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ParaHox Cluster Evolution — Hagfish and Beyond

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The ParaHox genes comprise three Hox-related homeobox gene families, found throughout the animals. They were first discovered in the invertebrate chordate amphioxus, where they are tightly clustered. In this paper we carry out a comparative review of ParaHox gene cluster organization among the deuterostomes, and discuss how the recently published hagfish ParaHox clusters fit into current theories about the evolution of this group of genes.

Key words: hagfish, cyclostome, ParaHox, homeobox cluster

INTRODUCTION TO THE PARAHOX GENE CLUSTER

The ParaHox gene cluster was first discovered in the cephalochordate amphioxus (Brooke et al., 1998), and is composed of members of three Hox-related homeobox gene families: Gsx, Xlox, and Cdx. Gsx is most similar to the anterior Hox genes (insect lab, pb), Xlox to group 3 Hox genes (insect zen), and Cdx to the posterior Hox genes (insect Abd-B). Only one member of each ParaHox gene family is found in amphioxus, and these are located in a single cluster with Gsx adjacent to Xlox in the same orientation, followed by Cdx on the opposite strand, the cluster spanning about 56kb (Ferrier et al., 2005) (Table 1). There is no doubt that the clustered organization of these genes as seen in amphioxus is ancestral; the common origin of the Hox and ParaHox gene clusters from a hypothetical ancestral “protoHox” cluster has been discussed extensively elsewhere (e.g., Brooke et al., 1998; Holland, 2001; Garcia-Fernandez, 2005; Chourrout et al., 2006) and will not be discussed in detail here. However, the protoHox hypothesis has also led to other parallels being drawn between the organization and control of the Hox and ParaHox clusters.

The Hox genes are recognised for their spatial and temporal colinearity; that is, the order of expression both during development and along the body axis echoes the order of genes within the cluster. In amphioxus, Gsx is expressed in the cerebral vesicle, Xlox is expressed transiently in the neural tube and in the presumptive gut, and Cdx is expressed in the posterior neural tube and gut. Thus the order of expression sites along the body axis echoes the order of ParaHox genes within the cluster and may be described as spatial colinearity (Brooke et al., 1998). It has been suggested that this expression pattern is modified from an ancestral spatial colinearity of gut expression present at the genesis of the ParaHox cluster (Holland, 2001). Furthermore there is evidence of reverse temporal colinearity, with Cdx expressed first, followed by Xlox and finally Gsx. But with only three genes in a cluster, there are only six possible orders of expression, two of which will be co-linear with the gene order. Could the colinearity simply have occurred by chance?

The cephalochordates (amphioxus) were long thought to be the closest invertebrate relatives of the vertebrates; however, recent studies have revealed that the urochordates form a sister group to the vertebrates (Delsuc et al., 2006; Bourlat et al., 2006; Vienne and Pontarotti, 2006). Amphioxus remains a useful model for its apparently archetypal genomic organization and gene complement (Garcia-Fernandez and Holland, 1994), but the genomes of other deuterostome animals may also reveal much about the evolution of ParaHox gene clustering and expression. For example, in the invertebrate urochordate Ciona intestinalis, surveys of the ParaHox genes indicated that their organization was at the very least much more dispersed than that in amphioxus, and that they may not be chromosomally linked (Ferrier and Holland, 2002; Wada et al., 2003). The same is currently true for ParaHox orthologs in other urochordate genomes: Ciona savignyi (CSAV 2.0 at http://wwwensembl.org) and Oikopleura dioica (V3, http://www.genoscope.cns.fr). However, examination of version 2.0 of the Ciona intestinalis whole genome shotgun (http://www.ensembl.org), in which scaffolds have been mapped to chromosome arms, shows that Gsx is on chromosome 2q, while Xlox and Cdx are found about 250 kb apart on chromosome 14q. The intervening region between Xlox and Cdx contains many predicted or novel genes, transposable elements, and at least one known gene (Ci-Orphan Fox-1, UniProt ID Q4H319). Whether this genomic organization can be termed a cluster seems rather doubtful.

In Ciona intestinalis, the first ParaHox gene to be expressed is Cdx, in the posterior epidermis and nervous system of the late gastrula (Imai et al., 2004). This is followed by Gsx in the sensory vesicle precursor during the neurula to tailbud stages (Hudson and Lemaire, 2001; Imai et al., 2004), followed by extremely weak expression of Xlox (IPF) in muscle cells in the tailbud stage (Corrado et al., 2001; Imai et al., 2004). Thus the presumed ancestral timings of expression have been rearranged, perhaps having some connection with the dispersal of the cluster (Ferrier and Minguillon, 2003).

