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The zebrafish homeobox gene *hox[zf-114]:* primary structure, expression pattern and evolutionary aspects

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ABSTRACT It is gradually becoming accepted that vertebrate homeobox genes, like their counterparts in *Drosophila*, are crucial for normal development of the embryo. Most vertebrate homeoboxes reported so far are related to the *Drosophila Antennapedia* (*Antp*) sequence, and here we describe *hox*[*zf*-114], a novel *Antp*-like homeobox gene from the zebrafish. The sequence of the *hox*[*zf*-114] homeodomain indicates that this gene could be a member of a subfamily defined by the mouse *Hox*-*1.5*/-*2.7*/-*4.1* genes. However, the evolutionary origin of *hox*[*zf*-114] is unclear and, based on the putative protein sequence, we conclude that it is not directly homologous to *Hox*-1.5, *Hox*-2.7 or *Hox*-4.1, or to other known mammalian homeobox genes. Nevertheless, as revealed by *in situ* hybridization, *hox*[*zf*-*114*] exhibits a spatial expression pattern typical for vertebrate *Antp*-like homeobox genes. Transcripts are detected in the posterior hindbrain, where a sharp anterior border of expression is observed, and throughout the spinal cord. The *hox*[*zf*-114] gene is also active in a region that gives rise to the pectoral fins. These findings suggest a role for *hox*[*zf*-114] in anteroposterior patterning of the neural tube and in pectoral fin development.

KEY WORDS: Brachydanio rerio, embryogenesis, homeobox gene, neural development, zebrafish

Introduction

Molecular studies on animal embryogenesis have been greatly facilitated by the discovery of the homeobox. This conserved gene element was originally identified in the fruit fly Drosophila melanogaster (McGinnis et al., 1984a; Scott and Weiner, 1984), but is probably present in all metazoans. In Drosophila, the homeobox is characteristically found in many of the genes regulating pattern formation and differentiation during early development (reviewed in Ingham, 1988; Hayashi and Scott, 1990). Similarly, most of the analyzed vertebrate homeobox genes have temporal and spatial expression patterns that suggest a regulatory function in embryogenesis (reviewed in Holland and Hogan, 1988; Kessel and Gruss, 1990). This function is likely to be mediated through the homeodomain, the precisely defined protein region encoded by the homeobox (Gehring et al., 1990). The homeodomain consists of some 60 amino acids and can bind to DNA in a sequence-specific manner (Desplan et al., 1988; Müller et al., 1988). Thus, it is now generally accepted that homeodomain-containing proteins act as transcription factors (Levine and Hoey, 1988).

Since the homeobox is evolutionarily conserved, similar sequences can be isolated from vertebrates by using probes from, for example, *Drosophila*. Thereby one can identify vertebrate genes potentially having control functions in embryogenesis without first isolating specific developmental mutants. In the mouse, nearly 40 different genes related to the Drosophila Antp sequence have been identified. These so-called Hox genes are organized into four large gene clusters and, within each Hox cluster, individual genes map in the same linear order as their most closely related Drosophila homeotic genes (Boncinelli et al., 1988; Graham et al., 1989). As in the fruit fly, the expression domain of each Hox gene is correlated with its position in the cluster, i.e., the more 3' a gene is located in the complex, the more anterior its transcripts extend in the embryo (Gaunt et al., 1988; Duboule and Dollé, 1989; Graham et al., 1989). However, the homology between Hox genes and Drosophila homeotic genes is even more intriguing: when transfected into Drosophila embryos, vertebrate Hox genes can substitute for at least some of the functions of their counterpart fly genes (Malicki et al., 1990; McGinnis et al., 1990). Moreover, it has recently been shown by mutation analyses that Hox genes, like their relatives in Drosophila, are essential for normal development of the embryo (Chisaka and Capecchi, 1991; Lufkin et al., 1991).

Abbreviations used in this paper: Antp, Antennapedia gene; Bp, base pairs; CNS, central nervous system; kb, kilobase pairs.

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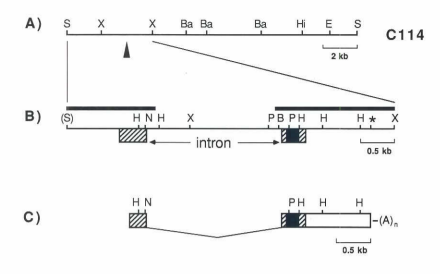


Fig. 1. Organization of the hox[zf-114] gene. (A) Restriction map of the genomic lambda clone C114 containing hox[zf-114]. The homeobox-containing Xbal-fragment is indicated by an arrowhead. The Sall sites are derived from the phage. **(B)** Enlargement of the genomic region where hox[zf-114] is located. The hatched boxes symbolize the open reading frame, of which the homeobox is painted black. The horizontal bars mark the region that has been sequenced and that is listed in Fig. 2. The positions of the poly(A)-signal (indicated by an asterisk) and the intron were deduced by analysis of the cDNA clone. **(C)** Structure of thehox[zf-114] cDNA clone aligned with **B**. Non-coding 3' sequence is indicated by a white box and (A)_n denotes the poly(A)-tail. Restriction enzymes: B, Bg/ll; Ba, BamHl; E, EcoRl; H, Hincll; Hi, Hindlll; N, Ncol; P, Pvull; S, Sal; X, Xbal.

Despite the striking similarities described above and the rapid progress of molecular biology, we still have only limited information about the developmental function of vertebrate Hox genes. Due to the great complexity of the mammalian embryo it will probably be necessary to exploit other systems before a profound understanding of the genetic control of vertebrate development can be reached. Considerable progress has been made with the zebrafish (Brachydanio rerio), which has a number of experimental advantages (Kimmel, 1989). We have initiated a study of zebrafish homeobox genes and have previously reported the cloning and characterization of seven such sequences (Eiken et al., 1987; Fjose et al., 1988a; Njølstad et al., 1988a, 1988b, 1990). For some of the genes, definite mouse homologues could be assigned (Njølstad et al., 1988b, 1990). In these cases, both the gene structure and the expression pattern were highly conserved considering that the fish and mammalian lines of evolution have been separated for more than 300 million years (Benton, 1990).

Here we describe the lambda clone C114, isolated from a zebrafish genomic library during a screening for homeobox genes (Eiken *et al.*, 1987). We present the sequence of the homeobox found in C114 and the molecular structure of the corresponding gene, *hox[zf-114]*. We also show by *in situ* hybridization that this gene seems to be expressed in a restricted anteroposterior domain of the central nervous system (CNS) and in the region where the pectoral fins will form. Finally, we discuss possible evolutionary relations between *hox[zf-114]* and other vertebrate *Hox* genes.

