Direct downstream targets of proneural activators in the imaginal disc include genes involved in lateral inhibitory signaling

Andrew Singson, Michael W. Leviten, Anne G. Bang, Xuequn Helen Hua, and James W. Posakony

Department of Biology and Center for Molecular Genetics, University of California San Diego, La Jolla, California 92093-0322 USA

In Drosophila imaginal discs, the spatially restricted activities of the achaete (ac) and scute (sc) proteins, which are transcriptional activators of the basic-helix-loop-helix class, define proneural clusters (PNCs) of potential sensory organ precursor (SOP) cells. Here, we report the identification of several genes that are direct downstream targets of ac-sc activation, as judged by the following criteria. The genes are expressed in the PNCs of the wing imaginal disc in an ac-sc-dependent manner; the proximal promoter regions of all of these genes contain one or two high-affinity ac-sc binding sites, which define the novel consensus GCAGGTG(T/G)NNNYY; where tested, these binding sites are required in vivo for PNC expression of promoter-reporter fusion genes. Interestingly, these ac-sc target genes, including Bearded, Enhancer of split m7, Enhancer of split m8, and scabrous, are all known or believed to function in the selection of a single SOP from each PNC, a process mediated by inhibitory cell–cell interactions. Thus, one of the earliest steps in adult peripheral neurogenesis is the direct activation by proneural proteins of genes involved in restricting the expression of the SOP cell fate.

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The body surface of the adult fly is covered with a complex array of multicellular sensory organs arranged in a largely invariant spatial pattern. In most cases, the cells comprising each of these organs are the progeny of a single sensory organ precursor (SOP) cell (Hartenstein and Posakony 1989). SOPs are determined during the late larval and early pupal stages within undifferentiated epithelial sheets, the imaginal discs and histoblast nests, that ultimately give rise to the cuticular structures of the adult. The process of establishing the pattern of cells committed to the SOP fate, and thus the pattern of sensory organs, consists of at least two major steps. First, the competence to acquire the SOP cell fate is conferred on small groups of cells by the spatially restricted expression and activity of certain proneural genes, which encode transcriptional activators of the basic-helix-loop–helix (bHLH) class. Then, within each of these proneural clusters, all but one of the competent cells are inhibited from expressing the SOP fate by local cell–cell interactions that depend on the activity of a second set of genes, the neurogenic genes. Following its stable determination, the SOP cell executes an invariant lineage that generates the component cells of the sensory organ (Hartenstein and Posakony 1989), a process that also requires neurogenic gene function (Posakony 1994).

The proneural genes include achaete (ac), scute (sc), and daughterless (da), which are required for the development of the external sensory organs of the adult. The spatial pattern of ac and sc expression is largely responsible for defining the pattern of proneural clusters for these organs (Cubas et al. 1991; Skeath and Carroll 1991), whereas da is expressed ubiquitously (Cronmiller and Cummings 1993). In a similar way, the expression of atonal (ato), a recently identified proneural gene that controls the development of chordotonal organs and photoreceptors, defines a distinct pattern of potential chordotonal SOPs and R8 photoreceptor precursors (Jarman et al. 1993, 1994).

In vitro, the ac, sc, and ato bHLH proteins each form hetero-oligomeric complexes with da that are active in sequence-specific DNA binding, and the da protein alone also exhibits DNA-binding activity (Cabrera and Alonso 1991; Van Doren et al. 1991; Jarman et al. 1993). Like other bHLH activators, these proneural proteins recognize specific sites that include a common core sequence
of Bearded (Brd) yield a highly penetrant bristle “tufting” phenotype that results from the commitment of multiple cells within the proneural cluster to the SOP fate (M.W. Leviten and J.W. Posakony, in prep.). We have found recently that the predicted protein products of Brd and E(spl)m4 share sequence and structural similarity, suggesting that they have related or overlapping functions (M.W. Leviten and J.W. Posakony, in prep.). Finally, the scabrous (sca) gene, which encodes a secreted protein (Baker et al. 1990), is expressed in the proneural clusters of the wing imaginal disc and is required for normal singularization of some adult bristles (Mlodzik et al. 1990; Lindsay and Zimm 1992).

Several members of the neurogenic gene family, including Notch (Cagan and Ready 1989), Delta (Parody and Muskavitch 1993), and E(spl) (Knust et al. 1987a; Tietze et al. 1992), are known to have important functions during the development of the compound eye of Drosophila, indicating that this process involves inhibitory cell–cell signaling events similar to those that take place during external sensory organ development. In both systems, these inhibitory interactions serve to restrict the expression of particular cell fates among two or more competent cells. In addition, sca is required nonautonomously within the R8 photoreceptor cell of most ommatidia to inhibit neighboring retinal cells from adopting this fate (Baker et al. 1990). Cells just anterior to the morphogenetic furrow of the eye imaginal disc appear to be part of a proneural territory that is analogous to the proneural clusters of the wing disc. ac and sc are not expressed in the early developing retina, and it is clear that they do not act as the proneural regulators for photoreceptors. Instead, this function is supplied by ato, which is expressed in a stripe on the anterior edge of the morphogenetic furrow (Jarman et al. 1994).

In this paper we provide evidence that the proneural proteins ac and sc directly activate several genes in the proneural clusters of the wing imaginal disc that are known or thought to play a role in the cell–cell signaling process termed lateral or mutual inhibition, by which the expression of the SOP fate is restricted to single cells. We show that these target genes are expressed in a proneural cluster pattern in the wing disc in an ac-sc-dependent fashion; that the proximal promoter regions of these genes contain one or more high-affinity binding sites for da/ac, da/sc, and da/da protein complexes; that these binding sites define a novel extended E-box consensus sequence; and that these binding sites are essential in vivo for the expression of reporter genes in proneural clusters.