In the genomic sequence of the echinoderm Strongylo-
**Table 1.** Gene lengths and intergenic distances in deuterostome ParaHox clusters, expanded and adapted from Ferrier et al. (2005). Lengths are given in base pairs, with the percentage of the total cluster shown in parentheses. The total length of the cluster is measured from the Gsx ATG start site to the Cdx ATG start site. Inferred gene lengths are measured from the ATG start site to the stop codon.

<table>
<thead>
<tr>
<th>Homo sapiens</th>
<th>Mus musculus</th>
<th>Monodelphis domestica</th>
<th>Xenopus tropicalis</th>
<th>Amia calva</th>
<th>Eptatretus burgeri</th>
<th>Branchiostoma floridae</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total length of ParaHox cluster</strong></td>
<td>176,315</td>
<td>118,471</td>
<td>197,041</td>
<td>113,674</td>
<td>34,661</td>
<td>57,537</td>
</tr>
<tr>
<td><strong>Gsx gene length</strong></td>
<td>1,257 (0.7%)</td>
<td>1,484 (1.3%)</td>
<td>1,126 (0.6%)</td>
<td>2,021 (1.0%)</td>
<td>1,002 (2.9%)</td>
<td>1,875 (3.3%)</td>
</tr>
<tr>
<td><strong>Intergenic distance Gsx-Xlox</strong></td>
<td>126,190 (71.6%)</td>
<td>80,244 (67.7%)</td>
<td>145,784 (74.0%)</td>
<td>58,054 (51.1%)</td>
<td>16,075 (46.4%)</td>
<td>19,602 (34.1%)</td>
</tr>
<tr>
<td><strong>Xlox gene length</strong></td>
<td>4,560 (2.6%)</td>
<td>4,585 (3.9%)</td>
<td>5,498 (2.8%)</td>
<td>24,844 (21.9%)</td>
<td>6,824 (19.7%)</td>
<td>23,122 (20.3%)</td>
</tr>
<tr>
<td><strong>Intergenic distance Xlox-Cdx</strong></td>
<td>38,416 (21.8%)</td>
<td>27,017 (22.8%)</td>
<td>38,906 (19.7%)</td>
<td>23,122 (20.3%)</td>
<td>7,116 (20.5%)</td>
<td>6,947 (12.4%)</td>
</tr>
<tr>
<td><strong>Cdx gene length</strong></td>
<td>5,892 (3.3%)</td>
<td>5,141 (4.3%)</td>
<td>5,756 (2.9%)</td>
<td>5,630 (5.0%)</td>
<td>3,644 (10.6%)</td>
<td>19,602 (34.1%)</td>
</tr>
</tbody>
</table>

**centrotus purpuratus,** there is a single member of each ParaHox gene family, and again each is dispersed onto a different scaffold with no current evidence for their linkage. Intriguingly, despite this lack of clustering, qPCR reveals that Gsx is expressed first, followed by Xlox (known as Splox) and finally Cdx (Arnone et al., 2006). This would be colinear with the presumed ancestral cluster organization as typified by amphioxus, despite being exactly opposite to the amphioxus order of expression. Furthermore, the domains of expression are also colinear with this order of expression and with the amphioxus genomic organization. Very little data are available for other invertebrate deuterostomes. In the hemichordate Ptychodera flava, four ParaHox genes have been sequenced (Gsx, Cdx, and two Xlox [Lox] genes) but their genomic organization and expression are unknown (Peterson, 2004). Further analysis of the hemichordate lineage awaits assembly of the Saccoglossus kowalevskii genome project currently underway.

In the invertebrate deuterostomes, therefore, it appears that tight clustering of the ParaHox genes may be the exception rather than the norm. While the presumed ancestral order of expression seems to be possible even in the absence of a tight cluster in *S. purpuratus,* in other organisms such as *Ciona intestinalis,* dispersal of the cluster may have occurred in concert with an uncoupling of colinear expression. There are no intervening genes in either the Hox or ParaHox clusters of amphioxus, but transposable elements have been characterised within the ParaHox cluster (Osborne et al., 2006). This suggests that the genomic raw material for rearrangement and relaxation of the cluster is present, but that selective pressure retains the tight cluster in this organism (Ferrier et al., 2005).

**PARAHOX CLUSTER DUPLICATION**

In humans a ParaHox cluster is found on chromosome 13q12.2 (Pollard and Holland, 2000; Ferrier et al., 2005); it contains the genes *GSH1, IPF1* (also known as *PDX1,* the ortholog of invertebrate Xlox; we use the name Xlox to refer to this gene family in this review since the alternative names imply functions which are not always known) and *CDX2.* While the intergenic distances differ, the order and orientation of the genes is identical to the single cluster of amphioxus and there are no intervening genes.

