetic studies of cell migration are particularly straightforward using the nematode *Caenorhabditis elegans*, because individual migrating cells can be observed directly in living, intact animals (Sulston and Horvitz 1977; Kimble and Hirsh 1979; Sulston et al. 1983). Many genes required for various *C. elegans* cell migrations have been identified (Chalfie et al. 1983; Kenyon 1986; Hedgecock et al. 1987, 1990; Desai et al. 1988; Manser and Wood 1990; Stern and Horvitz 1992; Clark et al. 1992, 1993; Wang et al. 1993).

With the goal of understanding in detail how a specific cell migrates, we are studying genes involved in the embryonic migrations of the hermaphrodite-specific neurons (*HSNs*), a pair of serotonergic motor neurons that control egg laying by the hermaphrodite (Desai et al. 1988). To date, 19 genes involved in these migrations have been identified (Desai et al. 1988; G. Garriga, L. Bloom, and H.R. Horvitz, unpubl.). Here we describe the genetic and molecular characterization of *egl-43*, a gene that plays a key role in HSN migration. Mutations in *egl-43* result in the most severe HSN migration defect of all HSN migration mutants. Genetic results demonstrate that *egl-43* is required for HSN migration but not for other cell migrations, indicating that this gene does
not function globally in cell migration. Molecular analysis demonstrates that egl-43 encodes two proteins containing zinc finger motifs that are related to the zinc fingers of the murine Evi-1 proto-oncoprotein (Morishita et al. 1988). Our working model is that the egl-43 gene encodes two transcription factors that control HSN migration, presumably by regulating other genes that function directly in cell migration.

Results

The egl-43 gene is required for HSN migration

The C. elegans HSNs are a pair of serotonergic motor neurons that innervate the vulval muscles and stimulate egg laying by hermaphrodites (Desai et al. 1988). The two bilaterally symmetric HSNs are generated during embryogenesis and migrate anteriorly to the middle of the embryo from their birthplace in the tail (Fig. 1) (Sulston et al. 1983). Differentiation occurs later: The HSNs extend axons during larval development and synthesize serotonin in young adult hermaphrodites (Desai et al. 1988; Garriga et al. 1993).

Of the HSN migration mutants, egl-43 hermaphrodites exhibit the most severe defects in migration; the egl-43 HSNs migrate, on average, only 5–10% of the distance normally traversed by wild-type HSNs (Desai et al. 1988). Figure 2 compares the positions of the HSNs in wild-type and egl-43(n1079) adult hermaphrodites. The egl-43 hermaphrodite shown in Figure 2b exhibits the most extreme migration defect: The HSNs have failed to migrate from their birthplace in the tail.

Although migration is the only HSN trait that is severely disrupted in the two existing egl-43 mutants, axonal guidance and serotonin expression are also affected. As in the wild type, in egl-43 animals, each displaced HSN cell body extends a single axon to the ventral nerve cord (Garriga et al. 1993). Once in the ventral nerve cord, each axon turns anteriorly and extends to the head where it enters the nerve ring. Although these aspects of HSN axonal outgrowth are relatively normal in egl-43 mutants, the HSN axons fasciculate abnormally within the ventral nerve cord and fail to branch at the vulva (Garriga et al. 1993). In wild-type hermaphrodites, most of the synapses with the egg-laying muscles are made on this branch (White et al. 1986). Because all HSN migration mutants exhibit similar fasciculation and branching defects, the axonal guidance abnormalities of the egl-43 mutants presumably are secondary consequences of the posterior displacement of the HSN cell bodies (Garriga et al. 1993).

egl-43 mutants also exhibit a weak neurotransmitter—expression defect: Approximately 10% of the HSNs fail to stain with anti-serotonin antisera; other serotonergic neurons stain normally (Desai et al. 1988). Like the axonal—outgrowth defect, the serotonin—expression defect also might be a secondary effect of the misplacement of the HSN cell bodies. Alternatively, egl-43 might play a direct role in the expression of HSN serotonin. Consistent with the former possibility, other mutations that cause severe HSN displacements also result in low penetrance defects in HSN—serotonin expression (G. Garriga, unpubl.). Thus, although egl-43 mutants are abnormal in HSN migration, axonal outgrowth, and serotonin expression, migration might be the only HSN trait that is affected directly by the absence of the egl-43 gene product.

