The *Drosophila* Jun-N-terminal kinase is required for cell morphogenesis but not for DJun-dependent cell fate specification in the eye

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We cloned and characterized the *Drosophila* homolog of mammalian Jun-N-terminal kinases (DJNK). We show that DJNK is encoded by basket (bsk). Like hemipterous (hep), which encodes the *Drosophila* JNK kinase, bsk is required in the embryo for dorsal closure, a process involving coordinate cell shape changes of ectodermal cells. Dorsal closure can also be blocked by dominant negative *Drosophila* cdc42, which has been shown to act upstream of JNK in vertebrates. Therefore it appears that the JNK pathway is conserved and that it is involved in controlling cell morphogenesis in *Drosophila*. Although DJNK efficiently phosphorylates DJun in vitro, bsk function is not required for the specification of cell fate in the developing eye, a process that requires MAP kinase and DJun function.

**[Key Words: ]** Jun-N-terminal kinase; dorsal closure; basket; *Drosophila*; MAPK

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Mitogen-activated protein kinase (MAPK) cascades transduce a variety of signals in eukaryotic cells in response to multiple extracellular stimuli. Depending on cell type, duration of stimulus, and pathway, they mediate a range of responses including proliferation, differentiation, and the regulation of metabolic pathways in differentiated cells (Marshall 1995). In yeast, for example, three different MAPK pathways have been identified, each one regulating different intracellular responses (Herskowitz 1995).

In vertebrate cells, much has been learned from biochemical studies of the extracellular-signal-regulated protein kinases (ERK) pathways and their involvement in growth control and cellular differentiation (for review, see Marshall 1994, 1995; Nishida and Gotoh 1993). In invertebrates, the corresponding MAPK pathway has been unraveled through the use of genetics in *Drosophila* (for review, see Dickson and Hafen 1993; Zipursky and Rubin 1994; Wasserman et al. 1995), *Caenorhabditis elegans* (for review, see Kayne and Sternberg 1995; Kenyon 1995), and yeast (Herskowitz 1995). In *Drosophila*, this MAPK pathway (R/MAPK) is required for many processes during development, such as in the specification of terminal structures of the larva in response to the Torso receptor, in the formation of wing veins in response to the *Drosophila* EGF receptor homolog, and in the differentiation of photoreceptor cells in the compound eye (for review, see Dominguez and Hafen 1996).

By contrast, the role of the Jun-N-terminal kinase (JNK) pathway, another MAPK pathway, is less well understood. In particular, there is a dearth of genetic studies of the JNK pathway in a multicellular organism, although recently a *Drosophila* JNK kinase has been reported (Glise et al. 1995). From studies in vertebrate cell culture systems the JNK pathway has been implicated in the response of cells to stress, growth factors, and Ras activation (Karin and Hunter 1995). By contrast, mammalian JNKs efficiently phosphorylate c-Jun on two serine residues (Ser63 and Ser73) in the amino-terminal domain of the protein. This phosphorylation correlates well with c-Jun activation. In contrast, ERKs seem to phosphorylate c-Jun at another site in the carboxy-terminal domain that correlates with inhibition of c-Jun DNA binding (Karin and Hunter 1995). Ras can activate both pathways, but to different potencies in several cell lines (Karin and Hunter 1995). Recent reports have suggested other interactions between the JNK and ERK pathways. For example, in rat PC-12 cells, the JNK pathway plays a role in apoptosis, whereas the ERK pathway promotes cell survival (Xia et al. 1995). In contrast, in anergic T cells both pathways need to be activated to mediate the response of T cells to antigen presentation (Su et al. 1994; Fields et al. 1996; Li et al. 1996). Because all these studies have been carried out in cultured cells, little is known about the role of the JNK pathway during the development of multicellular organisms and its interac-

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tion with other parallel MAP kinase pathways such as the ERK pathway in vivo.

To characterize the JNK pathway in a genetically amenable multicellular organism, we have cloned and characterized DJNK, the Drosophila homolog of mammalian JNKs. Here we show that DJNK is encoded by the basket (bsk) gene and that mutations in bsk, like mutations in the Drosophila JNK gene hemipterous (hep) (Glisé et al. 1995), disrupt dorsal closure, a process during embryonic development that involves cell shape changes and cytoskeletal reorganization (Young et al. 1993). We present evidence that the JNK and R1/MAPK pathways in Drosophila are largely independent in the embryo and the eye. Our results also suggest that DJNK function is not required for Jun function in the eye.