Results

ac-sc activity is required for the expression of multiple genes in imaginal disc proneural clusters

Using in situ hybridization to imaginal discs of late third-instar larvae, we investigated the spatial patterns of expression of a number of genes that are known or are likely to play a role in adult sensory organ development. As shown in Figure 1C,G,K,O, we have identified four
Figure 1. Spatial distribution of sc and ac target gene transcripts in wing and eye-antennal imaginal discs of late third-instar larvae. Whole-mount preparations were hybridized in situ with digoxygenin-labeled antisense RNA probes for sc [A,B], Brd [C–F], E(spl)m7 [G–I], E(spl)m8 [K–N], E(spl)m4 [O–R], and sco [S–V]. Wing discs are shown with the presumptive anterior wing margin to the left, and the future ventral wing surface at the top. Eye-antennal discs are shown with anterior at the top. Pattern of accumulation of sc transcripts in wild-type wing [A] and eye-antennal [B] imaginal discs is shown for comparison. Expression of the indicated genes in external sensory organ proneural clusters of the wing disc [C,G,K,O,S] is abolished in discs lacking ac and sc function [sc^+/Y, D,H,L,P,T]. In contrast, expression of these genes in the antennal disc and in the retinal field of the eye disc [E,I,M,Q,U] appears unaffected in discs lacking ac and sc function [F,J,N,R,V]. Arrows indicate the positions of clusters of cells that accumulate E(spl)m4 transcripts in the head capsule portion of wild-type eye discs [Q] but not of eye discs lacking ac and sc function [R], these appear to be precursors to the vibrissae.
genes \([Brd, E(spl)m4, E(spl)m7, \text{and } E(spl)m8]\), the transcripts of which accumulate in the wing disc in a pattern very similar to that of the proneural regulators \(ac\) and \(sc\) [Fig. 1A], the localized expression and activity of which define the proneural clusters of potential SOP cells (Cubas et al. 1991; Skeath and Carroll 1991). Hinz et al. (1994) have also reported the wing disc expression pattern of \(E(spl)m8\). In addition, Mlodzik et al. (1990) have shown previously that a fifth gene, \(sca\), is also expressed in this same proneural cluster pattern [Fig. 1S].

All five of these genes are also expressed in the eye–antennal imaginal disc [Fig. 1E,I,M,Q,U; Baker et al. 1990; Mlodzik et al. 1990]. The most notable feature of this expression is a stripe of accumulated transcript in the vicinity of the morphogenetic furrow of the eye disc. The exact pattern of retinal field expression is distinctive for each gene [Fig. 1] and will not be analyzed in detail here. Nevertheless, the presence of their transcripts near the furrow is consistent with the known or suspected roles of these genes in early cell fate decisions in ommatidial development, particularly those involving cell–cell interaction [see introductory section].

The similarity of the wing disc expression patterns of these five genes to those of the \(ac\) and \(sc\) proteins, which are known to function as transcriptional activators in auto- and cross-regulation in proneural clusters [Van Doren et al. 1992; Martinez et al. 1993], suggested that the former might be direct or indirect targets of \(ac-sc\) regulation. We tested the \(ac-sc\) dependence of the expression of these candidate target genes by examining their pattern of transcript accumulation in imaginal discs from larvae carrying the \(sc^{10-1}\) mutation, which inactivates both \(ac\) and \(sc\). Figure 1D,H,L,P,T shows that, for all five of these genes, the proneural cluster pattern of transcript distribution in wing discs is lost in \(sc^{10-1}\) discs. This result indicates that \(ac-sc\) activity is required to activate the normal expression of these genes in wing disc proneural clusters. In contrast, the patterns and levels of transcript accumulation in the retinal field and antenna appear unaffected in the \(sc^{10-1}\) background [Fig. 1F,J,N,R,V], indicating that transcriptional activation of the five genes in these territories is under \(ac-sc\)-independent control. We also find that certain of the proneural clusters at the margin of the eye disc express these genes under \(ac-sc\) control, whereas expression in other clusters is \(ac-sc\)-independent [Fig. 1Q,R].

Proneural protein complexes bind in vitro to specific sites in the proximal promoter regions of putative \(ac-sc\) target genes

The genetic results presented thus far demonstrate that \(ac-sc\) activity is required for the correct expression of \(Brd, E(spl)m4, E(spl)m7, E(spl)m8, \text{and } sca\) in proneural clusters of the wing imaginal disc. However, these experiments do not address the question of whether \(ac\) and \(sc\) are direct transcriptional activators of these genes. We first approached this problem by attempting to identify, in the upstream regions of the candidate target genes, specific in vitro binding sites for the proneural bHLH proteins.

The specific binding sites that have been identified previously for bHLH activator proteins include a common core sequence \(\text{CANNTG}\) called the E box. In particular, a subset of E-box sequences conforming to the more restricted consensus \(\text{CAG(G/C)TG}\) represent high-affinity in vitro binding sites for da/da, da/ac, and da/sc protein complexes [Cabrera and Alonso 1991; Van Doren et al. 1991; Jarman et al. 1993] and, where tested, these sites function in vivo to mediate transcriptional activation by da, ac, and sc (Cabrera and Alonso 1991; Van Doren et al. 1992; Martinez et al. 1993). We inspected the 5′-flanking sequences of the five candidate target genes for occurrences of the E-box consensus. Table 1 shows that the proximal upstream regions of all of these genes contain either one \([Brd, E(spl)m8, sca]\) or two \([E(spl)m4, E(spl)m7]\) E-box sequences with a GG or GC core, as well as one or more other E boxes (Fig. 2). Interestingly, with the exception of the \(E(spl)m7\) gene, the GG/GC-core E boxes are the most proximal to the transcription start site, even in genes (such as \(Brd\)) with multiple upstream E boxes (Fig. 2).

