FGF Is an Essential Mitogen and Chemoattractant for the Air Sacs of the Drosophila Tracheal System

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Summary

The Drosophila adult has a complex tracheal system that forms during the pupal period. We have studied the derivation of part of this system, the air sacs of the dorsal thorax. During the third larval instar, air sac precursor cells bud from a tracheal branch in response to FGF, and then they proliferate and migrate to the adepithelial layer of the wing imaginal disc. In addition, FGF induces these air sac precursors to extend cyto-neme-like filopodia to FGF-expressing cells. These findings provide evidence that FGF is a mitogen in Drosophila, correlate growth factor signaling with filo-podial contact between signaling and responding cells, and suggest that FGF can act on differentiated tracheal cells to induce a novel behavior and role.

Introduction

Many functions and mechanisms that program development are common to different tissues and diverse animals. Instructional regulators, signaling pathways, and general and specific growth factors constitute a shared vocabulary of development. Fibroblast growth factor is one example. It is produced extensively in the animal kingdom, and it regulates cell division and differentiation in many contexts (reviewed in Basilico and Moscatelli, 1992). In Drosophila, there is a single FGF gene, branch-less (bnl) (Sutherland et al., 1996). It plays a key role modeling the complex branched structure of the tracheal system of the Drosophila embryo and larva. We show here that, in unexpected ways, Bnl-FGF also plays a key role in the development of the tracheal system of the adult.

The tracheal system is a tubular epithelial network that delivers oxygen to tissues throughout the body (reviewed in Manning and Krasnow, 1993). In the embryo and larva, it has bilaterally symmetric trunks that orient rostrocaudally and open to the outside through anterior and posterior spiracles. With segmental periodicity along their lengths, these trunks connect to smaller tubes that branch and give rise, successively, to smaller and smaller branches that ramify throughout the body. The shape and size of the tubes are genetically programmed and stereotyped. Fine terminal branches (“tracheoles”) also form in many locations in response to local requirements for oxygen, creating a web that spreads over many tissues (Jarecki et al., 1999).

There has been much progress toward understanding the genetic and cellular mechanisms controlling branching morphogenesis in the embryo and larva. In the embryo, the transcription factors Tracheless (Trh) (Isaac and Andrew, 1996; Wilk et al., 1996) and Tango (Ohshiro and Saigo, 1997; Sonnenfeld et al., 1997) cooperate to select and direct the formation of 20 clusters of tracheal epithelial invaginations known as tracheal pits, each containing about 80 cells. Elongation, migration, and branching then create, without further cell divisions, a ramified array of about 500 branches from each cluster. FGF signaling is the critical determinant of the tracheal branching pattern, involving both expression of bnl in cells in the proximity of tracheae and expression of the FGF receptor Breathless (Btl) and other constituents of the FGF signaling transduction pathway in tracheal cells (reviewed in Metzger and Krasnow, 1999). Among the essential constituents is Stumps, a cytoplasmic protein whose activity is unknown (Michelson et al., 1998; Vincent et al., 1998; Imam et al., 1999). In performing its role as architect of branch morphogenesis, FGF signals between neighboring groups of cells, but it is not known over what distance these signals travel or how they move.

Metamorphosis presents special challenges to the tracheal system. The process that transforms the Drosophila larva into an adult fly consumes larval tissues and creates new organs using imaginal cells that were prepared during larval development. A new tracheal system that will satisfy the aerobic requirements of the specialized tissues of the adult must be built. But, in addition, the organs of the pupa must be kept oxygenated while the adult develops (reviewed in Whitten, 1980; Manning and Krasnow, 1993). Transformation of the tracheal system begins during the third larval instar, when imaginal tracheoblasts start to divide (reviewed in Manning and Krasnow, 1993). These proliferating tracheoblasts spread over the larval tracheal system, using it as a scaffold to form an extensive branching network before the larval cells histolyze (Matsuno, 1990). Some tracheoblasts elaborate coiled structures that are unique to the pupa. Others grow to form the air sacs of the adult. Air sacs are large reservoirs that are juxtaposed with major muscle systems and with the brain (reviewed in Miller, 1965; Whitten, 1980; Manning and Krasnow, 1993). These structures have been thought to form as dilations of the main tracheal trunks, which are the direct descendants of the main tracheal trunks of the embryo and larva. The work we describe here shows that, in the dorsal thorax, these air sacs originate independently from a distinct population of cells.

The majority of the adult thorax, including most of the dorsal thoracic epidermis, the wing, and flight muscle, is produced by the wing imaginal disc (reviewed in Cohen, 1993). This organ arises as a tubular invagination of the epidermis, and, when it grows and flattens during the larval periods, it develops four distinct cell types. It has squamous peripodial cells on one surface, columnar epithelial cells on the other surface, a distinct group of adepithelial cells that nestle against the most proximal columnar epithelial cells, and stalk cells that connect the disc to the epidermis. A large tracheal branch attaches in...
some manner to the columnar epithelial surface. It orients along the dorsal/ventral axis of the disc, but it does not ramify to generate multiple contacts with the disc cells (Jarecki et al., 1999). Therefore, no framework exists to serve as a template for the tracheolar network that provides oxygen to the thoracic cells of the adult. The mechanisms responsible for the branch formation and path finding that produce the extensive and complex tracheolar network in the adult thorax remain to be identified.

The work described here represents an effort to understand the role of FGF in wing disc development. These investigations led to the identification of a new cell type in the wing disc that migrates and proliferates in response to FGF. These cells contact FGF-expressing cells across at least one cell layer by extending long cytoneme-like filopodia (Ramirez-Weber and Kornberg, 1999). We found that these cells are destined to form the air sacs that associate with the flight muscles in the adult thorax, but they are distinct from the cells that form the larval trachea or from the group of imaginal precursors that are programmed to generate tracheae in the pupa and adult (Manning and Krasnow, 1993). There are intriguing implications of this novel strategy of tissue development.

