breathless, a Drosophila FGF receptor homolog, is essential for migration of tracheal and specific midline glial cells

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A Drosophila homolog of the fibroblast growth factor (FGF) receptor was isolated and structurally characterized. After EMS mutagenesis or imprecise excisions of marked P elements inserted upstream to the gene, a phenotypic series of mutations in the locus was isolated. The mutants exhibit defects in the two embryonic tissues in which the receptor is expressed: the tracheal system and the midline. The tracheal cells fail to migrate in severe mutants and remain within the tracheal pits. Hypomorphic alleles exhibit partial migration of all tracheal branches; thus, the locus was termed breathless (btl). In the midline of the mutant embryos, the posterior pair of midline glial cells begins to migrate anteriorly, but fails to reach the posterior commissure. Abnormalities in cell migration appear to be a common denominator for the btl defects in these two disparate tissues. In hypomorphic mutants the cells exhibit partial migration but follow the normal tracts, suggesting that the presence of this receptor is essential for the ability of the migrating cells to recognize external guiding cues.

Key Words: Receptor tyrosine kinase, FGF, Drosophila embryonic development; cell migration; trachea; midline glia

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Communication between cells during development is crucial to the determination of cell fate and cell movement. In most cases the cells are multipotent and must probe their environment to receive cues directing the developmental decisions that they have to make. Cell-surface receptors in Drosophila are essential elements in these processes, as they represent the components used by the cell to receive information from its immediate or more distant environment (for review, see St. Johnston and Nüsslein-Volhard 1992). For example, the Toll receptor transmits signals into the ventral part of the syncytial embryo, which set up the polarity of the dorso-ventral axis (Hashimoto et al. 1988, 1991). The torso receptor transmits the signals that determine the terminal structures of the embryo (Sprenger et al. 1989). Finally, activation of the sevenless receptor by a ligand on the neighboring R8 cell directs the differentiation of the photoreceptor R7 cell in the eye imaginal disc (Hafen et al. 1987; Krämer et al. 1991; for review, see Rubin 1991).

In addition to the determination of diverse cell fates, developmental processes are comprised of cell migration and extension events. In Drosophila, cell migration is necessary at several stages of development. For example, invagination of the mesoderm and its subsequent dorsal migration requires cell movements (Leptin and Grunewald 1990). Similarly, cell migrations are needed during the formation of the tracheal system. Tracheal cells migrate from the tracheal pits and generate the tracheal tree comprised of branches that reach all tissues of the embryo (Campos-Ortega and Hartenstein 1985; Bier et al. 1989). For the cell migration events that establish the central nervous system (CNS), the characteristic migration events of the midline cells have been described in great detail (Klämbt et al. 1991). Two of the three pairs of midline glial cells that exist in each segment (the anterior and middle pair) move posteriorly and, thereby, physically separate the anterior and posterior commissures. The third pair moves anteriorly into the next segment. No specific function has yet been attributed to the latter cells. Mutations that disrupt the normal differentiation or migration of the midline glial cells have been described and include spitz, rhomboid, Star, and faint little ball (flb) (Mayer and Nüsslein-Volhard 1989; Klämbt et al. 1991; Raz and Shilo 1992). Some of these loci encode cell-surface molecules (Scheiter et al. 1986; Bier et al. 1990). It is not possible to say, however, whether these cell-surface molecules are specifically involved in the process of cell migration, as the midline glial cells fail to differentiate or die in these mutant backgrounds.

One candidate for a cell-surface receptor that is required for cell migration is a Drosophila fibroblast
growth factor (FGF) receptor homolog that we have isolated (DFGF-R1). This locus at 70C/D encodes a receptor tyrosine kinase that has all the hallmarks of the FGF receptor subfamily; most notably, it possesses immunoglobulin-like domains at the extracellular region and a split cytoplasmic kinase with a small kinase insert sequence (Glazer and Shilo 1991). A second Drosophila FGF receptor homolog (DFGF-R2) has been isolated recently (M. Beiman, L. Glazer, and B. Shilo, unpubl.). It does not appear, however, to have overlapping functions with the DFGF-R1 protein. The extracellular domains of the two receptors show only 30% amino acid sequence identity, suggesting that they may recognize a different set of ligands. Furthermore, the two loci do not share common sites of expression during embryogenesis. DFGF-R2 is expressed in the mesoderm, whereas DFGF-R1 was shown to be expressed in the developing tracheal system and the midline cells (Glazer and Shilo 1991).

