Genetic analysis of the larval optic nerve projection in *Drosophila*

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**SUMMARY**

The *Drosophila* larval optic nerve, called Bolwig’s nerve (BN), projects into the central brain along a simple invariant path. The growth of the BN proceeds in three phases, during which the nerve changes direction at two intermediate targets, P1 and P2. Here we show that the projection of the BN is amenable to genetic dissection. In a mutagenesis screen, we have isolated mutations in 13 genes that disrupt the BN projection in distinct phases of its development. The mutant phenotypes in combination with the expression patterns of corresponding candidate genes define cellular components necessary for directing the growth of the BN toward P2 and for redirecting its growth at P2, and reveal developmental strategies employed in the establishment of the BN projection.

Key words: *Drosophila*, optic nerve, Bolwig’s nerve, genetic analysis

**INTRODUCTION**

How neurons establish connections with their often distant targets is one of the fundamental questions of developmental neurobiology (Dodd and Jessell, 1988; Goodman and Shatz, 1993). Most often, it is the axon that navigates this distance through directed growth at its tip. In some cases, however, the cell bodies themselves migrate toward their final location after axonal contacts have first been established (Simon and Lumsden, 1993). Navigation of axons – and possibly the migration of cell bodies – is widely believed to be steered by the spatial distribution of attractive and repulsive cues and by selective adhesion. The nature of these cues is diverse: gradients of diffusible chemicals, local extracellular signals and cell-associated cues (reviewed in Goodman, 1996). Cell-associated cues include guidepost cells, which may represent undifferentiated neurons or glial cells, as well as tracts of other fibers (Bentley and Keshishian, 1982).

In recent years, several families of molecules have been identified that are involved in axonal guidance, including Netrins (Serafini et al., 1994; Kennedy et al., 1994), Semaphorins (Kolodkin et al., 1993; Lou et al., 1993), Eph receptors and their ligands (Cheng et al., 1995; Drescher et al., 1995), and adhesion molecules (CAMs, Cadherins; Grenningloh et al., 1993; Grumet, 1991). While much has been learned about the molecular components involved in the establishment of neuronal connectivity, we are still far from a mechanistic understanding of even simple pathfinding paradigms in vivo. An important prerequisite for testing the central tenets of axon guidance is a simple experimental system, namely an invariant path of axonal growth and easily identifiable intermediate targets at which axons change the direction of their growth. Previously described axon guidance models in the *Drosophila* embryo include the CNS (Seeger, 1994), the motor neuron projection (VanVactor et al., 1993; Bate and Broadie, 1995) and the PNS (Salzberg et al., 1994; Kolodziej et al., 1995). Here we would like to advocate the larval visual system of *Drosophila* as a simple model to study the cellular and the molecular mechanisms of axonal growth and pathfinding using a combination of cellular, genetic and molecular approaches. The larval visual system offers the opportunity to study the projection of a small group of photoreceptor neurons both with respect to their pathfinding and their fasciculation into a nerve. The availability of an early and specific marker for the larval visual system furthermore permits study of these processes selectively and at very high resolution.

The main photoreceptive organs of the larval visual system of flies have first been described for *Musca domestica* by N. Bolwig (Bolwig, 1946) and are therefore referred to as Bolwig’s organs (BO) (Steller et al., 1987). The BOs are a pair of photoreceptive organs, which are located bilaterally in a dorso-medial position in the head of the *Drosophila* larva. Each BO consists of only 12 photoreceptors, unipolar neurons, which project ipsilaterally along an invariant pathway via a single optic nerve, the Bolwig nerve (BN). The BN extends posteriorly from the BO to the ipsilateral brain hemisphere, enters the brain in the vicinity of the optic lobe anlagen (OLA), the primordia of the adult optic lobes, and grows toward a small target area in the central brain (Fig. 1).

The embryonic development of the BO and the OLA are correlated both spatially and temporally. The two structures arise from the same location in the ectoderm of the embryonic head region (Green et al., 1993) and the projections from the BO contact the OLA early in development, while the two cell groups are in close proximity to one another. These initial axonal projections are thought to be maintained during the morphogenetic separation of BO and OLA due to the incorporation of the OLA into the ventral brain and the complex process of head morphogenesis. The growth of the BN toward its target in the central brain is thought to be guided by the
axons of the so-called optic lobe pioneer (OLP) cells (Campos et al., 1995).

Thus far, three genes have been implicated in the development of the BO/BN projection, Krüppel (Kr, Schmucker et al., 1992), glass (gl, Moses et al., 1989) and disconnected (disco, Steller et al., 1987; Heilig et al., 1991; Lee et al., 1991). All three genes encode transcription factors with Zn-finger motifs. gl and Kr are expressed in the BO precursor cells. While gl seems to be required for proper differentiation of BO cells into photoreceptors, Kr is likely to be involved in several different aspects of BO/BN development. In contrast, disco is not expressed in the cells of the BO, but rather in a large portion of the OLA, which represents the cellular substratum for most of the growth of the BN (Lee et al., 1991). In disco mutant embryos, a variety of BN projection defects are observed. The BN defasciclulates and either projects to the OLA but fails to recognize and connect with its appropriate synaptic target cells, or projects to ectopic locations far from the OLA. In addition, the cohesion of the BO is impaired (Steller et al., 1987). However, because of the wide expression of disco, its precise role in the establishment of the BN projection is not well understood (Campos et al., 1995).

Here we present a developmental and genetic analysis of the BN projection. We show that the establishment of the projection occurs in three distinct phases during which the nerve contacts two intermediate targets at which it redirects its growth. We then present the results from a mutagenesis screen designed to identify genes involved in axonal growth and pathfinding of the BN. We have identified 13 genes, which disrupt the BN projection in different phases of its development. We focus our description on the projection defects of representative mutants and the expression patterns of the genes that they likely disrupt. By comparing the mutant phenotypes with the expression patterns, we are able to identify candidates for essential cellular components and assess their role in the normal projection of the nerve.