**Results**

**Expression of FGF and FGF Receptors in the *Drosophila* Wing Imaginal Disc**

To investigate the role of FGF in *Drosophila* wing disc development, we examined the patterns of expression of *btl* (Sutherland et al., 1996) and the two FGF receptor genes, *btl* and *heartless* (*htl*) (Klämbt et al., 1992; Shishido et al., 1993; Shishido et al., 1997), in third instar discs. *btl* expression was restricted to a small group of cells in the columnar epithelium (Figures 1A–1C). An exact count of their number was problematic, since the apparent level of expression in many cells was very low, but we roughly estimate their number to be between 15 and 60 in early third instar discs and between 80 and 150 in late third instar discs. These cells straddle the anterior/posterior compartment border and are dorsal to the region of the prospective wing blade. The cells that express *bnl* most strongly are ventral to the progenitor of the aPA machrochaete (Sato et al., 1999b), as indicated by double staining for Achaete protein (data not shown). They are in a region that contributes to the adepithelial layer of the columnar epithelium. The adepithelial cells will give rise to the adult musculature; they also express *twist*, which controls *htl* (Shishido et al., 1997) and is a signature of all mesodermal cells and muscle precursors (reviewed in Bate, 1993). We confirmed that the wing disc adepithelial cells expressed both *Htl* (Figure 1D) and *Twist* (data not shown). In addition, we identified a small group of adepithelial cells that expressed a *btl* enhancer trap line but expressed neither *Htl* nor *Twist* (Figure 1D and data not shown). Stumps, a putative adapter protein required for both *Btl* and *Htl* signaling (Vincent et al., 1998), was expressed in both *btl*- and *Htl*-expressing adepithelial cells (Figure 1F).

The *btl*-expressing cells in the adepithelial layer had not been identified previously, and their presence was unexpected, since one of the principal domains of *btl* expression is the trachea (Klämbt et al., 1992). Tracheal cells are almost invariably associated with tubules with cuticle-lined lumen, and the adepithelial cells have no such distinct structures. Nevertheless, the adepithelial *btl*-positive cells appeared to maintain continuity with the cells of the main tracheal branch (Figures 1E and 1F). These results are summarized in Figures 1G–1I. We now present evidence that the *btl*-expressing adepithelial cells are the precursors of the adult tracheal air sacs.

**Origin of Air Sac Tracheoblasts**

To better understand the origin and fate of the *btl*-expressing adepithelial cells, we tracked them during larval and pupal development. In early third instar wing discs, no *btl* expression was detected in adepithelial cells; only tracheal branch cells were *btl* positive (Figure 2A). As third instar discs matured, however, *btl*-positive cells were detected budding from the tracheal branch that adheres to the wing disc. This bud formed at a stereotypical position just dorsal to the wing hinge progenitors and adjacent to the group of 15–60 *bnl*-expressing cells in the columnar epithelium (Figure 2B). During development of the third instar, the number of *btl*-expressing cells increased, and the bud expanded posteriorly toward the region of greatest *bnl* expression (Figures 2B–2H). In late third instar discs, the *btl*-expressing cells formed a coherent group surrounded by *Htl*-expressing cells. These *btl*-expressing cells did not express *Htl* (Figure 1D). Possible explanations for the complementarity of the patterns of *btl* and *Htl* expression are that *btl* cells displace *Htl*-expressing cells or that the expression of these genes is mutually exclusive. However, in the early third instar discs, *Htl* expression was already absent from the region that will be occupied by *btl*-expressing cells in older discs (Figures 2E and 2F); therefore, these are not likely to be sufficient explanations.

The number of *btl*-expressing cells populating the bud that originated from the tracheal branch increased during third instar (Figures 2E–2H). Since anti-phospho histone H3 antisera, which is specific to mitotic phase histone H3, stained nuclei among the *btl*-expressing cells (Figure 2I), we conclude that some, and perhaps all, of this increase in cell number is due to cell divisions. This duplicative capacity distinguishes these cells from cells of the tracheal branch, which do not proliferate in the postembryonic period (Manning and Krasnow, 1993; Samakovlis et al., 1996). Since the *btl*-expressing adepithelial cells also differ from tracheal cells by not forming a luminal cuticle, they are a distinct cell type.

To characterize these cells further, we examined their relationship to the larval tracheal system. The larval tracheal branch that adheres to the wing disc (Figures 1E and 1G–1I) is called the first transverse connective (Figure 3O). It has a small offshoot called the spiracular branch, where imaginal tracheoblasts, precursors of the adult trachea, are located. These imaginal tracheoblasts do not express *btl* (Samakovlis et al., 1996) but
Figure 1. Expression of FGF and FGF Receptors in Wing Imaginal Discs

Unless indicated otherwise, all discs in this and subsequent figures are late third instar and are oriented anterior left and ventral up. (A) btl (btl-Gal4 UAS-gapGFP, green) was expressed in tracheal cells and in adepithelial tracheoblasts. bnl-lacZ (light blue) was expressed in approximately 150 cells in the columnar epithelium. Different focal planes are merged in the left panel to show that bnl and btl were expressed in adjacent regions (arrows). bnl expression (right panel) was strongest among the more posterior cells. (B and C) bnl mRNA (arrows) detected by in situ hybridization in late (B) and early (C) third instar discs. Note that bnl mRNA was detected in a smaller area (approximately 80 cells in [B] and 15 in [C]) than bnl-lacZ (A). (D) btl-lacZ (green; left disc) was expressed in the tracheal branch, in a patch of adepithelial cells (arrow), and weakly in the wing pouch. Htl protein ([D] and [D’], red) was present in all adepithelial cells except those that express btl-lacZ (arrow). (E) GFP protein in btl-Gal4 UAS-gapGFP discs shows that tracheae (arrowhead) and tracheoblasts (arrow) are both btl-positive and contiguous but that tracheoblasts do not form a tracheal tube. (F) Stumps (red) was expressed in both btl-positive cells (arrow) and in the surrounding adepithelial (Htl positive; see Figure 1D) cells. btl-lacZ, green. (G–I) Schematic drawings showing FGF and FGF receptor expression in late third instar wing discs: TB, tracheoblasts (green spotted area); CE, columnar epithelium; AE, adepithelial cells; PM, peripodial cells; Tr, trachea; S, stalk. Scale bars are 100 μm (A–D) and 50 μm (E and F).

do express escargot (esg) (Samakovlis et al., 1996) and trachealless (trh) (Samakovlis et al., 1996; Ohshiro and Saigo, 1997) (Figures 2J and 2K, arrowheads). Since esg inhibits endoreplication of imaginal cells (Fuse et al., 1994) and the imaginal tracheoblasts are assumed to be diploid, expression of esg in these cells was not unexpected. The presence of Trh, a transcription factor that directly activates btl transcription (Ohshiro and
Figure 2. Tracheoblast Cell Dynamics

Migration of tracheoblasts in early (A and E), mid (B, C, F, and G), and late (D and H) third instar discs was revealed by expression of btl-Gal4 UAS-gapGFP ([A]–[D], green) and btl-lacZ ([E]–[H], green) in tracheoblasts and tracheae. bnl-lacZ expression was in columnar epithelial cells ([A]–[D], light blue). Htl protein was present in adepithelial cells (red) but was absent from the region of bnl-positive cells ([E] and [F], encircled areas).