Chromosomal deficiencies removing the DFGF-R1 locus give rise to mutant embryos in which the tracheal pits form, but the subsequent processes of cell migration and extension do not take place (Glazer and Shilo 1991). Because the chromosomal deficiency that was used removed several loci, it was not possible to conclude unambiguously whether the FGF receptor homolog itself is indeed crucial for tracheal cell migration. Here, we present the isolation of mutations and DNA excisions that specifically affect the DFGF-R1 gene. These mutants represent a phenotypic series of alleles in the locus. The mutants are defective in cell migration in two embryonic tissues in which the receptor is expressed. The tracheal cells display a reduced ability to migrate, and the tracheal tree is not formed. In the midline, the posterior midline glial cells do not complete their anterior migration to reach the posterior commissure. The CNS axon pattern also shows subtle abnormalities. The mutant phenotype demonstrates that the DFGF-R1 receptor is crucial for the migration capacity of different cell types and may even be directly responsible for receiving external cues guiding these intricate cell migration events.

Results

Structure of DFGF-R1

The sequence of genomic and cDNA clones of DFGF-R1 has been presented previously (Glazer and Shilo 1991). It showed a cytoplasmic tyrosine kinase domain that is most closely related to the vertebrate FGF receptor subfamily, as well as an extracellular region with three immunoglobulin-like domains. The extracellular sequence was not complete, however, as it did not contain the predicted signal peptide. The sequence was extended by isolation of cDNA clones that encode the amino-terminal domain and by sequencing the parallel genomic region. These sequences showed that the extracellular region of DFGF-R1 contains a hydrophobic signal peptide and two more immunoglobulin-like domains, in addition to the three domains that were identified previously (Fig. 1).

Although the presence of five immunoglobulin-like domains was described for the PDGF receptor subfamily (Yarden and Ullrich 1988), it has not been found previously for members of the FGF receptor subfamily, which typically contain only two or three immunoglobulin-like domains. Nevertheless, DFGF-R1 is clearly a member of the FGF receptor family, as demonstrated by the characteristic size and consensus sequences of the immunoglobulin domains and by the organization of the cytoplasmic kinase domain (Fig. 2).

Alternative splicing processes have been implicated in generating diversity in the number and structure of the immunoglobulin domains of several vertebrate FGF receptors (Reid et al. 1990; Yayon et al. 1992). This region of DFGF-R1, however, does not appear to be modified by alternative splicing, as the cDNA and genomic clones are colinear along the entire extracellular coding sequence. The only intron that was identified in the gene is 35.5 kb in the genomic clone.

Figure 1. Sequence of DFGF-R1 extracellular domain. The sequence of the amino-terminal three immunoglobulin-like domains of DFGF-R1 is shown. The hydrophobic signal peptide is underlined. The cysteine residues defining the borders of the immunoglobulin-like domains are circled, and the domain number is shown in Roman numerals. The putative initiator codon was designated as residue 1. Cysteine residue 262 was tentatively marked as 1 in Glazer and Shilo (1991). The putative protein contains 1051 residues, including the signal peptide.
Mutations in the DFGF-R1 locus

Two strategies were used to isolate mutations in the DFGF-R1 locus: imprecise excision of marked P elements and an 'F2 lethal screen' to isolate ethylmethane sulfonate (EMS)-induced mutations. The distinct expression pattern of the gene in the tracheal and midline cells has allowed the identification of two different enhancer trap elements that show a similar pattern of expression and map to the same chromosomal locus (Bier et al. 1989; C. Klambt and C.S. Goodman, unpubl.). Southern blot analysis was carried out on these two lines to determine their integration sites with respect to the DFGF-R1 locus. The analysis shows that both elements have integrated within several hundred bases of each other (on either side of the genomic BamHI site), 2 kb upstream of the transcriptional start site (Fig. 3). The integration of the elements upstream of the gene agrees with the observation that both 6-81 and H82 homozygous flies are viable. H82 homozygotes appear but at a reduced frequency.