The egl-43 gene is required for phasmid neuron development

Mutations in the egl-43 gene also affect the development of the PHA and PHB phasmid neurons, which are sensory neurons that are located in the tail. When wild-type animals are soaked in certain fluorescent dyes, such as the carbocyanine dye Dil [1,1-dioctadecyl-3,3',3'-tetramethylindocarbocyanine perchlorate, Molecular Probes, Inc.], PHA and PHB take up the dye through their exposed sensilla, and both their cell bodies and processes can be visualized by fluorescence microscopy (Fig. 2d) (Hedgecock et al. 1985; E. Hedgecock, pers. comm.). In contrast, egl-43 mutants, phasmid neurons often could not be visualized with these dyes (Fig. 2e). Although most egl-43 phasmid neurons failed to fill with Dil, those that took up the dye appeared normal: The cell bodies were in their usual positions, and axonal and den-
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Figure 2. egl-43 mutants are defective in HSN cell migration and in phasmid neuron staining. (a–e) Fluorescence photomicrographs and tracings of wild-type and egl-43 adult hermaphrodites either stained with an anti-serotonin antiserum [a,b,c] or soaked in the fluorescent dye Dil [d,e]. (a) Ventral view of a wild-type (N2) hermaphrodite stained with an antiserum to serotonin. The bright ovals, which are located near the middle of the animal, are the HSN cell bodies. (b) Ventral view of an egl-43(n1079) hermaphrodite stained with an antiserum to serotonin. The bright ovals, which are located in the tail, are the displaced HSN cell bodies. This animal illustrates the most extreme type of HSN migration defect, in which the HSNs have failed to migrate from the positions in the tail where they were born. (c) Ventral view of an egl-43(n1079) hermaphrodite carrying the extrachromosomal array gmEx2, after staining with an antiserum to serotonin. The bright ovals are the HSN cell bodies, which are located in their normal positions near the middle of the animal. The HSN migration defect of this egl-43 mutant was rescued by the extrachromosomal array gmEx2, which contains a portion of the egl-43 gene [p43BA-RR] that rescues the egl-43 migration defect. (d) Left oblique view of a wild-type hermaphrodite that was soaked in the fluorescent dye Dil. The bright ovals are the cell bodies of the two phasmid neurons. The phasmid neuron dendrites and axons are labeled in the tracing at right. The fluorescence anterior to the phasmid neuron cell bodies is caused by Dil that has been ingested by the worm and retained in the intestinal lumen. (e) Left oblique view of an egl-43(n997) hermaphrodite that was soaked in the fluorescent dye Dil. The single phasmid neuron that took up the dye exhibited normal dendritic and axonal morphology. The portion of the dendrite connected to the cell body is out of the plane of focus. Note that none of the phasmid nerves in the second animal in the field of view took up the dye. In wild-type animals, 99% (n = 56) of phasmid neurons took up Dil (n = number of animals), whereas in egl-43(n997) and egl-43(n1079) mutants, 33% (n = 47) and 30% (n = 40) of phasmid neurons filled with Dil, respectively. In unc-4(e120) egl-43(n997)/mnDf28 animals, 31% (n = 48) of the phasmid neurons filled with Dil.

dritic processes appeared wild type (Fig. 2e). Electron micrographs of cross sections though the tails of egl-43(n997) animals showed that the sensory ciliary structures of PHA and PHB, which are normally exposed to the environment, were variably missing [E. Hartwieg, G. Garriga and H.R. Horvitz, unpubl.]. This observation provides a structural basis for the dye-filling defect of the egl-43 mutants.

In contrast to PHA and PHB, 12 sensory neurons located in the head that also take up Dil [Hedgecock et al. 1985; E. Hedgecock, pers. comm.] appeared normal in egl-43 mutants [data not shown]. It is noteworthy that these head sensory neurons are born at the anterior tip of the embryo and migrate posteriorly during embryonic development [Sulston et al. 1983]. The wild-type positions of these sensory neurons in egl-43 mutants indicate that they have migrated normally.

The dye-filling deficit of egl-43 mutants might reflect...
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a general defect in phasmid neuron development, for example, in the ability of the phasmid neurons to assume their proper identity. Alternatively, this deficit might reflect a more specific defect in dendrite morphogenesis, for example, in the ability of the phasmid neuron dendrites to extend or connect to the phasmid opening. Besides the HSN and phasmid neuron defects, egl-43 males exhibit a reduced efficiency of mating (Desai and Horvitz 1989). We have not determined the cellular basis for this mating defect.

The egl-43 gene does not function globally in cell migration

Although HSN migration is severely defective in egl-43 mutants, all other cell migrations that occur during embryonic and larval development appear normal (see Materials and methods). Moreover, other than the dendritic defects described above for PHA and PHB, we have not observed any abnormalities in process morphology for 16 classes of neurons that were analyzed (see Materials and methods).

To determine whether the specific effects of the two existing egl-43 mutations on HSN but not other cell migrations could be an artifact of having only weak egl-43 alleles, we scored the final positions of migrating cells in animals that lack the egl-43 gene. There are several deficiencies that fail to complement egl-43 as well as flanking genes (Sigurdson et al. 1984; Desai and Horvitz 1989; Miller et al. 1992). Animals homozygous for several of these deficiencies (mnDf14, mnDf22, mnDf24, mnDf25, mnDf26, mnDf28, and mnDf56) progress through embryogenesis, hatch, and then arrest development as larvae. Because the homozygous deficiency animals hatch, Nomarski optics can be used to score the positions of the HSNs as well as the positions of other cells that migrate embryonically. Hermaphrodites homozygous for these deficiencies had HSN migration defects similar to those seen in the egl-43 mutants. We analyzed in detail the positions of migrating cells in animals that are homozygous for two of these deficiencies, mnDf24 and mnDf28.

We scored the positions of neurons (ALM, CAN, and HSN) and mesodermal cells (coelomocyte mother cells, intestinal muscle cells, M, Z1, and Z4) that migrate embryonically in larvae homozygous for the deficiencies mnDf24 and mnDf28 (see Materials and methods). As in egl-43 mutants, only the positions of the HSNs were abnormal in these homozygous deficiency animals. Moreover, the Q neuroblasts and their descendants, which migrate during the first larval (L1) stage, also migrated normally in animals homozygous for mnDf24 or mnDf28. Unfortunately, these animals did not progress past the L1 stage, so we were unable to determine whether the SM myoblasts and the distal tip cells of the somatic gonad, which migrate later during larval development, migrated normally. Similarly, we could not determine whether later aspects of HSN development were normal in these homozygous deficiency animals because the HSNs do not begin to differentiate until the second [L2] or third [L3] larval stages (Garriga et al. 1993). Because cells other than the HSNs migrated normally in animals lacking egl-43, the egl-43 gene product is not necessary globally for cell migration. Desai and Horvitz (1989) described a weak Egl phenotype of mnDf28/+ animals. Using anti-serotonin staining to visualize the HSNs, we showed that mnDf28/+ animals display a weak but significant defect in HSN migration (data not shown). This effect on HSN migration, which is not seen in egl-43/+, may be a consequence of reducing the dosage of egl-43 by 50%. Alternatively, reducing the dosage of multiple genes may lead to this defect.