Results

Structural analysis of the lambda clone C114

The screening of a zebrafish genomic library for homeobox sequences (see Materials and Methods) resulted in 30 independent clones. These clones could be classified into 10 groups on the basis of their cross-hybridization patterns with various homeobox probes (not shown). The lambda clone C114 (Fig. 1A) was the single member of one of these groups. The hybridization studies indicated that a 0.56 kb *Sau*3A subclone, which is located within the 3.0 kb *Xbal* fragment shown in Fig. 1A, contained a homeobox and this was confirmed by subcloning, followed by sequence analysis. The

corresponding gene has provisionally been named hox[zf-114] in accordance with the proposed nomenclature for zebrafish genes (Njølstad *et al.*, 1990).

Next, to determine the organization of the *hox*[*z*f-114] gene, we screened a cDNA library with a probe from C114 (see Materials and Methods). Only one positive clone, having a size of 1.55 kb, was obtained (Fig. 1C). Sequence analysis of this cDNA indicated that it was not full length at the 5' end (see below), but a canonical poly(A)-signal and a poly(A)-tail were present at the 3' terminus. Hybridization experiments together with a comparison of genomic and cDNA restriction maps revealed the orientation and structure of the *hox*[*z*f-114] gene depicted in Fig. 1B.

Characterization of the hox[zf-114] gene and predicted protein

The nucleotide sequence of *hox*[*zf*-114], based on the analysis of both genomic and cDNA subclones, is shown in Fig. 2. At a position 47 bp upstream of the homeobox (nucleotide number 2386, Fig. 2), the cDNA and genomic sequences diverge, the latter continuing into a consensus acceptor site for mRNA splicing. The 5'most 203 bp of the cDNA clone are colinear with the genomic sequence from position 385 (adjacent to a consensus donor splice site) and upstream. Thus, an intron is present in the *hox*[*zf*-114]gene, and its size was estimated at 2.0 kb.

In the cDNA clone, the open reading frame containing the homeobox region extends to the ultimate 5' end of the clone (corresponds to position 183 in Fig. 2). The only ATG codons in this frame are located very close to the splice site and would result in homeodomain proteins of unusually small sizes. We therefore conclude that the cDNA clone is not full-length. Consistent with this, analysis of the genomic sequence shows that the open reading frame continues approximately 180 bp upstream of the first nucleotide of the cDNA clone. In the genomic extension, there are two possible initiation codons. The 5'-most we have given the position number 1, the other is located 84 bp downstream (Fig. 2). If we assume that the longest open reading frame is the one translated in vivo, the corresponding hox[zf-114] protein has a length of 250 amino acids. The homeodomain is contained in the Cterminal part of the protein. Hence, it is encoded by the second exon, a common feature of the Hox genes. Common is also the

- / / /	GATCOTTIACTIGIGIGGAATTICCTIAACATTAAGAAACACATTITAGATAGGATGATGATGATGATGATGATGATGATGATGAT
-687	gctattattactaaagtattttattcgcaataaaacatcgctaaaatgtacatcactatttacgtaagattaaaaccggttaaaatacgc
-597	AATCGAAGAACATTGACAAGCGTAAGGAACTGTTGTTAGATCAGCAGAGTGTGAAATGAATAGTTATTTAACCATAAAGCTAAACAAGTC
-507	GATTATTTTTAAATACCTACCACAAGAGGACTGAAAAGTCAGTC
-417	CTGTATTCTTCAATCCATTTAAGCAAAACTAAGACGACGCGATCCCAGACATGTATTTTCTCTGTCCAAAATGGGTCTCTCTC
-327	CCCACCTCACGCTCTCTGAACTCGTGTCAGCACGATGACAGTGTAATGTATGCGCCCTGCTCATGGTTTTTCGTCTGTCCACAGGAAAAT
-237	CACTCGAAATGGCAAAATAGTCGAAGTATGCGGATGAAAACGGAATCACCGTTTGGCTGCTTACATGCAGTTTACAACTCAAACACCGAG
-147	TGCGCAAGGCTATTGGAGGAGGCCACCACGTGACTACATTGTTTACTCTTGTTGGCAAACAATŤAAAGACGCGAGTAAGTGTATTTTTTG
_57	*** MetAsnAsnSerPheHisGluTyrAsnPhe GCTTTCCTACACCTCGAATGAACAATTATATATTTCGGGTGATTAGGACGATTAGAA <u>ATG</u> AACAATAACTCGTTTCACGAATATAATTTT
12 34	AsnSerGluTyrLeuAsnLysIleCysCysAsnLysSerTyrValCysTyrMetGlyGlnHisPheSerProCysAlaLeuGlnGluAsn AACAGTGAATACTTAAACAAAATATGTTGCAATAAATCTTATGTTTGTT
<i>42</i> 124	SerSerThrTrpLysArgHisAlaGluGluSerSerSerGluAsnAspLysGlyAsnValTrpAsnPheGlnAsnLeuSerAsnValGln AGTTCTACGTGGAAACGACATGCTGAAGAAAGTTCATCAGAAAATGATAAAGGCAACGTGTGGAACTTTCAAAAACCTCTCGAATGTCCAA → Start cDNA
72 214	HisProTyrSerPheSerAspAspGlyAsnHisSerLeuAspProAlaAsnAlaLeuGluArgLysLysAlaCysGluLeuSerThrSer CATCCCTATTCATTCTCTGATGATGGCAACCACTCTTTGGACCCTGCGAATGCATTGGAAAGAAA
1 <i>02</i> 304	CysLeuSerThrMetLysTyrProTrpMetArgGluThrHisAlaProThrHisPheSerSerIleAsnAlaMetGluSerG TGCTTATCAACCATQ <mark>AAGTACCCTTGGATGAGA</mark> SAGACACATGCACCCACCACCACTTCAGCTCCATTAACGCCATGGAATC AGgtgaga ga
394	*** *** acacgactttgaagtaaattatgcatattgatttcatgactgtattgcttagaaatacccactaaatttaatcagtcag
334	acacyacticyaagtaaactacyacticgacticacyacticgactacyactaguattaccactactactactactactactactactactacta
484	gtctgggatgagag
2298	$\label{eq:tctgttcactgtctaaaactgtgcagcagtctctgtttgacagatctggtatgcctaaacacttctgtatatctgcatattttctag \texttt{GT}$
<i>130</i> 2388	AspSerLysTyrSerAsnGlyGluAlaValValArgAsnSerSerSerLysArgAlaArgValAlaPheThrSerSerGlnLeuLeuGlu GATTCTAAATACAGCAATGGTGAAGCAGTTGTACGGAATAGCAGGAGTAAACGGGCTCGTGTGGCGTTCACCTCCTCTCAGCTGCTGGAG
160 2478	LeuGluLysGluPheHisPheSerAlaTyrLeuCysArgAsnArgArgLeuGluMetAlaGluLeuLeuLysLeuThrAspArgGlnIle TTAGAGAAGGAGTTTCACTTCAGCGCTTACCTGTGCCGCAACCGGGAG <u>ACTTGAAATGGCTGAGCTGA</u>
190 2568	LysIleTrpPheGlnAsnArgArgMetLysTyrLysLysAspHisLysGluLysSerThrAlaLysSerSerTyrThrTyrLeuGlyThr AAAATCTGGTTCCAAAACCGTCGCATGAAGTATAAGAAGGACCACAAGGAAAAGTCAACGGCAAAGTCCTCTTACACCTACGCGAACA
<i>220</i> 2658	GluAsnGlnProLeuIleIleSerArgSerThrThrAspSerProValProLeuLysPheGlnAsnAsnTyrGluThrProSerMetAsn GAAAACCAGCCATTAATAATTAGCAGAAGCACAACAGATTCTCCGGTGCCATTGAAATTCCAGAATAACTATGAAACACCATCAATGAAC
250 2748	Trp*** TGGTGAAATTACAGAAAAACTGAAAACCGGTGACTTTTGCTTTTAGAACTTATATAATGATAGTAATCTGCATCAACCATAAAGACTGAA
2838	AATGAATGAATGAATGAATGAATGTGTGTGAATGAATGA
2928	GTAACTCTCTCTGACATG <u>ATTTATATTTA</u> AAAACATTGATGATTAATTGCCTCTGGATGATG <u>ATTTA</u> TGTGCAAAGTTAACTTGGAAGAG
3018	TTCAATAGGCAAGCTCGCTCTGAACCAGATTACATGATATCAGCTAGTACATTTCAGAGTTAGTCTCTATTGTGCATAAACTGCTGTAGT
3108	AATATCTTTGCAAAGCACCAAG <u>ATTTA</u> TTGCAACCCTACTTGTGCAAGTTTATCTCTTGGGCACAGAAAATACAATTTTAAAAGGGCATT
3198	CAAGACTTTGAATGGTCATTAACATCCTCTCAACTTTTTAAGTGCACAACATATTTTTTATTACTCTCAAAATGATATGCTACTTTTCCC
3288	AAGGCATGAGCTAAAATGTACAACTTGCAGCGTTGTTCATTATGATGGTCACAATTGTTCCATGTTAAAGCCTATTGACCTGCTCTACTA
3378	TTAACAATGAATTGAATGACTTTTTTTTCTGCTTTACACTGAAAGTAGGTGCGTTTAAGTCTGA <u>ATTTA</u> CAGCATATCTACTGACAAAAT
3468	GAATGAAAGAGCAAAGCAGGATTTTCACAATGCTATATCAATGC <u>ATTTA</u> TAGTGTAATTACTTTACCCATTCATGTACACAAAAGTGTCA
3558	ACATGGCACCTGCAATGTAAAATATGTGCATATTTGTA <u>ATTTA</u> TACCATTTCAATTATAACCTGTCCATATTTGACAAAAAAAA
3648	TTTGAAAATAATAAGTGGTTGTTTTTTTAAACTG <u>ATTTA</u> TTCCCTCTGGATAGTC AATAAA TAATCATTGTAATTCATGTAACTGACAATA
3738	ATGTTGTTTGTTTAGTTAAAACAGGTAATGGTGAGTCTAAATTTAGCTTAAAGGTGCTTCTGTGAATCTTACTTGATACTTTTGAATT
3828	AGGCAATACTGTACTTTGCATTCTTAAAACATAACCTTACATATATGCCTTCCTCCGCCTCCGCCTACCAACCTCGGAACATTAATCATT
3918	TAGACAAGGCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTC

Fig. 2. Genomic sequence (corresponding to Fig. 1B) and a translation of the protein coding region of hox[zf-114]. The two possible initiation codons are underlined and numbering starts at the more upstream position. Asterisks mark stop codons delimiting the open reading frame and in-frame stop codons within the intron. The hexapeptide and homeobox sequences are boxed. A canonical TATA-box upstream of the putative initiation codon and the consensus donor and acceptor splice sites are in bold face letters. For the intron, only the 100 bp closest to the splice sites are listed. In the 3'untranslated region, ATTTAstretches (see text) are underlined. The poly(A)-signal is shown in bold face and the asterisk indicates the site of poly(A)-addition. The sequence of hox[zf-114] appears in the EMBL nucleotide sequence database under the accession number X60095.

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presence of a short conserved sequence in the first exon, close to the splice site. It translates into a hexapeptide of unknown function and occurs in *hox*[*zf*-114], too (boxed in Fig. 2). The N-terminus of the predicted *hox*[*zf*-114] protein, however, differs significantly from typical N-termini of vertebrate *Hox* proteins (Schughart *et al.*, 1988).

TGTGCGCTTCTCTCCAACACACTTCTCTCCCAATACGGGAGCTGAAATACATCTAGA

-777

4008

Since we failed to obtain a full-length cDNA clone, conclusive data about the 5' structure of the *hox*[*zf*-114] gene is lacking. For example, additional upstream exons like those reported by Simeone *et al.* (1988) and Cho *et al.* (1988) cannot be ruled out. But we note that a canonical TATA-box, which might be used for transcription *in vivo*, occurs at position -444 (Fig. 2). Usage of this TATA-box would

result in a processed transcript of about 2.2 kb, which is in good agreement with the major band (estimated at 2.3 kb) detected on a northern blot (see below). However, *in vitro* mapping of transcription start points and isolation of additional cDNA clones are necessary for clarification of how the 5' region of *hox*[*zf*-114] is organized.

In the 3' region of the gene, there is a poly(A)-signal at position 3702, the actual site for poly(A)-addition following 20 bp downstream. This implies that the *hox*[*zf*.114] transcript has an untranslated 3' tail of nearly 1 kb. Long, non-coding 3' regions are often observed in the mRNAs of *Hox* genes and could serve reguŀ

					Reference:
		20	40	61	
	Antp	RKRGRQTYTRYQTLELEKEFHFNRYLTRRF	RRIEIAHALCLTERQIKIWFQ	NRRMKWKKENK	Scott and Weiner (1984)
	Hox-2.5	SRKK-CPKD-	H-V-RL-N-SV	MM-N	Bogarad et al. (1989)
	Hox-2.4	-RK-	VSGV	N	Hart et al. (1987)
	Hox-2.3	YY	T		Hart et al. (1987)
	Hox-2.2	GRYY		S-	Hart et al. (1987)
	Hox-2.1	GA-TA	S	D	Krumlauf et al. (1987)
	Hox-2.6	PS-TAQ-VYY	VS	DH-	Graham et al. (1988)
*	Hox-2.7	SA-TASA-LVC-P-	V-M-NL-N-S	YDQ-	Graham et al. (1988)
	Hox-2.8	SR-L-TANT-LKC-P-	VAL-DV-V	H-RQTQ	Rubock et al. (1990)
	Hox-2.9	PGGL-TNF-TR-LTKS-A-	VAT-E-N-T-V	QRER	Rubock et al. (1990)
*	Hox-1.5	STAP-LVM-P-	V-M-NL-N	YDQ-	McGinnis et al. (1984b)
*	Hox-4.1	SA-TASA-LVFV-P-	VQM-NL-N-S	YDQ-	Lonai et al. (1987)
hox	[zf-114]	SA-VAF-SS-LSAC-N-	L-M-EL-KD	YDH-	this report

Fig. 3. Comparison of the *hox[zf-114]* homeodomain with the homeodomains of all known genes in the mouse *Hox-2* cluster (top) and paralogs of *Hox-2.7* (bottom). The homeodomain of the Drosophila Antp gene is included as a common reference sequence. Hyphens represent amino acid identity as compared to Antp. The hox[zf-114] homeodomain is most related to that of Hox-2.7 and the Hox-2.7 paralogs Hox-1.5 and Hox-4.1 (marked with asterisks). See also Table 1.

latory functions. In this connection, it might be relevant that the untranslated part of the *hox*[*zf*-114] transcript contains several AUUUA motifs (Fig. 2), which have been suggested to influence message stability (Shaw and Kamen, 1986). Such motifs are also found in 3' non-coding regions of some other mRNAs transiently expressed during animal embryogenesis (e.g. Graham *et al.*, 1988; Njølstad *et al.*, 1988b; Nakano *et al.*, 1989).