Both elements were excised by dysgenic crosses (for details, see Materials and methods), and three excision mutants, which removed DNA sequences in the direction of the DFGF-R1 gene, were further characterized. Southern blot analysis to remove sequences 5' to the integration site, 2 kb upstream of the transcriptional start site (Fig. 3). The analysis shows that both elements have integrated within several hundred bases of each other (on either side of the genomic BamHI site), 2 kb upstream of the transcriptional start site (Fig. 3). The integration of the elements upstream of the gene agrees with the observation that both 6-81 and H82 homozygous flies are viable. H82 homozygotes appear but at a reduced frequency.

The tracheal phenotype of DFGF-R1 mutants: breathless

The process of tracheal differentiation and migration can be followed by several markers. Enhancer trap lines allow tracing of the number and position of the tracheal nuclei at all stages of embryogenesis (Bier et al. 1989). Alternatively, antibodies recognizing tracheal membrane antigens [e.g., DFGF-R1 or crumbs (Tepass et al. 1990; Glazer and Shilo 1991)] help to visualize the out-

Figure 2. Scheme of the DFGF-R1 protein. (a) The hydrophobic signal peptide and transmembrane domains; (b) the kinase domain. An acidic box typical to FGF receptors is located between the third and fourth immunoglobulin domains, and is stippled.

Figure 3. A map of the DFGF-R1 locus and the induced excisions. The breakpoints induced by the imprecise excisions of the enhancer trap elements are shown by arrows. The positions of the deletions in 6-81A1 and H82A3 that map to the 2-kb BamHI-NdeI fragment are shown in parentheses. We could not identify other restriction sites within this fragment that would allow the localization of the positions with greater accuracy. [B] BamHI, [E] EcoRI, [G] BglII, [N] Ndel, [X] XbaI. [P] The site of DFGF-R1 transcription initiation, as mapped by primer extension.
lines of the cells comprising the tracheal system. Finally, antibodies that recognize components secreted into the lumen of the tracheal tubes were identified in monoclonal antibody screens [N. Patel and C.S. Goodman, pers. comm.] or fortuitously in polyclonal antibody preparations (antiserum 84, N. Walker and B. Shilo, unpubl.). These antibodies are extremely useful in following the normal pattern generated by the tracheal tree and abnormalities that may occur in different mutant backgrounds.

The process of tracheal differentiation during embryogenesis has been divided into two phases [Campos-Ortega and Hartenstein 1985]. First, the cells that will form the tracheal system are generated at stages 10–11 by cell divisions in the ectoderm, which generate ~100 cells in each of the tracheal pits [G. Manning and M. Krasnow, pers. comm.]. In the second phase (encompassing stages 12–17), the entire tracheal system, on each side of the embryo, is generated by the apparent migration and extension of the ~900 tracheal cells that originated in the 10 tracheal pits.

Our earlier observations have indicated that tracheal pits form in embryos homozygous for a chromosomal deficiency of the 70C/D region [Glazer and Shilo 1991]. Thus, although DFGF-R1 is expressed from the early stages of tracheal formation, it is not required for the initial invagination of the tracheal precursor cells to form the tracheal pits. These embryos showed, however, that the phase of cell migration is defective. The availability of different alleles specifically affecting the DFGF-R1 locus provides an opportunity to examine directly the role of this locus in tracheal migration. Embryos homozygous for the different alleles were stained with an antibody that marks the lumen of the tracheal tubes (Fig. 5). These alleles define a phenotypic series. The most severe phenotype is observed for the two EMS-induced alleles. Even at late stages of embryogenesis, the tracheal cells do not undergo significant migration; they remain clustered at the site of the tracheal pit. On the basis of this phenotype, we termed the DFGF-R1 locus *breathless* (*btl*). Although no migration takes place in the severe alleles (LG18 and LG19), the tracheal lumen antigen that is normally detected by antiserum 84 is secreted into the cavity of the residual tracheal pits. The two EMS-induced alleles appear to represent the complete null phenotype. Embryos homozygous for the LG18 or LG19 alleles display a similar phenotype to embryos in which each of these mutations is carried over a deficiency for the 70C/D chromosomal region (not shown).