Results
Drosophila JNK is a protein kinase

From a staged embryonic cDNA library we isolated cDNAs whose deduced amino acid sequence exhibits high similarity to mammalian JNKs. Because of this, we named the encoded protein DJNK (Drosophila Jun-N-terminal kinase). Northern blot analysis probed with the DJNK cDNA revealed a single band of 1.5 kb, the size of which corresponds to the longest cDNA (1507 bp) (data not shown).

The degree of amino acid identity between the Drosophila sequence and its mammalian counterparts is ~70% (Fig. 1A). In the kinase domain, DJNK contains all the conserved residues of protein kinases, and the TPY motif between kinase subdomains VII and VIII that is phosphorylated on the threonine and tyrosine residues to activate JNKs. The amino acid sequence between kinase subdomains IX and X varies between the INKs, and determines the efficiency of binding to c-Jun (Kallunki et al. 1994). In this region, DJNK is most homologous to mammalian JNKs. Here we show that DJNK is encoded by the DJNKK gene and that mutations in DJNKK disrupt dorsal closure, a process during embryonic development before the separation of vertebrates and invertebrates.

Next, we characterized DJNK biochemically. To do so, we performed in vitro kinase assays with Jun and myelin basic protein (MBP) as substrates. Western blot analysis of Drosophila extracts of embryos, larvae, pupae, and adults probed with anti-JNK1 antibody revealed a single band with an apparent molecular weight of 49 kD, the predicted weight of DJNK (Fig. 2A). This antibody specifically recognizes DJNK and does not cross-react with the similarly sized R1/MAP kinase on Western blots (Fig. 2B). To test for DJNK protein kinase activity we immunoprecipitated DJNK and R1/MAP kinase from larval protein extracts and performed in vitro kinase assays on the immunoprecipitates using DJun as a substrate (Fig. 2B). Like their vertebrate homologs, both DJNK and R1/MAP kinase phosphorylated DJun.

Finally, we wanted to test whether DJNK can be induced by stress, as it has been shown for mammalian JNKs. Here we show that DJNK is encoded by the DJNKK gene and that mutations in DJNKK disrupt dorsal closure, a process during embryonic development before the separation of vertebrates and invertebrates.

Figure 1. Molecular characterization of DJNK. [A] Sequence alignment of DJNK with mammalian JNKs. The conserved TPY motif is marked with bold underlines, the kinase subdomains are marked with roman numerals above the sequences, the conserved residues within the kinase domain are underlined in the consensus sequence, and the "specificity-determining region" domain is underlined in the four listed sequences. [B] Molecular map of the 31B region where the DJNKK is located. The DJNK open reading frame encodes 354 amino acids and is split by five introns. It lies proximal to fs(2)A and ME31B, and distal to the receptor tyrosine kinase gene drar. Below the genomic restriction map are diagrammed the extent of the pWX rescue construct and the extent of DJNKK.
DJNK is expressed dynamically during embryonic development

To examine the pattern of DJNK expression during embryogenesis we hybridized whole-mount embryos with an antisense RNA probe synthesized from a DJNK cDNA template. DJNK is present homogeneously, albeit at low levels, in early syncitial stage embryos (Fig. 3A). Later, during gastrulation, high levels of DJNK mRNA are detected in groups of cells that will undergo shape changes during morphogenetic movements: the cephalic furrow, the anterior and posterior transverse folds, and the leading edge of the ventrolateral epidermis (Fig. 3C–F). Beginning at stage 8–9 of embryogenesis expression is also detected in the peripheral and central nervous system (CNS), and this expression is maintained in the CNS and imaginal disks during larval stages (Fig. 3B–F; data not shown). The dynamic and specific expression pattern of DJNK contrasts with the homogeneous expression pattern of other kinases, including Hep and RI. In vertebrates, JNK3 expression is also specific: It is expressed in subsets of pyramidal cells in the CNS (Mohit et al. 1995).

Mutations in basket disrupt DJNK function

The DJNK gene is located in the 31B region on the polytene chromosomes (data not shown). Molecular characterization of the region showed that DJNK is flanked by two transcription units corresponding to the fs(2)4 and the ME31B loci on the distal side, and the receptor tyrosine kinase gene dror (Wilson et al. 1993; A. Fritz unpubl.) on the proximal side (Fig. 1B). Among other mutations known to map to this region is basket (bsk) (Nüsslein-Volhard et al. 1984). Homozygous bsk mutant embryos die at the end of embryogenesis with a dorsal hole in their cuticle (Nüsslein-Volhard et al. 1984; Lindsley and Zimm 1992). Because mutations in hep, which encodes Drosophila JNKK, have a similar embryonic

