Specificity of HOX protein function depends on DNA–protein and protein–protein interactions, both mediated by the homeo domain

Vincenzo Zappavigna, Daniela Sartori, and Fulvio Mavilio

Laboratory of Gene Expression, Department of Biology and Technology [DIBIT], IstitutoScientifico H.S. Raffaele, Milan, Italy

Transcription of human HOX gene promoters in cultured cells is positively and negatively regulated by HOX proteins interacting with specific target sequences. The human HOXD9 protein activates transcription of the HOXD9 promoter by interacting with the HCR sequence and is antagonized by the HOXD8 protein. HOXD8 is not intrinsically a repressor, since it can activate transcription on different targets. Complete or partial HOXD8/HOXD9 homeo domain swapping indicates that the ability to recognize, and activate transcription from, the HCR target in vivo depends on the amino terminus and helix 1 of the homeo domain. The inhibitory activity of HOXD8 is not affected by deletion of the homeo domain helix 2/3 region, whereas it requires the amino terminus/helix 1 region and an additional, effector domain located at the protein amino-terminal end. This activity is therefore DNA-binding independent, and possibly mediated by protein–protein interactions. Affinity chromatography experiments show that the homeo domain amino terminus/helix 1 region is able to mediate direct interactions between HOX proteins in solution. These data indicate that specificity of HOX protein function in vivo depends on both DNA–protein and protein–protein interactions, mediated by the same sub region of the homeo domain.

[Key Words: Homeo box; homeo domain; transcriptional regulation; DNA–protein interaction; protein–protein interaction]

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Vertebrate homeo box-containing (Hox) genes, homologous to the Drosophila homeotic selector (HOM) genes, are organized in four related, unlinked clusters derived by duplication from a putative, ancestral Hox complex [Schubert et al. 1993]. Expression of Hox genes is developmentally regulated and regionally restricted into partially overlapping expression domains in the paraxial mesoderm, the neuroectoderm, and the limbs [for review, see Gruss and Kessel 1991; Duboule 1992, McGinnis and Krumlauf 1992; Krumlauf 1993]. This expression pattern suggests the existence of a Hox “code” specifying the identity of different regions or segments of the body, whereby progressively more posterior structures are defined by the activation of progressively more 5’ Hox genes in all four clusters [Hunt and Krumlauf 1991; Kessel and Gruss 1991]. The molecular mechanisms by which this putative code is translated into positional specification are still largely unknown.

Hox proteins belong to a large family of transcription factors whose DNA-binding domain, the homeo domain, is structurally related to the helix–turn–helix motif common to several prokaryotic regulatory proteins [Laughon and Scott 1984; Gehring et al. 1990]. Consistently, Hox proteins have been shown to bind DNA in vitro [Fainsod et al. 1986; Odenwald et al. 1989; Sasaki et al. 1990] and to regulate transcription through specific target sequences in cell culture [Zappavigna et al. 1991; Arcioni et al. 1992; Jones et al. 1992; Pöpperl and Featherson 1992; Jones et al. 1993]. The homeo domain and immediate flanking regions in both Hox and HOM proteins play a crucial role in DNA sequence recognition and binding in vitro [Desplan et al. 1988; Hoey and Levine 1988; Müller et al. 1988], and, at least in the case of HOM proteins, in the determination of the functional specificity in vivo [Kuziora and McGinnis 1989; Kuziora and McGinnis 1991]. However, because all known Hox and HOM proteins have similar, if not identical, DNA-binding specificity in vitro [for review, see Hayashi and Scott 1990], they are unlikely to achieve their specificity of action in vivo only on the basis of DNA–protein interaction. The homeo domain DNA-binding and high-affinity target recognition functions have been shown to be neither sufficient nor absolutely required for the biological activity of HOM proteins, both in vivo and in cell culture [Fitzpatrick et al. 1992; Ananthan et al. 1993;}

1Corresponding author.
Schier and Gehring 1993). Therefore, protein–protein interactions between different HOM (or Hox) proteins, or between these proteins and other components of the transcriptional machinery, could be critical factors in the regulation of HOM/Hox functional specificity (Hayashii and Scott 1990; McGinnis and Krumlauf 1992). In the case of other homeo domain-containing proteins, protein–protein interactions do play a role in enhancing DNA-binding specificity and/or regulating their functions [see Manak and Scott 1993], the best characterized examples being the yeast MAT [Smith and Johnson 1992] and the POU [Stern et al. 1989; Treacy et al. 1992; Verrijzer et al. 1992] transcription factors. So far, however, products of the HOM/Hox family genes were not known to establish protein–protein interactions among themselves and/or with other factors, even though such interactions have been postulated in several cases in order to explain their mechanism of action [Jaynes and O’Farrell 1988; Han et al. 1989; Jaynes and O’Farrell 1991; Fitzpatrick et al. 1992; Ananthan et al. 1993; Furukubo-Tokunaga et al. 1993; Schier and Gehring 1993].