The *btl* alleles generated by excision of the marked P elements are hypomorphs. The severity of the phenotypes is in accordance with the residual levels of *btl* transcripts that are synthesized by each of the alleles. In H82A3, only very limited migration of the tracheal cells takes place. The cells begin to migrate from the pits and form short stumps of the visceral branches and the dorsal trunk. In H82A11, the migrations are more pronounced. In several segments, the dorsal trunk is continuous and the branches that normally extend to the gut and CNS are partially formed. Finally, in 6-81A1 the dorsal trunk is continuous in most segments, and the dorsal, visceral, and head branches are more complete. The hypomorphic nature of these three alleles is also reflected by the fact that their phenotype over a deficiency is more severe than the homozygous phenotype (not shown).

Because expression of β-gal by the enhancer trap is retained in the hypomorphic *btl* alleles, it was possible to follow the fate of the tracheal nuclei in these mutant backgrounds. As judged by fluorescence microscopy, the number of nuclei in the mutants appears similar to wild-type embryos [Fig. 5h]. The partial migration that is observed for these hypomorphic alleles is reflected in a limited migration of the nuclei; however, no major loss of cells has occurred. In spite of the failure of these cells to migrate, they express the lumen-specific antigen at the proper time (shown above).

The hypomorphic *btl* alleles show defects in the formation of all tracheal branches; thus, they appear to be
defective in a process common to all migrating tracheal cells. It is important to note that when partial migration of the tracheal cells occurs, the branches appear to form in the normal positions. This would suggest that DFGF-R1 is necessary for the tracheal cells to follow the signals guiding their migration but does not affect the actual migration cues.

Expression of btl in the midline cells

In addition to btl transcription in the tracheal system, the gene is also expressed in the midline cells of the embryonic CNS (Glazer and Shilo 1991). This expression pattern can be visualized by β-gal localization in the two enhancer trap lines. Whole-mount in situ hybridization with a btl-specific probe reveals that expression starts just before the onset of gastrulation, as early as stage 5. Cells at the posterior pole, as well as two groups of ventral cells located at the anterior tip of the embryo, show hybridization signals (Fig. 6a).

During gastrulation, btl expression is detectable in two single-cell-wide stripes flanking the invaginating mesoderm anlage. These cell rows correspond to the mesectodermal precursor cells moving toward the midline, where they meet to form the single row of midline precursor cells (Fig. 6b,c). It is interesting to note that btl expression is modulated in a segmental manner from the onset of transcription in the midline. During stage 12, btl expression is most pronounced in the midline cells of the posterior half of the segment. This includes the ventral unpaired median (VUM) neurons and the posterior pair of midline glial cells. btl expression in these midline cells continues until the end of stage 12 (not shown).

CNS phenotype of btl

The formation of segmentally arranged commissures requires a diverse set of cell–cell interactions in the midline of the CNS (Klambt et al. 1991). Impairment in any of these interactions subsequently leads to defects in the
pressed in the midline cells, we analyzed the CNS axon pattern of different btl mutants [H82dA3 and LG19]. The homozygous mutant embryos were unambiguously identified on the basis of their tracheal phenotype and by the absence of staining contributed by the "Blue balancer" chromosome. Initially, both commissures are observed (Fig. 7). Instead of the well-defined space between the anterior and posterior commissures in wild-type embryos [where the RP neurons 1, 2, and 3 are located (Jacobs and Goodman 1989b)], a smaller irregular and more rounded space is seen in homozygous btl embryos (Fig. 7a,b). This commissural phenotype is slightly more pronounced in LG19 compared with H82dA3 [not shown], in accordance with the more severe tracheal phenotype of LG19. It is important to note that this axon pattern phenotype emerges after the separation of the segmental commissures has taken place.

