

A conserved region of engrailed, shared among all en-, gsc-, Nk1-, Nk2- and msh-class homeoproteins, mediates active transcriptional repression in vivo

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SUMMARY

The engrailed homeoprotein is a dominantly acting or 'active' transcriptional repressor both in cultured cells and in vivo. When retargeted via a homeodomain swap to the endogenous *fushi tarazu* gene (*ftz*), it actively represses it, resulting in a *ftz* mutant phenocopy. We have mapped functional regions of engrailed using this in vivo repression assay. In addition to a region containing an active repression domain identified in cell culture assays (K. Han and J. L. Manley (1993) *EMBO J.* 12, 2723-2733), we find that two evolutionarily conserved regions contribute to activity. The one of these that does not flank the HD is particularly crucial to repression activity in vivo. We find that this

domain is present not only in all engrailed-class homeoproteins but also in all known members of several other classes, including goosecoid, Nk1, Nk2 and msh. Thus engrailed's active repression function in vivo is dependent on a highly conserved interaction that was established early in the evolution of the homeobox gene superfamily. We further show using rescue transgenes that the widely conserved in vivo repression domain is required for the normal function of engrailed in the embryo.

Key words: active repression, *Drosophila*, embryonic development, homeodomain, transcriptional repressor, engrailed, *fushi tarazu*

INTRODUCTION

Transcriptional repressors that function at a distance, analogously to transcriptional activators, have been termed active repressors (Jaynes and O'Farrell, 1991; reviewed in Gray et al., 1995). Functional distinctions within this class of effectors have been proposed based on their ability to act over different distances on the DNA (Gray et al., 1994). However, mechanistic distinctions between short- and long-range repressors have yet to be made. One active repressor that has been well characterized both in cultured cells and in vivo is the product of the *engrailed* locus of *Drosophila*. The engrailed protein (*en*) contains a homeodomain (HD) related in DNA-binding specificity to that of the *Antennapedia* (*Antp*) class (Desplan et al., 1988), but representing a separate, conserved class with two known members in both insects and mammals. Several members of the *Antp* class have been shown to be transcriptional activators, including the *fushi tarazu* protein, *ftz*. *Ftz* is a strong, context-independent activator in cultured cells (Jaynes and O'Farrell, 1988; Winslow et al., 1989), and participates in a direct positive feedback on its own gene in *Drosophila* embryos (Schier and Gehring, 1992). *En* is an active repressor in cultured cells and dissection of this activity showed that the N-terminal half, which does not include the HD (Jaynes and O'Farrell, 1991; Jaynes et al., 1990), or a small subdomain of that region (Han and Manley, 1993), can confer repression activity to heterologous DNA-binding domains.

Gene-specific transcriptional regulators generally have two activities that determine their function. First, they interact with specific sets of target genes, usually through sequence-specific

DNA-binding domains. Then they affect the expression of those target genes through activation or repression domains that are often separable from domains involved in DNA binding. This distinction between targeting and effector functions can be blurred by cooperative interactions with other regulators. However, the modularity of most well-studied factors, along with the promiscuous activity of most activation and repression domains analyzed to date, suggests that the two functions may often act independently in vivo. This view is supported by recent manipulations of binding sites within the *even-skipped* stripe 2 enhancer (Arnosti et al., 1996). One way to test this notion and to separately address the requirements for targeting and effector function in vivo is to 'retarget' a regulator by exchanging targeting domains between proteins with distinct effector functions. Using such a strategy, dominant negative effects have been seen in several contexts when the active repression region from engrailed was retargeted using the DNA-binding domain of an activator. Both in mammalian cells and in transgenic mice, the DNA-binding region of the *myb* proto-oncogene fused with the N-terminal portion of *en* resulted in a dominant negative effect (Badiani et al., 1994). Similarly, by swapping HDs between *ftz* and *en*, it was shown that *en* domains can confer a dominant negative activity on the *ftz* HD. When produced ubiquitously from a heat-inducible promoter, this chimeric repressor can specifically override the activity of endogenous *ftz* protein, causing repression of the *ftz* gene and generating a *ftz*-mutant phenotype in *Drosophila* embryos (John et al., 1995). This effect was determined to be the result of active repression by three criteria. First, this action of the chimeric protein, termed

EFE, depended on an N-terminal region from *en* implicated in active repression in culture. Second, EFE without this repression domain, while unable to significantly repress endogenous *ftz* gene expression, could still interact with *ftz* HD target sites. This was apparent from its ability to inhibit the activity of the autoregulatory *ftz* enhancer driving a *lacZ* reporter in vivo, presumably by competing for *ftz* binding sites, as this enhancer was previously shown to be a direct target of the *ftz* HD (Schier and Gehring, 1992). A third indication that the repression is active was that EFE repressed another *ftz* target gene (the *engrailed* gene) outside the domain of endogenous *ftz* expression. Thus, the action of EFE could not be attributed solely to preventing *ftz* from binding to its target genes.

Here we show, using repression of endogenous *ftz* as the primary assay, that the repression function of EFE is contributed by several domains, including the C-terminal region flanking the HD, but that another conserved region found in the N-terminal repression domain is particularly important. This crucial region is distinct from the region found to be most active in culture (Han and Manley, 1993) and is homologous not only to all other *en* class homeoproteins, but to all known members of four other homeoprotein classes: *msh*, *Nk1*, *Nk2* and *gooseoid*. These results suggest that a highly conserved interaction, established early in homeoprotein evolution, mediates active transcriptional repression by *engrailed* in vivo. We further show that this domain is required for the function of wild-type *engrailed* protein expressed in its normal pattern in early embryos.

MATERIALS AND METHODS

Embryo preparation and staining

P-element transformations (Spradling and Rubin, 1982), cuticle preparations (Wieschaus and Nüsslein-Volhard, 1986) and in situ hybridization to fixed embryos (Edgar and O'Farrell, 1990) were performed as described previously. Antibody staining was performed essentially as described (Manoukian and Krause, 1992) using a polyclonal α -*en* antisera, a kind gift of Charles Girdham and Patrick O'Farrell, which had been prepared against full-length, partially purified GST-tagged *en* and affinity purified against a His-tagged peptide with the N-terminal 150 amino acids of *en*. Either alkaline phosphatase-(AP) or peroxidase-coupled secondary antibodies (Vector Laboratories) were used both for microscopic examination of fixed embryos, where either BCIP and NBT (for AP) or DAB substrates were used for staining (Boehringer Mannheim) and for quantitation of antibody signals, where the AP substrate p-nitrophenyl phosphate (Sigma) was used as described (Manoukian and Krause, 1992). Incubation times were determined to be in the linear range of the assay by incubating sets of embryos with different signal intensities for various times.

