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NEURONAL CONNECTIVITY IN THE LARVAL VISUAL SYSTEM OF DROSOPHILA MELANOGASTER: CELLULAR AND MOLECULAR ANALYSES

By

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A Thesis

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Abstract

In this study the *Drosophila melanogaster* larval visual system development was used as a model to identify cellular interactions and uncover molecular mechanisms that govern the formation of precise neuronal pathways during nervous system development.

The first part of this study demonstrates that the outgrowth of two serotonergic processes and their contact with the larval optic nerve at the larval visual center are dependent on the proper development of larval optic nerve (LON).

The second part of this study reports identification of molecular interactions that may be important for larval visual system development. A tissue-specific autoregulatory activity of a putative transcription factor DISCONNECTED (DISCO) has been shown to be important for proper larval visual system connectivity formation. To understand the molecular mechanisms that govern *disco* function in the visual system, two new proteins (DIP1 and DIP2; DISCO-interacting proteins 1 and 2) and a previously known protein that interact with DISCO in yeast and *in vitro* were identified. These proteins also show overlapping mRNA expression patterns with DISCO during embryogenesis.

This study has also led to the identification of a putative activation domain in the DISCO protein. Furthermore, the results of this study suggest a possible protein-binding function of the second zinc-finger domain of DISCO.
The dip1 gene encodes a protein with two putative dsRNA-binding motifs. An unusual repeated element is present in the 3′ untranslated region of dip1 cDNAs. Northern analysis revealed expression of three different size transcripts of the gene during embryogenesis. The genetic location of the dip1 gene is similar to that of a gene called flamenco.

The dip2 gene located at the polytene chromosome band 61B, codes for a highly conserved protein with presently unknown function. The amino acid sequence and mRNA expression pattern of the Drosophila DIP2 protein are very similar to its mouse homolog.

This study contributes towards understanding the cellular and molecular mechanisms of neuronal connectivity formation.
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*dip2*
Chapter 1: Introduction
Introduction:

The complex organization found in a functional nervous system requires that axons establish their correct pathways and find their appropriate targets during development. Precise neuronal connections in the nervous system occur in two steps: In the first step various molecular guidance cues guide a neuron to its target region which is initially broadly defined. The second step includes an exact point-to-point matching between each axon and the specific target neuron. The first step i.e. axon outgrowth, pathfinding and target selection do not require action potentials and synaptic transmission. However, the second step i.e., refinement and remodeling of synaptic connections almost always require neuronal activity (reviewed by Goodman and Shatz, 1993).

General strategies for neuronal pathway formation and cellular guidance

To make precise connections with their targets, axons need to grow long distances, as long as several centimeters or more than a thousand times the diameter of the cell body. To simplify this task the journey is fragmented into shorter steps where intermediate targets provide critical guidance cues directing growth cones to the next stage of the neurons trajectory. The intermediate targets are often provided by smaller, isolated groups of transient neurons, called 'guidepost cells'. In the absence of the 'guidepost cells' the earliest pioneer growth cones often make pathfinding errors (Klose and Bentley, 1989). Later
navigating neurons may be guided to and from choice points by selective fasciculation on earlier-developing axons (reviewed by Goodman et al., 1984).

Studies with vertebrate and invertebrate systems established that pioneer neurons play an important role in neuronal pathway formation (e.g. McConnell et al., 1989; Klose and Bentley, 1989). These neurons establish initial scaffolds of axon tracts that act as substrates for directed migration of later developing neuron growth cones. For example, the first neurons which differentiate in grasshopper limb, the sibling afferent Ti1 neurons, send out axons along a defined pathway, and later developing sensory neurons follow the pathway pioneered by the Ti1 neurons (Bate, 1976). Along the pathway at three locations, the pioneer axons contact specific cells (Fe1, Tr1 and the pair of Cx1 cells) which are likely to act as intermediate targets and guidance cues. The route taken by the pioneer axons always passes over the so called "guidepost" cells, which are immature neurons. Transient gap junctions between the pioneer fibres and the guidepost cells have also been reported. Ablation of the Cx1 cells with UV results in failure of the pioneer growth cones to leave the trochafter/coxa boundary and to grow proximally to the CNS (Bentley and Caudy, 1983).

Cells with characteristics of guidepost cells are likely to act as intermediate targets in the vertebrate nervous system as well. A set of early generated and transient neurons in the cortex, the 'subplate cells', may be involved in guiding the thalamic fibres toward and into their cortical target area. Ablation of the subplate neurons, by kainate injection, leads to a failure of thalamic fibres to recognize their normal cortical target areas (Ghosh et al., 1990). Thus subplate
cells may provide critical guidance cues which help thalamic axons to locate and recognize their correct target.

*In vitro* assays and *in vivo* perturbation studies have led to the proposal that selective fasciculation is an important determinant of precise neuronal connectivity (Letourneau, 1975; Raper et al., 1984; Bastiani et al., 1986). After pioneer neurons establish the first scaffold of different pathways, follower axons must choose which fascicles will bring them to their correct targets. Reaching a distant target may require a series of navigational choices, each followed by a period of fasciculation along a new pathway. The ordered neuronal pattern in the insect nervous system arises from the ability of individual central nervous system (CNS) neuron growth cones to discriminate between fascicles and make selective choices during outgrowth (Goodman et al., 1984). For example, in the grasshopper embryo, the growth cone of the aCC neuron selectively fasciculates with three U axons in the U fascicle and that of pCC neurons fasciculate with MP1 and dMP2 axons in the MP1 fascicle (Bastani et al., 1986; Du Lac et al., 1986). When the pioneer neurons (U axons, MP1 and dMP2) of these two pathways are selectively ablated, the follower growth cones (aCC and pCC respectively) stall and do not show affinity for any other neighboring axons. This data supports the “labeled pathway” hypothesis (Raper et al., 1984) which proposes that CNS pathways are selectively labeled with guidance molecules that allow axon growth cones to select and fasciculate with appropriate axon fascicles.

Monoclonal antibody screens in the developing insect nervous system and other studies in both invertebrate and vertebrate system led to the identification of molecules selectively expressed on subset of developing axon pathways (e.g.,
Bastiani et al., 1987; Grenningloh et al., 1991; Hynes, 1992). These studies have shown that axonal fasciculation and defasciculation are controlled by adhesion mechanisms mediated by surface glycoproteins. Genetic analyses has revealed the in vivo functional significance of at least one of these cell adhesion molecules (Fasciclin II in Drosophila) in selectively labeling a subset of CNS axon pathways (Grenningloh et al., 1991).

In vertebrate systems the 'Labeled Pathway' hypothesis may explain the ability of axons from chick motor neurons to select appropriate nerve routes and plexuses (Lance-Jones and Landmesser, 1981). This general principle may even apply to development of the mammalian cerebral cortex. In this system, early developing axons from cortical subplate cells pioneer the pathway from the cortex to the thalamus and other subcortical targets (McConnell et al., 1989). Later, follower axons from cortical neurons associate with these subplate fibres and may be guided along these pathways to their appropriate targets.

Non-neuronal cells have also been implicated in providing guidance cues to axon tracts. For instance, the role of glial cells in growth cone guidance has been recognized in both insects and vertebrates. In vertebrates glial cells appear, prior to neurite outgrowth, along the pathways that axons will subsequently follow. This observation led to the 'blueprint hypothesis' (Singer et al., 1979) which suggests that the paths recognized by growth cones are formed by glial cells before axon outgrowth. This hypothesis is supported by experiments in which this preformed pathway was physically ablated, leading to a disruption of the axon tract (Silver et al., 1982).
The initial evidence suggesting the role of glial cells as guidposts in insects was provided by experiments in grasshopper in which one glial cell type, the segment boundary cell (SBC) was ablated using a laser beam (Bastiani and Goodman, 1986). In this organism, the U axons, which pioneer the intersegmental nerve, contact the SBC prior to exiting the CNS. The growth cone of the aCC motor neurons follow the U axons while exiting the ventral nerve cord in the intersegmental nerve. After ablation of the SBC, both the U and aCC axons fail to exit the CNS.

Studies on the development of chick motor neuron projections to limb muscles have indicated that mesenchyme and connective tissue can directly or indirectly pattern nerve pathways in the nerve plexus region and in the limb (Lance-Jones and Dias, 1991; Tosney and Landmesser, 1984). In addition, somatic tissue can also specify axonal pathways (e.g., Oakley and Tosney, 1993; Keynes and Stern, 1984), as demonstrated by the fact that axons from spinal motor neurons project their axons only through the rostral half of each somite. Following rotation of the paraxial mesoderm such that the rostrocaudal polarity of the somites was reversed, there was a corresponding reversal of the pattern of outgrowth of motor axons (Keynes and Stern, 1984).

Molecular mechanisms of axon guidance and families of axon guidance molecules

At least four axon guidance mechanisms have been recognized; (1) short-range attractive or permissive cues, (2) short range repulsive cues, (3) long range attractive cues and (4) long-range repulsive guidance cues (Reviewed by
Goodman and Shatz, 1993; Keynes and Cook, 1995; Goodman, 1996). The short range cues are provided by non diffusible cell surface molecules and extracellular matrix molecules which either permit axon growth through selective adhesion or inhibit or repel axons on contact (e.g. Luo and Raper, 1994; Jessell, 1988). In contrast, long range cues are provided by diffusible molecules (Cajal, 1892; Tessier-Lavigne et al., 1988; Colamarino and Tessier-Lavigne, 1995; Ebens et al., 1996). Diffusible cues form gradients that originate from distant tissues and specify the direction of axonal growth (reviewed by Tessier-Lavigne, 1992). Although these definitions imply distinct mechanisms, in reality, the differences between these mechanisms are often blurred. For example, a secreted molecule in certain contexts can become immobilized by binding to cell surface or extracellular matrix and thus act as a short range signaling cue (e.g. Baier and Bonhoeffer, 1992). The importance of a balance between positive and negative cues in axon pathfinding have been recognized in both vertebrate and invertebrate systems (reviewed by Tessier-Lavigne and Goodman, 1996). The integration of multiple and often opposing molecular interactions at each site along the axon’s trajectory, specially at choice points, help to fine tune the directional response of its growth cones.

In recent years substantial progress have been made in the isolation of molecules involved in axon guidance. The molecular characterization of the guidance cues shows that they belong to different families which provide axon guidance in both vertebrates and invertebrates, supporting the view that both the molecules and the mechanisms of growth cone guidance are conserved phylogenetically (Hynes and Lander 1992, ; Goodman, 1994).
In the context of molecules that behave as short range cues, attention has focused on neural cell adhesion molecules of the immunoglobulin (Ig) superfamily (Reviewed by Walsh and Doherty, 1997). The Eph family of receptor tyrosine kinases have gained attention in recent years for their role in axon navigation, particularly in the context of target recognition (e.g. Callahan et al., 1995; Cheng et al., 1995; Xu et al., 1996).

The gene families which were originally identified on the basis of their role in long-range signaling are the netrin and semaphorin gene families. Interestingly, the first chemoattractant to be isolated, netrin-1, displays sequence similarity with the extracellular matrix protein laminin (Serafini et al., 1994). Similarly, the semaphorin family contains both cell surface and diffusible members (Kolodkin et al., 1993). Thus these families are implicated in both long- and short-range signaling. In addition, netrins are bifunctional, attractive to some axons and repulsive to others (reviewed by Culotti and Kolodkin, 1996). Thus individual molecules or families of molecules do not always fit into any one mechanistic category. Furthermore, how one axon will behave towards a guidance cue presumably depends on the receptors expressed by its growth cones (Colamarino and Tessier-Lavigne, 1995; Wehrle and Chiquet, 1990; Mukhopadhyay et al., 1994; Nose et al., 1994).

The netrins are homologs of the C. elegans UNC-6 protein, which plays a role in circumferential guidance (Hedgecock et al., 1990; Ishii et al., 1992; Serafini et al., 1994). In vitro experiments have shown that the floor plate at the ventral midline of the developing mammalian spinal cord provides chemotrophic guidance signals for the growth cones of commissural neurons whose axons
extend towards the floor plate (Tessier-Lavigne et al., 1988; Placzek et al., 1990). When a dorsal spinal cord explant is placed within a few hundred microns of a floor plate explant in a collagen gel matrix, commissural growth cones turn and extend towards the floor plate from a distance. Two vertebrate netrins were isolated and identified as the chemoattractants that emanate from the floor plate cells and guide the spinal commissural axons (Serafini et al., 1994; Kennedy et al., 1994). Experimental analysis suggests that vertebrate netrins and nematode UNC-6 are bifunctional molecules that can act both as chemoattractants and chemorepellents. These molecules secreted by ventral cells appear to attract some growth cones toward the ventral midline but to repel other growth cones dorsally from the ventral midline. While netrin 1 attracts commissural growth cones towards the ventral midline, it repels subpopulations of dorsally-projecting hindbrain motor axons including the trochlear motor axons (Colamarino and Tessier-Lavigne, 1995; Varela-Echavarria et al., 1997).

UNC-40 and UNC-5 are receptors for nematode UNC-6 that mediate chemoattractant and chemorepellent effects of UNC-6 respectively (Hedgecock et al., 1990). Based on DNA sequence similarity two vertebrate netrin chemoattractant receptors (DCC and neogenin) have been identified (Keino-Masu et al., 1996). These receptors are members of immunoglobulin superfamily. Functional evidence for the role of DCC came from in vitro experiment in which it was demonstrated that the floor plate dependent outgrowth of commissural axons can be blocked by administering antibodies raised against the extracellular domain of DCC (Keino-Masu et al., 1996).
The Semaphorins (Sema) are a family of cell surface and secreted proteins that in general function as chemorepellents or inhibitors of axon pathfinding, branching or targeting (reviewed by Goodman, 1996). The first identified member of the family, Semaphorin-1, (Kolodkin et al., 1992), is a transmembrane protein. Sema 1 is expressed on subsets of fasciculating axons in the developing CNS and on stripes of epithelial cells of the developing limb bud of grasshopper. Antibody blocking experiments in the limb bud show that Sema 1 inhibits branching and prevents steering of identified growth cones. In contrast to Sema I, Sema II and III and several other members of the family lack a transmembrane domain and are secreted. A member of the Semaphorin family, Collapsin (a homolog of human Sema III), was identified in chicken on the basis of its ability to cause the collapse of sensory growth cones in vitro (Luo et al., 1993).

Extracellular matrix molecules (ECM) such as collagen, fibronectin and laminin bind cell surface receptors that may be integrins or immunoglobulin superfamily members. Laminin is known to promote axon outgrowth in vitro (Gunderson, 1987) and is present in some axonal pathways, notably in that of the optic tract (Cohen et al., 1987). ECM molecules may have positive or negative effects on axon outgrowth. For example, tenascin-C has both axon outgrowth promoting and anti-adhesive domains (Gotz et al., 1996). It has been suggested that the function of ECM molecules is regulated by binding to proteoglycans. For example, tenascin binds to glypican, laminin to dystroglycan and so on (Vaughan, et al., 1994; Gullberg and Ekblom, 1995).

Cell adhesion molecules (CAMs) are an important class of axon guidance molecules. These are expressed on the cell surface and fall into several structural
classes, most notably the immunoglobulin superfamily, the cadherins and the integrins. Integrins are transmembrane proteins that bind ligands that are components of extracellular matrix, such as laminin, fibronectin, vitronectin, thrombospondin-1 and others (Clark and Brugge, 1995). For example, integrin mediates the thrombospondin-1 promoted outgrowth of sympathetic neurons from the central and peripheral nervous system. Cadherins are transmembrane glycoproteins that constitute the calcium-dependent adhesion system, in which molecules on adjacent cells bind homophilically (Takeichi, 1991). The neural-cadherins are implicated in the guidance of retinal axon outgrowth in vivo (Riehl et al., 1996). The CAMs of the immunoglobulin superfamily include neural CAM (NCAM), L1/NgCAM, NrCAM and TAG-1/axonin-1. NCAM is the first identified CAM. It constitutes the calcium-independent adhesion system (Cunningham et al., 1987).

Elegant studies on the role of immunoglobulin superfamily members have examined pathfinding of chick commissural interneurons across the floor plate (Stoeckli and Landmesser, 1995) and axon pathfinding in the muscle target area of the chick limb bud (Tang et al., 1992, 1994). During their growth across the floor plate the commissural axons express NgCAM and axonin-1, while the floor plate expresses NrCAM. Perturbation of axonin-1 function by injecting anti-axonin antibody caused axonal defasciculation of commissural neurons and many axons failed to cross the floor plate, and turned instead along ipsilateral floor plate border (Stoeckli and Landmesser, 1995). Injection of anti-NrCAM also resulted in a similar commissural axon phenotype suggesting that interaction between axonin-1 on growth cones and NrCAM on floor plate cells is
involved in the entry of commissural axons into the floor plate (Stoeckli and Landmesser, 1995).

The navigation of motor axons from the neural tube into the periphery has proved an interesting system for studying the function of cell adhesion and other molecules in axon guidance. The initial growth of motor axons through the anterior sclerotome was proposed to be due to repulsive interactions between peanut agglutinin binding protein and T-cadherin, expressed on the posterior sclerotome and motor axons respectively (reviewed by Keynes et al., 1991). Polysialic acid (PSA) associated with NCAM is involved in modulating interactions between axons (Tang et al., 1994). PSA expresses at low level in regions were motor axons are tightly fasciculated before reaching the plexus (the region where motor axons rearrange before entering the limb bud of chick embryo), and at higher level in regions of defasciculation. Specific removal of PSA by application of the enzyme endoN resulted in motor axon projection errors and increased fasciculation in the plexus region (Tang et al., 1992, 1994). The effect of endoN on axon pathfinding was reversed by treatment with antibodies against L1, implicating that PSA induces inhibition of L1-mediated fasciculation, which in turn facilitates normal motor axon guidance (Tang et al., 1994). Such balance of interactions mediated by NCAM and L1 is also involved in regulating patterns of nerve branching to fast and slow regions of embryonic muscle (Landmesser et al., 1988).

A variety of receptor protein tyrosine kinases (RPTKs) including fibroblast growth factor receptors, and the Trk family of neurotrophin receptors have been implicated in the regulation of axon outgrowth, target invasion, and axonal
branching (reviewed by Basilici and Moscatelli, 1992; McFarlane et al., 1995; Barbacid, 1995). The largest subfamily of RPTKs in vertebrates is the Eph family, with at least 14 distinct members, and 8 ligands, now termed the 'Ephrins' (reviewed by Gale and Yancopoulos, 1997). Ephrins are all membrane-anchored via either a phospholipid anchor or a transmembrane domain. Many of the Eph receptors and ligands are expressed in the developing nervous system. Several of the lipid-anchored Eph ligands have been implicated as contact repellents that regulate axon fasciculation (Callahan et al., 1995; Tessier-Lavigne, 1995) and retinotectal topographic map formation (e.g. RAGS and Elf-1; Cheng et al., 1995; Drescher et al., 1995).

Although much progress has been made in identifying and elucidating the properties of the different types and families of molecules that influence the direction of axon outgrowth, it is largely unknown how axon growth cones interpret these signals. This is primarily due to the fact that the molecular identity of the receptors that mediate growth cone responses to guidance cues are still unknown, with the exception of the Eph receptor tyrosine kinases. It is generally thought that receptor-ligand interactions on the surface result in activation of intracellular signaling pathways that induce changes in the cytoskeletal organization in the growth cone, leading to neurite extension in the proper direction. Little is known about the second messengers involved in the intracellular signaling pathway. Calcium has been implicated in the regulation of a wide range of growth cone behaviors (Kater and Mills, 1991; Cypher and Letourneau, 1992). Most Ca2+ signaling is transduced by calmodulin (CaM). In Drosophila embryos, a selective disruption of calcium-CaM function in a specific
subset of growth cones was achieved through tissue specific expression of CaM antagonists. This disruption resulted in dosage-dependent stalls in extension and errors in axon guidance (VanBerkum and Goodman, 1995). The results of this study demonstrated an in vitro function for calcium-CaM signaling in growth cone guidance.

It has been shown that CAMs stimulate neurite outgrowth by activating fibroblast growth factor receptors (FGFR; reviewed by Doherty and Walsh, 1996). Activated FGFR triggers a presently unknown second messenger system involving tyrosine phosphorylation and increased calcium influx into neurons. This presumably leads to the activation of calcium-calmodulin dependent protein kinase, which may in turn be responsible for the phosphorylation of proteins required for axon motility. The intracellular proteins which are strongly implicated in the regulation of cytoskeletal rearrangement are the Rho family of small GTP-binding proteins, which includes Rho, Rac and Cdc42 (Nobes and Hall, 1995).

Transcription factors and neuronal connectivity
A number of transcription factors have been identified that regulate specific pathway selections by navigating axons in the developing nervous system. This is achieved through temporal and spatial regulation of expression of external guidance cues and molecules that allow a growing axon to respond appropriately to the cues it encounters (Lundgren, et al., 1995; Mastick, et al., 1997; Retaux and Harris, 1996; Bossing, et al., 1996; Ginger, et al., 1994).
Axon outgrowth occurs following final cell division. Thus, a transcription factor expressed solely in postmitotic neurons is more likely to be involved in pathfinding decisions of a neuron than those expressed earlier in neural cell lineages. These factors either control expression of receptors, or intracellular signaling components that are activated by the external guidance cues. Examples of such regulatory factors include the family of LIM domain containing proteins that carry a homeodomain and a cysteine-histidine rich LIM domain (Tsuchida, et al., 1994; Tokumoto, et al., 1995; Lundgren, et al., 1995). Examples of transcription factors that are implicated in the regulation of expression of external guidance cues include Pax-6 (Mastick, et al., 1997), engrailed (en; Retaux and Harris, 1996), and others (Bossing, et al., 1996).

The family of LIM domain proteins have generated much interest as possible regulators of axon pathfinding because of a close correlation seen between the expression pattern of family members and the projection phenotype of motor neurons in the spinal cord of chick and zebrafish (Tsuchida, et al., 1994; Tokumoto, et al., 1995). The role of a LIM homeobox gene called apeterous (ap) in axon pathfinding has been examined in Drosophila (Lundgren, et al., 1995). ap is expressed in a subset of interneurons that normally choose a single projection pathway in the ventral nerve cord. These neurons send out axons that grow towards the midline, but prior to reaching the midline the axons fasciculate with each other and extend anteriorly in the ipsilateral connective. In the absence of ap function, these axons fail to fasciculate normally and extend along the appropriate pathway. This specific mutant phenotype together with a restricted pattern of expression of the ap gene seen in the postmitotic interneurons, led
investigators to propose a direct role of ap in regulating the expression of cell
surface molecules that mediate recognition events necessary for the formation of
the ap fascicle (Lundgren et al., 1995).

In the mouse CNS, the LIM and homeodomain containing protein Pax-6
regulates the formation of the first prosencephalic tract called t poc (Mastick, et
al., 1997). T poc originates as axons projecting caudally from a cluster of neurons
located at the base of the optic stalk. In wild-type embryos, the developing tract
comes in close contact with a superficial patch of Pax-6 expressing neurons. In
Pax-6 mutants, the projections of the tract forming axons are disrupted in the
region where these axons contact the Pax-6 expressing neurons. The
differentiation of the Pax-6 expressing neurons is not affected in the mutant.
These observations suggest a role for Pax-6 in providing local positional
information for proper guidance of axons that form the first prosencephalic tract.

The en gene codes for a homeodomain containing transcription factor. A
gradient of en expression is seen in the mesencephalon that correlates with the
topographic projections of optic axons (Retaux and Harris, 1996). A role for en in
guiding optic axons has been demonstrated through various manipulations of
the en gradient that lead to the disruption in the formation of retinotectal
topographic map. Interestingly, the guidance function of en appears to be
mediated by several candidate cell surface molecules, notably the Eph ligands
that are upregulated by en (Logan, et al., 1996).

Another transcription factor that regulates both axon growth and
guidance is longitudinals lacking (lola; Ginger, et al., 1994). The lola gene is a
member of the tramtrack family of proteins with two zinc finger domains.
Mutations in the *lola* gene leads to fused commissures in *Drosophila*. Two different isoforms of *lola* have been shown to be expressed in two different cell-types, the navigating axons and the tissue (midline glia) that provides guidance cues to these axons. Identifying the mechanistic difference in the mode of action of the two isoforms of *lola* will thus be important in understanding the difference between the two pathways regulating axon growth and guidance.

It is likely that in the near future the role of previously known transcription factors as regulators of neuronal pathway formation will emerge along with the identification of new transcription factors. Spatial and temporal control of the activity of these regulators along with characterization of their targets will provide valuable information on axon growth and guidance.

**Genetic analysis of growth cone guidance:**

The fact that the behavior of growth cones can be influenced by both attractive and repulsive factors, either by contact-mediated or diffusible mechanisms, is well established from *in-vitro* studies (reviewed by Goodman, 1996). An understanding at the molecular level of how these different forces are coordinated *in-vivo* to establish the stereotypical patterns of synaptic connections of a complex nervous system remains, however, largely elusive. A powerful approach to studying this problem is to use genetics to identify genes necessary for *in vivo* axon guidance. Two methods of genetic analysis can be used to identify and study *in vivo* function of molecules that control growth cone guidance. One approach is the so called reverse genetic approach which has been used in a variety of organisms to test the function of previously identified
molecules through disruption of gene function or misexpression studies. The other approach is the classical genetic approach, in which a large-scale mutagenesis is performed. Mutations that result in defects in axonal projections may define genes involved in establishing neuronal connectivity in the intact organism. By doing large-scale systematic screens it is theoretically possible to identify all genes involved in a given developmental pathway. The proteins identified in this fashion could function in any step in the process (i. e., regulators, signals, receptors, effectors). An innovative classical genetic approach is an interactive genetic screen in which one begins with a mutation in a particular gene that creates a sensitized background (using either partial loss-of-function or gain-of-function mutants) and then screens for mutations in other genes that either enhance or suppress this phenotype. Such screens can be used to identify upstream and downstream components in a genetic pathway (e. g. Simon et al., 1991). Until recently classical genetic approaches were restricted to invertebrate systems, most notably the nematode and Drosophila. However, large-scale mutant screens have now been conducted in the lower vertebrate Zebrafish. Using anatomical probes to search for defects in the retinotectal projection, a number of interesting mutants have been identified (reviewed by Kuwada, 1995).

**Genetic analysis in Drosophila:**

There are a number of reasons for using Drosophila as a model system to unravel complex biological processes. The most compelling is the power of the genetic approach. This is best exemplified in the genetic screen of Nusslein-Volhard and
Wieschaus (1980) from over a decade ago, which revolutionized our understanding of early embryonic pattern formation.

Behavioral assays have been used in *Drosophila* identifying mutations that disrupt axon pathfinding and cellular connectivity in the nervous system. Two such screens were based on the jumping behavior of flies in response to light-off stimulus (Thomas and Wyman, 1984) and grooming behavior (Phillis *et al.*, 1993). The screen for jumpless mutants led to the identification of two mutants (*bendless* and *passover*) that disrupt connectivity between an identified paired neuron (GF neuron) and a specific jump muscle motoneuron (the TTMmn, tergotrochauteral jump muscle motor neuron).

The *bendless* gene, that encodes a ubiquitine conjugating enzyme, is required for normal pathfinding/target recognition by the GF axons; while, the *passover* gene is required for function and maintenance of synaptic connections between GF and TTMmn neurons (Thomas and Wyman, 1984). The *passover* gene encodes a transmembrane protein similar to the *Drosophila* ogre and C. *elegans* Unc-7 proteins (Krishnan *et al.*, 1993). Unc-7 is also required for proper neuronal connectivity in C. *elegans* (Starich, *et al.*, 1993). Later studies have shown that the *passover* gene is also necessary for gap junction communication between neurons of the *Drosophila* giant fibre system (Crompton *et al.*, 1992; Krishnan *et al.*, 1993), the group of eight neurons that relay excitation from the eyes to the muscles of the thorax.

Using anatomical probes, three large-scale histological screens have been conducted in *Drosophila* to identify molecules required for axon guidance and neuronal connectivity. These screens were aimed at studying CNS axon
pathways (Seeger et al., 1993), neuromuscular connectivity (Van Vactor et al., 1993) and retinotopic connectivity (Martin et al., 1995).

The midline of bilaterally symmetric CNS from arthropods to vertebrates represents a boundary that has major influences on growth cone behavior. A systematic genetic screen using a general CNS axon marker (MAb BP102) was conducted to identify mutations that affect CNS patterning in Drosophila (Seeger et al., 1993). Two mutations, commissureless (comm) and roundabout (robo), were isolated that severely affected axonal growth across midline. Mutations in the comm gene resulted in absence of nearly all CNS axon commissures. Axons that normally project across the midline, such as RP1 and SP1 (two identified commissural neurons), failed to do so in these mutant embryos. Instead these neurons sent ipsilateral projections which are normal in all respects except for being on the wrong side of the embryo. The comm mutant does not cause abnormality in the midline structure or disrupt any other axon pathway that may be important for commissural axons outgrowth towards the midline. This suggests that the comm gene is a good candidate to act either as a signal or a receptor that guide axon growth cones towards the midline.

Mutations in the gene roundabout (robo) lead to a phenotype opposite to that of comm. In robo mutant embryos, axons that are close to the midline, but do not normally cross it, are inappropriately attracted towards the midline and their contralateral homologues on the other side. On the basis of this mutant phenotype a role of robo in directly or indirectly generating repulsive cues at the midline for the ipsilaterally projecting neurons has been proposed. Furthermore, the observation that the comm phenotype is suppressed in comm-robo double
mutant suggests that these two systems do not act independently. Instead, there may be a balance between attractive and repulsive cues that affect growth cone behaviour in the midline (Seeger, et al., 1993).

Mutations in two other genes (fasciclin I and abelson) isolated in the same screen led to the identification of two redundant pathways that attract commissural axons outgrowth towards midline (Elikins, et al, 1990). Mutation in either gene does not reveal any obvious commissural axon pathfinding defect; however, a phenotype similar to that of comm mutant is seen when both genes are mutated simultaneously.

The ability of motoneurons to find and recognize their appropriate muscle targets has long been an excellent model system for studying axon pathfinding and target recognition. Using an antibody probe that identifies motoneuron growth cones and axons (MAb 1D4), a screen of the second chromosome for mutations that disrupt motoneuron pathfinding and target recognition was conducted (VanVactor et al., 1993). The screen led to the identification of several genes that are necessary for motoneurons to navigate past particular choice points in the pathfinding process, e.g. beaten path, stranded and short stop. In addition, two genes (walkabout and clueless) required for target recognition by a group of motoneurons, have also been identified in this screen.