We and others had reported previously the existence of auto- and cross-regulatory interactions between mammalian Hox proteins and Hox gene regulatory elements, at least in cell culture [Zappavigna et al. 1991; Arcioni et al. 1992; Pöpperl and Featherstone 1992]. In particular, the products of the human HOXD9 [HOX4C or Hox-4.4 according to the old nomenclature] and the immediately 5′ HOXD10 [HOX4D or Hox-4.5] genes were shown to activate transcription of the HOXD9 promoter by binding to an evolutionary conserved, 100-bp upstream element, termed HOX cross-talk region (HCR). The product of the immediately 3′ gene HOXD8 [HOX4E or Hox-4.3] showed no activity on the HCR target but was able to antagonize the activating function of HOXD9 and HOXD10 in a cotransfection assay [Zappavigna et al. 1991]. Whether these interactions take place and have any role in regulating Hox gene expression patterns in embryos is still to be established. Nevertheless, cross regulation in transfected cells provides a very convenient model for analyzing the DNA-binding requirements and functional properties of Hox proteins in vivo. In this report we show that specific target recognition by HOXD8 and HOXD9 in vivo is determined by the amino terminus and helix 1 of the homeo domain. The HOXD8 transcriptional inhibitory function is not dependent on DNA binding, and requires both the presence of the homeo domain amino terminus/helix 1 and the activity of an amino-terminal effector domain. We show that the amino terminus/helix 1 sub region of the Hox homeo domain is able to mediate protein–protein interactions between Hox proteins.

Results

Specificity of target recognition by HOXD8 in vivo is determined by the amino terminus/helix 1 region of the homeo domain

The HOXD9 and HOXD8 proteins were expressed in the SV40 promoter-based constructs pD9 and pD8, respectively [nos. 1 and 2 in Fig. 1]. In a transient cotransfection assay in NIH–3T3 cells, HOXD9 was able to activate transcription from the regulatory sequence HCR fused to the −81 herpes simplex virus–thymidine kinase [HSV–TK] promoter in the pTHCR luciferase [luc] reporter construct [Fig. 1], whereas HOXD8 was inactive [Zappavigna et al. 1991, Fig. 2A]. To test whether HOXD8 is intrinsically unable to activate transcription, the amino-terminal 176 amino acids were replaced by the acidic activation domain of the yeast transcription factor GAL4 [amino acids 149–196] [Gill and Ptashne 1987] in the pGALD8 expressor construct [no. 3 in Fig. 1]. Cotransfection of pGALD8 with pTHCR resulted in no significant stimulation of luc activity [Fig. 2A], indicating that the lack of activating function in the HOXD8 protein is not attributable to the absence of an efficient activation domain. In parallel, pD8 and pGALD8 were cotransfected with the reporter construct pTCBS, in which a multimerized form of a homeo domain consensus-binding sequence [CBS; see Materials and methods] was linked to the −109 HSV–TK promoter [Fig. 1]. This construct is trans-activated in the cotransfection assay by a large number of HOX proteins [not shown], including HOXD9 [Fig. 2B]. The activity of pTCBS increased six- to eight-fold when cotransfected with pD8 or pGALD8 [Fig. 2B], showing that HOXD8 is able to activate transcription from an appropriate target in cell culture.

These results suggest that the HOXD8 protein is unable to recognize the HCR target in vivo, although it can bind to the same sequence in vitro [Zappavigna et al. 1991]. To test this hypothesis, we generated two chimeric constructs representing homeo domain swaps between the HOXD8 and HOXD9 proteins. In the pD8swD9 expressor construct [no. 4 in Fig. 1], 66 amino acids of the HOXD8 protein, 1 amino acid upstream to 5 amino acids downstream of the homeo domain, were replaced by the corresponding residues of the HOXD9 homeo domain (Fig. 2B), showing that HOXD8 is able to activate transcription from an appropriate target in cell culture.
Figure 1. Protein-expression and reporter constructs used in cotransfection experiments. Complete coding sequences of the HOXD9 (white) and HOX8 (shaded) genes are indicated. The regions of HOXD9 and HOX8 involved in the chimerical and/or mutant versions of both proteins (amino terminus, homeo domain, carboxyl terminus) are indicated by separate boxes. The homeo domains (darker in HOXD8) are further subdivided in two boxes, representing the amino terminus/helix 1 (amino acids 1–20) and helices 2–4 (amino acids 20–60). Helices 1, 2, and 3/4 are indicated schematically by small black bars on top of construct no. 1. (■) The acidic activator domain of the yeast GAL4 transcription factor, a hatched box indicates the coding sequence of the GST gene. The splicing signs indicate internal deletions in the HOXD8 amino terminus and homeo domain. In the reporter plasmids, the HOXD9 upstream control element (HCR), and an 8-mer repeat of a Hox CBS are indicated. The activity of each expression construct in trans-activating the pTHCR or pTCBS reporter is indicated by + and − signs (right), together with the ability to inhibit the HOXD9-induced trans-activation of pTHCR. (n.d.) Not determined.
Specificity of HOX protein function

likely to be attributable to a squelching effect, as transfection of excess pD8 caused no inhibition of the HOXD8-mediated trans-activation of the pTCBS reporter, nor of the activity of an unrelated transcription factor on its target, like that of the retinoic acid receptor β [RARβ] on the retinoic acid responsive element [RARE]–TK–luc reporter construct [data not shown]. Cotransfection of pD8ΔH2/3 [no. 7 in Fig. 1], which expresses a HOXD8 deletion mutant lacking the homeo domain DNA-binding helix 2/3 region and the protein carboxyl terminus [amino acids 214–289, see Fig. 3], caused the same dose-dependent antagonistic effect observed for pD8 [Fig. 4], indicating that the HOXD8 inhibitory activity does not require DNA binding. Conversely, cotransfection of pD8HD [no. 8 in Fig. 1], expressing only the homeo domain and carboxyl terminus of HOXD8 [amino acids 194–289, see Fig. 3], had no effect on the HOXD9-induced trans-activation of pTHCR [Fig. 4]. These results indicate that transcriptional inhibition by HOXD8 is not attributable to competition for binding to the HCR sequence. Production and correct nuclear targeting of the D8HD and D8ΔH2/3 mutants was checked by immunoblotting and immunofluorescence analysis [see Materials and methods, data not shown].