**MATERIALS AND METHODS**

*Drosophila* strains

Genetic markers and chromosomes are described in Lindsley and Zimm (1992). Lethal *P* enhancer trap mutations on the third chor-
some were provided by A. Spradling and are included in the Berkeley Drosophila Genome Project (BDGP).

**EMS mutagenesis and screening procedure**

Males of the transgenic marker line BS23 (Schmucker et al., 1992) were treated with 25 mM EMS using a standard protocol (Roberts, 1986). Mutagenized males were crossed en masse to e, DTS4/TM3, Sh, hhB females. Single F1 males BS23/TM3, Sh, hhB were then crossed to e, DTS4/TM3, Sh, hhB females. To facilitate the subsequent establishment of stocks, the crosses were subjected to a 2 day temperature shift to 29°C to kill the progeny bearing a DTS4 chromosome. F2 siblings of the genotype BS23/TM3, Sh, hhB were crossed to each other, to establish a balanced stock. 0-17 hour embryo collections of the balanced stocks were screened for defects in BO/BN development by histochemical staining for β-galactosidase activity (Klämbt et al., 1991). Embryos from each line were analysed under the dissecting microscope, and those showing any abnormalities in their BN projection were examined more carefully at higher magnification under a compound microscope (Zeiss Axiophot). Homozygous mutant embryos were easily recognizable by the absence of the TM3 hhB balancer, which harbors a hh-lacZ reporter gene.

Since the establishment of the BN projection is tightly linked to morphogenetic movements in the head, we carefully examined the mutations isolated in the primary screen for any defects in head segmentation and/or head involution as well as for general body pattern defects. As judged by morphological examination under Nomarski optics and 22C10 immunostainings, 27 mutations showed gross body pattern or nervous system defects. The remaining 45 mutations were tested for embryonic head defects by examining α-Engrailed stainings (Schmidt-Ott and Technau, 1992), and for defects in optic lobe formation by examining the expression of the markers Crumbs (Yanoussi-Hartenstein et al., 1993), FasII (VanVactor et al., 1993) and the disco enhancer trap line C50.1S1 (Heilig et al., 1991). Thus additional 27 mutations were eliminated because they showed either head involution defects that interfered with BO development or abnormally small OLAs or OLAs that failed to invaginate. Such mutations are likely to affect BN development only indirectly.

Mutations with similar phenotypes were tested for complementation of lethality. Members of three different complementation groups (II-F1, XI-E2 and V-D1) were mapped by meiotic recombination using the ru, th, st, cu, sr, e, ca marker chromosomes. II-F1 was found to be uncovered by the deficiency Df(3)XM3 and fails to complement the P enhancertrap insertion l(3)04505 (BDGP). The mutations XI-E2 and V-D1 failed to complement mutations in pointed and prospero, respectively. The chromosomal map positions of the other EMS mutations isolated in the screen have not yet been determined.

**Fig. 3.** The projection of the BN from P1 to P2 and into the central brain. (A,C,E) α-Kr (dark blue) and α-FasII (brown) double stainings of stage 14, 15 and 16 wild-type embryos, respectively. Apart from the BO, Kr expression is found in many cells of the CNS. (B,D,G,H) α-22C10 stainings of stages 14, 15, 16 and early first instar, respectively. (B,D) Whole-mount embryos; (G,H) paraffin sections. (F) α-Bsh/22C10 double staining of stage 16 wild-type embryo. (A,B) While the morphogenetic separation of BO (bold arrow) and OLA (arrowhead) proceeds, the projection of the BN to P1 (arrow) is maintained. (C,D) While the distance between BO and OLA continuously increases due to intercalary growth of the BN, a second phase of directed BN growth begins. (D) A few BN axons (arrows) emanating from a a thicker shaft (arrowhead) are visibly growing beyond P1 (arrowhead) and taking a sharp turn towards posterior (compare BN tip in B and D). (E) Concomitant with head involution, the BO (bold arrow) is displaced further anteriorly relative to the OLA (arrowhead; compare A, C and E). During this process, the BN elongates substantially, as does the FasII-expressing eOS. Note that FasII expression is lost from the eOS during stages 15/16 (compare A, C and E). (F) A single Bsh-expressing cell in the OLA (arrow) marks the intermediate target P2, which is contacted by the BN during stage 16. (G) The BN leaves the brain surface at P2 (arrowhead) and grows into the central brain (arrow indicates nerve tip). (H) In early first instar larvae, the BN has reached target cells in the central brain (arrow). The OLPs (arrowhead) are contacted by the BN, and their axons fasciculate. Scale bar, 16 μm (A-E); 10 μm (F-H).
Screening of P enhancer trap lines
Approximately 300 lethal P enhancer trap insertions on the third chromosome (A. Spradling) were examined for embryonic expression using α-β-galactosidase antibodies (S. Elend and H. Jäckle, unpublished data). Enhancer trap lines with expression near or in the BO and OLA were tested for phenotypic defects of the BN projection by 22C10 immunostainings. In this screen, we identified the P enhancer trap lines l(3)03550 (scd), l(3)03203 (pkl) and l(3)05845 (argos) (see Table 1).

P element reversions
P elements were mobilized in matings with a stable transposase source (Robertson et al., 1988). In each reversion experiment, 100 independent ry lines were established and tested for reversion of lethality and BN projection defects. The P enhancer trap line l(3)3203 is in fact semi-lethal; the semi-lethality is uncovered by the deficiency Df(3)pXT9. In the reversion experiment for l(3)3203, we also recovered lethal excision alleles. The mutant phenotype of several tested lethal excision alleles is similar to that of the original P insertion and is not enhanced in trans to the deficiency, suggesting that the original P insertion as well as the newly recovered excision alleles represent strong hypomorphic, if not amorphic, alleles of the pkl locus. In the case of l(3)03550, reversion of the P(ry+) element was also carried out by X-ray mutagenesis in order to recover deletions of the scd locus (Roberts, 1986). We screened 30,000 mutagenized chromosomes and found 7 chromosomes that had lost the ry+ marker and failed to complement the lethality of l(3)03550. 4 of these lethal revertants also fail to complement the lethality of neighboring lethal P insertions (l(3)03929; l(3)04083), and are therefore likely to represent deficiencies in the 88E-F region.

