Pbx proteins display hexapeptide-dependent cooperative DNA binding with a subset of Hox Proteins

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The human proto-oncogene \textit{PBX1} codes for a homolog of \textit{Drosophila} extradenticle, a divergent homeo domain protein that modulates the developmental and DNA-binding specificity of select HOM proteins. We demonstrate that wild-type Pbx proteins and chimeric E2a–Pbx1 oncoproteins cooperatively bind a consensus DNA probe with HoxB4, B6, and B7 of the Antennapedia class of Hox/HOM proteins. Specificity of Hox–Pbx interactions was suggested by the inability of Pbx proteins to cooperatively bind the synthetic DNA target with HoxA10 or \textit{Drosophila} even-skipped. Site-directed mutagenesis showed that the hexapeptide motif (IYPWMK) upstream of the Hox homeo domain was essential for HoxB6 and B7 to cooperatively bind DNA with Pbx proteins. Engraftment of the HoxB7 hexapeptide onto HoxA10 endowed it with robust cooperative properties, demonstrating a functional role for the highly conserved hexapeptide element as one of the molecular determinants delimiting Hox–Pbx cooperativity. The Pbx homeo domain was necessary but not sufficient for cooperativity, which required conserved amino acids carboxy-terminal of the homeo domain. These findings demonstrate that interactions between Hox and Pbx proteins modulate their DNA-binding properties, suggesting that Pbx and Hox proteins act in parallel as heterotypic complexes to regulate expression of specific subordinate genes.

[Key Words: Pbx; Hox; E2a–Pbx1; homeo domain proteins; proto-oncogene]

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Hox proteins are members of a large family of vertebrate transcription factors that share a similar DNA-binding motif known as the homeo domain (Scott et al. 1989). Hox proteins have many features in common with \textit{Drosophila} homeotic selector (HOM) proteins including primary sequence, chromosomal organization of their respective genes, and regionally and developmentally restricted expression profiles (Akam 1989; McGinnis and Krumlauf 1992). Some Hox proteins have also been shown to function similarly in vivo to their \textit{Drosophila} counterparts. For both Hox and HOM proteins, the homeo domain has been shown to mediate DNA binding in vitro and play an important role in the ability of these proteins to regulate transcription through specific target sequences (Desplan et al. 1988; Hoey and Levine 1988; Müller et al. 1988; Gehring et al. 1994a). The homeo domain has also been shown to make important contributions to the functional specificity of HOM proteins (Kuziora and McGinnis 1989, 1991; Dessain et al. 1992). However, recent data suggest that homeo domain proteins do not achieve their functional specificity on the basis of DNA–protein interactions alone, as best illustrated by studies of \textit{fushi tarazu} (ftz) showing that high affinity homeo domain–DNA interactions are neither sufficient nor necessary for its biological activity (Fitzpatrick et al. 1992; Ananthan et al. 1993; Schier and Gehring 1993).

Several studies suggest that the functional specificity of homeo domain proteins is determined by the combined effects of various individual interactions both within and outside of the homeo domain. Such interactions have been shown experimentally to influence the DNA-binding specificity of homeo domain proteins. For instance, combinatorial interactions between MCM1, MAT\textalpha{}1 and MAT\textalpha{}2 determine the ability of \textalpha{}1 and \textalpha{}2 to recognize specific operators that mark genes regulating mating type in yeast (Johnson 1992). Homeo domain proteins may influence the DNA binding of nonhomeo domain proteins such as serum response factor (SRF), a mammalian protein with similarity to MCM1 (Gruenberg et al. 1992). Protein–protein interactions may also influence the transcriptional properties of homeo domain proteins as shown for HoxD8 inhibition of HoxD9 activation, an effect dependent on contacts within helix

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1 of their homeo domains [Zappavigna et al. 1994]. Studies of the POU and Oct 1 proteins [Stern et al. 1989; Ingraham et al. 1990; Voss et al. 1991; Treacy et al. 1992; Verrijzer et al. 1992, Xue et al. 1993], vHNF1 [Rey-Campos et al. 1991], and Paired class homeo proteins [Wilson et al. 1993] provide additional evidence that cooperative interactions play an important role in regulating their DNA-binding and functional properties.

Previous studies have identified a distinctive subfamily of atypical homeo domain proteins that are potential candidates for interactions with Hox/HOM proteins. These include the mammalian Pbx proteins, Drosophila extradenticle (exd), and Caenorhabditis elegans ceh-20 [Monica et al. 1991; Bürglin and Ruvkun 1992; Rauskolb et al. 1993], notable for their extensive similarities both within and outside of their homeo domains. Pbx1 is a proto-oncogene that was originally discovered at the site of t(1;19) chromosomal translocations in pre-B cell acute leukemias [Kamps et al. 1990; Nourse et al. 1990]. In Drosophila, genetic studies suggest that the Pbx homolog exd serves a unique role in segment identity through parallel interactions with homeotic selector proteins such as ultrabithorax (Ubx) and antennapedia [Antp] (Peifer and Weischaus 1990; Rauskolb and Weischaus 1994). Recent studies support this role by demonstrating cooperative interactions between exd and select HOM proteins (or engrailed) in DNA-binding assays [Chan et al. 1994; VanDijk and Murre 1994] and in a yeast two-hybrid system [Johnson et al. 1995]. These studies are consistent with the thesis that biological specificity of homeotic selector proteins may be achieved, at least in part, through cooperative interactions that affect the manner in which they recognize and regulate downstream subordinate genes.