In the retinal connectivity screen (Martin et al., 1995), a lacZ reporter gene (glass-lacZ) specific for photoreceptor neurons was used to identify mutations affecting retinal axon outgrowth, retinotopy and target recognition. The screen led to the identification of four genes that play direct roles in neuronal connectivity. These include genes regulating axonal outgrowth (e.g. eddy), target
recognition (e.g. *limbo* and *nonstop*) and retinotopy (e.g. *limbo*). Some of these
genes appear to encode components of the growth cone itself (*limbo* and *diva*),
while others may act as extracellular cues for growth cone guidance (*eddy* and
*nonstop*).

Reverse genetics has provided a complementary approach in *Drosophila* to
identify guidance molecules. In the past, several molecules implicated in growth
cone guidance have been identified in *Drosophila* based on their pattern of
expression (using MAb or enhancer trap screens), or sequence similarity to
known guidance molecules (e.g., Grenningloh *et al.*, 1991; Nose *et al.*, 1992;
Klambt *et al.*, 1991). The specific function of these molecules was subsequently
studied using a reverse genetic approach. For example, three receptor-like
tyrosine phosphatases that are expressed in the embryonic CNS have been
cloned based on sequence similarity to vertebrate proteins (Nose *et al.*, 1992,
and semaphorin II) are expressed by subsets of muscles during the establishment
of neuromuscular connections. These proteins contain immunoglobulin-like
domains and fibronectin type III repeats in the extracellular domain, protein
motifs often associated with recognition/adhesion proteins. Additionally, the
obvious signaling potential of the cytoplasmic tyrosine phosphatase domain of
these proteins, raise intriguing questions about the function of these proteins
during growth cone guidance. Ectopic expression studies showed that all three
proteins can function in growth cone guidance: connectin as a bifunctional signal
(Nose *et al.*, 1994), fasciclin as an attractive signal (Chiba *et al.*, 1995) and
semaphorin as an inhibitory signal (Matthes *et al.*, 1995).
Genetic analysis in zebrafish:

Until recently, large-scale genetic screens have not been practical in vertebrates. Unfortunately, applying reverse genetics to study just the molecules which have already been identified limits the extent of the investigation. Additionally, when functions of multiple proteins overlap in guidance of a particular axon pathway, knock out of a gene corresponding to one of these proteins with overlapping function might not lead to an obvious pathfinding phenotype, even though the gene might be expected to play a role in axon guidance. By employing forward genetics in zebrafish it has become possible to overcome these problems and to analyze in vivo axon guidance systematically in a vertebrate system for the first time.

The convergence of biochemical approaches in vertebrates and genetic approaches in invertebrates has shown that some axon-guidance mechanisms are conserved to a remarkable degree (Goodman, 1994). At the same time, given the complexity of the vertebrate CNS, it seems likely that evolution has provided vertebrates with novel guidance mechanisms or has used conserved guidance cues in novel pathways. Thus it is important to study axon guidance systematically in vertebrates as has been done in invertebrates. The refinement of zebrafish as a genetic system (Mullins et al., 1994; Grunwald and Streisinger, 1992) has made it possible to apply a systematic genetic approach to the study of axon guidance in vertebrates.

A mutant screen was performed on zebrafish larvae in search for defects in axon projections between the eye and its main target in the brain, the optic
tectum (Baier et al., 1996; Trowe et al., 1996; Karlstrom et al., 1996). In addition to finding several genes specifically involved in retinotectal topography, a large number of genes were found that affect axon guidance more generally. Four classes of mutants were identified in this screen. In one class of pathfinding mutants, retinal axons fail to cross the ventral midline of the brain. These neurons connect to the ipsilateral rather than contralateral tectal lobe. Five of these mutations (chameleon, detour, iguana, umleitung and you-too) affect differentiation of midline structures. These mutants demonstrate the importance of midline cells in retinal axon guidance. Another class of mutations that affect the ability of retinal axons to turn toward the contralateral tectal lobe after they reach the midline include bashful, grumpy and sleepy. In these mutants, after axons cross the midline they often turn anteriorly and grow along the margin of the telencephalon.

Three mutations that affect sorting of retinal axons in the optic tract were identified. These are boxer, dackel and pinscher. In these mutants, dorsal axons grow through both dorsal and ventral nerve bundles instead of selectively using the ventral bunch and arrive at the dorsal side of the tectum. These axons then traverse the tectum to find their correct ventral targets. Further molecular analysis of these genes will elucidate the process of fibre sorting, a phenomenon that is poorly understood.

Four mutations, nevermind, who-cares, gnarled and macho affecting the mapping of the retinal fibres have also been identified in this screen. The first two affect the retinal axons mapping along the dorsoventral axis, while gnarled
(gna) and macho (mao) affect it along the anterior-posterior axis. In gna and mao Nasodorsal axons defasciculate and terminate too soon in the anterior tectum.

A general finding of the screen in zebrafish is that most of the retinotectal mutants show other developmental defects. This indicates that molecular mechanisms used for axon guidance are also used in other developmental processes. In addition, genes absolutely necessary for early development can not be found in the screen. Affected embryos do not develop to a stage where axon guidance could be assayed. A genetic screen designed to find conditional, temperature-sensitive mutations may provide an alternative way to find more of the genes involved in axon guidance.

Organization and neuronal connectivity pattern in the Drosophila visual system

The Drosophila visual system offers an excellent model for the investigation of neuronal patterning. The advantages include, relative simplicity, availability of numerous molecular markers which can be used to identify and analyse individual neurons (Bellen et al., 1989; Bier et al., 1989; Wilson et al., 1989; Zipursky et al., 1984), availability of behavioral tests (phototaxis and optomotor behavior), opportunity for genetic analysis and most importantly, the remarkable similarity in the mechanism of neuronal growth cone guidance in insect and mammals (Harrelson and Goodman, 1988; Klose and Bentley, 1989; McConnell et al., 1989).

Drosophila displays two distinct visual systems during larval and adult stages. The adult visual system consists of compound eyes and the optic lobes,
the portion of the visual system that receives and processes visual information from the eyes (reviewed by Wolff and Ready, 1993; Meinertzhagen and Hanson, 1993). The compound eyes are composed of approximately 800 ommatidia, each containing eight photoreceptor cells. The photoreceptor cells differentiate from the eye imaginal disc during third instar larval stage (Ready et al., 1976). These cells send their axons to the optic lobe primordium through a specialized epithelium tube called optic stalk. Retinotopy is seen in the adult visual system, i.e., retinal axons send their axons to the corresponding positions in the brain. An examination of photoreceptor projections in mutants containing fewer number of these cells and in mosaic animals containing patches of wild-type axons in a mutant background, has shown that each photoreceptor axon is independently guided to its target cell (Kunes et al., 1993). These findings suggest a model similar to Sperry’s chemoaffinity hypothesis where positional guidance and recognition cues label pathways and synaptic targets (Sperry, 1963).

All three components of the Drosophila visual system; the adult eye, larval eye and the optic lobe originates from a small region of the procephalic lobe (embryonic head; Green et al., 1993). The optic lobe and the larval photoreceptor cells arise from within a coherent region of the posterior head ectoderm, while the eye imaginal disc develops from two pouches of the anterior ectoderm. Each primordium of adult optic lobes develops from a mass of tissue that invaginates from the posterior portion of the procephalic lobe during stage 12 of embryogenesis. The optic lobes form in direct contact to the developing larval brain and by late embryogenesis this tissue completely incorporates into the
brain hemispheres. The optic lobe primordium remains undifferentiated until larval stage, when it resumes mitosis and differentiates to form the optic ganglia (Meinertzhagen, 1973; White and Kankel, 1978).

Although the tissue origin for the eye imaginal disc and the optic lobe primordium are different from each other, the two tissues interact extensively during development. The differentiation of the first order interneurons of the first ganglion layer (lamina) of the optic lobe is dependent on photoreceptor axon innervation (reviewed by Meinertzhagen and Hanson, 1993). However, the dependence of the second order neurons in the second optic ganglion (medulla) upon retinal innervation is less extensive. Retrograde interaction, that is, the dependence of photoreceptor integrity on connection of these cells with their synaptic partners of lamina ganglionaris has also been shown to be a characteristic of insect visual system development (Campos et al., 1992).

In comparison to the adult visual system, the larval visual system is extremely simple, consisting of two bilateral clusters of 12 photoreceptor cells. These cells extend their axons in a fascicle (the Bolwig's nerve; Bolwig, 1946) towards the larval visual system target area located within the anlage of the adult optic lobes (Green et al., 1993). The Bolwig's organ develops at the ventral tip of the invaginating optic lobe primordium. These cells send short axonal projections that contact the cells of the neighboring optic lobe invagination (Steller et al., 1987; Schmucker et al., 1992; Green et al., 1993). At stages 15-16, during head involution, the larval photoreceptor cells begin to migrate out of the head epidermis by delamination. While these cells migrate anteriorly their axons retain connection with the optic lobe primordium (Green et al., 1993). During
this period the Bolwig's nerve which is formed from fasciculation of the 12 photoreceptor axons undergo extensive elongation. Later in embryogenesis, the Bolwig's nerve terminal extends deeper into the brain, past the optic lobe primordium and terminates at the posterior margin of the optic lobe primordium (Green et al., 1993; Campos et al., 1995). The cells in the invaginating optic lobe primordia that contact the larval optic nerve act as intermediate targets for the larval photoreceptor cells, analogous to the guidepost cells discussed earlier (Campos et al., 1995).

Some of the cell types that are contacted by the larval optic nerve during larval visual system development have been identified, among the most interesting are the three optic lobe pioneer cells (OLPs; Tix et al., 1989; Campos et al., 1995). Two of these cells that originate from the optic lobe primordium itself are called the corner OLPs. The one that originates from outside this region is called the central OLP. This cell is contacted by the LON in the final stage of nerve pathway formation. The axon of this cell extends towards the central brain into the larval optic neuropil before it contacts the LON (Campos et al., 1995). The LON and the axons from the two corner OLPs fasciculate with the central OLP axon and together project towards the central brain. Thus the central OLP is thought to pioneer the route taken by the LON toward the central brain. On the other hand the corner OLP cells that contact the LON at initial stages of development before invagination of the optic lobe primordium are thought to act as intermediate targets for the LON and are thus analogous to the guidepost cells discussed earlier (Campos et al., 1995). The OLP cells are also incorporated into
the adult optic lobe and are thought to play a role in organizing the adult visual system as well (Fischbach and Technau, 1987).

The *Drosophila disconnected* gene and its role in the establishment of neuronal connections in the visual system

The *disconnected* (*disco*) gene codes for a putative transcription factor with two C2H2 type zinc finger motifs (Heilig *et al.*, 1991). The zinc finger proteins represent a major group of transcription factors that play an important role in many aspects of eukaryotic gene regulation. Proteins of this family contain tandem repeats of 30 amino acids zinc finger motifs enriched in Cysteine and/or Histidine residues (reviewed by Berg, 1990).

The importance of the zinc finger domains in *disco* function was demonstrated by the fact that mutations in the conserved Cysteine residues of either zinc finger motif result in a visual system phenotype similar to that seen in mutants in which the *disco* gene is deleted (Heilig *et al.*, 1991). Since two zinc finger domains are sufficient for sequence specific DNA binding activity (Keller and Maniatis, 1992), it is hypothesized that *disco* is a DNA binding protein. Nuclear localization of the DISCO protein and an autoregulatory activity of *disco* demonstrated in the cells of the optic lobe primordium support this hypothesis. A high affinity DNA binding site has also been identified in genomic DNA, 2.5 kb upstream of the *disco* transcription start site in an *in vivo* electrophoretic mobility shift assay (Kevin Lee, Ph. D. thesis, 1993).

*disco* gene function is required for proper neuronal connectivity formation in both larval and adult visual system of *Drosophila*. Mutations in the *disco* gene
cause failure of the larval and adult photoreceptor axons to make proper connections with their target cells (Figure 1; Steller, et al., 1987). In the majority of the adult disco mutant flies the compound eyes fail to innervate the brain and show a significant reduction in the volume of the optic lobes. This phenotype is referred to as the unconnected phenotype (Steller et al., 1987). In other adult mutant flies, the retinal innervation of the optic ganglia occurs, but the optic ganglia in these mutants are roughly normal in size although still grossly disorganized. This phenotype is called the connected phenotype. Since retinal innervation is required for proper development of the optic ganglia, the reduced optic ganglia in the unconnected disco mutant is at least partly due to the lack of retinal innervation. The presence or absence of retinal innervation again depends on presence or absence of the optic stalk, the tube that connects the eye disc to the optic lobe and through which retinal axons enter the optic lobe area of the brain. Individual mutant larvae can lack just one or both optic stalks.

The disco gene is expressed in several of the different cell types contacted by the larval optic nerve including a group of glial cells distributed along the LON and the OLP cells (Lee et al., 1991; Campos et al., 1995). Interestingly, mutation in disco leads to failure in the differentiation of corner OLP cells and mislocation of the glial cells (Campos et al., 1995). Thus, loss of disco function leads to defects in cellular interactions between the LON and the cells it contacts during pathway formation. The absence of disco expression in the photoreceptor cells at least during axon outgrowth indicates that the effect of a disco mutation on the LON phenotype is nonautonomous. The disco gene function is required at least in the OLPs or LON glial cells for proper LON connectivity. It may regulate
cellular guidance cues and recognition cues that are specific to the development of the larval optic nerve pathway and connectivity.

**Questions and aims of this thesis**

The goal of the work presented in this thesis is to study some of the cellular and molecular mechanisms underlying nervous system development. To that end I focused on the development of neuronal pathways in a simple model system, the larval visual system of *Drosophila melanogaster*. The simplicity of the larval visual system makes it amenable to cellular biological studies, and the classical genetic approaches that are uniquely applicable to *Drosophila* research together with a recent advent in molecular biological tools made the visual system of *Drosophila* an attractive model system for neurobiological studies.

The first part of my thesis described identification and characterization of specific cellular interactions at the larval optic center of *Drosophila*. The outcome of this study may suggest that changes in behavior of an organism that occur as a function of normal development require establishment of new neuronal connections. The second part of my thesis addressed the role of the *disconnected* gene in establishing proper neuronal connections in the visual system of *Drosophila*. The tissue specific autoregulatory activity of *disco* in the cells of the optic lobe primordium, and the differential effect of *disco* mutations on the *disco* expressing cells contacted by the LON, reveal the importance of tissue specific *disco* function in establishing proper LON connectivity. Therefore, identification of tissue specific factors regulating *disco* function will provide us with valuable
information on the role of the disco gene in Drosophila visual system development. In this part of my thesis, I used the yeast interaction trap approach to identify DISCO interacting proteins.

Chapter 2 of this thesis describes the identification of a serotonergic neuronal process and the influence of the LON on the outgrowth and establishment of arborization from the newly identified process at the larval optic center. This study led to the speculation that the establishment of connections between the LON and the serotonin process modulates larval photoresponse.

Chapter 3 describes the identification of three Drosophila proteins (DISCO-interacting proteins DIP1, DIP2 and DIP3) that interact with DISCO in yeast interaction trap and in vitro binding assays. Among these two are newly identified proteins, one of which contains two putative double stranded RNA-binding domains (DIP1) and the other is a highly conserved protein (DIP2) of unknown function. The known DISCO interacting protein (DIP3) is the putative ATPase component of the 26S proteasome. Expression of the DIP1 and DIP2 proteins partially overlap with that of disco, and therefore, these are good candidates to act as components of disco mediated pathway that regulate larval visual system connectivity formation.

Northern analysis and analysis of dip1 cDNAs isolated from various cDNA libraries indicate expression of three alternatively spliced forms of the dip1 mRNA. Repetitive sequences in the 3' untranslated region of these transcripts may confer translational regulation and therefore may suggest a less extensive expression pattern of the DIP1 protein than its mRNA. Based on the
chromosomal location and the deficiency map of the dip1 gene, a putative mutant (flamenco, flam) for this gene has been identified. The physiological role of the gene affected in flam mutant is presently unknown; however, the effect of the hypomorphic mutant allele on the expression and mobilization of a retroviral element in Drosophila ovary might support a RNA binding activity of the gene.

The human and C. elegans homologs of the dip2 gene have been cloned in the corresponding genome projects. The work presented in this thesis describes the isolation of the Drosophila and mouse homologs of this gene. The putative DIP2 protein lacks any known functional domain. Although no functional analysis has yet been done on any dip2 homologs, nervous system specific expression of the Drosophila and the mouse homologs during embryogenesis suggest an important role for dip2 in nervous system development.

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Chapter 2: Manuscript

THE LARVAL OPTIC NERVE IS REQUIRED FOR THE DEVELOPMENT OF AN IDENTIFIED SEROTONERGIC ARBORIZATION IN DROSOPHILA MELANOGASTER.

Preface

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The Larval Optic Nerve Is Required for the Development of an Identified Serotonergic Arborization in Drosophila melanogaster

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The larval visual system in the fruitfly Drosophila melanogaster consists of two bilateral clusters of 12 photoreceptor cells. These neurons send their axons in a fascicle, the Bolwig's nerve, toward the target area in the ventral lateral region of the brain hemispheres. We describe the development of a serotonergic arborization originating in the central brain found in the larval optic center in association with the larval optic nerve. This arborization is formed by processes from larval neurons born during embryogenesis. However, these neuronal processes do not reach their final destination, the larval optic center, until late in larval development. Using mutations that disrupt the connectivity and/or development of the larval photoreceptor cells, as well as mosaic analysis, we demonstrate that the innervation of the larval optic center by this serotonergic arborization depends upon contact with the larval optic nerve.


INTRODUCTION

A fundamental aspect of nervous system development is the precise interconnection between different cell types. That precise assembly of a complex network depends not only on the matching of pre- and postsynaptic cell populations but also on the development and maintenance of the appropriate axonal and dendritic arborizations.

The dependence for development and maintenance of neuronal arbors on connections with other neurons is well documented in several different model systems (e.g., Loer and Kristan, 1989; Antonini and Stryker, 1993; Byrd and Burd, 1993). While several molecules required for axonal guidance have been described (reviewed by Goodman and Shatz, 1993), comparatively little is known about the nature of the intercellular signals required for the local induction and maintenance of dendritic and axonal arbors.

The role of activity of afferent synaptic inputs in the development and remodeling of dendritic arbors has been extensively examined. Neuropeptides contained in afferent innervation have been shown to regulate dendritic growth (reviewed in Schwartz, 1992). Several reports demonstrate that patterned afferent excitatory activity results in structural changes of the corresponding dendritic arbors (Tieman and Hirsch, 1982 and Katz and Constantine-Paton, 1986). Activation of N-methyl-D-aspartate subtype of glutamate receptors is involved in motor neuron dendritic maturation in early postnatal life (Kalb, 1994). Endogenous electrical activity has also been shown to influence dendritic growth patterns in Purkinje cells in primary dissociated cultures of mouse cerebellum (Schilling et al., 1991).

Cell interactions are also likely to play a role in the development of neuronal arbors. Regulation of axonal and dendritic growth by glial cells has been demonstrated by a variety of coculture experiments (Frohnaltz et al., 1990; Rousselet et al., 1990; and Qian et al., 1992). Recently, it has been suggested that glia-mediated dendritic growth is promoted by a diffusible factor (Le Roux and Reh, 1994). Beyond its role in the control of cell number, NGF has also been shown to influence the degree of axonal branching locally both in vivo and in vitro (Campenot, 1982a; Campenot, 1982b; De Konnick et al., 1993; Van der Zee et al., 1995). The visual system of the holometabolous insect Drosophila melanogaster has been successfully used as a model system for the study of cell interactions during development. Drosophila displays two visual systems during its life cycle: the adult and the larval visual systems. The adult visual system consists of compound eyes containing photoreceptor neurons that project to the underlying optic lobes (reviewed by Wolff and Ready, 1993, and Meinertzhagen and Hanson, 1993). The larval visual system is considerably less complex, consisting of two bilateral clusters of 12 photoreceptor cells which send their axons in a fascicle (The Bolwig's nerve, Bolwig, 1946) toward the larval visual system target area located within the anlagen of the adult optic lobes.
Cell interactions have been shown to play a fundamental role in the development of the adult visual system in the fruitfly Drosophila melanogaster (reviewed by Meinertzhagen and Hanson, 1993, and Wolff and Ready, 1993). This assertion is well exemplified by reports on the dependence of adult optic lobes on afferent innervation (reviewed by Meinertzhagen and Hanson, 1993, and Power, 1943; Meyerowitz and Kankel, 1978; Fischbach, 1983; Fischbach and Technau, 1984; Hofbauer and Campos-Ortega, 1990; Selleck and Steller, 1991). In particular, the first optic ganglion, the lamina, requires retinal innervation not only for mitosis of the lamina precursor cells but also for proper morphogenesis of the lamina cartridges (Meyerowitz and Kankel, 1978, and Selleck and Steller, 1991). The dependence of the second order neurons in the medulla ganglion upon retinal innervation is less extensive than that found in the lamina ganglion. Several optic lobe neurons are able to survive deafferentation, although they display sprouting and compensatory innervation (Fischbach, 1983; Fischbach and Technau, 1984).

The larval optic nerve is apparently not required for the establishment of the optic stalk nor for the targeting of the retinal projections to the developing optic lobes, the OLPs (Moses et al., 1988; Kunes and Steller, 1991). The connectivity of the larval optic nerve to the optic lobe anlagen, however, may depend upon the presence of the optic lobe pioneer cells (Tix et al., 1988; Campos et al., 1995). The differentiation of OLPs precedes the final step in the establishment of connectivity of the larval optic nerve to the optic lobe primordium and is apparently not dependent upon contact with the larval optic nerve (Campos et al., 1995).

A role for the larval optic nerve in the development of adult optic lobe projections is suggested by the observation that certain identified neurons which project from the central brain toward the lamina (the LBO 5-HT, Nassel, 1987) follow the same pathway pioneered by the larval optic nerve during larval stages. The projections from these serotonergic cells as well as the OLPs occupy the posterior optic tract which contains axons of the lamina and medulla tangential cells (reviewed in Meinertzhagen and Hanson, 1993). Additionally, it is possible that larval retinal innervation is required for the development of the larval visual system target cells, similar to what is observed in the adult visual system. These target cells may in turn be incorporated into the developing adult optic lobes and there participate in the organization of the optic lobes during development.

Here, we report the consequences of deafferentation of the larval optic center. We focus our analysis on the development of an identified serotonergic arborization that projects from the central brain. Using mutant strains that impair the connectivity of the larval optic nerve to the larval neuropil center as well as mosaic analysis, we conclude that neither the cell viability nor the initial outgrowth of this projection depends on the presence of the larval optic nerve. However, the arborization of this projection within the larval optic neuropil requires the presence of the larval optic nerve. Interestingly, this serotonergic process does not contact the larval optic nerve until late in larval development.

**MATERIALS AND METHODS**

**Drosophila Strains and Culture**

Flies were grown at 25°C or 19°C in inactivated yeast, sucrose, and agar medium supplemented with fresh active yeast. Ten percent tegoept in ethanol was used to prevent microbial growth. Eggs were collected on molasses-agar egg-collection plates and aged as described by Ashburner (1989).

$gf^{om}$ Null mutant allele of the glass ($gf$) gene. Expression of the glass gene is disrupted by an insertion (>30 kb) in the coding region (Moses et al., 1989).

$disco$. Ethyl methanesulfonate-induced allele of disconnected (disco) gene (Fischbach and Heisenberg, 1984). In disco a point mutation (cys127 to ser) in the coding region of the disco gene resulted in a missense mutation (Heilig et al., 1991). In this mutant, the larval optic nerve cannot establish proper connections with its target cells during embryonic development (Steller et al., 1987).

$P[gf^* (10 kb Sal), rf^+]$; $gf^{om}$. The third chromosome is homozygous for the $gf^{om}$ mutation, while the first chromosome is homozygous for a P-element insert carrying a normal glass gene and a wild-type rosy gene (Moses et al., 1989).

**Delta 2-3, $gf^{om}$**. The third chromosome is homozygous for the $gf^{om}$ mutation and carries the gene for a transposable enzyme which can mobilize P-element inserts. However, the transposable element that carries the transposable gene is stable in this position (Laski et al., 1986; Robertson et al., 1988).

**C50.151: A disco-enhancer trap line, in which expression of a $\beta$-galactosidase gene is driven by the disco promoter. Thus, $\beta$-galactosidase expression is localized to the disco expressing cells and acts as a marker for disco expression (Cohen et al., 1991; Heilig et al., 1991, and Lee et al., 1991).**

**Immunocytochemistry of Larval Brain**

Larval brains of different larval stages were dissected in Drosophila Ringer's solution and fixed in 4% paraformaldehyde (pH 7.4) for 4 hr or overnight at 4°C.
Fixed brains were washed several times in 0.1 M phosphate buffer containing 0.3% Triton X-100 (PBT) and blocked with PBT containing 5% goat serum for an hour at room temperature. Incubation with primary antibody (monoclonal or polyclonal) diluted with PBT containing 5% goat serum was performed overnight at 4°C. After several washes with PBT for 2 hr at room temperature the brains were again blocked with PBT + goat serum and then incubated with secondary antibody for 5 hr at room temperature. This was followed by washing the samples several times in PBT for 2 hr. Enzymatic reactions were carried out for Horseradish peroxidase-conjugated secondary antibodies using diaminobenzidine and hydrogen peroxide as described by Steller et al. (1987). In some cases 2.5 mM CoCl2 was used to intensify the signal. For fluorescence-tagged secondary antibodies brains were mounted in 70% glycerol containing p-phenylenediamine.

The primary antibodies used were rat monoclonal anti-serotonin, clone YC5/45 (Consolazione et al., 1981), a mouse monoclonal anti-CHAOPTIN (Zippuraki et al., 1984), and a mouse monoclonal anti-GLASS (Ellis et al., 1993). The secondary antibodies used were Cy3-conjugated goat anti-rat IgG (Jackson ImmunoResearch Laboratories, Inc.), FITC-conjugated goat antimouse IgG (Cappel), and Peroxidase-conjugated goat anti-mouse IgG (Jackson ImmunoResearch Laboratories, Inc.).

The specimens were viewed under Nomarski optics in a Zeiss AxioScope. Confocal microscopy was performed on a Bio-Rad MRC 600 Krypton/Argon laser confocal microscope.

Generation of glas Somatic Mosaics

gl mosaics were created using the same scheme as described by Kunes et al. (1993). Somatic loss of a glas P-element (Moses et al., 1989) in a strain carrying a null mutation in the gl gene was induced by somatic expression of the Delta 2–3 transposase (Laski et al., 1986; Robertson et al., 1988). Females homogygous for the null allele gl and carrying a wild type gl gene in a transposable element inserted in the X chromosome (PIgl(10 kb Sal), rsy1); Moses et al., 1989) were crossed to males PIrsy+, Delta 2–3, gl and in the progeny of this cross the constitutive source of the transposase mobilized the P-element carrying the normal gl gene in some somatic cells, thereby uncovering the P-element mutation. Cells which retained the P-element remained wild-type. Excision occurs in 100% of the cases as seen by the presence of mosaic compound eyes in all adult flies.

RESULTS

Serotonin Immunoreactive Processes Are Found Innervating the Larval Optic Neuropil

Little is known about the influence of the larval photoreceptor innervation on the development of the larval visual system neuropil. In this investigation we sought to address this question by examining a newly identified arborization found in intimate association with the termini of the larval photoreceptor axons in the larval visual neuropil of the third instar larvae (Fig. 1). This serotoninergic arborization is found in the third instar larval brain, apparently associated with the larval photoreceptor axons (Fig. 1C–E). It can be easily and reproducibly identified in larval brain whole-mount preparations. It is the only serotoninergic process found in the larval optic lobes at this point in development. Additionally, it emerges at the same level where the LP1 axons enter the central brain. The LP1 cell bodies are located in a midlateral position at the margin of the optic lobes and can therefore be used as a landmark for the identification of the larval optic center serotoninergic innervation (Fig. 1C, empty arrow). Brains from late third instar wandering larvae showed, in addition to the larval optic center serotoninergic projection, fibers from the

![Figure 1: Localization of the 5-HT immunoreactive arborization in the third instar larval brain. All photomicrographs depict a horizontal view of the larval brain. (A) A line drawing of a third instar larval brain showing 5-HT immunoreactive neurons and their projections. There are four prominent serotoninergic cell clusters in the brain: SP1, SP2, LP1, and IP (Valies and White, 1983). The 5-HT arborization in the larval optic center (arrowhead) is formed by one or two cells of the SP2 cluster. The SP2 cell cluster is located just posterior to the anterior-must cell cluster SP1 and is composed of four cells. It is also the dorsal-must cell cluster. The LP1 cell cluster is composed of two cells that occupy a midlateral position in the brain. In a horizontal or frontal view of the larval brain the processes of the LP1 cells cross the arborisation forming neuronal processes (arrow) barely seen under conventional microscopy. The best represents the area of interest depicted in all subsequent figures. (B) A third instar larval brain hemisphere double-labeled with the anti-CHAOPTIN antibody (black/brown) to visualise the photoreceptor axons and an anti-5-HT antibody (brown) seen under Nomarski optics. The larval optic nerve which traverses the eye disc (ed) and the optic stalk (os) can be seen terminating (open arrow) in the developing optic lobe (ol). The solid arrow points to the thin neuronal process that forms the 5-HT arborisation in the larval optic center, shown in A inside the boxed area (solid arrowhead). (C, D, and E) Confocal micrographs of a third instar larval brain double-labeled with anti-5-HT (rhodamine channel) seen in red in C and E and with anti-CHAOPTIN (FITC channel seen in green in D and E) antibodies. (E) Merge of C and D. The 5-HT arborisation (arrowhead in C) can be seen overlapping with the LON (arrow in D) in E. The LP1 cluster (empty arrow in C) crosses the serotoninergic processes that arborise in the larval optic neuropil. Scale bar in B, 50 μm and in C, D, and E, 25 μm.]