We tested the ability of the proneural \(da, ac, \) and \(sc\) proteins, individually and in pairwise combination, to bind to a selected subset of these sites in vitro, using purified full-length glutathione S-transferase (GST) fusion proteins in an electrophoretic mobility shift assay (EMSA) with labeled E-box-containing oligonucleotides. The results are summarized in Table 1. Figure 3A shows that all of the probes containing E boxes with GG \([Brd \text{ E1}, E(spl)m7 \text{ E2 and E3}, sca \text{ E1}]\) or GC \([E(spl)m8 \text{ E1}]\) cores are strongly bound in vitro by the da+ac and da+sc combinations and less well by the da protein alone. In contrast, very little binding is detectable in this assay with ac alone, sc alone, or the ac+sc combination, and no binding is observed with the GST protein alone. Shorter autoradiographic exposures of the gels shown in Figure 3A [data not shown] indicate that the major shifted complex observed with the da+ac and da+sc combinations has a greater mobility than the complexes observed with da alone, consistent with the lower relative molecular masses of ac and sc compared with da. We interpret this to mean that da/ac and da/sc hetero-oligomeric complexes are principally responsible for the DNA-binding activity observed with these protein combinations (Cabrera and Alonso 1991; Van Doren et al. 1991). Similarly, the mobility of the complexes formed weakly on the \(E(spl)m8\) E1 probe by sc and the ac+sc combination is consistent with binding by sc homo-oligomers and ac/sc hetero-oligomers (Jiang and Levine 1993).

Both direct binding and competition assays demonstrate that the strong binding activities of the proneural
Table 1. Alignment of E-box sequences in ac-sc target gene promoters and summary of in vitro DNA-binding and competition experiments

| E box | Sequence                  | Binding | Competition |
|-------|---------------------------|---------|-------------|
|       |                           | WT mut  | WT mut      |
| Brd E1'(-88) | CCAGGGTGGT              | ++++  | ++++ -      |
| E2'(-188)  | TCCAGGGTGC              | +      | +++ -       |
| E3'(-490)  | AGAGGGTGGT              | ++     | +           |
| E4'(-672)  | CAGAGGGTGT              |        | ++          |
| E5'(-834)  | ACGAGGGTGA              |        |             |
| E6'(-877)  | GACAGGTGTA              |        |             |
| E7'(-955)  | GCCAGGTGGA              |        |             |
| E8'(-999)  | ACCAGGTGGA              |        |             |
| E9'(-1039) | GAGAGGGTGT              |        |             |
| E10'(-1117)| GACAGGGTCT              |        |             |
| E(spl)m7 E1(-41) | GCAGGGTGCA          |        |             |
| E2(-128)   | CCAGGGTGGT              | ++++  | +++ -       |
| E3(-256)   | AGAGGGTGGT              | ++     | +           |
| E4(-342)   | CAGAGGGTGT              |        | ++          |
| E(spl)m8 E1(-138) | TCCAGGGTGC          | ++++  | +++ -       |
| E2(-462)   | AGAGGGTGGT              | ++     | +           |
| E(spl)m4 E1(-162) | GCAGGGTGTT         | ++++  | +++ -       |
| E2(-250)   | ACCAGGTGGA              |        |             |
| E3(-492)   | GACAGGTGGA              |        |             |
| sca E1(-174) | GCAGGGTGGC              | ++++  | ++ + + +    |
| E2(-416)   | GACAGGGTGCT             |        |             |
| E3(-470)   | GACAGGGTGCT             |        |             |
| E4(-578)   | AGAGGGTGGT              |        |             |

All E-box sequences (CANNNTG, bold type) found in the sequenced upstream regions of the indicated genes are listed, along with two flanking nucleotides on each side. E boxes are numbered in order of increasing distance upstream of the transcription start site (negative numbers, in bp; see Fig. 2), sequences read from the antisense strand are indicated by a prime ('). The two core nucleotides of each E box are underlined, E boxes with a GG or GC core are given bold italic designations (e.g., Brd E1). The relative strength of da/ac and da/sc binding to each wild-type (CANNNTG) or mutant (AANNNTT) site tested, as well as the ability of each tested site to compete for binding to the wild-type Brd E1 site, is summarized from Fig. 3. (++) no detectable binding or no ability to compete specifically for binding to Brd E1. Sequence data for E(spl)m8 are from Klambt et al. (1989); data for sca are from Mlodzik et al. (1990). Nucleotide sequences of the 5'-flanking regions of Brd, E(spl)m7, and E(spl)m4 [this paper] have been deposited in the GenBank data base.

Proximal promoter fragments of the Brd and E(spl)m7 genes direct ac-sc-dependent expression of a reporter gene in a proneural cluster pattern

The results presented in Figure 3 establish that four of the five candidate target genes contain, in positions close to their transcription start sites, specific high-affinity in vitro binding sites for proneural proteins. This finding suggested that proximal promoter fragments from these genes might be sufficient to direct ac-sc-dependent ex-
expression in imaginal discs. We tested this possibility by establishing lines of transgenic flies carrying one of three fusion genes, in which the *Escherichia coli* lacZ gene is placed under the control of a 1.5-kb promoter fragment from *Brd* (-1449 → +42; Fig. 2), a 0.19-kb *Brd* fragment (-144 → +42; Fig. 2), or a 0.5-kb promoter fragment from *E(spl)m7* (-478 → +65; Fig. 2). Both of the *Brd* constructs include the high-affinity E1 proneural protein binding site, whereas the *E(spl)m7* construct includes both the E2 and E3 high-affinity sites.

