The Homothorax homeoprotein activates the nuclear localization of another homeoprotein, Extradenticle, and suppresses eye development in Drosophila

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The Extradenticle (Exd) protein in Drosophila acts as a cofactor to homeotic proteins. Its nuclear localization is regulated. We report the cloning of the Drosophila homothorax (hth) gene, a homolog of the mouse Meis1 proto-oncogene that has a homeobox related to that of exd. Comparison with Meis1 finds two regions of high homology: a novel MH domain and the homeodomain. In imaginal discs, hth expression coincides with nuclear Exd. hth and exd also have virtually identical, mutant clonal phenotypes in adults. These results suggest that hth and exd function in the same pathway. We show that hth acts upstream of exd and is required and sufficient for Exd protein nuclear localization. We also show that hth and exd are both negative regulators of eye development; their mutant clones caused ectopic eye formation. Targeted expression of hth, but not of exd, in the eye disc abolished eye development completely. We suggest that hth acts with exd to delimit the eye field and prevent inappropriate eye development.

[Key Words: homothorax; extradenticle; homeodomain protein; nuclear localization; eye; Drosophila]

Received October 16, 1997; revised version accepted December 2, 1997.

Homeotic genes determine the identity of body segments in the anterior—posterior axis in Drosophila. These genes contain a DNA-binding domain, the homeodomain (HD), and act as transcriptional regulators. It was expected that different HD proteins bind to a different set of DNA targets to exert their functional specificity. However, in vitro binding studies revealed that most Homebox (HOX) proteins bind very similar sequences with similar affinities (for review, see Mann 1995). Thus, it was proposed that the HOX proteins require the cooperation of cofactor proteins to enhance their DNA-binding specificity. The extradenticle (exd) gene and its mammalian homologs, the Pbx genes, were found to encode HD proteins that can act as cofactors to many HOX proteins.

Mutations in exd do not affect the expression of the homeotic genes Ultrabithorax (Ubx) and Antennapedia (Antp) but caused homeotic transformation of the embryonic segments (Peifer and Wieschaus 1990). exd also encodes a HD protein (Rauskolb et al. 1993). The suggestion that Exd forms a heterodimer with the HOX proteins and alters their DNA-binding specificity has been confirmed by in vitro and in vivo studies (for review, see Mann 1995). Similar results were shown for PBX (for review, see Lu et al. 1995; Mann 1995; Phelan et al. 1995). The HOX monomer and the HOX–Exd heterodimer can have different sequence specificity, and Exd binding to different HOX proteins selects different binding sites (Chan and Mann 1996; Chang et al. 1996). It was also proposed that the Exd–HOX interaction has a more general role in turning HOX proteins from repressors to activators (Pinsonneault et al. 1997).

Exd is not uniformly available to interact with the HOX proteins. Thus, the availability of Exd provides one way to modulate the HOX activity. Regulation of Exd can be achieved at three levels. The first is on transcription. Although exd maternal transcript and early zygotic transcript are uniformly distributed in early embryo, zygotic expression of exd is not uniform in late embryo (e.g., expression in the epidermis and CNS is higher in the thoracic segments than in the abdominal segments) (Flegel et al. 1993; Rauskolb et al. 1993). This expression

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pattern is regulated by HOX genes in the Bithorax complex (Rauskolb et al. 1993), probably at the transcription level. The second is probably at the translation level or affects protein stability. The exd mRNA is uniform in imaginal discs (Rauskolb et al. 1995), but Exd protein has a patterned distribution in the discs (Gonzalez-Crespo and Morata 1995, 1996). The third level is at the subcellular localization of Exd. Exd is not located in the nucleus in all cells. The distribution of nuclear Exd (Aspland and White 1997) correlates with the functional requirement for exd, indicating that only the nuclear Exd can exert its function. In the embryonic midgut, Exd nuclear localization in the endoderm is regulated by Decapentaplegic (Dpp) and Wingless (Wg) signaling (Mann and Abu-Shaar 1996). However, Dpp and Wg are not the only signals regulating Exd nuclear localization (Mann and Abu-Shaar 1996; Aspland and White 1997).

exd has other roles in addition to its homeotic cofactor function. exd mutant clones caused ectopic eye formation in the ventral head, ventral to dorsal transformation in the imaginal abdomen, affected bristle patterning in the thorax and wing, and affected the proximal leg development (Gonzalez-Crespo and Morata 1995; Rauskolb et al. 1995), all of which are not associated with the anterior-posterior segmentation function of the HOX genes. In vitro studies showed that Exd alone can bind to specific DNA sequences (Lu and Kamps 1996), suggesting that it can act as a transcription factor by itself in vivo.