(I) Proliferating cells stained with anti-phospho histone H3 antibody (red; arrows); btl-Gal4 UAS-gapGFP, green.

(J) Trh (red) and btl (green; btl-Gal4 UAS-GFP) were expressed in identical cells in tracheae and tracheoblasts (arrow). In the putative spiracular branch, Trh, but not btl, was expressed (arrowhead). Only Trh expression is shown in (J').

(K and L) esg-lacZ expression (light blue) was in a subset of tracheoblast cells at late third instar. Tracheae and tracheoblasts were visualized by btl-Gal4 UAS-gapGFP (green).

(K) esg-lacZ was hardly detectable in tracheoblasts until mid third instar (arrow). Note that esg was already expressed in the putative spiracular branch (arrowhead).

(L) At late third instar, esg was preferentially expressed at the tip of migrating tracheoblast nest (arrow). Note that esg was not expressed in proximal cells (arrowhead). Only esg expression is shown in the right panel (L'), which is a magnified view of an area enclosed by a white rectangle in the left panel. Scale bars are 50 μm (A–H and J–L) and 25 μm (I and L').

Saigo, 1997), may presage btl expression at a subsequent stage. The btl-expressing adepithelial cells also expressed Trh. They did not express esg early in the third instar; therefore, it is unlikely that they derive from the imaginal tracheoblasts. However, in mid third instar (Figure 2K and data not shown) and thereafter, esg expression was evident in the most posterior cells of the group (Figure 2L, arrow).

The capacity of tracheal branch cells to proliferate was unexpected because the cells lining the main tracheal branches had been considered to be both terminally differentiated and polyploid. Neither state is expected to be compatible with a mitotic cell cycle program. We examined the morphology and fluorescence of DAPI-stained discs to better understand the nature of the disc-associated tracheal cells. We observed that most of the cells that line the lumen of the main tracheal branch, the first transverse connective,
Figure 3. Tracheoblast Development in the Pupa

(A–E) In discs from late third instar to 12 hr APF, GFP protein (btl-Gal4 UAS-gapGFP; black; arrows) in tracheoblasts was in the posterior region between the notum and wing hinge. Dorsal, up; times, hours after puparium formation (APF). Wing hinge (arrowheads) was identified by wingless-lacZ (brown staining).

(F–L) During pupal development, tracheoblasts (btl-Gal4 UAS-GFP; green) grew, migrated dorsally, and generated three branches of thoracic air sacs: m, mediocutal sac; l, laterocutal sac; s, scutellar sac. Time-lapse images of a live pupa 12–32 hr APF (F–J) were taken on a stereo dissecting microscope. The wing blade, including the hinge region, is outlined by white dots. Tracheoblasts (arrows) migrated dorsally from 12 to 22 hr APF (F–H) and generated new branches 26–32 hr APF (I and J).

(F–K) Lateral view of pupal thorax; dorsal is up; (L–S) are dorsal views; anterior is up.

(K and L) Tracheoblasts did not migrate extensively after 32 hr APF but elaborated to become thoracic air sacs about 3 days APF (K). They are very similar to air sacs of black pupae (L). Pupal cases were removed in (K) and (L). Note that mediocutal sacs are positioned between dorsocentral macrochaetae (asterisks in [L]).

(M–O) Dissected and fixed air sacs of a black pupa. Despite sac-like appearance (M), tracheal bundles were found in and around air sacs (N and O).

(N and O) Magnified views of areas enclosed by rectangles in (M).

(P) Pupal tracheae (arrowhead) extended to developing flight muscle at 12 hr APF. They originated from dorsal branch 2 (see Figure 3R in which arrowheads also indicate pupal tracheae). Note that tracheoblasts (arrows in [P] and [R]) were found laterally at this stage (see also Figure 3P).

(Q–S) Schematic drawings of the formation of thoracic air sacs. Larva (Q), pupa (R), and adult (S) are shown. Tracheoblasts and air sacs, red; spiracular branches (SB), green; both air sac tracheoblasts and spiracular branches connect to transverse connectives (TC), which extend from dorsal trunks (DT) to lateral trunks (LT). Dorsal branches (DB) connect dorsal trunks. Bundles of tracheoles along dorsal branch 2 in (R) are pupal tracheae, which extend to the developing flight muscle prior to air sac formation (see Figure 3P). Scale bars are 100 μm (A–E and M), 162 μm (F–L and P), 10 μm (N), and 25 μm (O).
as well as the imaginal tracheoblasts, have nuclei that are similar in diameter (5–6 μm), are relatively small, and have a similar level of fluorescence. Some cells with large nuclei and >3× the fluorescence intensity of the smaller nuclei were observed populating other branches that connect to the first transverse connective. Assuming that the imaginal tracheoblasts are diploid, these observations suggest that the cells that respond to Bnl-FGF have a similar ploidy and that the tracheal branches that associate with the wing disc include both diploid and polyploid cells.

Development of Air Sac Tracheoblasts
To determine the fate of the btl-positive adepithelial cells, we examined their behavior and movements throughout pupal development. As shown above (Figure 1), they are located next to the posterior part of the prospective wing hinge in the late third instar disc. Analysis of fixed wing discs prepared at various times during the first 12 hr after puparium formation (APF) revealed that they remained in this location as a tight, rounded cluster of cells next to the prospective wing hinge (APF; Figures 3A–3E, arrows).