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Fig. 1. Localization of the 5-HT immunoreactive arborization in the third instar larval brain. All photomicrographs depict a horizontal view of the larval brain. (A) A line drawing of a third instar larval brain showing 5-HT immunoreactive neurons and their projections. There are four prominent serotoninergic cell clusters in the brain: SP1, SP2, LP1, and IP (Valies and White, 1983). The 5-HT arborization in the larval optic center (arrowhead) is formed by one or two cells of the SP2 cluster. The SP2 cell cluster is located just posterior to the anterior-must cell cluster SP1 and is composed of four cells. It is also the dorsal-must cell cluster. The LP1 cell cluster is composed of two cells that occupy a midlateral position in the brain. In a horizontal or frontal view of the larval brain the processes of the LP1 cells cross the arborisation forming neuronal processes (arrow) barely seen under conventional microscopy. The best represents the area of interest depicted in all subsequent figures. (B) A third instar larval brain hemisphere double-labeled with the anti-CHAOPTIN antibody (black/brown) to visualise the photoreceptor axons and an anti-5-HT antibody (brown) seen under Nomarski optics. The larval optic nerve which traverses the eye disc (ed) and the optic stalk (os) can be seen terminating (open arrow) in the developing optic lobe (ol). The solid arrow points to the thin neuronal process that forms the 5-HT arborisation in the larval optic center, shown in A inside the boxed area (solid arrowhead). (C, D, and E) Confocal micrographs of a third instar larval brain double-labeled with anti-5-HT (rhodamine channel) seen in red in C and E and with anti-CHAOPTIN (FITC channel seen in green in D and E) antibodies. (E) Merge of C and D. The 5-HT arborisation (arrowhead in C) can be seen overlapping with the LON (arrow in D) in E. The LP1 cluster (empty arrow in C) crosses the serotoninergic processes that arborise in the larval optic neuropil. Scale bar in B, 50 μm and in C, D, and E, 25 μm.
Fig. 4. Expression of the glas and the disco genes relative to 5-HT expression. (A) Third instar larval brain double-labeled with anti-GLASS (blue-black) and anti-5-HT (brown) antibodies (horizontal view where anterior is up in all panels). (B) Right brain hemisphere of the brain shown in A at higher magnification. The anterior glas cell cluster (arrow) is in focus. (C) Same brain hemisphere as in B at a different focal plane. The posterior glas cell cluster (arrow) is in focus. The 5-HT expressing LP1 cells (arrowheads) lie very close to the posterior glas cell cluster; however, there is no overlap of the two patterns of expression. The SP2 cluster which is involved in the formation of the 5-HT arborization at the larval optic center is shown by an empty arrow. (D) Third instar larval CNS of a disco-enhancer trap strain double-labeled with anti-
β-galactosidase antibody for the detection of the disco gene expression (blue-black) and anti-5-HT antibody (brown). (E) Left brain hemisphere of the CNS shown in D. The disco gene is expressed in several scattered cells and in a band of cells (arrow) that wraps around the brain lobe. The 5-HT arborization (arrowhead) at the larval optic center can be seen. (F) Right brain hemisphere of the CNS shown in D. The two 5-HT-expressing LP1 cells (arrowheads) lie very close to the band of disco-expressing cells (arrow). However, there is no overlap between disco and 5-
HT expression. The SP2 cell cluster is surrounded by faint disco-expressing cells (empty arrow). Scale bar in A and D, 100 μm. Scale bar in B, C, E, and F, 50 μm.
LBO 5-HT neurona described in more detail in the Fleshfly (Nassel et al., 1987) (data not shown).

Observations in earlier stages of larval development (see below) suggest that this arborization is formed by two neuronal processes. However, the level of resolution conferred by the confocal microscope does not allow us to determine if the 5-HT processes innervating the larval optic neuropil in the third instar larval brain correspond to two axons only or to assess the ultrastructural nature of the interaction between the larval optic nerve (LON) and the 5-HT arborisation. Our observations in whole-mount preparations suggest that this projection is formed by axons of the contralateral SP2 cell cluster located in the central brain dorsal rim, posterior to the SP1 cell (nomenclature according to Valles and White, 1988) (data not shown). Similar processes were also described in the Fleshfly by Nassel et al. (1987).

Development of the 5-HT Innervation of the Larval Optic Center

In order to determine the time course of the development of the 5-HT innervation of the larval optic neuropil, double labeling of dissected brains from different larval stages with anti-5-HT antibody and with anti-CHAOPTIN antibody was performed.

5-HT neuronal processes whose position is consistent with those that are later found innervating the larval optic neuropil were first visualized during mid first instar larval stage (Fig. 2A, solid arrow). At this time, only a single neuronal process is seen extending toward the larval optic center. During late first instar larval stage a second neuronal process was often seen joining the first one (Fig. 2C, solid arrow). These processes started branching during early second instar larval stage (Fig. 2E, solid arrow). Up to this time the terminus of the LON can be seen lying ventral to the very tip of the 5-HT projections (Figs. 2B, 2D, and 2F, empty arrow). It was during late second instar larval stage that the 5-HT processes were first seen on the same focal plane of the LON, apparently contacting each other (Figs. 2G and 2H, solid arrow and empty arrow, respectively). These 5-HT processes undergo further branching after the first contact with the LON up to the level seen in the middle third instar stage (Fig. 1C, arrowhead).

Absence of the 5-HT Arborization in Mutants Lacking LON Connection with the Larval Optic Center

In order to determine whether the development of the larval neuropil 5-HT arborization is dependent upon contact with the larval photoreceptor axons, we examined visual system mutants in which the connection of the LON to the target area was either absent or impaired. For this, we chose to examine larval brains of strains carrying mutations in either the glass (g) gene or in the disconnected (disco) gene. In larvae homozygous for the null glass gene allele gtm, only a few loosely clustered photoreceptor cells can be seen without formation of a LON (Moses et al., 1989; Schmucker et al., 1992; Campos et al., 1995). Mutations in the disco gene prevent the proper connection of the LON with its target area during embryonic development. In most cases the optic stalk which connects the larval brain to the anlagen of the adult eye is absent in disco mutants and the LON does not enter the larval brain (unconnected phenotype, Steller et al., 1987). A few of the disco mutant larvae carry normal optic stalk and the LON can be seen projecting to the brain. However, in these animals the LON is still unable to establish proper connections with the target area and is often found in abnormal locations (the connected phenotype, Steller et al., 1987; Campos et al., 1995).

In gtm mutant larvae no 5-HT immunoreactivity re-
FIG. 3. Absence of the 5-HT arborization in $p^{\text{im}}$ and $\text{dico}^\text{c}$ third instar larval brains. Confocal micrographs of 5-HT expression in third instar larval brains of $p^{\text{im}}$ and $\text{dico}^\text{c}$ mutant strains. In all panels the serotonergic larval optic neuropil arborization is indicated by an arrowhead at the LPI cell cluster by an arrow. (A) Wild-type. (B) $p^{\text{im}}$. Often in $p^{\text{im}}$ mutants the LPI cells (arrow) show abnormal neuronal processes (empty arrow) but no clear 5-HT arborization can be detected. (C) $p^{\text{im}}$. In a few specimens thin unpatterned 5-HT processes can be seen (arrowhead). Their location is consistent with that of the soma described for the serotonergic processes that innervate the larval optic neuropil of wild-type strains. (D) $\text{dico}^\text{c}$. The 5-HT arborization is mostly absent (arrowhead). This is an unconnected $\text{dico}^\text{c}$ mutant brain where no adult or larval photoreceptor projection was seen (Steller et al., 1987). (E) $\text{dico}^\text{c}$. Again, in this specimen very thin 5-HT processes (arrowhead) can be seen in a different focal plane adult and presumably larval photoreceptor axons were seen (connected phenotype, Steller et al., 1987). (F) $\text{dico}^\text{c}$. The two arrows point to the apparent duplication of the LPI cell cluster. No 5-HT arborization can be seen. Scale bar in A–E, 25 μm.
TABLE I
MOSAIC ANALYSIS

<table>
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<th>Total number of brain hemispheres</th>
<th>Number of brain hemispheres</th>
<th>Anterior glass cell cluster</th>
<th>Posterior glass cell cluster</th>
<th>Larval optic nerve</th>
<th>SHT arborization</th>
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</table>

Assembling the arborization seen in wild-type preparations (Fig. 3A, arrowhead) was found in the larval optic center (Figs. 3B and 3C). In a few cases, thin neuronal processes were seen (Fig. 3C, arrowhead). Additionally, the two 5-HT-expressing LP1 cells often displayed abnormal sprouting (Fig. 3B, open arrow).

Examination of disco3 third instar larval brains showed that regardless of absence or presence of LON, 5-HT immunoreactivity in the larval optic center was always absent (Figs. 3D and 3E, arrowheads). Furthermore, extra 5-HT-expressing cells were often found (Fig. 3F, solid arrows). These cells were seen in the midlateral portion of the brain at the margin of the optic lobes. Their location and their pattern of projection sug-

Fig. 3. Mosaic analysis for the pfr- allele. Putative mosaic larval brains were dissected and labeled with anti-GLASS, anti-5-HT antibodies. The anti-GLASS binding was detected by HRP-conjugated secondary antibodies. CoCl$_2$ was added during the reaction for the detection of glass expression. The anti-5-HT antibody was detected using a FITC-conjugated secondary antibody. Glass were analyzed using Nomarski optics while 5-HT expression was seen under the confocal microscope. The presence of larval visual system was determined by anti-CHAOPTIN antibody detected by a FITC-conjugated secondary antibody (not shown). (A) Wild-type third instar larval brain hemisphere, labeled with anti-GLASS antibody. There are two glass-expressing cell clusters: one anterior (arrow) and one posterior (arrowhead). Horizontal view where anterior is up in all panels. (B) One example of a pfr$-$ mosaic third instar CNS where the left brain hemisphere is completely mutant as seen by the absence of glass gene expression in both the anterior and posterior cell clusters. The larval visual system is present as determined by anti-CHAOPTIN expression (data not shown). In the eye disc (C) the developing adult photoreceptor cells are stained with the anti-GLASS antibody (arrow) demonstrating that (1) this specimen does contain a functional glass gene in areas of the nervous system other than the central brain and (2) immunostaining of the GLASS protein was successful in this sample. (C) Same mosaic brain as in B, showing anti-5-HT antibody labeling as seen in the confocal microscope. The 5-HT arborization (empty arrow) is present. Arrow points to the LP1 cells. This brain lobe is positioned on a slightly frontal plane, therefore the orientation of the LP1 cells appears different from the one shown in previous figures. Scale bar in A and B, 50 µm and in C, 25 µm.
suggested that these extra cells in disco mutants represented duplication of the LP1 cluster.

In order to determine if the absence of the 5-HT arborization in gl" and disco" mutants is due to the absence of the corresponding neurons, we counted the number of all 5-HT-expressing cells in 24 brain hemispheres of wild-type, gl" and disco" mutant strains. Except for the LP1 cells all other 5-HT clusters were found in equal numbers in wild-type and in mutant brains, including the SP2 cell cluster presumed to be the corresponding cell bodies for the 5-HT arborization. No difference was found regarding the general morphology or immunoreactivity level of these neurons in either one of the mutant strains. Thus the absence of the 5-HT larval optic neuropil arborization is not accompanied by a decrease in 5-HT positive cells in disco" or gl" mutants.

Expression of the glass and of the disco Gene Relative to 5-HT Expression

The results described above suggest that the presence of the LON connection is required for the proper innervation of the larval optic neuropil by 5-HT fibers originating in the central brain. It is possible, however, that in both mutants the absence of 5-HT immunoreactivity in the larval optic neuropil is due to an autonomous or nonautonomous effect of the mutant gene on the serotonergic cells rather than to the lack of interaction with the LON. In order to begin addressing this question we examined the expression of the disco and of the glass gene relative to the expression of 5-HT in third instar larval brains.

The glass gene is expressed in only two cell clusters in the third instar larval brain in addition to the larval photoreceptor cells (Ellis et al., 1993). In contrast, the disco gene is widely expressed in the central nervous system. disco expression is particularly extensive in the developing adult optic lobes, in which it is displayed as a solid band of cells that wraps around the brain hemisphere at the level of the optic lobes (Lee et al., 1991).

Double labeling of wild-type third instar larval brain with anti-GLASS and anti-5-HT antibodies showed that the posterior glass-expressing cell cluster lies very close to the 5-HT-expressing LP1 cells (Fig. 4C, solid arrow and arrowheads). However, there was no overlap in the expression of glass and 5-HT in this or in the other more anterior glass-expressing cell cluster (Fig. 4B, solid arrow).

In order to assess disco gene expression, the disco enhancer-trap line C50 IS1, previously shown to reflect disco expression, was used. (Lee et al., 1991). Double labeling with anti-βgal (for disco expression) and anti-5-HT antibodies showed that the 5-HT-expressing LP1 cells lie very close to the band of disco-expressing cells (Fig. 4F, arrowheads and solid arrow). However, there was no significant overlap between disco and 5-HT expression in any part of the brain at that point in development. Given that there is widespread expression of disco, some of which is faint, it is possible that some of the 5-HT cells express disco at low levels (Fig. 4F, open arrows). Additionally, the disco gene expression undergoes complex temporal regulation: the pattern of expression found in the late third instar brain is more widespread than that found in the CNS during embryogenesis (Lee et al., 1991). The expression of the glass gene in the third instar, however, reflects the expression described in the late embryo (Ellis et al., 1993).

The absence of overlap between both glass and disco expression with the 5-HT expression suggests that the absence of the 5-HT arborization in the larval optic center of gl" and disco" mutants is not due to the lack of GLASS protein or abnormal function of the DISCO protein in the 5-HT neurons. However, it is still possible that the glass and disco genes may have a nonautonomous influence on 5-HT neurons which can lead to absence of the serotonergic innervation of the larval optic neuropil.

Absence of the 5-HT Arborization Is Strictly Correlated with Absence of Larval Optic Nerve Innervation in gl" Mosaic Larvae

The results described so far suggest that proper innervation of the larval optic neuropil by serotonergic fibers is dependent upon contact with the LON. However, given that both mutations used in this study are likely to affect the central brain (see pattern of expression above), one cannot exclude the possibility of direct or indirect influence of these mutations on the development of the 5-HT arborization.

In order to address this question we decided to concentrate on the mutation in the glass gene because of its restricted expression in the central brain (circa 20 cells per brain hemisphere) as opposed to the widespread expression of the disco gene throughout the central nervous system, making the latter more likely to display a
pleiotropic phenotype. To that end, we examined mosaic larvae lacking glass gene expression in one or both glass-expressing clusters in the central brain and/or in the larval visual system. These were concomitantly analyzed for the presence of 5-HT immunoreactivity in the larval optic neuropil.

Mosaic animals were generated by somatic excision of a P-element transposon carrying a wild-type copy of the glass gene in a gl^m72 mutant background (Kunes and Steller, 1991). Given that the glass mutation employed does not express any detectable protein (Moses et al., 1989), reduction or absence of glass expression in the central brain indicated excision of the transposable element carrying the wild-type glass gene function and thus a mutant patch. In the case of the larval photoreceptor cells, we used the presence of the photoreceptor-specific protein CHAOPTIN (Zippursky et al., 1984) to assess absence or reduction of larval optic nerve which in turn was indicative of mutant patches in the larval photoreceptor cells. Additionally, anti-5-HT antibody was used to determine the presence of 5-HT processes arborizing within the larval optic center.

Table 1 depicts the distribution of mosaic patches and the phenotype of the corresponding 5-HT projection innervating the larval optic center in a total of 771 brain hemispheres. In spite of the low frequency of mosaic patches it is clear that absence of glass expression in the central brain cells is not correlated with lack of serotoninergic innervation of the larval optic center. Complete absence of glass expression in either the anterior or posterior cluster in the presence of the larval optic nerve was not sufficient to affect the serotoninergic projection. Absence of the 5-HT arborization was only seen in cases where the larval optic nerve was also absent. An additional 316 brain hemispheres from putative mosaic larvae were double labeled with anti-CHAOPTIN and anti-5-HT antibodies (data not shown). Of these, only two did not show any recognizable 5-HT immunoreactivity within the larval optic center. Both brains lacked innervation by the LON. An example of a larval brain with a normal LON but completely lacking glass gene function in the central brain, as seen by the absence of GLASS immunoreactivity in both anterior and posterior clusters, is shown in Fig. 5E. In this brain the 5-HT projections could be easily seen displaying the expected morphology (empty arrow in Fig. 5C). These results demonstrate that the absence of 5-HT arborization in the larval optic neuropil in glass mutants is solely due to lack of innervation of the LON and not to lack of gene function in the central brain.

Larval Optic Nerve Interaction Is Required for the Development but Not for the Initial Outgrowth of the 5-HT Projection

The absence of the 5-HT arborization in gl^m72 mutants could be due to arrest in the initial outgrowth or to disruption in the subsequent development and/or maintenance of the 5-HT neuronal processes. Developmental analysis of gl^m72 allele (Figs. 6A and 6B, solid arrow) showed that the initial outgrowth of the 5-HT processes during first instar larval stage is similar to that of the wild-type larvae. However, during second instar larval stage, at the time in which the first contact with the LON is made, the 5-HT arborization of gl^m72 mutant larvae was significantly different from that of wild-type (Fig. 6F, solid arrow). These results suggest that the role of the LON is in the development of the 5-HT arborization and not in the initial outgrowth of the arborization forming processes, however, we cannot exclude a possible role for the LON in the maintenance of the 5-HT arborization later in larval development.

**DISCUSSION**

In this study, the role of the larval optic nerve on the development of the larval visual system neuropil was examined. We focused our investigation on the fate of newly described serotoninergic arborization that innervates the larval optic center whose cell bodies are located in the central brain. We examined the consequences of absence of contact with the LON in the development of this arborization and the viability of the corresponding cell bodies. To that end two mutations that disrupt connectivity in the larval visual system were used: glass^o9(gl^m98) and disconnected (disco). In gl^m98 mutant flies the larval photoreceptors do not differentiate appropriately, resulting in absence of the LON (Moses et al., 1989). In flies carrying mutations in the disco gene the LON fails to maintain stable connections with the optic lobe primordium (Steller et al., 1987). In larvae mutant for either one of these genes the arborization of 5-HT processes could not be seen. No apparent effect on the viability of the cell bodies could be detected. Developmental studies in gl^m98 mutant larvae showed that the arborization of the 5-HT processes in the larval optic center, but not the initial outgrowth of these processes, is disrupted. The direct influence of the LON on the 5-HT arborization was further confirmed by mosaic analysis of the gl^m72 mutation. Our results demonstrated that, although a glass-expressing cell cluster lies very close to the 5-HT arborization, lack of glass gene expression in these cells does not affect the development of the 5-HT arborization. Although the presence of the 5-HT arborization is dependent on the presence of the LON, the opposite is not true. In Ddc mutants lacking serotonin expression the LON morphology is indistinguishable from wild-type, suggesting that the LON does not require exposure to 5-HT for development or maintenance (data not shown). Our results do not exclude the possibility that continuous contact with the LON is required...
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for maintenance of the 5-HT arborization. In order to address this issue we would need to ablate the larval visual system in the late third instar larva rather than during embryonic development as is the case with the mutations employed in this study, and then follow the fate of the serotoninergic arborization. Thus we conclude that the serotoninergic innervation of the larval optic center is dependent upon afferent innervation from the larval photoreceptor cells for development and possibly maintenance.

Similar cellular interactions have been suggested for the development of the adult neuropil in Drosophila. Previous studies demonstrated that adult photoreceptor innervation is required for the proliferation of lamina precursor cells (Selleck and Steller, 1991). It has been suggested that at least two different signals operate during retinal axon-mediated development of the lamina ganglionaris, one that directs the lamina precursor cells to the proliferative phase and the other that triggers the differentiation program of the precursor cells (Meinertzhagen and Hanson, 1993). Anterograde retinal–lamina interactions have also been reported in the adult visual system of Drosophila. Photostimulation of the afferent retinal innervation causes terminal degeneration and synaptic disassembly but no transsynaptic degeneration of the deafferented targets (Brandstatter et al., 1991). Our observations in the larval visual system are consistent with those of Brandstatter and colleagues in the adult visual system in that no overt signs of degeneration were observed in the serotoninergic cell bodies in the absence of larval optic nerve innervation.

The observations discussed above suggest that factors secreted by or associated with the photoreceptor membrane are responsible for anterograde retina–lamina interactions during or after development is completed. It is possible that neurotransmitters present in the photoreceptor neurons play a role in some of these interactions. In Drosophila and other insects, histamine is the only neurotransmitter known so far to be expressed in the adult photoreceptor neurons and presumably in the larval photoreceptor neurons as well (reviewed by Nasset, 1991). It is unlikely that activity plays a role in the anterograde interaction reported here, given that maintenance of larvae in the dark throughout development did not alter the morphology of the 5-HT projection in the larval optic neuropil (data not shown).

The adult visual system of Drosophila is invaded by extensive serotoninergic processes. In the Fleshy, absence of synaptic connection between the photoreceptor axons and the serotoninergic processes in the lamina suggested a paracrine action of serotonin in this system (Nasset et al., 1983). Whether the serotoninergic processes in the larval optic center make synaptic contact with the larval optic nerve terminal is yet to be determined.

Developmental analysis of the serotonin processes showed that the contact of the LON by the 5-HT arborization takes place during early third instar larval stage. During early pupal stages, concomitant with the degeneration of the LON, the 5-HT processes retract from the larval optic center and are not seen again projecting toward the developing adult optic ganglia (data not shown). These results suggest that this 5-HT projection has a primary role in the function of the larval visual system during the third instar stage.

Drosophila larvae display marked aversion to light during the first, second, and first half of the third larval instar (Sawin et al., 1994; Sawin-McCormack et al., 1995). At the onset of the wandering phase larval photobehavior becomes progressively less photonegative, achieving photo neutrality just before pupariation (Sawin-McCormack et al., 1995). In several invertebrates serotonin has been proposed to have a role in modulating the photosensitivity of the eye (Barlow et al., 1977; Eskin and Maresh, 1982; Arechiga et al., 1990). Thus, it is possible that the contact of the LON by this serotoninergic process mediates the transition in photobehavior observed during the mid third instar larval stage.

Identification of additional mutants displaying a normal larval visual system morphology but with defective larval photobehavior will aid in the investigation of the significance of the serotonin arborization in the developing optic lobe area of the larval brain. The presence of serotonin processes in intimate association with the LON and the simplicity of the larval visual system in Drosophila makes it an excellent model to study the role of serotonin in visual system function and development.

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Chapter 3

Identification of DISCO-interacting proteins.
Introduction

Disconnected (disco) gene is required for proper neuronal connections in both larval and adult visual system of Drosophila (Steller et al., 1987; Campos et al., 1995). The disco gene codes for a protein with two C2H2 type zinc-finger domains, a feature shared by many transcription factors (Pieler and Belfroid, 1994). A human gene called basonuclein, contains zinc finger motifs similar to the motifs present in DISCO (Tseng and Green, 1992). The striking conservation of the zinc finger sequences in evolution suggests an important role for this motif in the function of the DISCO protein. A portion of the DISCO protein including the zinc finger motifs has been shown in vitro to confer sequence specific DNA binding activity (Kevin Lee, Ph. D. thesis, 1993).

Expression of the disco gene is seen in several cell types including neuronal, nonneuronal and glial cells, throughout the life cycle of Drosophila (Lee et al., 1991). Of particular importance to the visual system development is its expression in glial cells that are intimately associated with the larval optic nerve (LON) and three so called “optic lobe pioneer cells” (OLPs), also contacted by the LON. The OLP cells are neuronal cells in the optic lobe primordium that are thought to guide the LON in the brain (Campos et al., 1995). It has been suggested that the disco gene activity may be required in the optic nerve glia and the OLPs for the development of proper connections between the LON and its target cells in the brain (Campos et al., 1995).
Among other cell types, *disco* expression is seen in visceral mesoderm, gnathal segments, thoracic segment, leg imaginal discs and cardioblasts. *disco* expression is differentially regulated in different tissues. For example, *disco* expression in the primordia of leg imaginal disc is dependent on the homeodomain-containing protein DISTALLELSS (Cohen *et al.*, 1991), whereas, *disco* expression in the cardioblasts is dependent on TINMAN (Bodmer, 1993), another homeodomain protein.

In two apparent null mutant alleles of *disco*, carrying changes in the highly conserved Cysteins of the zinc finger motifs, expression of a non-functional DISCO protein is seen in all embryonic tissues that normally express *disco* except the cells in the optic lobe region (Lee *et al.*, 1991). The absence of *disco* expression in the optic lobe tissue of these mutants has been shown to be due to a tissue specific autoregulatory activity of *disco* which is localized only to the cells of the optic lobe primordium (Kevin Lee, Ph. D. thesis, 1993). The tissue specific autoregulatory function of *disco* suggests that interaction of *disco* with other tissue specific factors restrict *disco* autoregulatory activity to the optic lobe primordium.

There are a variety of different approaches one might use to identify factors that interact in a tissue specific manner with DISCO. These include, traditional approaches, such as genetic screens or immunoprecipitation assay, as well *in-vitro* approaches using synthetic machinery such as that used in “interaction trap” assays. In our study we chose to use the latter approach (Gyuris *et al.*, in 1993) to identify DISCO interacting proteins.
The yeast interaction trap.

The yeast interaction trap used in this study is an implementation of the yeast two-hybrid system developed by Gyuris et al. in 1993. The method uses the transcription of yeast reporter genes as a synthetic phenotype to detect protein-protein interactions (reviewed by Finlay and Brent, 1995). It provides a convenient way of cloning cDNAs for interacting protein.

Most eukaryotic transcription factors contain two distinct domains; a DNA binding domain and an activation domain required for transcription activation. These domains can be exchanged between different transcription factors while retaining their function. For example, the DNA binding domain of Gal4 by itself is not sufficient to activate transcription of reporters with upstream Gal4 binding sites. However, when it is fused to activation domains of other proteins, the fusion protein is able to activate transcription of the same reporter gene. Another important feature is that the DNA binding domain and the activation domain need not be covalently attached to each other for activation to occur (Ma and Ptashne, 1988). If two proteins, one carrying the DNA binding domain and the other carrying the activation domain, interact with each other activation of reporter gene transcription can occur. This is the basis for the yeast interaction trap.

There are three basic components in any two-hybrid system: (i) a yeast vector for expression of a protein of interest fused to a DNA binding domain, (ii) a yeast vector for expression of an unknown cDNA (from a library) encoded protein fused to a transcription activation domain, and (iii) a yeast reporter gene that contains binding sites for the DNA binding domain. All systems utilize the DNA binding domain from either the yeast transcription factor Gal4 or from the
bacterial transcription factor LexA. The yeast interaction trap developed by Gyuris et al. (1993) uses the LexA system. Here, a LexA-fusion protein, containing the protein of interest fused to the LexA DNA binding domain, is referred to as a 'bait'. The activation domain B42 used in this system is derived from *E. coli* and is fused to the library proteins expressed from cDNA libraries made in the yeast interaction trap expression vector pJG4-5. The expression of the activation domain tagged library proteins is under the control of the galactose inducible promoter from the yeast Gal1 gene. The library vector not only provides the activation domain, but also a nuclear localization signal and an epitope tag (for convenient detection of the fusion protein) fused at the N-terminus of the expressed library proteins. Two reporter gene constructs are used in this system; a yeast *leucine2* derivative that has its normal upstream sequences replaced with *lexA* operator and a *lacZ* reporter construct with upstream LexA binding sites. Transcription of the *lexA*-operator-*leu2* gene is detected by the ability of a *leu*<sup>+</sup> yeast strain to grow in the absence of Leucine. While, transcription of the *lacZ* reporter is detected by the ability of the strain to form blue colonies on X-gal plates.

The yeast interaction trap is schematically represented in Figure 1. The bait proteins used in our study are different LexA-DISCO fusions carrying either N-terminal or C-terminal DISCO fragments. The bait protein is constitutively expressed. It binds to the *lexA* operator located upstream of the *lacZ* gene (or *leu2* gene) in the reporter construct. The activation tagged library protein is expressed conditionally from the Gal1 promoter. In glucose medium the Gal1 promoter is repressed. Only when the yeast is growing on galactose medium, the library protein expression is induced. Those library proteins that interact
with the bait activate transcription of the *leu2* and *lacZ* reporter genes. As a result, cells containing activation-tagged library proteins that interact with the bait form colonies on galactose medium lacking Leucine and form blue colonies on galactose X-gal plates.
Figure 1

The yeast interaction trap scheme:

The yeast interaction trap is schematically represented in this Figure. The bait (LexA-DISCO fusion protein) binds to the LexA operator (LexA OP) located upstream of the reporter genes (lacZ or leu2). The binding of the bait does not itself activate transcription of the lacZ reporter gene. Only when an activation domain-tagged library protein (X) interacts with DISCO, the activation domain fused to the library protein activates the transcription of the reporter gene.
AD = Activation Domain
X = Library protein that interacts with DISCO
Results:

I. The yeast interaction trap screening.

A. Testing the DISCO baits: Repression assay.

The size of an exogenous protein that can be efficiently expressed in yeast is limited. Since the expression of bait with the full-length DISCO protein (92 kDa) was poor in yeast, we used smaller disco fragments to construct baits for the "interactor hunt".

The middle portion of DISCO contains a 40 amino acids glutamate-rich acidic region (Heilig et al., 1991). Glutamate- and aspartate-rich acidic regions are known to act as activation domains in several transcription factors (e.g., Ma and Ptashne, 1987; Triezenberg et al., 1988). Therefore, presence of both a DNA binding domain and an activation domain in the bait may cause activation of the reporter gene in the absence of any interaction with activation tagged library proteins. Indeed a recent study in our lab has demonstrated that binding of the LexA promoter by the glutamate-rich region of DISCO as a LexA fusion protein was sufficient to cause activation of the lacZ reporter gene (Figure 3). Therefore, the glutamate-rich region was excluded from our study and only the N-terminal and C-terminal portions of DISCO were used as baits for the interaction trap assay (Figure 2).

The C-terminal DISCO bait used in our study contained a 153 amino acids DISCO fragment. The N-terminal DISCO fragment used for the interactor hunt was 186 amino acids long. To our surprise, this DISCO bait caused activation of
reporter gene transcription in the absence of a activation-tagged library protein (Figure 3).