We report the cloning and characterization of the homothorax (hth) gene in Drosophila. hth was found to act as a homeotic cofactor in embryonic development, similar to exd (Rieckhof et al. 1997). We found the gene to be a homolog of the mouse Mei1 (myeloid ecotropic viral integration site 1) proto-oncogene, which has a homeobox related to that of exd (Moskov et al. 1995). It is expressed in specific patterns in the embryo and imaginal discs. In this report, we focus on the role of hth in imaginal development. Its expression pattern in the imaginal discs and mutant clonal phenotypes in adults were virtually identical to those of exd, suggesting that hth and exd function in the same pathway. We show that Hth activates the nuclear localization of Exd. This is the first case of a HD protein regulating the subcellular localization of another HD protein. In addition, we present evidence that hth and exd act in the same pathway as a negative regulator of eye development. Their normal function in adult head development may be to delimit the eye field and to prevent inappropriate eye formation.

Results

Cloning of the hth gene

Two PlacW enhancer trap lines (P1 and P2) have been isolated on the basis of an interesting expression pattern of their w^m^ reporter gene, which was suppressed in the anterior two thirds of the developing eye and showed a posteriorly pigmented eye phenotype (Sun et al. 1995). The P[+]cDNA in the two lines have been mapped to 86C1-4. P2 is homozygous lethal because of the P[+]/cDNA insertion. The P1 homozygote has no apparent morphological defect, but imprecise excision of P[+]/cDNA from P1 can result in lethal mutations that are allelic to P2, and also allelic to hth and dorsotonal (dtl) mutations (Jurgenst al. 1984; Salzberg et al. 1994, 1997). Thus, these mutations define the hth locus. Another lethal locus, toothless (tss), has been mapped to the same region and also have P-tagged mutants with the posterior eye pigmentation phenotype (Pradip Sinha, pers. comm.). Complementation tests showed that tss and hth are different loci (data not shown). Mutant hth and tss embryos have similar head defects. Two P mutations (tss^47 and tss^470) were also defective for hth. These results suggested that the two genes are very closely located or overlapping, and may be under similar regulation and have similar developmental functions.

Genomic fragments flanking the P[+]/cDNA insert site in P1 were isolated and used as probes to screen an imaginal disc cDNA library. Two nonoverlapping groups of cDNAs were identified, thus defining two transcription units (Fig. 1A). Group I consists of 8 independent overlapping clones (Fig. 1B). The cDNAs hybridized to four separate genomic fragments (Fig. 1A), suggesting that the gene has at least four exons. The P[+]/cDNA in the hth^1224-4 and hth^1237 mutants inserted 63 and 347 bp, respectively, upstream of the 5' end of the group I cDNA clone 7 (Fig. 1A). Therefore, we suspected that the group I transcription unit corresponds to the hth gene. To check this possibility, we tested whether the cDNA can rescue the lethality associated with hth loss-of-function mutations.

hs–hth transgenic strains were made using the 1.6-kb ORF from clone 7 fused to a hsp70 promoter. In wild-type background, a 2-hr heat shock induction of the hs–hth transgene in the embryonic stage caused significant lethality, the effect in the early stages [0-4, 4-8 hr AEL after egg laying] being the strongest. However, the few flies that survived to adulthood were morphologically normal. Induction of hth overexpression during the third instar larval and early pupal stages caused severe shortening of the legs. The shortening was attributable to the lack or shortening of the tibia and tarsal segments. In hth mutants, hs–hth induction resulted in the rescue to adulthood of ~15% of the mutants. The rescued mutant flies have normal head, notum, and abdomen. Their legs were severely shortened, similar to the effect of hth expression in wild type. Although only the weak hth allele can be rescued to adulthood, its rescue demonstrated unambiguously that the group I transcription unit corresponds to the hth gene.

Group I cDNA consists of a single clone (clone 1), located in the intronic region of the group II transcription unit. The cDNA has 3' poly(A) and an ORF starting from the 5' end of the cDNA and without ATG, apparently missing the 5' end. The ORF encodes 88 amino acids, and has no significant homology in the genome databases. P1-K1-8 is a lethal hth mutant derived from P1 by imprecise excision of P[+]/cDNA. Southern blot and sequencing showed that the region hybridizing to the
The group I cDNA has been deleted (Fig. 1A). Complementation tests showed that \textit{P1–K1-8} is defective for \textit{hth}, but not for \textit{tss}. Therefore, group I cDNA is unlikely to correspond to the \textit{tss} gene. It is possible that there is a small \textit{hth} exon located in the region deleted in \textit{P1–K1-8}. Alternatively, the deleted intron may be important for the regulation of \textit{hth} expression.