Figure 5 (A,G,J) shows that all three of these constructs are expressed in a proneural cluster pattern in the late third-instar wing disc. The similarity of the expression patterns of the *Brd* 1.5-kb and 0.19-kb constructs is striking and indicates that a promoter fragment containing only the high-affinity *Brd* E1-binding site is sufficient to direct reporter gene expression in the proneural cluster pattern, though other sequences within this fragment may be required for this activity. All three of the promoter–reporter fusion genes exhibit a higher level of expression in the presumptive SOP than in the remaining cells of the proneural cluster, consistent with the higher level of ac and sc protein in SOPs (Skeath and Carroll 1991). While this behavior is clearly observed for the endogenous *sca* gene (Mlodzik et al. 1990), it is not certain from our in situ hybridization data whether the other genes are normally up-regulated in the SOP. It would perhaps be surprising for the *E(spl)m7* and *m8* genes to be expressed at higher levels in the SOP, given that their wild-type function is to antagonize the SOP cell fate within the proneural cluster. We think it likely that specific mechanisms normally operate to regulate the SOP expression or activity of such inhibitory ac-sc target genes. These might include transcriptional repression (requiring cis-regulatory sites not present in our promoter constructs) (Kramatschek and Campos-Ortega 1994) or post-transcriptional controls on RNA or protein stability (to which *lacZ* mRNA and protein would most likely not be responsive).

The distinct regulation of *Brd* and *E(spl)m7* expression in the antenna and in the retinal field of the eye versus the proneural clusters of the wing disc (Fig. 1) is emphasized further by the activities of the promoter–reporter constructs in the eye–antennal disc. The *Brd* 1.5-kb promoter fragment reproducibly drives a high level of lacZ expression near, and posterior to, the morphogenetic furrow (Fig. 5D). This pattern mimics that of the endogenous *Brd* transcript (Fig. 1E). In contrast, only a fraction of the *Brd* 0.19-kb or the *E(spl)m7* promoter fusion lines exhibit any detectable expression in the retinal field, and this is very weak by comparison (data not shown). This result implies that the cis-regulatory elements that direct expression of *Brd* and *E(spl)m7* in the retina are at least partially distinct from those that control the activation of these genes in wing disc proneural clusters.

We tested whether the activity of the three promoter–reporter fusion genes is dependent on the activity of the proneural regulators ac and sc by introducing them into the *sc10-1* background. As shown in Figure 5 (B,H,K), lacZ expression from these constructs in wing disc proneural clusters is reduced severely or abolished in the absence of *ac-sc* function. In contrast, the *Brd* 1.5-kb promoter remains active at apparently normal levels in the retinal field and antenna of *sc10-1* eye–antennal discs (Fig. 5E). Thus, the pattern of *ac-sc* dependency exhibited by the expression of the endogenous genes (Fig. 1C–J) is mimicked by the *Brd* and *E(spl)m7* promoter–lacZ fusion genes.

**Mutation of binding sites for proneural proteins abolishes activation of the *Brd* and *E(spl)m7* promoters in proneural clusters**

Because proximal promoter fragments from *Brd* and *E(spl)m7* are capable of directing *ac-sc*-dependent expression in wing disc proneural clusters, a critical question was whether the da/ac/sc-binding sites that we had identified in these promoters are necessary for this ac-
Figure 3. Electrophoretic mobility shift assays of da, ac, and sc binding to E box sequences in the proximal promoters of candidate target genes. (A) Oligonucleotide probes containing E boxes that conform to the consensus CAG(G/C)TG (GG- or GC-core E boxes; see Table 1) are bound efficiently by da/ac and da/sc hetero-oligomeric complexes, as well as by da homo-oligomers. ac alone, sc alone, and the ac + sc combination fail to show significant binding activity with these probes. (B) Relative to the Brd E1 probe, oligonucleotide probes containing E-box sequences with other than GG or GC cores (see Table 1) are bound much more weakly or not at all by da, ac, and sc, either alone or in pairwise combination. (C) Binding of proneural protein complexes to oligonucleotide probes containing GG- or GC-core E boxes requires the integrity of the E-box sequence. Mutation of the E box (CAGNTG to AANNTT) completely abolishes binding of da/ac, da/sc, and da/da protein complexes. All individual and pairwise combinations of the da, ac, and sc proteins were tested, though results for some combinations are not shown for the E(spl)m7 E2, E(spl)m7 E3, E(spl)m8 E1, and sca E1 sites. (D) Binding of proneural proteins to the Brd E1 probe is efficiently competed by probes containing strong binding sites (see A), but poorly or not at all by probes containing weaker or mutant binding sites (see B, C). Binding and competition activity of weaker sites may correlate with their similarity to high-affinity sites such as Brd E1 [e.g., the Brd E2 site represents only a 1-bp change from Brd E1 (GCACGTGT vs. GCAGGTGT)]. Competition by weaker binding sites was observed by using a 200-fold excess of competitor and poly[d(A,T)] as the nonspecific competitor (see Materials and methods).

tivity. We constructed three promoter-reporter fusion genes identical to those described above but including 2-bp mutations that change the Brd E1 and E(spl)m7 E2 and E3 E-box sequences from CAGGTG to CCGGTT. Figure 5 (C, I, L) shows that the mutant promoters fail to be activated in the proneural clusters of the third-instar wing imaginal disc, though the Brd 1.5-kb promoter fragment remains capable of directing apparently normal levels of expression in the antenna and retinal field of the eye-antennal disc (Fig. 5F). This result demonstrates
that the high-affinity proneural protein binding sites in the Brd and E(spl)m7 promoters are specifically required for the normal activation of these promoters in wing disc proneural clusters.