The egl-43 gene encodes two transcripts

To understand better the role of egl-43 in HSN and phasmid neuron development, we have begun a molecular analysis of this gene. The egl-43 gene maps ~0.08 map units to the right of the unc-4 gene and is removed by the deficiency mnDf28 but not by the deficiency mnDf60 (Fig. 3a) (Desai and Horvitz 1989; also, this study). The right endpoints of mnDf28 and mnDf60 are located within genomic DNA contained in the two cosmid Cosm W02B7 and R53 (Fig. 3b) (Miller et al. 1992; C. Shamus and M. Shen, pers. comm.), thus restricting the physical location of egl-43 to this region. To determine the precise location of egl-43, we used germ-line transformation to rescue the HSN migration, phasmid neuron, and egg-laying defects of the egl-43(n1079) mutant (Fire 1986; Mello et al. 1991). Each of the three cosmids, W02B7, R53, and C40F9, rescued the migration defect, placing the egl-43 rescuing activity within the 15 kb of genomic DNA shared by these cosmids (Fig. 3b). Furthermore, a 5.6-kb BamHI–ApaI fragment [p43BA], which is located between the right endpoints of mnDf28 and mnDf60, contained egl-43-rescuing activity (Fig. 3b,c).

Northern hybridization experiments using the 5.6-kb BamHI–ApaI fragment as a probe demonstrated that this region encodes poly[A]+ RNAs of 2.5, 1.8 , and 1.0 kb [Fig. 4]. Additional Northern hybridization experiments with T7 transcripts and DNA subclones of the 5.6-kb BamHI–ApaI fragment as probes showed that the 2.5- and 1.8-kb RNAs overlap and are transcribed from left to right and that the 1.0-kb RNA is transcribed from right to left (see Materials and methods). Additional germ-line transformation experiments suggested that the 2.5- and 1.8-kb transcripts correspond to the egl-43 gene: Removing sequences from the left end of the 5.6-kb BamHI–ApaI genomic DNA fragment, including sequences that encode the 1.8- and 2.5-kb transcripts, abolished rescue [Fig. 3c].

To define the structures of the 1.8- and 2.5-kb RNAs, we determined the sequences of genomic DNA from the region and then used oligonucleotides derived from the genomic sequence as primers to amplify cDNAs by the polymerase chain reaction (PCR) (see Materials and methods). These experiments demonstrated that the 2.5- and 1.8-kb transcripts encode 10 and 5 exons, respectively. The five exons of the 1.8-kb transcript are iden-
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Figure 3. Genetic and physical maps of the egl-43 region of chromosome II. (a) A genetic map of the egl-43 region. The genetically determined extents of the chromosomal deficiencies mnDf28 and mnDf60 are shown below the map. The position of the right endpoint of mnDf60 was previously placed within the egl-43 gene on the basis of its partial complementation of egl-43(n997) [Miller et al. 1992]. In our experiments, however, mnDf60 complemented both alleles of egl-43 [see Materials and methods]. (b) Physical mapping of the egl-43 gene. The top line represents a restriction map of the genomic DNA contained within the egl-43-rescuing cosmid R53 (data not shown). The three cosmid clones W02B7, R53, and C40F9 rescued (+) the HSN migration defects of egl-43 mutants when maintained as an extrachromosomal array after germ-line transformation. The fraction of independently derived transformed lines rescued for the HSN migration defect is presented in parentheses for each cosmid clone. The approximate positions of the right endpoints of mnDf28 and mnDf60 are indicated by arrows. (c) An expanded restriction map of the egl-43 region, showing the structure of the 2.5-, 1.8-, and 1.0-kb RNAs and subclones that either rescued (+) or failed to rescue (−) the HSN migration defects of egl-43 mutants when maintained as an extrachromosomal array after germ-line transformation. The fraction of independently derived transformed lines rescued for the HSN migration defect is presented in parentheses for each cosmid clone. (SL1) Trans-spliced SL1 leader sequences; [ATG and TAG] the predicted start and stop sites of translation, respectively; [AAA] the site of poly(A) addition. Solid boxes represent coding sequences; open boxes represent untranslated sequences of the 2.5- and 1.8-kb transcripts. Northern hybridization experiments demonstrated that the 1.0-kb RNA is transcribed from right to left, and its 3' exon(s) is contained within the 0.9-kb EcoRI fragment. Southern hybridization experiments and partial sequencing of a cDNA of the 1.0-kb transcript suggest that the 3' exon is spliced to sequences to the right of the Apal site; the precise structure of the 1.0-kb transcript has not been determined. The structures of the plasmids used in the rescue experiments are described in Materials and methods. p43BA-FS is a derivative of p43BA that contains a frameshift mutation in the third exon of the 1.8-kb transcript. Southern analysis of DNA prepared from five roller lines that had been injected with p43BA-FS showed that all five lines contained injected egl-43 sequences (data not shown). p43BA-RR is a derivative of p43BA-FS that has a restored reading frame. Abbreviations: [A] Apal; [B] BamHI; [E] EcoRI; [N] NotI; [S] SalI; [S] SphI.
The asterisk (*) indicates the position of the TAG amber stop codon. The positions of introns are indicated by arrowheads. A good candidate for the poly(A) + addition signal is underlined with a thick line (Proudfoot and Brownlee 1976). The sequences of the two transcripts are generated by splicing of SL1 to the 5' end of the exon containing this methionine (the position of the 5' of the exon is indicated by the arrowhead immediately upstream). The two highly acidic regions are underlined. Each of the zinc fingers is boxed. The first ATG [amino acid position 1] of the 2.5-kb transcript is shown as the first methionine of the larger protein, although the second methionine at amino acid 68 might represent the initiator methionine. The arrow indicates the proposed initiator methionine of the 1.8-kb transcript; this methionine is located within the third zinc finger encoded by the 2.5-kb transcript. The 5' end of the 1.8-kb transcript is apparently generated by splicing of SL1 to the 5' end of the exon containing this methionine [the position of the 5' of the exon is indicated by the arrowhead immediately upstream]. The two highly acidic regions are underlined. Each of the zinc fingers is boxed. The asterisk (*) indicates the position of the TAG amber stop codon. The positions of introns are indicated by arrowheads. A good candidate for the poly(A)+ addition signal is underlined with a thick line (Proudfoot and Brownlee 1976). The sequences of the two egl-43 cDNAs were determined by sequencing genomic DNA (not shown) and confirming the structure of the transcripts by PCR of cDNAs, RACE, and primer extension (see Materials and methods).
2.5- and 1.8-kb transcripts could encode proteins of 581 and 360 amino acids, respectively (Fig. 5). The larger Egl-43 protein contains six zinc finger motifs of the C2H2 type (such zinc fingers were first discovered in the Xenopus transcription factor IIIA; Miller et al. 1985) and two acidic regions that could function as transcriptional activators (Ma and Ptashne 1987). The smaller protein contains the 360 carboxy-terminal amino acids of the larger protein, including three of the six zinc finger motifs and one of the acidic regions. The first methionine of