\textit{Hth} is a homolog of the mouse \textit{Meis1} HD protein

The \textit{hth} cDNA clones reveal an ORF encoding a polypeptide of 487 amino acids (Fig. 2A). It is highly homologous to the mouse \textit{Meis1a} protein (Moskow et al. 1995) in two large regions (Fig. 2B). The C’ homology region (88\% identity in 83 residues) includes a HD (93\% identity in 60 residues). The HD has three additional residues in helix 1, thus belonging to the TALE (three-amino-acid loop extension) HD superfamily (Burglin 1994). It is homologous to the HD encoded by a number of genes from diverse phylogenetic groups, including human, mouse, chicken, Xenopus, Caenorhabditis \textit{elegans}, Arabidopsis, and yeast (Fig. 2C legend). These define a HD family (the \textit{Meis1} family) that is distinct from three other related TALE families: Knotted, PBX, and \textit{Mat\alpha2} (Fig. 2C). The N’ homology region (87\% identity in 119 residues) defines the MH domain for \textit{Meis1} and \textit{Homothorax}; Fig. 2D). The MH domain is found only in \textit{Hth} and the mouse, human, and \textit{Xenopus Meis1} homologs. A stretch of basic residues (KKNQKKR) at the amino terminus of the HD might serve as the nuclear localization signal (NLS) (Nigg 1997).

Tissue distribution of \textit{Hth} in imaginal discs is similar to that of \textit{Exd}

We have determined the expression pattern of \textit{hth} by in situ hybridization and by anti-\textit{Hth} immunofluorescence staining of imaginal discs. The distribution of \textit{hth} RNA and protein in the wild-type imaginal discs have the same pattern and are also the same as the lacZ expression pattern in the enhancer trap line \textit{hth1422-4}. (Fig. 3). The other \textit{hth} enhancer trap lines have lacZ expression patterns that are subsets of the \textit{hth} pattern (data not shown), suggesting that each is responding to a different subset of regulatory elements. In the wing disc, the expression is present in the regions corresponding to notum, wing hinge, and ventral pleura (Fig. 3B,E,H). In the leg discs, the expression is in the periphery region, corresponding to the proximal segments of the legs (Fig. 3A,D,G). In the antennal disc, the expression is in all but the arista region (Fig. 3C,F,I). In the eye disc, the expression is strong in the anterior region surrounding the eye field, including the regions corresponding to ptilinum, ocellus, and head capsules, and weak in the posterior and lateral margins of the eye disc (Fig. 3C,F,I). Very weak expression is detected in the posterior region composed of mature photoreceptors. \textit{Hth} is also expressed in all cells of the peripodial membrane in the eye disc (not
shown). These patterns are very similar to those of nuclear Exd protein (data not shown; Gonzalez-Crespo and Morata 1995; Aspland and White 1997). Closer examination showed that the distribution of Hth and nuclear Exd generally coincide (Fig. 3J,K,L). Outside of the Hth expression domain, Exd protein is present in low level and in the cytoplasm (Fig. 3K,L).

The mutant clonal phenotypes of hth resemble those of exd

All hth alleles, except 1422-4, are embryonic lethal with defects in segmentation, head involution, CNS, and PNS (Kurant et al. 1998). To examine the role of hth in imaginal development, clones of hth mutant cells were generated by FLPase-induced mitotic recombination (Xu and Rubin 1993). Three hth alleles (P2, P1-Δ15, P1-K6-1) were studied. All of them showed similar results. Clones on ventral head tissue (from antenna to postorbital bristles) resulted in ectopic eye formation (Fig. 4C,D), some of which are at the tip of a tubular outgrowth. However, within the compound eyes (Fig. 4D, arrow) as well as in the dorsal head structures, hth mutant clones do not show morphological phenotypes. When clones crossed the eye border, the shape of the eye could become distorted (not shown). In the eye-antenna discs of late third instar larvae bearing hth mutant clones, ectopic photoreceptor differentiation and local overgrowth could be detected. Consistent with the adult phenotypes, these ectopic photoreceptors were found only in the ventral margin of the eye-antenna disc (Fig. 4E,F). Clones in the dorsal margin of the eye disc did not lead to ectopic photoreceptor development (Fig. 4F, arrowheads).

Mutant clones on the second or third antennal segments resulted in transformation to leg-like structure, with the larger clones giving a clear claw structure, indicative of a distal leg (Fig. 4A,B). Mutant clones on the coxa, femur, or tibia often cause fusion of these leg segments, whereas those on the tarsal segments were morphologically normal. The structures affected by hth mutant clones are limited to the proximal antenna and leg segments, consistent with the expression of hth in the proximal region but not in the distal region of the antenna and leg discs. Clones on the mesonotum and abdomen did not have significant morphological phenotypes.

