Independent Hox-Cluster Duplications in Lampreys

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Abstract.

The analysis of the publicly available Hox gene sequences from the sea lamprey *Petromyzon marinus* provides evidence that the *Hox* clusters in lampreys and other vertebrate species arose from independent duplications. In particular, our analysis supports the hypothesis that the last comman ancestor of agnathans and gnathostomes had only a single *Hox* cluster which was subsequently duplicated independently in the two lineages.

Keywords. Hox clusters, lamprey, phylogenetic footprints

1. Introduction

Hox genes code for homeodomain containing transcription factors which are homologous to the genes in the Drosophila homeotic gene clusters (McGinnis and Krumlauf, 1992; Schubert et al., 1993). There is good evidence that the common ancestor of sharks, bony fish, and tetrapods, had four clusters homologous to the mammalian ones (Holland and Garcia-Fernandez, 1996; Prohaska et al., 2003b). An additional duplication event in the teleost lineage increased the number of distinct clusters to at least 7, e.g. in zebrafish (Amores et al., 1998; Stellwag, 1999).

The agnathan vertebrates, lampreys (*Hyperoartia*) and hagfishes (*Hyperotreti*), as the most primitive extant true vertebrates, occupy a phylogenetically intermediate position between the cephalochordates, such as amphioxus, with a single Hox cluster (Garcia-Fernández and Holland, 1994) and the gnathostomes with four or more clusters. PCR surveys (Pendleton *et al.*, 1993; Sharman and W., 1998) and recent genomic mapping data (Force *et al.*, 2002; Irvine *et al.*, 2002) indicate that lampreys have at least three and possibly four Hox clusters, Fig. 1.

Despite recent efforts, the evolutionary history of the lamprey Hox genes and their relationship with the quadruplicate mammalian Hox clusters is far from being resolved. Irvine $et\ al.\ (2002)$ conclude that they have "insufficient data to determine with confidence the identities and evolutionary histories of the lamprey Hox clusters." Amores $et\ al.\ (1998)$ argue for a two-step duplication scenario, with a duplication of both ancestral agnathan clusters, possibly simultaneously by genome duplication, to produce the four cluster ancestral gnathostome arrangement. Force $et\ al.\ (2002)$ report that "in general, the lamprey Hox genes do not appear to be orthologues of specific Hox genes in gnathostomes" and conclude that the most likely scenario is one genome duplication in the vertebrate ancestor producing a HoxAB and a HoxCD cluster with subsequent divergence of the agnathan and gnathostome lineages and independent subsequent duplications in each linage. Ample evidence from other gene families (Escriva $et\ al.\ 2002$), including Dlx (Neidert $et\ al.\ 2001$) and Otx (Germot $et\ al.\ 2001$) confirms at least one independent duplication in the agnathan and

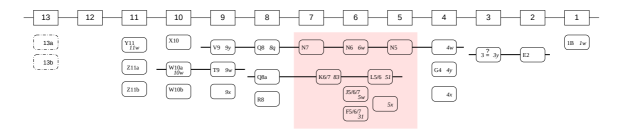


Figure 1. Petromyzon marinus Hox clusters. Summarized from (Force et al., 2002), Fig.1, and (Irvine et al., 2002), Fig.1 and Table 1. Hox13 genes identified in the PCR survey (Force et al., 2002) but for which no cDNA or cosmid was reported in (Force et al., 2002; Irvine et al., 2002) are indicated by dashed boxes. The corresponding sequences are not available. Physical linkage is indicated by a line. The sequences of paralog groups 5, 6, and 7 are insufficient to resolve their mutual relationships, and are therefore excluded from further analysis.

the gnathostome lineages. In this letter we report on a re-evaluation of the publicly available lamprey Hox sequences.

2. Materials and Methods

The available lamprey *Hox* sequences are compiled (together with their accession numbers) in Table 1 in the Appendix. Only short sequences of the homeobox region are available in almost all cases. In contrast to the previous studies we use the nucleic acid sequences rather than the sequences of the *Hox* proteins because of the weak phylogenetic signals in the short and highly conserved amino acid sequences. The sequence from the PCR survey of *Lampetra planeri* (Sharman and W., 1998) are much shorter (82nt) than the *Petromyzon marinus* sequence reported by Pendleton *et al.* (1993) (180nt) and Irvine *et al.* (2002) (240nt). In almost all cases it was possible to identify the homology between the *Lampetra planeri* sequences and their *Petromyzon marinus* counterparts, see Table 1. We therefore use the data from Irvine *et al.* (2002) where possible.