Immunostainings
Immunohistochemistry of embryos was carried out using standard procedures (Schmucker et al., 1992). The following primary antibodies were used: rabbit α-β-galactosidase (1:1000; Cappel); rabbit α-Kr (1:500; (Hoch et al., 1994); mouse mAb α-Fasciclin II (1:20; VanVector et al., 1993); mouse mAb 22C10 (1:50; Fujita et al., 1982); mouse mAb 24B10 (1:10; VanVector et al., 1988); rabbit α-Repo (1:500; Campbell et al., 1994). Detection of primary antibodies was carried out using the ABC-Elite kit (secondary antibodies 1:1000; Vectastain). In double label experiments, we used alkaline phosphatase- and peroxidase-conjugated secondary antibodies (1:2000; Jackson Immunoresearch), or, two peroxidase-conjugated secondary antibodies (1:2000; Vector Immunoresearch). Embryos were mounted in Vectashield (Vector) and immunofluorescent images were collected on a Bio-Rad MRC 1992). For confocal microscopy, primary antibodies were detected by FITC- or CY3-conjugated secondary antibodies (1:100; Jackson Immunoresearch). Embryos were mounted in Vectashield (Vector) and immunofluorescent images were collected on a Bio-Rad MRC 600.

RESULTS
In the first part of this section, we describe the development of the BO and the establishment of the BN. We begin with a description of the events that lead to cell fate determination and organogenesis of the BO, and continue with a detailed account of the development of the BN projection, including the initial outgrowth of axons and their fasculation, the elongation of the BN, and its navigation and ingrowth into the central brain. In the second part, we present the results from our mutagenesis screen, which reveal a number of genetic components required for different aspects of the establishment of the BN projection.

Development of BO and BN projection
Cell fate determination and organogenesis
The BO neurons derive from a coherent group of approximately 30 cells in the posterior head region of a stage 12 embryo (all staging according to Campos-Ortega and Hartenstein (1985), and can be visualized by the expression of the cell membrane antigen Fasciclin II (FasII; Grenningloh et al., 1991) (Fig. 2A). Four to six cells located at the ventral lateral margin of this cluster of FasII-positive cells represent BO precursors, while the remainder represent precursors of the optic lobe anlagen (OLA) (Fig. 2A), which will become part of the adult optic lobes. At late stage 12, BO and neighboring OLA precursor cells are morphologically indistinguishable, but can be differentiated by their expression of specific molecular markers. The BO precursors are marked by expression of the Kr gene (Schmucker et al., 1992), while the neighboring OLA cells (Fig. 2B-D) are marked by expression of the disco gene (Lee et al., 1991). From this stage onwards, BO development can also be followed by the expression of a lacZ reporter gene driven by a BO-specific enhancer element of the Kr gene (BS23 reporter gene; Hoch et al., 1990), which exclusively labels the cell bodies and axons of the BO (Fig. 2E-G; see below).

During stage 13 the BO precursors elongate, become polarized and form a distinct cell cluster (Fig. 2E). At the same time the BO cells separate from the OLA precursors. The OLA cells detach from the superficial ectoderm and invaginate in a process called optic lobe invagination (Campos-Ortega and Hartenstein, 1985). The BO precursor cells, as well as a few of the adjacent FasII-positive cells, remain at the surface ectoderm and form a cellular bridge between BO neurons and the invaginating OLA cells (Fig. 2E,G). We refer to this cellular bridge as the ‘embryonic optic stalk’ (eOS). During the morphogenetic separation of BO and OLA, the relative position between the two structures changes such that the BO comes to lie anterior to the OLA (Fig. 3A-E compare with Fig. 2B-D). Whether this rearrangement is a secondary consequence of optic lobe invagination or due to an active migration of the BO is unclear.

During stages 12 to 14, the number of Kr-expressing cells in the BO increases from 4-6 in the anlage to the final complement of 12 cells in the differentiated organ (compare Figs 2D,G and 3B). Whether this increase is due to cell division or involves recruitment of additional cells is not clear. During stage 13/14, the cell bodies of the BO form the tightly packed rosette characteristic of the mature organ (Fig. 3A,B). This morphogenetic event is preceded by the neuronal differentiation of BO cells during stage 13 as shown by expression of the 22C10 antigen (Fig. 3B). The final differentiation of BO cells into photoreceptors becomes apparent at stage 15 by their expression of Chaoptin, a membrane protein exclusively expressed in photoreceptor cells (VanVector et al., 1988) and detected by the monoclonal antibody 24B10.

Axonal outgrowth and nerve fasciculation
A single posteriorly projecting axon can be visualized in the BS23 line during early stage 13 (Schmucker et al., 1992). As additional BO cells undergo axonogenesis, their axons join a leading axon and fasciculate to form the BN (Fig. 2E). The sequential outgrowth of BN axons is paralleled by a sequential
expression of the 22C10 antigen in the corresponding BO cells (Schmucker et al., 1992).

The nascent BN initially extends posteriorly (Fig. 2E) and then ventrally as the separation between BO and OLA proceeds (Fig. 3A-D). During stage 13, the BN grows along the FasII-expressing cells of the eOS (Figs 2E, 3A), suggesting that the eOS represents a cellular substratum for the posteriorly directed growth of BN axons. By the end of stage 13, the distal tip of the fasciculated BN has reached the base of the eOS at the margin of the invaginated OLA. The position at which the distal tip of the BN contacts the OLA margin will be referred to as P1 (Fig. 2E).

Nerve elongation and directional growth
With the onset of stage 14, the process of head involution begins (Campos-Ortega and Hartenstein, 1985), which leads to an extensive movement of cells in the embryonic head. This complex process continues until the end of embryogenesis and is poorly understood. Concomitant with the morphogenetic movements of head involution, the BO is first displaced to a more dorsal and then to a more anteromedial location (Fig. 3A-F).