Heat shocks were administered to embryos on 35 mm collection plates by floating the plates on 37°C water inside a sealed container, in order to minimize evaporative cooling. Standard heat-shock conditions employed a 15 minute incubation followed by return to a 25°C humidified environment.

Plasmid constructions and *Drosophila* strains

Expression plasmids for EFE derivatives were modifications of a P-element transformation vector capable of providing inducible expression of EFE in transformed *Drosophila* from a heat-shock promoter (described in John et al., 1995). Modifications were made using either PCR-based methods (for $\Delta 5$ and $\Delta 6$), synthetic DNA

adaptors to create deletions adjacent to unique restriction sites (for $\Delta 234$, $\Delta 23$, $\Delta 34$, $\Delta 3$ and $\Delta 4$), or a combination of the two ($\Delta eh1$, F \rightarrow E, Meh1). Resulting deletion end points and amino acid substitutions are described in figure legends and the text. All regions containing synthetic or PCR-synthesized DNA were subsequently sequenced (automated) to confirm the expected structure. Appropriate restriction fragments were combined to generate the combined deletion plasmid $\Delta 46$. Details are available on request. These plasmids were introduced into flies using standard methodologies (Spradling and Rubin, 1982). Homozygous viable insertions on either the second or third chromosome were used in all analyses of repression activity.

The embryonic *en* expression/rescue construct p31Rg-*en* was constructed by combining elements from three parent plasmids: *en* downstream-SV40 poly(A) region and vector sequences from pRK232 (Heemskerk et al., 1991), *en* intron and partial coding region from pen7.5pUC (a subclone made by J. Kassis from *en* genomic clones described in Kuner et al., 1985) and partial *en* coding, promoter and upstream regions from P[*en/lac*] (DiNardo et al., 1988). Additional details are either contained in figure legends or text or are available on request.

Transfections

Cell culture assays were performed using *Drosophila* S2 cells as described previously, with 2 μ g per 60 mm culture dish of the reporter gene T₃N₆D-33Cat_B (Jaynes and O'Farrell, 1991) and 0.3 μ g of *ftz* expression plasmid pP_{Ac}-*ftz* (Jaynes and O'Farrell, 1988; Winslow et al., 1989). CAT assays, as well as β -galactosidase assays for expression of the cotransfected reference gene pLac82SU (Dorsett et al., 1989), were performed as described (Jaynes and O'Farrell, 1991). Co-transfected plasmids used to express EFE and its derivatives were the same as those used for P-element transformation, wherein expression is driven by the *hsp70* promoter.

Database searches and sequence comparison

Searches of GenBank and EMBL databases were performed using the BLAST (at the NCBI using the BLAST network server), FASTA and TFASTA programs of the Wisconsin Package, version 8 (Sept. 1994, Genetics Computer Group, 575 Science Drive, Madison, WI). Additional sequence analysis utilized GCG software, including PileUp, used to order the sequences of Fig. 4, and MacVectorTM version 4.1.4.

RESULTS

Repression activity in vivo is determined by multiple *en* domains

Previous results showed (John et al., 1995) that a chimeric protein, consisting of *en* with its HD replaced by that of *ftz*, termed EFE (Fig. 1), was capable of specifically repressing *ftz* target genes in vivo, including its own gene, *ftz*. This repression required not only the *ftz* HD, but also a portion of *en* implicated in active repression in cultured cells. This was demonstrated by characterizing a deletion derivative of EFE, which removed both of the *en*-derived regions 3 and 4. This deletion derivative (EFE $\Delta 34$) retained the ability to localize to nuclei in embryos, as well as the ability to repress transcription when competing for binding sites in cultured cells (passive repression), but had lost most of its active repression activity in culture. Consistent with the involvement of active repression in the action of EFE in vivo, this derivative was unable to effectively repress *ftz* gene expression in embryos. Despite this loss of activity, it could still substantially reduce the activity of a transgenic copy of a *ftz* autoregulatory enhancer driving a *lacZ* reporter in embryos. This *ftz* enhancer has been implicated

as a direct *in vivo* site of action of the endogenous *ftz* protein (Schier and Gehring, 1992), indicating that EFE Δ 34 could still interact with a *ftz* HD target site *in vivo*. The difference in response between the endogenous *ftz* gene and the isolated upstream enhancer may be due to endogenous *ftz* expression



Fig. 1. Features of the *Drosophila* engrailed-fushi tarazu chimera, EFE. The diagram indicates which portions of the coding sequence derive from *en* and which from *ftz*, our numerical designations of regions of *en* (1-6, not including the *ftz* HD), and the locations of known features within those regions (eh1, eh2, eh5 and R). eh1, eh2 and eh5 are peptide sequences found in all known *en* homologs (Logan et al., 1992) from widely divergent species, including insects and mammals, and R is an autonomous active repression domain identified in cell culture studies (Han and Manley, 1993). Homology eh1 is described further in Fig. 4, while eh2 and eh5 are part of the conserved regions flanking the *en* HD, which also include a sequence termed eh3 (immediately flanking the N terminus of the *en* HD) that has been implicated in nuclear localization (S. J. Poole, unpublished observation), and so was left intact in our analyses. Locations of region boundaries in the amino acid sequence are given below the line. Deletions and other alterations of these regions are described in detail in subsequent figures or in the text.

becoming independent of direct autoregulation once it is established, while it remains subject to active repression by EFE (probably acting through the upstream enhancer). A further indication that EFE is not simply competing for *ftz* target sites came from the observation that another *ftz* target gene, the *engrailed* gene, was repressed not only where *ftz* is present (and required to activate *en*), but also outside the *ftz* domain, in the *ftz*-independent (odd-numbered) *en* stripes (again, EFE may be acting through normal *ftz* binding sites, but counteracting the effects of other activators). Thus the ability of EFE to repress *ftz* gene expression *in vivo* appears to require an activity in addition to targeting. The region removed in EFE Δ 34 contains two interesting features. One is a previously identified homology found in all known *en*-class homeoproteins (Logan et al., 1992), located in region 3, and the second is a minimal active repression domain identified in cultured cell assays (Han and Manley, 1993), located in region 4 (see Fig. 1).