The hox[zf-114] homeodomain is related to Hox-2.7 paralogs

The Hox complexes are thought to have arisen by duplications of a single ancestral gene cluster (Boncinelli et al., 1988; Graham et al., 1989). This hypothesis is based upon the fact that a given Hox gene is more related to one specific gene (denoted paralog) in each of the other clusters than it is to any gene in its own cluster. It was therefore of interest to see whether hox[zf-114] can be assigned to any subfamily of Hox genes. A comparison with all known genes of the murine Hox-2 complex reveals that the hox[zf-114] homeodomain displays the highest similarity to that of Hox-2.7 (78.7%, Table 1). The similarity is confirmed by relating these two homeodomains to the Antp sequence (Fig. 3). In a set of paralogous genes, the corresponding homeodomains tend to have amino acid substitutions (as compared to Antp) in identical positions and also to have the same substituting amino acids in those positions. Such a relationship is seen between hox[zf-114] and Hox-2.7; of the 21 positions in the hox[zf-114] homeodomain that are different from the Antp sequence, 17 are also different in Hox-2.7. Of the latter, 10 are identical substitutions. The only other Hox genes that show a similar level of identity to hox[zf-114] are the paralogs of Hox-2.7, namely Hox-1.5 and Hox-4.1 (Table 1 and Fig. 3). We therefore conclude that the hox[zf-114] homeobox appears to be a member of the subfamily of homeobox sequences defined by Hox-1.5, -2.7 and -4.1. However, hox[zf-114] is probably not the zebrafish homologue of any of these genes, and we will elaborate on this issue below.

Expression of hox[zf-114] in the neural tube

The zebrafish *hox* genes that we have studied are all expressed during embryogenesis (see, for example, Njølstad *et al.*, 1988a). They are characteristically turned on 11-12 hours after fertilization, around the time when somites start to form and differentiation of the CNS is initiated. We have previously reported the temporal distribution of *hox*[*zf*-114] transcripts in embryos (Fig. 3 in Fjose *et al.*, 1990), and the pattern is similar to that of the other zebrafish *hox* genes. On the northern blot, one major transcript of 2.3 kb was detected, and there were also weaker bands of 2.8, 1.9 and 1.5 kb (Fjose *et al.*, 1990). Multiple transcripts and complex expression patterns are in fact often observed for vertebrate *Hox* genes (Graham *et al.*, 1988; Condie *et al.*, 1990).

Having established that *hox*[*zf*-114] has a temporally restricted expression during embryogenesis, we investigated the spatial

TABLE 1

HOMEODOMAIN HOMOLOGIES

		Homology to hox[zf-114]	Homology to Antp
	Antp	65.6%	
	Hox-2.5	59.0%	67.2%
	Hox-2.4	55.7%	83.6%
	Hox-2.3	63.9%	96.7%
	Hox-2.2	62.3%	93.4%
	Hox-2.1	68.9%	90.2%
	Hox-2.6	67.2%	82.0%
*	Hox-2.7	78.7%	68.9%
	Hox-2.8	65.6%	63.9%
	Hox-2.9	57.4%	59.0%
*	Hox-1.5	75.4%	73.8%
*	Hox-4.1	73.8%	65.6%
	hox[zf-114]	-	65.6%

Amino acid homology (based on 61 residues) between the homeodomain of *hox*[*z*f-114] and those listed in Fig. 3. A comparison has also been performed to the *Antp* sequence. The *hox*[*z*f-114] homeodomain exhibits the closest relationship to the homeodomains encoded by *Hox-1.5, Hox-2.7* and *Hox-4.1* (marked with asterisks).

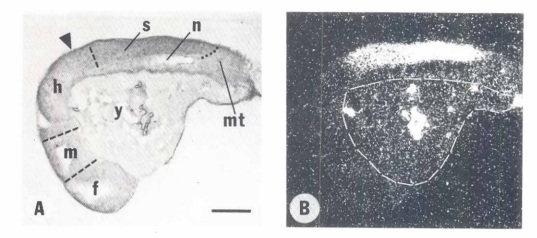


Fig. 4. A parasagittal section of a zebrafish embryo (22 h old) hybridized with a probe from the hox[zf-114] gene. (A) The section in bright-field illumination. The arrowhead points to the anterior limit of hox[zf-114] expression in the posterior hindbrain. The borders between the main regions of the developing CNS are indicated by dashed lines. Abbreviations: f, forebrain; h, hindbrain; m, midbrain; mt, muscle tissue; n, notochord; s, spinal cord; y, yolk sac. (B) Dark-field view of A. Signals are present in the posterior part of the hindbrain and in the spinal cord. The apparent posterior limit of expression is simply due to spinal cord leaving the oblique section and muscle tissue entering it (at the stippled line in A). From other sagittal sections (not shown) and from the cross-sections in Fig. 5, it is clear that expression in the spinal cord continues into the tail region. Scale bar (A and B): 0.2 mm.

distribution of its transcripts. Typical for the Hox genes is that each is active in a particular subdomain of the CNS (reviewed in Holland and Hogan, 1988; Kessel and Gruss, 1990), having a discrete anterior boundary of expression. From in situ hybridization studies performed on sagittal sections of 22 h zebrafish embryos, it is clear that the same holds for hox[zf-114] (Fig. 4). Transcripts are detected in the developing neural tube, and there is a sharp anterior border of expression in the hindbrain, posterior to the otic vesicle. This is confirmed by the horizontal section in Fig. 5A,B. More posterior sections (Fig. 5C-H) reveal that spinal expression continues into the tail region of the embryo. Apparently, hox[zf-114] hybridization signals are uniformly distributed across each transverse section of the spinal cord. In contrast, Graham et al. (1991) observed a dorsoventral restriction of expression in the neural tube for several murine Hox-2 genes. This pattern was developmentally regulated, and further analyses of additional embryonic stages in the zebrafish will be necessary to see whether similar restrictions exist for hox[zf-114].