Between early stage 14 and midstage 16, the length of the BN between the somata of the BO and P1 (Fig. 3A,B) increases from 10 μm to over 70 μm (Fig. 3E). Considering that this elongation occurs after the distal end of the BN has already established its contact with OLA cells, this severalfold increase in the length of the BN is most likely due to intercalary growth rather than axonal growth at the tip of the BN. Based on their electron microscopic studies, Green et al. (1993) have suggested that this substantial elongation of the BN is driven by the displacement of the BO during head involution. However, whether the displacement of the BO is entirely a consequence of head involution or at least partially due to an active migration of the BO cell bodies is presently unknown.

During stage 14, the BN begins with a directed growth beyond P1, while the intercalary growth between BO cell bodies and P1 is still continuing (Fig. 3C,E). This process is distinct from the nerve elongation described above, since now the distal tip of the BN is growing by extending axonal processes beyond P1. As shown in Fig. 3D, axonal extensions emanate as relatively thin processes from a comparatively thick shaft which remains attached to P1. These axonal extensions grow in a sharp turn (60°-90°) towards posterior. The axons grow on the OLA surface in posterior/medial direction for about 35-40 μm. At stage 16, they reach a position marked by a single cell, which expresses the brain-specific homeobox (bsh) gene (Fig. 3F) during stages 15 and 16 (Jones and McGinnis, 1993). We refer to the position marked by the Bsh-expressing cell as P2.

Ingrowth into the central brain
Upon reaching position P2, growth of the BN stalls for some time and the distal tip of the nerve changes its morphology; while the tip of the nerve is rather blunt during its growth on the OLA surface, it now shows a more elaborate growth cone with extended filopodia-like processes at P2. These processes are very thin and cover an area of no more than two cell diameters (not shown). When nerve growth later resumes during stage 16, the distal end of the BN leaves the brain surface in a sharp turn (>90°) and grows into the central brain (Fig. 3G) to reach a small group of synaptic target cells (Fig. 3H).

At late stage 16, three 22C10-expressing cells are seen at the periphery of the OLA, which are termed optic lobe pioneer (OLP) cells (Fig. 3H; Tix et al., 1989). One of the OLPs is identical with the Bsh-expressing cell at position P2 (Campos et al., 1995). During its growth into the central brain, the BN remains in close contact with the OLP axons. We do not know, however, whether the OLP axons become part of the BN from P2 onwards or whether they in fact pioneer the projection of the BN into the central brain. A careful comparison of the time course of the OLP projection into the brain with that of the central projection of the BN will be necessary to distinguish between these two alternatives.

Isolation of mutations affecting BO and BN development
The identification of characteristic cellular landmarks along the BN pathway and the availability of the BS23 marker line, which permits the visualization of the entire projection in a simple histochemical procedure, encouraged us to use the BO/BN system to identify genes specifically required for axonal growth and pathfinding with a mutagenic approach.

In order to facilitate the visualization of the BN, flies

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**Table 1. Mutations affecting the BN projection**

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<tr>
<th>Alleles</th>
<th>Locus</th>
<th>Mutation</th>
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<tr>
<td>VI-G6</td>
<td>aloof (alf)</td>
<td>EMS</td>
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<tr>
<td>l(3)03550</td>
<td>scattered (scd)</td>
<td>P</td>
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<tr>
<td>VII-II</td>
<td>III-A2</td>
<td>EMS</td>
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<tr>
<td>VI-C4</td>
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<td>EMS</td>
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<tr>
<td>III-A4, III-C3, V-D1</td>
<td>prospero (pros)</td>
<td>EMS</td>
</tr>
<tr>
<td>l(3)05845</td>
<td>argos</td>
<td>P</td>
</tr>
<tr>
<td>II-F1, l(3)04505</td>
<td>abstrakt (abs)</td>
<td>EMS, P</td>
</tr>
<tr>
<td>XI-E2</td>
<td>pointed (put)</td>
<td>EMS</td>
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<td>XII-F3, I-D8</td>
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<td>EMS</td>
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<tr>
<td>l(3)03203</td>
<td>pumuckel (pkl)</td>
<td>P</td>
</tr>
<tr>
<td>VIII-E7</td>
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<td>EMS</td>
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<tr>
<td>II-E9</td>
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<td>EMS</td>
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<td>VII-C7</td>
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harboring the BS23 transgene were mutagenized using ethylmethanesulfonate (EMS). We generated 2800 lines bearing mutations on the third chromosome and examined these lines for BN projection defects. In addition, we examined 300 lethal P element insertions on the third chromosome (obtained from A. Spradling; Drosophila genome project) for BN defects. Among these lines, we identified 76 mutations (72 EMS alleles and 4 P-element insertions) that affect BO and BN development (see Materials and Methods). We then used a set of morphological criteria (see Materials and Methods) to assess the specificity of these mutations. Of the 76 mutations, 54 were excluded from further consideration, because they showed either a disruption of the segmented body pattern, abnormalities in head or brain morphology, or defects in optic lobe formation (not shown). The remaining 22 mutations fell into 16 different complementation groups (see Table 1). Four of the mutations could be mapped to the known genes gl, pointed (pnt), prospero (pros) and argos (Moses et al., 1989; Klämbt, 1993; Doe et al., 1991;...
Fig. 6. pnt causes misprojections and defasciculation of the BN. (A–C) α-Bsh/22C10 double stainings. (D,E) α-β-galactosidase stainings of pnt P enhancer trap lines. (A) In stage 16 wild-type embryos, the BN projection has reached P2, which is marked by a single Bsh-expressing cell (arrow). Additional Bsh-expressing cells are found outside the OLA in the embryonic brain (arrowhead). (B) pnt858 mutant embryo at stage 16. No Bsh-positive cell is found in the OLA and the projection of the BN ends without ever reaching a Bsh-expressing cell. The aberrant BN does not project to its normal ventromedial position (cross). Note that the other Bsh-expressing cells outside the OLA are present (arrowhead). (C) pnt858 mutant embryo at stage 15. At the OLA, the BN has split into two branches (arrow). (D,E) α-β-galactosidase/22C10 double stainings of the pnt P enhancer trap lines rM254 (D) and l(3)7825 (E) at stage 16. rM254 is an insertion in the P1-promoter and l(3)7825 is an insertion in the P2-promoter of pnt. In the rM254 line, several β-galactosidase-expressing cells are found along the BN (arrows, D). In the l(3)7825 line, a few β-galactosidase-expressing cells are found close to P2 in the posterior portion of the OLA (E). At least one of these cells is in direct contact with the BN (arrow in E). Scale bar, 16 μm (A,C,E); 10 μm (B,D).