We mapped the domains responsible for the active repression activity of EFE *in vivo*, using the ability to repress endogenous *ftz* gene expression as an assay. A set of transgenic flies was constructed, each expressing a deletion derivative of EFE from a heat-inducible promoter. As described previously, a brief heat pulse induces ubiquitous expression from the transgene. Such expression of EFE causes rapid and persistent (Fig. 2B) loss of *ftz* expression in the trunk region (stripe 7 recovers). This results in the generation of pair-rule deletions

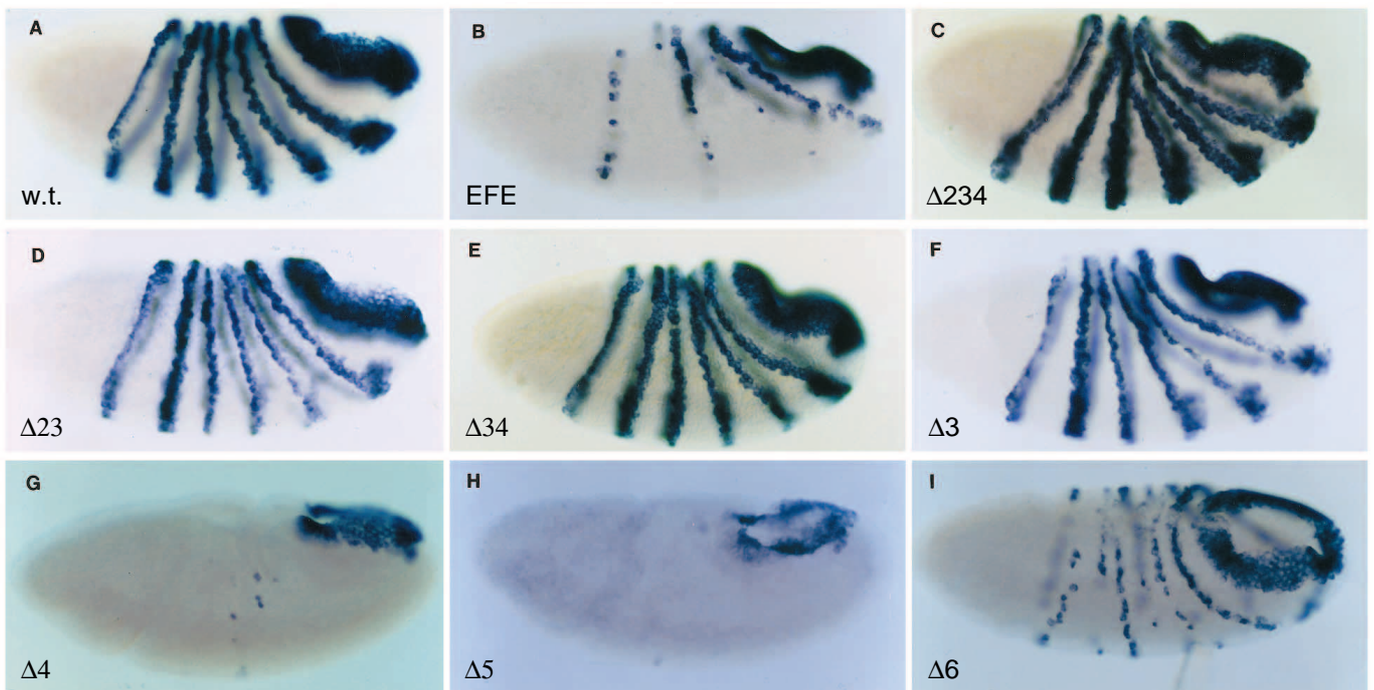


Fig. 2. Repression of endogenous *ftz* gene expression by EFE and derivatives. The pattern of *ftz* RNA expression was monitored in embryos after induction of the indicated EFE derivative from a heat-inducible transgene. Embryos were fixed 40 minutes after induction and stained by *in situ* hybridization (as described in Materials and Methods) using a probe to *ftz* RNA sequences outside the HD. Expression was induced by giving a 15 minute heat pulse at 37°C (see Materials and Methods) beginning 2 hours 40 minutes after egg collection. Derivatives are named for the regions deleted. They remove the following amino acids of the 552 total (the *ftz* HD consists of amino acids 454-513): 82-399 (Δ 234), 82-221 (Δ 23), 169-399 (Δ 34), 172-216 (Δ 3), 229-399 (Δ 4), 407-440 (Δ 5) and 523-531 (Δ 6). Thus Δ 23 removes the entire eh1 homology plus additional sequences, Δ 3 removes the most conserved portion of eh1 plus additional sequences, Δ 5 and Δ 6 are partial deletions of the most conserved portions of the homologies eh2 and eh5 (see Fig. 1 legend), Δ 4 removes the R domain plus additional sequences and Δ 34 removes both R and the most conserved portion of eh1.

in the cuticle pattern at the end of embryogenesis that mimic those seen in *ftz* mutants (John et al., 1995). Such heat treatment had no effect on endogenous *ftz* expression in wild-type embryos (Fig. 2A). In testing derivatives deleted for enderived portions of EFE, we discovered that multiple regions contribute to activity. First, each of the deletions that remove region 3 caused substantial loss of activity (Fig. 2C-F). However, EFE Δ 3 retains sufficient activity to cause significant repression of *ftz* expression (Fig. 2F), as the remaining stripes were discontinuous either laterally or dorsally. The additional deletion of region 2 caused no further loss of repression activity (Fig. 2D). In contrast, additional deletion of region 4 did cause a further reduction in activity, to the point that EFE Δ 34 produced no lasting repression of the *ftz* gene (Fig. 2C,E; see John et al., 1995 for description of a mild, transient effect of EFE Δ 34). Nonetheless, EFE Δ 34, as described above, is still able to repress the autoregulatory *ftz* upstream enhancer, indicating that it retains targeting activity *in vivo*. In contrast, deletion of either region 4 or 5 alone resulted in increased repression activity (Fig. 2G,H; note that Δ 5 is a partial deletion of region 5: Figs 1 and 2 legends). In the case of EFE Δ 4, this may be attributed to increased protein stability (for EFE Δ 5, see below and Discussion). In region 6, a deletion of the most conserved 9 amino acids within the en C-terminal tail, which are the core of the C-terminal extension of the en HD, caused a partial loss of repression activity (Fig. 2I; for simplicity, we refer to this directed deletion as Δ 6). The significance of this loss of activity was tested by examining both hatching rates of embryos and the cuticle defects caused by induction of EFE Δ 6. Relative to lines expressing the parental EFE, the hatching rate was significantly increased and the severity of cuticle defects was clearly reduced in EFE Δ 6 lines (data not shown), confirming the loss of activity due to this small deletion.