The HOXD8 inhibitory activity requires the presence of the amino terminus/helix 1 region of the homeo domain

To identify the region[s] of the HOXD8 protein required for the inhibitory function, two additional constructs were generated which express deletion mutants lacking either the whole homeo domain [pD8ΔHD, no. 9 in Fig. 1, encoding the amino-terminal portion of HOXD8 from amino acids 1–193, see Fig. 3], or only the amino terminus and helix 1 [pD8ΔH1, no. 10 in Fig. 1, deleted in the region corresponding to amino acids 196–214, see Fig. 3]. Cotransfection of either pD8ΔHD or pD8ΔH1 had no effect on the HOXD9-mediated trans-activation of pTHCR [Fig. 4], showing that the 19-amino acid region that contains the amino terminus and helix 1 of the HOXD8 homeo domain is necessary for the inhibitory activity. Production and nuclear targeting of the D8ΔHD and D8ΔH1 deletion mutants were checked as described above [not shown]. Interestingly, expression of the HOXD8 mutant lacking the DNA-binding region of the homeo domain [pD8ΔH2/3, no. 7 in Fig. 1] caused inhibition of the wild-type HOXD8 activity on the pTCBS reporter construct. This suggests that the D8ΔH2/3 protein, unable to bind DNA and therefore to activate transcription from the pTCBS reporter, behaves as a dominant–negative mutant with respect to the HOXD8/CBS interaction. Excess HOXD8 protein caused no inhibition of its own activity on the CBS target [not shown], suggesting that the inhibitory activity of HOXD8 is alternative to target recognition, and unmasked by the deletion of part of its DNA-binding domain.

HOXD8 inhibits the activating function of HOXD9 on the HCR target

Cotransfection of increasing amounts of pD8 [from equimolar to 2:1 excess] with pD9 and pTHCR resulted in progressive inhibition of the reporter gene trans-activation induced by HOXD9 [Fig. 4]. This inhibition is un-
The amino terminus and helix 1 of the HOXD8 homeo domain mediate protein–protein interactions between HOXD8 and HOXD9

The experiments described above showed that the inhibitory activity of HOXD8 is DNA binding independent and requires the presence of the amino terminus/helix 1 sub region of the homeo domain. To provide evidence for a possible role of this sub region in mediating protein–protein interactions between HOXD8 and HOXD9, we used a protein–protein affinity chromatography technique. A fusion protein between the HOXD9 homeo domain (amino acids 270–342, see Fig. 3) and the carboxyl terminus of the glutathione-S-transferase (GST) enzyme was expressed in bacteria, purified, and immobilized onto glutathione–Sepharose resin. The HOXD8 protein and the deletion mutants D8AHD and D8AH2/3 were translated and 35S-labeled in vitro, batch-adsorbed to the GST–HOXD9-homeo domain resin (GST–D9HD in Fig. 5A) or to a GST control resin (GST in Fig. 5A), and then eluted and analyzed by SDSPAGE. As shown in Figure 5A, the full-length HOXD8 protein [D8] and the D8AH2/3 mutant lacking the homeo domain helix 2/3 region were specifically retained by the GST–D9HD resin, whereas the D8AH2/3 mutant lacking the entire homeo domain was not retained. These data show that HOXD8 and HOXD9 are able to establish protein–protein contacts in solution in the absence of DNA through their homeo domains and that the amino terminus/helix 1 of the HOXD8 homeo domain is necessary for this interaction. The in vitro-translated HOXD9 protein was able to associate as efficiently as HOXD8 to the GST–D9HD resin [Fig. 5A], indicating that the HOXD9 homeo domain has a potential ability to establish both heterophilic and homophilic interactions in solution.

A GST fusion protein containing only the amino terminus/helix 1 of the HOXD9 homeo domain [amino acids 270–294, see Fig. 3] was then generated [GST–D9H1 in Fig. 5B], coupled to glutathione–Sepharose, and used in protein–protein affinity chromatography experiments. As shown in Figure 5B, the D8 and the D8AH2/3 mutant associated specifically with the GST–D9H1 resin, whereas the D8AH2/3 mutant failed to bind. Binding of HOXD8 to the GST–D9H1 resin was specifically competed by a ~20-fold excess of unlabeled protein [D8c] produced in bacteria [Fig. 5B]. In a reciprocal experiment, the in vitro-translated HOXD9 protein was specifically retained by a resin coupled to a fusion between GST and the amino terminus/helix 1 (amino acids 194–214, see Fig. 3) of the HOXD8 homeo domain [GST–D8H1, Fig. 5B, right]. On average, 10–25% of the labeled, input proteins were retained by the GST fusion resins. These results indicate that the HOXD8/HOXD9 protein–protein interaction in solution is mediated by the amino terminus/helix 1 sub region of both homeo domains.

Binding of HOXD9 to HCR in vitro is not affected by HOXD8

Because HOXD8 and HOXD9 are able to establish pro-
protein–protein interactions in the absence of DNA, we
tested whether such interactions may affect the binding
of HOXD9 to the HCR target. The full-length HOXD8
and HOXD9 proteins were expressed in NIH-3T3 cells
fused to an amino-terminal, 6-histidine tag and partially
purified by affinity chromatography under native condi-
tions (see Materials and methods). The amount of active
protein was titrated in both preparations by in vitro bind-
ing to a 32-bp oligonucleotide containing two HOX-bind-
ing sites located in the 5’ part of HCR (see Materials and
methods), followed by a gel-shift assay [Fig. 6A]. A fixed
amount of active HOXD9 protein was then mixed with
increasing amounts of HOXD8, incubated with the
HCR-derived oligonucleotide and then analyzed by the
same assay. The concentration of HOXD8 and HOXD9
in the binding reactions was in the same order of mag-
nitude as in the GST-affinity-chromatography experi-
ments (see Materials and methods), potentially allowing
the establishment of protein–protein interactions. As
shown in Figure 6A, the two proteins bound independ-
ently to DNA, and the presence of excess HOXD8 in
the binding reaction caused neither significant decrease
in the binding of HOXD9 to the oligonucleotide, nor the
formation of new DNA–protein complexes.