Figure 2. Biochemical characterization of DJNK. [A] Western blot of protein extracts of the following developmental stages probed with the JNK1 antibody, embryos [lane 1], first instar larvae [lane 2], second instar larvae [lane 3], early third instar larvae [lane 4], late third instar larvae [lane 5], pupae [lane 6], male flies [lane 7], and female flies [lane 8]. For A the molecular weight markers are 52 kD, 35 kD, 32 kD; for D, 84 kD, 62 kD, and 52 kD. (B) Kinase assay of larval lysates with DJun as a substrate. The lysate was split in two: One-half was precipitated with anti-JNK1 antibody [lane 1], and the other half with anti-Rolled antibody [lane 2]. The kinase assay blot was then probed with JNK1 antibody. The upper panel represents the anti-JNK1 antibody Western, showing that anti-JNK1 antibody does not recognize Rolled, and the lower panel the corresponding kinase autoradiogram. The arrowhead marks the position of the DJNK band. (C) Normalized kinase assays with either 5-min UV-treated larvae, or unirradiated controls. Irradiated and unirradiated lysates were split in two, and treated as in B. Normalization is to unirradiated controls, first column of each pair (n = 5). [D] Western blot with JNK1 antibody of embryonic lysates from the following mutants: bsk1/bsk1 [lane 1]; bsk1/ + [lane 2]; bsk2/bsk2 [lane 3]; bsk2/ + [lane 4]; Df(2L)flp147E/Df(2L)flp147E [lane 5]; Df(2L)flp147E/+ [lane 6]. The big arrowhead marks DJNK. As a control, the blot was also developed with anti-hsp83 antibodies [small arrowhead]. (E) Normalized DJNK kinase activity in embryonic lysates of bsk1, bsk2, and Df(2L)flp147E homozygotes normalized to their balancer siblings (first columns of each pair). n = 4 for bsk1 and Df(2L)flp147E, and n = 3 for bsk2. (F) Normalized DJNK kinase activity in male/female wild-type larvae compared with male hep+/sibling wild-type females larvae. n = 5.

Figure 3. Localization of DJNK mRNA in whole mount embryos. In all panels anterior is left, and in A, B, and C dorsal is up. D and E are ventral views, and F is a dorsal view. Arrows in C, E, and F mark the leading edge of the ventrolateral epidermis. (A) Stage 5 embryo; (B,D) stage 7–8; (C,E,F) stage 12. For nomenclature, see Campos-Ortega and Hartenstein (1985).
phenotype, we wanted to test whether bsk encodes DJNK. To this end we performed transgenic rescue experiments in bsk mutant backgrounds with a 5-kb genomic construct (pWX, Fig. 1B) that only includes the DJNK transcript. A single copy of the pWX construct was able to rescue to viable, fertile adults the lethality of heteroallelic combinations of bsk mutants and deficiencies uncovering bsk. Because pWX does not include dror, fs(2)4, or ME31B sequences, we conclude that the bsk gene codes for DJNK.

The bsk alleles and deficiencies can be ordered in an allelic series with increasing strength of the cuticular phenotypes: \( Df(2L)J27 < bsk^1 < bsk^2 < Df(2L)flp147E \) (Fig. 4B–E). \( Df(2L)flp147E \) is a small, 3-kb deletion generated by an imprecise excision event of the P-element insertion in \( fs(2)4 \) (K. Beckingham, pers. comm.). We found that \( Df(2L)flp147E \) breaks proximal to the bsk gene, deleting most of the bsk coding sequence including the kinase domain, and part of the \( fs(2)4 \) transcript. Therefore \( Df(2L)flp147E \) is a null allele for bsk. In addition, we found that \( bsk^1 \) is temperature-sensitive. At 18°C, egg lays show weaker phenotypes (Fig. 4F).

Next we examined the expression and activity of DJNK by Western blot analysis and in vitro kinase assays on extracts of bsk mutants. bsk1 mutant embryos still have wild-type levels of DJNK, whereas both bsk2 and \( Df(2L)flp147E \) show strong reductions in DJNK compared with their respective heterozygous siblings (Fig. 2D). Kinase activity from bsk1 and bsk2 mutant embryos was significantly reduced (60% and 40% of heterozygous siblings, respectively), whereas \( Df(2L)flp147E \) mutant embryos had only 13% kinase activity of heterozygous siblings (Fig. 2E). The residual DJNK activity observed in the \( Df(2L)flp147E \) homozygous embryos could be due to maternal contribution of Bsk product in the egg.