Canonical split decomposition (Bandelt and Dress, 1992), as implemented in the splitstree package (version 3.1) by Huson (1998), is used for the reconstruction of the phylogeny. The split-based methods are particularly suitable for our purposes because they are known to be very conservative in that they tend to produce multifurcations rather than poorly supported edges (Semple and Steel, 2003). For comparison we compute exact maximum parsimony trees using the program dnapenny which is part of the phylip package (Felsenstein, 1989). We use a variety of Hox genes from mammals (Homo sapiens and Rattus norvegicus), shark (Heterodontus francisci), coelacanth (Latimeria menadoensis), and amphioxus (Branchiostoma floridae) for phylogeny reconstruction. All sequences were downloaded from genbank. Alignments were constructed using dialign (Morgenstern, 1999). Since split-based methods tend to lose resolution with increasing number of taxa we use different combinations of lamprey and sequences from other taxa instead of using all sequences together.

An independent line of evidence is derived from the analysis of conserved non-coding DNA. The 30kb PAC clone Pm18 containing the HoxW10a region of Petromyzon marinus was sequenced by Irvine et al. (2002), accession number AF464190. Here we use the tracker program (Prohaska et al., 2003a) to search for phylogenetic footprints in the non-coding parts of this sequence by comparing it with the corresponding regions of the publicly available sequences of human, fugu (Takifugu rubripes, sequences obtained from the JGI database¹, release 3.0), and shark Hox clusters. In the case of the HoxB clusters, which lack Hox-10, Hox-11 and Hox-12 gene, we use the complete inter-genic region from Hox-13 to Hox-9 for the tracker run. The output is then restricted to the region between the first and the last footprint that the lamprey sequence shares with another cluster to account for the fact that Pm18 does not span the entire range to the neighboring genes.

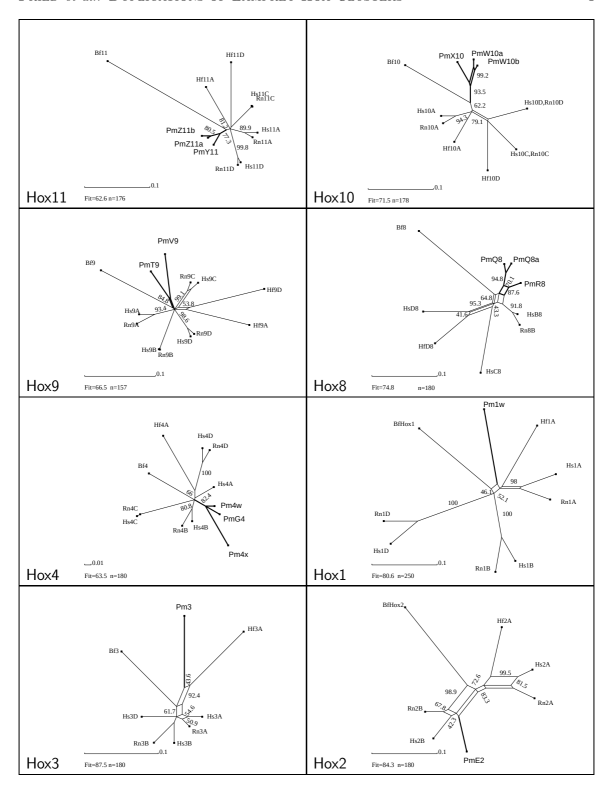


Figure 2. Buneman graphs of the homeobox sequences for paralog groups 1, 2, 3, 4, 8, 9, 10, and 11. We show here the comparison with Human, rat, shark, and amphioxus sequences. Using Teleost fish or coelacanth sequences instead of mammalian data yield qualitatively the same results (data not shown).

3. Results

Only the paralog groups 1, 2, 3, 4, 8, 9, 10, and 11 could be used for our purposes because (i) only a single short *Hox-13* sequence from *Lampetra planeri* was found in the databanks, (ii) there does not seem to be a *Hox-12* gene at all in lampreys, and (iii) the available sequences are too short and too conserved to distinguish unambigously between members of the paralog groups 5, 6, and 7, see also (Force *et al.*, 2002; Irvine *et al.*, 2002).