The human genome also contains additional paralogs of the Gsx and Cdx families (*GSH2, CDX1, CDX4*), each on a different chromosome. The ParaHox genes of mouse show an equivalent genomic organization (Ferrier et al., 2005). It has long been known that humans and mice possess these additional members of the Gsx (Singh et al., 1991) and Cdx gene families (Duprey et al., 1998; James and Kazenwadel, 1991; Gamer and Wright, 1993), but not until 2000 was it suggested that they may in fact be the remains of duplicated ParaHox gene clusters Coulier et al., 2000; Pollard and Holland, 2000). Studies of the chromosomal regions containing these genes have revealed that a single prevertebrate ParaHox gene cluster was duplicated on bloc, giving rise to the four ParaHox paralogons seen in the human and mouse genomes (Ferrier et al., 2005). These en-bloc duplications were probably part of two rounds of genome duplication ("2R") at the base of the vertebrate lineage.

However, neither human nor mouse has four intact ParaHox clusters; rather, they have undergone the loss of at least six genes to retain only a single cluster of Gsx1, Pdx1, and Cdx2 with un-clustered Gsx2, Cdx1, and Cdx4 spread over three other chromosomes. Examination of the genome assemblies for the opossum *Monodelphis domestica* (MonDom5; Mikkelsen et al., 2007) and frog *Xenopus tropicalis* (v4.1) at http://www.ensembl.org show a similar organization, suggesting that these losses occurred prior to the divergence of tetrapods. Mulley et al. (2006) have shown that this organization is also present in basal actinopterygian fish such as *Amia calva* and *Polypterus senegalus.* Therefore we can infer that the common ancestor of all bony fish had undergone both duplications and losses in the ParaHox clusters (Fig. 1).

The complete genome sequences of several species of teleost fish have revealed that the entire teleost lineage is derived from a tetraploid ancestor, the so-called "3R" duplication (Jaillon et al., 2004; Meyer and Van de Peer, 2005). This genome duplication event resulted in the loss of the ParaHox cluster in this lineage (Mulley et al., 2006; Prohaska and Stadler, 2006; Siegel et al., 2007). In all species of teleost examined to date, the usually clustered *cdx2* gene has been lost, and *gsh1* and *pdx1* are on two paralogous chromosomes. Of the remaining ParaHox genes, *gsh2* and *cdx4* are found elsewhere in the genome, while duplicated *cdx1* genes have been retained and may compensate for the loss of *cdx2* in these species (Fig. 1).

**THE PARAHOX COMPLEMENT OF HAGFISHES**

As previously mentioned, it has been hypothesised that two whole-genome duplications ("2R") occurred on the vertebrate stem after the divergence of cephalochordates and urochordates and before the divergence of bony fish (Dehal and Boore, 2005; Holland et al., 1994; Dehal and Boore, 2005). Furthermore it has been suggested that these
genome duplications preceded and may have permitted the evolution of the innovations seen in vertebrate developmental patterning (Shimeld and Holland, 2000). The timing of these duplications is therefore key to our understanding of vertebrate development and evolution, and the en-bloc duplication of ParaHox gene clusters may be symptomatic of a larger-scale duplication event.

Only two extant groups diverged around the time of these putative duplications. These are the jawless fish, hagfishes and lampreys. The origins and relationships of these two groups are rather contentious; while molecular phylogenetic analysis strongly suggests that they form a monophyletic sister group to the vertebrates (e.g., Kuraku and Kuratani, 2006; Vienne and Pontarotti, 2006), their morphology argues that they are paraphyletic, and that hagfish form a “craniate” sister group to the true vertebrates, including lampreys (e.g., Forey and Janvier, 1993; Forey, 1984). Furthermore, the timing of their divergences with respect to the 2R duplications is also undetermined. While some cyclostome gene families appear to have diverged prior to the 2R duplications, other cyclostome genes appear to have undergone 1R or 2R duplications, grouping with vertebrate subfamilies rather than falling basally (e.g., Escriva et al., 2002; Bridgham et al., 2006; Zhang and Cohn, 2006). This suggests that the speciation and duplication events may all have occurred in such quick succession that the phylogenetic signal is impossible to resolve. After genome doubling, the organism must undergo the process of diploidization, during which most gene duplicates are lost and the sequences of the surviving paralogs diverge (Wolfe, 2001). Speciation during this diploidization process may confound phylogenetic analysis (Furlong and Holland, 2002). Genome doubling may also occur by hybridization rather than duplication (allopolyploidy); if one parental species contributes to several different tetraploids, their phylogenetic relationships may again be rather difficult to disentangle.