Figure 4. Allelic series of bsk/DJNK mutant phenotypes. Cuticular preparation of embryos of the following genotypes: (A) Wild-type; (B) bsk1/bsk1; (C) bsk2/bsk2; (D) Df(2L)flp147E/\( Df(2L)flp147E \); (E, I) bsk1/\( Df(2L)flp147E \) derived from bsk1/bsk1 germ-line clones; (F) bsk1/bsk1 reared at 18°C; (G) hep1; and (H) UAS-Deb424817+\(^+/+\) B69-Gal4/+ . I was photographed before digestion with Hoyer’s to show the embryonic tissues. In all panels anterior is \( \text{left} \), and dorsal is \( \text{up} \). (I) Ventrolateral view.

basket shows maternal contribution and disrupts initiation of dorsal closure

We examined the effect of removal of the maternal contribution by making germ-line clones using mitotic recombination. Embryos lacking both maternal and zygotic bsk activity show an extreme dorsal open phenotype: Cuticle is formed only in the ventral and ventrolateral parts of the mutant embryo (Fig. 4E and I). This phenotype is stronger than phenotypes observed by removing only the zygotic bsk activity. We conclude that bsk has a maternal contribution. However, bsk1 embryos derived from bsk2 germ-line clones are rescued by zygotic bsk* function to fertile adults. This indicates that the zygotic bsk expression can compensate for the lack of maternal product.

Dorsal closure occurs during mid-embryogenesis and involves cell shape changes but not cell division (Campos-Ortega and Hartenstein 1985). The cells of the leading edge of the ventrolateral epidermis elongate and stretch in the dorsoventral axis (followed by rows of epidermal cells underneath) until the two edges meet in the dorsal midline and close the embryo, effectively internalizing the cells of the amnioserosa. The epidermal cells stretch over the surface of the amnioserosa cells. Apical accumulation of actin and myosin in the leading edge cells initiates the final steps of dorsal closure (Young et al. 1993). The latter process is disrupted in strong zipper mutants. zipper codes for a nonmuscle myosin heavy chain (Young et al. 1993).

We stained bsk mutant embryos with antibodies against Spectrin and Coracle (Pesacreta et al. 1989; Feigon et al. 1994). Coracle is the Drosophila homolog of the vertebrate band 4.1 cytoskeletal protein (Feigon et al. 1994). Both anti Spectrin and Coracle antibodies mark the profiles of the cells. bsk2 embryos show the initial features of dorsal closure: The leading edge cells change shape and elongate, but the process fails to come to completion, because the first rows underneath the leading edge cells either show only a partial change in shape, or fail to change cell shape completely (Fig. 5B). In wild-type embryos, the epidermal cells underneath the leading edge change shape and elongate after the leading edge cells do (Fig. 5A). \( Df(2L)flp147E \) embryos show a more extreme phenotype: The leading edge cells elongate even less than in bsk1 mutants (Fig. 5C). Embryos derived from bsk1 germ-line clones with no paternal contribution show no change in cell shape and dorsal closure never initiates (Fig. 5D). This result shows that bsk is required for the initiation of dorsal closure.

In bsk mutants, the spreading defect of the cells is more pronounced in anterior cells. In the posterior part of the embryo in weak mutants, some stretching epider-
vertebrates (Coso et al. 1995; Hill et al. 1995; Minden et al. 1995). In Drosophila, the homologs have been cloned [Eaton et al. 1995]. We tested whether interfering with Dced42 function would also give a dorsal closure phenotype. We used a transgene that encodes a dominant negative form of Dced42 (UAS-Dced42N17) [Eaton et al. 1995] and expressed it under the control of Gal4 in the Gal4-69B line [Brand and Perrimon 1993]. This line drives Gal4 expression in epithelial cells during embryogenesis [Brand and Perrimon 1993]. Embryos with one copy of UAS-Dced42N17 and 69B showed a dorsal open phenotype (Fig. 4H). This result suggests that Dced42 like DracA acts in dorsal closure regulation. Therefore it is possible that Dced42 and DracA function in the DJNK pathway like their vertebrate homologs.