Figure 5. (See facing page for legend.)
the 1.8-kb transcript begins within the third zinc finger motif of the larger protein.

The six zinc finger motifs encoded by egl-43 are similar to 6 of the 10 zinc finger motifs encoded by Evi-1, a gene that was originally identified as the site of retroviral insertions that lead to myeloid leukemia in mice (Morishita et al. 1988). The Egl-43 zinc-finger motifs 1–6 correspond to the zinc finger motifs 5–10 of the murine Evi-1 protein (Fig. 6). Both the order and sequences of the Egl-43 and Evi-1 zinc finger motifs are conserved, and as in the murine gene, the zinc finger motifs of Egl-43 are clustered into two domains. The Egl-43 and Evi-1 zinc finger motifs of the second domain are more highly conserved (67 of the 85 amino acids are identical) than those of the first (45 of the 85 amino acids are identical) [Fig. 6b]. The third Egl-43 zinc finger, which is the least conserved of the zinc fingers, also contains the presumptive initiator methionine of the smaller 360-amino-acid Egl-43 protein; this methionine replaces a conserved phenylalanine found in most zinc fingers of this class [Fig. 6b]. There are differences, however, in both the number and arrangement of zinc fingers motifs between Evi-1 and Egl-43; only three of the seven Evi-1 zinc fingers of the first domain are present in Egl-43, and the 47 amino acids that separate the fifth and sixth zinc fingers of Egl-43 are absent in Evi-1 on the basis of sequenced cDNAs (Fig. 6a). Outside the zinc finger motifs there is no obvious similarity between the Evi-1 and Egl-43 protein sequences.

**Discussion**

The *C. elegans* gene egl-43 plays a key role in the migration of the HSNs and also functions in the development of the PHA and PHB sensory neurons. Our genetic results argue that egl-43 functions specifically in HSN migration rather than globally in cell migration: Animals homozygous for egl-43 mutations or deficiencies that span egl-43 are defective in HSN migration but normal for all other long-range cell migrations. egl-43, however, might also function in the outgrowth of the PHA and PHB dendrites [see Results]. This possibility is noteworthy, because cell migration and nerve process outgrowth are phenomenologically and mechanistically similar, requiring similar cytoskeletal elements as well as similar cell surface and extracellular matrix proteins (Singer and Kupfer 1986; Reichardt and Tomaselli 1991). The specificity of egl-43 activity for HSN and phasmid neuron development differs from the pleiotropic activity of most cell migration and axonal outgrowth genes that have been identified in *C. elegans*, which function in the migrations of many cells and axons (Hedgecock et al. 1985, 1987, 1990; Desai et al. 1988; Manser and Wood 1990; Siddiqui and Culotti 1991; McIntire et al. 1992). Such genes define components that function more generally in cell migration and process outgrowth than does egl-43. Our molecular analysis indicates that egl-43 encodes two proteins that contain zinc finger motifs, suggesting that egl-43 controls HSN migration indirectly, by transcriptionally regulating other genes.

Genetic analysis in *C. elegans* has revealed several other genes that appear to function as transcriptional regulators in the control of cell migration. The *C. elegans* genes *lin-39*, *matb*, and *egl-5* encode proteins that contain homeo domains and appear to specify positional information in *C. elegans* [Kenyon 1986; Costa et al.

**Figure 6.** Comparison of the *C. elegans* egl-43 and murine Evi-1 genes. (a) The orders of the Evi-1 and egl-43-related zinc fingers are similar. The lengths and structures of the murine Evi-1 and the *C. elegans* egl-43 cDNAs are represented. The positions of the zinc fingers are indicated by boxes. The Evi-1 transcript encodes 10 zinc fingers in two clusters [1–7 and 8–10], whereas the egl-43 transcript encodes 6 zinc fingers in two clusters [1–3 and 4–6]. Zinc fingers 5–7 of Evi-1 are similar in sequence to zinc fingers 1–3 of egl-43, and zinc fingers 8–10 of Evi-1 are similar in sequence to zinc fingers 4–6 of egl-43. (b) Sequence comparisons between the zinc fingers of Evi-1 and egl-43. Vertical dashes indicate identical amino acids; dots indicate conserved amino acids.
in the process of cell migration. Thus, it seems likely that the egl-43 transcriptional targets might be found in the remaining 18 genes known to function in HSN migration.

Materials and methods

Strains and genetics

Strains were grown at 20°C and maintained as described by Brenner (1974). This paper uses standard C. elegans genetic nomenclature (Horvitz et al. 1979).