Discussion

Identification of direct downstream targets of ac and sc in wing disc proneural clusters

The evidence presented in this paper strongly supports the conclusion that the Brd, sca, E(spl)m4, E(spl)m7, and E(spl)m8 genes are directly activated in proneural clusters of the late third-instar wing imaginal disc by protein complexes that include the ac and sc bHLH proteins. Transcripts of these genes accumulate in a spatial pattern in the wing disc that is very similar to the proneural cluster pattern of ac and sc activity, and their expression is reduced severely or abolished in ac− sc− mutant discs (Fig. 1). The proximal promoter regions of all of these genes contain one or two specific, high-affinity binding sites for the proneural proteins da, ac, and sc (Figs. 2 and 3; Table 1). Fragments of the Brd and E(spl)m7 promoters that include these sites are sufficient to direct ac-sc-dependent expression of a reporter gene in wing disc proneural clusters (Fig. 5). Finally, mutation of the high-affinity da/ac(sc) binding sites in these two promoters reduces severely or abolishes reporter gene activity in proneural clusters (Fig. 5).

These data do not strictly rule out the possibility that another transcription factor, the expression or activity of which is ac-sc-dependent, is actually the direct activator of these genes in vivo and that ac and sc act indirectly. The activity of the hypothetical factor would be dependent on the proneural protein-binding sites that we have shown are required in vivo for the activities of the Brd and E(spl)m7 promoters. For example, ac and sc could control the expression of a factor that potentiates transcriptional activation by da, so that da alone would be the direct activator. However, we believe the simplest interpretation of the evidence is that ac and sc participate directly in transcriptional activation of the genes that we have studied.

Previous experiments indicate that ac and sc are highly active in DNA binding in vitro and in transcriptional activation in yeast and tissue culture cells only in the presence of da (Cabrera and Alonso 1991; Van Doren et al. 1991, 1992) or a non-bHLH partner such as dorsal (Gonzalez-Crespo and Levine 1993; Jiang and Levine 1993). These findings, along with the demonstration that da protein is apparently present in all cells of the wing imaginal disc (Cronmiller and Cummings 1993), motivate our working hypothesis that da/ac and da/sc heterooligomers are the active complexes in mediating the transcriptional regulatory functions of ac and sc in imaginal disc proneural clusters. However, it is possible that post-translational modifications (for example) allow ac and sc to activate transcription in vivo independently of da or other factors.

Proximal promoter regions of ac-sc target genes contain high-affinity binding sites with a characteristic extended E-box consensus

We have found that the proximal upstream regions of ac-sc target genes contain either one or two E-box sites to which ac and sc, in combination with da, bind with high affinity in vitro; the da protein alone also binds to these sequences. Three features of the binding sites are noteworthy. The first is their location. In all six of the ac-sc proneural cluster target genes we have defined (including ac itself), a high-affinity binding site is found relatively close to the TATA box and transcription start site [between −58 (ac) and −174 (sc)]. With one exception [E(spl)m7], the high-affinity site or sites are the most proximal E box (CANNTG) consensus sequences found in the promoter. Second, all of the high-affinity binding sites belong to a subset of E boxes described by the consensus ACAG[C/T]G; E boxes with other than a GG or GC core are bound much more weakly by da, ac,
Figure 5. Promoter–reporter fusion genes require da/ac/sc binding sites for expression in external sensory organ proneural clusters. All panels show imaginal discs from late third-instar larvae stained for β-galactosidase activity. Brd promoter fragments of 1.5 or 0.19 kb and an Esplm7 promoter fragment of 0.5 kb [see Fig. 2] were fused to the E. coli lacZ gene in the CaSpeRlacZ P-element transformation vector. Lines of transgenic flies carrying either the wild-type reporter constructs [P[Brd 1.5-lacZ], P[Brd 0.19–lacZ], and P[m7 0.5–lacZ]], or the same constructs with their high-affinity da/ac/sc binding sites mutated [P[Brd 1.5M–LacZ], P[Brd 0.19M–LacZ], and P[m7 0.5M–LacZ]], E-box sequences changed from CAGGTG to CCGGTT, were established.

The wild-type constructs are expressed in a proneural cluster pattern in the wing disc: (A) P[Brd 1.5-lacZ]-1 [4 of 4 lines], (G) P[Brd 0.19–lacZ]-10 [10 of 15 lines], and (J) P[m7 0.5–lacZ]-6 [3 of 7 lines]. The proneural cluster expression of these wild-type reporter genes is abolished in discs lacking the function of the endogenous ac and sc genes: (B) sc10Y, P[Brd 1.5–lacZ]-1/P[Brd 1.5-lacZ]-1, (H) sc10Y, P[Brd 0.19–lacZ]-10/P[Brd 0.19–lacZ]-10, and (K) sc10Y, P[m7 0.5–lacZ]-6/P[m7 0.5–lacZ]-6. The same lack of proneural cluster expression is observed in wing discs from transgenic animals carrying the mutant promoter–reporter constructs: (C) P[Brd 1.5M–lacZ]-8 [7 of 8 lines], (D) P[Brd 1.5M–lacZ]-8 [7 of 8 lines], (E) P[Brd 0.19M–lacZ]-4 [10 of 10 lines], and (L) P[m7 0.5M–lacZ]-10 [15 of 15 lines]. In contrast, construct- or insertion-specific ectopic expression is unaffected by the sc10Y background or by mutation of proneural protein-binding sites [e.g., A–C], and provides a control for the staining reaction. Of the three promoter fragments tested, only the 1.5-kb Brd fragment (D: P[Brd 1.5-lacZ]-1) exhibits consistent expression in the retinal field of the eye–antennal disc [4 of 4 lines]. This expression pattern is not detectably altered in discs lacking ac and sc function [E: sc10Y, P[Brd 1.5–lacZ]-1/P[Brd 1.5–lacZ]-1] or in wild-type discs carrying the 1.5-kb Brd promoter fragment with the high affinity da/ac-sc DNA-binding site mutated [F: P[Brd 1.5M–lacZ]-8]. β-Galactosidase staining in the retinal fields of these discs (D–F) extends more posteriorly than transcript accumulation from the endogenous Brd gene [cf. Fig. 1E]. This is likely because of the perdurance of β-galactosidase protein: When reporter gene expression is assayed using a digoxigenin-labeled antisense probe for lacZ RNA [inset in D], the observed pattern of transcript accumulation is comparable to that of endogenous Brd RNA [cf. inset in D with Fig. 1E].
and sc in vitro (Van Doren et al. 1991; Jarman et al. 1993). Finally, an alignment of the most proximal high-affinity da/ac/sc binding sites in these genes defines a novel extended E-box consensus sequence, GCAG-GTG{T/G}NNNY. In genes with more than one high-affinity da/ac/sc binding site—E(spl)m4, E(spl)m7, and ac—the site nearest the transcription start fits this consensus, in E(spl)m7, both sites match. We propose the name proximal proneural response element (PPRE) to refer to these particular [i.e., most proximal] sites and to the consensus sequence that they define (Fig. 4). The predictive value of this consensus is illustrated by the fact that we did not obtain the sequence of the E(spl)m4 upstream region until after the consensus had been defined by in vitro DNA-binding studies with the other promoters.