Expression of hox[zf-114] in the pectoral fin buds

Many *Hox* genes are expressed in the limb buds of mouse and chicken embryos (see Tabin, 1991 for a review). Genes in the *Hox*-4 cluster (previously named *Hox*-5) have been studied most extensively in this regard, and each *Hox*-4 gene shows a unique pattern of limb bud expression, which correlates to its position within the complex. This finding suggests that vertebrate homeobox genes play a key role during limb development. In a previous report, we have shown that antigens related to the *Xenopus* homeoprotein XIHbox1 (probably the zebrafish *hox*-3.3 gene product) are distributed in an anteroposterior gradient in fish pectoral fin buds (Molven *et al.*, 1990), which corresponds to the anterior limb buds of tetrapods. Fig. 6 reveals that in 22 h zebrafish embryos, *hox*[zf-114]-derived transcripts have accumulated in the region where the pectoral fin buds will appear 6-7 hours later. The section in Fig. 6 was cut slightly oblique and it should be noted that there is a

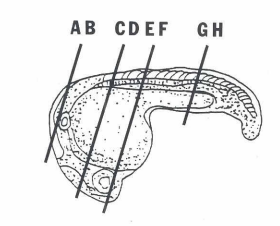
difference in signal intensity between the two sides of the embryo. This indicates that *hox*[*z*f-114] could be expressed as a gradient in the pectoral fin buds, lending support to the hypothesis that this gene is involved in their specification.

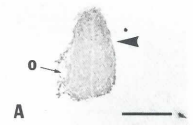
Discussion

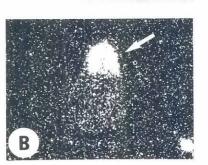
Classification of hox[zf-114]

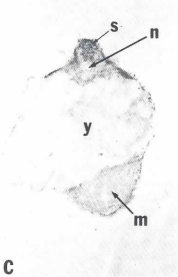
Several lines of evidence suggest that the gene described in this report relates to the growing family of vertebrate *Hox* genes. For example, the genomic organization of *hox*[*zf*-114] points to this conclusion. The open reading frame of the gene spans two exons, with the homeobox located in the second. Close to the splice site, the first exon encodes a conserved hexapeptide. Also, a long untranslated 3' sequence is present in the mRNA. These are all features commonly found in the *Hox* genes. Moreover, the size of the predicted *hox*[*zf*-114] protein is in the range (220-300 amino acids) typical for *Hox* gene products.

Our proposed classification of hox[zf-114] is further strengthened by its temporal and spatial expression pattern. The gene is turned on shortly after gastrulation is finished, and transcripts are detected in the developing CNS, exhibiting a sharp anterior limit of expression. This is highly reminiscent of the pattern observed for the murine Hox genes. However, Hox transcripts are usually detected also in mesodermally-derived tissues of the mouse embryo, in domains posteriorly displaced from the anterior expression borders in the CNS (e.g., Gaunt et al., 1988). We have obtained preliminary evidence that hox[zf-114] is expressed in somitic mesoderm (not shown), but further analyses are necessary for verification and for comparison with the CNS expression domain. Another observation linking hox[zf-114] to the murine Hox genes is that hox[zf-114] transcripts accumulate in the pectoral fin fields; many genes of the Hox-3 and Hox-4 complexes are active in the limb buds of mice (Oliver et al., 1988; Dollé et al., 1989). Since hox[zf-114] shares so many general properties with the vertebrate Hox

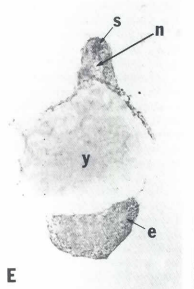




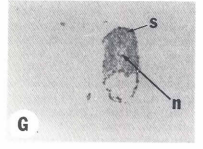


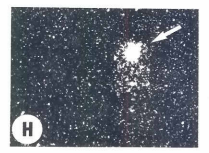












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genes, it is tempting to speculate that *hox*[*zf*-114] is a member of a gene complex. By performing a chromosomal walk in both directions from the C114 clone it should be possible to verify or disprove this assumption.

The anterior expression border of hox[zf-114]

In mouse embryos, it has now been firmly established that the Hox genes are expressed in the embryonic CNS according to their positions in the gene clusters (see Introduction). It still remains to be proven, though, that this is the case also for the zebrafish. In light of the considerable conservation of some zebrafish hox genes as compared to their mammalian counterparts (Njølstad et al., 1988b, 1990), it would not be surprising if the same correlation between anterior borders of expression and chromosomal localization were to apply to fish. An indication that it might be so is obtained by comparing the expression domain of hox[zf-114] with that previously reported for the putative hox-3.3 gene product (Molven et al., 1990). Based on analysis of the homeobox sequences, hox[zf-114] is likely to be positioned more 3' in a cluster than hox-3.3. Consistent with this, the anterior limit of hox[zf-114] expression reported here (posterior hindbrain) lies more anterior than the border of hox-3.3 expression at the 22 h stage (spinal segment number 2).

Wilkinson *et al.* (1989) have performed a detailed investigation of the expression limits of murine *Hox-2* genes in the developing hindbrain. They found that *Hox-2.1*, *-2.6*, *-2.7* and *-2.8* transcripts are distributed in a segment-specific pattern with successive genes having borders at two-segment intervals. Since *hox*[*zf*-114] transcripts extend into the posterior hindbrain, it is pertinent to ask whether the anterior restriction coincides with a segment border. The resolution of our *in situ* hybridization experiment did not allow us to answer this question unambiguously, but if such a relationship exists for *hox*[*zf*-114], the anterior border is at the 7th or 8th hindbrain segment (Ca1 or Ca2, see Hanneman *et al.*, 1988).

Relationships between hox[zf-114] and other Hox genes

Among the *Hox* sequences reported so far, the homeodomain of *hox*[*zf*-114] is most similar to that of *Hox*-2.7 (Table 1 and Fig. 3). Nevertheless, when we compared the whole coding region of *Hox*-2.7 (R. Krumlauf, personal communication) with the open reading frame of *hox*[*zf*-114] no obvious sequence similarity outside the homeodomains was observed. For example, the hexapeptide of *hox*[*zf*-114] (Lys-Tyr-Pro-Trp-Met-Arg) shares only 3 residues with the *Hox*-2.7 hexapeptide. The sizes of the gene products are also different: the protein encoded by *Hox*-2.7 is considerably larger than the predicted *hox*[*zf*-114] protein. We conclude that *hox*[*zf*-114] appears not to be the fish equivalent of *Hox*-2.7.