In this report we demonstrate that the DNA-binding affinity of Pbx proteins is markedly stimulated through interactions with a subset of Hox proteins under conditions where they demonstrate little or no inherent DNA-binding affinity alone. Cooperative DNA binding required the homeo domains and specific flanking amino acids in both the Hox and Pbx proteins. The conserved YPWM amino acid motif upstream of the homeo domain in many Hox proteins was essential for cooperativity, thereby establishing a role for this highly conserved element that may define a subset of Hox proteins capable of cooperative interactions with Pbx proteins. These studies support a model in which Pbx and Hox proteins act in parallel as heterotypic complexes to regulate expression of specific subordinate genes.

**Results**

*Interaction of Pbx1 with HoxB7 in a yeast two-hybrid assay*

A search for proteins with the potential to interact with Pbx was initially conducted in yeast by use of a two-hybrid assay [Harper et al. 1993]. HoxB7 was selected for testing interactions with Pbx1 as it is a cognate homolog of Ubx [Scott et al. 1989] that interacts with exd under comparable conditions [Johnson et al. 1995] and could be stably coexpressed with Pbx1 in yeast cells. The homeo domain and flanking amino acids of HoxB7 were fused to the GAL4 DNA-binding domain in yeast vector pAS2 [construct pAS-ypHDB7] and various portions of Pbx1 were fused to the GAL4 activation domain contained in pACT2 [Fig. 1]. Yeast cells cotransformed with pACT–Pbx1b and pAS–ypHDB7 showed activation of lacZ reporter gene expression under the control of the GAL4 UAS, whereas no β-gal activity was observed in cells transformed with each of the plasmids individually [Fig. 1, data not shown]. The interaction between Pbx1 and HoxB7 required the homeo domain of Pbx1 as its deletion abrogated activation of β-gal expression [Fig. 1]. A minimal portion of Pbx1 consisting of 88 amino acids spanning the homeo domain and immediate carboxy-terminal flanking amino acids was sufficient for interaction with HoxB7. These observations suggested the potential for interaction between Pbx1 and HoxB7 proteins mediated, at least in part, by the homeo domain of Pbx1.

*In vitro interaction of Pbx and Hox proteins is stimulated by the presence of Pbx-binding site DNA*

The potential interaction of Pbx1 with HoxB7 was investigated further by immunoprecipitation analyses of in vitro-synthesized proteins. These experiments employed wild-type Pbx1a with an epitope tag at its amino terminus. In vitro translated Pbx1a–flag and HoxB7 were mixed and subjected to immunoprecipitation analysis with antibodies against the epitope tag. In parallel, the translocation-derived E2a–Pbx1a fusion protein was also tested for potential interaction with HoxB7 with an anti-E2a monoclonal. Precipitation of Pbx1a or E2a–Pbx1a from in vitro translated showed little or no coprecipitated HoxB7 [Fig. 2]. However, when an oligonucleotide (5’-CGAATTCATGATCACTAATTGGAG-3’) containing consensus Hox (Pellerin et al. 1994) and Pbx (VanDijk et al. 1993; LeBrun and Cleary 1994; Lu et al.

### Table 1

| pACT-Pbx Constructs | β-Gal Activity |
|---------------------|----------------|
| pACT                | -              |
| pACT–Pbx1b          | +              |
| pACT–Pbx1C          | +              |
| pACT–Pbx1a3N        | +              |
| pACT–Pbx1HD         | +              |
| pACT–Pbx1N          | -              |
| pACT–Pbx1aC         | -              |
| pACT–Pbx1a3HD       | -              |

*Figure 1.* Interaction of Pbx1 and HoxB7 in a yeast two-hybrid assay. Yeast was cotransformed with pAS–ypHDB7, and the indicated pACT plasmids, and then tested for activation of a lacZ reporter gene. β-Galactosidase (β-gal) activity was determined by use of a filter blot assay in which conversion of the substrate [X-gal] was assessed after incubation for 2 hr at 37°C. Constructs showing no β-gal activity were also negative after 18 hr at 37°C.
Cooperative DNA binding by Hox and Pbx proteins

Protein complexes were specifically precipitated with a monoclonal antibody against an epitope-tagged Pbx1a or a monoclonal anti-E2a (Yae) antibody (denoted below gel lanes). Precipitated proteins were analyzed by SDS-PAGE and autoradiography. Labeling of lanes: [Translate] in vitro-translated proteins prior to precipitation; [−DNA] no input oligonucleotide DNA; [H + P site] oligonucleotide with intact consensus Hox and Pbx sites; [Mut. Pbx] oligonucleotide with mutant Pbx site; [Mut. Hox] oligonucleotide with mutant Hox site.

1994) sites was added to the translated proteins, almost equimolar amounts of HoxB7 were coprecipitated with Pbx1a or E2a–Pbx1a. Mutation of the Pbx site but not the Hox site in the oligonucleotide reduced the amount of coprecipitated HoxB7 to residual levels (Fig. 2). Because a large molar excess (1000-fold) of binding-site DNA was present in the reactions, coprecipitation likely resulted from cooperative protein interactions and not simply independent binding of both proteins to the same DNA molecule. Taken together, the in vitro and yeast data were consistent with potential interactions between Pbx and HoxB7 that were enhanced in the presence of DNA containing an intact Pbx consensus binding site.

Pbx proteins bind DNA cooperatively with HoxB7

To determine the effect of Pbx and Hox interactions on their respective DNA-binding properties, electrophoretic mobility shift assays (EMSA) were performed. We used the oligonucleotide containing both Hox and Pbx consensus binding sites, which facilitated coprecipitation of a Hox–Pbx DNA complex. When this oligonucleotide was added to binding reactions containing in vitro-synthesized Pbx1a or HoxB7 alone, no shifted complexes were observed under our EMSA conditions other than those corresponding to endogenous factors in the reticulocyte lysates (Fig. 3A). However, when both Pbx1a and HoxB7 were incubated together with the probe, a unique complex was detected whose formation was dependent on increasing amounts of either Pbx or Hox protein (Fig. 3A, lanes 4–9). Calculations based on the percent of shifted probe (5%–10%) suggested that most of the input protein (<0.3 ng each) was bound to DNA. Thus, the observed DNA binding occurred at very low concentrations (<0.5 nM) of Pbx and Hox proteins suggesting a highly specific interaction.