To continue following the fate of these cells, five individual pupae that expressed btl-Gal4 UAS-GFP were observed during the pupal period, and photographs were taken at regular intervals. The pupal case is transparent to the GFP fluorescence; therefore, no surgical manipulations were necessary. Figures 3F–3K show lateral views of a pupa that expresses GFP under btl control. The btl-positive adepithelial cells were identified by their proximity to the wing hinge (dotted line) at 12 hr APF (Figure 3F), consistent with the observations of dissected, fixed discs (above). These btl-positive cells migrated dorsally between 12 and 23 hr APF then anteriorly and posteriorly to form three branches (Figures 3F–3J). At 32 hr APF, they ceased their migrations and began to elaborate into air sacs (Figures 3K and 3L).

We conclude that the btl-positive adepithelial cells are the precursors of the adult air sacs and that the air sacs of the adult thorax are derived from cells that are distinct from the imaginal tracheoblasts.

Although the air sac tracheoblasts did not form a tubular structure at 12 hr APF (Figure 3E and data not shown), these pupae did have numerous tracheae that projected to the developing muscles (Figure 3P). These tracheae are clearly distinct from the air sac tracheoblasts. They derive from the second dorsal branch (Figure 3R; DB2) and are present only during the pupal period (reviewed in Manning and Krasnow, 1993). Air sacs are prominent and extensive in older pupae and in adults (Figures 3L and 3S) and are associated with numerous bundles of tracheal that extend from the air sacs and extensively interdigitate with flight muscle cells (Figures 3M–3O). We do not, at present, understand enough about the structure of the air sacs to know how these tracheae connect with, or contribute to the function of, air sacs.

Multiple Roles of FGF Signaling
During embryonic development, Bnl-FGF is a guidance cue for tracheal branching and migration (Sutherland et al., 1996). btl-expressing tracheal tips branch out and migrate toward bnl expression to elaborate the fine structure of the embryonic tracheal system. Similarly, migration of the air sac tracheoblasts in the third instar wing disc appears to be directed to the region of bnl expression (Figures 2A–2H). To test whether Bnl-FGF is a guidance cue for air sac tracheoblasts, bnl was ectopically expressed under the control of dpp-Gal4. btl-expressing adepithelial cells migrated over the wing pouch along the region of ectopic expression (Figures 4A–4C). Like the air sac tracheoblasts, these cells lacked a tubular structure (Figure 4C) but were Stump positive (data not shown).

We then sought to determine whether Bnl-FGF could induce tracheoblasts to bud from the larval trachea at sites separate from the region in which bnl is normally expressed. We generated clones of cells expressing bnl in the columnar epithelial disc cell layer (Ito et al., 1997) and assayed tracheoblasts by the presence of Trh and btl. Trh-, btl-expressing cells migrated toward bnl-expressing clones that were generated at various locations in this disc. These migratory cells formed ectopic branches of tracheoblasts (Figures 4E and 6) and arose even at significant distances from the endogenous branch (Figure 4E). In dppGal4 UAS-bnl discs, esg-expressing cells were present in the most posterior aspect of the ectopic tracheoblast branch, but esg expression was negligible in more-anterior cells and in tracheal cells (Figure 4D). This suggests that esg-positive cells found in ectopic tracheoblast branches did not derive from endogenous tracheoblasts and that esg was activated de novo in response to bnl. The growth of these ectopic branches indicates that, in the wing disc, Bnl-FGF is sufficient to induce tracheoblast migration and that cells that are competent to respond to Bnl-FGF are not uniquely positioned at the site where air sac tracheoblasts normally arise. Rather, such Bnl-FGF-responsive cells are distributed more broadly. As discussed below (see Discussion), this suggests that Bnl-FGF may function locally in the third instar disc to reprogram tracheal cells.

To determine whether FGF signaling is necessary for tracheoblast migration, Bnl-FGF and Btl were inactivated. First, large bnl mutant clones were induced in the columnar epithelial cell layer. Tracheoblast migration was significantly reduced in discs with large bnl clones (Figure 4F). Second, dominant-negative (BtlDN) and constitutively active (Btl) forms of Btl were expressed under the control of btl-Gal4. Ectopic expression of UAS-btlDN significantly reduced migration of tracheoblasts (Figure 4G). This phenotype is similar to bnl loss of function (see Figure 4F). Ectopic expression of UAS-btl caused a significant increase in the number of tracheoblast cells, but, unlike ectopic bnl expression (Figures 4B–4E), UAS-btl did not cause abnormal branching (Figure 4H). We suggest that spatially regulated Btl activity is required to induce cell migration and proliferation (see Lee et al., 1996), and we conclude that, in the wing disc, Bnl-FGF is both necessary and sufficient for directed tracheoblast migration.

In the experiments described above, the number of tracheoblast cells significantly increased in response to ectopic Bnl-FGF/Btl signaling (Figures 4B–4E and 4H). In contrast, reduction of Bnl-FGF/Btl signaling resulted in fewer or no tracheoblasts (Figures 4F and 4G). Thus, Bnl-FGF/Btl signaling appears to activate tracheoblast
Figure 4. Role of FGF in Tracheoblast Maturation

(A) Location of tracheoblast nest in wild-type (btl-lacZ; green).
(B–D) UAS-bnl/+; dpp-Gal4 UAS-GFP/+; btl-lacZ/+ expressed bnl (light blue) along the anterior/posterior compartment border. btl-lacZ-positive tracheoblasts (green; arrows) track along area of ectopic bnl expression. Expression of dpp-Gal4 in the notum was present along the posterior edge (arrowhead in [B]).
(C) A magnified DIC image of the wing pouch. Left panel, DIC image isolated. Note that ectopic tracheoblasts are morphologically distinct from tracheal cells and are attached to the basal surface of columnar epithelial cells (arrows).
(D) esg-lacZ expression (light blue) was activated by ectopic bnl (arrows). Trh protein, red.
(E) Ectopic bnl expression (AyGal4; UAS-bnl UAS-GFP; light blue; arrow) in dorsal-most notum induced new tracheoblast branches. Trh protein (red).
(F) Tracheoblast migration stalled in discs with large bnl loss-of-function clones (arrow). bnlP1 clones in the columnar epithelium were marked by the absence of GFP (light blue).
(G) Dominant-negative Btl (btl-Gal4; UAS-btl DN UAS-gapGFP; green) inhibited tracheoblast formation and/or migration. Trh protein (red). Few or no migrating tracheoblasts were observed (arrow).
(H) Constitutively active Btl (btl-Gal4; UAS-Btl UAS-gapGFP; green) increased the number of tracheoblasts. Trh protein, red. Cell migration was not significantly affected (arrow). Scale bars are 100 μm (A and B), 20 μm (C), and 50 μm (D–H).

proliferation. To substantiate this possibility, we counted the number of anti-phospho histone H3-positive cells after a brief induction of bnl, λbtl, and btlKR. Compared with control (2.44 cells, n = 18), bnl and λbtl expression stimulated proliferation (5.17 cells, n = 18 and 3.05 cells, n = 22). Fewer proliferating cells were observed after btlKR expression (0.77 cells, n = 13).