We also tested whether the presence of HOXD8 affects
the dissociation rate of HOXD9 from the HCR target.
The HOXD9 protein was allowed to bind to equilibrium
to the labeled oligonucleotide, in the absence or in the
presence of an equimolar amount of HOXD8. The DNA–
protein complexes were then challenged by addition of a
250-fold excess of unlabeled oligonucleotide and loaded
on a running gel after short time intervals. As shown in
Figure 6B, the off-rate of HOXD9 from the target DNA
was very rapid and was not significantly affected by the
presence of HOXD8.

The inhibitory activity of HOXD8 requires
an amino-terminal effector domain

As described above [Fig. 4], the D8HD mutant (no. 8 in
Fig. 1), containing the entire homeo domain and carboxyl
terminus of HOXD8, had no inhibitory activity on
HOXD9. This suggested that the homeo domain amino
terminus/helix 1 region, although necessary, is not suf-
ficient for the inhibition, which apparently requires the
presence of an additional function located in the amino-
terminal portion of the HOXD8 protein. To test whether
the HOXD8 amino terminus is specifically required for
the inhibitory function, we generated two expression
constructs in which either the amino-terminal portion of
HOXD9 (amino acids 1–273) or the GST protein were
fused to the amino terminus/helix 1 of the HOXD9 or the
HOXD8 homeo domains [GST–D9H1 and GST–D8H1, respectively]. The binding reaction was carried out also in the presence of a 20-fold excess of unlabeled, bacterially made HOXD8 protein [D8 + D8c]. Relative molecular mass standards (M) are in kilodaltons.

Figure 5. Protein–protein interaction in solution between HOXD8 and HOXD9, as analyzed by protein–protein affinity chromatography and SDS-PAGE. (A) Glutathione–Sepharose coupled to GST–HOXD9 homeo domain fusion (GST–D9HD) was batch-adsorbed to 35S-labeled full-length HOXD9 (D9) or HOXD8 proteins (D8), or HOXD8 deletion mutants lacking the homeo domain helix 2/3 (D8ΔH2/3) or the entire homeo domain (D8ΔHD). The same proteins were batch-adsorbed to a control resin coupled to GST only. One microliter of each in vitro translation reaction containing the labeled, input proteins (10 μl) was loaded as a control (C). (B) The same experiment was carried out with a GST fusion containing only the amino terminus/helix 1 of the HOXD9 or the HOXD8 homeo
domains (GST–D9H1 and GST–D8H1, respectively). The binding reaction was carried out also in the presence of a 20-fold excess of unlabeled, bacterially made HOXD8 protein (D8 + D8c). Relative molecular mass standards (M) are in kilodaltons.

D8 D8ΔH2/3 D8ΔHD D8 + D8c
M C GST–D9H1 C GST–D9H1 C GST–D9H1 C GST–D9H1
M C GST–D8H1 C GST–D8H1 C GST–D8H1 C GST–D8H1

Figure 6. Analysis of interaction between HOXD8 and HOXD9 proteins. (A) Autoradiogram of a 5% polyacrylamide gel showing the DNA shift analysis of HOXD8 and HOXD9 proteins. The HOXD9 protein was allowed to bind to equilibrium to the HCR target. The full-length HOXD8 and HOXD9 proteins were expressed in NIH-3T3 cells fused to an amino-terminal, 6-histidine tag and partially purified by affinity chromatography under native conditions (see Materials and methods). The amount of active protein was titrated in both preparations by in vitro binding to a 32-bp oligonucleotide containing two HOX-binding sites located in the 5’ part of HCR (see Materials and methods), followed by a gel-shift assay (Fig. 6A). A fixed amount of active HOXD9 protein was then mixed with increasing amounts of HOXD8, incubated with the HCR-derived oligonucleotide and then analyzed by the same assay. The concentration of HOXD8 and HOXD9 in the binding reactions was in the same order of magnitude as in the GST-affinity-chromatography experiments (see Materials and methods), potentially allowing the establishment of protein–protein interactions. As shown in Figure 6A, the two proteins bound independently to DNA, and the presence of excess HOXD8 in the binding reaction caused neither significant decrease in the binding of HOXD9 to the oligonucleotide, nor the formation of new DNA–protein complexes.

We also tested whether the presence of HOXD8 affects the dissociation rate of HOXD9 from the HCR target. The HOXD9 protein was allowed to bind to equilibrium to the labeled oligonucleotide, in the absence or in the presence of an equimolar amount of HOXD8. The DNA–protein complexes were then challenged by addition of a 250-fold excess of unlabeled oligonucleotide and loaded on a running gel after short time intervals. As shown in Figure 6B, the off-rate of HOXD9 from the target DNA was very rapid and was not significantly affected by the presence of HOXD8.