To assess the presence of both proteins in the shifted complex, supershift analyses were performed with specific antibodies. The complex formed by epitope-tagged Pbx1a and wild-type HoxB7 was supershifted by a monoclonal antibody against the epitope tag (Fig. 3B, lane 2)

Figure 2. Coprecipitation of Pbx proteins with HoxB7 in the presence but not absence of DNA. In vitro-synthesized, radio-labeled proteins were mixed and allowed to associate in the presence or absence of the indicated oligonucleotide DNAs. Protein complexes were specifically precipitated with a monoclonal antibody against an epitope-tagged Pbx1a or a monoclonal anti-E2a (Yae) antibody (denoted below gel lanes). Precipitated proteins were analyzed by SDS-PAGE and autoradiography. Labeling of lanes: [Translate] in vitro-translated proteins prior to precipitation; [−DNA] no input oligonucleotide DNA; [H + P site] oligonucleotide with intact consensus Hox and Pbx sites; [Mut. Pbx] oligonucleotide with mutant Pbx site; [Mut. Hox] oligonucleotide with mutant Hox site.

Figure 3. Pbx proteins bind DNA cooperatively with HoxB7. (A) EMSA demonstrating cooperative DNA binding by HoxB7 and Pbx1a proteins. Various in vitro-translated proteins were incubated in DNA-binding reactions in the presence of radiolabeled probe and then subjected to EMSA. The oligonucleotide probe contained adjacent, consensus binding sites for Hox and Pbx proteins (Pellerin et al. 1994; LeBrun et al. 1994). The volumes of translates added to individual binding reactions were 4 μl of unprogrammed lysate (lane 1); 3 μl of HoxB7 + 1 μl of lysate [lane 2]; 3 μl of Pbx1a + 1 μl of lysate [lane 3]; 1 μl of HoxB7 + 1 μl of Pbx1a + 1 μl of lysate [lane 4]; 2 μl of HoxB7 + 1 μl of Pbx1a + 1 μl of lysate [lane 5]; 3 μl of HoxB7 + 1 μl of Pbx1a [lane 6]; 1 μl of HoxB7 + 1 μl of Pbx1a + 2 μl of lysate [lane 7]; 1 μl of HoxB7 + 2 μl of Pbx1a + 1 μl of lysate [lane 8]; 1 μl of HoxB7 + 3 μl of Pbx1a [lane 9]. (lys) Endogenous complexes present in the reticulocyte lysate that bind to the DNA probe; [Hox + Pbx] Complexes resulting from cooperative binding of Hox and Pbx proteins. (B) Complexes resulting from cooperative DNA binding contain both Hox and Pbx proteins. Different combinations of in vitro-translated proteins [as indicated above the gel lanes] were incubated in DNA binding reactions and subjected to EMSA. For supershift analyses, monoclonal antibodies were added to binding reactions as indicated above the gel lanes. Binding reactions contained 2 μl of each specifically programmed lysate. Supershifted and Hox/Pbx complexes are indicated respectively. (lys) Endogenous complexes present in the reticulocyte lysate.
confirming the presence of Pbx1a in the complex. The presence of HoxB7 was demonstrated by shifted migration with the anti-tag monoclonal antibody of a complex formed by coincubation of epitope-tagged HoxB7 and wild-type Pbx1a [Fig. 3B, lane 4]. Cooperative binding of a similar magnitude was also observed with HoxB7 and Pbx1b, which differs from Pbx1a in its carboxy-terminal portions, and with other Pbx family members Pbx2, Pbx3a, and Pbx3b [data not shown]. These findings demonstrated that various wild-type Pbx proteins were able to bind DNA cooperatively as heteromeric complexes with HoxB7.

Pbx1 cooperatively binds DNA with HoxB4 and HoxB6 but not HoxA10

The spectrum of potential Pbx/Hox interactions was addressed by determination of whether other Hox proteins could cooperatively bind DNA with Pbx1. These studies employed HoxA10, HoxB4, and HoxB6, which are members of paralog groups different from HoxB7. Similar to results with HoxB7, incubation of HoxB6 alone with the Hox/Pbx combined-site probe resulted in no shifted complexes other than the endogenous lysate complex. Addition of increasing amounts of HoxB6 resulted in formation of a shifted complex similar but slightly slower in its migration to that observed for HoxB7–Pbx1a [Fig. 4, lanes 1–5]. The shifted complex formed by co-incubation of HoxB6 and Pbx1a contained both proteins as demonstrated by supershift analyses with specific antibodies [data not shown].

In contrast to HoxB6 and B7, incubation of HoxB4 with the combined-site probe showed a faint shifted complex in the absence of added Pbx1a [Fig. 4, lane 7] indicating that HoxB4 alone bound the probe under these conditions, consistent with earlier studies on its inherently higher affinity for the consensus Hox site (Pellerin et al. 1994). In the presence of increasing amounts of Pbx1a, a new, more abundant, complex was observed, whose migration was slower than that observed with HoxB4 alone [Fig. 4, lanes 8–13]. Enhanced formation of the HoxB4–Pbx1a complex compared to the HoxB4 complex was consistent with a cooperative effect on DNA binding, and supershift analyses [data not shown] confirmed the presence of both proteins in the complex.