The adult clonal phenotypes are nearly identical to those of exd (Gonzalez-Crespo and Morata 1995; Rauskolb et al. 1995). Together with the above observation that the Hth and nuclear Exd have nearly identical distribution in the imaginal discs, these results suggest that hth and exd act in the same regulatory pathway.
Hth activated Exd protein accumulation and nuclear localization

We checked the epigenetic relationship of hth and exd by making hth mutant clones and examining whether exd expression was affected. In hth mutant clones in the eye disc, Exd protein was not detected (Fig. 4G–I). Thus, hth acts upstream of exd and is required for Exd protein expression or accumulation.

Ectopic expression experiments were performed to further study the relationship between hth and exd. The ectopic expression was achieved using the glass multimer reporter (GMR)-gal4 to drive UAS-hth or UAS-exd expression. The GMR-gal4 contains the yeast gal4 gene driven by the GMR enhancer, which activates the gal4 gene in the region posterior to the morphogenic furrow (MF) of the eye disc (Hay et al. 1994), a region with no or very low levels of endogenous Exd or Hth expression. Subsequently, the Gal4 protein activates the expression of the transgene fused to the Gal4-responsive UAS (Brand and Perrimon 1993). GMR-gal4-induced expression of either Hth or Exd had no significant effect on the adult eye morphology. This provided an ideal situation to study the relationship of these two genes with the least complication.

When Exd was expressed ectopically, it was located in the cytoplasm rather than in the nucleus, where it is expected to exert its function as a transcriptional regulator (Fig. 5A–F). The cytoplasmic localization of Exd was not attributable to cloning artifact that prevented its nuclear import, as the ectopic Exd was transported in the nucleus when Exd and Hth were both induced (Fig. 5G–L). Thus, Hth is sufficient to promote the nuclear localization of Exd protein.

Interestingly, when only UAS-hth was induced, Exd protein level was also elevated (Fig. 5M–R). The Hth-
induced Exd appeared several rows of cells posterior to the Hth-expressing front, and its level increased toward the posterior side (Fig. 5M–R). The gradual increase is consistent with the temporal order of GMR–gal4 expression; cells near the MF have just begun to express GMR–gal4, whereas more posterior cells have expressed GMR–gal4 for longer time. This is in contrast to the situation when both Hth and Exd were induced directly by GMR–gal4, where both proteins appeared simultaneously (Fig. 5G–L). Thus, the gradual increase of Exd indicated that the ectopically expressed Hth is able to induce the nuclear accumulation of Exd protein.

Ectopic hth expression suppress normal eye development

Mutant hth or exd clones in the ventral head can result in ectopic eye formation. Both hth and exd are expressed in the region of the eye disc that develops into the head cuticle surrounding the compound eye (see Fig. 3C; see fate map in Haynie and Bryant 1986). Their expression seems to define the boundary of the eye field and therefore, one of their normal functions may be to prevent inappropriate or excessive eye development. To test this hypothesis, we expressed hth or exd cDNA ectopically (UAS–hth or UAS–exd) using dpp–gal4 (Staehling-Hampton et al. 1994), which induced expression of the UAS fusion gene at the posterior margin of the eye disc where MF (the front of the posterior-to-anterior progression of retinal development) initiates. As expected, hth expression driven by dpp–gal4 suppressed eye development completely (Fig. 6E). With weaker UAS–hth lines, the eyes were smaller and rough. Not surprisingly, expression of exd driven by the same dpp–gal4 did not have any effect (data not shown), presumably because the Exd protein cannot get into the nucleus in the absence of Hth.

We then checked whether Hth inhibited eye development by inhibiting MF movement. Ectopic clonal hth expression was induced using the flip-out and Gal4/UAS system (Brand and Perrimon 1993; Struhl and Basler 1993; Pignoni and Zipursky 1997). Thirty-one adult flies with clonal phenotype were analyzed. The hth-expressing clones located anterior to the MF disrupted the propagation of MF efficiently (Fig. 6). The disruption could be detected in the adult compound eyes as a scar running in the posterior–anterior direction (Fig. 6A–D).
and also in the eye discs of late third instar larvae (Fig. 6E). In two cases, an eye was split into a dorsal and a ventral part, suggesting that a clone at the center of posterior margin can block the normal initiation of furrow, which may then result in the ectopic initiation of MF from the lateral margins. These results suggest that hth can suppress MF progression and possibly also MF initiation.