The comparison of mammalian, shark, lamprey, and amphioxus sequences for a given paralog group presents a striking pattern. We find that the lamprey sequences cluster together outside the gnathostome Hox sequences for paralog groups 11, 10, 9, 8, and 4 according to the split decomposition analysis, Fig. 2. Paralog group 1 is at least consistent with this picture. The single paralog group 3 sequence shows affinity with the shark HoxA sequence but is well separated from the mammalian HoxA-3 genes in the split data. The PmE2 sequence, which is physically linked to Pm3, is more similar to the mammalian HoxB-2 genes. Replacing the rat sequences by coelacanth sequences from the work of Koh $et\ al.\ (2003)$ yields very similar results (data not shown).

The same picture is obtained from maximally parsimonious trees, see Table 2 in the Appendix, for groups 11, 10, 9, and 8. In contrast to the split decomposition method, the best trees for both paralog group 3 and 2 place the lamprey and amphioxus sequences together and as outgroup to the gnathostome clusters. Furthermore, the single Hox-13 sequence of $Lampetra\ planeri$ reported by Sharman and W. (1998) branches outside the other vertebrate genes. Paralog group 1 yields one tree that shows the 1w sequence outside the mammalian cluster and two alternative trees placing 1w with mammalian A clusters. In paralog group 4 the lamprey sequences also lie outside the mammalian clusters but form two separate branches. In no case do we find a clear assignment of the lamprey clusters to either a single or a pair of mammalian and/or fish clusters.

At present the genomic context of only a single lamprey Hox gene, Hox-W10a from $Petromyzon\ marinus$, has been published. Irvine $et\ al.\ (2002)$ report footprint clusters shared with both HoxA and HoxC clusters. The footprint cliques detected by the tracker program in a comparison with Fugu, Shark, Human, and Ciona Hox clusters are summarized in Table 3 in the Appendix. Non-colinear cliques have been removed because they are most likely not homologous (Prohaska $et\ al.\ 2003a$). There is no clear evidence that the non-coding part of the Pm18 sequence is more closely related to either a particular single gnathostome cluster or pair of clusters. The total length of available footprints is unfortunately insufficient for an independent reconstruction of the phylogeny. The most significant footprint cliques are those shared with the HoxA and HoxC clusters, in particular, and an element designated pp that is most likely the proximal promotor of the Hox-10 genes and also appears in the HoxD clusters. The elements A1, A2, C1, and C3 are described already in the work of Irvine $et\ al.\ (2002)$. Both A1, and A2 were also detected in comparisons of HoxA clusters only by Chiu $et\ al.\ (2002)$. It is interesting to note that both A2 and the C1, C2 motifs

¹http://genome.jgi-psf.org/fugu6/fugu6.home.html

also have their counterparts in the Human HoxB cluster, even though it lacks the HoxB-10 gene.

4. Discussion

The re-evaluation of the available lamprey hox genes strongly supports an independent origin of the three (or four) lamprey Hox clusters and suggest that the common ancestor of agnathans and gnathostomes had only a single Hox cluster. This is consistent with the Dlx gene phylogeny described by Neidert $et\ al.\ (2001)$. These authors proposed that a tandem duplication of an ancestral Dlx gene predated the divergence of lampreys from gnathostomes, which was then followed by independent chromosomal or genome duplications and gene loss in each lineage. Our evaluation of the Hox clusters supports this hypothesis. Similar patterns have been reported for other developmentally important gene families. The neural crest marker AP-2, for which no duplicates have been found in lampreys, also fails to group with any one gnathostome AP-2 isoform (Meulemans and Bronner-Fraser, 2002). Consistent with an independent duplication history it is impossible to assign any one of the lamprey (and hagfish) Otx sequences to one of the three classes identified in gnathostomes (Germot $et\ al.$, 2001).

The phylogenetic signal in the Hox clusters is not as strong as one would like so that a definitive result will have to await more complete sequencing. This will in particular allow the unambiguous identification of the genes of paralog group 5, 6, and 7, and their use as additional phylogenetic information. At present, at least, the publicly available sequence information does not contain evidence for a Hox-cluster duplication preceding our common ancestor with the lampreys.