Air Sac Tracheoblasts Extend Actin-Based Filopodia

To better understand the process that induces tracheal cells to migrate and proliferate as air sac tracheoblasts, we examined these cells in isolated discs using fluorescent probes. Gap-GFP was expressed in btl-containing adepithelial cells and was observed to delineate numerous filopodia with a cytoneme-like appearance (Ramirez-Weber and Kornberg, 1999) extending in the direction in which these cells migrate (Figures 5A and 5B). Filopodia were occasionally observed extending in other directions, but these nonoriented filopodia were less numerous and significantly shorter than the oriented ones, which extended up to 50 μm. The morphology of the extensions was not preserved after paraformaldehyde or methanol fixation (data not shown).

We investigated the fine structure and composition of the filopodial extensions by characterizing the distribution of several different GFP fusion proteins. We noted that the fluorescence of Gap-GFP was nonuniform and included bright spots along the length of the extensions (Figure 5B). We also expressed GFP derivatives of actin (Verkhusha et al., 1999), Tau (Jarecki et al., 1999), which binds microtubules, and Btl. Actin-GFP illuminated the extensions uniformly (Figure 5D). The fluorescence observed with Tau-GFP illuminated only spike-like short protrusions (Figure 5E). Btl-GFP fluorescence delineated the extensions and concentrated in bright spots along their length and at their apparent termini (Figure 5F). We conclude that tracheoblasts have mainly actin-based filopodia whose tips or shafts may have the capability to sense a Bnl-FGF ligand.

Bnl-FGF enhances or regulates the filopodia of air sac tracheoblasts. As described in the previous section, this conclusion is based on the effects of Bnl-FGF, the dominant-negative Btl protein (BtlDN), and the constitutively active Btl protein (λBtl). To correlate the relationship
Figure 5. Actin-Based Filopodia at the Tip of Tracheoblast Nest and the Involvement of FGF Signaling

Filopodia were observed in nonfixed discs prepared in hanging drops (A, F, H, J, K, M, and N) or under coverslips (B, D, and E). btl-Gal4 UAS-gapGFP was used to visualize filopodia in (A), (B), (G), (H), (J), (K), (M), and (N). MAP kinase activation in tracheoblasts was observed by double labeling with Δ-MAPK antibody (red) and btl-Gal4 UAS-gapGFP (green) in (C), (I), (L), and (O).

(A and B) Filopodia produced by btl-expressing cells oriented toward putative bnl-expressing cells (large arrows). A greater number of filopodia were observed in flattened preparations under coverslips (B) than in hanging drops (A). Short filopodia were infrequently observed with nonspecific orientations (arrowheads). Small punctate structures were found along filopodia (small arrows).

(C) MAPK activation at the tip of the tracheoblast nest (arrow) was greater than in the more proximal regions (arrowhead).

(D) Filopodia were labeled by actin-GFP fusion protein (btl-Gal4 UAS-GFPact; arrows).

(E) Filopodia were not effectively labeled by Tau-GFP fusion protein (btl-Gal4 UAS-tauGFP; arrow).

(F) Filopodia (large arrow) and small punctate structures (small arrows) contained Btl-GFP fusion protein (btl-Gal4 UAS-btlGFP).

(G) Dominant-negative Btl inhibited filopodia formation (hs-btlKR) as well as MAPK activation 3 hr after heat shock (I, arrow), but, after 24 hr recovery from heat shock, filopodia returned to their normal distribution (H, arrow). Constitutively active Btl (hs-λbtl) did not increase the number of filopodia (arrows) 5 hr (J) or 24 hr (K) after heat shock. Note that filopodia in (J) appeared to be shorter than normal. MAPK activation was enhanced and present in proximal cells (arrowhead) 3 hr after heat shock (L). Ubiquitous bnl expression induced ectopic, nonoriented filopodia (M) 5 hr after heat shock induction of hs-bnl. Twenty-four hours after heat shock, both the number of btl-expressing cells and nonoriented filopodia were increased (N). MAPK was activated in proximal cells 3 hrs after heat shock (O). Scale bars are 30 μm (A–C and G–O), 12 μm (D), 20 μm (E), and 16 μm (F).

between FGF signaling and filopodia, Btl<sup>KR</sup> and λBtl were expressed ubiquitously, and the number and distribution of filopodia, as well as the activity of MAPK, were assessed. MAPK is known to be activated by Bnl-FGF signaling (Gabay et al., 1997), and, in wild type, it was activated in those air sac tracheoblasts that are closest to the bnl-expressing columnar epithelial cells (Figure 5O). Three hours after induction of btl<sup>GR</sup>, MAPK activation
in tracheoblasts was completely abolished (Figure 5I). In contrast, induction of \(btl\) and \(bnl\) caused MAPK activation throughout tracheoblast and tracheal cells (Figures 5L and 5O).

In discs in which Gap-GFP was expressed in \(btl\)-positive tracheoblasts, a brief pulse of \(Btl^{cr}\) expression caused the disappearance of all filopodia within 3–5 hr (Figure 5G and data not shown). Their absence was temporary, as, without further induction of \(Btl^{cr}\), many filopodia were once again present 24 hr later (Figure 5H). In contrast, ubiquitous expression of \(bnl\) caused a large increase in the number of filopodia, and, unlike in the wild type, these filopodia extended from the \(btl\)-positive tracheoblasts in every direction (Figures 5M and 5N). The number of \(btl\)-expressing cells also increased, probably due to stimulated cell proliferation and migration (Figure 5N).