bsk regulates puckered expression

puckered [puc] mutations disrupt the formation of a sutu between the two sides of the embryo in the dorsal midline. puc is expressed in the leading edge cells during dorsal closure, as revealed by lacZ expression of an enhancer trap insertion in puc [pucE69] [Ring and Martinez-Arias 1993]. In hep1 mutants, puc–lacZ is not expressed in the dorsal rim cells during dorsal closure, suggesting that expression of puc–lacZ is dependent on hep function [Glise et al. 1995]. To test whether puc expression is also dependent on bsk function we stained embryos from a cross of Df(2L)flp147E/+ ; puc–lacZ/+ and bsk1/+ flies. From this cross half of the progeny carry the puc–lacZ transgene and should show staining of the dorsal rim cells if bsk function were not required for puc expression. We found, however, that of 324 embryos at the stage of dorsal closure, only 92 showed detectable puc–lacZ staining. This number deviates significantly from the expected half that was observed in control crosses not involving bsk alleles. In fact, these numbers fit a distribution of one-quarter staining and three-quarters nonstaining embryos (χ2 = 1.99) and suggest that not only bsk1/Df(2L)flp147E embryos but also the embryos that are heterozygous for the bsk null allele [Df(2L)flp147E] lack detectable levels of lacZ staining. These results suggest that puc–lacZ expression in the dorsal rim cells is dependent on hep and on bsk function. It is interesting to note that it has been shown recently that puc encodes a member of the CL100 family of dual specificity MAP kinase phosphatases [A. Gampel, E. Martin-Blanco, and A. Martinez-Arias, pers. comm.]. In vertebrates it has been shown that CL100 is transcriptionally induced by the activation of the MAP kinase pathway and stress [Keyse and Emslie 1992; Sun et al. 1993]. Because puc expression is similarly regulated by the DJNK pathway, it appears that yet another element of this pathway has been conserved.

We have also noted that in older bsk mutant embryos (scored by the bsk cuticular phenotype) heterozygous for puc–lacZ, there is an ectopic expression of puc–lacZ, including cells of the amnioserosa [data not shown]. This indicates that bsk function might be required later to represses puc–lacZ expression. In this regard, it is interesting to note that DJNK activity is increased in extracts.

Figure 5. Cell shape changes during dorsal closure in bsk mutant embryos. (A) Wild-type larvae showing the ventrolateral epithelia during dorsal closure over the amnioserosa cells. (B) bsk1 embryo, where the leading edge cells (arrows in B, C, and D) and some underlying epithelial cells have begun elongation, but the process stopped mid-way because the epithelium is partially detached from the underlying amnioserosa cells [top, at left]. (C) Df(2L)flp147E embryo showing only partial elongation of the leading edge cells. (D) bsk1 germ-line clone where the leading edge cells have not undergone cell elongation [cf. D with B]. A and D were stained with anti coracle antibody, and B, C, and D with anti spectrin antibody.

Dominant negative Dced42 also shows a dorsal open phenotype

The small GTPases Rac and Cdc42 have been implicated together with JNKK and JNK in the JNK pathway in vertebrates [Coso et al. 1995; Hill et al. 1995; Minden et al. 1995]. In Drosophila, the homologs have been cloned [Dced42 and Drac], but no mutations have been isolated [Eaton et al. 1995]. However, expression of a dominant negative form of DracA under the control of the heat shock promoter resulted in dorsal closure phenotypes
of hep1 mutant larvae (Fig. 2F). This result suggests that DJNK activity is not properly regulated in hep1 mutants.

bsk function is not required for cell fate specification in the eye

We have shown that DJNK phosphorylates DJun in vitro. Because DJun has been implicated in the specification of the R7 photoreceptor cells in the developing eye (Bohmann et al. 1994; Treier et al. 1995), we wanted to investigate a possible role of bsk in photoreceptor cell specification. In situ hybridization experiments with a bsk RNA probe showed expression in the eye antennal disk (data not shown). We generated clones of homozygous bsk1 cells in a heterozygous background normally because the clone size is similar to that of control clones. Furthermore, within the bsk1 mutant clones of young flies ommatidia had developed normally. Only occasionally mutant ommatidia with altered number of photoreceptor cells are seen (Fig. 6). The specification of the R7 photoreceptor cell, which is most sensitive to the expression of a dominant negative DJun protein (Treier et al. 1995), is not affected in bsk mutant clones. Because bsk2 is not a complete loss-of-function allele of bsk, it is possible that in the bsk1 cells there is still sufficient JNK activity for normal Jun activation. We therefore tested also clones of Df(2L)flp147E. Cells homozygous for Df(2L)flp147E lack not only bsk function but also fs(2)4 function, which affects cell size (J. Riesgo-Escovar and E. Hafen, in prep.). Despite their reduced size, homozygous Df(2L)flp147E cells were able to form normal ommatidial units with eight photoreceptor cells (data not shown). These results indicate that in absence of bsk function specification of photoreceptor cells is normal.