To determine whether the observed changes in activity could be attributed to differences in expression levels from the transgenes, we stained embryos following heat induction with an anti-en antiserum that reacts with the N-terminal region, including region 1 and part of region 2 (John et al., 1995). As previously shown for EFE Δ 34 (John et al., 1995), each derivative retained its ability to localize to nuclei (not shown) and was expressed initially at levels comparable to EFE (Fig. 3). In addition, while most derivatives decayed with a time course very similar to that of EFE, both EFE Δ 34 and EFE Δ 4 were significantly more stable. This was confirmed in two other experiments, in which the apparent half-life of EFE Δ 4 was estimated to be about two-fold longer than that of EFE. These experiments also suggested that EFE Δ 3 has a very slightly increased half-life (data not shown). Thus, the increased activity of EFE Δ 4 may be accounted for by its increased stability, but the increased activity of EFE Δ 5 is apparently not due to increased expression or stability. Likewise, the loss of activity of other derivatives cannot be explained by differences in expression or subcellular localization.

Region 3 contains similarity to other classes of homeoproteins

An obvious candidate for mediating the activity of region 3 was a previously identified region of homology among en class homeoproteins (Logan et al., 1992). In database searches using this 'eh1' homology region, we found that a similar motif is present in several other classes of homeoproteins (Fig. 4),

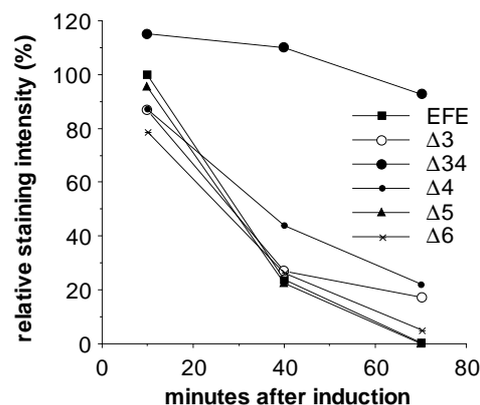


Fig. 3. Protein levels following induction of transgene expression. Embryos were fixed at the indicated times after induction and stained with α -en antibodies (affinity purified using the N-terminal 150 amino acids of en, present in each of these derivatives) and with alkaline phosphatase (AP) coupled 2^o antibodies. AP activity was quantified using the soluble-product-producing substrate PNP (Manoukian and Krause, 1992). The assay was within the linear range, since longer incubation times resulted in further linear increases in signal and both longer heat shocks with the same number of embryos and the inclusion of more embryos in parallel reactions gave approximately proportional increases in signal.

which is always located N-terminal to the HD, with a variable length stretch of non-conserved amino acids intervening between the two (see Fig. 4). For the major classes, *msh*, *gooseoid*, *Nk1*, *Nk2* and *engrailed*, all known members (for which complete sequence information is available) contain the motif (J. B. Jaynes, unpublished observation). In addition, two 'novel' class members, *XANF1* and *HESX1*, as well as *Nkx5.1*, were also found to contain it (Fig. 4). In general, the apparent relatedness of these motifs parallels that of the homeodomain class to which the protein belongs, although there may be exceptions. Interestingly, there are apparently several subclasses of the motif conserved independently between vertebrates and invertebrates, including one conserved to the flatworm *S. mansoni* that is closely related among *Nk2*-class members (Fig. 4). This suggests the possibility that the different versions of the motif mediate interactions with a family of other factors. Homology within each class extends a variable distance to either side of the core motif, but similarity between the classes falls to an undetectable level within 10-20 amino acids. No reliable indications either of secondary structure or of similarity with other known domains was detected.

The eh1 homology mediates repression of *ftz* in vivo by EFE

In order to test whether eh1 is required for repression by EFE, we constructed both a small deletion within eh1 and a single point mutant at the most conserved position. Both a 15 amino acid deletion removing the most conserved portion of eh1 and a change of the invariant Phe to Glu (F \rightarrow E) resulted in derivatives of EFE with strongly reduced ability to repress *ftz* in early embryos. The levels of *ftz* RNA were reduced only slightly relative to wild type following induction of each of these derivatives (Fig. 5). Thus each of these changes in eh1 had an effect on EFE activity indistinguishable from that of removing region

D.mel.	<i>en</i>	EPFLAFSISNLSDRFGDVQKPG
R.nov.	<i>TTF-1</i>	HTTFFSVDILSPLEESYKKG
B.rer.	<i>eng-2</i>	HRITNFFIDNLRPDRGRKKEAN
B.rer.	<i>eng-3</i>	HRITNFFIDNLRPDRGRRKEGS
H.sap.	<i>En-2</i>	HRITNFFIDNLRPDRGRKRDAG
M.mus.	<i>En-1</i>	HRITNFFIDNLRPDRGCKKEQP
Trib.	<i>en</i>	NGNLKFSIDNLRKADFGRRITDP
D.mel.	<i>inv</i>	ERALKFSIDNLRKADFGSRRLPKI
M.mus.	<i>Hlx1</i>	SIDLKFGIDRILSAEDDPKVKEG
H.ror.	<i>AHox1</i>	STVLKFGIDSILKNKNEKVPKG
C.ele.	<i>en</i>	EMILKFGIERILSPPCPSPTI
D.mel.	<i>S59</i>	TRFLAFSISNLRPDRGRKKEAN
M.mus.	<i>Nkx1.1</i>	HKKSFSSYLDLIDPQKFTBAAPP
X.lav.	<i>XANF-1</i>	SPSSFSIHLGIDDKKTDVASS
M.mus.	<i>HESX1</i>	PAPCSFSIKSILGIDDKTCTTS
X.lav.	<i>Nkx2.3c</i>	VTTFSSYKDLNLRQQGQPPA
M.mus.	<i>Nkx2.5</i>	LTPTPFSYKDLNLRQQQRSLAS
D.mel.	<i>msh-2</i>	LNTTFFSYKDLNLRVNTQTEAYEG
B.rer.	<i>NK-2.2</i>	NIKTGFSSYKDLNLRPDNDDEEBS
S.man.	<i>smox-2</i>	XXANGFEVYKDLNLRDKHVIIRQ
X.lav.	<i>gsc</i>	MPGCMFSIDNLRARPRCESYL
M.mus.	<i>gsc</i>	MPAGSMFSIDNLRARPRCDAVLL
B.rer.	<i>gsc</i>	MPAGMFSIDSILGRPSCDASVLL
D.mel.	<i>gsc</i>	AAASLFTIDSILGSRQQGGGTAP
X.lav.	<i>msx-1</i>	PGILPFSYKDLNLRDRKPGDRDLD
M.mus.	<i>msx-1</i>	ASLPLFSYKDLNLRADHRKPGAKES
B.rer.	<i>mshC</i>	ALPLPFSYKDLNLRDRTSSRTLYE
D.mel.	<i>H2.0</i>	SVKLSFSYKDLNLRGSPPEESYKQ
D.mel.	<i>msh</i>	SRLSNFSYKDLNLRDTRPRTPPNQ
M.mus.	<i>Nkx5.1</i>	EGAAGFALSQVGDNAFPRFEIPL
	Consensus:	HRALPFSIDNLRPDRGRKKG