In addition to the standard wild-type strain [N2], strains of the following genotypes were used in this work: egl-43(n997), egl-43(n1079), mnDf14/mnC1, mnDf24/mnC1, mnDf25/mnC1, mnDf26/mnC1, mnDf28/mnC1, mnDf56/mnC1, and mnDf60/mnC1. The isolation and initial characterization of the two egl-43 alleles n997 and n1079 were described by Desai and Horvitz (1989) and by Desai et al. (1991). The LG II deficiencies are described by Sigurdson et al. (1984). We observed that mnDf60 complements the egg-laying and HSN migration defects of egl-43(n997) and egl-43(n1079) mutants; this result differs from the partial complementation described by Miller et al. (1992).

mmC1 is a LG II balancer that contains the mutations dpy-10(e128) and unc-52(e444) [Herman 1978].

The strains mnDf24/mmC1 and mnDf60/mmC1 segregated ~50% wild-type animals [mnDf/mmC1], 25% sterile Dpy Unc animals [mnC1/mmC1], and 25% arrested larvae [mnDf/mmDf]. In contrast, mnDf28/mmC1 animals not only segregated arrested larvae but many dead eggs. The embryonic lethality seen in this strain is surprising, because genetically mnDf28 behaves like a smaller deficiency than mnDf24 [Sigurdson et al. 1984]. Moreover, the mmC1 balancer chromosome in this strain breaks down generating Unc non-Dpy and Dpy non-Unc animals. These results suggest that mmC1 is abnormal in this strain and/or that mnDf28 is not a simple deficiency.

Neither of the two deficiencies that remove egl-43 nor the egl-43 mutants exhibit maternal rescue: Homozygous deficiency or egl-43 mutant animals of heterozygous mothers display an HSN migration defect.

Analysis of cell migrations

All long-range migrations were determined to be normal or defective on the basis of the final positions of the migrating cells as observed using Nomarski optics. For each genotype the position of each cell was scored in at least 10 animals. The final positions of cells that migrate during embryonic development were determined in newly hatched first larval (L1) stage animals; these migrations are described by Sulston et al. (1983).

The Q neuroblasts and their descendants, as well as the sex myoblasts [SMs] and the distal tip cells [dtcs] migrate during postembryonic development (Sulston and Horvitz 1977; Kimble and Hirsh 1979). The final positions of the left and right Q descendants were scored in late L1 animals. The final positions of the SMs were scored in third larval (L3) stage animals. In the male the SMs and their progeny migrate posteriorly to various positions in the tail. Because the diagonal muscles, which are derived from the SMs in males, appeared normal in egl-43 mutants, we inferred that the male SMs migrated normally. The pathways taken by the dtcs during their migrations was inferred from the shape of the gonad in late L4 hermaphrodites (Kimble and Hirsh 1979).

In animals homozygous for the two egl-43 mutations n997 and n1079, the Q neuroblasts and their descendants, as well as the sex myoblasts [SMs] and the distal tip cells [dtcs] migrate during postembryonic development (Sulston et al. 1983; Kimble and Hirsh 1979). The final positions of the left and right Q descendants were scored in late L1 animals. The final positions of the SMs were scored in third larval (L3) stage animals. In the male the SMs and their progeny migrate posteriorly to various positions in the tail. The migration of the sex myoblasts is inferred from the shape of the gonad in late L4 hermaphrodites (Kimble and Hirsh 1979).

If the Egl-43 proteins do function in transcriptional control, what genes do they regulate? Presumably they regulate, either directly or indirectly, genes that function
and n1079 and the two deficiencies mnDf24 and mnDf28, only the HSNs were misplaced; all other migrating embryonic cells and the descendants of the QL neuroblasts were in their normal positions. The HSNs were similarly displaced in the egl-43 mutants and the homozygous deficiency animals. In animals homozygous for the deficiency mnDf60, which complements the egl-43 mutations, all embryonic migrating cells, including the HSNs, and the descendants of the QL neuroblasts were in their normal positions. The average positions of the HSNs were scored in >10 animals for the egl-43 mutants and mnDf24 homozygotes using the method described by Desai et al. (1988). When normalized to the positions of the HSNs in wild-type animals (100% migration; n = 48), the average HSN position was 5% (n = 158) for egl-43(n997)/egl-43(n997) animals, 10% (n = 136) for egl-43(n1079)/egl-43(n1079) animals, and 9% (n = 71) for mnDf24/mnDf24 animals. We were unable to score the migrations of the SMs and the dts in animals homozygous for mnDf24, mnDf28 and mnDf60, because these migrations occur after the stage L1 in which the homozygous deficiency animals arrest development.

Homologous deficiency animals were generated by allowing mnDf24/mnC1, mnDf28/mnC1, or mnDf60/mnC1 adult hermaphrodites to lay eggs for 1–2 hr and then removing the adults. After 24–36 hr the arrested Lls on the plate were scored for cell migration defects. The positions of the SMs and the dts in the two egl-43 alleles and of all migrating cells in egl-43(n997)/mnDf28 animals were normal. To score egl-43(n997)/mnDf28 animals, unc-4(e120) egl-43(n997)/+ males were crossed with mnDf28/mnC1 hermaphrodites, and the Unc-4 cross progeny [unc-4(e120) egl-43(n997)/mnDf28] were scored for cell migration defects. unc-4(e120) egl-43(n997) animals were also normal for all cell migrations except the migrations of the HSNs.