Discussion

We have isolated and characterized a novel Jun kinase in Drosophila. Sequence analysis reveals high homology to its vertebrate counterparts, especially to JNK2. As is the case for its mammalian homologs, DJun is a good substrate for DJNK in vitro. We show that DJNK is encoded by the bsk gene. The embryonic lethality associated with bsk loss-of-function alleles is fully rescued by a genomic rescue fragment encompassing only the DJNK transcription unit. DJNK protein levels and kinase activity are severely reduced in embryos homozygous for either the strong bsk2 allele or the Df(2L)flp147E. Mutations in bsk and in hep (Glise et al. 1995) affect the process of dorsal closure, suggesting that both of these kinases are required together in this process.

The role of Bsk in dorsal closure

A large number of embryonic lethal mutations disrupt the process of dorsal closure during embryogenesis and display a common “dorsal open” phenotype (Jürgens et al. 1984; Nüsslein-Volhard et al. 1984; Wieschaus et al. 1984). Cloned genes in this group can be grouped into two main classes: genes involved in the regulation of dorsal closure, and genes coding for structural proteins required for dorsal closure.

The first class is represented by members of the JNK pathway: bsk and hep (Glise et al. 1995) and possibly DracA (Harden et al. 1995) and Dced42. Cleared cuticles of mutant embryos in this class show only dorsal closure phenotypes, whereas the body plan seems unaffected. Mutations in decapentaplegic (dpp), which encodes a TGF-β homolog, or in the genes coding for its receptors, thick veins and punt, or schnurri, which encodes a transcription factor acting downstream of the Dpp receptors, also show dorsal open phenotypes (Jürgens et al. 1984; Nüsslein-Volhard et al. 1984; Affolter et al. 1994; Rüperte et al. 1995). It is unclear at present how the Dpp and the JNK pathway cooperate in this process.

The second class of mutants identifies genes coding for structural elements, mostly associated with the cytoskeleton, that are needed in dorsal closure. This group includes zipper, which encodes a nonmuscle myosin (Young et al. 1993); inflated, coding for an α-integrin subunit (Wilcox et al. 1989); lethal(1)myospheroid, coding for a β-integrin subunit (Wieschaus et al. 1984); and coracle, which encodes a band 4.1 homolog (Fehon et al. 1994).

When is Bsk required during dorsal closure? We observe a wide range of dorsal open phenotypes depending on the strength of the mutant allele and the presence or absence of maternally derived Bsk. The strongest phenotype is observed when both maternal and zygotic Bsk are reduced. The resulting embryos fail to initiate the closing process. When only the zygotic component is removed the process initiates normally but is not completed. This argues strongly for a sustained requirement of bsk during dorsal closure, which normally lasts ~2 hr.

Little is known about the exact role that hep and bsk play in the process of dorsal closure. Specifically, it is not clear what the target of JNK phosphorylation is in dorsal

Figure 6. Analysis of bsk function in the developing eye. Tangential section, at the level of the R7 photoreceptor cell, through the eye of a bsk1/ +/+ fly carrying a clone of homozygous bsk1 cells (marked by the absence of pigment granules). Note that most ommatidia in the clone are wild-type, typically only a few have abnormal numbers of photoreceptors (arrowhead).
closure. JNK could directly phosphorylate and modify cytoskeletal components involved in dorsal closure such as zipper, coracle, inflated, and I(1)mysopheroid. Alternatively, JNK could modify the activity of transcription factors that are known to be involved in dorsal closure. Mutations in genes coding for several transcription factors have dorsal open phenotypes, like pannier (Ramain et al. 1993; Heitzler et al. 1996) and serpent (Abel et al. 1993; Frank and Rushlow 1996), two GATA transcription factors, and anterior open (Nüsslein-Volhard et al. 1984), an ETS domain protein (Rogge et al. 1995). The fact that both hep and bsk mutants affect the expression of puc as monitored by the puc-lacZ staining suggests that JNK acts by regulating transcription factors rather than by directly modifying cytoskeletal components involved in the actual process of cell shape change. Genetic and biochemical experiments are needed to identify the putative transcription factors whose activity is modulated by the JNK pathway as well as further target genes whose expression is modified by the pathway.