Fig. 4. The engrailed eh1 region has similarity to other classes of homeoproteins. Selected members of each of the homeoprotein classes found to contain an eh1-homologous motif are shown. These were selected to show the range of variation within each class between vertebrate and invertebrate species. The consensus shows the most highly represented amino acid at each position. Dark shading in the consensus indicates similarity in at least 30% of the individual sequences. Dark shading in the individual sequences indicates exact match to the consensus, while lighter shading indicates similarity to the consensus. The invariant Phe (F) is residue 175 of the *D. melanogaster* engrailed sequence. For each major class represented, all known members for which the sequence of the N-terminal region is available contain the homologous region. Species designations are followed by gene names, either as given in Duboule (1994), if included therein, or otherwise as in the GenBank and EMBL databases. *S. mansoni* (a flatworm) is the most divergent species found to contain a homologous sequence. *C. elegans* (roundworm), *D. melanogaster* (fruit fly), *Tribolium* (flour beetle), *H. roretzi* (an ascidian), *B. rerio* (zebrafish), *X. laevis* (frog), *R. norvegicus* (rat), *M. musculus* (mouse) and *H. sapiens* (human) are also represented. The *msx-1s* contain an *msh*-class homeobox (HB), while *msh-2*, a.k.a. *Nk-4* or tinman, contains an *Nk2*-related HB, and *XANF-1* and *HESX1* are in a novel HB class. *invected* (*inv*) is the second *en*-class gene that is found in some insects, *engs* contain an *en*-class HB (there are three in zebrafish, Ekker et al., 1992), while *S59* contains an *Nk1*-related HB and *TTF1* contains an *Nk2*-related HB. See Duboule (1994) for evolutionary relationships based on HB sequence and more detailed descriptions.

3 entirely. To test whether the conservation of this region from flies to mammals had preserved function, we replaced the 15 amino acid region of the *Drosophila* protein with the corresponding region from the mouse *En1* protein. This resulted in 4 non-conservative, 3 neutral and 1 conservative substitution within the region. As shown in Fig. 5, this replacement fully restored the ability of EFE to repress the endogenous *ftz* gene in *Drosophila* embryos, indicating that the function required for this activity, presumably active repression, is conserved. In contrast to the drastic effect of mutating region 3, combining two deletions that each reduce active repression in culture, $\Delta 4$

and $\Delta 6$ (Han and Manley, 1993; data not shown), resulted in a protein (EFE $\Delta 46$) still capable of repressing *ftz* (Fig. 5D). Examination of protein levels produced in embryos showed that, for those mutated in region 3, the less active proteins were produced at slightly higher levels than were the more active ones, while all were about equally stable (Fig. 6A). For EFE $\Delta 46$, the levels were slightly higher initially and the protein was considerably more stable than EFE, perhaps contributing significantly to its repression activity. However, EFE $\Delta 46$ is less stable than EFE $\Delta 34$ (Fig. 3), but nonetheless is a better repressor of *ftz* (compare Fig. 5D with Fig. 2E), indicating that it still has considerable active repression function in vivo, consistent with the strong in vivo activity of region 3.

To test whether the alterations in region 3 might be affecting the ability of the *ftz* HD to bind to DNA, we tested several derivatives for their ability to compete for *ftz* binding sites in cultured cells. As shown in Fig. 6B, neither altering the eh1 domain nor removing region 3 affected the ability to repress transcription by competing for *ftz* binding sites. Since each of these derivatives also localize to nuclei in embryos (data not shown) and, in addition, retain the ability to partially repress endogenous *ftz* gene expression (seen more clearly at earlier times after induction than that shown in Fig. 2), it is likely that they retain their DNA-binding activity in vivo as well. Thus eh1 appears to mediate the effector function of EFE, that is, its ability to actively repress the endogenous *ftz* gene, rather than its targeting function.

eh1 mediates normal *en* function in the embryo

To test whether the eh1 domain is required for the function of wild-type *en* in vivo, we used a partial rescue assay. Although a *cis*-acting region capable of fully complementing an *en* mutant has not been identified, we were able to achieve good partial rescue of the embryonic pattern elements lost in the trunk region of *en* mutants by combining two previously characterized *cis*-acting regions. These were a portion of the *en* upstream region previously shown to direct part of the normal *en* expression pattern (DiNardo et al., 1988), and the *en* introns, the larger of which drives a slightly more complete *en* pattern in the embryo than the upstream fragment alone (Kassis, 1990). A construct containing these *cis*-elements driving normal *en* protein expression, in a strong *en* mutant background (*en*^{CXI}, Heemskerk et al., 1991), prevents the fusion of ventral denticle bands in the thorax and abdomen that is characteristic of *en* mutants (Fig. 7C,D versus B; Nüsslein-Volhard and Wieschaus, 1980). This function of *en* is known to be mediated by the maintenance of *wg* expression in the cell row adjacent to each *en* stripe in the early embryo. The rescue construct also restores many, but not all, of the pattern elements within ventral denticle bands that are lost in *en* mutants. This aspect of pattern is thought to require a relatively late function of *en* that maintains expression, in *en*-expressing cells, of the hedgehog signaling molecule (for a review of these interactions, see DiNardo et al., 1994). This partial rescue of late function is consistent with the above described patterns of expression driven by each of the *cis*-acting elements in our construct, which fade about half way through embryogenesis, while normal *en* expression persists to very late stages. Thus our construct provides good rescue of early *en* function and partial rescue of late *en* function when it is used to express a wild-type *en* protein.

To test whether the eh1 region is required for this rescue activity, we used the same *cis*-elements to drive expression of two altered *en* proteins. The first is *en* Δ eh1, which is deleted for the 15 amino acid core of the homology region, and the second is the point mutated *en*F \rightarrow E derivative. Both of these constructs gave similar profiles of partial rescue that were much weaker than that produced by the wild-type rescue construct. Each resulted in restoration of some naked cuticle between ventral denticle bands (Fig. 7E,F and data not shown), but the degree of rescue did not approach that shown by a high percentage of embryos with the wild-type construct (Fig. 7C,D). A comparison of RNA levels expressed by the rescue constructs showed similar levels and patterns of expression for each, with the *en* Δ eh1 line showing a slightly longer persistence of expression than the *en*F \rightarrow E and the wild-type rescue lines, which were indistinguishable (data not shown). When we compared the range of phenotypes shown by heterozygous rescue lines (in which about 2/3 of the rescued embryos have a single copy of the rescue construct and 1/3 have two copies) we see little, if any, overlap between the phenotypes of the eh1-mutated and the wild-type constructs. In contrast, the range of phenotypes of *en* mutant embryos does overlap that of embryos rescued by the eh1-mutated constructs. This suggests that one copy of the wild-type construct provides more rescue activity than two copies of the eh1-mutated construct. Thus, eh1 contributes substantially to the ability of *en* to carry out its normal embryonic functions.