Discussion

hth activates Exd nuclear localization

We report the cloning and characterization of the hth gene in Drosophila. The Hth protein has an HD that in Drosophila is most similar to the HD of Exd. We had expected it to have functions similar to Exd (i.e., as a cofactor to HOX proteins), perhaps interacting with a different subset of HD proteins or have different effects on the DNA-binding specificity. It was surprising to find that the hth mutant phenotypes revealed by clonal analysis were almost identical to those of exd mutant clones (Gonzalez-Crespo and Morata 1995; Rauskolb et al. 1995). The results suggested that hth and exd act in the same pathway. Our finding that hth activated Exd protein accumulation and nuclear localization provided a clear explanation of why the two genes have nearly identical mutant phenotypes.

The nuclear localization of Exd was shown to be regulated (Mann and Abu-Shaar 1996; Aspland and White 1997). Here we show that the Exd nuclear localization can be controlled by hth. The expression domain of Hth correlates with the domain of nuclear Exd. Outside of the Hth expression domain, the low level of Exd appears to be present in cytoplasm (Fig. 3; Aspland and White 1997). When Exd is expressed ectopically without Hth,
The situation may be different in the leg discs. Exd is largely cytoplasmic (Kurant et al. 1998). These results indicated that Exd is generally required for Exd nuclear localization. When Exd and Hth are coexpressed, both proteins were located in the nucleus (Fig. 5). These results indicated that at least in the leg disc, Exd is also sufficient for Exd nuclear localization.

Exd has a basic region at the amino terminus of its HD that may serve as the NLS. Cytoplasmic retention of proteins with NLS can occur by modulation of phosphorylation at sites within or near the NLS, or through interaction with an inhibitor protein. Phosphorylation by CAMP-dependent protein kinase (PKA) and casein kinase II (CKII) have been shown to cause nuclear import of several mammalian proteins (for review, see Vandromme et al. 1996). We note that Exd has a potential phosphorylation site for PKA and for CKII, just upstream of its HD.

Because the high level of Exd correlated with Exd being nuclear, it is possible that the only posttranscriptional regulation is on nuclear localization. The effect of hth on Exd protein level may simply be a consequence of activated nuclear localization, which concentrated the cytoplasmic Exd into the nucleus. Alternatively, hth may regulate exd at an additional level: transcription, translation of endogenous exd mRNA, or post-translational events that stabilize the Exd protein. These alternatives remain to be resolved.

**hth and exd suppress eye development**

hth and exd mutant clones in the ventral part of the head caused growth of ectopic eye, suggesting that both genes normally function in suppressing inappropriate eye formation outside of the eye field. When hth was expressed in the developing eye (with the subsequent induction of Exd), the eye development can be abolished completely, confirming the suppressive role of hth in eye development. When exd was expressed similarly, there was no effect, probably because the Exd protein cannot get into the nucleus in the absence of Hth. Our results suggested that exd and hth act in the same regulatory pathway suppressing eye development. Hth may function solely through Exd, or Exd and Hth may act cooperatively in the nucleus.

One possible mode of Hth/Exd action in suppressing eye development is by activating Wg signaling, which suppresses the initiation of the MF in the eye disc (Ma and Moses 1995; Treisman and Rubin 1995) or by suppressing Dpp signaling, which is required for MF initiation (Chanut and Heberlein 1997; Pignoni and Zipsrursky 1997). Hth is present in the periphery of the eye disc,
weaker in the posterior margin, and stronger in the anterior half of the lateral margins. The distribution is consistent with a role in suppressing dpp expression (expressed in the posterior and lateral margins but not in the most anterior of the lateral margins). hth mutant clones located in the ventral region of the head can cause ectopic eye formation, whereas clones in the dorsal region have no effect. This correlates with the stronger expression and inhibitory effect of wg in the dorsal margin (Treisman and Rubin 1995). hth-Expressing clones can block the propagation and possibly the initiation of the MF (Fig. 6). These effects are consistent with either a loss of Dpp function or an enhanced Wg function.

Another possible mode of hth/Exd action is by interacting negatively with a nuclear protein, which is required for eye development. Similar mode of action has been proposed for proboscipedia (pb; Bennassayag et al. 1997). Ubiquitous expression of a mutant Pb protein can suppress eye development. The effect is independent of DNA binding by Pb and was suggested to be attributable to perturbed interaction with other proteins. The many nuclear factors required for eye development (eyeless, dachshund, sine oculis, and eyes absent; see Bonini and Choi 1995) are candidates for this suppressive interaction with Exd or Hth.