Analogous experiments with HoxA10 showed that it also bound weakly to the combined-site probe in the absence of added Pbx1a [Fig. 4, lane 7] indicating that HoxB4 alone bound the probe under these conditions, consistent with earlier studies on its inherently higher affinity for the consensus Hox site (Pellerin et al. 1994). In the presence of increasing amounts of Pbx1a, a new, more abundant, complex was observed, whose migration was slower than that observed with HoxB4 alone [Fig. 4, lanes 8–13]. Enhanced formation of the HoxB4–Pbx1a complex compared to the HoxB4 complex was consistent with a cooperative effect on DNA binding, and supershift analyses [data not shown] confirmed the presence of both proteins in the complex.

Figure 5. Pbx1 does not cooperatively bind DNA with HoxA10. In vitro-translated proteins were incubated in DNA-binding reactions in the presence of radiolabeled probes and then subjected to EMSA. The relative volumes of specifically programmed translates [4 μl total] added to individual binding reactions are indicated above the gel lanes. Nucleotide sequences of the oligonucleotide probes are indicated below the autoradiograms; nucleotide differences in the Pbx + A10 probe are underlined. (lys) Endogenous complexes present in the reticulocyte lysate that bind to the consensus DNA probe. Complex resulting from HoxA10 alone binding to the probe is indicated at right.
trasted with the robust cooperative binding observed for HoxB4, B6, and B7 in the presence of Pbx1a with either the Hox–Pbx (Figs. 3 and 4) or the AbdB–Pbx (data not shown) probes. An element of specificity to Hox–Pbx cooperativity was also suggested by the inability of Pbx1a to bind cooperatively to either DNA probe in the presence of Drosophila even-skipped (data not shown). A reversed orientation of the Hox and Pbx sites in the Hox–Pbx probe yielded reduced but not complete loss of cooperativity (data not shown). These observations indicated that Pbx varied in its ability to cooperatively bind the synthetic DNA target with this set of Hox proteins. Specificity could not be entirely accounted for by differences in the binding affinities of the Hox proteins for the DNA probe, suggesting that Hox–Pbx protein–protein interactions were playing an important role in delimiting the observed cooperativity.

Chimeric E2a–Pbx1 oncoproteins bind DNA cooperatively with Hox proteins

Chimeric E2a–Pbx1 oncoproteins result from fusion with the heterologous E2a protein following chromosomal translocations in pre-B cell leukemias (Kamps et al. 1990; Nourse et al. 1990). Because fusion with E2a occurs in a highly conserved amino-terminal portion of Pbx1, we assessed whether chimeric E2a–Pbx1 proteins retain their ability to bind DNA cooperatively with Hox proteins. Incubation of E2a–Pbx1a with HoxB7 resulted in a shifted DNA–protein complex whose migration was slower than Hox–Pbx complexes formed with wild-type Pbx (Fig. 6A, lanes 3–8); no complex was seen when E2a–Pbx1a alone was incubated with the probe under these conditions (Fig. 6A, lane 2). Similar results were obtained with E2a–Pbx1a and HoxB4 and B6 (data not shown).

The presence of both E2a–Pbx1a and HoxB7 in the DNA-bound complex was confirmed by supershift analysis. A monoclonal antibody against E2a resulted in complete conversion to a slower migrating complex as did a monoclonal antibody directed against an epitope tag in HoxB7 (Fig. 6B, lanes 2,4). Thus, replacement of the amino-terminal region of Pbx1a with E2a does not impair cooperative DNA binding with Hox B4, B6, or B7.

Cooperative DNA binding requires an intact Pbx site

To determine whether Pbx–Hox cooperativity required the presence of both Hox and Pbx sites, mutant oligonucleotide probes (Fig. 7) were used for DNA binding and EMSA. These studies compared the relative cooperative DNA binding of Pbx1a with HoxB4, B6, and B7, respectively on each oligonucleotide probe. Analyses employing a probe with a mutant Pbx site resulted in complete abrogation of DNA binding by each of the Hox–Pbx complexes (Fig. 7, lanes 2,5,8). In contrast, various mutations of the Hox site showed a reduction but not complete loss of cooperative complex formation (Fig. 7, lanes 3,6,9). Probes Mut1–Hox and Mut2–Hox (data not shown) gave similar results. Quantitation of the shifted bands showed that the reduction in cooperative binding was greater for
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Figure 7. The Pbx site is required for cooperative DNA binding. (A) EMSA with mutant DNA probes. In vitro-translated proteins were incubated in DNA binding reactions in the presence of radiolabeled probe and then subjected to EMSA. Specifically programmed lysates (4 μl) and DNA probes added to individual binding reactions are indicated above the respective gel lanes. A second endogenous lysate band that is usually obscured by Pbx-Hox complexes appears more prominent in lanes with the Mut. Pbx probe. (B) Sequences of oligonucleotides used for EMSA. The consensus Hox and Pbx sites are indicated above the aligned sequences. Mutant nucleotides that deviate from the consensus are underlined.

Figure 8. The Pbx homeo domain (HD) is necessary but not sufficient for cooperative DNA binding with Hox proteins. (A) Pbx1 deletion mutants used for EMSA are shown schematically. Solid boxes indicate homeo domains. Numbers correspond to amino acid positions (Monica et al. 1991) at deletion end points. (B) EMSA of cooperative DNA binding by HoxB7 and mutant Pbx proteins. In vitro-translated proteins were incubated in DNA-binding reactions in the presence of radiolabeled probe and then subjected to EMSA. Specifically programmed translates (2 μl of Hox protein plus 2 μl of Pbx mutant) added to individual binding reactions are indicated above the gel lanes. Endogenous complexes present in the reticulocyte lysate that bind to the DNA probe. Complexes resulting from cooperative binding of Hox and Pbx proteins are indicated by brackets. (C) SDS-PAGE of in vitro-synthesized proteins. Various constructs encoding mutant Pbx proteins were translated in vitro and the radiolabeled proteins analyzed by SDS-PAGE.