Could hox[zf-114] instead correspond to any of the Hox-2.7 paralogs (i.e. Hox-1.5, Hox-4.1 or a predicted, but yet unidentified, gene in the Hox-3 cluster)? At least two observations make this idea

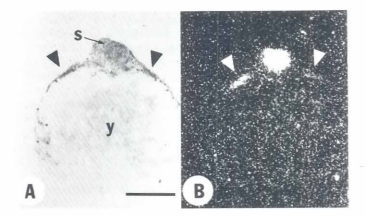


Fig. 6. The *hox*[zf-114] gene is expressed in the developing pectoral fins. (A) A transverse section of a 22 h zebrafish embryo at the level of the pectoral fin fields (arrowheads) hybridized with the hox[zf-114] probe (bright-field illumination). (B) Dark-field view of A. The fin field signals are indicated by white arrowheads, but due to oblique sectioning the signal over the right side of the embryo is barely visible. Abbreviations: s, spinal cord; y, yolk sac. Scale bar (A and B): 0.2 mm.

less attractive. Firstly, the homeodomains of *Hox-1.5*, *Hox-2.7* and *Hox-4.1* have 89-95% mutual similarity, which is substantially higher than the 74-79% identity they exhibit to hox[zf-114]. And secondly, the hexapeptides of the three mouse paralogs are identical (P. Lonai, personal communication), but quite different from the hox[zf-114] hexapeptide (see above).

We have also compared the predicted *hox*[*zf*.114] protein with the sequences of the other *Hox* gene products reported thus far. Except for the homeodomain and the hexapeptide, no regions of obvious similarity were revealed (not shown). It is therefore difficult to draw conclusions about the evolutionary origin of *hox*[*zf*.114], especially since we have no information about a possible cluster containing this gene.

However, a clue might come from the pattern of *hox*[*zf*-114] expression in the neural tube. Paralogous genes in the 3' part of the *Hox* complexes tend to have nearly identical anterior borders of expression in the embryonic CNS (Gaunt *et al.*, 1989; Hunt *et al.*, 1991). *Hox-2.7* transcripts are detected in all hindbrain segments from the 5th and caudally (Wilkinson *et al.*, 1989), while *hox*[*zf*-114] is active from the 7th or 8th segment and caudally. Assuming that there is a one-to-one relationship between the hindbrain segments of mammalian and fish embryos and that the segmental expression pattern of *Hox* genes in the hindbrain is highly conserved, this

Fig. 5. Sections of a zebrafish embryo (22 h old) hybridized with a probe from the hox[zf-114] gene. The positions of the sections are illustrated in the schematic drawing at the top of the figure. Note that hybridization signals (indicated by white arrows) are present both in the posterior hindbrain and in the spinal cord, and that there are no signals in the forebrain/midbrain regions. (A, B) A section through the hindbrain at the level of the otic vesicle. An arrowhead points to the anterior border of expression (posterior is to the top). This section is almost horizontally oriented with respect to the anteroposterior axis. (C, D) A transverse section through the cervical part of the spinal cord (upper) and the midbrain region (lower). (E, F) A transverse section through the thoracic part of the spinal cord (upper) and the eyes/forebrain (lower). (G, H) A transverse section through the tail region. Panels A, C, E, G: bright-field illumination. Panels B, D, F, H: dark-field illumination. Abbreviations: e, eye; m, midbrain; n, notochord; o, otic vesicle; s, spinal cord; y, yolk sac. Scale bar (A-H): 0.2 mm.

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indicates that *hox*[*zf*-114] relates to a slightly more 5' or "posterior" subfamily than *Hox*-2.7 does. Support for this contention comes from analysis of the *hox*[*zf*-114] hexapeptide: it is most similar (5 of 6 residues) to that encoded by *Hox*-1.3 and some other genes located in the middle of the *Hox*-1 and *Hox*-2 complexes (Schughart *et al.*, 1988).

Thus, whereas the homeodomain of *hox*[*z*f-114] hints at a connection to the *Hox*-2.7 subfamily, the hexapeptide and anterior expression border suggest that *hox*[*z*f-114] is a member of one of the more 5' subfamilies. Interestingly, for the murine *Hox*-1.3/-2.1/-3.4 group a paralog remains to be identified in the *Hox*-4 cluster. If it exists, *hox*[*z*f-114] may be the zebrafish counterpart of that gene. A high sequence divergence compared to the other members of the subfamily could then explain why the murine *Hox*-4 member has been difficult to trace by hybridization analysis.

Alternatively, it is not the hexapeptide and expression border, but the homeodomain that reflects the true ancestry of hox[zf-114]. We note that there is a much smaller number of nucleotide differences between the homeoboxes of Hox-1.5, -2.7 and -4.1, than there is within genes of the other Hox subfamilies (Schughart et al., 1989). This observation can be explained by gene conversions (see Lewin, 1990), and it is conceivable that such events took place after the evolutionary lines leading to fish and mammals had separated. Hence, *hox*[*zf*-114] might represent an ancient gene which in higher vertebrates was replaced because of gene conversion. The original function(s) of the hox[zf-114] ancestor could even have been taken over by more than one gene, as additional genes were recruited during the evolution of an increasingly complex body pattern. One should also keep in mind the possibility that hox[zf-114] itself evolved after the divergence of fishes and tetrapods (for example by a duplication event within a cluster), thus being a fish-specific gene. Careful mapping and analysis of all zebrafish hox complexes will clarify both this issue and other questions concerning the evolution of the vertebrate Antp-like genes.

Materials and Methods

Embryos

Zebrafish were maintained and bred essentially as described in Stuart *et al.* (1988). Developmental age is given as 'h', hours after fertilization at 28.5° C, which was the temperature of incubation.

Isolation of hox[zf-114]-containing clones

The construction of a zebrafish genomic library in the EMBL3 vector is described in Eiken *et al.*, 1987. The library was screened under low-stringency hybridization and washing conditions (McGinnis *et al.*, 1984a) with a mixture of homeobox probes from *Drosophila* (*Antp:* McGinnis *et al.*, 1984a; *Sex combs reduced:* Kuroiwa *et al.*, 1985; *engrailed:* Fjose *et al.*, 1985) and Atlantic salmon (S12-B: Fjose *et al.*, 1988b). The screening resulted in 30 independent clones, one of them was the lambda clone C114 which contained the *hox*[*z*f-114] gene reported here.

After confirming by sequence analysis that a homeobox was present in C114, a zebrafish cDNA library (Njølstad *et al.*, 1988b) was screened under high-stringency hybridization and washing conditions (Sambrook *et al.*, 1989) with a 0.56 kb *Sau*3A homeobox-containing fragment from C114. A single, positive clone was isolated.

The restriction fragments used as probes in the above experiments were all purified on agarose gels and labeled with $[\alpha^{-32}P]dCTP$ using nick-translation.

Subcloning and sequencing

Restriction fragments from C114 and from the cDNA clone were subcloned into various pGEM-vectors (Promega Biotec). Double-stranded DNA was

sequenced by the chain termination method using the Sequenase system (USB Corporation) and oligonucleotide primers complementary to the plasmid SP6 and T7 promoters. Where necessary, specific sequencing primers were employed or sets of overlapping clones were generated by digestion with exonuclease III ('Erase-A-Base System', Promega Biotec).