Function of Hth

Several pieces of evidence suggest that hth can interact with Hox genes, similar to exd. In the hth mutant embryo, the expression of a number of Hox genes were not affected, but their functions were affected; thus hth can function as a homeotic cofactor (Rieckhof et al. 1997; Kurant et al. 1998). Hth and Dfd1 (a gain-of-function mutation of the Deformed HOX gene) double mutants have synergistic effects on eye development (C.-Y. Pai and Y.H. Sun, unpubl.), suggesting that the two proteins may interact. In retrovirus-induced myeloid leukemia in the BXH-2 mice, Meis1 is often coactivated with either Hoxa7 or Hoxa9, suggesting that Meis1 may cooperate with these two Hox proteins in leukemia formation (Nakamura et al. 1996b). The mammalian link suggests that in Drosophila Hth may interact with Antp and AbdB (the homologs of Hoxa7 and Hoxa9, respectively; Burglin 1994).

Because Hth activates Exd nuclear localization, it may exert the above described functions solely through Exd. In vitro studies have shown that Exd can have DNA-binding activity and can interact with HOX proteins, all in the absence of Hth. Does Hth have its own function other than bringing Exd into the nucleus? Hth has a HD similar to that of Pbx and Exd. Eight of the 13 highly conserved residues in all homedomains are conserved in Hth. Another three residues are identical to those in the Pbx/Exd family (Fig. 2). The high degree of conservation suggests that the HD is functional. Whether Hth can serve as a homeotic cofactor or transcription factor independent of Exd is an interesting question awaiting further analysis.

Hth and Meis proteins define a new HD family and a MH domain

The hth gene is a member of the Meis gene family, including three genes in human, mouse, and Xenopus (Fig. 2; Nakamura et al. 1996a; Chen et al. 1997; Smith et al. 1997; Steelman et al. 1997). Our study of hth is the first demonstration of a physiological function for this highly conserved gene family. Hth has two regions highly homologous to other members of the Meis family. The N’ homology region defines the MH domain, which is found only in the vertebrate Meis family and in Hth. This domain has previously eluded recognition because of the high degree of overall homology in the vertebrate Meis gene family. Its high conservation from vertebrates to Drosophila suggests an important, but as yet unknown, function. The C’ region includes and extends slightly beyond the HD. The HD defines a Meis family in the TALE superfamily of HDs, characterized by three extra amino acids in the first α helix.

Meis1 was isolated as a hot spot of retroviral insertions in myeloid leukemia arising in the BXH-2 mice (Moskow et al. 1995). Activation of Meis1 expression is correlated with leukemia formation (Moskow et al. 1995; Nakamura et al. 1996b). The human MEIS1 is also expressed in a subset of myeloid leukemia cell lines (Smith et al. 1997). The exd homolog PBX1 was similarly identified as a fusion partner with the transcription factor E2A found in human pre-B-cell leukemias (Kamps et al. 1990; Nourse et al. 1990). Our results suggest that the vertebrate Meis1 genes may have a function similar to its Drosophila homolog in regulating Pbx nuclear localization, and that misexpression of Meis1 may cause leukemia by activating the nuclear localization of Pbx proteins.

Materials and methods

Fly stocks

P1 and P2 enhancer trap lines have been described previously (Sun et al. 1995). P1;115, P1;K-1-8, P1;K-6-1 are lethals derived from P1 after induction of P[lacw] mobilization. H321 and J186 are two EMS-induced alleles of dfl (Salzberg et al. 1994). 1422-4 and 1323-7 are two P[lacw] tagged dfl alleles, also with the posterior eye pigmentation phenotype (Salzberg et al. 1997). A P[PBX1] tagged lethal mutation (I1305745) was mapped to 86C by the Berkeley Drosophila Genome Project (Gelbart et al. 1997). Complementation tests showed that all of these lethal mutations are allelic to hth (Jurgens et al. 1984). hth mutants were provided by Richard Mann (Columbia University, New York, NY).

P[GMR-gal4] was from Hui-yun Chang (Don Ready, Purdue University, West Lafayette, IN). The FLPase source y w P[Py; hs-flp]U2; Cx/TM3, Sb Ser e was from Tze-bin Chou (National Taiwan University, Taipei). y w; P[Py; hs-neo; FRT]82B P[w+]; hs-m90E Sb P[Py]; y]TM6B (Xu and Rubin 1993) and dpp-gal4 (Staehling-Hampton et al. 1994) were from the Bloomington Stock Center (IN). P[UAS-exd] was from G. Mo-
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Cloning and molecular analysis of hth