Different developmental processes are controlled by the JNK pathway and the MAP kinase pathway in Drosophila

With the identification and cloning of bsk and hep (Glise et al. 1995), the JNK pathway is firmly established in Drosophila. Dominant negative versions of DracA (Harden et al. 1995) and Ddc42 give also dorsal closure phenotypes. It is thus possible that they form part of the pathway, as in the vertebrate JNK pathway. A more definitive placement of Drac and Ddc42 in the DJNK pathway requires the availability of loss-of-function mutants of these genes and gain of function mutations in hep and bsk. In the absence of such mutants we have tested whether bsk mutants act as dominant suppressors of the gain of function phenotype obtained by expressing activated Ddc42 (UAS-Ddc4212) in ectodermal cells in the embryo. Preliminary results suggest that removal of one copy of bsk suppresses partially the dominant phenotype of activated Ddc42 (data not shown). These results are consistent with a model in which Ddc42 acts via DJNK during embryogenesis.

The phenotypes of mutations in genes of the JNK pathway are different from those of mutations in the Rl/MAP kinase pathway in both the embryo and the adult. In the embryo, bsk is involved in dorsal closure and rl is required for signaling in the Torsao pathway (Brunner et al. 1994b). In larvae, in contrast to rl, bsk appears not to be required for cell proliferation and for cell fate specification in the eye. Therefore, it appears that during Drosophila development the JNK and MAP kinase pathway are largely independent.

In vertebrate cell culture systems, the main substrate for JNK is the transcription factor Jun. We have shown that DJun can also phosphorylate Djun efficiently in vitro. Whether DJun is also a major in vivo target of DJNK, however, is questionable. Mutations in DJun have not been identified yet, but through the use of dominant negative and constitutively active Jun proteins expressed selectively in the developing eye it has been shown that DJun may function in the specification of neural fate in the R7 equivalence group in response to the Sevenless signaling pathway (Treier et al. 1995). If indeed bsk were the major kinase involved in the activation of DJun, we would expect impairment of photoreceptor cell specification in clones of homozygous bsk cells. However, bsk mutant cells differentiate largely into normal-looking ommatidia. Therefore it appears that DJNK is not involved in photoreceptor cell specification and is not required for Jun activation in the eye. This is consistent with recent biochemical evidence suggesting that DJun phosphorylation is dependent on the activity of the rolled/MAP kinase in Drosophila (Peve-rali et al. 1996).

Materials and methods

Genetics

Df(2L)flp147E was a gift of K. Beckingham (Rice University, Houston, TX). bsk1 and bsk2 were obtained from the Tübingen stock center, hep1, hep75, and puc609 from S. Noselli (Centre National de Recherche Scientifique, Toulouse, France), the Gal4-698 line from K. Basler (University of Zürich, Switzerland), and Df(2L)J2 and Df(2L)J27 from the Indiana Stock Center. The UAS-Ddc42V12 and UAS-Ddc42V17 transgenic flies were from L. Luo (University of California, San Francisco).

Beside Df(2L)flp147E, Df(2L)J2 and Df(2L)J27, with approximately the same breakpoints (31B-32A), had been reported to uncover bsk (Nüsslein-Volhard et al. 1984; Clegg et al. 1993). We performed complementation tests between the deficiencies and the extant bsk alleles in all pairwise combinations. Failure of complementation was observed in all cases except with Df(2L)J27. Viable transheterozygous with the bsk alleles and Df(2L)J27 were recovered. Egg layers derived from these crosses and from the balanced Df(2L)J27 stock, though, gave rise to a fraction of dead embryos with dorsal open phenotypes, indistinguishable from weak to intermediate bsk phenotypes. We conclude that Df(2L)J27 shows partial complementation with bsk behaving as a weak bsk allele. It is possible that Df(2L)J27 breaks distally near bsk, and disrupts regulatory sequences. The bsk2 allele has a stronger dorsal open phenotype than bsk1 but could not be used for the clonal analysis because it contains at least two additional closely linked lethal mutations on the same chromosome.

For rescue experiments we used three different insertions of pWX: one insertion on the second chromosome, and two on the third chromosome. The pWX line with the insertion on the second was recombined onto the Df(2L)flp147E chromosome and then homozygous. For the third chromosomal lines, flies heterozygous for a bsk allele or Df(2L)J2/CyO, pWX + were crossed to Df(2L)flp147E/CyO and the progeny scored for viable transheterozygotes. We rescued the following mutant combinations: Df(2L)flp147E/bsk1 and Df(2L)flp147E/Df(2L)J2 for one insertion, and Df(2L)flp147E/bsk2 for the other.

From the crosses with pWX insertions, rescued Df(2L)flp147E/ Df(2L)J2, pWX + and pWX, Df(2L)flp147E also uncover fs(2)4 mutations and homozygous females were sterile, as expected. We also used a longer rescue construct (10 kb) in the X chromosome, pWAX. pWAX contains the fs(2)4 transcript and bsk. pWAX is sufficient to rescue both bsk and fs(2)4 alleles (J.R. Riesgo-Escovar and E. Hafen, in prep.).