Sectioning and in situ hybridization

The embryos were fixed and sectioned as described in Molven *et al.* (1991). The *hox*[*z*f:114] cDNA fragment (Fig. 1C) was labeled with [α .³⁵S]dATP, and *in situ* hybridization and autoradiography were carried out essentially as described in Njølstad *et al.* (1990). The exposure time was 2 weeks.

Acknowledgments

We thank Michele McDowell for making cryostat sections of zebrafish embryos and Robb Krumlauf and Peter Lonai for communicating data prior to publication. We are also indebted to the Department of Medical Genetics, Haukeland Hospital for providing us with necessary facilities, to Rein Aasland for assistance with the computer analyses and to Chuck Kimmel and Monte Westerfield for valuable comments on the manuscript. A.M., P.R.N. and A.F. were supported by grants and fellowships from the Norwegian research councils NAVF and NFFR. Financial support was also obtained from the Nansen Foundation.

References

- BENTON, M.J. (1990). Phylogeny of the major tetrapod groups: morphological data and divergence dates. J. Mol. Evol. 30: 409-424.
- BOGARAD, L.D., UTSET, M.F., AWGULEWITSCH, A., MIKI, T., HART, C.P. and RUDDLE, F.H. (1989). The developmental expression pattern of a new murine homeobox gene: *Hox-2.5. Dev. Biol.* 133: 537-549.
- BONCINELLI, E., SOMMA, R., ACAMPORA, D., PANNESE, M., D'ESPOSITO, M., FAIELLA, A. and SIMEONE, A. (1988). Organization of human homeobox genes. *Hum. Reprod.* 3: 880-886.
- CHISAKA, O. and CAPECCHI, M.R. (1991). Regionally restricted developmental defects resulting from targeted disruption of the mouse homeobox gene *Hox-1.5*. Nature 350: 473-479.
- CHO, K.W.Y., GOETZ, J., WRIGHT, C.V.E., FRITZ, A., HARDWICKE, J. and DE ROBERTIS, E.M. (1988). Differential utilization of the same reading frame in a *Xenopus* homeobox gene encodes two related proteins sharing the same DNA-binding specificity. *EMBO J.* 7: 2139-2149.
- CONDIE, B.G., HEMMATI-BRIVANLOU, A. and HARLAND, R.M. (1990). Most of the homeobox-containing Xhox 36 transcripts in early *Xenopus* embryos cannot encode a homeodomain protein. *Mol. Cell. Biol.* 10: 3376-3385.
- DESPLAN, C., THEIS, J. and O'FARRELL, P.H. (1988). The sequence specificity of homeodomain-DNA interaction. *Cell* 54: 1081-1090.
- DOLLÉ, P., IZPISUA-BELMONTE, J.-C., FALKENSTEIN, H., RENUCCI, A. and DUBOULE. D. (1989). Coordinate expression of the murine *Hox-5* complex homoeobox-containing genes during limb pattern formation. *Nature* 342: 767-772.
- DUBOULE, D. and DOLLÉ, P. (1989). The structural and functional organization of the murine HOX gene family resembles that of *Drosophila* homeotic genes. *EMBO J. 8*: 1497-1505.
- EIKEN, H.G., NJØLSTAD, P.R., MOLVEN, A. and FJOSE, A. (1987). A zebrafish homeoboxcontaining gene with embryonic transcription. *Biochem. Biophys. Res. Commun.* 149: 1165-1171.
- FJOSE, A., EIKEN, H.G., NJØLSTAD, P.R., MOLVEN, A. and HORDVIK, I. (1988a). A zebrafish engrailed-like homeobox sequence expressed during embryogenesis. *FEBS Lett.* 231: 355-360.
- FJOSE, A., EIKEN, H.G., NJØLSTAD, P.R., MOLVEN, A. and HORDVIK, I. (1990). Homeobox sequences of Atlantic salmon (*Salmo salar*) and zebrafish (*Brachydanio rerio*). *Aquaculture 85:* 51-60.
- FJOSE, A., McGINNIS, W.J. and GEHRING, W.J. (1985). Isolation of a homoeoboxcontaining gene from the *engrailed* locus of *Drosophila* and the spatial distribution of its transcripts. *Nature* 313: 284-289.
- FJOSE, A., MOLVEN, A. and EIKEN, H. G. (1988b). Molecular cloning and characterization of homeobox-containing genes from Atlantic salmon. *Gene 62*: 141-152.
- GAUNT, S.J., KRUMLAUF, R. and DUBOULE, D. (1989). Mouse homeo-genes within a subfamily, *Hox-1.4*, -2.6 and -5.1, display similar anteroposterior domains of expression in the embryo, but show stage- and tissue-dependent differences in their regulation. *Development 107*: 131-141.

- GAUNT, S.J., SHARPE, P.T. and DUBOULE, D. (1988). Spatially restricted domains of homeo-gene transcripts in mouse embryos: relation to a segmented body plan. *Development 104 (Suppl.):* 169-179.
- GEHRING, W.J., MÜLLER, M., AFFOLTER, M., PERCIVAL-SMITH, A., BILLETER, M., QIAN, Y.Q., OTTING, G. and WÜTHRICH, K. (1990). The structure of the homeodomain and its functional implications. *Trends Genet. 6*: 323-329.
- GRAHAM, A., MADEN, M. and KRUMLAUF, R. (1991). The murine Hox-2 genes display spatially and temporally dynamic dorsoventral patterns of expression during central nervous system development. Development 112: 255-264.
- GRAHAM, A., PAPALOPULU, N. and KRUMLAUF, R. (1989). The murine and Drosophila homeobox gene complexes have common features of organization and expression. *Cell* 57: 367-378.
- GRAHAM, A., PAPALOPULU, N., LORIMER, J., McVEY, J.H., TUDDENHAM, E.G.D. and KRUMLAUF, R. (1988). Characterization of a murine homeobox gene, *Hox-2.6*, related to the *Drosophila Deformed* gene. *Genes Dev. 2*: 1424-1438.
- HANNEMAN, E., TREVARROW, B., METCALFE, W.K., KIMMEL, C.B. and WESTERFIELD, M. (1988). Segmental pattern of development in the hindbrain and spinal cord of the zebrafish embryo. *Development* 103: 49-58.
- HART, C.P., FAINSOD, A. and RUDDLE, F.H. (1987). Sequence analysis of the murine Hox-2.2, -2.3 and -2.4 homeoboxes; evolutionary and structural comparisons. Genomics 1: 182-185.
- HAYASHI, S. and SCOTT, M. P. (1990). What determines the specificity of action of Drosophila homeodomain proteins? Cell 63: 883-894.
- HOLLAND, P.W.H. and HOGAN, B.L.M. (1988). Expression of homeo box genes during mouse development: a review. Genes Dev. 2: 773-782.
- HUNT, P., GULISANO, M., COOK, M., SHAM, M., FAIELLA, A., WILKINSON, D., BONCINELLI, E. and KRUMLAUF, R. (1991). A distinct *Hox* code for the branchial region of the vertebrate head. *Nature* 353: 861-864.
- INGHAM, P.W. (1988). The molecular genetics of embryonic pattern formation in Drosophila. Nature 335: 25-34.
- KESSEL, M. and GRUSS, P. (1990). Murine developmental control genes. Science 249: 374-379.
- KIMMEL, C.B. (1989). Genetics and early development of zebrafish. Trends Genet. 5: 283-288.
- KRUMLAUF, R., HOLLAND, P.W.H., McVEY, J.H. and HOGAN, B.L.M. (1987). Developmental and spatial expression of the mouse homeobox gene *Hox-2.1*. *Development* 99: 603-617.
- KUROIWA, A., KLOTER, U., BAUMGARTNER, P. and GEHRING, W.J. (1985). Cloning of the homeotic Sex combs reduced gene in Drosophila and in situ localization of its transcripts. EMBO J. 4: 3757-3764.
- LEVINE, M. and HOEY, T. (1988). Homeobox proteins as sequence-specific transcription factors. Cell 55: 537-540.
- LEWIN, B. (1990). Genes IV. Oxford University Press, Oxford, pp. 515-516.
- LONAI, P., ARMAN, E., CZOSNEK, H., RUDDLE, F.H. and BLATT, C. (1987). New murine homeoboxes: structure, chromosomal assignment, and differential expression in adult erythropolesis. DNA 6: 409-418.
- LUFKIN, T., DIERICH, A., LEMEUR, M., MARK, M. and CHAMBON, P. (1991). Disruption of the *Hox-1.6* homeobox gene results in defects in a region corresponding to its rostral domain of expression. *Cell 66*: 1105-1119.
- McGINNIS, N., KUZIORA, M.A. and McGINNIS, W. (1990). Human Hox-4.2 and Drosophila Deformed encode similar regulatory specificities in Drosophila embryos and larvae. Cell 63: 969-976.
- McGINNIS, W., HART, C.P., GEHRING, W.J. and RUDDLE, F.H. (1984b). Molecular cloning and chromosome mapping of a mouse DNA sequence homologous to homeotic genes in *Drosophila. Cell* 38: 675-680.