8A] were used in binding reactions and analyzed by EMSA. Each of the mutant constructs was translated and analyzed by SDS-PAGE (Fig. 8B) to ensure that equivalent amounts of deleted Pbx proteins were added to the individual binding reactions. Data are only shown for cooperativity with HoxB7 although similar results were also obtained for HoxB4 and B6. A requirement for the Pbx1 homeo domain was demonstrated by the inability of construct ΔHDI to cooperatively bind the probe with HoxB7 [Fig. 8C, lane 6]. However, the homeo domain alone was not sufficient because a construct containing only the 64 amino acids spanning the Pbx1 homeo domain was incapable of cooperative binding [Fig. 8C, lane 5], suggesting that at least some of the flanking amino acids were necessary. Cooperative DNA binding by construct ΔN but not ΔC implicated Pbx amino acids downstream of the homeo domain [Fig. 8, lanes 2,4]. Deletion of the Pbx carboxy-terminus with preservation of 25 amino acids flanking the homeo domain [construct ΔC+25] did not abrogate cooperativity [Fig. 8C, lane 3] thereby localizing the required downstream amino acids to conserved residues immediately flanking the homeo domain. A minimal construct containing the homeo domain plus 15 downstream amino acids cooperatively bound DNA in the presence of HoxB7, B6, or B4 [Fig. 8B, lanes 9–11] and weakly bound DNA in their absence [lane 8]. Interstitial deletion of 10 amino acids from the implicated downstream region significantly reduced cooperativity by Pbx1α [Fig. 8C, lane 7]. Taken together, these results show that the determinants for cooperative DNA binding with this subset of Hox proteins are contained in the Pbx1 homeo domain and a limited number of highly conserved carboxy-terminal flanking amino acids.

The Hox homeo domain and conserved YPWM motif are required for cooperative binding with Pbx proteins.

A similar approach was employed to determine the por-
tions of HoxB7 required for cooperative DNA binding with Pbx. Various deleted HoxB7 proteins (Fig. 9A) were tested by EMSA for cooperative binding with wild-type Pbx1A. A mutant lacking the carboxyl terminus and helix III of the homeo domain ΔH3 + C was incapable of cooperativity with Pbx1A, suggesting that HoxB7 needed to bind DNA through its homeo domain, consistent with the observed partial dependence on an intact Hox site in the DNA probe. Specific carboxy- and amino-terminal truncations of HoxB7 that left the homeo domain intact did not affect cooperativity, thereby defining a minimal fragment [HD + YPWM] spanning amino acids 120–198 capable of cooperative DNA binding (Fig. 9C, lane 4). Deletion of 10 amino-terminal amino acids from this construct [HD−YPWM] resulted in loss of cooperativity in spite of the fact that this HoxB7 mutant could bind DNA weakly by itself in the presence [Fig. 9C, lane 5] or absence (data not shown) of Pbx1A. A construct containing HD−YPWM plus the carboxy-terminal tail of HoxB7 also showed weak binding alone but no cooperativity (data not shown). These studies suggested that in addition to the HoxB7 homeo domain, additional flanking amino acids amino-terminal of the homeo domain were required and that there was no requirement for residues carboxy-terminal to the homeo domain.

The region amino-terminal to the homeo domain of HoxB7 required for cooperativity contains the so-called hexapeptide motif that is highly conserved in a large subgroup of Hox/HOM proteins (Bürglin 1994; Gehring et al. 1994a). To test directly whether the hexapeptide motif was necessary for cooperative DNA binding, it was altered by site-directed mutagenesis in HoxB6 and HoxB7 proteins. In vitro-translated proteins were incubated in DNA-binding reactions in the presence of radiolabeled probe and then subjected to EMSA. Specifically programmed translates [2 μl of Pbx1A plus 2 μl of Hox mutant] added to individual binding reactions are indicated above the gel lanes. (lys) Endogenous complexes present in the reticulocyte lysate that bind to the DNA probe.
10B, lanes 1–4). These data indicated that the hexapeptide motif plays a critical role in the observed Hox–Pbx cooperativity and likely accounts for the requirement of residues amino-terminal of the Hox homeo domain. Because HoxB7 proteins with hexapeptide mutations were incapable of binding DNA in the absence of Pbx1a (data not shown), the YPWM motif does not have the characteristics of a negative regulatory element that inhibits HoxB7 binding on its own.

The role of the hexapeptide was tested further by determination of whether it was sufficient to enhance the ability of HoxA10 to cooperatively bind DNA with Pbx1a. A mutant [hex-A10 in Fig. 11] was constructed in which amino acids 280–316 upstream of the HoxA10 homeo domain were replaced by 15 amino acids (120–134) from HoxB7 spanning its hexapeptide. EMSA showed that both wild-type HoxA10 and hex-A10 mutant bound the DNA probe in the absence of Pbx1a [Fig. 11, lanes 1, 5]. However, only hex-A10 displayed robust cooperativity in the presence of Pbx1a [Fig. 11, lanes 2, 6] comparable to that observed for HoxB7 [Fig. 11, lane 8]. Most of hex-A10 was present in a complex with Pbx1a in contrast to wild-type HoxA10, which bound alone to the probe even in the presence of Pbx1a. Cooperativity by mutant hex-A10 was directly attributable to the hexapeptide acquired from HoxB7 and not deletion of an inhibitory domain because a mutant HoxA10 with only a partial HoxB7 graft lacking the YPWM motif (del-A10) displayed no cooperativity [Fig. 11, lane 4]. These data demonstrate that the hexapeptide motif is both necessary and sufficient for Hox proteins to cooperatively bind the synthetic probe with Pbx proteins.