To address the question of whether any of the midline cells may be responsible for this phenotype, we analyzed the midline cell lineages using antibodies and enhancer trap lines as cell-specific markers (Klämbt et al. 1991). The anterior and middle pair of midline glial cells [MGA and MGM cells, respectively; Jacobs and Goodman 1989a] are present in homozygous btl mutant embryos, but their nuclei sometimes appear slightly displaced. The posterior migration of the MGA and MGM cells, as well as the midline migration of the RP motor neurons, occurs normally in the mutant embryos [data not shown]. These glial cells are the midline cells that express the lowest level, if any, of the DFGF-R1 transcript. The VUM neurons form normally and show no defects when stained with the monoclonal antibody 22C10, which labels their cell surfaces, in addition to other cells. Subtle defects in the position of individual neurons of the VUM cluster or in their final projection pattern, however, may have escaped our attention.

The most striking cellular defect in btl embryos is seen in the posterior pair of midline glial (MGP) cells in each segment. These two glial cells are born just anterior to the MGM cells and then migrate during stage 12 anteriorly into the next segment, where they reach the posterior commissure at the beginning of stage 13 when the segmented commissures have already been separated (Klämbt et al. 1991). In homozygous btl mutant embryos this migration is defective, and the MGP cells remain closer to the anterior commissure of their original segment. As the CNS condenses during later stages of embryonic development, these glial cells are displaced dorsally and the CNS axon pattern phenotype becomes clearly visible (Fig. 7d). The defects in the MGP cells appear to be cell autonomous, as these cells express the btl gene.

The MGP migration defect and the axon pattern abnormalities are the only CNS phenotypes that we could detect during embryonic development, in spite of the identification of the btl transcript in several midline cell types. Whole-mount in situ hybridization experiments with a rhomboid-specific probe, which labels the mesectodermal precursor cells (Bier et al. 1990), did not reveal any defects in their number or position in btl mutant embryos.

Discussion
The role of breathless in tracheal migration
The process of tracheal formation represents an amenable biological system in which the intricate regulation of cell migration can be dissected experimentally. The tracheal tree is formed by cell migration and extension in a period of only several hours (stages 12-17, 8-16 hr after egg lay). Finally, the structures are generated from a finite number of cells (~900 cells for each side of the embryo).

Several key issues remain unanswered. It is not known whether cell lineage plays a role in tracheal morphogenesis or whether all tracheal cells are equivalent and choose their migration path stochastically. Although it appears that the cues for migration must be provided by the microenvironment, it is not known whether these signals are diffusible or anchored to cell membranes or to the extracellular matrix. Finally, the cues for the formation of the different tracheal branches may be identical.
Alternatively, each branch may be guided by different signals that could be recognized by a diverse set of cell-surface receptors.

The btl alleles described in this work represent a phenotypic series in which tracheal cell migration is defective. No migration takes place in the severe alleles, whereas partial migration of all branches is observed in the weaker alleles. In situations of partial migration, the tracts that do form appear normal. This observation suggests that DFGF-R1 activity is crucial for the tracheal cells to recognize the cues for migration, rather than generation of these migration signals. It is plausible to envisage the cues for tracheal migration as external signals that are presented by the tissues that are in contact with the migrating tracheal cells, such as the ectoderm or mesoderm.

The ability of mutations in a single locus to alter the migration of all tracheal branches is dramatic and suggests two possible models for the role of btl in tracheal migration. One option is that btl is required by all tracheal cells to respond to the external cues for migration. According to this model, the receptor is not directly responsible for detecting these cues, which may differ for each subset of branches. For example, activation of the DFGF-R1 protein may be necessary to trigger a set of transcription factors that would allow the expression of receptors that will detect the external cues for migration. According to this possibility, the potential ligand of DFGF-R1 need not be localized, as it is only necessary for priming the tracheal system to respond to the external cues.