Together, these observations suggest that \(Bnl-FGF/\)
\(Btl\) signaling activates filopodia formation in the air sac tracheoblasts. Interestingly, ubiquitous expression of \(\lambda Btl\), a constitutively active form of the \(Bnl-FGF\) receptor, did not activate filopodia formation. Rather, shorter filopodia were observed (Figures 5J and 5K). A possible interpretation is that regulated \(Btl\) activity is necessary for filopodia formation. Under our experimental conditions, \(\lambda Btl\) would be expected to be dimerized and activated uniformly on the surface of expressing cells, and this condition may not lead to filopodia formation. Another possibility is that \(\lambda Btl\) does not activate all aspects of \(Btl\) signaling. It may be relevant that regulated \(Btl\) activity is also essential for normal tracheal migration in embryos (Lee et al., 1996).

**Cytoneme-like Filopodia Are Attracted to \(bnl\)-Expressing Cells**

As described above, ectopic expression of \(Bnl-FGF\) in cell clones induces novel regions of \(btl\)-positive tracheoblasts. We sought to determine whether these ectopic sites of tracheoblast migration and cell proliferation are also associated with filopodial extensions. We designed a “flip-out” construct to mark clones that express \(bnl\) in discs in which \(btl\)-expressing cells contain GFP (Figure 6A). We found that ectopic filopodia extended towards clones expressing \(bnl\) (Figures 6B and 6C). Clones were a distance of several cell diameters from the tracheoblasts and in an adjacent cell layer. Interestingly, the ectopic filopodia did not extend in all directions. For example, in the clone illustrated in Figure 6C, they extended dorsally and posteriorly toward the clone of \(bnl\)-expressing cells, but not ventrally away from the clone. We conclude that, in the wing disc, Bnl-FGF is both necessary and sufficient to induce oriented filopodia formation in tracheoblast cells.

Small punctate structures with a diameter of approximately 1.2 \(\mu m\) were occasionally found along filopodia very close to \(bnl\)-expressing clones (Figures 6D–6F). Note that their distribution coincides with the shape and location of clones. Considering their close proximity to \(bnl\) expression and the presence of \(Btl\) in filopodia (see Figure 5F), their formation may relate to the reception of Bnl by filopodia. Similar structures were found in the filopodia extending from normal air sac tracheoblasts (Figure 5B).

**Discussion**

The wing imaginal disc produces a variety of structures and cell types. These include the many different cuticular elements, the organs of the peripheral nervous system, and muscles (reviewed in Bate, 1993; Cohen, 1993). Our studies add the tracheal air sacs to this list. In studies of the role of FGF in the wing disc, we detected \(bnl\) expression in a small group of columnar epithelial cells during the third instar and pupal periods. Although we did not establish their fate in the adult, these cells are small in number and therefore cannot produce more than a small part of the adult cuticle. Nevertheless, their effect on the adult is profound. Through the action of Bnl-FGF, they induce a group of tracheal cells to initiate a program of proliferation and migration and to join with the cells in the disc adip epithelial layer. Despite this intimate association with these mesodermal progenitors, the FGF-responsive cells retain their tracheal identity and go on to form the prominent adult tracheal air sacs that extend throughout much of the dorsal thorax.

**Genesis of the Tracheal System in the Dorsal Mesothorax of the Adult**

The tracheal system of the *Drosophila* embryo has ten interconnected metameric units on either side of the animal; one derives from the second, or mesothoracic, segment. This mesothoracic component consists of portions of the dorsal and lateral trunks, a transverse connective that links these trunks to each other, a dorsal branch that connects the left and right sides, and numerous branches that radiate out to various tissues. During larval development, this general structure is retained, and, although the tracheal cells do not divide, many new branches form, and the diameter of the more proximal tracheae increase (reviewed in Manning and Krasnow, 1993; Samakovlis et al., 1996; Beitel and Krasnow, 2000). The wing imaginal disc attaches to the transverse connective. Imaginal tracheoblast precursors of the adult tracheae populate a small spiracular branch at a location just dorsal to the disc attachment.

In constructing the adult tracheal system, the imaginal tracheoblasts use parts of the larval framework as templates and, in effect, remodel the dorsal and lateral trunks, the transverse connective, the dorsal branch, and the main pupal branches to the wing and leg (reviewed in Manning and Krasnow, 1993). In contrast, the large and extensive air sacs do not correspond to earlier branches in any obvious way and have no apparent antecedent. We have shown that the airs sacs of the dorsal thorax form de novo from a small group of wing imaginal disc cells. We chronicled the transformation of these air sac tracheoblasts from a tight cluster of adip epithelial disc cells to sculpted air sacs. We made these observations by expressing GFP under \(btl\) control and by following the GFP-containing cells through the pupal period. We could directly account for three branches of the dorsal thoracic air sacs (e.g. the mediascutal, lateroscutal, and scutellar sacs) as products of the wing disc air sac tracheoblasts. It was our impression that all of the air sacs in the notum contained GFP in these animals, but the resolution of our analysis was not sufficient to make this a definitive conclusion nor
did it allow us to conclude that all of the air sacs derive from disc tracheoblasts. Nevertheless, our study did establish that the disc tracheoblasts generate air sacs and, by some process that is as yet unknown, form a tracheal lumen and tracheal network. It will be interesting to identify the intrinsic and extrinsic systems that direct the genesis of the air sacs, since they apparently develop in the absence of a preexisting framework.

**Imaginal Disc Tracheoblasts, a General Mechanism for Air Sac Development**

We have presented evidence for air sac tracheoblasts in the wing imaginal disc. Data not shown suggest that there may be similar strategies to make air sacs in other regions of the fly. We base this statement on the presence of nontracheal btl-positive cells and bnl-expressing cells in other imaginal discs. In leg discs, bnl expression was found in the stalk region, and the pattern of expression became more extensive and complex in the disc epithelium of pupal discs. In third instar discs, btl-positive cells were localized to an offshoot of a tracheal branch that attaches near the stalk and adjacent to the bnl-expressing cells. After puparium formation, the btl-positive cells migrated along the basal surface of the disc columnar epithelium to a position that roughly correlates with the region where air sacs will later form (reviewed in Manning and Krasnow, 1993). In eye-antenna discs, bnl was expressed in cells surrounding the ocelli progenitors. Although we did not find btl-expressing cells in larval eye-antenna discs, we did observe that, in early pupae, btl-positive cells assumed a position underlying the presumptive ocelli cells. Air sacs in the adult head underlie most of the medial head cuticle and encircle the region where the ocelli form. These observations lead us to suggest that the process that induces the air sac tracheoblasts in the wing disc may be common to other discs as well.