Indirect immunofluorescence histochemistry and filling sensory neurons with DiI

Indirect immunofluorescence histochemistry to stain animals for serotonin was performed using the procedure described by Garriga et al. (1993). Rabbit anti-serotonin antiserum was from J. Steinbusch [Free University, Amsterdam, The Netherlands]. FITC-conjugated goat anti-rabbit antiserum was from Cappel, Inc. The cell bodies and axons of the non-HSN serotonin neurons [ADF, NSM, and RIIH neurons in hermaphrodites and males, and the serotonergic CP neurons in males] appeared normal in egl-43 animals. The egl-43 mutants were also scored for defects in axonal outgrowth by staining animals with an antiserum to the neurotransmitter GABA [McIntire et al. 1992]. Rabbit anti-GABA antisera was from J. Steinbusch. The cell bodies and axons of all six classes of GABA-ergic neurons [McIntire et al. 1993] appeared normal in egl-43 hermaphrodites. The amphid (sensory neurons in the head) and phasmid (sensory neurons in the tail) neurons were filled with the fluorescent dye DiI using a modification of the procedure developed by Hedgecock et al. (1985). Both classes of phasmid neurons [PHA and PHB] and 6 of the 12 classes of amphid neurons (ADL, ASH, ASI, ASJ, ASK, and AWB) take up DiI [E. Hedgecock; C. Bargmann, both pers. comm.]. Only the PHA and PHB neurons appeared abnormal in egl-43 mutants.

Germ-line transformation

Germ-line transformation [Mello et al. 1991] was achieved by injecting the DNA to be tested (20 µg/ml) and the plasmid pRF4 [50 µg/ml], which contains the dominant rol-6(su106) roller marker, into the gonads of young adult egl-43(n1079) hermaphrodites. Transgenic animals typically carry such co-injected DNAs as an extrachromosomal array and are identified by the roller phenotype conferred by pRF4 [Mello et al. 1991]. Although egl-43(n1079) animals exhibit defects in egg laying, HSN migration, and phasmid neuron development, only the HSN migration defect is fully penetrant; the HSNs are never in their normal positions in egl-43(n1079) mutants (G. Garriga, unpublished). Because rescue of the HSN migration defect is the most sensitive and unambiguous assay for egl-43 rescue, we defined an egl-43-rescued population as a roller line (derived from an F2 roller) in which >20% of the HSNs were in wild-type positions. The positions of the HSNs were determined in L4 hermaphrodites using Nomarski optics [Desai et al. 1988]. If a roller line appeared to be partially rescued (the HSNs occasionally migrated 50–80% of the distance traveled by wild-type HSNs, which rarely occurs in egl-43 mutants), the positions of the HSNs were determined by staining the lines with anti-serotonin antiserum. In addition, some of the lines were scored for rescue of the egg-laying [Egl] and phasmid neuron defects. Ninety percent of egl-43(n1079) hermaphrodites are Egl, and roller lines that were <40% Egl were considered to be rescued for the Egl defect. Wild-type animals almost always have two phasmid neurons per side that take up the dye DiI, in egl-43(n1079) animals, only 10% of the sides had two phasmid neurons that filled with DiI [data not shown]. When tested, roller lines with >40% of the sides containing two phasmid neurons that filled with DiI were considered to be rescued for the phasmid neuron defect. Roller lines containing p43BA, p43BA-FS, and p43BA-RR were assayed for rescue of the Egl and phasmid neuron defects. By use of the above criteria for rescue, lines containing p43BA and p43BA-RR were rescued for these defects, and lines containing p43BA-FS were not. In these experiments, the lines rescued for the HSN migration were also rescued for the egg-laying and phasmid neuron defects [see Fig. 3c for the number of lines rescued for the HSN migration, PHA and PHB dye-filling, and egg-laying defects].

In our first germ-line transformation experiments, we used cosmid DNA. To refine the position of egl-43, we used plasmids containing DNA subcloned from the cosmid R53 into pBlueScript KS−. The structures of the subclones are illustrated in Figure 3c.

p43BA-FS contains a BglII linker ligated into the SphI site of p43BA. The BglII linker added eight nucleotides and created a frameshift mutation, the methionine at position 403 in Figure 5 was changed to glutamate followed by 20 amino acids and an ochre stop codon, which is upstream of zinc fingers 4–6. To verify that the lack of egl-43-transforming activity of p43BA-FS was caused by the frameshift mutation, we constructed p43BA-RR by digestion of p43BA-FS with BglII and filling in the 5′ overhang using the Klenow fragment of DNA polymerase I [New England Biolabs] according to the procedure described by Sambrook et al. [1989]. p43BA-RR lacks the BgII site, and DNA sequencing confirmed that the reading frame had been restored, replacing an original methionine by four different amino acids. The restoration of egl-43-rescuing activity in p43BA-RR indicated that the lack of rescuing activity in p43BA-FS was caused by the frameshift mutation.

Both egl-43 alleles contain a similar 0.5-kb deletion located at the right end of the 5.6-kb BamHI–ApaI fragment between the EcoRI site closest to the Apal site in Figure 3c and the Apal site, a region that lacks detectable transcripts on Northern blots [data not shown]. It seems unlikely that these deletions cause the egl-43 phenotypes by affecting the 1.0-kb transcript, because deleting the region containing the 1.0-kb transcript from p43BA...
did not abolish rescuing activity [data not shown]. In addition, the 2.5-, 1.8-, and 1.0-kb transcript levels appeared normal in RNA prepared from egl-43(n997) embryos [data not shown]. Thus, we do not know whether the deletion is the egl-43 mutation and affects the expression of the 2.5- and/or 1.8-kb transcripts or a polymorphism that was present in the original strain from which the two egl-43 alleles were derived.