The inhibitory activity of HOXD8 requires an amino-terminal effector domain

As described above [Fig. 4], the D8HD mutant (no. 8 in
Fig. 1), containing the entire homeo domain and carboxyl
terminus of HOXD8, had no inhibitory activity on
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terminus/helix 1 region, although necessary, is not suf-
ficient for the inhibition, which apparently requires the
presence of an additional function located in the amino-
terminal portion of the HOXD8 protein. To test whether
the HOXD8 amino terminus is specifically required for
the inhibitory function, we generated two expression
constructs in which either the amino-terminal portion of
HOXD9 (amino acids 1–273) or the GST protein were
fused to the amino terminus/helix 1 of the HOXD9 or the
HOXD8 homeo domains (pD9swD8H1 and pGSTD8H1,
respectively, nos. 11 and 13 in Fig. 1). Cotransfection of
these two constructs, and of pGALD8 in which the
HODXD8 amino terminus is replaced by the GAL4 ac-
tivator domain (no. 3 in Fig. 1), resulted in no change in
the HOXD9 activity on pTHCR (Fig. 7, and summary in
Fig. 1). Conversely, an expressor construct in which the
HOXD8 amino terminus (amino acids 1–193) was fused
to the amino terminus/helix 1 of the HOXD9 homeo
domain (pD8swD9H1, no. 12 in Fig. 1) showed the same
inhibitory activity on HOXD9 [Fig. 7A] as the pD8ΔH2/3
construct [Fig. 4]. This indicates that the inhibitory func-
tion of the amino terminus (the effector function) is spe-
cific for HOXD8, whereas the function of the homeo
domain amino terminus/helix 1 region (the protein–pro-
tein interaction function) can be exerted also by the cor-
responding region of the HOXD9 protein.

To narrow down the region in the HOXD8 amino ter-
minus responsible for the inhibitory function, we gen-
erated a number of deletion mutants within this region.
Internal deletions of a polyalanine repeat (amino acids
15–33, see Fig. 8), or of a histidine–proline- and a proline-
rich repeat (amino acids 67–79 and 106–120, see Fig. 8),
represented by the pD8ΔRP1 and pD8ΔRP2/3 construct,
respectively (nos. 14 and 15 in Fig. 1), had no effect on
the inhibitory activity [Fig. 7B]. Moreover, cotransfec-
tion of an expression construct for the HOXC8 protein
HOXD9 +
HOXD8

4). HOXC8 is highly homologous to HOXD8 in the amino-terminal region, although it contains none of the three repeats (see Fig. 8), which are therefore not necessary for the inhibitory activity. Conversely, a deletion of the amino-terminal 117 residues, represented by the pD8Δ[1–117] construct (no. 16 in Fig. 1), caused complete loss of the HOXD8 inhibitory activity (Fig. 7C). Interestingly, the D8Δ[1–117] mutant was also unable to activate the pTCBS reporter construct (Fig. 7D), indicating that both the inhibition and the activation domains of HOXD8 are contained in the amino-terminal 117 residues. A deletion of the first 46 residues [pD8A(1–46), no. 17 in Fig. 1] was instead unable to inhibit the HOXD9 activity (Fig. 7C) while it showed an activity comparable with that of the full-length protein on the pTCBS reporter (Fig. 7D). These results indicate that the HOXD8 inhibitory effector domain can be separated from the activation domain and is located in the amino-terminal 46 residues, the region of highest homology between the three proteins belonging to the paralogous group 8 (see Fig. 8). To strengthen this conclusion, the amino-terminal 46 residues of HOXD8 were swapped with the corresponding region of HOXD9, in the pD8NTD9 construct (no. 18 in Fig. 1). Cotransfection of this construct had no effect on the HOXD9 activity (Fig. 7C), confirming that the HOXD8 amino-terminal 46 residues are specifically required for the inhibitory function. The pD8NTD9 mutant was active on the pTCBS reporter (Fig. 7D), confirming that the HOXD8 activation domain is located between amino acids 46 and 117.

The HOXD8 homeo domain is able to establish protein–protein interactions with several members of the HOX gene family

To test whether the ability to establish protein–protein interactions is unique to HOXD8 and HOXD9 or rather reflects a more general potential of the HOX homeo domain, we used the affinity-chromatography technique to test for interactions between the HOXD8 homeo domain and the products of other HOX genes belonging to different clusters and/or paralogous groups. The in vitro-translated, 35S-labeled products of HOXB3, HOXC4, HOXC5, HOXC8, and HOXD10 genes were batch-absorbed to the GST–D8HD resin or to a GST control resin. Retained proteins ranged between 10% and 30% of the total input. As shown in Figure 9, all tested proteins were able to associate specifically with the GST–D8HD resin, indicating that homeo domain-mediated protein–protein interaction is possible between different members of the HOX gene family.

Discussion

Specificity of target recognition by Hox proteins in vivo depends on the amino terminus/helix 1 region of the homeo domain

A regulatory element located in the HOXD9 upstream genomic region (the HCR sequence), was reported previ-
Specificity of HOX protein function

Figure 7. Characterization of the HOXD8 amino terminal effector domains. 

(A–C) NIH-3T3 cells were cotransfected with 10 μg of either pTS10 luc [C] or pTHCR [HCR] and 5 μg of pD9 (first two bars in each panel), or with 10 μg of pTHCR, 5 μg of pD9, and increasing amounts (1, 5, and 10 μg) of expressor constructs for different HOXD8 amino-terminal deletion mutants or fusion proteins. 