Southern blots showed that there is a single P\[acW\] in the enhancer trap P1 line. A 2.1-kb fragment (pSY46) was cloned by plasmid rescue (Bier et al. 1989) from EcoRI-digested P1 genomic DNA. Southern blots using this fragment as a probe showed a different pattern in P1 from wild type. The fragment was used to screen a fly genomic \( \chi \)Ch4A library (Maniatis et al. 1978), from which a clone (SY46) containing a 13.4-kb insert was isolated (Fig. 1A). The restriction map of SY46 is consistent with the map deduced from genome blot analysis. Genome blot also showed that the \( \chi\)SY46 clone does not contain repetitive elements. It was used to screen an imaginal disc cDNA library provided by N. Brown (Brown and Kafatos 1988). Eleven positive clones were isolated. Restriction mapping and partial sequencing classified the cDNA clones into two nonoverlapping groups. Group I is represented by a single clone (clone 1) of 1068 bp. Its 3' end is -260 bp from the P[acW] insert site. Group II consists of 10 overlapping clones, 8 of which are independent clones. Clones 1, 4, 5, and 7 have been sequenced completely in both strands, the others were checked by partial sequencing, restriction mapping, and PCR. The TNT lysate coupled transcription/translation kit (Promega) was used to check the protein-coding capability of the cDNA clones. Additional genomic clones were isolated using the group II cDNA as a probe.

The group II cDNA clones contain an ORF of 1461 bp. One of the first six in-frame ATGs have flanking sequences that match the consensus for ribosome binding site in Drosophila (C\( _{IA} \) A\( _{IC} \) A\( _{IC} \) A\( _{IC} \) AUG; Dalphin et al. 1996) or in vertebrates (Kozak 1984). The predicted molecular masses of the encoded polypeptides are 52.8, 51.2, 50.5, 47.9, 46.9, and 43.9 kD. In vitro translation of clones 5 and 7 gave a protein band of -55 kD (data not shown), suggesting that the first ATG may be used as the translation start site.

The 5' end sequence of clone 7 (GACAGGTGT) matched well with the consensus of Drosophila transcription initiator (Arkhipova 1995). However, its upstream genomic sequence (data not shown) did not have a TATA box. Sequence at the 5' end of clone 5 did not match with the initiator element but has a consensus TATA box (TTTAAACC, -32 to -24 relative to the 5' end of clone 5) (Arkhipova 1995). It is not clear whether there are two potential transcription initiation sites. The 5' end sequence of the other cDNA clones did not match with the TATA box or initiator and therefore, probably resulted from incomplete cDNA synthesis. The group II cDNA clones revealed four different polyadenylation sites, none affecting the ORF.

By comparing the sequences of cDNAs with parts of the genomic fragments, one exon-intron junction has been identified after the +14 position in the ORF (Fig. 1A,B, arrow). A 45-bp segment was present in clones 3, 4, 7, 8, 9, and 10, but absent in clones 2 and 5 (Fig. 1B). The boundary sequences did not fit the consensus of 5' and 3' splice junctions (GT and AG, respectively; Mount 1982), nor those of the minor intron class (AT and AC; Jackson 1991; Hall and Paddget 1994). We do not know whether it is an alternatively spliced exon. Interestingly, several cDNA clones isolated from an embryonic library were analyzed and all contained 45 bp (data not shown). Clone 4 is unique in that it contained two segments (Fig. 1B, open bar) that do not hybridize with the other clones. The sequence at the junctions (ATTG\( _{GT} \)GA ...ACAG\( _{GT} \)TC ...CAG\( _{GT} \)AAA ...; Fig 1B, arrowheads) matched perfectly with the consensus splice junction sites, suggesting that these are unspliced intron. The first unspliced intron immediately generated a stop codon after residue 415; therefore, the protein encoded by clone 4 cDNA is truncated in the HD. Whether this isoform is just an aberrant splice product or plays a regulatory role is not clear.

A 2.2-kb plasmid can be recovered from the P1-K1-8 mutant by plasmid rescue using EcoRI digest. Sequence and Southern blot analysis showed that a region starting 75 bp 3' from the EcoRI site to the proximal end of the P[acW] (including the HindIII site) has been deleted in P1-K1-8 (Fig. 1A).

UAS-hth and hs-hth transgenic flies

A PCR-amplified fragment containing the complete hth ORF was first inserted into the BamHI site in pBlueScript (SK+) vector (Stratagene), then was recut by EcoRI and XbaI and inserted between EcoRI and XbaI sites of p[\( \mu \)UAST] (Brand and Perrimon 1993) and p[CaSpeR-hs/act] (Thummel and Pirtrotta 1992). They were used to generate transgenic flies using the traditional germ-line transformation procedure (Rubin and Spradling 1982). Multiple independent transformants were obtained for each construct. Two lines with p[CaSpeR-hs-hth] on the X chromosome, p[hth-hth]\( ^{0} \), p[hth-hth]\( ^{X} \), and one line with p[CaSpeR-hs-hth] on the second chromosome, p[hth-hth]\( ^{Y} \), were tested for their ability to rescue the lethality of several hth alleles (P2, 1422-4, P1–\( \Delta 15 \), K1–\( 8 \), P[hth-hth]/hth/TM3, Sb Ser e flies were maintained at 25°C. Eggs from these flies were collected every 12 hr and were heat shocked in 37°C water bath for 15 min every 12 hr throughout development. Rescued flies were detected as non-Sb flies. Rescue to adulthood was detected only in the weak hth\( ^{14222} \) allele. Whether the other alleles can be rescued partially has not been examined.