Discussion

Evolutionary conservation of a molecular code that regulates Hox–Pbx interactions

The studies reported here provide strong support for the thesis that Pbx proteins are selective cofactors for Hox proteins. Hox proteins have been assigned to paralog groups according to the degree of sequence conservation among their respective homeo domains (Gehring et al. 1994a). On this basis HoxB6 and B7 are in paralog groups that share the highest similarity with the Drosophila proteins Antp, Ubx, and Abd-A (the so-called Antp class). Both Ubx and Abd-A have recently been shown to cooperatively bind DNA with exd, the Drosophila homolog of Pbx (VanDijk and Murre 1994; Chan et al. 1994). Thus, in addition to structural conservation within their homeo domains, Hox/HOM proteins of the Antp class also demonstrate a conserved capacity to function as heterotypic DNA-binding complexes with Pbx/exd proteins. HoxB4 belongs to the Deformed class on the basis of sequence similarities inside and flanking the homeo domain. Because HoxB4 bound DNA cooperatively with Pbx1, our data indicate that the capacity for cooperative interactions extends beyond the Antp class of homeo domain proteins to include other paralog groups. Cooperative interactions, however, were not observed with HoxA10, which is a member of the Abd-B class, consistent with previous observations that exd does not cooperatively bind DNA with Abd-B (VanDijk and Murre 1994). Thus, Pbx/exd proteins appear to interact specifically with a select subset of homeo domain proteins.

Cooperative interactions between Hox and Pbx proteins appear to be determined at least in part by sequences outside of their respective homeo domains. For both HoxB6 and B7, amino acid residues immediately amino-terminal to their homeo domains were essential for cooperative DNA binding with Pbx1. Site-directed mutagenesis of the conserved hexapeptide motif in this region completely abrogated cooperative DNA binding with both HoxB6 and B7. The highly conserved hexapeptide motif defines a distinct superclass of homeo domain proteins that includes the Antp homologs HoxB6 and B7 as well as the Dfd family member HoxB4 and several other groups of homeo domain proteins outside of the HOM/Hox clusters (Bürglin 1994) whose representatives were not examined here. On the basis of our observations and those of others on Ubx (Johnson et al. 1995) it seems reasonable to propose that the hexapeptide motif...
constitutes a key molecular determinant of the ability of select homeo domain proteins to cooperatively interact with Pbx/exd proteins.

Previous studies were unable to demonstrate cooperative DNA binding by Antp and exd (Chan et al. 1994), a surprising result given that Antp is a member of the hexapeptide superclass with a homeo domain that is highly conserved with HoxB7. However, the Antp construct employed by Chan et al. (1994) did not contain the hexapeptide motif. On the basis of our data, we predict that Antp and exd should cooperatively bind DNA, a prediction supported by genetic studies indicating that regulated expression of *teashirt* by Antp is dependent on exd function [Rauskolb and Weischaus 1994]. It is also apparent that cooperativity with Pbx/exd proteins is unlikely to be limited to the hexapeptide superfamilies as engrailed, which lacks a hexapeptide motif, was shown to cooperatively bind DNA with exd [VanDijk and Murre 1994], although it has been noted that the engrailed EH2 motif contains residues that are somewhat reminiscent of the hexapeptide [Bürglin 1994]. We cannot rule out the possibility that proteins such as HoxA10 may bind cooperatively with Pbx1 to different DNA targets as a result of interactions that do not require a hexapeptide motif. Additional studies are necessary to establish the complete spectrum of potential cooperative Hox–Pbx interactions but our studies suggest that the hexapeptide motif constitutes one critical component of a conserved molecular code governing at least some of these interactions.

Although previous studies suggested the importance of the Ubx carboxy-terminal tail for cooperativity with exd [Chan et al. 1994], the comparable region of HoxB7 was neither essential nor sufficient in our studies for cooperativity with Pbx1a. A potential requirement for the carboxy-terminal tail of Ubx contrasts with the lack of sequence conservation in this region of Ubx, HoxB6, and HoxB7, which show similarity only in their homoeo domains and hexapeptide motifs. However, the studies by Chan et al. (1994) employed Ubx constructs that lacked the conserved hexapeptide motif and were conducted at protein concentrations at least several hundred-fold over those employed by us. Weak cooperativity was previously observed for Ubx alone on a multisite probe and proposed to be mediated by sequences carboxy-terminal to the Ubx homeo domain [Beachy et al. 1993].

Our studies also define a domain within Pbx proteins that may regulate their DNA-binding properties and serve as a contact site for cooperative interactions with Hox proteins. This domain flanks the Pbx1 homeo domain and is highly conserved in Pbx family members and their homologs exd and ceh-20 [Monica et al. 1991; Bürglin and Ruvkun 1992; Flegel et al. 1993; Rauskolb et al. 1993]. By deletional mutagenesis we showed that removal of 10 conserved amino acids carboxy-terminal to the Pbx1 homeo domain severely compromised cooperative DNA binding, and a minimal construct containing the Pbx homeo domain demonstrated no cooperativity in the absence of downstream residues. These observations suggest that the downstream flanking amino acids play a critical role in stabilizing interaction of the Pbx homeo domain with DNA. Solution and crystal structures of homeo domains show that carboxyl flanking amino acids partially contribute to helix IV, an elongation of recognition helix III that may assume a flexibly disordered conformation in solution [Gehring et al. 1994b]. We posit that Hox proteins may interact with the amino acids downstream of the Pbx homeo domain thereby inducing a conformational change in helix IV and flanking amino acids that stabilizes homeo domain binding to DNA. There is precedent for the role of carboxy-terminal amino acids in regulating heterotypic interactions between homeo domain proteins as reported previously for cooperative interactions between MEC-3 and Unc-86 [Xue et al. 1993].