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Appendix (or Supplement)

Table 1. Lamprey Hox sequences used in this study.

Hox			etromyzon m	narinus			empetra planeri		
		genomic cl			PCR :	surveys			
	Irvine	Force	Acc. No.	Pendleton	Acc. No.	Sharman	Acc. No.		
13						Lp13A	AF044814		
11	Y11	11w	AF410923	11.1					
	Z11a		AF410924	11.8					
	Z11b		AF410925						
				11.6					
10	X10	Hx13(9)	AF410922	10x	L14900				
	W10a	10w	AF410920	10w	L14895	Lp10B	AF044812		
	W10b		AF410921						
9	V9	9y	AF410919	9v	L14889	Lp9A	AF044809		
				9s	L14911				
	Т9	9w	AF410918	9t	L14894	Lp9B	AF044810		
		9x		9u	L14910	Lp9C	AF044811		
8	R8		AF035588	8r		Lp8A	AF044807		
	Q8		AF035591						
	Q8a		AF035589	8q	L14901	Lp8B	AF044808		
4	G4	4y	AF410911	4g	L14912				
		4w	AF434666	4n	L14896	? Lp4-7B	AF044803		
		4x	AY056469	41	L14891	? Lp4-7A	AF044802		
						? Lp4-7E	AF044806		
				(4h	L14909)				
3	3	3y	AF410909			Lp3A	AF044801		
2	E2		AF410908	2e		Lp2A	AF044800		
1	1B	1w	AF434665	1b	L14902	Lp1B	AF044798		
				1a	L14893	Lp1A	AF044797		
				1c	L14908	Lp1C	AF044799		
				(1d	L14904)				

The sequence shown in parentheses are not included because we could not confirm their assignment to a paralog group based on their nucleic acid sequence.

Table 2. Maximum parsimony trees of the homeobox sequences obtained with the program dnapenny from the phylip package Felsenstein (1989). Petromyzon marinus, **Pm**, sequences are indicated in bold. For Hox-13 we used the short Lampetra planeri, **Lp** sequence. Grey boxes indicated that all available lamprey paralogs form a subtree, dark gray boxes are used when all lamprey and the amphioxus sequence are separated from the vertebrate hox clusters. Horizontal lines indicate the two exons of the Hox-10 in the HoxA, HoxB, and HoxD, as well as the Ciona Hox cluster Dehal et al. (2002).

Hox	Maximum parsimony tree
13	$(\mathbf{Lp13}, (((\mathit{Bf13}, (RnC13, HsC13)), ((RnD13, HsD13), ((RnA13, HsA13), (HfD13, HfA13)))), (HsB13, RnB13)))$
	$(\mathbf{Lp13}, (((\mathit{Bf13}, (RnC13, HsC13)), ((RnD13, HsD13), (HfD13, ((RnA13, HsA13), HfA13)))), (HsB13, RnB13)))$
11	$(\ (Bf11, (PmY11, (PmZ11a, PmZ11b))), (((Hf11D, (Hs11C, Rn11C)), ((Rn11A, Hs11A), Hf11A)), (Rn11D, Hs11D)))$
10	$(\ (Bf10, (PmX10, (PmW10b, PmW10a))), ((Rn10A, Hs10A), (Hf10A, ((Rn10C, Hs10C), ((Hs10D, Rn10D), Hf10D)))))))$
9	((BfHox9, (HoxT9, HoxV9)), (((Hs9D, Rn9D), ((((Rn9A, Hs9A), Hf9A), (Hs9B, Rn9B)), ((Hs9C, Rn9C), Hf9D))))))
	$((\mathit{BfHox9}, (\mathbf{HoxT9}, \mathbf{HoxV9})), (((((Hs9D, Rn9D), ((Hs9C, Rn9C), Hf9D)), (Hf9A, (Hs9B, Rn9B))), (Rn9A, Hs9A))))$
8	$((Bf8, (\mathbf{PmQ8}, (\mathbf{PmR8}, \mathbf{PmQ8a}))), ((HsC8, (HsB8, Rn8B)), (HfD8, HsD8)))$
4	$((\mathit{Bf4},\mathbf{PmG4}),(((\mathbf{Pm4x},\mathbf{Pm4w}),(\mathrm{Hs4A},(\mathrm{Hf4A},(\mathrm{Rn4D},\mathrm{Hs4D})))),(\mathrm{Hs4B},\mathrm{Rn4B})),(\mathrm{Hs4C},\mathrm{Rn4C}))$
	$((\mathit{Bf4},\mathbf{PmG4}),((\mathbf{Pm4x},\mathbf{Pm4w}),((\mathrm{Hs4B},\mathrm{Rn4B}),(\mathrm{Hs4A},(\mathrm{Hf4A},(\mathrm{Rn4D},\mathrm{Hs4D}))))),(\mathrm{Hs4C},\mathrm{Rn4C}))$
3	(Bf3,Pm3),(Hf3A,(Hs3D,((Hs3A,Rn3A),(Hs3B,Rn3B)))))
2	$((Bf2,\mathbf{PmE2}),(((Hf2A,(Rn2A,Hs2A)),(Rn2B,Hs2B))))$
	$((Bf2,\mathbf{PmE2}),((((Hf2A,(Rn2A,Hs2A)),Hs2B),Rn2B)))$
	((Bf1, Pm1w), ((((Hs1D, Rn1D), Hf1A), (Rn1A, Hs1A)), (Hs1B, Rn1B)))
1	$(\mathit{Bf1},(((Hs1D,Rn1D),Hf1A),((Hs1B,Rn1B),((Rn1A,Hs1A),\mathbf{Pm1w}))))$
	$(\mathit{Bf1},(((Hs1D,Rn1D),(Hs1B,Rn1B)),(Hf1A,((Rn1A,Hs1A),\mathbf{Pm1w}))))$