The similarity in both location and sequence of functional da/ac/sc binding sites in such a diverse set of target genes is very likely to be of regulatory significance. It is possible that the proximity of these high-affinity sites to the TATA box serves to maximize the activating capacity of proneural protein complexes bound to them. In this case, the PPRE may not function effectively at great distances (>500 bp) from the promoter. Nevertheless, it is clear that PPREs can function in vivo in either orientation with respect to the direction of transcription [cf. Brd E1 and E(spl)m7 E2/E3]. The extended sequence restriction of the PPRE on either side of the E-box hexamer may help to confer greater specificity on the activation of these target genes by proneural proteins.

Distinct cis and trans control of ac-sc target gene expression in proneural clusters and in the retina

The results presented here indicate clearly that both the cis-regulatory elements and the trans-activators that control expression of ac-sc target genes in late third-instar wing disc proneural clusters are at least partially distinct from those that control their expression in the antenna and retinal field of the eye–antenna disc at the same stage. First, ac and sc are not detectably expressed in the late third-instar retina (Fig. 1B), and we have shown here that the expression of several proneural cluster target genes in the vicinity of the morphogenetic furrow is correspondingly independent of ac-sc function [Fig. 1]. Second, certain proximal promoter fragments that contain high-affinity da/ac/sc binding sites and are capable of directing ac-sc-dependent reporter gene expression in wing disc proneural clusters give only very weak and variable expression in the retinal field of third-instar eye discs [data not shown]. This includes both the Brd 0.19-kb fragment [Fig. 2] and the E(spl)m7 0.5-kb fragment [Fig. 2], as well as a 0.9-kb fragment of the ac promoter that contains three E-box sites required in proneural clusters for ac autoregulation and cross-regulation by sc [Martinez and Modolell 1991; Van Doren et al. 1991, 1992, Martinez et al. 1993]. Finally, mutation of the single high-affinity E-box site (Brd E1) in the Brd 1.5-kb promoter fragment (Fig. 2; Table 1) results in the loss of promoter activity in wing disc proneural clusters [Fig. 5A,B], but apparently does not affect its activation in the retina [Fig. 5C], although we cannot rule out the loss of a small subset of the expression pattern (e.g., a narrow anterior stripe). These last observations demonstrate that high-affinity da/ac/sc binding sites are neither necessary nor sufficient for high-level expression of ac-sc target genes in the retinal field of the third-instar eye disc.

The proneural gene ato is required for the development of both chordotonal organs and photoreceptors (Jarman et al. 1993, 1994). The predicted ato product is a bHLH protein closely related to ac and sc and when combined with da, can bind in vitro to oligonucleotides containing a GG-, a GC-, or a GA-core E-box from the asense gene [Jarman et al. 1993]. Unlike ac and sc, ato is expressed at the anterior edge of the morphogenetic furrow in the third-instar eye disc and in a large ring-shaped territory in the antenna disc [Jarman et al. 1993, 1994]. In addition, it is expressed in two small clusters of cells in the nonretinal (head capsule) portion of the eye disc. The pattern of ato transcript accumulation in these regions is quite similar to the distribution of ac-sc-independent transcripts from the ac-sc target genes that we have defined here, particularly E(spl)m4 and E(spl)m7 [Fig. 1]. Moreover, ato activity is required for the expression of sca protein in the retina [Jarman et al. 1994]. These observations raise the possibility that ato might function as a direct transcriptional activator of these genes in the retina, in Johnston’s organ, and in chordotonal organs of the head capsule. If this is the case, our results suggest that ato may function through promoter-binding sites distinct from those utilized by ac and sc in proneural clusters, or that an additional factor is required for its activation of genes in the retina and the antenna.