Clonal analysis and ectopic expression

To generate mosaic flies with hth mutant clones, y w [\( ry^{+}\); hsFLP]; P[\( ry^{+}\); hs-neo; FRT]82B P[w\( ^{\mu}\); hs–m] Sb\( ^{6b} \) P[\( ry^{+}\); y\( ^{1}\) CxD were crossed with y w; P[\( ry^{+}\); hs-neo; FRT]82B hth/TM3, Sb Ser e. Embryos were collected every 12 hr and were incubated for 1 hr in 37°C water bath 24–36 hr AEL. The hth alleles used included P2, P1–\( \Delta 15 \), and P1–K6-1. All gave similar results. A total of 271 mutant clones on the adult head were scored: 63 on the second or third antennal segment, 120 on the ventral head structures including vibrissiae (22), gena (36), postgena (38), and lower postorbital bristles (24), 72 within the compound eye, and 16 on the dorsal head structures including vertex bristles (11), ocellus (2), and upper postorbital bristles (3). To induce random hth expression clones, y w P[\( ry^{+}\); hsFLP]\( ^{P2} \); UAS-hth/TM3, Sb Ser e flies were crossed with P[act–FRT–CD2–FRT–gal4]. Larvae 24–36 hr AEL were incubated for 1 hr in a 37°C water bath. About 300 adults were examined and 31 had clonal phenotype affecting the eye. Because of the lack of marker, the exact extent of the hth-expressing clones cannot be determined in the adult.

Histochernistry

Digoxigenin-labeled RNA probe was used for whole mount in situ hybridization of imaginal discs (Theisen et al. 1996). Rat anti-Exd antibody was from G. Morata (Gonzalez-Crespo and Morata 1995). Rabbit-anti-Hth antisera was prepared from the Escherichia coli expressed Hth protein with an amino-terminal
poly-(His) tag. The antisera detected only one protein species (∼55 kDa) on the immunoblot of Drosophila embryo extract. For anti-Exd and anti-hHth immunofluorescence staining, imaginal discs were dissected from late third instar larval in PBS (137 mM NaCl, 2.68 mM KCl, 8.1 mM Na₂HPO₄, 1.47 mM KH₂PO₄). Discs were incubated in fixation solution (4% paraformaldehyde, 0.2% Triton X-100, PBS) for 20 min. After several washes with PBT (0.2% Triton X-100, PBS), discs were transferred into PBT (10% bovine serum, PBT) for 1 hr. Then, discs were incubated with staining solution (primary antibody, PBT) overnight at 4°C. After several washes with PBT to remove primary antibodies, discs were incubated with Cy5- or Cy3-labeled secondary antibodies for 1 hr. In those experiments that have DNA counterstaining, SYTOX (Molecular Probes Co.) was added to the secondary antibody staining solution to 0.1 µM. After several washes with PBT, discs were mounted in mounting solution (90% glycerol, 0.22 M 1,4-diazabicyclo-[2.2.2]octane) and viewed under a Zeiss confocal microscope.

Acknowledgments

We thank Hui-yun Chang, Francesca Pignoni, Tze-bin Chou, Pradip Sinha, Gines Morata, and the Bloomington Stock Center for providing the fly stocks, and Gines Morata for antibody. We are grateful for valuable suggestions from Cheng-Ting Chien and Don Ready, to Chen-shu Hsieh for preparing fly food, and to Chiu-Chung Tang for maintaining fly stocks. This study was supported by grants to Y.H.S. from the National Science Council (NSC-84-2311-B-001-087, NSC-86-2313-B-001-003, NSC-87-2311-B-001-106) and the Academia Sinica, Taiwan, Republic of China, and a grant from the Israel Cancer Association to A.S. C.-Y.P. is supported by a National Science Council postdoctoral fellowship. The publication costs of this article were defrayed in part by payment of page charges. This article must therefore be hereby marked “advertisement” in accordance with 18 USC section 1734 solely to indicate this fact.

Note added in proof

Sequence of the hth cDNA clones 7 and 5 have been deposited in GenBank (accession nos. AF035825 and AF036584, respectively).

References


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