An alternative option is that DFGF-R1 itself is the receptor through which the external cues guiding migration are transmitted. This notion is supported by the observation that in the btl alleles generated by imprecise excision of marked P elements there is a strict correlation between the level of DFGF-R1 expression and the extent of migration of the tracheal cells. The ligand triggering DFGF-R1 may be soluble or it could be anchored to the cell membrane, the extracellular matrix, or to heparin–sulfate proteoglycans. FGF-like molecules in vertebrates were shown to be anchored in a nonsoluble form [for review, see Ruoslahti and Yamaguchi 1991]. Furthermore, in several cases, FGF can be displayed in a biologically active form only when it is associated with heparin–sulfate proteoglycans [Yayon et al. 1991]. The implication of this model is that the ligand of DFGF-R1 would be made [in a diffusible or insoluble form] by neighboring cells in a prepattern that precedes the migration of the tracheal cells. Although the identities of the cells marking each of the migration tracts at different positions need not be identical, they may be producing the same signaling molecule (i.e., the putative ligand of DFGF-R1).

FGF has been implicated in the regulation of cell migration in vertebrate systems. It was shown to be a potent angiogenic factor, capable of stimulating division and migration of endothelial cells toward high local concentrations of FGF [Folkman and Klagsbrun 1987]. Because FGF is released as a soluble factor, the growing blood vessels that are attracted to the region develop in a random fashion, rather than in a highly organized and stereotypic structure. In this respect, the formation of the tracheal tubes in the embryo may be more similar to
the construction of blood vessels during embryogenesis or organogenesis in vertebrate organisms, where the vessel pattern is very similar in all individuals.

The role of btl in glial migration

Although the btl gene is expressed from the early stages of mesectoderm differentiation, no phenotype is seen in these cells during gastrulation of mutant embryos. The absence of any phenotypes at this stage may be accounted for by postulating that the function of DFGF-R1 at this stage is redundant or nonessential. Alternatively, although the gene may be transcribed in the mesectoderm precursor cells during gastrulation, some time may elapse until the processing of the protein and its translocation to the plasma membrane is completed. The DFGF-R1 protein could be detected by immunohistochemistry only after the mesectodermal cells meet at the midline, after the completion of gastrulation (Glazer and Shilo 1991).

A characteristic defect was detected, however, in the MGP cells of btl embryos. These cells do not complete their migration toward the posterior commissure of the next segment, and stay closer to the anterior commissure, where they were originally born and eventually become displaced dorsally. It is not clear how defects in the migration of MGP cells may lead to the commissure phenotype. During the initial formation of the commissures, the MGP cells are not thought to play an instructive role, as they are located at a distance from the commissures (Klämbt et al. 1991). The defects that are observed in btl embryos suggest that the MGP cells may be needed for the stabilization of the final commissure pattern during later stages. For example, these cells may normally stop the posterior migration of the MGC cells preventing them from growing around the posterior commissure. Therefore, when the MGP cells do not reach their final destination in the mutant embryos, development of the commissures is initially normal. As the nerve cord condenses, however, the MGC cells may overgrow the posterior commissure. Instead of separating the two commissures, they now “pull” them together, thus generating an irregular spacing between them. Alternatively, the axon pattern defects could be the result of a change in the VUM cluster of midline neurons or in the MGA and MGM cells. No major positional changes were detected in the VUM neurons, which express the btl gene. Similarly, no alteration in the positions of the MGA and MGM cells, which do not express detectable levels of btl RNA, was observed. It is possible, however, that subtle changes in the interaction between these cells, which may be crucial for late commissural morphogenesis, could also contribute to the commissural defects of btl embryos. Therefore, the precise causal relationship between the altered MGP cell migration and the commissural phenotype remains to be demonstrated.

Possible common themes

Although the embryonic tracheal and midline cells represent completely different lineages, it is striking that btl appears to play a similar role in both cases. The migration of the tracheal cells, as well as that of the MGP cells is not completed in btl mutant backgrounds. Thus, DFGF-R1 emerges as a central player in processes of cell migration for both tissues.