(D) Cells were transfected with 5 μg of either pT109 luc or pTCBS, and 5 μg of pD8 or expressor constructs for two HOXD8 deletion mutants and a HOXD8–HOXD9 fusion protein. Identification of the expression constructs is given in Fig. 1. The total amount of transfected DNA was kept constant by complementation with up to 10 μg of the pSG5 plasmid. One microgram of pRSVpgal was added as an internal standard. Luciferase activity is expressed in arbitrary light units. Bars, mean ± S.E.M. of at least five independent experiments.

ously to mediate both positive and negative regulation in response to Hox proteins. The HOXD9 and HOXD10 proteins are transcriptional activators of the HCR sequence in a transient transfection assay, whereas HOXD8 is inactive although it can bind HCR in an in vitro assay (Zappavigna et al. 1991). HOXD8 is nonetheless active on a different HOX gene promoter (Arcioni et al. 1992) or on a heterologous promoter linked to a multimerized Hox CBS [this paper]. We show that homeo domain swapping between HOXD8 and HOXD9 is sufficient to transfer the target specificity to each other protein, indicating that specific target recognition in vivo depends almost entirely on the homeo domain structure. Therefore, this specificity cannot be predicted by in vitro binding studies, which are either unable to provide appropriate stringency conditions or lack additional/accessory factors favoring DNA–protein recognition in a cell context. The HOXD8 and HOXD9 homeo domains differ in 20 of 60 amino acids, 11 located in the amino terminus/helix 1 region, and 9 in the DNA-binding, helix 2/3 segment [see Fig. 3]. We show that swapping of the amino terminus/helix 1 region is sufficient to provide HOXD8 with the in vivo target specificity of HOXD9. In Drosophila embryos, much of the functional specificity of the Dfd, Ultrabithorax [Ubx], Antp and Sex combs reduced [Scr] homeo domains was reported to be conferred by the amino-terminal region, a flexible arm establishing contacts with the DNA minor groove, and particularly by the amino acids 1, 4, 6, and 7 [Lin and McGinnis 1992; Furukubo-Tokunaga et al. 1993]. These positions are occupied by different residues in the HOXD8 and HOXD9 homeo domains [Fig. 3]. In particular, the K residue at position 4 and the C-P residues at position 6–7 are invariant in all Hox homeo domains belonging to the AbdB-like paralogous groups 9, 10, and 11, and characteristically different from the G and Q–T residues respectively conserved in the homeo domains belonging to the AbdA/ Ubx/Antp-like groups 7 and 8 (see Mavilio 1993). We therefore suggest that these residues might play a role in the appropriate recognition of, and/or interaction with, the HCR target by the HOXD9 and HOXD10 homeo domains in vivo.

The amino terminus/helix 1 region of the Hox homeo domain mediates protein–protein interactions

As reported previously, the HOXD8 protein is able to inhibit the activating function of HOXD9 and HOXD10 on the HCR sequence in a cotransfection assay (Zappavigna et al. 1991). We now show that this activity is DNA binding independent and requires the same amino terminus/helix 1 region of the homeo domain that allows specific target recognition in vivo. Affinity chromatography experiments showed that this sub domain is a protein–protein contact interface, which can mediate direct interactions between Hox proteins in solution. It
of several classes of homeo domain-containing proteins. The yeast MATα1 and MATα2 transcription factors make protein–protein contacts with a cofactor (MCM1) that influences their ability to discriminate between different targets [Smith and Johnson 1992]. Interestingly, interaction between α2 and MCM1 is mediated by the flexible extension of the α2 homeo domain [Vershon and Johnson 1993]. This region contains sequences related to the YPWM motif conserved upstream of all Hox homeo domains belonging to the paralogous groups 1–8 [see Mavilio 1993], although our data indicate that this motif is not sufficient per se to mediate protein–protein contacts, at least in the case of two interacting Hox proteins. The Drosophila POU domain-containing transcription factor Cfl-a can establish high-affinity heterodimeric interaction with a second POU protein, I-POU, which is unable to bind DNA but can prevent Cfl-a from binding to its target. The heterodimeric interaction between the two factors is specifically mediated by the amino-terminal part of the I-POU homeo domain [Treacy et al. 1991, 1992]. It seems therefore likely that the amino-terminal region of different types of homeo domain share the conserved function of mediating selective protein–protein interactions, which might be an essential requirement for regulating their specificity of action as activators, corepressors, or repressors. Our data extend this function to the amino terminus of the homeo domain of the Hox proteins, which were not yet reported to establish, and be functionally regulated by, protein–protein interactions. The Hox homeo domain is potentially able to establish interactions with a number of different partners, as HOXD8 can interact with at least six HOX proteins belonging to at least three different clusters and five paralogous groups.

**Amino-terminal effector domains are required for the function of Hox proteins**

The amino terminus/helix 1 region of the HOXD8 homeo domain, although necessary, is not sufficient for...
inhibition of the HOXD9 activity on HCR, which requires the presence of a second (in turn) nonsufficient domain consisting of the amino-terminal 46 residues. We interpret these findings by proposing that the negative activity of HOXD8 requires the combined presence of a protein–protein interaction function, located in the homeo domain, and an effector function, located in the amino terminus (see model in Fig 10). The former does not necessarily require HOXD8-specific sequences, as it can be exerted also by the corresponding, nonidentical region of HOXD9, whereas the latter is specifically encoded in the HOXD8 amino terminus. The amino-terminal end is extremely conserved among Hox proteins belonging to the paralogous group 8 (see Fig. 8). Interestingly, HOXC8 has the same negative activity of HOXD8 in our system, whereas the product of the mouse Hoxb-8 gene was reported to inhibit the activating function of Hoxb-9 on the N-CAM promoter, a putative Hox gene target [Jones et al. 1992]. Inhibition of Hox gene function seems therefore to be a common characteristics of Hox proteins belonging to the paralogous group 8.