Northern blot hybridization

C. elegans was grown in liquid culture as described by Meyer and Casson [1986], and poly[A]' RNA was isolated as described by Nonet and Meyer [1991]. Electrophoresis, blotting, and probing of the RNA was as described by Sambrook et al. [1989]. The DNA probes used in these experiments were p43BA [which recognizes the 2.5-, 1.8-, and 1.0-kb transcripts], p43BE [which contains sequences from the BamHI site to the nearest EcoRI site of p43BA and hybridized to the 2.5- and 1.8-kb transcripts], p43E2 [which contains the 0.9-kb EcoRI fragment of p43BA and hybridized strongly to the 1.0-kb transcript and weakly to the 2.5- and 1.8-kb transcripts], and p43EA [which contains sequences from the EcoRI site to the Apal site of p43BA and did not hybridize to any transcripts] (see Fig. 3c). The RNA probes were digested before eDNA synthesis in 100 μl of a solution containing 0.2 μg p(dN)6 [Pharmacia], 50 mM Tris HCl [pH 8.3], 75 mM KCl, 3 mM MgCl2, 1 mM dNTPs, 20 units of RNasin [Promega Biotech], and 200 units of M-MLV reverse transcriptase (BRL Gibco). The reaction was incubated at 4°C for 5 min, 22°C for 10 min, 37°C for 60 min, and 95°C for 10 min.

cDNAs were amplified in 100 μl of a solution that contained the cDNA mixture described above, 10 mM Tris HCl [pH 9.0 at 25°C], 50 mM KCl, 0.1% Triton X-100, 25 μM primer 1 [see appended list of primers], 25 μM primer 2, and 3 units of Taq DNA polymerase [Promega Biotech]. The mixture was denatured for 5 min at 94°C followed by 35 cycles of 1-min denaturation at 94°C, 1-min annealing at 55°C, and 3-min extension at 72°C. The reaction ended with an additional 10-min extension at 72°C. The PCR products were cloned into the EcoRV site of pBluescript KS, and their DNA sequences were determined.

The 3' ends of the 2.5- and 1.8-kb transcripts were cloned by use of a variation of the 3' RACE method described by Frohman et al. [1988]. mRNA was incubated with reverse transcriptase, and the dT17 adapter primer was used in place of the random primer p(dN)6. The 3' ends were PCR-amplified as described above with the Eg5 and adaptor primers. The PCR products were cloned into the EcoRV site of pBluescript K5-, and their DNA sequences were determined. The sequence of the RACE product mapped the position of the 3' ends of the 2.5- and 1.8-kb transcripts to the positions shown in Figure 5. Table 1 shows a list of oligonucleotides used as primers. Table 2 shows the primer combinations used in the PCR experiments to determine the structures of the egl-43 2.5- and 1.8-kb transcripts.

DNA sequencing

Deletions of clones used for sequencing were prepared using exonuclease III and S1 nuclease as described by Sambrook et al. [1989]. Electrophoresis, blotting, and probing of the RNA was as described by Sambrook et al. [1989]. Digestion of p43BA with Sall and transcription with T7 polymerase generated a transcript that hybridized to the 1.0-kb transcript and weakly to the 2.5- and 1.8-kb transcripts. Digestion of p43BA with Smal and transcription with T3 polymerase generated a transcript that recognized the 1.0-kb transcript.

PCR of cDNAs and 3' RACE

Residual genomic DNA present in poly[A]’’ RNA preparations was digested before cDNA synthesis in 100 μl of a solution containing 40 mM Tris HCl [pH 7.9], 10 mM NaCl, 6 mM MgCl2, 0.1 mM CaCl2, and 5 units of RQ1 DNase [Promega Biotech] at 37°C for 30 min. The RNA was extracted with phenol–chloroform twice and precipitated with ethanol. Complementary DNA was prepared by incubating 25 ng of DNA-free poly[A]’’ RNA in 20 μl of a solution containing 0.2 μM p(dN)6 [Pharmacia], 50 mM Tris HCl [pH 8.3], 75 mM KCl, 3 mM MgCl2, 1 mM dNTPs, 20 units of RNasin [Promega Biotech], and 200 units of M-MLV reverse transcriptase (BRL Gibco). The reaction was incubated at 4°C for 5 min, 22°C for 10 min, 37°C for 60 min, and 95°C for 10 min.

cDNAs were amplified in 100 μl of a solution that contained the cDNA mixture described above, 10 mM Tris HCl [pH 9.0 at 25°C], 50 mM KCl, 0.1% Triton X-100, 25 μM primer 1 [see appended list of primers], 25 μM primer 2, and 3 units of Taq DNA polymerase [Promega Biotech]. The mixture was denatured for 5 min at 94°C followed by 35 cycles of 1-min denaturation at 94°C, 1-min annealing at 55°C, and 3-min extension at 72°C. The reaction ended with an additional 10-min extension at 72°C. The PCR products were cloned into the EcoRV site of pBluescript K5-, and their DNA sequences were determined.

The 3' ends of the 2.5- and 1.8-kb transcripts were cloned by use of a variation of the 3' RACE method described by Frohman et al. [1988]. mRNA was incubated with reverse transcriptase, and the dT17 adapter primer was used in place of the random primer p(dN)6. The 3' ends were PCR-amplified as described above with the Eg5 and adaptor primers. The PCR products were cloned into the EcoRV site of pBluescript K5-, and positive clones were identified by probing colony lifts with the 5.6-kb BamHI–Apol fragment body labeled with [α-32P]dCTP using random oligonucleotides as primers [Sambrook et al. 1989]. The sequence of the RACE product mapped the position of the 3' ends of the 2.5- and 1.8-kb transcripts to the positions shown in Figure 5. Table 1 shows a list of oligonucleotides used as primers. Table 2 shows the primer combinations used in the PCR experiments to determine the structures of the egl-43 2.5- and 1.8-kb transcripts.