The proneural regulators ac and sc activate antagonists of the SOP cell fate in proneural clusters

One of the most important findings of our study concerns the function of the genes that are directly activated in the proneural clusters of the wing disc by the proneural regulators ac and sc. With the possible exception of E(spl)m4, genetic evidence indicates that all of the genes we have studied play a role in the process by which expression of the SOP cell fate is inhibited in all but one cell of the proneural cluster [see introductory section]. This observation has important implications for our understanding of pattern formation in adult sensory organ development. It appears that ac and sc have two oppositely directed activities with respect to neural cell fate determination. First, they promote sensory organ development by conferring on groups of imaginal disc cells the potential to become SOPs, presumably by activating (directly or indirectly) genes required for this fate, particularly within the SOP itself. This first activity constitutes the genetically defined proneural function of ac and sc, and includes direct auto- and cross-regulation within the proneural clusters and perhaps in the SOPs [Van Doren et al. 1991, 1992, Martinez et al. 1993]. Second, ac and sc
activate, within the proneural clusters and within SOPs, other genes that act instead to antagonize the SOP fate, apparently as part of the cell—cell signaling process referred to as lateral or mutual inhibition. These diverse regulatory functions of ac and sc are summarized in Figure 6. We conclude that ac and sc are not strictly proneural genes but, instead, have a complex role in establishing local conditions of gene expression in the imaginal disc epithelium that lead to the stable determination of individual cells as SOPs.

Consistent with our results, ectopic activation of genes involved in lateral inhibition has been observed previously to accompany the ectopic expression of lethal of scute, a proneural gene not normally active in imaginal discs [Hinz et al. 1994]. In addition, Kramatschek and Campos-Ortega [1994] recently reported evidence that proneural genes function to activate the expression of the E(spl)m5 and m8 genes in the ventral neuroectoderm of the embryo, from which the larval CNS arises. It is unclear whether this activation in the embryo is direct. Although we have shown here that E(spl)m8 E1 is the only high-affinity E-box-binding site for proneural proteins in the E(spl)m5 proximal promoter, Kramatschek and Campos-Ortega [1994] find that mutation of this site only slightly reduces reporter gene expression in the neuroectoderm and does not mimic the strong qualitative effect of loss-of-function mutations in proneural genes. Nevertheless, it seems likely that proneural gene-dependent regulation of certain genes involved in lateral inhibition is a common element of neurogenesis in both the embryonic ventral ectoderm and the imaginal discs.

The observations reported here offer an important addition to our understanding of the proneural cluster, the group of cells from which the single SOP is selected. Many of the genes required for the process of lateral inhibition, including Notch [Fehon et al. 1991], Delta [Kooh et al. 1993], Suppressor of Hairless [Schweisguth and Posakony 1992], Hairless [Bang and Posakony 1992], and groucho [Delidakis et al. 1991], A.G. Bang and J.W. Posakony, unpubl.), are broadly or ubiquitously expressed in third-instar larval imaginal discs, including the territories in which proneural clusters arise. Thus, these and other elements of the inhibitory signaling machinery are available to imaginal disc cells independently of their proneural potential. However, our results indicate that only in the proneural clusters—that is, only in those cells in which ac and sc are active as transcriptional activators—will the complete ensemble of gene functions required to stably determine a single SOP be expressed.

Materials and methods

Drosophila stocks

Flies were raised on standard yeast/cornmeal/molasses/agar medium. Transformant lines carrying promoter-reporter gene fusions [Brd 1.5–lacZ, Brd 1.5M–lacZ, Brd 0.19–lacZ, Brd 0.19M–lacZ, E(spl)m7 0.5–lacZ, and E(spl)m7 0.5M–lacZ] were characterized as homozygous stocks, except where the chromosome carrying the transgene insertion was homozygous lethal. All other mutations and chromosomes are described in Lindsley and Zimm [1992].

General molecular biology procedures

Molecular biology techniques not described in detail below are described in Ausubel et al. [1987] and Sambrook et al. [1989].

In situ hybridization

In situ hybridization was performed on imaginal discs dissected from late third-instar larvae as described by Tautz and Pfeifle [1989], with modifications by Jiang et al. [1991] and Schweisguth and Posakony [1992]. Digoxigenin-labeled antisense RNA probes for sc, Brd, E(spl)m7, E(spl)m8, E(spl)m4, sea, and lacZ were prepared as described by the manufacturer [Boehringer Mannheim].

β-Galactosidase activity staining

Imaginal discs dissected from late third-instar larvae were stained for β-galactosidase activity as described by Romani et al. [1989].

Isolation and sequencing of promoter fragments

Brd

A 1491-bp EcoRI–SalI Brd promoter fragment containing 42 bp of 5′-untranslated sequence was isolated from a Lambda FIX II [Stratagene] genomic DNA clone [M.W. Leviten and J.W. Posakony, in prep.], and subcloned into the XbaI site of pBluescript KS[+][Stratagene] using XbaI linkers, to create pBSBrd-1.5.

E(spl)m7

A 543-bp XbaI–XmnI E(spl)m7 promoter fragment containing 65 bp of 5′-untranslated sequence was isolated from a 1.6-kb genomic DNA clone (provided by D. Kosman, R. Park, and M. Levine, University of California, San Diego) and subcloned into the XbaI site of pBluescript KS[+][p][p] using XbaI linkers, to create pBSm7-0.5.

E(spl)m4

A 450-bp DNA fragment containing the E(spl)m4-coding region
was amplified by the polymerase chain reaction (PCR) from genomic DNA prepared from Canton S flies using the following primers (Operon Technologies): M4A, 5'-CTCTGGAATTCATGTGCCAGAACAAGATC-3' and M4B 5'-CTCTGTCGACTTAGGCCCTGAACCCCAAGC-3'. The PCR product was ligated into EcoRI/Sall-digested pBluescript KS(+) vector. This fragment was used as a probe for isolation of a genomic DNA clone containing the E(spl)m4 gene from an EMBL3 library (kindly provided by R. Blackman, University of Illinois, Urbana). A 2.0- kb Xhol–HindIII fragment from the EMBL3 clone containing 503 bp of promoter sequence was subcloned into pBluescript KS(+) vector.