In order to be able to use the N-terminal portion of the disco gene in the interaction trap screen, a DISCO bait was constructed where the cysteine (Cys) residue at position 127 was replaced with a serine (Ser) residue. The presence of the Cys residue at position 127 of the second zinc-finger domain of DISCO is crucial, since, replacement of this residue with Ser in the disco1 mutant leads to a complete loss of disco function (Heilig et al., 1991). The Cys127 to Ser substitution also abolished the transcription activation property of the N-terminal DISCO bait, which gave us the opportunity to use the mutated DISCO bait for our yeast interaction trap screening. The mutated DISCO bait was generated simply by PCR amplification of the N-terminal disco fragment from the chromosomal DNA of disco1 mutant that carries the above mutation. The construct was subsequently sequenced to confirm the presence of the desired mutation.

In order to activate the transcription of the reporter genes, baits need to be localized in the nucleus (Finley and Brent, 1995). To examine nuclear localization and the ability of these baits to bind LexA operators, transcription repression of a lacZ reporter gene was monitored. Expression of the lacZ gene from the reporter gene construct pJK101 (Brent and Ptashne, 1984) is driven by yeast Gal1 promoter. The construct pJK101 also contains one LexA operator located between the Gal1 promoter and a TATA box. lacZ expression is thus galactose inducible. For reasons not completely understood, binding of a bait to the LexA operator causes 2-20 fold repression of the lacZ gene expression in galactose medium. This property of the reporter construct is used in the
repression assay to confirm nuclear localization and LexA operator binding ability of an experimental bait (Brent and Ptashne, 1984).

The repression assay was done on X-gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside)-galactose plates and the extent of blue coloration produced by the yeast colonies with or without the bait constructs were compared. Yeast colonies carrying the DISCO bait constructs showed distinct reduction in the expression level of the lacZ reporter gene compared to that produced by the colonies lacking the DISCO baits. The results of the repression assay show that both N- and C-terminal DISCO baits are able to localize to the nucleus and are able to bind LexA operator (data not shown).

B. Interactor hunt using the C-terminal DISCO bait:

A Drosophila melanogaster 0-12 hr. embryonic cDNA library made in the yeast interaction trap vector pJG4-5 was kindly provided by Dr. Brent (Harvard Medical school). This library was amplified and screened for all interactor hunts. In two trials, 1X10⁶ and 0.5X10⁶ colonies were screened with the C-terminal DISCO bait. No cells survived on galactose plates lacking Leucine. Thus, no interactor was isolated using the C-terminal DISCO bait.

C. Interactor hunt using the N-terminal DISCO bait:

About 2.5X10⁶ colonies from the 0-12 hr. Drosophila embryonic library were screened using the N-terminal DISCO bait carrying a Cys127-Ser127 mutation. 150 colonies survived on galactose plates lacking Leucine. To check the expression of the lacZ reporter gene, these 150 colonies were plated on galactose medium containing X-gal. 20 colonies appeared blue on the X-gal
plate. The other 130 colonies therefore may represent nonspecific interactions. Since, the library vector expression is induced by galactose and repressed by glucose, the above 20 colonies were replica plated on glucose medium containing X-gal. Out of 20 colonies 18 showed repression of lacZ expression on glucose X-gal medium, indicating that expression of the activation tagged library protein was necessary to activate transcription of the lacZ reporter gene in those colonies. Two colonies that failed to show repression of lacZ expression in glucose plate again represent non-specific interaction.

D. Classification and isolation of the cDNAs encoding the interacting proteins:

Since the Drosophila genome consists of approximately 12,000 genes (Miklos and Rubin, 1996), screening of 2.5X10^6 cDNAs should lead to the isolation of several copies of a single interacting cDNA. The number of copies of a single cDNA present in 2.5 million clones depends on the abundance of the corresponding mRNA in the embryo. In order to classify the isolated positive yeast clones, the library cDNAs were PCR amplified from yeast plasmids isolated from the positive clones. The amplified products were digested with HaeIII and AluI enzymes and the restriction patterns were compared. From the restriction pattern, the 18 cDNAs were classified into three groups: 16 out of 18 showed a similar restriction pattern with both enzymes (group 1) while the other two showed distinct restriction patterns (group 2 and 3).

The plasmids isolated from positive yeast colonies represented a mixture of three different plasmid constructs; the library construct, the bait construct and the reporter gene construct. The selectable marker on the library plasmid was tryptophan. In order to isolate the library plasmid containing the interacting
cDNA, the plasmid mixture isolated from a positive yeast clone was transformed into a trp⁻ E. coli strain MC1066. The colonies that survived on M9 minimal medium lacking Tryptophan represented colonies carrying the library plasmid. Yeast library plasmids isolated from 4 out of the 16 interacting colonies from group 1 and the two single interacting colonies from group 2 and 3 were transformed into the trp⁻ E. coli strain. The library plasmids from the positive trp⁺ bacterial colonies were isolated and transformed back in yeast together with the N-terminal DISCO bait to confirm the interactions detected previously (Figure 4).
Figure 2

Diagrammatic representation of baits used in the interaction trap assay:
The baits used in the interactor hunt are schematically shown in this
Figure. The entire DISCO protein and the fragments used in the baits are
represented with horizontal bars. A 186 amino acids N-terminal DISCO
fragment (amino acids 1-186) carrying a Cys127 to serine127 substitution,
and a 153 amino acid C-terminal DISCO fragment (amino acids 415-568)
were used for the interaction trap screening. The red box shows the
location (amino acids 89-149) of the two repeats of the zinc finger motif
present in DISCO. The yellow box shows the 40 amino acids glutamate
rich region (from residue 311 to 351) located in the middle portion of the
DISCO protein.
Figure 3

Transcription activation ability of the N-terminal and glutamate-rich regions of DISCO:

This Figure shows the ability of the zinc finger domain and the glutamate rich region of DISCO to activate lacZ reporter gene transcription from the reporter construct used for interactor hunt. Panel A shows a schematic representation of the DISCO protein. The zinc-finger domains and the glutamate-rich region are indicated with different shades of blue.

Panel B shows the effects of different DISCO baits on reporter gene expression. The ability of the DISCO baits with the N-terminal (From amino acids 1-186, including the zinc-finger domain) and Glu-rich regions of the protein (From amino acids 186-414) to activate the lacZ reporter gene expression is revealed from the blue colour of the colonies containing these baits. Substitution of cysteine residue at position 127 by a serine residue in the mutated DISCO bait (indicated with an asterix) abolishes the transcription activation ability of the N-terminal bait as revealed from the white colour of the colony. The C-terminal bait is also unable to activate the reporter gene expression.
Figure 4

Results of the interaction trap screening:
Screening of a Drosophila cDNA library constructed in a yeast expression vector with the N-terminal DISCO bait led to the identification of three interacting proteins. This Figure shows three positive yeast clones (blue colonies = lacZ reporter gene expression). All of the three interacting proteins; a putative ATPase subunit of the Drosophila 26S proteasome complex, the DIP1 protein and the DIP2 protein activate lacZ gene expression in galactose X-gal plate (blue colonies) in the presence of mutated N-terminal DISCO bait. The lacZ gene expression is activated only when both the DISCO protein and one of the interacting proteins are present simultaneously. Presence of either of these proteins alone is not sufficient to activate the transcription of the reporter gene. LexA alone without the DISCO fragment is also unable to produce blue colonies in the presence of any of the interacting proteins. The LexA-DISCO fusion protein in the presence of the activation domain alone without an interacting protein, also does not activate transcription of the lacZ reporter.
LEC A-DISCO (N-Term.)

LEC A-DISCO (C-Term.)

LEC A alone

Proteasome+AD  DIP2+AD  DIP1+AD  AD alone

AD=activation domain
E. Sequence analysis of the cDNAs encoding interacting proteins:

The cDNAs corresponding to 4 interacting clones from group 1 and single interacting clone from both group 2 and group 3 were sequenced. Sequence analysis confirmed that all members of group 1 represent the same cDNA. We denoted the interacting protein encoded by the group 1 cDNA as "DISCO interacting protein 1" (DIP1).

The partial dip1 cDNA isolated from the interacting clone contained a 1.1 kb fragment corresponding to the 3' end of dip1 mRNA. Sequence analysis using 'BLAST' tool (Basic Local Alignment Search Tool; Altschul et al., 1990) revealed that the putative DIP1 protein contains a region that shows sequence similarity to dsRNA binding domains (dsRBDs) (Figure 5). A consensus for dsRBDs has been determined through mutational analysis, in-vitro binding experiments and comparison of dsRBD sequences from several dsRNA-binding proteins (St. Johnston et al., 1992). The putative dsRBD sequence of DIP1 matches well with the C-terminal short domain consensus but shows poor similarity with the N-terminal long domain consensus (Figure 13). The putative dsRBD of DIP1 shows extensive similarity to the dsRBDs of mammalian glutamate receptor editases (Melcher et al., 1996; Lai et al., 1997). The similarity extends beyond the conserved residues of the dsRBD consensus (Figure 14). One striking feature of the dip1 cDNA is the presence of 4 repeats of a 124 bp sequence at the 3'end of the cDNA (Figures 5, 12A.). Nucleotide sequences of the two 124 bp middle repeats are exactly equivalent, while, the first repeat contains two extra base pairs and the last repeat (120 bp) is interrupted by an EcoRI site (Figure 12B).

The group 2 cDNA is also a newly identified cDNA. We will refer to the interacting protein encoded by this cDNA as "DISCO interacting protein 2"
(DIP2). The cDNA clone contains a 1.2 kb 3' end fragment of dip2 cDNA (Figure 6). Sequence analysis through 'BLAST' search (Altschul et al., 1990) using the NCBI BLAST search server showed a high degree of similarity with a human cDNA isolated in the Human Genome project (Accession # D80006). The C-terminal 100 amino acids segment and N-terminal 155 amino acids segment of the predicted partial Drosophila DIP2 protein share 82% and 66% identities respectively with that of the human homolog. Both the predicted human protein and DIP2 share a high degree of similarity with a protein sequence predicted from a Caenorhabditis elegans genomic DNA sequence (Accession # 2088703, Wilson et al., 1994). The degree of similarity shared between these proteins (Figure 26) suggests that these proteins are human and C. elegans homologs of Drosophila DIP2.

Sequence analysis showed that the group 3 cDNA codes for an ATPase, the fourth subunit of the Drosophila 26S proteasome complex. This cDNA was originally cloned based on its similarity to the ATPase component of the mouse 26S proteasome (Hoyle and Fisher, 1996).
Figure 5

Partial dip1 cDNA sequence.

This Figure shows the sequence of the 1.1 kb partial dip1 cDNA isolated from the yeast interaction trap screening. The predicted partial protein sequence is also shown. The cDNA lacks the 5' end of the corresponding mRNA including a portion of the coding region. The putative dsRNA-binding domain in the partial protein sequence is highlighted. The first 3 nucleotides of all four 124 bp repeats in the 3' UTR are shown in boldface. The stop codon (denoted with an asterix) lies at the beginning of the first repeat. The EcoRI site (GAATTC) that interrupts the fourth repeat is also shown in boldface and is underlined. The EcoRI site is followed by a 48 bp sequence and a polyA tail.
Partial dip1 cDNA sequence and the predicted protein sequence:

GCCTGCGAGAAGGCTTTGGCGCGATTTTTATATTGCAAAAAATGACCCCCAAGCCGGCCCGT 60
ACEKAWRDFIIIAKMTPKPPR
ATTACCAGGTGGAGATGGGCTCAGAGCCTATGATCAACAGGAGATGAGGCCTAGCA 120
IHQVEMGSEPMIDINEDEADA
CCGGATGATCATCTGCCATGTTAATCTGGCTGCTTGGGACATCTCAAAGCTTGTCGCG 180
FDDLPMLNLASFAIYKLFA
GAGTTGGAACGGGAAGGCTATGTCGTGCAGCAACCTTCGCGCAATGCTGCAGCA 240
EWEREYVPEMHPSANAQQ
CAGGCAGGGAGGATGCCGAACCTCAAGTTCCCCCCCGGCGAAAAGGACCCAGCG 300
QAGGDAGTPVPVPVPKPPK
CCAGTGCGACACGAAGCTACCCCTCGTCGGGTGAGACAGAGAAGCCGACCGG 360
PVRTELPWSGWE-developer-PATILC
ATTATGCAGTCCGGAGACTCAACTACGTGGACTACGGGTATCTGGCGACAAAGCCAACCG 420
IMRPGLNYVDYGSSGDKTN
ATGGCAGTATCTTCAGGACACCATGGGACACCTACGCCACCCGAGATCAAAAG 480
MQHLMIMVNDQEFHANGRSK
AAAATCGGGCCTGCAGCTGCTGAAAGTGTCACCTCTCTGTKGCGACACACTTC 540
KIARRPVKVCNSLFGTNF
ACCTACAGCACACACTTAAGACACACACACTCCAACCGAGACTCCAAGGACGACA 600
TYSDDT*
ACACACATAACGGGACACTCTTTACGGGACAGGAGACATTTGGGACCTGCAGTCAT 660
TGCGCAACTTTAAGGGACACACTCTGAAAGCACCACACCTCCCAACCCGAGACTCCAAGGCGC 720
ACGACAGCAGACACCTATGCTCTCTTAACGGGACAGCAAATCGAGAGATTGGCAGCCCGATG 780
CAATGGGCGACACTTTAAGGGACACACTCTGAAAGCACCACACCTCCCAACCCGAGACTCCAAGGCGC 840
GCCACAGCAGACACACCTATGCTCTCTTAACGGGACAGCAACTAGGGAGATTGGCAGCCCGATG 900
ACGTACATGCGGCAACCTTTAAGGGACACACTCTGAAAGCACCACACCTCCCAACCCGAGACTCCAAGGCGC 960
AACTGCCGAGCAGACACACACTCTATGCTCTCTTAACGGGACAGCAACTAGGGAGATTGGCAGCCCGATG 1020
CCTGACATTGCCGCAACCTTTAAGGGACACACTCTTCTGCTCAAAAGTGAGAGGACATATTATA 1080
AACAAAGATTCTCCATAAAGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG
Figure 6

Partial *dip2* cDNA sequence.

This Figure shows the sequence of the 1.2 kb partial *dip2* cDNA isolated in yeast interaction trap screening. The partial amino acid sequence (337 residues) of the predicted protein is also shown. It contains no known functional domain. The stop codon on the cDNA is indicated with an asterix. The cDNA fragment belongs to the 3’ end of the corresponding mRNA. The 53 amino acids sequence that is unique to the *Drosophila* DIP2 homolog is highlighted.
Partial dip2 cDNA and the predicted amino acids sequence:

AGAGTGAACCTGACCCGACGATCCTGACAGCTATTTTTCAAGCCCTTTGCTGATATGACGC

RVQLTQQFCFKLFQALGLNTR

TGCGTCTGCAACTTCTTGTGATGTGCTGTAAATCCTGCGGATATGGCTGTTTGCT

CVSTSFGRVPNAPICVQGAS

TCTGCAGAGAGTCTAGTATATGAAGATATAGTGGAGACTTGAAATATGGCTGTT

SAESAOQPVDVMRARLRRNNRVA

CTGTTAGCGAGGCGGCAAAATTGGTGTTGTAATGGATACGATTAACCTTTACCAA

LVERAPNSLCVIESGKLPLP

GGCCGAAAGTTGAAATTTCCCGAAAATCTAAGGCGCCACGCTGGCGACGTGGAT

GVKVIANPETKGHCGDSHL

GGAGAAACTTGGTTGTCAGCTCCTCAACACGACATGTGTACTTTAAACCTTTTGAAC

EIAWVQPHNAHGYFTIYGD

AAATCGACTACAAATGTACACTTCAACGCACAAATTGGTAACCTGGGGCCACCTCGAGACTA

ETDYNDHFNAKLVGTATSEL

TATGCACGCACCTGGTTATTTAGATTCTTTCGCGCACCAGAATGCCTGCAATACGGCAT

480

YARTGYLEFLLRRTESQNAS

CTGGTTGAGACGAGCACCACCAAAATGGTGACGTGCAGATGTGATACGAGATTTTGAAT

540

LLDETTTPSVASRDSTDTELSN

TCGATAAGCTACTATGAAATTTCATTTCCATGT GCCGAAATTCCGACGAT

600

SISQLQLNFSNVSLLGGNSEH

AGGCTGGTAGGCGCGCGACATACGCTATAGCAAGAATACAGCGCGACGTGTTATGTA

660

SLVGGNANDQELHDAYV

GTCGGAGCTGTGGATAAGCTCTTTTAGCGACATGACTACCCCAAATTTGTATATC

720

VGDEMISLRRMGNYHPIDI

GAAATTCGGTAAATGTGGTGTCACAATAATTTTGGTACCTGCCGTGGTTTTCACCTGAGCT

780

ENSVMRCVKKAIAECACAVFTWT

AACTATTAGCTGATTTCTGATGAGCCTGGCAGGATACGAGCGCTTTTGATTTGTT

840

NLIVVVELDGNESEALDLV

CCCTGGTCACAAAAACAGTATAGCGATACGCTATAGCCTGCCTGGCTGTTAGTGTC

900

PLVNTNVTLEHDQQLIVGVVVV

GTGATCCAGGCTGTGGTGGCTATTAAATGTGCACCCGCTAATACGATCATGATACAC

960

VDPGVVVIPINSRGEEKQRMLR

GATGTTTTCTGCACGCAAAATTGGATCCAATATACGATGATCATATACATGTAATACAC

1020

DGFLADQLPDNIPYVAYN

AAATTACATATTACATTTCATTTTAGATTAGCAGCTACAAAAAGATGGTTAACATCTTTAACA

1080

ATAATTACAAACACGAGAAATCAGCAGAAGATGGCTATTTTGTGTAACCGCA

1140

GGGCTAATTACCATATTTTTTAGATTAGCGAACAGATGGTTAACATCTTTAACAAAATG

1200

AAATGACTATTATCTTTATATAGCATCACCATAT

1230
II. Verification of the interactions between DISCO and its interacting proteins DIP1 and DIP2 seen in the interaction trap assay.

A. In-Vitro Binding Assay.

The specificity of interaction between DISCO and the proteins isolated from the interaction trap screening were examined *in-vitro*. For this study, the individual proteins were prepared as follows; *disco* was expressed as a GST-fusion protein in *E. coli*, the *dip1* and *dip2* cDNAs were transcribed and translated *in-vitro*.

The 186 amino acids N-terminal DISCO fragment used in the yeast interaction trap was also used for the *in-vitro* binding assay. Since we were unable to use the wild-type DISCO protein for the yeast interactor hunt, to rule out the possibility of a non-specific interaction between the mutated DISCO protein and the library proteins, we used both wild-type and mutated N-terminal DISCO fragments for the *in-vitro* analysis.

For *in vitro* expression of the *dip1* and *dip2*, the corresponding cDNAs were cloned into the *in vitro* expression vector pSPUTK (MBI Fermentas). SP6 RNA polymerase was used to drive transcription of these cDNAs. The translation was done in rabbit reticulocyte lysate (MBI Fermentas). [S$^{35}$]Methionine was used to radiolabel the translated protein.

Purified GST-DISCO fusion proteins (bound to glutathione beads) were allowed to bind to the *in-vitro* expressed DIP1 and DIP2 proteins. Bound proteins were eluted from the glutathione-agarose beads and analyzed on denaturing SDS-polyacrylamide gel. Only if *in-vitro* translated DIP1 and DIP2 are able to bind DISCO will these proteins be present in the eluant, and the radiolabeled
proteins can be detected by autoradiography. Results of this experiment are shown in Figure 7. Both wild-type and mutated DISCO fragments were able to bind the in-vitro expressed DIP1 (lanes 6 and 7) and DIP2 (lanes 2 and 3) proteins to the same extent. GST alone, without the N-terminal DISCO fragment was unable to bind either one of these proteins (lanes 8 and 4). Therefore, the in-vitro binding assay showed a direct physical association between DISCO and DIP1/DIP2.

B. The mRNA expression pattern of disco overlaps with that of dip1 and dip2:

The expression pattern of the genes encoding the interacting proteins in Drosophila embryo should overlap with that of disco at least partially if these are true in-vivo interactors of DISCO. To examine this we determined the mRNA expression pattern of dip1, and dip2 during various stages of embryogenesis. For in-situ expression studies, a digoxygenin labeled RNA probe was synthesized from a dip2 cDNA and non-repeated portion of the dip1 cDNA.

The dip1 mRNA shows a ubiquitous expression pattern at all stages of embryogenesis. However, the expression is not homogeneous, rather, more concentrated in the developing CNS (Figure 8). Expression pattern of the dip1 mRNA overlaps with that of disco in the developing optic lobe. The cells of the optic lobe primordium invaginate from the posterior portion of the procephalic lobe (embryonic head) during stages 12. This tissue remains attached to the larval brain until stages 15-16, when it is completely incorporated into the brain hemispheres (Green et al., 1993). Figure 10 shows stage 13 embryos to compare overall expression patterns of dip1 and disco. The dip1 expression also overlaps with disco in the visceral mesoderm and the epidermis of gnathal and thoracic
segments. Visceral mesoderm staining becomes evident during stages 12 and 13, the last stages of germ band retraction. Following these stages, the cells of the visceral mesoderm elongate and contact the midgut primordia and eventually form a sheet around the gut (Campos-Ortega and Hartenstein, 1985).

The *dip2* mRNA shows extensive expression in the CNS throughout embryogenesis (Figure 9). Expression is seen in the optic lobe (OL) primordium starting from stage 12. Expression in the OL is seen at later stages as well. Staining in this tissue is shown in a stage 15 embryo, at which stage the OL has already been incorporated in the larval brain. A lower level of expression of the *dip2* mRNA may also be present at least in a portion of the peripheral nervous system. However, due to the lower resolution of the *in-situ* procedure, we were unable to unambiguously detect the peripheral nervous system expression pattern of the *dip2* mRNA. In addition, expression of the *dip2* mRNA is also seen in the visceral mesoderm during stages 12 and 13 (expression in a stage 12 embryo is shown in Figures 9 and 10). The expression pattern of the *dip2* mRNA partially overlaps with that of *disco* in the optic lobe primordium, and the visceral mesoderm (Figure 10).
Figure 7

Results of the *in vitro* binding assay:

Panel A depicts the results of the *in-vitro* binding assay. *In-vitro* translated partial DIP2 (40 kDa) and DIP1 (29 kDa) proteins are shown in lane 1 and 5 respectively. Both the wild-type and the mutated DISCO proteins are able to bind the *in-vitro* translated DIP2 (lanes 2 and 3) and the DIP1 (lanes 6 and 7) proteins. GST alone is not able to bind either protein (shown in the control lanes 4 and 8).

Panel B shows the expression of the GST-DISCO fusion proteins (48 kDa) expressed and purified from *E. coli* BL21 strain. Lane 1 shows expression of the GST protein alone. Lane 2 and 3 show expression of GST-fusion proteins (48 kDa) containing the wild-type N-terminal DISCO fragment and a mutated N-terminal DISCO fragment respectively. The mutated DISCO fragment contains a Cys127 to Ser127 substitution.
Figure 8

mRNA expression pattern of the dip1 gene:
The results of the in situ mRNA expression study are shown in this Figure. Expression in three different stages of embryogenesis (stages 9, 12 and 14) are shown. Only lateral views are shown. Anterior is to the left and dorsal side is up. Embryo staging was done according to Campos-Ortega and Hartenstein, 1985. A ubiquitous expression pattern of dip1 mRNA is seen at all stages of embryogenesis. Expression of the dip1 mRNA in the optic lobe primordium (arrow) is shown in the stage 12 embryo, the last stage of germ band retraction. Because of the superficial view of the embryo, expression in the invaginated optic lobe primordium can not be seen clearly. At this stage expression of the dip1 mRNA is evident in the visceral mesoderm (VM) and in a variety of other tissues, including the epidermis of gnathal (GN) and thoracic segments (T). At all stages, the expression appears to be more concentrated in the developing CNS than other parts of the embryo. In the stage 14 embryo, the CNS is indicated with an arrow with white arrowhead.
mRNA expression pattern of the *dip2* gene:

This Figure shows the *dip2* mRNA expression pattern during different stages (stage 11, 12 and 15) of embryogenesis. Expression in the CNS (arrows with white arrowheads) is seen at all stages of development. At around stage 12 expression in the visceral mesoderm (VM) is clearly visible. The location of the optic lobe primordium is indicated with an arrow. Because of the superficial view of the embryo, the OL can not be clearly seen.
Figure 10

Expression patterns of *dip1* and *dip2* overlap with that of *disco*.

In this Figure, expression patterns of the *dip1* and *dip2* mRNAs are compared with that of the *disco* at embryonic stage 12. *disco* expression is seen in the optic lobe primordium (arrow), in the gnathal (GN) segments, in the ectoderm of thoracic segments (T) and in the visceral mesoderm (VM). The expression pattern of *dip1* and *dip2* mRNAs overlap with that of *disco* in the optic lobe primordium (arrow) and the visceral mesoderm (VM). *dip1* and *disco* expression patterns also overlap in the ectoderm of gnathal and thoracic segments.
III. Characterization of the *dip1* cDNA:

A. Cloning of the full-length *dip1* cDNA:

A partial 1.1 kb 3' end fragment of the *dip1* cDNA was isolated in the interaction trap screen. In order to isolate the full-length *dip1* cDNA, another 12-24 hours *Drosophila* embryonic cDNA library made in λgt11 phage vector was amplified and screened. The library (obtained from Dr. Roger Jacobs, McMaster University) was made by cloning cDNAs synthesized from embryonically expressed *Drosophila* mRNAs into the EcoRI site of the phage-vector.

About 2 X 10^6 plaques from the library were screened with a probe made from the *dip1* cDNA fragment isolated from the yeast interaction trap library. Three positive plaques containing *dip1* cDNA fragments were isolated. The sizes of the cDNA inserts were examined by digesting the phage DNA with EcoRI. Two of the positive plaques contained only a small fragment, about 1 kb in length. The third plaque contained two EcoRI fragments of approximately 2.5 kb and 0.5 kb length. Both fragments were subcloned into the EcoRI site of the SK^- pPBluescript vector and sequenced. Since, we used only a 1.1 kb 3' end *dip1* cDNA fragment as probe, the two smaller cDNAs (∼ 1kb) are presumably part of the larger cDNA isolated in this screen. However, we can not exclude the possibility that the smaller cDNAs may also represent the differentially spliced *dip1* transcript.

B. Sequence analysis of the *dip1* cDNA:

Sequence analysis of the 2.5 kb and 0.5 kb fragments of the *dip1* cDNA showed that the 0.5 kb fragment corresponds to the 3' end and the 2.5 kb
fragment corresponds to the 5' end of the cDNA. The 2.5 kb fragment contained the full coding region of the *dip1* cDNA (Figure 11).

The number of 124 bp repeats present in the two cDNAs isolated from the yeast interaction trap library and the λgt11 library differ from each other. The newly isolated cDNA contains estimated 8-10 repeats as opposed to 4 repeats present in the previously isolated cDNA (Figure 12A.). Following the repeated region, an EcoRI site and a 48 bp sequence are common in both cDNAs. The 48 bp sequence is followed by a poly A tail in the *dip1* cDNA isolated from the yeast vector. On the other hand, the cDNA isolated from the λgt11 vector contained an extra 348 bp sequence between the 48 bp sequence and the polyA tail. This difference in sequence might indicate that these cDNAs represent two alternatively spliced forms of the *dip1* mRNA. The differences between the two cDNAs are diagrammatically represented in Figure 12A. The sequence of each individual repeat is shown in Figure 12B.

Considering the presence of 8 repeats, the cDNA isolated from the λgt11 vector is about 2.6 Kb. The exact number of repeats present, and as a result the exact size of the cDNA, was difficult to determine due to difficulty in generating primers with unique sequences. There is a putative start codon 263 bp downstream from the 5' end of the cDNA fragment and a 927 bp open reading frame that may represent the full coding sequence for the DIP1 protein. It is not certain if this cDNA contains the complete 5' untranslated region of full-length *dip1* cDNA. Sequence analysis through 'BLAST' search (Altschul et al., 1990) showed that the putative 309 amino acids DIP1 protein contains an additional dsRNA-binding motif N-terminal to the motif predicted before. Double stranded RNA binding domains (dsRBD) are usually about 70 bp long. A
consensus sequence for dsRNA binding domains has been determined (St. Johnson et al., 1992) from seven copies of the dsRBD motifs in Xlrbp, Staufen and human TRBP proteins. Each full-length motif consists of an N-terminal large domain and a C-terminal short domain. Two types of dsRBDs have been recognized and are denoted as type A and type B (St. Johnston et al., 1992). The type A dsRBDs show strong similarity to the entire length of consensus sequence defined for dsRBDs. While type-B dsRBDs match well with the basic C-terminal short domain consensus, but show only poor similarity to the N-terminal long domain consensus. The putative DIP1 protein contains two type B dsRBDs.