Our observations open new doors for investigating how Hox proteins may achieve some of their functional specificity in spite of extensive similarities in their DNA binding homeo domains. For a subset of Hox proteins containing the hexapeptide motif this would occur through interactions with Pbx proteins. Although our initial observations of Hox–Pbx interactions support a model in which Pbx proteins serve accessory roles as parallel effectors of Hox activity they do not provide an answer to the question of how Hox proteins generate different outputs. Interactions with Pbx only allowed a distinction between the Hox4–7 and Hox10 paralog groups, which bind slightly different sequences. Furthermore, under our experimental conditions, Pbx appeared to contribute most of the binding specificity of the complex, an apparent weak means of achieving specificity given the general expression of Pbx versus Hox proteins [Monica et al. 1991]. Clearly, additional studies are necessary to dissect further the molecular code governing Hox–Pbx cooperativity and its potential contribution to specificity of Hox function. It remains to be determined whether DNA binding to physiologically relevant sites will reveal a greater role for Hox proteins in specificity of DNA binding within the context of Pbx cooperativity. It is also likely that additional combinatorial interactions with other transcription factors contribute to the functional specificity of Hox proteins.

The observed role of Pbx in cooperative binding in vitro shares some similarities with that of a1, the homeo domain protein that interacts with a2 to repress genes involved in determination of yeast mating type [Johnson 1992]. Most of the DNA-binding specificity derives from a1 which alone has no intrinsic DNA-binding activity. Contact of a1, with the carboxy-terminal tail of a2, a low specificity DNA-binding protein by itself, creates a highly specific DNA-binding complex [Stark and Johnson 1994]. Similar to a1, full-length Pbx1 has little or no intrinsic DNA binding ability even at high concentrations of 200 nM [C.-P. Chang and M.L. Cleary, unpubl.] but is highly dependent on cooperative interactions to specifically bind its consensus DNA site. Like a2, the Hox proteins in our studies have relatively low intrinsic DNA-binding specificity [Pellerin et al. 1994], and amino acids outside of the Hox homeo domain appear to be
critical for cooperative DNA binding with Pbx1. Thus, Pbx–Hox cooperativity may adhere to the a1/a2 model, where two DNA-binding proteins with minimal intrinsic DNA binding ability can together create a specific DNA-binding complex. In spite of the similarities between Pbx–Hox and a1–a2 we have been unable to demonstrate cooperative binding of Pbx and Hox proteins on hsg operator sites (C.-P. Chang and M.L. Cleary, unpubl.). These and other observations suggest that Pbx–Hox heterodimers may prefer a configuration of half-sites different than a1–a2, and additional studies are required to define optimal sequences for Hox–Pbx cooperative DNA binding.

Hox proteins may be essential cofactors for E2a–Pbx1 chimeric oncoproteins

Our studies demonstrate that the cooperative effects of Hox proteins on Pbx DNA binding also extend to E2a–Pbx1 chimeric oncoproteins that result from chromosomal translocations in acute pre-B cell leukemias. E2a–Pbx1 cooperatively bound the consensus Pbx site with Hox proteins under conditions that yielded no DNA binding by E2a–Pbx1 alone. The portion of Pbx1 required for cooperative DNA binding is comparable to the minimal portion of Pbx1 that when fused to E2a is sufficient for transformation of NIH-3T3 cells [Monica et al. 1994]. These findings suggest that E2a–Pbx1 may be dependent on interactions with heterologous proteins for specific recognition of subordinate genes whose expression is responsible for oncogenesis. Although it is not clear from our studies which Hox proteins may cooperate with E2a–Pbx1 in t(1;19)-bearing leukemias, we note that the gene for HoxC4, which contains the YPWM motif, appears to be the major Hox gene expressed in lymphoid cells [Lawrence and Largman 1992; Lawrence et al. 1993]. Additional studies characterizing endogenous lymphoid E2a–Pbx1 complexes that bind the consensus Pbx site may help address this issue.

Although the Pbx1 homeo domain and its flanking sequences are sufficient for transformation when fused to trans-activation elements from the E2a protein, the homeo domain, paradoxically, is not essential for transformation by E2a–Pbx1. Recent studies have shown that an E2a–PbxXH mutant transforms fibroblasts in vitro and induces malignant lymphomas in transgenic mice [Monica et al. 1994] that are indistinguishable from those induced by E2a–Pbx1 [Dedera et al. 1993]. These findings suggested that E2a–Pbx1 may interact with cellular proteins that assist or mediate its effects on gene expression responsible for oncogenesis even in the absence of Pbx homeo domain–DNA interactions. A potential molecular explanation for transformation by the homeo domain-lacking mutant is suggested by the observed cooperativity between Pbx and Hox proteins. The deletion construct that maintained transforming activity lacked the homeo domain but preserved Pbx1 amino acids outside the homeo domain that were shown in the present study to be necessary for cooperative DNA binding with Hox proteins. It is possible that the E2a–PbxXH mutant protein retains some of its ability to interact with a heterologous Hox protein forming a complex that can specifically bind a Hox DNA site and regulate the expression of critical subordinate genes analogous to activation of an engrailed-derived promoter by PRD and a ftz mutant lacking its homeo domain [Ananthan et al. 1993]. Our inability to demonstrate a comparable interaction of Hox proteins with Pbx lacking its homeo domain may reflect limitations of the binding assay or the particular Hox proteins employed. Clearly, additional studies are necessary to further elucidate the role of Hox proteins in contributing to and perhaps defining the oncogenic effects of chimeric Pbx proteins. The studies reported here suggest a molecular model for understanding how wild-type and chimeric Pbx proteins may function in parallel with Hox proteins to regulate gene expression in normal and neoplastic cells.