DISCUSSION

A conserved repression domain

Analysis of *en* repression function in vivo has shown that one domain is of critical importance. In this assay, *en* is retargeted in vivo to the endogenous *ftz* gene, by replacing the *en* HD with that of *ftz*, resulting in repression of the *ftz* gene by the chimera (called EFE). Previous work indicates that targeting activity is provided by the *ftz* HD, since normal *en* protein is unable to significantly affect *ftz* gene expression. EFE appears to repress *ftz* expression by an active mechanism, as a derivative containing the *ftz* HD but deleted for *en* domains 3 and 4 is unable to repress *ftz*, but is still capable of interacting with a *ftz* upstream enhancer in a reporter transgene. That EFE is utilizing an active mode of repression in vivo is also indicated by its ability to repress another *ftz* target gene (*en*) outside the region of *ftz* expression (John et al., 1995). The *en*-derived region that is most crucial to this repression activity (region 3) contains the single conserved domain not closely associated with the HD in the primary sequence. Either deleting the core of this homology region, which was previously noted in all *en*-class homeoproteins, or mutating

the most conserved amino acid, Phe 175, strongly reduces repression activity in vivo, equivalent to deleting all of region 3 (Figs 5, 3).

The repression activity apparently conferred by the eh1 homology region suggested that active repression is a conserved function of the *en* homeoprotein family and led us to conduct a directed search for similarity to other protein domains in the database. We found that regions homologous to eh1 are present in all known members of several other classes of HD-containing proteins. In each case, the similarity is found upstream of the HD, with a variable-length stretch of non-related amino acids between, typically 80 or more. Representative members of each class are shown in Fig. 4. In each class for which several members are known, a distinct subtype of the eh1 motif is recognizable. For example, within the classes expressed predominantly in mesodermal derivatives, *msh* and *Nk2*, a Val is found at the 2nd position after the invariant Phe and, while the *Nk2* members have a basic residue at the next position (3rd following the Phe), most other classes, including *msh*, contain an acidic residue at this position. Such patterns of conservation suggest independently conserved functional divergence among the classes, such as might occur in conjunction with the evolution of a family of interacting proteins, with specific members of this family diverging to interact with specific members of the eh1 family. This view allows that there may be significant divergence of function associated with each distinct partner in the putative interacting family. It may be that as the interacting partner of each eh1 class diverged, the transcriptional activity was modified and perhaps reversed. Such a possibility is underscored by the recent finding that one of the proteins containing eh1 similarity, *Nkx2.5*, can function as an activator of transcription when its C-terminal region is deleted (Chen and Schwartz, 1995). However, it is likely that the specific transcriptional activity of

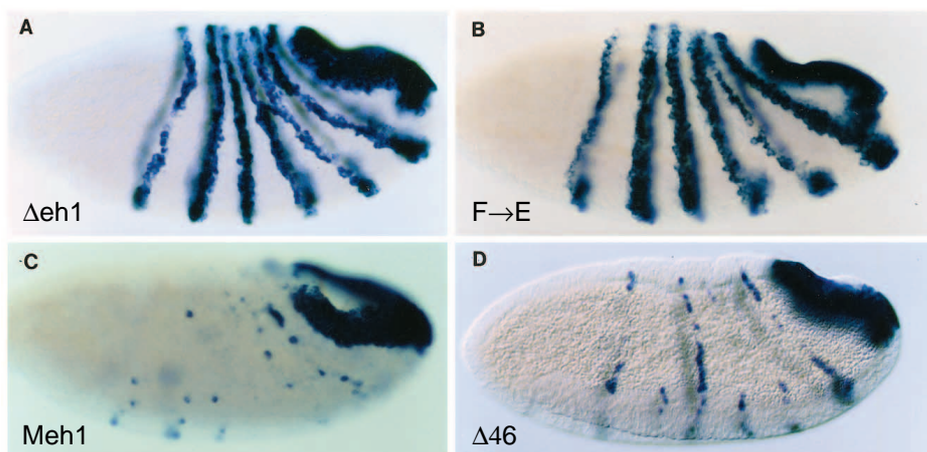


Fig. 5. A single point mutation in eh1 reduces, and the mouse eh1 region restores, repression activity in vivo. The ability of derivatives of EFE mutated in region 3 to repress endogenous *ftz* gene expression was compared to activity of a derivative that removes two other domains implicated in active repression in culture, regions 4 and 6. Endogenous *ftz* RNA levels were detected as in Fig. 2. (A) EFE Δ eh1, which removes amino acids 172-186, the core of the eh1 homology, and (B) point mutation of the most conserved amino acid, Phe 175, to Glu each cause a drastic reduction in the ability to repress *ftz*, while replacement of the Δ eh1 region by the corresponding mouse *En1* region restores repression function (C). In contrast, combined deletion of regions 4 and 6 (D) does not reduce activity as much as the point mutation in region 3.