Figure 13 shows alignment of the putative dsRNA binding domains of DIP2 with that of other proteins known to bind dsRNA. The two dsRNA-binding motifs of the DIP1 protein share significant similarity with the C-terminal consensus, but, match poorly with the N-terminal large domain consensus. The dsRNA binding domains of DIP1 share striking similarity with that of mammalian glutamate receptor editases (Melcher, T. et al., 1996; Lai et al., 1997). The similarity extends far beyond the consensus for dsRNA binding motifs (Figure 14). In addition, the predicted protein also contains a putative bipartite nuclear localization signal similar to that present in the glutamate receptor editases. Bipartite nuclear localization motifs are composed of: two basic amino acids, a spacer region of any ten amino acids followed by a basic cluster of five amino acids, three of which must be basic (Dingwall and Laskey, 1991). Figure 11. shows that the putative nuclear localization signal in the predicted DIP1 protein (KRKKSSERTRDKKLRQN) meets this criterion. The sequence similarity between the nuclear localization signal and the dsRBDs of the predicted DIP1 protein with mammalian glutamate receptor editases are shown in Figure 14.
Figure 11

Sequence of the dip1 cDNA that contains the full coding region:
This Figure shows the sequence of the dip1 cDNA (about 2.6 kb) isolated from the λgt11 library. Eight repeats in the 3'UTR are shown. The first three nucleotides of each repeat are shown in (boldface). The repetitive region ends with an EcoRI site (GAATTTC, shown in boldface and is underlined). The EcoRI site is followed by a 396 bp sequence and a polyA tail. The 348 bp sequence preceding the polyA tail is unique to this dip1 cDNA. The predicted protein sequence is also shown. The putative start codon is located 263 bp downstream from the 5' end of the cDNA. The stop codon (*) is located at the very beginning of the first repeat (at nucleotide position 1188). The putative bipartite nuclear localization signal (NUC) in the predicted protein (from amino acids 45-62) is underlined. The consensus for NUC consists of two basic amino acids, a 10 amino acids spacer region and a cluster of five amino acids out of which at least three should be basic. The basic amino acids in the bipartite signal are shown in boldface. Two putative dsRNA-binding domains of the protein are highlighted.
Sequence of dipl cDNA and the predicted DIP protein:

ACGAATTTCTTCTTGCTGACTGAGAGCTGAAAAATTAACTTACAAAGGGAAAGAACATT 60
ACAAAGCCGCAATGTCAGAGGACAGAGCTCTCTCTCGGTGCTTTATATCCGAGATCG 120
GCTCTCCGCTGCGGCCATCAGCCACAGCGGACATTTTTGCTGTCGCTGGCGCAAGG 180
GCTTAGTCGGCCCGTCCTGACGCGACAGATACTGGCCAATCCGAGATGGCGAGAC 240
GCCATTGCTCTGCAATTGCGCAATGCGGAGATGCTCTGCGCAAGATTCAACCAAGCCAAGGCGATGCT 300
MPVIKKEEPISVN 13
GATGAAACCGATCGCGACAAATATAGAGGACAAATACCGGCTGCTGCTAGTGCCGACG 360
DEPSVINDTSASTSASG 33
ATCCGGGGCAAGATCCCTTATAAAAAAAAACTTTCCAAAGCGCAAGATGCTGTCGACG 420
IGGKIKIFKIKKKSSER 53
ACTCGGGACAAAGAGATTTGCGACAGAACCCGAGCTGGCAGATCCATCAGCTCCAGAGC 480
TRDKRLQNLRLKSKMLPKN 73
GCTCTGATGGCCCTTCACAAAGGTGGAAGGCGGCTGACATCGAGATCTACAGCGAGAC 540
ALMNEVKGVGTISDFTIDS 93
AATAACGAGCGGGGATTCAGCGCCTGTCATGCTACTGACATCGAGATCTACAGCGAGAC 600
NTDGFTAVVTNVSNQYEGK 113
GCTACCTGAAATGACGACCAAGACACGGCCTGCGGAAAGCTGCTGCGGCGATTTATT 660
GTKMTKMTKNAACEKAWRDFI 133
ATTGAACAAATAGCCCGGCGATTCCAGGTGGAGATGGTGTTTCGGAGCA 720
IAKMTKPMPPRIHQVEMGSEP 153
ATGGATATCAACGGAGATGAGCGTCAGACGATTGATGACTGGCTGCATGCAAAATGCAACG 780
MDINEDEADAPDLPLMNL 173
GCCCTGCTGGCTACATCAGGCCTGCAGTGCAGTGAGCTGCTGCTGCTGCTGCTGCTGCTGCTGCT 840
ASFAIYLFADMREWEAREGYVVP 193
GAGATGCACCTTTGGCACCATAATCTGCGCCCCAAACAGCGGAGGAGTAGTCGGCGAATCTCA 900
EMHPSANAAQAAGGDAGTPV 213
CCCCCCTGTCGGACAGGGACCAAAAGAGACCGGCGATGCCGCAACCCGCTAGCTCGCTCGG 960
PPVPKPPVKPPVRTELPSGW 233
GAGACCATGCACTCGGGCAGCATTTTCTGCTACTATGCTGCGGACTCAACTACGATGC 1020
ETMPATILCIMRPGLNYVD 253
TACGGGCTATCCTGCGGACAGACCACGCTGACATCGTGGGAAATCGTGGTGACCAAC 1080
YGSSGDKTNGMQHLGIMVDN 273
CAGGAGTCAGCAGGCAAAGGCAGATCAAAGAGATTGCCGGCGACCTGCGCGCA 1140
QEFHANGRSSKIIARNVAVK 293
GTGTGCACTCTCTCTCTCTGCGACCAACCATCTTCCACACGCGACACCTTTAAAGCAACACAAC 1200
VCNSLFGNTFYSDT 309
CTCCACCACCAGACTTCCACGGGCAGACCAAAATCATAGACAACTCTACCTTAACGCAGAC 1260
CAATGCAAGAATTGCGACCCTTGTGCTGACATTGGCGCAACTTAAAGCAACACATTAAGAC 1320
ACCAACTTACAGGACGTCCAGCTGGCGGACGCAACACACCTACCTGGCTGGCCTTAACCCG 1380
ACAGCAATCTAGAGGTGGAGCCCTCTCGACTGCGTGCCGCAACTTAAAGCAACACCTTGT 1440
AGACACAACTCCAACAGTGCGGACAGCACACACATCGTGGCTGGCCTTAACCCG 1500
ACAGCAGACAAATCAGAGGATTGCGACCCTGCTGACATTGGCGCAACTTAAAGCAACACACAGC 1560
CTTAGAACAACCTCAACCCAGCGGACTCAAACACCGGACGCAACACACATCGTGGCCTTAACCCG 1620
CTTACAGGAGGCAATCAGAGGATTGCGACCCTGCTGACATTGGCGCAACTTAAAGCAACACACAGC 1680
ACCACCTTAAAGCAACCCAGCGGACTCAAACACCGGACGCAACACACATCGTGGCCTTAACCCG 1740
CTTACACAACTCCCGAGCAATCAGAGGATTGCGACCCTGCTGACATTGGCGCAACTTAAAGCAACACACAGC 1800
CGACACACTCTTAAAGCAACCCAGCGGACTCAAACACCGGACGCAACACACATCGTGGCCTTAACCCG 1860
ATGGCGCTCTTAAACGGGACAGCAATCAGAGGATTGCGACCCTGCTGACATTGGCGCAACTTAAAGCAACACACAGC 1920
TAAGCAGACACTTAAAGCAACCCAGCGGACTCAAACACCGGACGCAACACACATCGTGGCCTTAACCCG 1980
ACCTTAAGCAGACACTTAAAGCAACCCAGCGGACTCAAACACCGGACGCAACACACATCGTGGCCTTAACCCG 2040
ACCTTAAGCAGACACTTAAAGCAACCCAGCGGACTCAAACACCGGACGCAACACACATCGTGGCCTTAACCCG 2100
ACACACCTATGGCCTCTTTAAACGGGACAGCAATCAGAGGATTGCGACCCTGCTGACATTGGCGCAACTTAAAGCAACACACAGC 2160
CGACACCTTAAAGCAACCCAGCGGACTCAAACACCGGACGCAACACACATCGTGGCCTTAACCCG 2220
AATTTAGTTTGTATTTGGGACAGCTTTACCTTTAGGGGTTAGTGTCGGCCGAGCCGTT 2280
TTCAACACGAAATAATTTTGAAACTGCAAGAGCCGATAGGTTACCCGATATCTAAACC 2340
AGGATTTCTTTGGATATCTGATAATTTTTTTCCGTTATTTTTGGACACACTGTTGTTT 2400
GGTTTTTTGAAAACGCTTCACAGAAGAGGATTTTCGATGCGAGTTGGGTATCAATAT 2460
TAATTATGCTTACCGTGAATTAATTTTTGTCGTAACAGTTAGTAAGCTAAATGGGGG 2520
TATAATTACTTTGATTTTTAAAAACATGAGACATAAGTGACATGTATTTGATATAAAAA 2580
Figure 12

Difference in the 3′UTR of the dip1 cDNAs:

Panel A shows the features of the two dip1 cDNAs. In this Figure, the cDNA (1104 bp) isolated from the yeast interaction trap library is denoted as cDNA\(^1\). It is a partial cDNA that lacks the 5′ end of the corresponding mRNA. The cDNA isolated from the λgt11 library is denoted as cDNA\(^2\). The exact size of the cDNA is not known. It contains at least 8 repeats. The first and the last repeats of cDNA\(^1\) and cDNA\(^2\) are identical. The last repeat is interrupted with an EcoRI site. The numbers of middle repeats are different in the two cDNAs. All of the middle repeats in either cDNA have exactly same nucleotide sequence. In cDNA\(^2\), the location of the putative start codon at 263 bp and the stop codon at 1188 bp from the 5′ end of the cDNA are indicated with vertical lines. The location of the putative nuclear localization signal coding sequence (nuc, nucleotides 396-447 and the two dsRNA-binding motif coding sequence (nucleotides 462-660 and 1020-1164 respectively) are shown in cDNA\(^2\). The position of the C-terminal dsRNA-binding domain coding sequence present in cDNA\(^1\) is also shown. Panel B shows the sequences of the individual repeats. The first repeat is 126 bp long. The nucleotide sequences of all the 124 bp middle repeats in either cDNA are exactly equivalent. The last repeat is 121 bp long. The nucleotide sequence of the first repeat is different from that of other repeats in the region that is underlined.
B

First repeat (126 bp)
CACCACCTTAAGACACCAACTCTCAAACCGCAGACTCTCAACTGCCGACGACAACACACACATAACCGGACA
CTCTTAACGGACAGCAATCAGAAGATTGCAGACCCCTGACGTCAATGGCAGCAACCTTTAAGCGA

Middle repeats (124 bp)
CACCACCTTAAGACACCAACTCTCAAACCGCAGACTCTCAACTGCCGACGACAACACACACACCTATGGCCTTCTC
TTAACGGACAGCAATCAGAAGATTGCAGACCCCTGACGTCAATGGCAGCAACCTTTAAGCGA

Last repeat (121 bp)
CACCACCTTAAGACACCAACTCTCAAACCGCAGACTCTCAACTGCCGACGACAACACACACACCTATGGCCTTCTC
TTAACGGACAGCAATCAGAAGATTGCAGACCCCTGACGTCAATGGCAGCAACCTTTAAG
Figure 13.
Alignment of the putative dsRNA binding domains of DIP1 with the RNA-binding domain consensus:
The top portion shows the predicted protein sequence for dip1. The positions of the two dsRNA-binding domains (67-135 and 234-301) are shown in blue.

The bottom part shows the alignment of the putative RNA binding domains of DIP1 with those of known dsRNA binding proteins and the consensus for dsRNA binding domains (dsRBD). In the consensus, the uppercase letters indicate the most frequently found conserved residues, while the lower case amino acids are conserved in relatively fewer number of dsRBDs. The consensus sequence (about 70 amino acids long) has two distinct segments, denoted as N-terminal long domain consensus (shown in green), and C-terminal short domain consensus (shown in red). The dsRNA-binding domains of DIP1 represent type-B dsRBDs that match well with the C-terminal short domain consensus (orange), but show only a weak similarity with the N-terminal long domain consensus. The dsRNA-binding domains used for the alignment are as follows: first dsRBDs of Drosophila Staufen-1 and human TAR-binding protein hTRBP1; the dsRBD of the mammalian transcription factor NF-NT90 and two dsRBDs of the glutamate receptor editases hRED1-1 and hRED1-2.
Figure 14
Alignment of the predicted DIP1 protein with the N-terminal region of mammalian glutamate receptor editases:

This Figure shows the alignment of the putative nuclear localization signal (nuc) and dsRNA binding domains of DIP1 with those of human and rat glutamate receptor editases (hRED1, and rRED1 respectively). The putative DIP1 protein shares similarity with the bipartite nuclear localization signals (nuc) of hRED1 and rRED1. Identical amino acids are shown in blue (purple), conserved changes are shown in violet (pink) and similar amino acids are shown in yellow. The consensus for dsRNA-binding domains (shown in red) is underlined. It is evident that the similarity between the dsRBDs of DIP1 and glutamate receptor editases extends beyond the conserved amino acids.
C. Northern analysis for dip1 transcripts:

To examine the size of the dip1 transcript(s) and detect expression of alternatively spliced forms (if any) of dip1, northern analysis was done. Northern analysis was done with polyA⁺ RNA isolated from 0-22 hours embryos of Drosophila.

Two different probes for dip1 cDNA (with and without the repeated region) were used for northern analysis. Three bands of different size were obtained with the probe lacking the repeated region, indicating the expression of at least three different transcripts for dip1 during embryogenesis. The sizes of these transcripts are about 3 kb, 2.2 kb and 1.8 kb (Figure15). The cDNA isolated from the λgt11 vector is larger in size than the middle transcript seen on the northern blot. It might represent a partial form of the largest transcript.

Similar sized bands were detected in the northern blots hybridized with probes made from the dip1 cDNA containing both the repeated and the unique regions. However, the signal was stronger and more diffused when the probe with the repeated region was used (data not shown), indicating the presence of other RNA species that can be detected with the repeated region only.

D. Chromosomal location of the dip1 gene:

To understand the physiological role of the dip1 gene, isolation of mutants in which dip1 gene function is disrupted is essential. In order to identify existing mutants with mutation in a particular gene, the knowledge of its chromosomal location is important. Due to the presence of large polytene chromosomes in Drosophila, the chromosomal location of a gene can be readily determined by hybridizing the corresponding probe to a polytene chromosomal preparation.
made from salivary glands of third-instar larvae. There are about 5000 polytene bands and the limit of cytological resolution with in-situ hybridization is about 50 kb (Sorsa, 1988). The in-situ hybridization to polytene chromosome was performed with a biotin labeled *dip1* probe. The 1.1 kb *dip1* cDNA fragment isolated from the yeast interaction trap library was used as template to make the biotin labeled probe.

The in-situ hybridization studies localized the *dip1* signal to two locations on the polytene chromosome; band 20A at the base (adjacent to centromere) of the X-chromosome and band 91F on right arm of the third chromosome (Figure 16A). Since the results of northern analysis suggested the possibility of presence of the repeated element of the *dip1* cDNA in other *Drosophila* messages as well, one of the two hybridized regions on the polytene chromosome may represent location of another gene containing the same repeated sequence. To test this hypothesis, an in-situ probe was generated from the non-repeated region of the *dip1* cDNA. This probe hybridized to only band 20A (Figure 16B), confirming that the repeated sequence seen in the *dip1* cDNA is not unique to the *dip1* gene, but, is also present in at least another gene located on the right arm of the third chromosome (Figure 16A). These results implicated the location of the *dip1* gene to be on the band 20A of X-chromosome of *Drosophila*. 
Figure 15

Northern analysis for dip1 mRNA:

This Figure shows the result of northern analysis for dip1 mRNA. Poly A⁺ RNAs isolated from Drosophila embryos were hybridized to a probe made from the nonrepetitive region of the dip1 cDNA. Three distinct bands of approximate molecular weights 3.0 kb, 2.2 kb and 1.8 kb can be seen.
**Figure 16**

Localization of the *dip1* gene on *Drosophila* polytene chromosome:

Panel A shows that the probe made from the *dip1* cDNA containing both the nonrepetitive and repetitive regions hybridizes to two polytene bands; the band 20A located at the base of the X-chromosome (empty arrow) and band 91F on the right arm of *Drosophila* chromosome III (solid arrow).

Panel B (next page) shows that the probe made from the nonrepetitive region of the *dip1* cDNA hybridizes to only one region of the polytene chromosome: band 20A of the X-chromosome.
E. Deficiency mapping of the dip1 gene:

In order to determine the genetic location of the dip1 gene, deficiency mapping was done. This was done by southern analysis of deficiency lines that carry a deletion in the 20A region of the X-chromosome. The presence or absence of the dip1 gene was followed by a restriction fragment length polymorphism that exists in this region. Since homozygosity of deficiency chromosomes lead to lethality, heterozygous lines in which deficiency chromosomes are kept either under a wild-type or a balancer chromosome were used. For southern analysis the chromosomal DNA was digested with EcoRI or BamHI and hybridized to a probe made from the dip1 cDNA. Due to the polymorphism, the wild-type, balancer and deficiency chromosomes generated distinct restriction patterns on southern blots.

The results of this study are shown in Table 1. Here, '-' indicates deletion and '+' indicates inclusion of the gene in the deficiency chromosome. The genotypes of the deficiency lines are shown in the 'Materials and Methods' section of this chapter.

**Table 1: Results of the deficiency mapping for dip1:**

<table>
<thead>
<tr>
<th>Deficiency</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Df(1)DCB1-35b</td>
<td>-</td>
</tr>
<tr>
<td>Df(1)17-137</td>
<td>-</td>
</tr>
<tr>
<td>Df(1)16-2-13</td>
<td>-</td>
</tr>
<tr>
<td>Df(1)R22</td>
<td>-</td>
</tr>
<tr>
<td>LB6</td>
<td>-</td>
</tr>
<tr>
<td>Df(1)S54</td>
<td>-</td>
</tr>
</tbody>
</table>
Df(1)17-257 | -  
Df(1)A209 | +  
Df(1)HM430 | +  
Df(1)R21 | +  
Df(1)B12 | +  
Df(1)B1-35C | +  
Df(1)R38 | +  
Df(1)GA22 | +  

The southern blots from this study are shown in Figure 17. Deficiency map of the dip1 gene is shown schematically in Figure 18. The deficiency map of the dip1 gene is similar to that of a gene called flamenco (Prud'homme et al., 1995). Furthermore, the deficiency maps of the genes located on either side of flamenco (extra organ (eo) and touch insensitive larvae B (tilB)) do not match with that of the dip1 gene. These findings raise the possibility that flamenco and dip1 represent the same gene (additional evidence is presented in the discussion section). Therefore, through deficiency mapping we were able to narrow down the genetic location of the dip1 gene in between eo and tilB.
Figure 17

Results of the deficiency mapping of dip1 by southern analysis:

This Figure shows three southern blots that determined the genetic location of the dip1 gene. Panel A, B and C show EcoRI digested chromosomal DNA isolated from flies heterozygous for deficiency chromosomes and wild-type (CS, Canton S) or balancer chromosomes. The CS, balancer and deficiency chromosomes produced restriction fragments of distinctly different sizes when hybridized to a $^{32}$P labeled dip1 probe. In the control lane containing only EcoRI digested DNA from homozygous wild-type flies (CS), a single band is hybridized to the dip1 probe. In each of the two lanes, R21/CS, and B12/CS two bands, one corresponding to the CS chromosome and one corresponding to the deficiency chromosome are hybridized, indicating that these three deficiencies do not delete the dip1 gene. In the lane B1-35c/CS, the bands corresponding to the CS and the deficiency chromosomes show almost same electrophoretic mobility. However, in the B1-35c/balancer lane two distinct bands, one corresponding to the deficiency chromosome and one corresponding to the balancer chromosome are seen indicating that this deficiency does not delete the dip1 gene either. In the lanes, 16-2-13/CS, R22/CS, LB6/CS, S54/CS, B1-35b/CS, R22/CS, 17-257/CS, only one band corresponding to the CS chromosome can be seen. Similarly, in lanes where the same deficiencies are over a balancer chromosome, only one band corresponding to the balancer chromosome can be seen. Therefore, the above deficiencies delete the dip1 gene. The balancer chromosome used for the 16-2-13 deficiency is different from the FM7 balancer used for
other deficiencies. The electrophoretic mobility of the EcoRI fragment from this chromosome is close but distinctly different from that of CS fragment. The electrophoretic mobilities of the R38 and GA22 deficiency chromosomes are identical to that of CS. However, presence of two bands in the R38/FM7 and GA22/FM7 shows that these deficiencies do not delete the dip1 gene.

**Panel D** shows another southern blot which was done with chromosomal DNA digested with two different enzymes, EcoRI (lanes 1-5) and BamHI (6-10). Both EcoRI digests and BamHI digests of B1-35b/balancer and 17-137/balancer show only one band corresponding to the balancer chromosome. Therefore, B1-35b and 17-137 delete the dip1 gene. Whereas, A209/balancer and HM430/balancer show two bands each, one corresponding to the balancer chromosome and the other corresponding to the deficiency chromosome. Therefore, these deficiencies do not delete the dip1 gene.
Figure 18

Deficiency map of the dip1 gene:

This Figure shows a schematic representation of the deficiency map of the 20A region of the X-chromosome. The relative locations of several genes that map to the 19A to 20D region are shown. These are, extra organ (eo), flamenco (flam), touch-insensitive larvae B (tilB), wings apart (wap), introvert (intro), l(1)20Ad, stoned A (stnA), suppressor of forked [su(f)], A112, LB20, tumorous head (tuh1), maroon-like (mal), melanized (mel) and l(1)carot2. The extent of the deleted region for each deficiency chromosome is indicated with a horizontal line. The deficiencies that delete the dip1 gene are indicated with asterix. The map shows that the genetic location of the dip1 gene matches well with that of the flam gene.
F. Expression of the dip1 mRNA in the follicle cells of the ovary:

The possibility of dip gene being flamenco was examined by determining its expression in the follicle cells of the ovary, the previously identified site of flam gene function (Song et al., 1997). To that end the wild-type expression pattern of the dip1 mRNA was examined. This experiment was done in collaboration with Dorothy DeSousa, a graduate student of the lab.

During oogenesis (reviewed by Spradling, 1993), Drosophila ovarioles contain a series of progressively older egg chambers that represent 14 successive developmental stages. Each egg chamber contains cells from germ line origin as well as somatic cells. The cells of germ line origin consists of 16 interconnected sister cells, the posterior-most of which is the oocyte (OO) and the other 15 differentiate into giant polyploid nurse cells (N). The cells of somatic origin are called follicle cells. These cells form an epithelial sheath that completely covers the egg chamber. Ovarioles containing egg chambers from various stages were dissected from young Drosophila females (2-3 days old) to study dip1 expression pattern. In wild-type ovarioles we were able to detect dip1 expression in the follicle cells (arrow) and nurse cells of the ovary (Figure 19, Panel B). The arrow shows the localization of the dip1 gene expression at the basal level of the columnner follicle cell layer just adjacent to the oocyte.
Figure 19

Expression pattern of *dip1* mRNA in the ovary:

Panel A shows one ovariole that contains egg chambers from different stages of oogenesis. Expression of the *dip1* gene can be seen in the nurse cells and the follicle cells at all stages. Panel B shows one egg chamber from stage 10 at higher magnification. Expression of the *dip1* mRNA is clearly seen in the nurse cells (N) and at the base of the columnar follicle cells (arrow with a white arrowhead), that surround the oocyte.
G. Screening of genomic P1 and cosmid clones for the dip1 gene:

A number of genomic P1 and cosmid clones that map to the 20A region of the polytene chromosome of Drosophila were screened in an attempt to sequence the genomic region of the dip1 gene.

The genomic P1 clones for Drosophila were generated by infecting its bacterial host E. coli NS3145 with P1 phage vector containing Drosophila genomic DNA fragments (Hartl et al., 1994). This modified P1 phage is lysis defective. It is replicated and maintained inside its host as a plasmid vector. The P1 vector can hold foreign DNA as large as 100 kb. Due to this advantage, a Drosophila melanogaster genomic DNA library was constructed in this vector (Hartl et al., 1994).

A physical map of the Drosophila melanogaster genome has also been constructed with about 20,000 cosmid clones and the available P1 clones (Siden-Kiamos et al., 1990). Screening of 32 P1 clones and four cosmid clones that map to the 20A region of Drosophila polytene chromosome showed absence of the dip1 gene in any of those clones.

The results of this study are shown in Table 2. The ‘−’ sign indicates absence of the dip1 gene in the genomic clones.

Table 2: Results of the screening for dip1 genomic clones.

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IV. Characterization of dip2

A. Screening of λgt11 and λEXlox cDNA libraries for dip2 cDNA:

A partial 1.2 kb 3' end fragment of the dip2 cDNA was isolated from the yeast interaction trap library vector. In an attempt to isolate the 5' portion of the cDNA we screened two other Drosophila cDNA libraries made in λ phage vectors. A 12-24 hours Drosophila embryonic cDNA library made in λgt11 phage vector and 20-22 hours embryonic cDNA library made in λEXlox (Novagen) vector were screened. These libraries were screened with a radiolabeled probe made from the 1.2 kb dip2 cDNA fragment isolated from the yeast interaction trap library. Five positive plaques from the λgt11 library and four positive clones from the λEXlox library were isolated. The size of the cDNA inserts were examined by restriction digestion of the phage DNA from the positive plaques isolated from the λgt11 library with EcoRI, and that of λEXlox library with ApaI and SacI. All five positive plaques from the λgt11 library contained only a small fragment, about 1 kb in length corresponding to the 3' end of the dip2 cDNA. One of the four positive plaques isolated from the λEXlox library contained a 2.5 kb fragment also from the 3' end of the dip2 cDNA. The other three positive plaques contained cDNA inserts of larger that 2 kb, but smaller that 2.5 kb, all corresponding to the 3' end of the dip2 cDNA. The largest fragment isolated from these libraries was 2.5 kb, which lacked the 5' end of the dip2 cDNA. Thus, none of the three libraries (including the yeast interaction trap library) screened contained a full-length cDNA clone for the dip2 gene.

In collaboration with an undergraduate (Peter Pelka) in the lab the mouse homolog of the dip2 gene was also cloned. Screening of a 8.5 day old mouse
embryonic cDNA library made in λgt10 led to the isolation of a single clone with a partial 1.1 kb 3' end fragment of the mouse dip2 cDNA. The probe used for this study was made from a 4.6 kb partial human dip2 cDNA that was cloned in the human genome project from the human cell line KG1 (Accession# D80006). Sequence analysis of the 1.1 kb mouse cDNA fragment showed that the clone isolated contained a recombinant cDNA with only a 382 bp sequence from the middle portion of the 1.1 kb fragment representing the desired cDNA. This 382 bp fragment corresponded to the 3' end of mouse dip2 cDNA. A 124 amino acids sequence predicted from the 382 bp 3' end fragment showed 100% amino acid identity with that of its human homolog (Figure 26). The mouse and the corresponding Drosophila fragments shared 82.3% amino acid identity and 97.6% positives. The sequence of the partial mouse cDNA is shown in Figure 25.

B. Northern analysis for dip2 transcript:

Northern analysis was done to determine the size of the dip2 transcript expressed during Drosophila embryonic development. The analysis was done with polyA+ RNA isolated from 0-22 hours embryos. The probe used for this study was made from the 1.2 kb dip2 cDNA fragment isolated in the yeast interaction trap. The probe hybridized to a single band. As judged from the northern blot, the size of the single dip2 transcript is about 5 kb (Figure 20).
Figure 20

Northern analysis for dip2 mRNA:

Result of the northern analysis done with polyA⁺ RNA isolated from 0-22 hours Drosophila embryos is shown. The dip2 probe hybridized to a single band of approximately 5 kb in length.
C. Chromosomal location of the dip2 gene:

In order to understand the physiological role of dip2, knowledge of its chromosomal location is essential. In-situ hybridization to polytene chromosomes of Drosophila was done to determine the chromosomal location of the dip2 gene. This was done using a 1.2 kb biotin labeled dip2 probe isolated from the yeast interaction trap library.

The dip2 probe hybridized to a single location on the polytene chromosome, the band 61B, at the tip of the left arm of the third chromosome (Figure 21).

D. Deficiency mapping of the dip2 gene:

In order to narrow down the chromosomal location of the dip2 gene its genetic location was determined through deficiency mapping. This was done through southern analysis of chromosomal DNA from deficiency lines from the 61B region, the location of the dip2 gene on polytene chromosomes. Only three deficiency lines were available in Drosophila stock centers with breakpoints in the vicinity of the 61 B region. These were; Df(3L)Ar12-1, Df(3L)Ar11 and Df(3L)emc-E12. Unlike the dip1 locus we did not detect a polymorphism at the dip2 locus. Therefore, to identify deficiencies that delete the dip2 gene, the amount of DNA loaded in each lane was estimated by rehybridizing the experimental blot with a control probe (disco probe). This allowed us to determine if one copy of the dip2 gene was deleted in any of the heterozygous deficiency lines. Absence of one copy of the gene led to a reduction in band intensities in the experimental blot as compared to the control blot. Since the disco gene is an X-linked gene, only one copy of the gene is present in the male as
opposed to two copies in females. Therefore, in order to be able to use disco as a control probe, only chromosomal DNA of heterozygous female flies from the above deficiency lines was used.

The deficiencies Ar12-1 and Ar11 carry deletions from bands 61C to 61F. Whereas, Df(3L)emc-E12 is the only one in which the 61B region is deleted. As expected from the in-situ result, Df(3L)emc-12 was the only deficiency from the 61 region that deleted the dip2 gene. The results obtained with these deficiencies are shown in table 3, where ‘-‘ indicates deletion, and ‘+‘ indicates inclusion of the dip2 gene in the deficiency.

Table 3: Results of the deficiency mapping for dip2:

<table>
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<th>Result</th>
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<tr>
<td>Df(3L)Ar11</td>
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</table>

Due to the absence of smaller deficiencies from the 61B region we were unable to further narrow down the genetic location of the dip2 gene. The southern blots from this study are shown in Figure 23. The deficiency map is schematically represented in Figure 22.
Figure 21

Localization of the dip2 gene on Drosophila polytene chromosome:

This Figure shows the in situ chromosomal localization result for the dip2 gene. The dip2 probe hybridized to band 61B (arrow) on the left arm of chromosome III of Drosophila.
Figure 22

Results of the deficiency mapping of dip2 by southern analysis:

This Figure shows the southern blots done to identify a third chromosome deficiency (emc-E12) that delete the dip2 gene. Panel A shows the southern blot done with the EcoRI and BamHI digested chromosomal DNA isolated from the heterozygous deficiency lines that carry one third chromosome deficiency from the 61 region and one balancer chromosome. The intensity of the bands hybridized to the dip2 probe appears to be lower in the lanes corresponding to the deficiency line emc-E12 compared to other lanes.

Panel B shows the same blot hybridized to a disco probe as a control to examine the amount of DNA loaded in each lane. The amount of DNA loaded in the lane containing chromosomal DNA from the emc-E12 line does not appear to be less than that loaded in other lanes. This shows that the lower band intensities of the emc-E12 lane in the blot in panel A is due to the absence of one copy of the dip2 gene in that deficiency line. In other words, the deficiency emc-E12 deletes the dip2 gene.
Figure 23

Deficiency map of the dip2 gene:

The genetic location of the dip2 gene is shown in the deficiency map of the 61 region on the left arm of the third chromosome of Drosophila. The extent of the deleted region for each deficiency chromosome is indicated with a horizontal line. The deficiencies that were used for southern analysis are Df(3L)Ar12-1, Df(3L)Ar11 and Df(3L)emcE12. Only the deficiency emc-E12 deletes the dip2 gene (shown with an asterix). Mutations that map to the 61A-D region are shown. These are tracheless (trh), marble (marb), extra macrochaete (emc), smell impaired 61A(smi61A), pointless (ptl), forewheel drive (fwd), l(3)122 and l(3)06240.
E. Isolation of genomic P1 clones for the dip2 gene:

In order to sequence the genomic region of the dip2 gene, three genomic P1 clones that map to the 61B region of Drosophila polytene chromosome, were screened. These are shown in Table 4. Of the three, two (DS05688 and DS03215) contained the chromosomal DNA for dip2. Because of technical difficulties only a portion of the dip2 genomic DNA was sequenced (Figure 24). A portion of the P1 clone sequencing was done partly by Peter Pelka, a graduate student of the lab. The results of the sequence analysis are presented following this section.