There is one additional tissue expressing the gene. During stages 11–13, a small group of ectodermal cells that will eventually form the duct of the salivary gland show btl expression (not shown). In forming the duct, these cells normally migrate a short distance to connect the salivary gland with the pharynx (Campos-Ortega and Hartenstein 1985). No apparent defects in the formation of salivary duct have been observed, however, when the development of the duct cells was followed in H82Δ3 mutant embryos stained with anti-β-gal antibodies.

Several receptor tyrosine kinases in Drosophila have been shown to play a pivotal role in different contexts of cell determination (for review, see Shilo 1992). The torso protein transmits into the syncytial embryo the signals that define the position of the terminal structures (Sprenger et al. 1989). Drosophila epidermal growth factor receptor (DER)/fib has a diverse set of roles during embryogenesis, which include the determination of the identity of the ventral ectoderm (Schechter and Shilo 1989, Raz and Shilo 1992). In the egg chamber, alleles of the DER locus (torpedo) have demonstrated that the receptor is also required to transmit signals that determine the fate of the dorsal follicle cells (Schüpbach 1987, Price et al. 1989). Finally, the sevenless receptor is required in the eye imaginal disc for the future R7 photoreceptor cell to initiate neuronal differentiation, after triggering by the environment (Hafen et al. 1987; Krämer et al. 1991, Rubin 1991).

The functions of btl described in this discussion appear different from the functions of the previously characterized Drosophila receptor tyrosine kinases. In btl mutants, the identity of the cells that normally express the receptor does not appear to be altered. The tracheal cells continue to display tracheal markers, such as an antigen that is secreted into the tracheal lumen or the specific β-gal expression from the enhancer trap inserted upstream to the btl gene, which marks the tracheal cells throughout their development. Likewise, all midline cells express normally the diverse differentiation markers that were tested. Although the identity of the tracheal and midline cells is normal, these cells fail to follow their typical migration routes. Thus, DFGF-R1 appears to represent a crucial element in the migration processes of these cells.

In mice, the involvement of a receptor tyrosine kinase in processes of embryonic cell migration has been suggested on the basis of the phenotype of the White spotting mutations in the c-kit locus. The phenotype of these mutations affects three populations of cells: neural crest-derived melanocytes, primordial germ cells, and hematopoietic stem cells. The same tissues are similarly altered by mutations in the Steel locus, which encodes the ligand of the kit receptor (for review, see Silvers 1979). Steel was shown to be expressed during embry-
genesis in cells associated with migratory pathways and homing sites of melanoblasts, germ cells, and hematopoietic stem cells [Matsui et al. 1990]. The Steel protein exists in a soluble as well as a cell-surface form. A mutation encoding a Steel protein in which the transmembrane and cytoplasmic domains are deleted, but the soluble form of the protein is intact [S^Fl], results in developmental defects in the same three lineages typically affected by Steel and White mutations [Brannan et al. 1991, Flanagan et al. 1991]. Thus, the membrane-associated form of Steel may be crucial for guiding the migration and homing of embryonic cells, by triggering the kit receptor at very restricted sites.

In conclusion, this work has demonstrated the central role of DFGF-R1 in diverse processes of embryonic cell migration. Many questions concerning the precise function of the receptor in regulating these processes remain open. The key to understanding some of these issues may be the ligand. Future studies, including the identification and characterization of the ligands triggering DFGF-R1, should tell whether activation of the receptor is a prerequisite for tracheal cells to follow different migration tracts. Alternatively, the signal provided by the ligand may mark different paths of cell migration, which could be recognized directly by activation of the receptor encoded by btl.

Materials and methods

Fly strains

Two chromosomal deficiencies uncovering the 70C/D region, kindly provided by the Bloomington stock center, were used. [fz GF3b [70B-70D6] and fz GS1a [70C6-15-70E4-6]]. The fz enhancer trap line [Bier et al. 1989] was kindly provided by L. Jan and Y.N. Jan [University of California, San Francisco]. The H2B line containing the P-lwB element was generated by the enhancer trap screen carried out in C.S. Goodman’s laboratory (C. Klambt and C.S. Goodman, unpubl.). The following enhancer trap lines were used in this study to follow the differentiation of the midline cells: Aul, which labels the MGP cells; AA142, which labels all midline glial cells; and AE60, which labels the VUM neurons [Klambt et al. 1991].