Table 3. Summary of co-linear footprint cliques produced by the tracker program in the range of the *Petromyzon marinus* Pm18 sequence. Hs *Homo Sapiens*, Hf *Heterodontus fransisci*, Tr *Takifugu rubripes*, Ci *Ciona intestinalis*. Numbers in parentheses are non-colinear with the footprints in this species. The last column marks previously described footprints. *pp* is the proximal promotor of the *Hox-10* gene, numbers in sans serif font are cliques listed in Prohaska *et al.* (2003a) for a comparison of *HoxA* clusters. PFC, "phylogenetic footprint cluster", names from Chiu *et al.* (2002) are given in normal text font.

#	PMa	a	HsA	1	HsC	7	HsD		HfA		HfT)	TrA	a	TrA	b	TrD		TrC	a	Ci		TrBa	ı	$_{\mathrm{HsB}}$	Remark
69	1150	85	8425	50	9641	88	7528	161	11071	223	8404	162	4984	94	3360	139	8112	126								<i>pp</i> , 42
124																	9229	58	7258	69						
70	1535	48																	10137	42						
71					11172	36																			6867 36	
73	2977	50			12879	49																			39418 55	C1
77	9139	35			12879	70																			39418 47	C2
74	5107	52																							39793 52	
75	5671	44									10460	44														
76	6509	30																							39919 30	
96			9475	37																			6722	37		
106					13364	73	8657	76																		
80											11114	37											15608	37		
125							10744	52									10133	50								
83											12657	27													59050 27	
88											13213	26											19306	26		
90	10538	33									15207	33														
93	15694	23																			22400	23				
94	17246	19																							60196 19	
95	17946	43																			22578	43				
98									12558	29															63422 29	
99			12127	39					14376	43																10-9a, 43
100			12248	24					14522	24																10-9a, 44
101	21911	64	12292	187					14566	189			7525	108												<i>A1</i> , 10-9a , 45
109									15213	42															60500 42	
105					13293	36																			78949 36	
102	23635	27																					22252	27		
103	25443	21																							80312 21	
104	26904	26																					25684	26		
81																			19399	39					82476 39	
108					15006	169													19761	170						
110	27436	105	13224	116	16328	105			15518	116			7896	99					20562	106			(18814)	(83)	94203 98	<i>A2</i> , 10-9b , 46
85													8219	11	6226	10							,			
86													8438	82	6389	78										
92																					(11940)	(51)	26975	26	98721 29	
122			14160	49					16310	77			8304	99	6287	73										10-9c , 48
126			15372	59			10890	63																	1	