Table 4: Results of the screening for dip2 genomic clones: In this table ‘+’ indicates inclusion and ‘-’ indicates absence of the dip2 gene in the P1 clones.

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<td>DS00583</td>
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F. Sequence analysis of the dip2 genomic DNA:

A 2.7 kb fragment of the dip2 gene was sequenced from the P1 clone DS03215. This 2.7 kb sequence corresponds to the 3’ end of dip2 mRNA. BLAST search analysis of the dip2 genomic DNA sequence determined from the clone DS03215 showed sequence homology with a human and a C. elegans gene, both isolated from the corresponding genome projects.

The 2.7 kb sequence has a continuous open reading frame that stops at the nucleotide position 2509 (Figure 24). Both, the genomic sequence and the partial
dip2 cDNA sequence of Drosophila showed the presence of an extra 159 bp fragment (124 bp upstream of the stop codon) which was absent in any other dip2 homolog. Absence of a putative splice acceptor site (Padgett et al., 1986) adjacent (5') to the 159 bp genomic sequence, and a splice donor site (usually G) at the 3' end of this sequence also indicates that this region of the dip2 gene constitutes part of dip2 cDNA coding sequence. Thus, the predicted Drosophila DIP2 protein contains an extra 53 amino acid sequence that is lacking in other DIP2 homologs.

The protein predicted from the putative open reading frame show no significant similarity to any known functional domain. BLAST search analysis (Altschul et al., 1990) for DIP2 homologs show a weak similarity to the AMP-binding motif (PROSITE database, accession number PDOC00427) found in several enzymes such as insect luciferase, acetyl CoA ligases, 4-coumarate-CoA ligase and others. 'Block' database (Henikoff and Henikoff, 1994) searches show that the amino acid sequence of the C. elegans DIP2 homolog (for which the complete protein sequence is known) shows weak similarity to an N-terminal 30 amino acids fragment of adenosine deaminases and a similar fragment from AMP-deaminases (BLOCK number BL00485A). The sequence of the dip2 gene determined from the P1 clone is shown in Figure 24.
**Figure 24**

Partial *dip2* genomic sequence:

The sequence of the 2.7 kb partial *dip2* genomic region determined from the P1 clone DS03215 is shown in this Figure. It contains a continuous open reading frame that stops at the nucleotide position 2509 of the fragment. The amino acids sequence predicted from the open reading frame is also shown. A 53 amino acids region (residues 659-712) that is unique to the *Drosophila* DIP2 homolog is highlighted.
Partial genomic sequence for dip2

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<td>GACATTTCTCCTCTTATATTGTTATAGTGTACAAAGCTCTCAGCAAATCTTATA</td>
<td>2355</td>
</tr>
<tr>
<td>DTFC G SV IE L CTK ALSNSI</td>
<td>2405</td>
</tr>
<tr>
<td>CTTTCCTAAAGCAGAAACATAGATTTGGAGATGCTACGAGTGTCGTCCTGTTGGGCA</td>
<td>2455</td>
</tr>
<tr>
<td>PSKLQNRDLRCLCVRTCVVVA</td>
<td>2505</td>
</tr>
<tr>
<td>GAAGAGCGCCGCGAGATGCAACTGACCCACGAGACTTTCTGCAAGCTATTTCACCAAGGGCCTTGTG</td>
<td>2555</td>
</tr>
<tr>
<td>EERPVR VQL TQQQFCKLFQLG</td>
<td>2605</td>
</tr>
</tbody>
</table>
Figure 25

Partial mouse dip2 cDNA sequence:

This Figure shows the partial cDNA sequence corresponding to a 382 bp 3' end fragment of the mouse dip2 homolog. The predicted amino acids sequence corresponding to this fragment is shown. The stop codon is indicated with an asterix.
Partial mouse cDNA sequence and the predicted amino acids sequence:

GATGCACGTATGTCGTTGGTCTCTGGATGAAACGCTAGAAGAGGACATGCCTAT
D A L Y V V G S L D E T L E L R G M R Y
CACCACATGACATAGAAACCTCCGGATCCTGTCACACACAGGACATTCGAGATGTGC
H P I D I E T S V I R A H R S I A E C A
GTGTTCACCTGGACCAACCTGCTGGTTGTTGGTGCTGGAGCTGGATGGGCTGGAGCAGGAC
V F T W T N L L V V V V V E L D G L E Q D
GCCCTGGACCTGGTGGCCCTGGGACCAACGTCGTCGGAGGAGCACCTACCTGTGGTG
A L D L V A L V T N V V L E E H Y L V V
GGCGTGGTGGTCCATCGTGGATCCCCGGAGTGATCCCCATCAACTCTCGGCGGAGAAGCA
G V V V I V D P G V I P I N S R G E K Q
CCATGCAACCTCCGAGATGGCTCCTGGCTGACACGCTGGACCCCATCTACGTGGCCTAC
R M H L R D G F L A D Q L D P I Y V A Y
AATATGTGAGCCACAGCTG
N M *
Figure 26

Alignment of the predicted DIP2 homologs:

This Figure shows the sequence homology between the predicted DIP2 homologs from *Drosophila*, mouse, human and *C. elegans*. The top portion shows a schematic representation. The bottom portion shows actual protein sequences. Except for the *C. elegans* protein, all other DIP2 sequences presented here are partial. The protein alignment was done using Clustal W alignment program. The C-terminal 124 amino acids sequence is identical in all DIP2 homologs. Just adjacent to the 124 amino acids C-terminal fragment, there is a 53 amino acids sequence (shown in orange) that is unique to the *Drosophila* DIP2 protein.
Diagramatic representation of predicted DIP2 homologs

853 a.a. Predicted protein from 6539 b.p. partial dipi human cDNA

124 a.a. Predicted protein from 493 b.p. partial dipi mouse cDNA

337 a.a. COOH Predicted protein from 1233 b.p. partial dipi Drosophila cDNA

Predicted protein for dipi C. elegans homolog

---

Human Mouse Drosophila C. elegans

Human Mouse Drosophila C. elegans

Human Mouse Drosophila C. elegans

Human Mouse Drosophila C. elegans

Human Mouse Drosophila C. elegans

Human Mouse Drosophila C. elegans

Human Mouse Drosophila C. elegans

Human Mouse Drosophila C. elegans

Human Mouse Drosophila C. elegans

---

* Denotes 100% amino acid conservation

: Denotes a conserved amino acid change in at least one species
G. Search for P-element insertion mutants at the *dip2* locus:

To search for mutations that affect *dip2* gene function, P-element insertion mutants from the 61 region of the polytene chromosome were screened. In these mutants, disruption of gene function is achieved through insertion of the *Drosophila* P-element (transposable element) in the coding or the regulatory regions of a gene. The nine P-element insertion mutant lines were screened for this study include insertion in the genes denoted as *forewheel drive*, *marbles*, *pointless* and several other lethal mutations. None of these genes have yet been cloned and might represent *dip2*. For the detailed genotype of these lines please refer to the materials and methods.

In order to identify lines(s) with insertion inside the *dip2* locus, size difference among the EcoRI and BamHI digested chromosomal DNA bands from different lines that hybridized to a *dip2* probe were examined. Insertion of a P-element should change the length of a restriction fragment from the region of the insertion. The probe used for this study was synthesized from the 1.2 kb *dip2* cDNA fragment isolated in the yeast interaction trap assay.

None of the nine P-element insertion lines from the 61 region showed any difference in restriction pattern when compared to each other and to the wild-type chromosomal DNA restriction pattern. In these lines, no P-element insertion was detected at the *dip2* locus, using a probe made from the 3' portion of the *dip2* cDNA. However, these results do not exclude the possibility of an insertion at the 5' end or in the regulatory region of the gene. The results of the screen are shown in Figure 27.
Figure 27

Results of the screening of P-element insertion mutants for dip2:
This Figure shows the result of the southern analysis done on the P-element insertion mutant lines to screen for mutations at the dip2 locus. The P-element insertion lines are denoted with their Bloomington stock center numbers. For the actual genotype of these lines please refer to the materials and methods section of this chapter. None of the EcoRI or BamHI digested chromosomal DNA from any mutant line shows any difference in restriction pattern when hybridized to a dip2 probe made from the 3' 1.1 kb dip2 cDNA fragment as template. Bands hybridized to the dip2 probe from all mutant lanes are of same size as that obtained from the control lanes (digested chromosomal DNA from wild-type flies). For all mutant lines and the wild-type line, EcoRI digestion produced two bands of about 1.6 kb and 1.3 kb in length and Hind III digestion produced two bands of about 4.5 kb and 2 kb in length.
Discussion

The *disco* gene codes for a putative transcription factor:

The *disconnected* gene is required for proper development of the *Drosophila* larval visual system, and codes for a putative transcription factor with two Krüppel-like zinc-finger domains (Steller et al., 1987; Heilig et al., 1991). DISCO contains one complete C\textsubscript{2}H\textsubscript{2} and one incomplete and unusual CCH Krüppel-like fingers.

The C\textsubscript{2}H\textsubscript{2}-type zinc-fingers are known to bind DNA in a sequence specific manner (reviewed by Bernstein et al., 1994). The consensus for the C\textsubscript{2}H\textsubscript{2}-type motif is (Y,F)-X-C-X\textsubscript{2,4}-C-X\textsubscript{3}-F-X\textsubscript{3}-L-X\textsubscript{2}-H-X\textsubscript{3,5}-H-X\textsubscript{2,6}. In the 30 amino acids long motif, the seven conserved residues and the spacing between them are vital for its structure and DNA-binding activity (Bernstein et al., 1994; Berg, 1990). The two conserved cysteine and two histidine residues make direct contact with a zinc atom. Substitution of any of these four residues is sufficient to abolish the DNA binding activity of the domain. The *disco* gene function is completely disrupted by mutations that lead to substitution of one of the two conserved Cys residues in the zinc-finger domains of the encoded protein.

The possible role for DISCO as a transcription factor was first recognized from in-situ expression studies (Kevin Lee, Ph. D. thesis, 1993, Lee et al., 1999). These studies demonstrated an autoregulatory function of *disco* in maintaining its own mRNA expression in the optic lobe primordium (the target area for photoreceptor axons). Subsequent in vitro studies have demonstrated a direct binding of the zinc-finger domain of *disco* to its own promoter region. Furthermore, substitution of the first conserved Cys residue of the N-terminal

Although these studies were significant in understanding the disco gene function, the transcription regulatory function of DISCO has not yet been conclusively demonstrated due to the lack of any knowledge on DISCO targets. The results of the yeast two-hybrid assays reported in this chapter provide additional evidence that support the proposed transcription regulatory function of disco. It was observed that a LexA-DISCO fusion carrying the glutamate-rich (acidic) portion of DISCO, was able to activate transcription of the lacZ reporter gene from a yeast interaction trap-reporter gene construct (Figure 3). Activation of reporter gene transcription from this construct would otherwise need an activation domain. This result supports previous findings that have demonstrated the transcription activation ability of glutamate/aspartate-rich acidic domains in several transcription factors (e.g., Ma and Ptashne, 1987; Triezenberg, et al., 1988). Identification of a putative transcription activation domain in disco argues in favor of the proposed transcription regulatory function of DISCO.

The N-terminal DISCO bait generated for the yeast interaction trap assay uncovered an interesting property of the second zinc-finger domain of DISCO. It was observed that the wild-type zinc-finger domain of DISCO was sufficient to activate transcription of the reporter gene in the absence of any activation domain provided by tagged library proteins. The transcription activation ability of this bait was abolished when one of the conserved Cys residues of the second zinc-finger domain was substituted with a Ser residue.
A possible mechanism by which the second zinc-finger domain of DISCO allows activation of reporter gene expression is suggested by studies on other zinc-finger containing transcription factors (reviewed in Mackay et al., 1998). A number of these transcription factors contain several zinc-finger domains, some of which are involved in DNA binding, while others are dedicated to protein-protein interactions (Georgopoulos et al., 1997; Tsang et al., 1997; Merika and Orkin, 1995). For example, the transcription factor GATA-1 contains two CCCC-type zinc-finger domains, the C-terminal of which binds DNA, while the N-terminal domain interacts with zinc finger domains of other transcription factors including the sixth CCHC (an unusual Krüppel-like domain) zinc-finger domain of FOG (Friend of GATA) (Tsang, 1997) and the Krüppel-like zinc-finger domains of the transcription factors Sp1 and EKLF (Merika and Orkin, 1995). The Krüppel-like fingers in Sp1 and EKLF proteins appear to bind both DNA and protein. These observations suggest the possibility that the second zinc-finger domain of DISCO interacts with zinc-finger domains of other proteins to activate transcription. One may speculate that binding to an endogenous yeast transcription factor by the zinc-finger domain of DISCO may allow activation of the reporter gene expression.

Identification of DISCO partners:
Characterization of mutants in which disco gene function is disrupted has revealed a role for disco in photoreceptor axon pathfinding (Steller et al., 1987). Since disco expression is absent in photoreceptor cells, disco expression in the photoreceptor target area, the optic lobe, appears to be important for proper neuronal connectivity formation in the visual system. In fact, the differentiation
of optic lobe pioneer cells (OLPs) that are contacted by the larval optic nerve (LON) during larval visual system development, is disrupted in disco mutants (Campos et al., 1995). It has been proposed that expression of disco in these cells and in a subpopulation of glial cells, also contacted by the LON, direct the LON pathway formation. A tissue-specific autoregulatory activity of disco localized to the cells of the optic lobe primordium (Kevin Lee, Ph. D. thesis, 1993), and the importance of disco expression in this region (Lee et al., 1999; Campos et al., 1995), suggest that dissecting the disco autoregulatory pathway will be an important step towards understanding the role of disco in visual system connectivity.

To address the molecular mechanism of tissue-specific autoregulation of disco, we focused our study on identifying proteins that interact and cooperate with disco. To that end we used the yeast interaction trap technique (Gyuris et al., 1993). The expectation is that some of the proteins identified in the screen may represent molecules that regulate tissue specific disco function.

Using the interaction trap assay, two new genes and one previously known gene (dip1, dip2 and dip3 respectively) were identified (Figure4). Sixteen independent cDNA clones were isolated for dip1 alone. This suggests that the corresponding message is highly abundant in the embryo. Sequence analysis of four members of group 1 confirmed the single identity of the members of that group. All of them contained the same dip1 fragment, corresponding roughly to the 3' half of the dip1 mRNA and code for C-terminal half of the predicted DIP1 protein. This observation demonstrates that the C-terminal half of the DIP1 protein is sufficient for interaction with DISCO. The DIP1 fragment identified in the interaction trap screen contains a 111 amino acid region with no known functional domain that precedes the putative dsRNA-binding domain present in
the C-terminal portion of the predicted protein. Whether the 111 amino acid region contains a domain that mediates interaction with DISCO, has to be determined through deletion analysis. The specificity of the DIP1-DISCO interaction seen in yeast was further tested in an in-vitro binding assay. A direct physical association was demonstrated between GST-DISCO fusion proteins expressed in *E. coli* and DIP1 protein synthesized in vitro (Figure 7).

Unlike *dip*1 cDNA, the *dip*2 cDNA was isolated only once in the same interaction trap screen. The interaction was nevertheless quite strong as judged by the intensity of the blue colour produced on X-gal plates. Isolation of a single positive clone for *dip*2 may indicate a lower relative abundance of the *dip*2 transcript in *Drosophila* embryo compared to that of *dip*1 transcripts. Furthermore, the library used in this study was amplified several times. Presumably these amplifications led to a further reduction in the percentage of low copy number cDNAs. However, direct association between DISCO and DIP2 in-vitro strengthened the argument that DIP2 is a true interactor of DISCO.

The third DISCO-interacting protein (DIP3) isolated in the interaction trap assay is a putative ATPase, the fourth subunit of the *Drosophila* 26S proteasomal complex (Hoyle and Fisher, 1996). The *dip*3 cDNA was also represented once among the 18 interacting clones.

The *dip*1 gene expresses three different size transcripts with repeated sequences at the 3' untranslated region:

A nearly full-length *dip*1 cDNA was isolated from a *Drosophila* cDNA library made in λgt11 vector. This *dip*1 cDNA clone contains the full coding region for the putative DIP1 protein. The size of the protein, predicted from the
open reading frame of the cDNA, is approximately 36.6 kDa. The yeast interaction trap screening led to the isolation of only a 1.1 kb partial cDNA clone that lacked the 5’ end of the corresponding mRNA. Both cDNAs contain an unusual 124 bp repeating element in the 3’ untranslated region (UTR). Sequence comparison of the two cDNAs isolated from two different libraries showed a difference in the 3’ UTR (Figure 12). The larger cDNA contains 8-10 repeats, while the shorter cDNA has only 4 repeats. The larger cDNA also contains an additional 348 bp sequence at the 3’ end. The difference in the 3’ UTR indicates that these cDNAs represent alternatively spliced transcripts for dip1. The possibility of alternative splicing of nascent dip1 mRNA is supported by the results of the Northern analysis that showed expression of at least three different transcripts during embryogenesis (Figure 15). We were unable to determine the exact number of repeats and as a result the exact size of the dip1 cDNA isolated from the λgt11 library. From the size estimated from agarose gel electrophoresis (~2.6 kb), the dip1 cDNA appears to have 8 repeats and may represent the largest transcript (~3.0 kb) seen on the northern blot (Figure 15). It has not yet been determined if the above cDNA lacks a portion from the 5’ end of the corresponding transcript.

The presence of 124 bp repeating elements of almost identical sequence in the 3’ UTR is a unique feature of the dip1 cDNAs. Results of our Northern analysis suggest the possibility that other RNA species expressed in Drosophila embryos may also contain the same repeat. This possibility is strengthened from the results of in situ chromosomal localization studies that showed hybridization of a dip1 probe which included the repeated region to an additional location on the polytene chromosome. These observations suggest the presence of at least
one other gene in the *Drosophila* genome that contains the repeating element present in *dip1* cDNAs.

The striking identity in the nucleotide sequence of the individual repeats suggests an important regulatory role in gene regulation. The 3' UTRs in several mRNAs have been shown to be involved in translational regulation, subcellular localization and stability (Gavis *et al.*, 1996; Singer, 1992; Sachs, 1993; Rastinejad and Blau, 1993). These processes are in turn important for regulation of gene expression during cell growth and development.

Repetitive sequences in the 3'UTR have been shown to mediate translational regulation of mRNAs. An example has been demonstrated for the gene 15-*lipoxygenase* (*loxA*; Ostareck-Lederer *et al.*, 1994). The *loxA* mRNA codes for an enzyme that degrades phospholipids during breakdown of internal membranes of mitochondria during reticulocyte maturation. The 15-*loxA* mRNA is transcribed in bone marrow during early stages of erythropoiesis, but is translated only when reticulocytes reach the peripheral blood. It was demonstrated that a 19 nucleotide sequence repeated 10 times in the 3'UTR of *loxA* mediates the translational repression of the mRNA during early stages of erythropoiesis (Ostareck-Lederer *et al.*, 1994). The fusion of *loxA* repeated elements to the 3' UTR of a reporter gene was sufficient to cause translational repression of the reporter genes. Whether the varying number of the 124 bp repeats present in different *dip1* transcripts are involved in differential regulation of translation of the corresponding mRNAs and whether translational regulation confer spatial restriction to *dip1* gene function are important questions to be addressed in future.
Analysis of secondary structure of the repeated region of the dip1 cDNAs using 'Mfold' software (Walter et al., 1994) predicts formation of stem-loop structures in the 3' UTR of the corresponding mRNAs. The analysis showed that the number of stem-loops that form depends on the number of repeats present in the cDNA. Approximately two repeats are required for each stem-loop. For example, analysis using four repeats led to the formation of 2 stem loops, while eight of them formed 4 stem-loops (results of the analysis are shown in the appendix). This secondary structure analysis thus predicts the formation of double-stranded RNA regions in the 3' UTR of dip1 mRNAs. Double-stranded RNA regions formed by repeated elements may provide a mechanism for translational regulation of mRNAs by dsRNA-binding proteins.

The dip1 gene codes for a putative double stranded RNA-binding protein:

Sequence analysis of the dip1 cDNA that appear to contain the full coding sequence revealed the presence of two type B double stranded RNA-binding domains (dsRBDs) in the predicted DIP1 protein. The type B dsRBDs in general match well with the C-terminal short domain consensus (23 amino acids), but fit the N-terminal long domain consensus (47 amino acids) only poorly (St. Johnston et al., 1992; Figure 13).

Examples of type B proteins that are known to bind dsRNA substrates include the PAC1 protein of fission yeast S. pombe (Iino et al., 1991). PAC1 shares similarity with RNaseIII and is known to degrade dsRNA substrates when expressed in E. coli. Several other dsRNA-binding proteins are known to contain both type A and type B dsRBDs. These include, the Xenopus protein Xlrbpa, its human homolog TRBP (TAR binding protein), and the STAUFEN protein of

The dsRBDs of dip1 are similar to those of mammalian glutamate receptor editases:

The dsRNA binding domains of DIP1 share significant similarity with those of human and rat glutamate receptor editases, hRED1 and rRED1 respectively (Lai et al., 1997; Melcher et al., 1996), which extends beyond the conserved residues of the dsRBD consensus. Figure 14 shows the alignment of the dsRBDs of DIP1 with that of the glutamate receptor editases. On average, the amino acids sequences of DIP1 and the mammalian glutamate receptor editases share 36% identity and 72% similarity within their dsRNA binding motifs. This similarity raises the question of whether these proteins recognize and bind similar RNA sequences.

Although the predicted DIP1 protein shares extensive similarity with the 300 amino acid N-terminal region of hRED1/rRED1, it lacks the C-terminal catalytic deaminase domain of the glutamate receptor editases. Editing (deamination) adenine residues of glutamate receptor mRNAs by these editases causes an adenine to inosine conversion. In the case of the glutamate receptor-B subtype, such a site-specific A to I conversion results in the substitution of an arginine for a glutamine in the protein (Lomeli et al., 1994; Melcher, et al., 1996;
Lai et al., 1997). This in turn alters the calcium conductance properties of the glutamate receptor gated ion channel (Lomeli, et al., 1994).

Although the predicted DIP1 protein lacks an editase domain, we can not entirely exclude the possibility of its involvement in an editing event. At present the genomic sequence for dip1 is not known. In addition, the full coding sequence for only one alternatively spliced dip1 transcripts is available. Therefore, one may consider the possibility that one of the two uncharacterized dip1 transcripts contain a potential editase domain.

Another alternative view regarding the function of the predicted DIP1 protein is the possibility of it being a component of a multiprotein editase complex, where the editase activity is provided by a separate protein. One such example is seen in the case of apolipoprotein B mRNA editing (Navaratnam et al., 1993; Mehta and Driscoll, 1998). Editing of the apolipoprotein B mRNA leads to the production of two forms of the protein, apo-B100 and apo-B48. Apo-B100 and apoB-48 perform distinct functions in lipoprotein structure and metabolism. Apo-B48 is translated from an edited RNA in which cytidine (position 6666) is deaminated to uracil. This results in a premature termination of translation. Interestingly, the editing is done by a multiprotein complex, called the editase complex. In this complex, the RNA-binding domain that binds the apolipoprotein B mRNA and the editase catalytic domain are located on two different proteins.

In addition, to the dsRBDS, the DIP1 sequence also shares similarity with a bipartite nuclear localization signal present in these editases (Figure 14). Nuclear targeting of the DIP1 protein would be necessary for it to function as in-vivo interactor of the transcription factor DISCO. Preliminary studies in the lab with
antibody raised against the DIP1 protein indeed shows its nuclear localization (data not shown).

**The dip1 gene maps close to or represents flamenco:**

Southern analysis of 20A deficiencies showed that the genetic location of the dip1 gene is very close or may be identical to the flamenco (flam) locus (Prud’homme et al., 1995). flam is known to control the expression of gypsy retrovirus elements of *Drosophila melanogaster*. These viruses exist in the *Drosophila* genome as endogenous provirus. Establishment of the endogenous provirus arises from infection of germ cells by the viral particles and subsequent insertion of DNA corresponding to the RNA genome of these viruses in the *Drosophila* genome. In homozygous flam mutants, insertion rate of the retroviral elements in the germ line is increased several fold. Although, in the absence of the flam gene function, increased gypsy mobilization is seen in the germ line, the site of flam gene function is not the germ-line cells, but, the follicle cells of somatic origin that surround the germ-line cells (nurse cells and oocyte) (Song et al., 1997). This was shown by demonstrating that the expression of the retroviral genome encoded proteins occur only in the follicle cells of the ovary. The complete virion particles expressed in the follicle cells of the flam mutant ovaries then infect the underlying oocyte (Pelisson et al., 1994). Expression of the dip1 mRNA in the follicle cells, the site of flamenco gene function, is consistent with the possibility that the dip1 gene represents flamenco.

The production of complete infectious gypsy viral particles require expression of three transcripts, gag, pol and env (Marlor et al., 1986; Pelisson et al., 1994)). Gag and Pol proteins are expressed from the first and second open
reading frames of full-length gypsy mRNA. Env is expressed from an alternatively spliced transcript produced from the same mRNA. Interestingly, it was noted that the wild-type flam gene function inhibits expression of the alternatively spliced transcript (env) of the gypsy retroviral mRNA (Pelisson et al., 1994). Since a functional flam gene prevents alternative splicing of the gypsy mRNA, complete virions are produced only in the absence of flam gene function. These observations suggest that the mechanism by which the wild-type flam gene function prevents expression of env is by blocking a splice acceptor site in the full-length gypsy mRNA. Since dip1 codes for a putative RNA binding protein, a role for this gene in alternative splicing events may not be surprising.

A previous study has demonstrated intimate associations between transcription and splicing events in the nucleus (Jimenez-Garcia and Spector, 1993, reviewed by Neugebauer and Roth, 1997). Furthermore, several examples are known that show direct in vivo associations between transcription complexes and splicing factors. These include interaction of the transcription factor Spi-1/PU.1 with the splicing factor TSL (Translocated in Liposarcoma) and interaction of RNA polymerase II with SRrp129/CASP11, a member of the SR (serine-, arginine-rich domain) family of splicing factors, (Hallier et al., 1998; Tanner et al., 1997). The TSL protein promotes the use of a 5′ splice acceptor site during alternative splicing of the adenovirus E1A nascent mRNA in a IW1-32 erythroid cell line cotransfected with tsl and E1A genes (Hallier et al., 1998). The transcription factor Spi-1 appears to interrupt the splicing process by trapping the splicing factor TSL. In addition to its role in alternative splicing, TSL has also been shown to interfere with the transcription regulatory function of Spi-1. These observations suggest the possibility that DIP1 may have a role in splicing
while being a component of the DISCO transcription machinery. The example of TSL and Spi-1 show how interaction between an RNA-binding protein and a transcription factor can regulate more than one biological process.

Although, proteins containing dsRBDs are generally known to bind dsRNA substrates, one exception is the mammalian transcription factor NF90, a component of the heterodimeric nuclear factor of activated T cells (Corthesy, 1994; Kao et al., 1994). The normal physiological function of NF90 is to regulate transcription of the interleukin-2 gene by binding to its promoter (the ARRE element). However, during adenovirus infection, this protein may also be able to bind the highly structured single stranded RNA of these viruses (Liao et al., 1998). It has been proposed that adenoviruses exert their control on cellular function through interaction with various RNA-binding proteins including NF90 (Liao et al., 1998). The dual role of NF90 raises the possibility that the putative dsRBDs of the DIP1 protein may also have a dual function. On the other hand, under normal physiological conditions these domains may act as DNA-binding motifs. While, during retroviral infection of the Drosophila ovary, they may function as RNA binding motifs to activate the cellular defense mechanism that prevents infection of the germ line by the gypsy elements.

The example of NF90 suggests the possibility that the DIP1 protein might exert its effect on alternative splicing by binding to a single stranded region of the nascent gypsy mRNA. At the same time, the high similarity seen in the dsRBDs of DIP1 and the mammalian glutamate receptor editases might suggest similar substrate-binding specificity. The editases bind dsRNA substrates formed through base pairing between intronic and exonic sequences of newly transcribed target mRNAs (Lomeli et al., 1994; Higuchi, et al., 1993). Therefore,
we can not exclude the possibility that DIP1 binds to similar dsRNA substrates in order to mask certain splice acceptor sites.

The dip2 gene codes for a highly conserved protein expected to play an important role in nervous system development:

Sequence analysis of the partial 1.2 kb dip2 cDNA (Figure 6) revealed that it has the potential to code for a highly conserved protein of unknown function. The C. elegans and human homologs of the dip2 gene have already been cloned in the corresponding genome projects. A partial 382 bp 3' end fragment of mouse dip2 cDNA has also been cloned in collaboration with another member of the lab (Figure 25). Alignment of the predicted amino acids sequences of all DIP2 homologs showed that the partial C-terminal mouse sequence shares 82.3% identity and 97.6% similarity in amino acid sequence with that of the Drosophila DIP2 protein (Figure 26). The C-terminal 31 amino acids of the DIP2 proteins from all species are identical. The predicted Drosophila protein contains an extra 53 amino acids segment 124 amino acids upstream from the stop codon, which is not present in any other DIP2 homolog. Whether this extra 53 amino acid sequence in the Drosophila protein confers any unique property to it compared to homologs in other organisms will be a subject for future investigation.

Northern analysis of dip2 transcript revealed expression of a single mRNA approximately 5 kb in length during embryogenesis (Figure 20). The size of the full-length C. elegans DIP2 protein predicted from its genomic sequence is about 1414 amino acids in length. The protein coding sequence therefore corresponds to about 4.2 kb of mRNA. The size of the dip2 transcript on the Northern blot
suggests that the size of the *Drosophila* DIP2 protein may be comparable to the size of its *C. elegans* homolog.