Materials and methods

DNA constructs

In vitro expression clones for wild-type and chimeric Pbx proteins under control of the SP6 promoter have been described in previous studies [Nourse et al. 1990; Monica et al. 1991]. Deletion constructs of E2A–PBX1 have been reported elsewhere [Lebrun and Cleary 1994; Monica et al. 1994]. For the current studies, additional deleted forms of PBX1 and HOX7 cDNAs were constructed by use of standard cloning techniques and PCR. An epitope-tagged [FLAG] Pbx1a was constructed by replacement of the nine amino-terminal amino acids with a synthetic oligonucleotide encoding the amino acids MDYKD-SS. In vitro expression constructs for human HoxB4, HoxB6, HoxB7, and HoxA10 consisted of the respective human cDNAs [Simeone et al. 1987; Peeverali et al. 1990; Shen et al. 1991] cloned downstream of the SP6 promoter. An epitope-tagged form of HoxB7 consisted of 14 additional amino-terminal amino acids [MDYKD-DDKKSL] fused in-frame to the HoxB7 initiating methionine. An epitope-tagged version of HoxB6 was constructed by deletion of amino acids [MDYKD-DDKKSLG] to its amino terminus. Grafting of the HoxB7 hexapeptide onto HoxA10 involved deletion of amino acids 280–316 of HoxA10 and replacement of them with HoxB7 amino acids 120–134 [mutant hex-A10] or 131–134 [mutant del-A10]. An in vitro-expression construct for Drosophila even-skipped [pAR-EVE] in the pET3a vector has been described elsewhere [Hoey et al. 1988].

EMSAs

Various Pbx and Hox proteins were produced in vitro from SP6 expression plasmids by use of a coupled reticulocyte lysate system under conditions recommended by the supplier [Promega, Madison, Wis.]. Hox and Pbx proteins were translated separately and then added to binding reactions. To ensure that approximately equal amounts of in vitro-translated proteins were present in binding reactions, parallel translation reactions were performed in the presence of 35S-methionine, and labeled proteins were subjected to SDS-PAGE, quantitated on a PhosphorImager [Molecular Dynamics], and normalized for the number of methionine residues in each protein. DNA-binding reactions were performed at 4°C for 30 min and contained up to 4 µl of reticulocyte lysate proteins [<0.5 ng of Hox or Pbx] in a total volume of 15 µl containing 2 µg of poly[d(I-C)], 75 mM NaCl, 1 mM EDTA, 1 mM DTT, 10 mM Tris-HCl [pH 7.5], 6% glycerol, 2 µg BSA, and 50,000 cpm of DNA probe [1 ng]. Thirteen mi-
trolters of the binding reactions was subjected to EMSA under conditions reported previously [Jacobs et al. 1993] with 6% polyacrylamide gels (0.75 mm thickness) in 0.25x TBE buffer. DNA probes (50,000 cpm/binding reaction) consisted of gel-purified, end-labeled, double-stranded oligonucleotides whose sequences are described in the text and fig. legends. The combined-site DNA probe used for EMSA (5'-CGAATTGATTGTAGCACC-TAAATTGGAC-3') was designed to resemble yeast haploid-specific gene operator sequences based on similarities in the Pbx and Hox binding sites [TGATGTGT vs. TGATGTGA, respectively] and assuming a tandem orientation [Goutte and Johnson 1994] of the Pbx and Hox proteins. DNA complexes were supershifted with commercially prepared antibodies directed against the FLAG epitope tag or a monoclonal directed against E2a [Jacobs et al. 1993]. Antibodies (1 µg) were added to DNA-binding reactions and incubated on ice for 2 hr prior to addition of oligonucleotide probes.

**Immunoprecipitations**

In vitro-translated proteins in reticulocyte lysates (10 µl each of HoxB7 and Pbx translates) were added to 180 µl of 1× binding buffer (10 mM Tris-HCl at pH 7.5, 75 mM NaCl, 1 mM EDTA, 1 mM DTT, 10% glycerol, 1% BSA, 0.05% NP-40, 2 µg of poly[I-C]) in the presence or absence of oligonucleotide DNA (1 µg) and incubated overnight at 4°C in the presence of antibodies. For precipitation of Pbx1a, an anti-FLAG monoclonal antibody (1 µg/200 µl) was used with protein G sepharose beads. For precipitation of E2a-Pbx1a, an anti-E2a monoclonal antibody [Jacobs et al. 1993] cross-linked to sepharose beads was employed. Beads were washed 10 times with 1× binding buffer and precipitated proteins analyzed by SDS-PAGE.

**Yeast two-hybrid analysis**

Various fragments of PBX1 cDNAs were cloned into pACT2 [Harper et al. 1993] in-frame with the GAL4 activation domain. A NcoI-SacII fragment of HoxB7 cDNA encoding amino acids 110–198 spanning the hexapeptide motif and homeo domain was cloned into pAS2 [Harper et al. 1993] in-frame with the GAL4 DNA-binding domain. Yeast strain Y190 was transformed to trp and leu prototrophy with the pACT–PBX1 and pAS–HOXB7 constructs. Activation of the lacZ reporter gene was determined by assessing fPgal activity with a filter lift assay [Breeden et al. 1985].

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