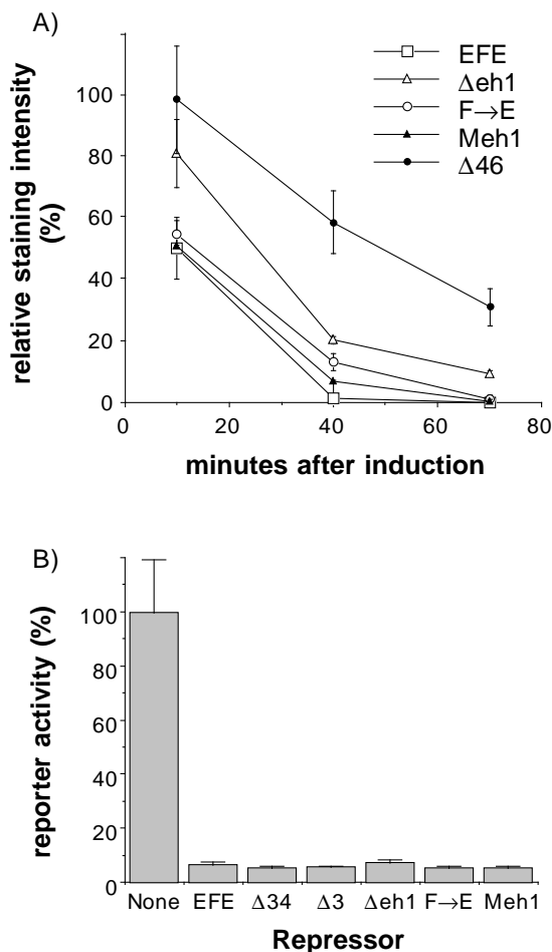


Fig. 6. (A) Changes in eh1 do not reduce protein levels in vivo. Embryos from each transgenic line were stained for transgene-produced protein following induction and the results quantified, as in Fig. 3. Note that the alterations in region 3 that reduce activity (Δ eh1 and F→E) do not reduce either protein levels or stability, while EFE Δ 46 is significantly more stable than EFE. (B) Competition for binding sites by EFE and derivatives in cultured cells. *Drosophila* S2 cells were cotransfected with a CAT (chloramphenicol acetyltransferase) reporter plasmid, which contains binding sites for the ftz HD upstream of a basal promoter and a plasmid that expresses ftz (see Materials and Methods for details). Ftz-activated reporter expression about 1000-fold above the basal level (activated level shown as 100%). The ability of EFE and the indicated derivatives to repress this activated transcription (by competing with ftz for binding sites in the reporter) was determined by cotransfection of 3 μ g of the appropriate expression plasmid, which in each case is the same P-element transformation vector used to construct the corresponding transgenic fly line shown in Fig. 2. The non-repressed level was determined by cotransfection of 3 μ g of 'empty' parental vector. CAT activity was determined and normalized to the activity of a cotransfected reference gene (see Materials and Methods for details). The graph represents the average and range of at least 2 independent transfections.

eh1 is conserved among the en homologs, since replacing the *Drosophila* eh1 with mouse eh1 restores repression function in *Drosophila* (Fig. 5). Active repression function may extend at least as far as the goosecooid class, as the *Drosophila* gsc protein (Gorieli et al., 1996) was recently found to be an active

repressor in cultured cells (C. Mailhos and C. Desplan, personal communication).

We note that there are possible similarities between eh1 and other conserved motifs, some of which have been proposed previously. For example, eh1 resembles the conserved octapeptide found downstream of some paired domains (reviewed in Noll, 1993; this was noted in Allen et al., 1991, along with the similarity between Hlx, H2.0 and en eh1, which was termed the 'Hep' motif). It also is similar to the first helix of the HD (not part of the helix-turn-helix region directly involved in DNA binding), as well as the last helix (helix 6) of the paired domain. However, in our database searches, these other potential homologies were not reliably recognized as significant against a background of apparently random similarities.

While it remains possible that eh1 has an effect on targeting to the *ftz* gene, we consider this unlikely in light of the following. First, it would be doing so in the context of the ftz HD, which has been implicated in direct targeting to the *ftz* upstream enhancer without the aid of an eh1-homologous region (that is, in the context of normal ftz protein). The ftz HD seems to be sufficient for such targeting, since EFE Δ 34, which does not contain eh1, is still able to repress the activity of that enhancer in vivo, presumably by competing for ftz binding sites (John et al., 1995). Second, en is not targeted to the *ftz* gene in vivo, since ectopic en expression does not repress *ftz* (John et al., 1995) and since *en* mutants do not affect *ftz* expression (Carroll et al., 1988). Therefore, eh1 would have to be aiding the ftz HD in targeting to the *ftz* gene in some non-specific way. Third, eh1 does not affect DNA binding in cultured cells, since derivatives that remove it repress transcription in a passive repression assay to the same extent as the parental EFE, that is when competing for binding sites with the activator, ftz (Fig. 6B). In addition, each of the derivatives altered in region 3 retain the ability to partially repress endogenous *ftz* expression (Figs 2, 5), suggesting that they still interact with the *ftz* gene in vivo. This repression activity is seen more clearly immediately after induction of transgene expression, before *ftz* expression is allowed to recover (our unpublished observations). Taken together, the results indicate that eh1 is providing an effector function in vivo (active repression) in the context of EFE, rather than a targeting function.

Multiple en domains contribute to active repression

In addition to eh1, a conserved region that normally flanks the C terminus of the en HD (and thus flanks the ftz HD in EFE) also contributes to repression activity (Fig. 2). This is interesting in light of the involvement of conserved regions flanking the HDs of HOX gene products in determining their functional specificities in vivo (Mann and Hogness, 1990; Lin and McGinnis, 1992; Zeng et al., 1993). Differences in these regions might affect transcriptional activity, leading to different activities on common target genes, rather than, or in addition to, providing selective targeting to distinct sets of target genes.

The requirements for repression by EFE in vivo may be somewhat different from those in cultured cells, where region 3 was not found to contribute significantly to repression activity (although it appeared to confer an extremely weak activity when fused to regions 5 and 6; Han and Manley, 1993). However, region 4, which contains the single strong repression domain previously localized in en (Han and

Manley, 1993), also appears to contribute to repression activity in vivo. Removing this region in addition to region 3 clearly reduces repression activity more than removing region 3 alone (Fig. 2; John et al., 1995). Removing region 4 alone results in a more stable protein (Fig. 3), possibly masking a reduction in potency, as EFE Δ 4 represses *ftz* more completely than the parental EFE. The increased stability of EFE Δ 4 relative to EFE, while it is only about two-fold, may well account for the increased repression activity, since effective repression of *ftz* probably involves the destabilization of a positive feedback loop. This is suggested by the all-or-none decision that each cell makes following EFE induction with regard to *ftz* expression, as seen in the discrete pattern of expressing and non-expressing cells in the embryos (Fig. 2). Thus, it may be necessary to keep *ftz* levels below a threshold for a period of time in order to disengage the feedback loop and a two-fold increase in protein stability may therefore have a strong impact on the number of cells that lose *ftz* expression. Since region 3 is clearly crucial for effective repression in vivo, it is likely that in vivo repression requires a function in addition to that identified in the cell culture transfection studies. Further analyses will be required to determine whether this additional activity is mechanistically distinct from that operating in the transfection assays.

The conserved region that flanks the N terminus of the HD has an interesting effect in vivo. Without increasing the stability of the protein, deleting this region (Δ 5) actually increases repression activity (Fig. 2). As this region does not appear to provide either activation or anti-repression function in cell culture assays (our unpublished observation), it may have an effect on targeting in vivo. Perhaps region 5 mediates an interaction normally involved in targeting by *en* and this targeting function interferes with that of the *ftz* HD. Then removing it might allow increased *ftz* repression by EFE.