Genetic manipulations

To generate mutations in the DFGF-R1 locus, excisions of two enhancer trap elements inserted upstream of the gene were induced, and the events of imprecise excision were scored. Males homozygous or heterozygous for the 6-81 or H82 insertions of the enhancer trap elements were crossed to females carrying the Xa/+. 1000 females were mated over a hundred pair matings were followed, and lines without fzGF3b/TM3 5b females were scored as imprecise excision mutations in the DFGF-R1 locus.

Molecular biology

To isolate cDNA clones containing the 5' region of the DFGF-R1 transcript, an embryonic 3- to 12-hr cDNA library (Poole et al. 1985) was screened with a 7-kb XbaI genomic fragment of DFGF-R1, labeled by random priming. One clone, with an overlap to the previously isolated cDNA clones, was identified. This clone was shown to include the entire 5'-coding sequence and a stretch of 5'-noncoding sequence. Using this probe we have also identified a clone containing the 3' region of a transcript encoded by a gene located upstream to the DFGF-R1 locus.

cDNA and genomic clones of DFGF-R1 were sequenced according to the Sanger protocol by use of the Promega or Sequenase (U.S. Biochemical) kits, and synthetic primers. Primer extension was performed according to Sambrook et al. [1989] using as a template 10 mg of total embryonic RNA and a synthetic primer located 60 bp downstream to the DFGF-R1 initiator codon.

To determine the sites of insertion of the enhancer trap elements, as well as the size and location of the excisions, Southern blot analysis was carried out. Total genomic DNA (1–2 µg) was digested with restriction enzymes and analyzed by probes prepared from different genomic or cDNA fragments of the DFGF-R1 gene. The position of each breakpoint within a restriction fragment on the DFGF-R1 map could therefore be determined.

RNA in situ hybridization and immunostaining

To determine the residual level of DFGF-R1 transcripts in embryos homozygous for various excisions, the excision chromosome was maintained over a TM3 balancer chromosome containing a lacZ gene driven by the hunchback promoter (kindly provided by W. Driever). Embryo collections were stained for 10 min with X-gal. This short staining was sufficient to allow the detection of lacZ expression contributed by the Blue balancer, but did not stain the sites of DFGF-R1 expression by the intact lacZ gene within the residual enhancer element. After the short exposure to X-gal, embryos were washed in PBS and 0.1% Triton X-100, and processed further for RNA in situ hybridization according to standard protocols (Tautz and Pfeifle 1989). Embryos were probed with a digoxigenin-labeled 2.2-kb DFGF-R1 cDNA fragment. During the staining of alkaline phosphatase, the reaction was carried out for a limited time, in an attempt to retain a linear relationship of color intensity.

Embryos were immunostained according to standard procedures (Ashburner 1989), followed by a secondary antibody conjugated to either horseradish peroxidase or rhodamine (Jackson Laboratories). The tracheal-specific antibody [48] was generated fortuitously by injection of acrylamide into guinea pigs. To visualize β-gal expression, a monoclonal antibody (Promega) or a rabbit polyclonal antibody (Cappel, USA) was used. The CNS

Grigliatti (1986). They were subsequently crossed to ru cu Pr ca/TM6B females. F1 ru cu ru cu Pr ca progeny males were pair-mated to Df. fzGF3b/TM3 5b females. One thousand six hundred pair matings were followed, and lines without ru cu/ fzGF3b progeny were characterized further. They were mated to fzGS1a/TM3 5b females to identify lethal mutations on the ru cu chromosome, which map to the chromosomal region defined by the overlap between the two deficiencies. Four such lines were identified and shown to define three complementation groups (LG18 and LG19, LG12 and LG9). The complementation group defined by the LG18 and LG19 mutations was shown to be allelic to the excision mutations in the DFGF-R1 locus.
was visualized by mAb BP102 kindly provided by N. Patel and C.S. Goodman.

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