Vaessen et al., 1991; Freeman et al., 1992; Kretzschmar et al., 1992. Mutations in three genes (including gl) lead to a strongly reduced cell number of the mutant BO. These mutations most likely affect early cell fate determination or cell proliferation processes (D. S., unpublished data).

In the present study, we analyse the remaining 18 mutations (corresponding to 13 genes, see Table 1) that do not alter the size of the BO or their photoreceptor identity. These mutations, which affect BO and BN development, fall into two major phenotypic groups. The first group of mutants show defects in both the formation of the BO and the projection of the BN, while the second only show defects in the projection of the BN. In the following description, we focus on representative examples from both groups.

Mutations that affect both the formation of the BO and the projection of the BN
Mutations in six different genes cause aberrations in BO formation (Table 1; Fig. 4). In these mutants, the BO cells fail to form a tight cluster of 12 photoreceptor cells, but retain their photoreceptor identity as revealed by their expression of the photoreceptor-specific marker Chaoptin (Fig. 4A-D, H-L).

In embryos homozygous for any of these mutations, the BO cells are either only loosely associated or displaced along the BN pathway (Fig. 4A,H). Interestingly, all these mutations also show characteristic aberrations in the BN projection. In particular, they show defasciculation of the BN, which results in the formation of ectopically projecting nerve branches. In one mutant of this group, aloof (afl) axonal projections fail to project to P1. In all the other mutants of this group, defasciculation of the BN occurs after P1 has been reached; scattered (scd) is shown as an example.

The scattered (scd) mutation disrupts cohesion of the BO and pathfinding of the BN.

The phenotype of scd is exemplary for the group of mutations

Fig. 7. pkl causes sprouting and misprojections of the BN at P2. (A–E) BN projection in stage 16 embryos visualized by α-Bsh/22C10 double stainings (A–C) and 22C10 single stainings (D,E). (F) α-β-galactosidase/22C10 double staining of a stage 16 embryo heterozygous for the pkl enhancer trap insertion P3203. Shown are ventral (A,C,F) and ventrolateral (D,E) views of the OLA area. (A) In a wild-type embryo, the BN projects as a single fasciculated nerve to the Bsh-expressing cell at P2. Sometimes, individual axons can be found directly contacting the Bsh-expressing cell (arrowhead). (B,C) In pkl mutant embryos, the BN grows beyond the Bsh-expressing cell. (B) The BN often splits (arrowheads) and the nerve branches or, as in C, the entire nerve grow around and beyond the Bsh-expressing cell. (D,E) The sprouting of the BN results in various branching patterns, however, none of the branches grow into the central brain. Note that ingrowth into the central brain should occur perpendicular to the optical plane shown here.

However, the aberrant BN branches, visible in B-E, extend parallel to the OLA/brain surface. (F) In embryos heterozygous for the pkl enhancer trap insertion P3203, several β-galactosidase-expressing cells (arrow and arrowhead) are found in the OLA. The cells appear to be in direct contact with the BN. Scale bar, 10 μm (A–C); 16 μm (D–F).
that affect both the formation of the BO and the projection of the BN (see Table 1). In scd mutant embryos a variable number of photoreceptors is displaced posteriorly along the path of the BN (Fig. 4A,B), and the normally tight fasciculation of BN axons is disrupted. The projection to P1 is still intact, but aberrant nerve branches are observed in the projection to P2 (Fig. 4C,D). In addition, we observe the formation of bifurcated nerve branches, which split and rejoin in the region of the OLA (Fig. 4C,D). Based on α-FasII stainings, both the eOS and the OLA are morphologically normal in scd mutant embryos (not shown).

The scd mutation is a lethal P element insertion (l(3)3550) at chromosomal position 88E9-9 (BDGP; Hartl et al., 1994). The lethality of the mutation as well as the disruption of the BN can be reverted by P element excision, indicating that the mutation is caused by the P element insertion. As the genetic locus is not uncovered by an existing chromosomal deficiency, we could not assess whether the mutation scd represents a complete loss-of-function allele for the locus. We generated four new X-ray alleles, which fail to complement the original scd mutation and are likely to represent small deficiencies including the scd locus (D. S. and U. G., unpublished data, Materials and Methods). In embryos that carry the scd mutation in trans over the X-ray alleles, the expressivity but not the severity of the BN phenotype is enhanced compared to the homozygotes. Thus, the scd mutation represents a strong hypomorphic rather than a complete loss-of-function allele of the locus.

The scd enhancer trap line P3550 shows β-galactosidase reporter gene expression during embryogenesis. During stages 15 and 16, expression is found in some cells of the eOS and OLA (Fig. 4E-G). These β-galactosidase-expressing cells are in direct contact with the BN along the major part of its pathway (Fig. 4G). Based on co-expression of the glia-specific marker, Repo, the P3550-positive cells in the eOS are glia cells (not shown), whereas the P3550-positive cells of the OLA, which are located at the base of the eOS are likely to represent neuronal cells as judged by their morphology. In addition, β-galactosidase expression is observed in cells of the developing nervous system and appears to be most prominent in glial cells associated with axonal projections, such as the exit glia and midline glia of the embryonic CNS (not shown). Barring the possibility that the P3550 enhancer trap is reporting the expression of a gene other than the one it disrupts (scd), this finding suggests that scd gene activity is in fact required in cells of the eOS rather than in the BO neurons themselves. Considering the BO and BN defects observed in the scd mutant, the finding suggests that cells of the eOS play a vital role in the formation of the BO and the pathfinding of the BN.