The *dip2* mRNA shows a highly tissue-specific expression pattern during embryogenesis. Extensive expression is seen in the developing nervous system at all stages of embryogenesis. Recently, in collaboration with Dr. M. Rudnicki’s lab (McMaster University) the expression of the mouse *dip2* mRNA has also been determined. Interestingly, expression in mouse is also localized to a subset of cells in the developing nervous system. These results may implicate an important role for *dip2* homologs in nervous system development.

The location of the *dip2* gene on *Drosophila* polytene chromosomes was determined to be in band 61B located at the tip of the left arm of chromosome III (Figure 21). Results of the deficiency mapping agree with the *in-situ* data (Figures 3.22, 3.23). Based on this chromosomal location we identified two genomic P1 clones each of which contains at least a portion of the *dip2* gene. Because of technical difficulties in sequencing DNA from large P1 clones (80-100 kb), so far only a 2.7 kb portion of the *Drosophila dip2* gene has been sequenced. There is a continuous open reading frame in this 2.7 kb sequence with a stop codon at nucleotide position 2509 (Figure 24). The predicted amino acid sequence shares a high degree of similarity with that of other DIP2 homologs along its entire length (836 amino acids). These observations suggest the absence of any intronic sequence in this 2.7 kb portion of the *Drosophila dip2* gene.

Search for protein domains in the ‘Motif’ database (ICR, Kyoto U) failed to detect the presence of any known functional domain in the predicted DIP2 protein. However, a weak similarity to an AMP binding domain consensus (PROSITE database, accession number PDOC00427) was detected. AMP binding
domains are normally found in a variety of enzymes that include insect
luciferase, acetyl-CoA ligase, 4-coumarate-CoA ligase as well as others.
Sequence analysis in the ‘Blocks’ database (Henikoff and Henikoff, 1994)
revealed weak similarity to a 30 amino acid sequence from members of the
adenosine- and AMP-deaminase families. The role this 30 amino acid domain
plays in the function of these enzymes has not yet been elucidated. Whether the
weak similarities with the AMP binding domain normally found in enzymes for
energy-dependent catalysis indicate an enzymatic function for the predicted DIP2
protein is not yet known.

Submission of the C. elegans DIP2 sequence in the ‘SOSUI’ database
(Hirokawa et al., 1998) failed to find evidence for the presence of any
transmembrane domain in the predicted protein. Analysis of the DIP2 sequence
for subcellular localization in the ‘PSORT’ database (Nakai and Kanehisa, 1992;
Horton and Nakai, 1997) predicted the presence of an SV40 large T-antigen type
nuclear localization signal (reviewed by Raikhel, 1995). However, the total basic
amino acid content of the DIP2 protein is less than 20%, a value lower than that
found in most nuclear proteins. Our efforts to express a C-terminal DIP2 protein
fragment as GST- or His- fusion proteins in bacteria and as haemaglutinin-
fusion protein in Drosophila S2 cells in culture failed. However, fusion of the same
DIP2 fragment with a nuclear localization signal allowed expression of the
protein in yeast. This observation might indicate that the dip2 gene codes for a
protein that is stable only inside the nucleus.

To assess the role of the dip2 gene product in the regulation of DISCO
function, it is essential to determine the consequence of disruption of dip2 gene
function. In an effort to identify existing mutations that might disrupt dip2 gene
function, we examined several P-element insertion mutants from the region of the third chromosome where the dip2 gene is located. Using a probe corresponding to the 3′ end of dip2 cDNA, available at that time, we were unable to identify any mutant line that carried a P-element insertion inside the dip2 gene. However, a probe corresponding to the 3′ end of the gene may not be able to detect an insertion at the 5′ end of the gene. Isolation of mutations at the dip2 locus and also their characterization will shed light on the possible role played by this gene during nervous system development.

The dip3 gene codes for the ATPase subunit of the Drosophila 26S proteasome complex:

The third DISCO interacting protein (DIP3) isolated in the interactor hunt is a putative ATPase, the fourth subunit of the Drosophila 26S proteasomal complex (Hoyle and Fisher, 1996. 26S proteasomes are known to regulate a variety of cellular processes through targeted degradation of cytoplasmic and nuclear proteins (reviewed by Goldberg, 1992). It consists of at least two components; a proteolytic core known as the 20S proteasome, and an ATPase complex (Orlowsky, 1990; Goldberg, 1992). 26S proteasomes are responsible for ubiquitin- and ATP-dependent degradation of proteins. The normal functioning of eukaryotic cells depends on these pathways to remove regulatory proteins such as cyclins and signal transduction molecules at appropriate times to ensure normal cell division and development.

Analysis of results from almost 100 hunts has revealed that the proteasomal subunits are often isolated as false positives in interactor hunts using different baits (reviewed in Brent and Finley, 1997). However, the current
literature suggests that functionally relevant interaction between transcription factors and ATPase subunits of 26S proteasomes is not uncommon. For example, the ATPase component of the 26S proteasomal complex has been shown to interact with a variety of transcription factors including Gal4 in yeast and c-FOS in mammalian cells (Swaffield et al., 1992; Wang et al., 1996). Mutation in SUG1, the ATPase subunit of the yeast 26S proteasome suppresses a mutation in the transactivation domain of Gal4 (Swaffield, et al., 1992). To show an in-vivo interaction between DISCO and DIP3, it will be necessary to generate mutations in the dip3 gene. Since dip3 mRNA has a ubiquitous expression pattern (data not shown), tissue specific modification of dip3 gene function will be necessary.

How to interpret yeast interaction trap results:
The most difficult problem encountered by investigators using a yeast interaction trap assay is to recognize the interactions that have physiological relevance as opposed to false positives (reviewed in Brent and Finley, 1997). False positive interactions that satisfy all the selection criteria can be of different types. For example, some false interactions never occur in nature, since, the spatial and temporal expression pattern of the interacting proteins do not overlap. These interactions may still be informative in the sense that they may suggest the existence of similar interactions between related molecules. Some false positives may result with “sticky” proteins that activate the reporters in some other ways, for example by interacting with other components of the transcription machinery. Examples of such proteins include heat shock proteins, ribosomal proteins and proteasome subunits. However, the interactions that are clearly false occur fairly infrequently (reviewed by Brent and Finley, 1997).
Interactions that may occur in nature but are not informative include those between baits and members of ubiquitin-dependent proteolytic pathway. Although it may provide a clue as to the degradation process of the bait, it is difficult to follow up the interaction to show its physiological relevance.

Another disadvantage of the interaction trap screening is its inability to detect certain physiologically relevant interactions (reviewed by Brent and Finley, 1997). This system is unable to detect weak interactions with equilibrium dissociation constants weaker than 10-50 µM. Expression of library proteins that are toxic for yeast cells can also lead to false negatives. Furthermore, both interacting proteins in this system contain bulky N-terminal multidomain moieties fused to their N-terminus. These can block interactions that require the N-termini of the partners. Interactions between two proteins also require proper folding of the proteins in the aqueous environment of the yeast nucleus. This is particularly problematic to those proteins that require hydrophobic environment for proper folding, such as, membrane spanning proteins.

The disadvantages of the interaction trap assay suggest that the interactions seen in yeast should be interpreted carefully and reexamined with properly designed follow up experiments to show physiological relevance. The studies presented in this thesis and proposed in the future direction (Chapter 4) outline some of the experiments that can be done to address the above issue.
Materials and methods:

Constructs:

Bait construct with full length disco coding region:
In this construct the disco coding sequence was cloned in frame with the lexA coding sequence located 5’ to the multiple cloning site of the yeast expression vector pEG202. pEG202 (Gyuris et al., 1993) is a yeast-E. coli shuttle vector that contains a E. coli origin of replication, the ampicillin resistance gene, a yeast origin of replication (2 mm ori) and a yeast selectable marker (HIS3). It also contains the promoter from the yeast alcohol dehydrogenase gene (ADH1), a coding sequence from the bacterial repressor protein LexA, a polylinker and the transcription termination sequence from the yeast ADH1 gene. The disco coding region was PCR amplified with primers containing EcoRI (GAATTC) sites. Pfu polymerase (Stratagene) with proof reading activity was used for PCR to eliminate mutations in the amplified product. Absence of any mutation was confirmed through sequence analysis. The amplified product was digested with EcoRI and cloned at EcoRI site of pEG202. The orientation of the disco fragment was examined by digesting the construct with XhoI and confirmed through sequence analysis. Sequences of the two primers used for PCR amplification are as follows:
AB6530 (sense primer): 5’ CCC ACC ACA GAA TCC ATG GAG CAC 3’.
AB6531 (antisense primer): 5’ T CGC CAT GGA TCA GAA TTC TGG AC 3’.

Bait construct with 5’ disco sequence:
For N-terminal DISCO bait, a 5’ disco cDNA fragment corresponding to 186 N-terminal amino acids was PCR amplified with primers containing EcoRI sites and
cloned at EcoRI site of the vector pEG202. The DISCO sequence was cloned in frame with the LexA coding sequence. Sequences of the two primers used for PCR amplification are as follows:

AB6530 (sense primer): 5’ CCC ACC ACA GAA TCC ATG GAG CAC.
AB7045 (antisense primer): 5’ G CAA GCG GAA TTC GTC CGG C 3’.

**Bait construct with mutated 5’ disco sequence:**
For the mutated DISCO bait, a 5’ disco cDNA fragment corresponding to 186 N-terminal amino acids of DISCO, but, with a mutation that substitutes Cys at position 127 to Ser (Heilig et al., 1991), was PCR amplified from genomic DNA extracted from disco1 mutant flies. The primers that were used to amplify the wild type 5’ disco fragment were also used to amplify the mutated 5’ disco fragment and subcloned into pEG202.

**Bait construct with 3’ disco sequence:**
For the C-terminal DISCO bait, a 3’ disco cDNA fragment corresponding to C-terminal 153 amino acids was PCR amplified with primers carrying EcoRI sites and the restriction fragment was cloned at the EcoRI site of pEG202 in frame with the lexA coding sequence. Sequences of the two primers used for PCR amplification are as follows:

AB7044 (sense primer): 5’ AGC ATC AAA ATG GAA TCC GAC CCG 3’.
AB6531 (antisense primer): 5’ T CGC CAT GGA TCA GAA TTC TGG AC 3’.

**lacZ reporter construct:**
The lacZ reporter construct pSH18-34 was kindly provided to us by Dr. Roger Brent (Harvard Medical School). The construct contains a TATA box, transcription start site and a small part of the Gal1 coding sequence fused to lacZ gene. In this construct, the Gal1 upstream activating sequence was replaced with
four LexA operators (Hanes and Brent, 1989). The reporter construct pJK101 (Brent and Ptashne, 1985) used for the repression assay contains most of the Gal1 upstream activating sequence (UASg), and one LexA operator located between the UASg and the Gal1 TATA box. pSH18-34 and pJK101 are both yeast-E. coli shuttle vectors and contain the yeast selectable marker URA3.

**GST-DISCO fusion constructs:**
The same wild-type and mutated 5' disco fragments that were cloned into the yeast expression vector pEG202 were also used to generate GST-DISCO fusion constructs. These 5' disco fragments were cloned into the EcoRI site of the vector pGEX-1 (Pharmacia). The pGEX-1 vector contains the GST coding sequence upstream of the multiple cloning site.

**In-vitro expression constructs for dip1 and dip2:**
The in-vitro expression vector used for this study was pSPUTK. The pSPUTK is a vector with high expressivity that contains the 5' untranslated region of Xenopus β globin gene and Kozak translation initiation sequence (ANNATGG; Kozak, 1996). It also contains SP6 and T7 promoter sequences. This vector was constructed in Dr. David Andrews's laboratory at McMaster University and was kindly provided to us. Both dip1 and dip2 cDNAs with the haemagglutinin epitope tag were PCR amplified from the yeast expression vector pJG4-5 with primers carrying restriction sites. The sense primer had a SalI site, and the antisense primer had a XbaI site. The amplified products were cloned at SalI and XbaI sites of the pSPUTK expression vector. The primers used for this purpose are as follows:

AB10133 (sense): 5' A GCC TCT TGC TGA GTC GAC ATG CC 3'.
AB10134 (antisense): 5' GC GAA GAA TCC AAA GCT TCT AGA G 3'.
Strains used:

Yeast strain: The yeast strain S. cerevisiae EGY48 used in this study was kindly provided by Dr. Roger Brent. The genotype of the strain is *ura3, his3, trp1, LexAop-leu2*. The LexAop-leu2 reporter gene construct is integrated into the genome of EGY48. In this reporter gene construct, regulatory region of the *leu2* gene was replaced with three high-affinity LexA binding sites (LexA operators).

Bacterial strains:

MC1066: The genotype of the strain is *trp1 leu2 ura3*. It was obtained from Dr. Alok Sil (Pennsylvania State University).

BL21 DE3: The genotype of the strain is F' *ompT hsdSb (r' m') gal dcm* (DE3). It is a protease negative *E. coli* strain. Is was obtained from Dr. Andre Bedard (York University).

Repression Assay (Brent and Ptashne, 1984):

The *lacZ* reporter construct used for repression assay was pJK101. This assay was performed to verify nuclear localization of different baits used in this study and to check their ability to bind LexA operator (LexAop). The *lacZ* gene expression from this construct is driven by Gal1 promoter. There is one LexAop site located in between a TATA box and the Gal1 promoter in the reporter construct. Binding of the LexAop by the bait represses *lacZ* expression by 2-20 fold.

Procedure:

The yeast strain EGY48 was transformed with three sets of plasmids; (i) pJK101 and a bait construct containing the auxotrophic markers URA3 and HIS3 respectively, (ii) pJK101 and the HIS3 control plasmid pRFHM1 (positive control
for repression of lacZ expression), and (iii) pJK101 alone. Double transformants were selected on glucose media lacking Uracil and Histidine, and single transformants were selected on glucose media lacking Uracil.

Four individual colonies from each double transformation were streaked onto galactose/raffinose, X-gal media lacking Uracil and Histidine. Four transformants from single transformation were streaked onto galactose/raffinose, X-gal media lacking Uracil. The blue colour intensities produced by these colonies were compared to each other to determine repression of lacZ expression by baits.

**Yeast cell transformation:**

Except for library transformation, all other yeast cell transformations were done using the protocol described below (Geitz _et al._, 1992):

Yeast cell cultures were grown in 50ml volume overnight at 30°C in appropriate medium. Overnight cultures were pelleted by centrifugation at 2000g for 5 minutes at 4°C. Cells were washed twice in TE buffer (10 mM Tris.Cl pH 7.5, 1 mM EDTA), resuspended in 2 ml of LA buffer (100 mM LiOAc, 0.5 X TE) and incubated at 30°C for 1 hour with gentle shaking. To 0.2ml of the cells in LA buffer 0.5 mg of transforming DNA was added. To the transformation mixture 0.7ml of PEG solution (100mM LiOAc, 40% PEG-3350, 1 X TE) was added, gently mixed and incubated at 30°C for 1 hour. Cells were then heat shocked at 42°C for 10 minutes, allowed to cool down to room temperature and plated on selective medium. The plates were incubated at 30°C for 3 days to allow generation of yeast colonies of detectable size.
Interaction trap screening:

Library used: A *Drosophila melanogaster* 0-12 hr. embryonic cDNA library constructed in the yeast expression vector pJG4-5 was kindly provided by Dr. Roger Brent. The library was made with unidirectional cDNAs with EcoRI sticky ends at the 5' end and XhoI sticky ends at the 3' end. The library plasmid is a 2mm plasmid with the TRP1 auxotrophic marker and the Gal1 promoter. Downstream of the Gal1 promoter there is an ATG codon, followed by 105 codons, encoding 9 amino acids from the SV40 Large T nuclear localization signal, 87 amino acids activation domain called B42 (derived from *E. coli*), and 9 amino acids haemaglutininn epitope tag.

Amplification of the library: To amplify the *Drosophila* embryonic cDNA library constructed in pJG4-5, the library DNA were transformed into *E. coli* Sure cells using CaCl₂ heat shock method. Transformants were plated on LB plates with 50 μg/ml of ampicillin. 2.0 X 10⁷ independent transformants were scraped from the plates with LB, and plasmid DNA was extracted using Qiagen plasmid extraction kit.

Screening procedure:
The screening procedure is essentially as described in Finley and Brent, 1995. EGY48 yeast cells containing the bait and the lacZ reporter plasmid were grown overnight in glucose ura'his' medium at 30°C in a 5 ml culture volume. It was subcultured starting with 1/1000 th dilution in 80 ml volume. The culture was grown overnight at 30°C to O.D600 =1. The culture was next diluted to O.D600=0.3 in a 200 ml final volume and grown for 3 hours at 30°C to approximately O.D600=0.6. The cells were pelleted at 2500 rpm at room temperature, washed in 125 ml TE (10 mM Tris.Cl pH 7.5, 1 mM EDTA) and
resuspended in 5 ml LA (100 mM LiOAc, 0.5 X TE). To 250μl cells 125 μg of salmon sperm DNA and 5 μg of library DNA were added and mixed gently. To this mixture 1.7 ml of PEG solution (100mM LiOAc, 40% PEG-3350, 1 X TE) was added, and mixed again. The transformation mixture was then incubated at 30°C for 30 minutes without shaking. After 30 minutes 0.22 ml DMSO was added to increase the transformation efficiency and the cells were immediately heat shocked for 10 minutes at 42°C. An aliquot was used to check the transformation efficiency. The remaining transformation mixture was diluted 10 fold with galactose/raffinose media lacking Uracil, Histidine, and Tryptophan.

Expression of library cDNAs from the Gal1 promoter was induced by growing the transformants in galactose medium for 4 hours at 30°C. The culture was then pelleted at 2000g for 4 minutes at room temperature. Cell pellet was resuspended in 1ml of galactose/raffinose media lacking Leucine, Uracil, Histidine and Tryptophan to select for colonies that activate leu2 reporter gene expression. A maximum of 10^6 transformants was plated on one 30mm plate.

Activation of the lacZ reporter gene expression:
The transformants that survived on the galactose / raffinose leu`ura`his`trp` plates, were next plated on galactose / raffinose leu`ura`his`trp` plates supplemented with 80 μg/ml of X-gal. The colonies were tested for their ability to produce blue color on the X-gal plates.

Isolation of plasmids from yeast cells (Hoffman and Winston, 1987):
Yeast cells containing DISCO interacting protein coding cDNAs were grown overnight at 30°C to saturation in selective medium (glucose medium lacking Uracil, Histidine and Tryptophan). 1.5 ml overnight culture was pelleted and
resuspended in 200μl of solution A (2% triton X-100, 1% SDS, 100 mM NaCl, 10 mM Tris pH 8.0, 1mM EDTA). To the resuspended culture, 200 μl of phenol : chloroform : isooamyl alcohol mixture (25:24:1), and 0.3 g glass beads (acid washed, prebaked) were added. The mixture was vortexed for 2 minutes and then centrifuged at maximum speed for 5 minutes in a bench top centrifuge. Aqueous layer (140 μl) was transferred to a new eppendorf tube. To the aqueous layer 420 μl of ethanol and 4 μl of 1 M MgCl₂ were added to precipitate the plasmid DNA. The mixture was incubated on dry ice for 10 minutes to ensure maximum precipitation. The plasmid DNA was collected by centrifugation at maximum speed for 15 minutes at 4°C, washed with 70% ethanol, dried and resuspended in 20 μl of H₂O.

**In-vitro binding experiment:**

*In-vitro* expression of DIP1 and DIP2:
The DIP1 and DIP2 proteins were *in-vitro* transcribed and translated from the vector pSPUTK using the expression system from MBI Fermentas. The *dip1* and *dip2* cDNAs were transcribed from the SP6 promoter using SP6 RNA polymerase following manufacturer’s protocol. The transcription was done for 30 minutes using non-radiolabeled nucleotides at 37°C in a 20μl reaction volume. Only 2μl of the transcription mixture was used to translate the corresponding protein in rabbit reticulocyte lysate in a 20 μl reaction volume. For translation, [³⁵S] methionine was used to label the translated product. 5 μl of the translated product was analyzed on SDS-polyacrylamide gel for *in-vitro* expressed proteins. Expression and purification of GST-DISCO fusion proteins:
Both wild-type and mutated GST-N-terminal DISCO fusion constructs, as well as GST alone without a fusion protein were expressed from the pGEX vector (Pharmacia) in the protease negative E. coli strain BL21 following manufactures protocol. BL21 cells containing the fusion constructs were grown overnight in LB medium with 100 μg/μl ampicillin. Overnight cultures were diluted 100 fold with fresh LB-ampicillin medium to a final culture volume of 30 ml. The culture was grown to O. D_{600} = 0.6. IPTG was added to a final concentration of 0.1 mM to induce expression of the fusion proteins. The induction of the cultures was allowed to proceed for 3 hours at 37°C. The induced cultures were pelleted, washed three times with PBS (sodium phosphate buffer pH 7.4), and resuspended in 10 ml PBS. The resuspended cultures were sonicated 1 ml at a time in polystyrene tubes in a refrigerated cup-horn type sonicator (model XL2020, Heat Systems Inc.). Sonication was done in three cycles of 15 seconds sonication and 30 seconds incubation on ice. The sonicated sample was centrifuged in a microcentrifuge at maximum speed for 15 minutes at 4°C to remove cell debris. The supernatant was carefully transferred to a new tube and Triton was added to a final concentration of 1%. To the supernatant 200 μl of 50% glutathione-agarose beads (Sigma) in PBST (1X phosphate buffer saline with 1% triton) was added. The GST-fusion proteins were allowed to bind to the glutathione-agarose beads for 1 hour at 4°C with gentle shaking. After 1 hour the beads were pelleted by centrifugation for 1 minute. Beads were washed 3 times with 1 ml PBST and one time with 20 mM Hepes buffer pH 7.4.

*In-vitro* binding experiment:

The protocol for *in-vitro* binding is essentially same as that described in Gekakis *et al.*, 1995. Briefly, 50μl of hydrated glutathione-agarose beads bound to GST-
DISCO fusion proteins, or GST alone, were mixed with 200μl of a binding buffer (20mM Hepes pH 7.4, 100mM KCl, 5mM EDTA, 5mM EGTA, 10% glycerol, 5% Bovine serum albumin (BSA), 0.4% NP40, 1mM dithiothreitol) and 10μl of in-vitro expressed DIP1 or DIP2 protein. The mixture was incubated at room temperature for 1 hour, with occasional mixing. After 1 hour, the GST-beads were pelleted by centrifugation, washed twice with 1 ml binding buffer with BSA and twice with 1 ml binding buffer without BSA. Bound proteins were eluted from the beads by boiling with 30 μl of 2 X Laemmli buffer for 5 minutes in a boiling water bath. The eluted samples were electrophoresed on denaturing 12% SDS-polyacrylamide gels. The gels were fixed/destained for 2 hours in 35% methanol and 10% acetic acid to remove unincorporated [35S]methionine. The gels were then dried, exposed to X-ray film overnight and developed after 12 hours.

**In-situ expression studies:**

**Preparation of digoxigenin-labeled probes:**

For in-situ expression studies digoxigenin-labeled antisense mRNAs were used as probes. The antisense mRNAs were synthesized in-vitro from either T3 or T7 promoter of the SK pBluescript vector (depending on the orientation of the cDNA fragments) using respective RNA polymerase.

**The dip1 probe:**

A 900 bp BamHI fragment of the dip1 cDNA without the repeated region was PCR amplified with Pfu polymerase using primers with BamHI sites and cloned into the BamHI site of the pBluescript vector (Stratagene). The plasmid with the dip1 cDNA was linearized with XhoI and digoxigenin labeled antisense RNA
probe was transcribed using dig-labeled UTP and T3 RNA polymerase (Boehringer Mannheim). The primers used to generate the BamHI fragment of the dip1 cDNA were:
AB11916 (sense probe): 5' T CAT CAA GGA GGA TCC GAT C 3'.
AB 11917 (antisense probe): 5' GAA GT GGA TCC GAA CAG AGA 3'.

The dip2 probe:
The digoxigenin labeled antisense dip2 mRNA was synthesized from a 700 bp EcoRI fragment from the 5' end of the dip2 cDNA isolated from the yeast interaction trap vector. The 700 bp EcoRI fragment was cloned at the EcoRI site of the SK' pBluescript vector and the probe was synthesized essentially as described in the previous section.

The disco probe:
The 1.7 kb EcoRI disco fragment used to construct the full-length disco bait was cloned at the EcoRI site of the SK' pBluescript vector for the expression study. The vector containing the disco cDNA was linearized and digoxigenin labeled antisense disco mRNA was synthesized following the procedure described above.

Preparation and fixation of embryos (Kobayashi et al., 19--):
0-22 hours wild-type (Canton-S) Drosophila embryos were collected. Embryos were dechorinated with 50% bleach for five minutes, and then washed with Embryo Wash (7% NaCl, 0.05% Triton X-100). Dechorinated embryos were fixed in a solution containing 5 ml of 40% embryo wash, 4% formaldehyde, 1X fixation buffer mixture, and 5 ml of heptane (5X fixation buffer composition: 800 mM KCl, 200 mM NaCl, 20 mM EGTA, 5 mM spermidine HCl, 2 mM spermine, 150 mM Pipes pH 7.4). Fixation was done for 20 minutes on an orbital shaker. Fixed embryos sank to the interface, that is, the bottom of the top organic phase. The
lower aqueous phase was removed. 5-10 ml of methanol was added and embryos in methanol-heptane mix (1:1) were shaken vigorously for 20-30 seconds to dislodge their vitelline membranes. Devitellinized embryos sank to the bottom of the tube. Embryos were washed several times with methanol and transferred to fresh tube to remove last trace of heptane. Embryos were next rinsed in a 50% ethanol/50% xylene mixture followed by soaking in 100% xylene for 2-3 hours. The xylene soaked embryos were rinsed in a mixture of 50% ethanol/50% xylene, followed by several rinses in 100% ethanol and finally rinsed in a mixture of 50% methanol/50% PBT (0.1 M sodium-phosphate buffer with 0.1% Tween 20). Embryos were transferred to a microcentrifuge tube, post fixed for 10 minutes in 8ml PBT + 2 ml 10% formaldehyde for 10 minutes with constant shaking. Unused fixative was removed from embryos by washing several times with PBT. Embryos were next treated with proteinase-K (50mg/ml of PBT) for precisely 3 minutes with constant mixing. Proteinase-K treatment makes the embryos more permeable to the probe. The proteinase-K reaction was stopped by rinsing 3 times with a solution containing 2 mg glycine/ml of PBT. Embryos were rinsed twice with PBT and post-fixed again for 10 minutes on a shaker. The fixative was removed by rinsing embryos four times in PBT.

**Preparation and fixation of ovaries** (Kobayashi et al., 1997):

Ovaries were dissected from 2-3 days old females in PBS and fixed with 4% paraformaldehyde in PBS + 1% DMSO for 20 minutes. The fixative was removed with one PBS and two PBT washes, 5 minutes for each step. Proteinase K treatment was done next for about 20 minutes with 50 μg/ml of the enzyme. The exact timing for this treatment was determined empirically to maintain
integrity of the tissue and to ensure optimal signal intensity. The proteinase K reaction was stopped by rinsing with 2mg/ml of a glycine solution. Ovaries were refixed and washed as before. To improve signal intensity, ovaries were incubated in 90% methanol + 10% DMSO for 1 hour at -20°C, followed by rehydration in PBT. Three PBT washes were done, 5 minutes for each wash.

Prehybridization and hybridization:
Fixed and proteinase-K treated embryos or ovaries were rinsed once in a mixture of 50% PBT/50% prehybridization solution and once in prehybridization solution (50% formamide, 300 mM NaCl, 10 mM Tris. HCl pH 6.8, 10 mM Na-phosphate buffer pH 6.8, 1X Denhardt’s solution, 5 mM EDTA and 1 mg/ml of yeast t-RNA), 5 minutes each. These tissue were next prehybridized for 1 hour at 53°C in an orbital rotator. Hybridization was done overnight at 53°C without shaking in hybridization solution (50% formamide, 300 mM NaCl, 10 mM Tris.HCl pH 6.8, Na-phosphate buffer, pH 6.8, 5 mM EDTA, 1X Denhardt’s solution, 10% dextran sulfate) containing antisense mRNA probes. For hybridization, a 1/10 th dilution of each probe was denatured at 60°C for 20 minutes. The denatured probe was further diluted to 1/1000 fold in the hybridization solution before adding to the tissue. After overnight hybridization, the embryos/ovaries were washed for 20 minutes each with a series of washing solutions containing a mixture of a post hybridization wash solution (same as prehybridization solution but without t-RNA) and PBT in the ratios 80%/20%, 60%/40%, 40%/60% and 20%/80%, followed by two washes in 100% PBT. The unhybridized probe was removed with RNAse A at a concentration of 20 mg/ml for 25 minutes at 37°C. RNAse A was removed with several PBT washes.
Detection of the in-situ signal:
To detect the hybridized signal, the embryos/ovaries were first incubated for 1 hour with alkaline phosphatase conjugated anti-DIG antibody (1 to 2500 dilutions in PBT, Boehringer Mannheim). Excess antibody was removed with four PBT washes, 20 minutes each. Embryos/ovaries were rinsed once in the detection solution for 20 minutes (100 mM NaCl, 50 mM MgCl₂, 100 mM Tris.HCl pH 9.5, 0.1% tween-20). The in-situ signal was detected with the alkaline phosphatase substrates NBT (4-nitro blue tetrazolium chloride) and BCIP (X-phosphate/5-bromo-4-chloro-3-indoly1-phosphate). The alkaline phosphatase enzyme catalyses reaction between NBT and BCIP leading to the production of a purple color complex which is deposited at the site of hybridization. The substrate mixture was made with 45 μl of NBT and 35 μl of BCIP in 10 ml of detection buffer. The enzymatic detection reaction was done in dark. Signal was detected within 4-5 hours of incubation. The reaction was stopped with several PBT washes and embryos/ovaries were stored in 70% glycerol in 1X PBS or cleared in methyl salicylate after dehydration.