A chi-square test was used to assess the distribution of stage
8–15 embryos stained for lacZ expression from the cross Df(2L)Fpl147E/+; puc<sup>606</sup>/+ males to bsk<sup>1</sup>/CyO females.

Clonal analysis

Germ-line clones of bsk<sup>1</sup> were generated by use of the "dominant female sterile" technique [Hou et al. 1995] with the FLP-FRT recombinase system [Xu and Rubin 1993]. For bsk<sup>2</sup> and Df(2L)Fpl147E, the same procedure was followed, but eggs were not recovered. Df(2L)Fpl147E also partially deletes a female sterile gene [J.R. Riesgo-Escovar and E. Hafen, in prep.]. bsk<sup>2</sup> has additional mutations on the chromosome [J.R. Riesgo-Escovar and E. Hafen, unpubl.], this could account for the female sterility observed. We also generated bsk<sup>1</sup> and Df(2L)Fpl147E mutant clones marked by the absence of the white gene marker using the FLP-FRT recombinase system of [Xu and Rubin 1993].

Germ-line transformations

A 10-kb genomic fragment [positions 1.2–11.5 in Fig. 1B] cloned in the pW8 transformation vector [Klemenz et al. 1987] was digested partially with PstI and the appropriate fragment ligated to generate pWX. Transgenic flies were generated according to Basler et al. [1991].

Cuticle preparations

Embryos were dechorionated in 50% chlorox, devitellinized, washed, and mounted in Hoyer’s medium. Cuticle preparations were photographed on a Zeiss Axioptem under bright and dark fields.

Histology

For antibody stainings, embryos were treated as described [Young et al. 1991]. Anti Spectrin antibodies were used at a 1:500 dilution, anti-Coracle at 1:250. Fluorescently labeled secondary antibodies [Jackson, Inc.] were used at a 1:100 dilution. Coracle protein distribution in bsk mutant embryos is normal when both maternal and zygotic bsk function is removed. Mounted embryos were viewed on a Multimicroscope laser confocal microscope [Molecular Dynamics, Inc.]. Eye sections were done according to Brunner et al. [1994b]. Images were assembled using Adobe Photoshop 3.0 software.

In situ hybridizations were performed essentially as described [Tautz and Pfeifle 1989] for embryos and larval brains using an antisense RNA probe encompassing the entire DJNK cDNA. Sense probes were used in parallel as controls.

Molecular techniques

Standard molecular techniques were used [Sambrook et al. 1989]. cDNAs and the corresponding genomic region were isolated, mapped, and sequenced on both strands using Sequenase 2.0 (U.S. Biotechnology, Inc.).

Kinase assays

Anti-JNK1 antibody was from Santa Cruz Biotechnology, Inc. MBP immune complex kinase assays were done essentially as described [Brunner et al. 1994a], except that for DJNK assays, 5 µl of JNK1 antibody was used per sample, and protein A–Sepharose beads, antibody, and lysates were added at the same time. Larval, pupal or embryonic extracts were homogenized in a homogenizer adapted with a plastic Eppendorf tube pestle by six steps in RIPA buffer [phosphate buffered saline (PBS), 1% Nonidet P40, 0.5% sodium deoxycholate, and 0.1% SDS], to which was added every time fresh 10 mg/ml phenyl methylsulfonyl fluoride (PMSF) in propanol [10 µl], aprotonin [30 µl/ml] and 100 mM sodium orthovanadate. For Westerns, we used a 1:1000 dilution of the JNK1 antibody.

DJNK immunoprecipitated activity is concentration-dependent, and saturates between 20 and 30 min of incubation [data not shown]. For UV treatment experiments, larvae were collected in empty glass vials with moist filter discs, and subjected to UV light from a transilluminator for 5 min at a 1-cm distance. The vials were then removed, and larvae allowed to recover [45 min] before processing. Harsher UV treatments [15 min] inhibit DJNK by 50%, and enhance Rolled activity [100%] [n = 3 assays] [data not shown]. For kinase assays in a hep<sup>1</sup> background, we took advantage of the fact that hep<sup>1</sup> hypomorphs are semilethal. We made a stock of hep<sup>1</sup> and an attached X chromosome balancer, such that males are always hep<sup>1</sup>, and females are always wild-type. We separated male and female third instar larvae from the same cultures for kinase assays.

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Note

The DJNK cDNA sequence has been submitted to GenBank under accession no. U73196.

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