The aloof (alf) mutation affects cell clustering and leads to random projections of the BN

In alf mutant embryos the BO cell clustering is abnormal. However, the defects are different from those observed in the other mutants from this group (compare Fig. 4A and H,K). In most alf mutant embryos, we find two groups of cells, one of which represents a loose assembly of cells lying anteriorly or laterally to a relatively well organized cluster of BO cells (Fig. 4H,K). In addition, the BN projection is severely affected in alf mutant embryos. Only some of the photoreceptor axons fasciculate, project to P1 and grow toward P2, while a subset of BO axons (2-4) show random projections (Fig. 4I,J). In 10% of the mutant embryos, we observe photoreceptor axons that project anteriorly and explore areas far remote from the brain. Some of these axons develop enlarged axonal tips, most likely exaggerated growth cones (Fig. 4I). In some cases, it is clear that the randomly projecting axons originate from the abnormally oriented and slightly displaced cells, which are part of a mutant BO (Fig. 4J,K). It is important to note that these neurons have not lost their photoreceptor identity, as they still express Chaoptin (Fig. 4H-L). As observed with the scd mutant, the eOS and the OLA appear intact, consistent with the observation that in all alf mutant embryos some of the BN axons are still capable of projecting to P1 and toward P2. However, even these axons show defasciculation (Fig. 4L), while still in contact with the OLA. A detailed genetic analysis of this locus has not yet been undertaken.

Mutations that affect only BN projection

Mutations in seven complementation groups cause BN projection defects without affecting BO formation (Table 1; Figs 5-7). The mutants of this group all show defasciculation and aberrant projections of the BN. These mutants differ, however, with respect to the developmental time at which phenotypic aberrations are first observed and with respect to the position at which these defects occur along the path of the BN. The earliest defects can be detected in abstrakt (abs) mutants, which display abnormalities as soon as axonogenesis begins (Fig. 5). Mutations in two genes (including pnt) show defects later, when the BN is growing toward P2 (Table 1; Figs 6, 7). Mutations in pumuckel (pkl) and three other genes (Table 1) display defects still later, when the BN reaches P2.

abs mutations affect early axonal growth and BN navigation

abs mutant embryos form an intact BO (Fig. 5A,B). Defects in BN development become evident shortly after the onset of axonal outgrowth (Fig. 5C; compare with Fig. 3B,D). While in wild-type embryos the outgrowing BN axons rapidly fasciculate and project as a tightly fasciculated nerve straight to position P1, abs mutant embryos show 2-3 axon bundles. These aberrant bundles explore a much broader brain surface area (Fig. 5C-G). In more than 50% of mutant embryos, the nerve branches of mutant BNs deviate between 45° and 60° from their normal direction (Fig. 5D), resulting in severely branched nerves. While, in all mutant embryos, the nerve branches remain close to or in contact with the OLA, most nerve branches fail to reach position P2 (Fig. 5D). The few BN branches that eventually reach this position do not follow the normal path. Instead, they show abrupt directional changes along their projection (Fig. 5C,F). We often observe that the distal-most portion of the BN branches are significantly thicker than in wild type (Fig. 5C,F,G) and that they contain fine filopodia-like axonal extensions (Fig. 5F,G). These extensions cover an area of several cell diameters in the region of the OLA (Fig. 5G), suggesting that the BN axons of abs mutant embryos display an enhanced searching behavior as compared to wild-type BN axons. As judged by α-FasII- and α-Bsh-stainings, the eOS, the OLA and P2 are morphologically intact in abs mutant embryos.

The abs locus is represented by an EMS allele (l(3)4505; BDGP; Hartl et al., 1994). The BN
phenotype of the EMS allele IIF-1 is not enhanced in trans to a deficiency for the region (Materials and Methods), suggesting that IIF-1 represents a strongly hypomorphic if not complete loss-of-function allele of the abs locus. l(3)4505 contains a single lethal P enhancer trap insertion at chromosomal location 82A1 (BDGP, Hartl et al., 1994), which can be reverted to wild type by P element excision. Unfortunately no embryonic reporter gene expression pattern is seen in the l(3)4505 P enhancer trap line. Thus, we have no indication whether the abs gene activity is required in cells of the BO or in the surrounding cells, such as the eOS or the OLA. Nevertheless, our results demonstrate that abs is essential for the directed and fasciculated outgrowth of the BN early on, as well as its navigation at later stages.

pnt causes defasciculation of the BN between P1 and P2

In pnt mutant embryos, the BN initially fasciculates and projects normally. However, during its growth from P1 to P2, defasciculation of the BN and misrouting (Fig. 6A-C) of the two to three resulting nerve branches (Fig. 6C) are observed. The aberrant BN branches grow on the surface of the OLA and their endings reach various ectopic positions during mid-stage 16. Interestingly, we find that, in 80% of the examined mutant brain hemispheres, the Bsh-expressing cell, which is normally found at position P2, is either missing or mislocalized, while other Bsh-expressing cells appear in their normal positions (Fig. 6B,C). The aberrant BN branches never reach any of the other Bsh-expressing cells. As judged by α-FasII stainings, the eOS and the OLA are present in pnt mutant embryos, although some OLA cells appear to express lower levels of FasII (not shown). In all our experiments, we have used the complete loss-of-function allele pnt(k0) (Klämbt, 1993).