Screening of the λgt11 cDNA library:
Amplification of the λgt11 cDNA library:
Amplification of the library was done essentially as described in Stratagene manual. The titre of the λgt11 library was determined through serial dilution of the phage stock in SM buffer (5.8 g NaCl, 2 g MgSO4. 7H2O, 50 ml 1 M Tris-HCl pH 7.5, 5 ml 2% gelatin). Each dilution was used to infect the host strain E. coli Y1090r'. The infected culture was plated on 7% agarose and the titre was estimated from the number of plaques obtained on each plate. Based on the titre
value, approximately $10^6$ plaques were plated onto twenty large plates (100 cm). Plates were incubated at 37°C for less than 8 hours to prevent overgrowth and merger of individual plaques. Each plate containing approximately 50,000 plaques as covered with 9 ml of SM buffer and stored overnight at 4°C with gentle rocking to allow the phage to diffuse into the buffer. The bacteriophage suspension was recovered from each plate and pooled together. Chloroform was added to this to 5% final concentration and incubated 15 minutes at room temperature to ensure complete lysis of the host cells. Cell debris was removed by centrifugation. The supernatant was stored at 4°C in 0.3% chloroform and/or at -70°C in 7% DMSO.

**Screening of the λgt11 library with a dip1 probe:**
The amplified library was screened twice for full-length dip1 cDNA. About $7.5 \times 10^5$ and $10^6$ plaques were screened in each individual trials. The host cells were infected and plated at a density of 30,000-50,000 plaques per plate. Nitrocellulose replica filters were prepared from these plates essentially as described in Stratagene manual (1991). The replica filters were hybridized to $[^{32}P]$dCTP labeled dip1 probe synthesized using random primed DNA labeling kit (Boehringer Mannheim). The kit uses random hexanucleotides as primers.

Characterization of the recombinant phage DNA was done following large scale phage DNA isolation procedures of as described in Sambrook *et al.*, 1995. The cDNA fragments from the recombinant phage DNA were subcloned into pBluescript (SK, Stratagene). The cDNA inserts were sequenced from the plasmid vector using an automated sequencing apparatus PD1 Biosystems of the Mobix core facility of McMaster University.
Screening of λgt11 library and λEXlox cDNA libraries for dip2 cDNA:
The probe used for the screening was synthesized from the 1.2 kb dip2 cDNA isolated from the Drosophila library used in the yeast interaction trap screening. The λgt11 library was screened twice and a total of about two million plaques were screened. Five positive plaques were isolated, all of which contained approximately a 500 bp fragment corresponding to the 3’ and of the cDNA. Million plaques from the λEXlox library were also screened. DNA from 4 positive plaques were isolated and digested with ApaI and SacI to determine the size of the inserts. All of these plaques had cDNA inserts of size larger than 2 kb. The largest fragment was about 2.5 kb. Restriction mapping showed that all of these fragments correspond to the 3’ end of the dip2 cDNA.

Northern analysis:
Isolation of total RNA:
Northern analysis was done with 0-22 hour Drosophila embryos. Embryos were collected from plates onto a nitex membrane with DEPC treated water, dried on kimwipes, frozen quickly in liquid nitrogen and stored at -80°C. About 1 g of the frozen embryos was homogenized with a sterile plastic rod in 1 ml of SDS-RNA buffer (10 mM Tris pH 7.5, 100 mM NaCl, 1 mM EDTA and 0.5% SDS). To the homogenate an equal volume of phenol/chloroform mixture was added, vortexed well and centrifuged at 3500 rpm for 10 minutes at room temperature. Aqueous phase was transferred to a fresh tube. Phenol/chloroform extraction step was repeated several times until the interface disappeared. The aqueous phase was collected and RNA was precipitated with 1/15 volume of 3 M NaCl
and 2.5 volume of 100% ethanol. Precipitation was done for 40 minutes at -80°C. The precipitate was collected by centrifugation at 6000 rpm for 30 minutes. The pellet was washed with 70% ethanol, dried and resuspended in 500 µl water.

Purification of polyA⁺ RNA:

Poly A⁺ RNA was purified from total cellular RNA using an oligo-dT column. A 1 ml oligo-dT column was prepared in a pasteur pipette with DEPC treated water. The column was washed with one column volume of 0.1 N NaOH followed by 2-3 volumes of water or until the pH of the eluant became 7. The column was next equilibrated with TSN (12.5 mM Tris pH 8.0, 250 mM NaCl and 0.125% SDS).

To the total RNA suspended in water an equal volume of TSN was added. RNA was denatured by heated at 65°C for 2 minutes. The solution was cooled on ice and poured into the oligo-dT column. The eluant was collected and ran through the column 3 more times. The column was rinsed with 5 ml of TSN. The polyA⁺ RNA was eluted with 3 ml of DEPC-treated water. To the eluant 200 ml of 3M NaCl and 7.5 ml of ethanol were added, and incubated for 30 minutes at -80°C to precipitate the polyA⁺ RNA. The precipitate was collected by centrifugation for 45 minutes at 7000 rpm. The pellet was washed in 70% ethanol and resuspended in 100 µl water. The O.D of the isolated polyA⁺ RNA was measured at 260nm and 5 µg of the RNA was loaded in each lane of an agarose/formaldehyde gel. The gel electrophoresis, transferring RNA to nitrocellulose membrane and hybridization with dip1/dip2 probe were done essentially as described in Sambrook (1995).
**In situ hybridization to *Drosophila* polytene chromosome** (Ashburner, 1989):

**Probe used:**

For in situ localization of the *dip1* gene on *Drosophila* polytene chromosome, biotin labeled DNA probes were made using biotinylated-dATP. A 1050 bp. EcoRI fragment of the *dip1* cDNA including the repeated region and a 343 bp. Sau3A fragment without the repeated region were used as templates to make probes for the *in situ* hybridization study. The biotin-labeled probes were synthesized using BRL nick translation kit. The reaction was done in a 50 μl volume containing 5 μl of solution A, 7.5 μl of 0.4 mM biotynylated-dATP, 0.5 mg/ml of the template DNA and 5 ml of solution C. The probe was synthesized for 90 minutes at 15° C. The reaction was stopped with 5 μl of 0.5 M EDTA. To inactivate the enzyme the reaction mixture was incubated in a boiling water bath for 10 minutes. The probes were precipitated with 5 μl of 3M Na-acetate and 150 μl of ice cold 100% ethanol. The ethanol precipitation was done for 30 minutes at -80° C. The precipitates were collected by centrifugation for 15 minutes at 4° C. Pellets were washed with 70% ethanol, dried at room temperature and resuspended in the hybridization solution (2X SSC, 0.8 mg/ml salmon sperm DNA, 10% dextran sulphate and 50% formamide) at a concentration of 20 mg/ml. The probes in hybridization solution were denatured by heating for 10 minutes in a boiling water bath, immediately cooled on ice and stored at -20° C for extended period.

Composition of solution A: 0.2 mM of dCTP, dGTP and dTTP; 500 mM Tris-HCl pH 7.8; 50 mM MgCl2; 100 mM 2-mercaptoethanol and 100 mg/ml nuclease free BSA, bovine serum albumin.
Composition of solution C: 0.4 u/ml DNA polymerase I, 40 pg/ml DNaseI, 50 mM Mg-acetate, 1 mM 2-mercaptoethanol, 0.1 mM PMSF, 50% (v/v) glycerol and 100mg/ml nuclease free BSA.

For in-situ localization of the dip2 gene, a probe was synthesized following the same procedure as described above. The 1.2 kb dip2 cDNA fragment isolated from the yeast interaction trap vector was used as template. Pretreatment of microscope slides with SSC-Denhardt's solution: Microscope slides were incubated for 2.5 hours at 65° C in SSC-Denhardt's solution (20X SSC: 3M NaCl, 0.3 M Na-citrate, 1% polyvinylpyrrolidone, 1% ficoll, 1% nuclease free bovine serum albumin). These were then quickly rinsed in distilled water, and fixed in ethanol-acetic acid fixative (3:1) for 20 minutes at room temperature. After fixation slides were air dried and stored in the refrigerator.

Preparation of chromosomes from salivary glands:
Wild-type (Cantos-S) Drosophila third-instar larvae were raised in a less crowded environment in Mother Parkers medium. Salivary glands of these larvae were dissected with fine forceps in 45% acetic acid, removing as much of the fat bodies, salivary gland membrane and salivary ducts from the glands as possible. The glands were transferred to a fresh drop of 45% acetic acid for approximately two minutes. The glands were then transferred to a drop of "1:2:3" solution on a siliconized coverslip and left there for 4-5 minutes. The coverslip containing the salivary glands was picked up with a slide treated in SSC-Denhardt's solution. Chromosomal spreading was done by first tapping the coverslip gently with the rubber end of a pencil, and then pressing it down with thumb. Slides with nicely spread chromosomes were refrigerated overnight to further flatten the salivary
chromosomes. The coverslips were removed from the slides by first dipping the slides in liquid nitrogen for 15 seconds and then popping the coverslip immediately with a razor blade. The slides were immediately placed in 95% ethanol for 15 minutes and then transferred and kept in fresh 95% ethanol for 1.5 hours. The slides were then air dried and stored in the refrigerator until use.

Prehybridization treatment:
The slides with the polytene chromosomes were treated in 2X SSC at 65° C for 30 minutes to remove basic proteins. These were then treated twice with 70% ethanol (for 5 and 10 minutes respectively) and once with 95% ethanol (for 10 minutes). Following ethanol treatment, slides were air dried in the fume hood. The polytene chromosomes on the slides were denatured in 0.07 N NaOH for precisely 2.5 minute, at room temperature. Rinsing was done sequentially in 3 baths of 2X SSC for 5 minutes each at room temperature. Slides were dehydrated again in 2 baths of 70% ethanol (5 minutes each) followed by 2 baths of 95% ethanol (5 minutes and 10 minutes respectively) and air dried in the fume hood.

Hybridization:
The probe was denatured in a boiling water bath for 5 minutes followed by immediate chilling on ice. 15 µl of the probe was applied to a cover slip, and the coverslip was picked up with the slide at the position where the salivary chromosomes were located. Hybridization was done overnight at 37° C in a moist chamber in order to prevent the preparation from drying out.

Washes and detection:
Following overnight hybridization, the slides were washed sequentially in 2 baths of 2X SSC for 10 minutes each at 37° C, followed by 2 baths of 2X SSC for
10 minutes each at room temperature. Slides were next rinsed in PBS for 5 minutes at room temperature. The detection was done using the ABC kit from Boehringer Mannheim and following the manufacturer’s protocol. Briefly, the biotin in the probe was allowed to bind streptavidin conjugated to horse radish peroxidase (HRP) enzyme overnight. The biotin-streptavidin complex was detected by using 3,3’-diaminobenzidine tetrachloride (DAB), the substrate for the HRP enzyme. HRP catalyzes a reaction between DAB and hydrogen peroxide that generates a brown colored product which is visualized as the in situ signal. The ABC kit provides other components that enhance the overall signal intensity. The chromosomes on the slides with the in situ signal were stained for 1 minute with a 8% solution of Giemsa (in 0.05 M Tris buffer, pH 7.6) to visualize the polytene chromosome banding pattern.

**Deficiency mapping / Southern analysis:**

**Table 5: Genotypes of the 20A deficiency lines:**

<table>
<thead>
<tr>
<th>Deficiency</th>
<th>flybase ID#</th>
<th>Stock#</th>
</tr>
</thead>
<tbody>
<tr>
<td>Df(1)DCB1-35b/FM6/Dp(1;Y)y[+]mal[106]</td>
<td>Fbab0000370</td>
<td>BL977</td>
</tr>
<tr>
<td>Binsn/Df(1)17-137/Dp(1;Y)mal[+]</td>
<td>FBab0000190</td>
<td>UM1800</td>
</tr>
<tr>
<td>Df(1)A209/FM7a</td>
<td>FBab0000317</td>
<td>BL3714</td>
</tr>
<tr>
<td>Df(1)HM430/FM6/Dp(1;Y)y[+]mal[+]</td>
<td>FBab0000440</td>
<td>UM15810</td>
</tr>
<tr>
<td>Binsn/Df(1)16-2-13/Dp(1;Y)mal[+]</td>
<td>FBab0000184</td>
<td>UM1450</td>
</tr>
<tr>
<td>C(1)DX,y¹w¹fl¹ / Df(1)R21/Dp(1;Y)y[+]mal[+]</td>
<td>FBab0000626</td>
<td>UM8100</td>
</tr>
<tr>
<td>Df(1)B12/FM6/Dp(1;Y)y[+]mal[106]</td>
<td>Fbab0000329</td>
<td>UM14350</td>
</tr>
<tr>
<td>Chromosome Line</td>
<td>stock/ID</td>
<td>stock/ID</td>
</tr>
<tr>
<td>-----------------</td>
<td>----------</td>
<td>----------</td>
</tr>
<tr>
<td>Df(1)DCB1-35c/FM7/Dp(1;Y)y[+]mal[126]</td>
<td>FBab0000371</td>
<td>UM14955</td>
</tr>
<tr>
<td>Df(1)R22,y¹/FM7a</td>
<td>FBab0000627</td>
<td>UM17700</td>
</tr>
<tr>
<td>Df(1)LB6/FM7</td>
<td>FBab0000496</td>
<td>UM16590</td>
</tr>
<tr>
<td>Df(1)S54/FM6/Y</td>
<td>FBab0000682</td>
<td>UM18550</td>
</tr>
<tr>
<td>Binsn/ Df(1)17-257/Dp(1;Y)y[+]mal[106]</td>
<td>Fb0000193</td>
<td>UM1955</td>
</tr>
<tr>
<td>Df(1)R38,y¹/FM7</td>
<td>FBrf0036523</td>
<td>UM78160</td>
</tr>
<tr>
<td>Df(1)R44,y¹w¹f¹/C(1)DX,y¹f¹/Dp(1;Y)mal[+]</td>
<td>FBrf0036523</td>
<td>UM8250</td>
</tr>
<tr>
<td>Df(1)GA22/FM6</td>
<td>FBab0000397</td>
<td>UM15120</td>
</tr>
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</table>

Deficiency lines are maintained in heterozygous conditions over balancer chromosomes. Balancer chromosomes represent special chromosomes that carry multiple inversions (ref). Due to the presence of multiple inversions, recombinations between two heterologous chromosomes are prevented in a heterozygous deficiency line. Absence of recombinations stabilizes the genotype of deficiency chromosomes. The X-chromosome balancers present in the deficiency lines used for this study are; FM6, Binsn, FM7 and FM7a. These balancer chromosomes carry dominant visible markers. As a result a fly with a balancer chromosome is easily recognized. Deficiency lines are lethal under homozygous and hemizygous conditions. X-chromosome deficiencies are maintained in females in heterozygous conditions. Since deficient chromosomes are lethal in homozygous or hemizygous conditions, males do not carry X-chromosome deficiencies. However, there were special strains in which part of the X-chromosome is duplicated on the Y-chromosome. In those lines, males are able to carry X-chromosome deficiencies from the region of duplication. Two such deficiency lines used in our study are D(1)R21 and Df(1)R44. One of the X-
chromosomes in both lines is an attached X-chromosome [C(1)DX], in which two X-chromosomes were attached at the centromere. Since in Drosophila, sex type depends on the ratio of X-chromosome to autosomes, in these lines, flies carrying one attached X-chromosome and a Y-chromosome with duplication develop as females, whereas flies with one attached X-chromosome and one deficient X-chromosome die. In these lines, flies that carry a deficient X-chromosome and Y-chromosome with the duplication develop as males. Therefore, in these lines the deficiency chromosome is carried only in males.

For deficiency mapping of the dip1 gene, females heterozygous for a deficiency chromosome and a balancer chromosome were crossed hemizygous males with a wild-type X-chromosome. From the progeny the female flies heterozygous for the deficiency chromosome and the WT chromosome were selected for southern analysis. The females from the parental line that are heterozygous for the deficiency chromosome and the balancer chromosome were also used for southern analysis. From each of the two lines where a portion of the X-chromosome is duplicated on the Y chromosome, male flies with the deficiency chromosome were used to cross to virgin females with wild-type X-chromosomes. All the female progeny from this cross were heterozygous for the deficiency chromosome and WT chromosome. These females were used for southern analysis.

For southern analysis, the chromosomal DNA from flies of desired genotype were isolated using a rapid small-scale isolation procedure. 50 flies from each line were homogenized with a plastic rod in an eppendorf tube in 500 μl solution A (0.1 M Tris. HCl pH 9.0, 0.1 M EDTA and 1% SDS). The homogenized tissue was incubated at 70°C for 30 minutes for lysis of the cells.
To the homogenate 70 μl of 8 M K-acetate was added and kept on ice for 30 minutes to precipitate cellular proteins. The cell debris and the precipitated proteins were removed by centrifugation at high speed for 15 minutes at 4°C. The supernatant was transferred to a fresh tube and the chromosomal DNA was precipitated with 0.5 volume of isopropanol at room temperature. The precipitate was collected by centrifugation at maximum speed for 5 minutes at room temperature. The pellet was washed with 70% ethanol, dried and dissolved in 100 μl of TE.

The isolated DNA was quantitated in a spectrophotometer at OD_{260} nm. 10 μg of each DNA was digested overnight with EcoRI and BamHI restriction enzymes separately. The digested DNA was electrophoresed on 0.7% agarose gels. After electrophoresis, the gels were used for southern analysis. Southern analysis was done essentially as described in Maniatis et al., 1995. Gels were denatured in 1.5 M NaCl and 0.5 M NaOH for 45 minutes, briefly rinsed in distilled water and then neutralized in 1.5 M NaCl and 0.5 M Tris-HCl pH 7.5 for 45 minutes. DNA from the gels was transferred to nitrocellulose membranes using 10X SSC as transfer buffer. DNA was immobilized by baking the membranes at 80°C for 1 hour. Hybridization was done with [³²P]dCTP labeled dip1 probe made from the non-repetitive region of the dip1 cDNA using random primed DNA labeling kit.

Table 6: Genotypes of the 61A-F deficiency lines:

<table>
<thead>
<tr>
<th>deficiency</th>
<th>breakpoint</th>
<th>Flybase ID#</th>
<th>Stock#</th>
</tr>
</thead>
<tbody>
<tr>
<td>Df(3L)emc-E12/TM6B</td>
<td>061A; 061D03</td>
<td>Fbab0002367</td>
<td>BL2577</td>
</tr>
<tr>
<td>Df(3L)Ar11</td>
<td>061C3-C4; 61E</td>
<td>Fbab0022342</td>
<td>BL1479</td>
</tr>
<tr>
<td>Df(3L)Ar12-1</td>
<td>061C1-C4; 61F3</td>
<td>Fbab0022343</td>
<td>BL1478</td>
</tr>
</tbody>
</table>
Chromosomal DNA from the above three heterozygous deficiency lines (shown below) was isolated. Isolated DNA were digested with EcoRI and BamHI in separate reactions and electrophoresed on 7% agarose gel. The gel was blotted for southern hybridization. The [32P]dCTP labeled probe used for this study was essentially same as that used for northern analysis. The [32P]dCTP labeled disco probe used in this study as control probe, was synthesized from a 1.7 kb EcoRI fragment of the disco cDNA.

**Screening of the genomic clones for dip1 and dip2 genes:**
The host for the P1 phage is the *E. coli* strain NS3145. Screening of the *Drosophila* genomic P1 clones were done following the colony hybridization procedure described in Sambrook *et al.*, 1989. The kanamycin resistant genomic P1 clones were first allowed to adhere to a nitrocellulose membrane. The cells were then lysed on the membrane with 0.5 N sodium hydroxide and 1.5 M NaCl, followed by neutralization in Tris-HCl buffer. The liberated DNA was immobilized on the membrane by baking the membrane at 80°C for 1 hour. The prehybridization and hybridization steps were done as described in Sambrook *et al.*, 1989, using radiolabeled *dip1* or *dip2* probes. The P1 clones screened are shown in the result section in tables 2 and 4.
Screening of P-element insertion mutants from the 61 region of the polytene chromosome:

**Table 7:** Genotype of the P-element insertion mutants screened for mutations at the *dip2* locus:

<table>
<thead>
<tr>
<th>Stock #</th>
<th>genotype</th>
<th>site of insertion</th>
</tr>
</thead>
<tbody>
<tr>
<td>P1206</td>
<td>P[ry[+t7.2]=lArB].A55.1M3 ry[506]</td>
<td>061A</td>
</tr>
<tr>
<td>P1235</td>
<td>P[ry[+t7.2]=23.2]T125 ry[506]</td>
<td>061A</td>
</tr>
</tbody>
</table>

Screening procedure: Chromosomal DNA was isolated from P-element insertion lines following the method described previously. These DNA were digested with EcoRI and BamHI in separate digestion reactions. Southern analysis was
done with the digested DNA using standard procedure (Sambrook et al., 1989). The [32P]dCTP labeled probe used for the southern blot was prepared from the 1.2 kb dip2 c-DNA isolated from the yeast interaction trap vector. The probe was synthesized using a random primed DNA labeling kit as described earlier.

Acknowledgement

I thank Dorothy DeSousa, a Ph. D. student of the lab, for her contribution in the in-situ expression studies for the dip1 gene. I also thank Peter Pelka, a Masters student of the lab for his contribution in cloning of the mouse dip2 homolog (part of his 4C9 thesis), in sequencing of the Drosophila dip2 genomic region and in the demonstration of transcription activation ability of the glutamine-rich region of DISCO in yeast. I thank Jhilik De, a former member of the laboratory for her help in screening genomic clones for dip1 and dip2 genes.
References:


Chapter 4: Conclusion and future direction
Conclusion and future direction

The studies presented in this thesis are aimed at elucidating the mechanisms underlying the establishment of appropriate neuronal connections in the nervous system. To achieve this goal the *Drosophila* larval visual system development was used as a model. The present work shows that not only proper development of neuronal connections in the developing larval visual system is essential for visual system functioning, but, establishment of new neuronal connections at late larval stages may also be important for proper visual response.

The first part of my work shows that a contact between two serotonergic neuronal processes and the LON is established at the larval optic center during third instar larval stage that precedes a change in larval photoresponse from negative phototaxis to neutral (Sawin *et al*., 1994; Sawin-McCormack *et al*., 1995). Serotonin plays a role in modulating photosensitivity of eyes in several organisms including *Aplysia*, crayfish and others (Eskin and Maresh, 1982; Arechiga *et al*., 1990; Barlow *et al*., 1977). Thus, it seems likely that in *Drosophila*, establishment of connections between the serotonergic processes and the LON may modulate larval photobehavior during late third instar larval stage. However, to conclusively demonstrate such a role for serotonin in *Drosophila*, it will be necessary to examine if the absence of serotonin at the larval optic center has any effect on the change of larval photoresponse during late larval stage. One of the ways to eliminate serotonin from the larval optic center would be to disrupt function of the gene encoding the enzyme Dopadecarboxylase required for serotonin synthesis (Livingstone and Tempel, 1983). Disruption of serotonin
synthesis specifically in the neurons that project to the larval visual center can be achieved through generation of genetic mosaics for the *dopadecarboxylase* gene. This study will allow an opportunity to study the *in vivo* role for serotonin in visual system functioning.

The second part of my thesis addressed the molecular mechanisms of neuronal connectivity formation in the visual system. Since optic lobe specific autoregulatory function of *disco* is important for proper LON connectivity and pathway formation, the study was focused on identifying proteins that regulate *disco* activity in the visual system or act in the *disco* mediated pathway in the optic lobe. Yeast interaction trap screening led to the identification of two previously unknown proteins (DIP1 and DIP2) and a known protein (the ATPase component of the 26S proteasomal complex) that interacted with DISCO in this assay.

The *dip1* cDNA codes for a protein with two putative double stranded RNA-binding domains (dsRBDs). The data presented in this thesis raises the possibility that the DIP1 protein may be the same as that encoded by the *flamenco* gene.

The *dip2* cDNA codes for a highly conserved protein with homologs existing in organisms as divergent as *C. elegans* and humans. Sequence comparison and mRNA expression studies have shown that the *Drosophila* and the mouse *dip2* homologs not only share extensive sequence homology, but also a similar embryonic expression pattern mainly localized to the CNS. These findings may suggest a vital role for the *dip2* homologs in nervous system development.
To determine the *in vivo* significance of the interaction between DISCO and its interacting proteins (DIP1/DIP2), a number of approaches may be taken. The suggested initial approach will be to generate antibodies against the interacting proteins and determine if indeed their spatial and temporal expression pattern overlap with that of DISCO. The cells coexpressing these genes in the visual system may be the ones in which *disco* expression is necessary to direct the formation of LON pathway. Generating antibodies will also be necessary for coimmunoprecipitation studies from embryonic tissue to detect *in vivo* physical associations between DISCO and DIP1/DIP2. However, these studies are often tricky due to low physiological levels of cellular proteins and experimental conditions are often not appropriate to maintain the physical association.

Other approaches, such as ectopic expression studies have been traditionally used as an indirect way to determine the function of a gene, can also be used to study interactions between two proteins. The *disco* gene activity has been shown to cause a severe nervous system connectivity phenotype when expressed ectopically in the nervous system (Kevin Lee, Ph. D thesis, 1993). If DISCO needs a cellular partner (such as DIP1 or DIP2) to exert its effects at ectopic locations, disrupting the expression of its cellular partner(s) may revert the connectivity phenotype produced by *disco* ectopic expression. Since appropriate mutations are presently unavailable, deficiencies can be used to disrupt *dip1* or *dip2* gene functions in such studies. Preferably, two different deficiencies need to be used in transheterozygous conditions to ensure deletion of as small number of genes as possible, thereby eliminating the effect of deletions of other irrelevant genes.
A variety of approaches may be taken to elucidate the function of interacting proteins and their roles as DISCO partners. However, the physiological role of dip1 and dip2 genes can be best understood through studies of mutants that disrupt the function of these genes. If suitable existing mutants are unavailable, new mutations may be generated at the dip1 and dip2 locus using P-element insertion or chemical mutagenesis. PCR technology has made the P-element mutagenesis easier for genes that have already been cloned. Approximately 50-100 mutant lines can be screened in a single PCR reaction using two primers, one corresponding to the P-element and the other corresponding to the gene to be mutated (Ballinger and Benzer, 1989).

Another effective way to disrupt gene functions is to inject animals with dsRNA corresponding to the gene of interest (Fire et al., 1998). For reasons not clearly understood, dsRNAs are more effective than single stranded antisense RNAs in interrupting endogenous gene functions. Interestingly, only dsRNAs without intronic sequences have the potency to disrupt gene function. The double stranded RNAs cause disappearance of the gene products in the injected organism and in the F1 progeny, but, not in the F2 progeny. These results argue against a nonreversible gene modification. Furthermore, a small amount of the dsRNA can have a profound effect on the gene function, therefore, masking the function of endogenous mRNAs through sense-antisense RNA-hybrid formation is not a likely explanation for dsRNA mediated inactivation of gene functions. A present hypothesis is that the dsRNAs exert their effect by suppressing transcription from the corresponding endogenous genes (Montgomery et al., 1998). The remarkable efficiency of a small amount of dsRNA in silencing gene expression is proposed to be due to either amplification of a gene silencing agent
inside the cells of the injected organism or establishment of a highly catalytic RNA degradation process (Fire et al., 1998; Ngo et al., 1998). The dsRNA-mediated disruption of gene function has been successfully used in several organisms including C. elegans, Trypanosoma brucei and Drosophila (Fire et al., 1998; Ngo et al., 1998; Kennerdell and Carthew, 1998). The two methods described above are examples of a variety of different ways a gene function can be addressed. Once the in vivo functional role of dip1 and dip2 genes become clear, models explaining the mechanisms of interactions between these proteins and DISCO can be hypothesized and examined.

Information presently available (please refer to the discussion section of chapter 3) suggest several alternative models for DIP1 and DISCO interaction. The first hypothesis is that DIP1 acts as a transcription factor. From the example of NF90, the RNA binding domains of DIP1 can be hypothesized to bind DNA rather than dsRNA substrates. This activity of DIP1 may either stimulate or interfere with the transcriptional regulation function of DISCO. If protein expression studies show coexpression of the DIP1 and DISCO proteins in the optic lobe primordium, the tissue where disco autoregulates its own mRNA expression, a negative effect of dip1 on disco function can be ruled out. Previous studies have shown that expression of a nonfunctional DISCO protein is induced in the optic lobe of the disco1 mutant by restoring wild-type disco gene function in this tissue. If DIP1 indeed functions as a transcription factor that promote disco autoregulation, in a disco-dip1 double mutant background, the above induction will be disrupted.
An alternative view is that the DIP1 protein functions as a dsRNA-binding protein with a role in RNA editing. As discussed previously dip1 may act as a component of an editase complex in which the role of DIP1 is to bind the substrate to be edited, while the catalysis is carried out by a different protein with editase activity. Although a putative editase enzyme that interacts with DIP1 in yeast interaction trap assays has recently been identified, knowledge on the dsRNA substrates for DIP1 is essential in order to further explore its role in mRNA editing. A yeast three-hybrid system that has recently been developed (SenGupta et al., 1996) can be used to identify target mRNAs edited by the hypothetical DIP1 editase complex.

The possible identity of dip1 and flamenco genes may suggest a role for dip1 in alternative splicing (Pelisson, et al., 1994). To examine this hypothesis, it will be necessary to identify DISCO targets that coexpress with disco and dip1 in the same tissue. The expectation will be that, in a disco mutant background expression of the target gene will be completely absent, while in a dip1 mutant background, only expression of certain alternatively spliced transcripts will be absent. The quickest way to identify a disco target will be to screen available enhancer trap lines which show reporter gene expression in the optic lobe region. Absence of the reporter gene expression in any such line, in a disco mutant background, may suggest that the gene whose promoter drives the reporter gene expression is a disco target. Since dip1 shares extensive similarity with the glutamate receptor editases, it will be worthwhile to examine if any of the glutamate receptor subtypes act as in vivo DISCO target.
These proposed studies will provide further insight that will help elucidate some of the general rules that govern proper neuronal connectivity formation in the nervous system.

References


APPENDIX

Structural analyses of the repeated region of dip1 cDNAs.
Results of the structural analyses of the repeated region of dip1 cDNAs:

This Figure shows the results of the structural analyses for dip1 cDNAs using ‘MFOLD’ database. Only repeated sequence in the 3’ untranslated region (UTR) of the dip1 cDNAs were used for the structural analyses. These analyses predicted formation of stem-loop structure in the 3’ UTR of the corresponding mRNAs. Approximately two repeats are needed to form one stem-loop. Panel A shows that structural analysis with the four repeats present in the smaller dip1 cDNA leads to the formation of two stem-loops. Panel B shows that eight repeats (the approximate number of repeats present in the larger dip1 cDNA) form four stem-loops.