**Bnl-FGF and the Genesis of Air Sacs**

Bnl-FGF is the key determinant of tracheal branching as the preadult tracheal system matures. In the embryo, it guides the migration of tracheal cells to form primary branches, induces secondary branches as the primary
branches approach the cells expressing bnl, and regulates the process that generates terminal branches. To produce these different outcomes, the FGF signalling pathway acts through different, but related, mechanisms, but its role is, in effect, a single one—to mold the tracheal cells and to influence where and how they extend. The roles that it plays during the early stages of air sac morphogenesis are different. As a signal in the wing imaginal disc, Bnl-FGF functions as a chemoattractant, inducing these cells to migrate from outside the imaginal disc to a location within the epidermal layer. And it acts as a mitogen, inducing the tracheoblasts to proliferate. These properties were not manifested by Bnl-FGF during earlier stages of tracheal development. Thus, the morphogenic process that generates the air sacs is distinguished both by its independence from the tracheal framework of the embryo and larva and by the roles that FGF plays.

Perhaps the most surprising behavior this work identified is the response of the larval tracheal cells to Bnl-FGF. These cells form a tripartite tube that consists of an external basal lamina, a squamous epithelium, and a complex, multilayered luminal cuticle. They had been considered to be terminally differentiated and incapable of proliferation, having ceased cell division in the early embryo. We found, instead, that many of the wing disc-associated tracheal cells can respond to Bnl-FGF by migrating out of the tracheal branch to embark on a program of proliferation and morphogenesis. Remarkably, this latent capacity for conversion to proliferative tracheoblast is shared by many (and perhaps all) of the tracheal cells that populate the disc-associated branch. These cells require only the action of Bnl-FGF to initiate the process. Although we cannot exclude the possibility of a separate and distributed population of tracheal stem cells, we feel that the widespread capacity of tracheal cells to adopt a program of proliferation and migration makes this possibility unlikely. Instead, we propose that Bnl-FGF acts as an instructional determinant, reprogramming the cells in the tracheal branch to become air sac tracheoblasts. To our knowledge, the activity of Drosophila FGF to induce cells to dedifferentiate has no precedent.

FGF Signalling and Cytoneme-like Filopodia

Actin-based filopodia have been observed in many cells that send or receive signals (reviewed in Ramirez-Weber and Kornberg, 2000). For purposes of illustration, we mention just four here. (1) Neuronal growth cones are populated with many active filopodia that appear to probe their environment for guidance cues (reviewed in Kater and Rehder, 1995). (2) Long and highly dynamic filopodia extend from primary mesenchyme cells in the interior of early sea urchin embryos, apparently to contact and explore the overlying ectoderm. They are thought to relay information from the peripheral ectoderm that patterns internal skeletal elements (reviewed in McClay, 1999). (3) Dendritic cells, professional antigen-presenting cells of the vertebrate immune system, are defined in part by their Medussa-like morphology. Their unusually large number of finger-like projections may maximize the likelihood that antigen presentation finds a suitable target cell (Raghunathan et al., 2001). (4) Drosophila wing imaginal disc cells have thin filopodial extensions, “cytonemes,” that appear to connect cells with the organizer region at the anterior/posterior compartment border (Ramirez-Weber and Kornberg, 1999). Although the filopodia produced by these various cell types are similar in gross morphology, it is not known whether they all have a role in trafficking signals, have a common mechanism that receives and transduces signals, or are regulated in a similar manner.

The filopodia that the wing disc tracheoblasts produce have a number of properties in common with cytonemes. Both are actin-based, have a comparable size and appearance, and are similarly sensitive to standard conditions of fixation. Functional analysis indicates that the tracheoblast filopodia are dependent on Bnl-FGF and upon the ability of the tracheoblasts to carry out FGF signal transduction. We do not have direct evidence that signalling cannot occur if they are absent, but the correlation between these structures and active signalling is strong. Moreover, their presence offers a possible mechanism to move the FGF signal from its source in the columnar epithelial cell layer across the epithelial cell layer to the tracheal target cells. We do not know what alerts the target cells to the presence of a source of FGF. Two possibilities are that nonspecific cytoneme-like filopodia explore the extracellular environment for potential sources (see Figures 5A and 5B) or that less efficient signalling can occur in the absence of direct contacts. We would assume that, if this second possible mechanism is operative, once filopodia are induced and make appropriate contacts, signalling will be accelerated and more productive. The question of how the filopodia are induced and oriented is certainly important, but the very presence of these cellular extensions that make contact across these distances offers a mechanism to facilitate the ordered movement of signals and the concerted and directed migration of cells.

Experimental Procedures

Plasmid Construction

**UAS-btlGFP**

The C terminus of *btl* was fused to *GFP<sup>act</sup>* and inserted into pUAST. The stop codon of *btl* was mutated with a NotI site, Sall and KpnI sites were added to the N and C termini, respectively, of GFP, and the two genes were joined and inserted between the NotI and KpnI sites of pUAST.

**hs-bnl**

The HindIII-BglII fragment of the *bnl* cDNA fragment was inserted into HpaI/BglII-digested pCaSpeR-hs after the HindIII site had been repaired with Klenow enzyme.