The activating function of HOXD8 on the appropriate target is also mediated by an effector domain, located somewhere between residues 46 and 117 in the protein amino terminus. This region, relatively less conserved among paralogous genes, can be replaced by a different activator domain, such as the yeast GAL4 acidic activator, but not by a nonspecific polypeptide of comparable size. An intact amino-terminal domain is necessary for the activating function of HOXD9 as well. These results show that positive and negative transcriptional activity of Hox proteins are mediated by specific effector domains located in the amino terminus. Based on our data, a model can be therefore proposed in which a general, protein–protein contact interface can be utilized to mediate combinatorial and/or hierarchical functional relationships among Hox gene products, the function of which is exerted by effector domains localized in different, specific regions of the proteins. The biochemical nature of these effector functions is unknown, although it may conceivably involve interactions with effector domains of other Hox proteins, and/or with other still unidentified components of the transcriptional machinery. It will be of interest to test whether these multiple interactions are functionally significant in the translation of the Hox “code” into positional specification in developing embryos.

Materials and methods

Protein expression and reporter plasmids

The pD9 and pD8 expression constructs (no. 1 in Fig. 1) were generated by cloning the complete open reading frame of the HOXD9 and HOXD8 cDNAs [Izpisua-Belmonte et al. 1990; Zappavigna et al. 1991] into the BamHI site of the pSG5 expression vector [Green et al. 1988]. The pGALD8 construct (no. 3) was generated by cloning an EcoRI–BamHI fragment containing the yeast GAL4 activation domain [amino acids 149–196] fused to a BamHI–EcoRI fragment containing the homeo domain and carboxyl terminus of HOXD8 into the EcoRI site of pSG5. Expression vectors for the truncated forms of HOXD8, and the hybrid HOXD8/HOXD9 proteins, were constructed as follows: [No. 4] pD8swD9 was generated by ligating a Asp718 [Klenow-filled]–Eagl, PCR-amplified fragment containing the HOXD9 homeo box to a SacI–AccI [Klenow-filled] fragment encoding the HOXD8 amino terminus and to a SacI–Eagl fragment.
encoding the HOXD8 carboxyl terminus. The resulting con­
struct codes for a protein containing amino acids 1–193 of 
HOXD8 followed by amino acids 274–339 of HOXD9 and 
amino acids 260–289 of HOXD8. [No. 5] pD8swD8 was con­
bstructed by ligation a PCR-generated fragment containing the 
HOXD8 homeo box fused to the 3-amino acid HOXD9 carboxyl 
terminus to the BamHI site 5' to the homeo box in the HOXD9-
coding region. [No. 6] pD8H1D9 was constructed by fusing a 
pD8swD9-derived EcoRI fragment encoding the HOXD8 amino 
terminus and the HOXD9 homeo domain amino terminus/he­
lix 1 to an EcoRI fragment encoding the HOXD8 homeo domain 
helix 2–4 and carboxyl terminus. [No. 7] pD8AH2/3 (represent­
ing a deletion of amino acids 215–289) was obtained by reclon­
ning in pSG5 a pD8 EcoRI fragment extending from a site 5’ to 
the ATG codon to a site at position +637 from the ATG in the 
HOXD8 coding sequence [Izpisua-Belmonte et al. 1990]. A stop 
codon [TAA] located 16-bp 3' from the EcoRI site in pSG5 was 
thereby put in-frame into the deleted HOXD8 protein. [No. 8] 
pD8HD was generated by digestion at two SacI sites located at 
position +3 and +520, resulting in an in-frame internal dele­
tion of amino acids 2–176. [No. 9] pD8AH2/3 was generated by 
cutting the HOXD8-coding sequence at an Acl site located 5' 
to the HOXD8 homeo box, resulting in a deletion of amino acids 
195–289. [No. 10] pD8AH1 (internal deletion of amino acids 
196–214) was generated by ligation of an Acl–BamHI-digested, 
PCR-generated fragment containing the homeo domain second 
and third helices and the carboxyl terminus of the HOXD8 pro­
tein to an EcoRI-Accl fragment encoding the amino terminus of 
the HOXD8 protein. pD8swD8H1 (no. 11) pD8swD9H1 (no. 12) 
were derived from the corresponding full-length constructs by 
recloning the amino-terminal EcoRI fragments in pSG5, as de­
scribed for pD8AH2/3. [No. 13] pGSTD8H1 was generated by 
cloning an EcoRI–BamHI-digested, PCR-amplified fragment 
containing the complete coding region of the GST gene fused to 
the amino terminus/Accl site of pD8, into the EcoRI and 
BamHI sites of pSG5. [No. 14] pD8ARP1 (internal deletion of 
amino acids 15–33) was obtained by deletion of nucleotides 
+43 to +100 through PCR splicing by overlap extension. [No. 15] 
pD8ARP2/3 (internal deletion of amino acids 67–79 and 
106–120) was similarly constructed by deleting nucleotides 
+198 to +268 and +316 to +378. [No. 16] pD8A(1–117) was con­
bstructed by cutting the HOXD8-coding sequence at the 
SmaI site at position +255. The D8A[1–117] protein was trans­
lated starting from the ATG codon at position +352. [No. 17] 
pD8A(1–46) was generated by cutting the pD8 construct at a 
SacI site 3' to the ATG codon, and ligating it to a NotI–BglII 
fragment encoding amino acids 46–289 of the HOXD8 protein. 
[No. 18] pD8NTD9 was obtained by cloning into pSG5 a PCR-
amplified EcoRI–NotI fragment encoding the HOXD9 amino 
terminus (amino acids 1–46) and a NotI–BglII fragment from 
pD8 encoding amino acids 47–289 of HOXD8. All constructions 
including PCR-amplified regions were controlled by DNA se­
quencing.