**Table 1. Oligonucleotides used to determine the structure of egl-43 2.5- and 1.8-kb transcripts**

<table>
<thead>
<tr>
<th>Name</th>
<th>Sequence</th>
<th>Start</th>
<th>End</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adaptor</td>
<td>5'-GACTCGAGTGCACTCGAC-3'</td>
<td>28</td>
<td>45</td>
</tr>
<tr>
<td>Eg2</td>
<td>5'-GACTGCAAGCAAGCTTCC-3'</td>
<td>1664</td>
<td>1680</td>
</tr>
<tr>
<td>Eg5</td>
<td>5'-CGGTACGAGACAAAAATC-3'</td>
<td>96</td>
<td>945</td>
</tr>
<tr>
<td>E4</td>
<td>5'-GGTTATCAATCTCGTGCT-3'</td>
<td>1241</td>
<td>1225</td>
</tr>
<tr>
<td>E10</td>
<td>5'-GTTGTGATTTTCACTTGCC-3'</td>
<td>1962</td>
<td>1946</td>
</tr>
<tr>
<td>E11</td>
<td>5'-GCCGCTGAGACATGCCCA-3'</td>
<td>1172</td>
<td>1188</td>
</tr>
<tr>
<td>E12</td>
<td>5'-CGGTTTTTTTTTGAGAAA-3'</td>
<td>566</td>
<td>541</td>
</tr>
<tr>
<td>E30</td>
<td>5'-CGACATACTCGCTGTC-3'</td>
<td>705</td>
<td>682</td>
</tr>
<tr>
<td>E31</td>
<td>5'-ATGAAATAGGAGTTTTTGTGAGAC-3'</td>
<td>-13</td>
<td>20</td>
</tr>
<tr>
<td>Kpe2</td>
<td>5'-GCTCGAAGTCAGACAT-3'</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Start and end refer to the positions of the 5' and 3' ends of the oligonucleotide in the 2.5-kb transcript, the sequence of which is shown in Fig. 5.

*E4 is an oligonucleotide corresponding to sequences within the fifth intron of the 2.5-kb transcript and upstream of the most 5' splice site, which is spliced to SL1, of the 1.8-kb transcript. The sequence of this oligonucleotide corresponds to nucleotides -43 to -27 upstream of the 1.8-kb transcript 5' splice site.

The first 13 nucleotides of this oligonucleotide are a linker containing Xhol, Sall, and ClaI restriction sites.
products. The labeled oligonucleotide (2.5 fmoles) was hybridized to 5 µg of poly(A)§ RNA in 10 µl of a solution containing 0.4 M NaCl, 40 mM PIPES (pH 6.4), and 1 mM EDTA at either 56°C or 65°C for 6 hr. The primer was extended with 200 units of T4 polynucleotide kinase as described by Sambrook et al. (1989). The labeled oligonucleotide used in this experiment (Kpe2) and a primer (E4) located just upstream from the 5' splice site within the fifth exon of E11 lies within an exon shared by both the 2.5- and 1.8-kb transcripts was labeled at its 5' end with [γ-32p]ATP and 1.8-kb transcripts was labeled at its 5' end with [γ-32p]ATP and T4 polynucleotide kinase as described by Sambrook et al. (1989). The labeled oligonucleotide (2.5 fmoles) was hybridized to 5 µg of poly(A)§ RNA in 10 µl of a solution containing 0.4 M NaCl, 40 mM PIPES (pH 6.4), and 1 mM EDTA at either 56°C or 65°C for 6 hr. The primer was extended with 200 units of T4 polynucleotide kinase as described by Sambrook et al. (1989). The labeled oligonucleotide used in this experiment (Kpe2) and a primer (E4) located just upstream from the 5' splice site within the fifth intron. Consistent with this interpretation, the oligonucleotide used in this experiment (Kpe2) and a primer (E4) located just upstream from the 5' splice site within the fifth intron could be used to amplify a PCR product from cDNAs [data not shown].

Table 2. Primer pairs used for PCR amplification of cDNAs

<table>
<thead>
<tr>
<th>Primer 1*</th>
<th>Primer 2</th>
<th>Size of PCR product[b] [bp]</th>
</tr>
</thead>
<tbody>
<tr>
<td>SL1</td>
<td>E31</td>
<td>565 (+)</td>
</tr>
<tr>
<td>SL1</td>
<td>E11</td>
<td>1241 [P] and 624 [P]†</td>
</tr>
<tr>
<td>Eg2</td>
<td>E11</td>
<td>1213 [P]</td>
</tr>
<tr>
<td>E4</td>
<td>E11</td>
<td>629 (+)</td>
</tr>
<tr>
<td>E30</td>
<td>E12</td>
<td>790 (+)</td>
</tr>
<tr>
<td>Eg5</td>
<td>dT17 adapter</td>
<td>853 (+)</td>
</tr>
</tbody>
</table>

*Replacing the SL1 primer with a primer (5'-TCTAGAATTC-CGCCGTTTTAACCCAGTACT-3') for SL2, a second 22-nucleotide RNA that is trans-sliced to C. elegans transcripts (Huang and Hirsh 1989), yielded no detectable RT–PCR products.

†The DNA sequence of either the whole PCR product (+) or of the ends of the PCR product was determined [P].

Because E11 lies within an exon shared by both the 2.5- and 1.8-kb transcripts, amplification of cDNAs by PCR using this oligonucleotide and the SL1 oligonucleotide generated two products and 1.8-kb transcripts was labeled at its 5' end with [γ-32p]ATP and T4 polynucleotide kinase as described by Sambrook et al. (1989). The labeled oligonucleotide (2.5 fmoles) was hybridized to 5 µg of poly(A)§ RNA in 10 µl of a solution containing 0.4 M NaCl, 40 mM PIPES (pH 6.4), and 1 mM EDTA at either 56°C or 65°C for 6 hr. The primer was extended with 200 units of T4 polynucleotide kinase as described by Sambrook et al. (1989). The labeled oligonucleotide used in this experiment (Kpe2) and a primer (E4) located just upstream from the 5' splice site within the fifth intron could be used to amplify a PCR product from cDNAs [data not shown].

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