The pnt gene encodes an ets-domain containing transcription factor. In the embryonic CNS, pnt is required in the glial cells that are involved in the formation of the anterior and posterior commissures (Klämbt, 1993). We examined pnt expression by monitoring β-galactosidase expression in the pnt P enhancer trap lines l(3)7823 and Rm254, which represent insertions in the two different promoters of the pnt gene (O’Neill et al., 1994; Klaes et al., 1994). During stages 14-16, a few β-galactosidase-positive cells are found along the entire path of the BN (Fig. 6D). In addition, a few β-galactosidase-positive cells are found close to P2 (Fig. 6E). Interestingly, one of these cells near P2 is contacted by the tip of the nerve at stage 16 (Fig. 6E). Based on their morphology, some of the pnt-expressing cells are likely to represent glial cells. We cannot exclude the possibility that pnt is expressed in additional cells in the environment of the BN, since the two P enhancer trap lines together do not report the entire pattern of pnt gene expression (C. Klämbt, personal communication). These results suggest that pnt gene function is required in cells along the path of the BN, as well as in the P2 region for the proper pathfinding of the nerve.

pkl affects the projection of the BN at P2

In embryos mutant for the pkl mutation, as well as three other mutations (see Table 1), the BN is properly fasciculated and follows its normal path until it approaches position P2. However, instead of taking its normal turn into the central brain, the BN undergoes terminal sprouting in the P2 area. This leads to the formation of several independent nerve branches (Fig. 7A-E), which remain on the brain surface (Fig. 7B-E). In about 20% of the mutant embryos defasciculation of the BN is visible prior to reaching P2. Based on α-FasII stainings, we do not find any morphologically visible abnormalities in the tissues associated with the BN, such as the eOS or the OLA (not shown), and the Bsh-expressing cell is present in its normal position as revealed by α-Bsh/22C10 double stainings. Considering that the wild-type BN displays searching behavior in the P2 area, these findings suggest that an essential spatial cue is missing at P2 or that the BN has lost its ability to recognize this cue.

pkl is a single semilethal P element insertion at chromosomal position 85D1-2 (l(3)3203; BDGP, Hartl et al., 1994), which likely represents a strong hypomorphic if not complete loss-of-function allele of the pkl locus (Materials and Methods).

The pkl enhancer trap line P3203 shows weak expression in the posterior head of stage 11 embryos (not shown), covering the region that will give rise to OLA and BO precursor cells during stages 12 and 13. During stages 14-16, the β-galactosidase reporter gene is also strongly expressed in cells that are associated with the BN close to position P2 (Fig. 7F). Based on their morphology, these P3203-positive cells are likely to represent glial cells. α-β-galactosidase/α-Bsh double stainings confirm that none of the P3203-positive cells are identical with the Bsh-positive cell at P2 (not shown). However, we have not been able to determine whether any are identical with the pnt-positive cells close to position P2. In any case, the expression of the pkl enhancer trap line P3203 in cells located in the P2 area correlates well with the appearance of phenotypic defects at P2 in the homozygous mutant embryos, suggesting a role for pkl in providing a permissive substratum or a directional cue for axonal growth at position P2.

DISCUSSION

The developmental and genetic analysis of the BO/BN system has led to the identification of several cellular components that are part of the BN pathway and revealed developmental strategies employed in the establishment of the BN projection.

Development of the BO and the eOS

Several lines of evidence suggest that the cells of the BO and the OLA are derived from a common pool of precursor cells. At embryonic stage 12, the Kr-positive BO precursors are found immediately adjacent to disco-positive OLA precursor cells in the posterior head. However, the two cell populations co-express both the sine oculis (so) gene (Cheyette et al., 1994; Serikaku and O’Tousa, 1994) and the cell surface marker FasII (Grenningloh et al., 1991). Furthermore, the fate of OLA precursors is readily switched to that of BO cells as is observed in patched (ptc) mutant embryos (Schmucker et al., 1994), suggesting the existence of a common origin for the two cell types. During the morphogenetic separation of BO and OLA first by optic lobe invagination and later by head involution, the two structures remain in contact through a cellular bridge, which we term the embryonic optic stalk (eOS). Some of the eOS cells may later differentiate into glial cells, because a string of Repo-positive glia is found lining the path of the BN along the
eOS at stages 16 and 17 (see Fig. 8 and Campos et al., 1995). Alternatively, the precursors of these glia may migrate to the eOS from elsewhere. However, since the lineage of the glia in the embryonic head has not been traced, we cannot discriminate between these two possibilities. Furthermore, we have not followed the fate of the eOS through postembryonic stages of development. It will be very interesting, however, to examine whether the cells of the eOS persist into larval stages and how they relate to the larval optic stalk (Meinertzhagen and Hanson, 1993), whose function it is to funnel the developing adult photoreceptor axons into the optic lobe.

**Development of the BN projection**

The establishment of the BN projection occurs in three distinct steps and involves at least two different modes of axonal growth. BN axons first project toward the intermediate target P1 by directed growth at their tip. Once the distal tip of the BN has reached P1, a substantial elongation of the nerve occurs, which likely is brought about by intercalary growth. Concomitant with the elongation of the nerve, the BN resumes its directed growth at the tip and projects first to P2 and then into the central brain. During this second phase of directed growth, the BN takes two sharp turns, the first at P1 and the second at P2.

The fidelity of the first step, the projection of the BN to P1, is ensured by the formation of the eOS, which serves as a cellular substratum for growth of the BN. The formation of a cellular bridge provides an effective developmental strategy to facilitate axonal pathfinding which has not been described for any other nerve pathway in the Drosophila embryonic PNS or CNS. Later on, the cells of the eOS may be important for stabilizing the axon bundle or for anchoring the axons to the OLA during the subsequent massive elongation of the BN.

During the second phase of directed growth, the OLA provides the cellular substratum for growth. However, the analysis of the wild type did not provide any specific insights as to which cues are responsible for reorienting the growth of the BN at P1. In contrast, P2 likely serves a function as a guidepost directing fibers into the central brain, since nerve growth stalls at P2 and individual processes are found exploring the area.

The oriented growth of the BN to P1 and then to P2 occurs in a sequential fashion, that is, one or very few axons grow first, while others follow. In a number of systems, leading fibers have been reported to serve as pioneers for the pathfinding of the follower axons (Goodman and Shatz, 1993; Bentley and Keshishian, 1982). However, in the case of the BN, we have not been able to determine whether the leading fibers are invariably generated by (a) specific neuron(s) of the BO, since subset-specific fiber markers have not been identified for the BN. Thus, we cannot exclude the possibility that the axons simply grow at different rates, and perhaps lead alternately. Interestingly, all axons stall for some time upon reaching P2, suggesting that P2 not only serves to guide the axons into the central brain, but also to synchronize their growth.