The nucleotide sequences of all fragments described above were obtained or verified by double-stranded DNA sequencing. Sequences of the 5'-flanking regions of Brd, E(spl)m7, and E(spl)m4 have been deposited in the GenBank data base.

Germ-line transformation

The 1491-bp Brd and the 543-bp E(spl)m7 Xbal-linkered promoter fragments were cloned into the CaSpeRlacz P-element transformation vector described in Margolis et al. (1994) and Van Doren et al. (1992) to create pBrd 1.5- lacZ and pm7 0.5-lacZ, respectively. pBrd 0.19–lacZ was created by digesting pBrd 1.5-lacZ with PstI and self-ligating the vector-containing fragment. The mutant forms of these constructs, pBrd 1.5M-lacZ, pm7 0.5M-lacZ, and pBrd 0.19M-lacZ (see below), were created identically to their wild-type counterparts. Transgenic flies were generated by P-element-mediated germ-line transformation techniques (Rubin and Spradling 1982) using w1118 as the recipient strain.

Site-specific mutagenesis

Site-specific mutagenesis was performed using pBSBrd–1.5, a mutant oligonucleotide, convenient restriction sites, and PCR to generate pBSBrd–1.5M. The high-affinity da/ac-sc binding site Brd E1 (Fig. 2, Table 1) in this plasmid was changed from GCAGGTGTG to GCCGGTTT and verified by DNA sequencing. pBrd 1.5M-lacZ and pBrd 0.19M-lacZ were then constructed as described above. pBSm7–0.5M was generated using a Site-Directed Mutagenesis Kit (Clontech). The two high-affinity da/ac-sc binding sites m7 0.5M-lacZ (Table 1, Table 1) in pBSm7–0.5M were changed from GCAGGTGTG to GCCGGTTT and from GCAGGTGTG to GCCGGTTT, respectively. After sequence verification, pm7 0.5M-lacZ was constructed as described above.

DNA-binding assays

Plasmid constructs for bacterial expression of full-length ac, sc, and da proteins will be described elsewhere [P.A. Powell and J.W. Posakony, in prep.]. GST fusion proteins were isolated from bacterial cultures that had been grown at 37°C to an OD600 of 0.2, shifted to 30°C and grown to an OD600 of 0.6–0.7, and induced with 400 μM final IPTG at 30°C for 1 hr. The cells were washed in PBS and resuspended in lysis buffer [25 mM K-HEPES (pH 7.5), 0.5 M KCl, 12.5 mM MgCl2, 1 mM EDTA, 0.1% NP-40, 10% glycerol, 1 mM DTT, 1 mM PMSE, 2 μg/ml of aprotinin, 2 μg/ml of leupeptin, and 1 μg/ml of pepstatin] to which lysozyme was added to 0.625 mg/ml. Following a 15-min incubation on ice, cells were sonicated [Fisher Dismembrator model 300] for 5×30 sec at 60% power at 30-sec intervals. The supernatant from the lysate was mixed with glutathione–Sepharose beads [Pharmacia] and gently agitated for 30 min at 4°C. The beads containing the fusion proteins were washed 4 times in lysis buffer without glycerol, and then 3 times in elution buffer lacking glutathione [25 mM K-HEPES (pH 8.0), 1 mM DTT, 1 mM PMSE, 2 μg/ml of aprotinin, 2 μg/ml of leupeptin, and 1 μg/ml of pepstatin]. Fusion proteins were eluted from the beads by incubation in elution buffer containing 10 mM glutathione. The beads were then pelleted and the supernatant was dialyzed [25 mM K-HEPES (pH 7.5), 50 mM KCl, 5 mM MgCl2, 0.1 mM EDTA, and 1 mM DTT]. 3×4 liters, overnight at 4°C. Eluted proteins were frozen in aliquots in 10% glycerol. Double-stranded DNA oligonucleotide probes and specific competitor oligonucleotides were synthesized by Operon Technologies. Each probe or competitor corresponds to the sequence surrounding a single wild-type or mutant E-box site (bold letters) found in the proximal promoters of Brd, E(spl)m7, E(spl)m8, or scd [see Fig. 2]. The sequences are as follows:

For electrophoretic mobility shift assays, oligonucleotide probes were end-labeled and annealed as described in Van Doren et al. (1991). Bacterial fusion proteins [0–400 ng] were mixed in a final volume of 6 μl of dialysis buffer plus 10% glycerol and incubated for 20 min at 25°C. After incubation, the following were added: 1 μl of probe [0.025 μM at 1–8×105 cpm/μl] in TE buffer, 1 μl of 1 mg/ml poly[dI⋅dC] or poly[dA⋅dT] (Sigma, 1 μl of 4× dialysis buffer [or for competition experiments, 1 μl of 2.5 μM (100×, Fig. 3D, last nine lanes) or 5 μM (200×, Fig. 3D, first eight lanes) unlabeled competitor oligonucleotide in 4× dialysis buffer] and 1 μl of 0.275 mg/ml denatured salmon sperm DNA. Presumably because of the high GC content of da/ac-sc binding sites, poly[dA⋅dT] was found to be a better nonspecific competitor than poly[dI⋅dC], so the former was used in the low-affinity binding site competition experiment (Fig. 3D, first eight lanes). The mixture was incubated for 40 min at 25°C. The reactions were electrophoresed on prerun 0.5×TBE/4% acrylamide gels (2 mm thick × 15 cm long) for 3 hr at 125 V. Gels were
fixed in 25% methanol/7% acetic acid for 30 min and dried prior to autoradiography on X-Omat AR film (Kodak) at −80°C with an intensifying screen.

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Note added in proof

The GenBank accession numbers for the promoter sequences described in this paper are U13067 (Brd), U13068 (E(Spl)m4), and U13069 (E(spl)m7).

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