The involvement of multiple domains in repression by *en*, the lack of apparent activation activity in any of our derivatives (our unpublished observations; however, Han and Manley, 1993, found that an *en* derivative containing regions 2, 5 and 6 could weakly activate in culture) and the involvement of highly conserved motifs in repression activity in vivo, taken together, suggest that repression may be the primary effector function of *en*. Genetically, *en* is a repressor of several genes in *Drosophila*, including *even-skipped* (Harding et al., 1986; John et al., 1995), *cubitus interruptus* (Eaton and Kornberg, 1990; Schwartz et al., 1995), *wingless* (Heemskerk et al., 1991), *patched* (Hooper et al., 1989), *Ultrabithorax* (Carroll et al., 1988) and *decapentaplegic* (Sanicola et al., 1995). However, the idea that *en* might be a dedicated repressor in vivo conflicts, superficially, with results from ectopic expression assays in embryos, in which *en* has been shown to induce expression of its own gene (Heemskerk et al., 1991), as well as with the positive regulatory effects of *en* on *hedgehog* (Tabata et al., 1992) and *polyhomeotic* (Serrano et al., 1995). That these latter interactions might be indirect, through repression of a repressor, is suggested by our results. However, it remains possible that protein-protein interactions allow *en* to have a net positive regulatory effect on some direct target genes. It is worthy of note in this context that a similar

positive autoregulatory effect of *even-skipped*, a repressor in both cell culture assays (Jaynes and O'Farrell, 1988; Han et al., 1989) and in vitro (Biggin and Tjian, 1989), has recently been explained by indirect effects in vivo, involving repression of other repressors (Fujioka et al., 1995).

eh1 is required for normal *en* function in embryos

To test whether the conservation of *eh1* in *en* is related to known functions in patterning the *Drosophila* embryo, we used an in vivo assay for function of the normal *en* protein. Since a *cis*-regulatory region sufficient to fully complement an *en* mutant has not been identified, we combined two known regulatory regions to generate early embryonic expression, sufficient to complement the major pattern defects of *en* mutants in the trunk region. This is the place and time in development where *en* function has been most extensively studied and where

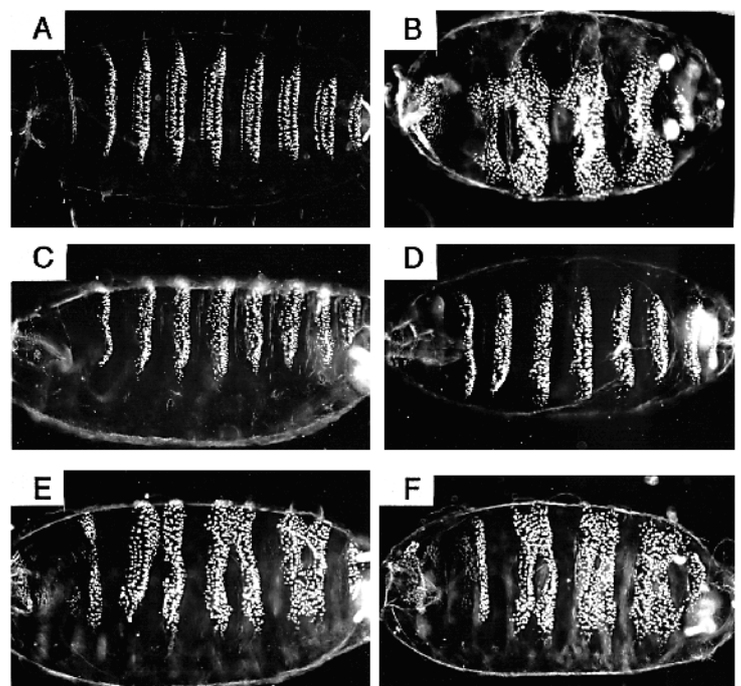


Fig. 7. *eh1* contributes to normal *en* function in embryos. The function of *eh1* in the context of wild-type *en* protein was tested using a partial rescue assay. Transgenes expressing either normal *en* (C,D) or the point-mutated derivative *enF*→*E* (E,F) in the normal *en* domain (see text) were crossed into a strong *en* mutant (*CXI*) background. Cuticles were prepared from unhatched eggs 40 hours after egg laying (normal embryos hatch after about 24 hours). Mutant embryos were identified by their characteristic phenotype, which is partially but not completely rescued. (A) A normal cuticle pattern of 3 thoracic and 8 abdominal denticle bands arrayed along the ventral surface. (B) The *en* mutant has pair-wise fusions of these denticle bands. Several transgenic lines expressing wild-type *en* showed a similar range of phenotypic rescue (average rescue in two independent lines homozygous for the rescue transgene is shown in C,D), characterized by a restoration of naked cuticle between fused bands, as well as near-normal denticle patterns within the bands (row 1 denticles are rarely rescued). In contrast, lines expressing *enF*→*E* showed only very weak rescue, with the range of phenotypes (E,F) apparently overlapping that seen in mutant-only embryos, based on the relative numbers of the various phenotypes seen in collections from heterozygous rescue lines (e.g. *en^{CXI} / CyO*; [*P(enF*→*E)*] / *TM3-Sb*, compared with *en^{CXI} / CyO*; *TM3-Sb* / D). Each panel is a ventral or ventrolateral view, with anterior to the left.

en is known to affect its own expression and expression of other downstream genes, including *hedgehog* (in *en*-expressing cells) and *wingless* (in the adjacent cell row), which encode signaling molecules important in the patterning processes that are disrupted in *en* mutants (reviewed in DiNardo et al., 1994). When wild-type *en* protein is expressed in strong *en* mutants using these *cis*-elements, it rescues the severe defects in the trunk region, as indicated by the reappearance of evenly spaced denticle bands on the ventral surface of the larval cuticle (Fig. 7). However, when either the *en*Δ*eh1* deletion mutant or the single point mutant *en*F→E is similarly expressed, very weak rescue is obtained, indicating the importance of *eh1* in the normal function of *en* during embryogenesis. The correlation of *eh1* activity in the context of EFE, where it is required for strong repression activity *in vivo*, to its function in normal patterning processes indicates that active repression is required for the normal functioning of *en* in the embryo.

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Note added in proof

Consistent with our interpretation of $\Delta 5$, which deletes essentially the eh2 homology, as removing a targeting function that interferes with targeting by the ftz HD in EFE, L. T. C. Peltenburg and C. Murre (1996, *EMBO J.* **13**, 3385-3393) have recently shown that the en eh2 region mediates interaction with extradenticle, another HD protein.