- McGINNIS, W., LEVINE, M.S., HAFEN, E., KUROIWA, A. and GEHRING, W.J. (1984a). A conserved DNA sequence in homoeotic genes of the *Drosophila* Antennapedia and bithorax complexes. *Nature* 308: 428-433.
- MALICKI, J., SCHUGHART, K. and McGINNIS, W. (1990). Mouse Hox-2.2 specifies thoracic segmental identity in Drosophila embryos and larvae. Cell 63: 961-967.
- MOLVEN, A., NJØLSTAD, P.R. and FJOSE, A. (1991). Genomic structure and restricted neural expression of the zebrafish wnt-1 (int-1) gene. EMBO J. 10: 799-807.
- MOLVEN, A., WRIGHT, C.V.E., BREMILLER, R., DE ROBERTIS, E.M. and KIMMEL, C.B. (1990). Expression of a homeobox gene product in normal and mutant zebrafish embryos: evolution of the tetrapod body plan. *Development 109:* 279-288.
- MÜLLER, M., AFFOLTER, M., LEUPIN, W., OTTING, G., WÜTHRICH, K. and GEHRING, W.J. (1988). Isolation and sequence-specific DNA binding of the Antennapedia homeodomain. EMBO J. 7: 4299-4304.
- NAKANO, Y., GUERRERO, I., HIDALGO, A., TAYLOR, A., WHITTLE, J.R.S. and INGHAM, P.W. (1989). A protein with several possible membrane-spanning domains encoded by the *Drosophila* segment polarity gene *patched*. *Nature* 341: 508-513.
- NJØLSTAD, P.R., MOLVEN, A., APOLD, J. and FJOSE, A. (1990). The zebrafish homeobox gene hox-2.2: transcription unit, potential regulatory regions and in situ localization of transcripts. EMBO J. 9: 515-524.
- NJØLSTAD, P.R., MOLVEN, A., EIKEN, H.G. and FJOSE, A. (1988a). Structure and neural expression of a zebrafish homeobox sequence. *Gene* 73: 33-46.
- NJØLSTAD, P.R., MOLVEN, A., HORDVIK, I., APOLD, J. and FJOSE, A. (1988b). Primary structure, developmentally regulated expression and potential duplication of the zebrafish homeobox gene ZF-21. *Nucleic Acids Res.* 16: 9097-9111.
- OLIVER, G., WRIGHT, C.V.E., HARDWICKE, J. and DE ROBERTIS, E.M. (1988). A gradient of homeodomain protein in developing forelimbs of *Xenopus* and mouse embryos. *Cell* 55: 1017-1024.
- RUBOCK, M.J., LARIN, Z., COOK, M., PAPALOPULU, N., KRUMLAUF, R. and LEHRACH, H. (1990). A yeast artificial chromosome containing the mouse homeobox cluster *Hox-2. Proc. Natl. Acad. Sci. USA* 87: 4751-4755.
- SAMBROOK, J., FRITSCH, E.F. and MANIATIS, T. (1989). Molecular Cloning. A Laboratory Manual (Second ed.). Cold Spring Harbor Laboratory, Cold Spring Harbor, New York.
- SCHUGHART, K., KAPPEN, C. and RUDDLE, F.H. (1989). Duplication of large genomic regions during the evolution of vertebrate homeobox genes. *Proc. Natl. Acad. Sci.* USA 86: 7067-7071.
- SCHUGHART, K., UTSET, M.F., AWGULEWITSCH, A. and RUDDLE, F.H. (1988). Structure and expression of *Hox-2.2*, a murine homeobox-containing gene. *Proc. Natl. Acad. Sci. USA 85*: 5582-5586.
- SCOTT, M.P. and WEINER, A.J. (1984). Structural relationships among genes that control development: sequence homology between the Antennapedia, Ultrabithorax, and fushi tarazu loci of Drosophila. Proc. Natl. Acad. Sci. USA 81: 4115-4119.
- SHAW, G. and KAMEN, R. (1986). A conserved AU-sequence from the 3' untranslated region of GM-CSF mRNA mediates selective mRNA degradation. *Cell* 46: 659-667.
- SIMEONE, A., PANNESE, M., ACAMPORA, D., D'ESPOSITO, M. and BONCINELLI, E. (1988). At least three human homeoboxes on chromosome 12 belong to the same transcription unit. Nucleic Acids Res. 16: 5379-5390.
- STUART, G.W., McMURRAY, J.V. and WESTERFIELD, M. (1988). Replication, integration and stable germ-line transmission of foreign sequences injected into early zebrafish embryos. *Development 103*: 403-412.
- TABIN, C.J. (1991). Retinoids, homeoboxes, and growth factors: toward molecular models for limb development. Cell 66: 199-217.
- WILKINSON, D.G., BHATT, S., COOK, M., BONCINELLI, E. and KRUMLAUF, R. (1989). Segmental expression of *Hox-2* homeobox-containing genes in the developing mouse hindbrain. *Nature* 341: 405-409.

Accepted for publication: February 1992