**tubulin**<sup>-DsRed2</sup>, y<sup>-btl</sup>, y<sup>-bnl</sup>

The *hedgehog* cDNA fragment of pKB798 (Zecca et al., 1995) was removed and replaced with a KpnI-XbaI fragment containing the full-length *bnl* cDNA (pKBBnl). NotI and XbaI sites were added to the N and C termini, respectively, of DsRed2 cDNA (Clontech) by PCR. DsRed2 was inserted into pUAST using the NotI and XbaI sites to generate pUAS-DsRed2. An EcoRI-BamHI fragment containing DsRed2 cDNA and an SV40 polyA signal was inserted into pBlue- script. Using pBlue-script as a shuttle vector, a NotI site was added just downstream of the SV40 polyA signal. The NotI fragment containing DsRed2 and SV40 polyA signal was replaced with the CD2 cDNA fragment of FC15 (Zecca et al., 1995) to generate a flip-out cassette containing DsRed2 and yellow<sup>+</sup> between two FRT sites. The flip-out cassette was inserted into the unique KpnI site of...
pKBbnl between the tubulin promoter and the bnl cDNA. Trans- 
formants were identified by rescue of the yellow mutant phenotype. 
pKB798 and FC15 were provided by K. Basler.

Fly Strains
Enhancer trap strains btl(zecl) (Klämbt et al., 1992), bnl(zecl) (Sutherland et al., 1994), wp5G (Sato et al., 1998a), and escl(zecl) (Samakovlis et al., 1996) were used to visualise btl, bnl, wp5, and escl expression, respectively. dpp-Gal4 (kindly provided by Y. Jan) and btI-Gal4 (Shiga et al., 1996) were used to drive ectopic gene expression (Brand and Perrimon, 1993). UAS-GFPcret (UAS-GFP; Fly Base), UAS-gapGFP (Ritzenthaler et al., 2000), UAS-GFPact (Verkruysse et al., 1999), UAS-lauGFP (Jarecki et al., 1999), and UAS-btlGFP (this work) were used to express various forms of GFP, Bnl/FGF ligand, dominant-negative Btl, and constitutively active Btl were expressed using UAS-bnl (Sutherland et al., 1996), UAS-btl(zecl) (Reichman-Fried and Shilo, 1995), or UAS-btl (Lee et al., 1996), respectively, or by using hs-bnl (this work), hs-btl(zecl), or hs-btl (Lee et al., 1996). btl(zecl) encodes a dominant-negative Btl receptor without functional cytoplastic domain (Reichman-Fried and Shilo, 1995). btl encodes a constitutively active form of Btl with an extracellular domain replaced by the dimerization domain of λ repressor (Lee et al., 1996). hs-flp (Zeca et al., 1995), AyGal4 (Tito et al., 1997), UAS-GFP, and UAS-bnl were used to induce clones expressing bnl marked with GFP, hs-flp, btl(zecl) FRT82 (Sutherland et al., 1996), and ubi-nlsGFP M(3R)6w FRT2B (Fly Base) were used to generate large bnl mutant clones marked by the absence of GFP. hs-flp and tubulin–DsRed2, y > bnl (this work) were used to generate clones expressing bnl marked by the absence of DsRed2. Germine transformation was performed as described in Bartoszewski and Gibson (1994).

Histology
Immunostaining and in situ hybridization were performed as described previously (Sato and Saijo, 2000), except that imaginal discs were fixed for 30 minutes in 8% parafomaldehyde for anti-dpMAPK staining. The following primary antibodies were used: mouse anti-lacZ (40-1a; Developmental Studies Hybridoma Bank), rabbit anti-lacZ (Cappel), mouse anti-GFP (B34; provided by P. O’Farrell), rabbit anti-Stumps (provided by M. Leptin), rabbit anti-Htl (provided by K. Saijo), rat anti-Trh (provided by S. Crews), mouse anti-dpMAPK (Sigma), rabbit anti-phospho-Histone H3 (Upstate Biotech), and anti-Digoxigenin-AP (Boehringer Mannheim). Secondary antibodies used are anti-mouse Cy3, anti-rabbit Cy3, anti-rat Cy3, anti-rabbit FITC, anti-mouse Biotin, and anti-rabbit Biotin (Jackson). ABC kit (Vector) was used for DAB staining.

Fluorescence intensities of nuclei were quantified after staining fixed discs with 1 μg/ml DAPI for 15 min and the collection of a 3D stack of images using a Zeiss fluorescence microscope. Image intensities of individual nuclei were summed using the EditPolygon program of Z. Kam (personal communication).

Numbers of proliferating cells were estimated by fixing wing discs of late third instar larvae and staining with anti-phospho-Histone H3 after heat shocking hs-bnl, hs-btl, or hs-btl(zecl) for 60 min at 37°C followed by 3 hr incubation at 25°C. The number of dividing cells in the tracheoblast branch, which was marked by btI-GAL4 UAS-gapGFP, were counted.

Mounting Unfixed Imaginal Discs
When placed under coverslips, imaginal discs flattened, and the appearance of their filopodia was highly variable. Better preparations were achieved by first gently adhering a disc to a coverslip by allowing it to settle for several minutes in a drop of PBS and then suspending the disc by mounting the coverslip in an inverted depression slide. Halocarbon B. Shilo for antisera, fly strains, and plasmids, the Developmental Studies Hybridoma Bank for anti-sera, and the Bloomington Stock Center for strains. This work was supported by a Research Fellowship of the Japan Society for the Promotion of Science for Young Investigators (to M.S.) and by a grant from the National Institutes of Health (to T.B.K.).

Received: March 3, 2002
Revised: May 22, 2002

References
Klämbt, C., Glazer, L., and Shilo, B.Z. (1992), breathless, a Drosophila FGF receptor homolog, is essential for migration of tracheal and specific midline glial cells. Genes Dev. 6, 1668–1678.

Acknowledgments
We thank J. Gu for technical assistance, Z. Kam, J. Sedat, and J. Vazquez for help in interpreting the DNA content of nuclei, M. Krasnow, G. Martin, G. Schubiger, and members of the Kornberg lab for helpful discussions and comments on the manuscript, K. Basler, S. Crews, K. Hill, C. Hosono, Y.N. Jan, M. Krasnow, M. Leptin, R. Matsuda, D. Montel, K. Saijo, and B. Shilo for antisera, fly strains, and plasmids, the Developmental Studies Hybridoma Bank for anti-sera, and the Bloomington Stock Center for strains. This work was supported by a Research Fellowship of the Japan Society for the Promotion of Science for Young Investigators (to M.S.) and by a grant from the National Institutes of Health (to T.B.K.).