**Mutations affecting both the formation of the BO and the projection of the BN**

The mutations that we isolated in our screen fall into two broad phenotypic groups. In the first group of mutants, both the BO and the projection of the BN are affected, while in the second group only the BN projection is abnormal.

The fact that we find an entire group of mutations (Table 1) that affect both the formation of the BO and the pathfinding of the BN emphasizes the close link between the two processes. Based on the observation that phenotypic defects in the BO only become visible while the massive elongation of the BN is taking place during stages 14 to 16, we favor the following interpretation. The disruption of the BO may largely be a consequence of the mechanical stress applied to both the BN and the BO during their morphogenetic separation in head involution. It is possible that this mechanical stress exacerbates developmental defects that would otherwise be less apparent such as reduced cohesion within the BN bundle or reduced adhesion to P1. The expression of *scd* enhancer trap line P3550 in glial cells of the eOS and in cells of the OLA at the base of the eOS is consistent with this interpretation. The glia surrounding the BN along the eOS may play a crucial role in establishing or maintaining proper cohesion of the nerve, while the cells at the base of the eOS are required for its adhesion to P1. Whether the projection defects of the BN on the OLA surface observed in the *scd* mutant could also be attributed to defects in the eOS glia or the cells at the base of the eOS is less obvious. However, it is interesting to note that some of the phenotypic defects observed in *disco* mutant embryos are strikingly similar (Steller et al., 1987) to those found in *scd*. *disco* is also expressed in the eOS and required there for the proper differentiation of glial cells (Campos et al., 1995).

In a different example of this group of mutants, *alf*, a subset of BO cells is not properly integrated into the main cluster and projects randomly, while the major part of the BO projects toward P1 and P2. It is quite likely that some of the BO cells fail to obtain their proper orientation in the cluster and therefore fail to project along the eOS. However, the cause of the projection defects observed in the portion of the BN that succeeds in projecting to the OLA is presently unclear. Remarkably, this is the only mutation from our screen that
shows random projections. In all other cases, abnormal projections retain contact either with the OLA or the brain surface.

**Mutations affecting only the BN projection**

The phenotypic defects of the mutants that affect only the BN projection share similarities, in that they all show defasciculation of the BN and aberrant projections of the resulting nerve branches. However, in the different mutants, phenotypic defects occur at different stages of the BN projection, suggesting that distinct cellular and molecular mechanisms are involved in each of the three steps of BN development.

In *abs* mutants, defasciculation of the BN occurs with the onset of axonogenesis, when independent nerve branches are established. However, since at later stages one may find a single nerve bundle, branches may either be retracted or refasciculate. The latter possibility would suggest that the system is capable of compensating for mistakes. In addition, the morphology of the nerve tip is highly abnormal in *abs* mutants and shows enormous growth-cone-like structures and filopodial extensions, which we have never observed in either wild-type or other mutant backgrounds. These abnormally large extensions may reflect defects in the capacity to withdraw extensions or may reflect an exaggerated search behavior. As we have not gained any insights into the expression pattern of *abs*, we cannot assess whether the gene is required in the cells of the BO or in the cells of the OLA.

In the majority of the mutants in this group, phenotypic defects are found later, either on the way between P1 and P2 (*pnt*), or at P2 (*pkl*). The phenotypic defects observed in *pnt* mutants correlate with the location of *pnt*-positive cells. In the wild type, *pnt* is expressed in presumed glia along the path of the BN, mostly between P1 and P2, which is where the BN axons defasciculate in the homozygous mutant. *pnt* is also expressed in presumed glia in the P2 region. In the homozygous mutant, there is often no Bsh-expressing cell at P2, suggesting that *pnt* function is required in the neighboring cells for induction of Bsh-expression in the cell at P2. Similarly, *pnt* has been reported to be required in glial cells of the embryonic CNS for the induction of the neuronal marker 22C10 in the MP2 cells (Klaes et al., 1994).

In *pkl* embryos, both the Bsh-positive cell and the putative pkl-positive cells, which likely represent glial cells, are still present at their normal position. However, the axons of the BN do not seem to recognize P2. Instead of growing straight toward the Bsh-positive cell and stopping there, the BN axons grow beyond the Bsh-cell and begin with an abnormal search behavior in the *pkl* mutant. These phenotypic defects corroborate the notion that P2 plays a dual role of growth inhibitor and guidepost. The putative expression pattern of *pkl* furthermore suggests that the P2 position consists of several cellular components, including the Bsh- and the putative pkl-positive cells. It is possible that these components are required individually for proper pathfinding of the nerve. Alternatively, the putative pkl-positive glial cells may be required for proper development or differentiation of the Bsh cell as guidepost.

**Concluding remarks**

An interesting result from our mutant analysis is that all the mutants that we found show defasciculation of the nerve in conjunction with misprojection of the nerve’s branches. We believe that the defasciculation of the nerve observed in our mutants is mainly a consequence of impaired pathfinding. As a consequence of missing cues or an impaired capacity to sense cues, individual fibers or groups of fibers undertake an independent exploration of the environment, thereby foregoing fasciculation with the bundle. It is quite possible that the fibers of the nerve bundle communicate to ensure concerted growth. Leading axons would signal to following axons that their pathfinding is successful, thereby reinforcing the fasciculated growth of the followers. Such positive feedback would not occur if the leading fibers fail to receive the proper cues.

The fact that in each subgroup of mutants projection defects are observed at different stages of BN development strongly suggests that distinct mechanisms are involved in the different steps of BN pathfinding. This interpretation is further corroborated by our finding that the genes disrupted by these mutations are likely to be expressed in different cellular components along the path of the BN. These cellular components include glial cells in the eOS, a few, presumably, neuronal cells at the base of the eOS, presumptive glia on the OLA and, finally, a complex of neuronal and glial cells at the nerve’s entry point into the brain. The presumptive glia on the OLA and the complex of cells at P2 may be required for proper pathfinding of the BN on the OLA surface and for its entry into the brain. The molecular characterization of the mutants promises to provide important insights into the mechanisms of BN pathfinding and nervous system development.

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**REFERENCES**


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