The lamprey genome project is as yet incomplete, and little is known about the ParaHox genes of these organisms.
However, ParaHox genes from two species of hagfish have recently been characterised (Furlong et al., 2007). In *Eptatretus burgeri*, a Gsx and a Cdx gene are contained in a single BAC in tail-to-tail configuration. No intervening genes are present between them. In *Myxine glutinosa*, an orthologous Gsx gene is linked to a pseudo Xlox sequence, suggesting that the functional Xlox gene has been lost in both species. Transposable elements are prevalent in the published sequences. The “complete” cluster of *E. burgeri* is around 57 kb, similar in size to that of amphioxus and rather smaller than that of many vertebrates. It has previously been suggested that the ParaHox clusters of amphioxus and mammals maintain not just gene order, but also have similarities in relative gene size and spacing (Ferrier et al., 2005). Expanding previous analyses to include all known deuterostome ParaHox clusters (i.e., those in which linkage has been retained and no intervening genes are found) reveals that the relative intergenic distances do remain fairly constant throughout evolution, with the distance between Gsx and Xlox being around three times that between Xlox and Cdx (Table 1; Supplementary Information). Without an Xlox gene, these distances obviously cannot be calculated for *E. burgeri*; however, the relative gene lengths can be examined. This shows that relative gene length is rather variable across the chordates, and there are no obvious trends.

While the tight linkage of the hagfish ParaHox cluster shows some similarities with the amphioxus cluster, it also shares some novel features with the vertebrate clusters. For example, a conserved non-coding element associated with the Gsx gene in vertebrates (Mulley et al., 2006) is present in hagfishes but not in amphioxus (data not shown). This is in keeping with the suggestion of Woolfe et al. (2005) that such regions are not found in invertebrates.

Loss is prevalent among the ParaHox genes, and in all the vertebrate species studied to date, a maximum of one intact cluster remains. The scheme shown in Fig. 1 indicates that, even if hagfishes diverged after the 2R duplications, they would have a maximum of two intact clusters. A phylogenetic analysis of the hagfish ParaHox genes indicates that in fact the hagfishes did not undergo any genome duplications at the base of the vertebrate lineage (Furlong et al., 2007). However, studies of cyclostome Hox genes suggest that both hagfish and lampreys have multiple Hox clusters (Ferrier and Holland, 2002). The genome also contains no Xlox gene (Edvardsen et al., 2005), but again the Cdx and Gsx genes have not yet been found to be chromosomally linked. It seems that, in deuterostome organisms, the three-gene cluster may be maintained in its entirety, but that any loss of genes is also associated with dispersal. The close linkage of *E. burgeri* Gsx and Cdx seems to be the only current example of a tightly linked partial cluster. An obvious question is whether these genes are expressed in a colinear manner; with the recent advances in hagfish embryology (Ota et al., 2007) it may soon become possible to test this.

There are several possible explanations for the maintenance of a ParaHox cluster in animal genomes over hundreds of millions of years. Firstly, the arrangement of the genes in the cluster may affect their spatial and temporal expression during development (as is the case in the hagfish gene clusters) if order and location of gene expression were found to follow the order of genes along the chromosome (Ferrier and Minguillon, 2003). Evidence from echinoderms would seem to counter this argument; however, since they do not have a tight cluster but appear to have spatial and temporal colinearity of expression, suggesting that apparent colinearity may be an artefact of the order of tissue development (Frobius and Seaver, 2006; Arnone et al., 2006). Secondly, the ParaHox genes may share overlapping regulatory sequences which trap the genes next to each other, allowing the genes to become dispersed only if these regulatory regions are duplicated as well — this hypothesis is supported by the dispersal of ParaHox genes to different chromosomes in teleost fish following whole genome duplication (Mulley et al., 2006). Finally, the ParaHox genes may be trapped together as “bystander genes” (Becker and Lenhard, 2007; Kikuta et al., 2007) by the action of long range regulatory sequences across them. In this case, there is no functional significance to the presence of Gsx, Xlox, and Cdx next to each other and the cluster is only maintained through the detrimental effects of genomic rearrangement to neighboring genes and their enhancers. This third hypothesis may also explain the break-up of the ParaHox cluster in teleost fish. At the present time there is insufficient data available to be able to choose between these options. More data on the timing and location of ParaHox gene expression in vertebrates are required, as well as a thorough analysis of conserved non-coding sequences associated with the ParaHox cluster which may be acting on it, or across it on adjacent genes. The supplementary information for this article can be found online at http://dx.doi.org/10.2108/zsj.25.955.s1.

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REFERENCES