Production and nuclear localization of proteins produced by 
expression constructs containing partial HOXD8 sequences 
were tested by immunofluorescence and immunoblotting on 
transfected NIH–3T3 cells using a series of monoclonal antibo­
dies directed against the amino terminus or the homeo do­
main of HOXD8 [V. Zappavigna et al., in prep.], and either a goat 
anti-mouse fluorescein-conjugated secondary antibody [Coulter 
Immunology, Hialeah, FL] or 125I-labeled protein A.

pTHCR was constructed as reported previously [Zappavigna 
et al. 1991]. pTCBS was generated by cloning a eightfold mul­
timerized form of the oligonucleotide 5'-TCGAGTCATCTC-
TACATTGACCCAGTCGA-3' into the SalI and XhoI sites of the 
pT109luc luciferase reporter vector [Nordean 1988].

Cell culture, transfection, and luciferase assay

NIH–3T3 cells were maintained in Dulbecco’s modified eagle’s 
medium [DMEM] supplemented with 10% fetal calf serum 
[GIBCO]. Transfections were carried out by standard calcium– 
phosphate precipitation using 10 μg of reporter plasmid, 5 μg of 
expression construct, and 1 μg of pRSV-β-gal as an internal 
control [Bonnerot et al. 1987] per 9-cm dish. Cells were har­
vested 48–60 hr after transfection, lysed with three cycles of 
freezing and thawing, and centrifuged at 4°C in an Eppendorf 
microcentrifuge for 15 min. Luciferase and β-galactosidase 
asays were carried out as described previously [Zappavigna et al. 
1991].

Protein–protein affinity chromatography

GST fusion proteins were produced using the pGEX 
bacterial expression vector system [Pharmacia, Uppsala, Sweden]. 
The pGEXDH9HD construct was obtained by cloning a PCR-gener­
ated, BamHI-digested fragment containing the HOXD9 homeo 
domain into the polylinker of the pGEX vector. pGEXDH9H1 
was generated from pGEXDH9HD by cutting/religating at EcoRI sites 
located within the homeo box and at the 3’ end of the pGEX 
polylinker. pGEXD8HD was constructed by ligating a BamHI 
fragment from pD8HD, containing the HOXD8 homeo domain, 
into the BamHI site of pGEX. pGEXDH9H1 was derived from 
pGEXD8HD by digestion/religation at EcoRI sites located 
within the homeo box and at the 3’ end of the pGEX polylinker. 
Bacteria transformed with pGEX constructs were grown in 100 
ml of Luria broth to an OD of 0.3–0.5, induced with 0.5–1.0 mM 
IPTG, and harvested after 2 hr. Bacterial pellets were resus­
pended in 500 μl of PBS and sonicated. Insoluble material 
was removed by centrifugation and the supernatant incubated 
with glutathione–Sepharose 4B resin [Pharmacia] for 30 min at 4°C 
on a rotating wheel. Beads were washed four times with PBS and 
once with HND buffer [20 mM HEPS at pH 7.4, 50 mM NaCl, 
0.1% NP-40, 5 mM DTT, 10 mg/ml of BSA]. 35S-Labeled pro­
teins were produced in rabbit reticulocyte lysates [Promega] 
from RNAs transcribed in vitro from the T7 promoter present in 
the pGEX expression vector. Ten microliters [packed volume] of 
the resin bearing equal amounts of either GST or the fusion 
proteins (~5 μg, or 0.15 nmol] were batch-adsorbed for 1 hr at 
+4°C on a rotating wheel to 35S-labeled proteins (~45 ng, or 1 
pmol] in 200 μl of HND buffer [final concentrations: ~1 μM for 
the GST fusion proteins and ~5 nm for the labeled proteins]. 
The resin was washed four times in a 100-fold [vol/vol] excess 
of MTPBS [150 mM NaCl, 16 mM Na2HPo4, 4 mM NaH2PO4, 
0.1% NP-40], resuspended in Laemmli sample buffer, and ana­
yzed by SDS-PAGE.

DNA–protein binding studies

A Ncol–BamHI fragment derived from the pTrHisA bacterial 
expression vector [Invitrogen], encoding a 35-amino acid region 
containing a 6-histidine tag, was fused to the amino terminus of 
the HOXD8 and HOXD9 proteins by ligation into the BamHI 
site of pD8 and pD9, to generate the pTHisD8 and pHisD9 eu­
karyotic expressor constructs. These plasmids were transfected 
into NIH–3T3 cells by standard calcium–phosphate precipita­
tion. Cells were harvested after 60 hr, nuclear proteins were 
extracted, and histidine-tagged, native HOXD8 and HOXD9 
proteins purified by affinity-chromatography on a Ni2+–nitrilo- 
triacetic acid [Ni2+–NTA]–agarose resin [Qiagen] according to 
the manufacturer’s specifications.

Binding of purified Hox proteins was analyzed by gel retarda­
tion as described previously [Zappavigna et al. 1991]. Dissozia-
tion rate was determined by adding a 250-fold excess of cold oligonucleotide to the incubation mixture at 0°C, and loading aliquots on a running 4% polyacrylamide gel at short time intervals. Gels were dried and exposed to a Kodak XAR film at −70°C. The double-stranded oligonucleotide used as probe (5'-GGGACACATTAATCTATAATCAAATACACCTC-3') corresponded to the 5' portion of the HCR region [Zappavigna et al. 1991].

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V Zappavigna, D Sartori and F Mavilio

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