# Comparative genomics provides evidence for an ancient genome duplication event in fish 

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#### Abstract

There are approximately 25000 species in the division Teleostei and most are believed to have arisen during a relatively short period of time ca. 200 Myr ago. The discovery of 'extra' Hox gene clusters in zebrafish (Danio rerio), medaka (Oryzias latipes), and pufferfish (Fugu rubripes), has led to the hypothesis that genome duplication provided the genetic raw material necessary for the teleost radiation. We identified 27 groups of orthologous genes which included one gene from man, mouse and chicken, one or two genes from tetraploid Xenopus and two genes from zebrafish. A genome duplication in the ancestor of teleost fishes is the most parsimonious explanation for the observations that for 15 of these genes, the two zebrafish orthologues are sister sequences in phylogenies that otherwise match the expected organismal tree, the zebrafish gene pairs appear to have been formed at approximately the same time, and are unlinked. Phylogenies of nine genes differ a little from the tree predicted by the fish-specific genome duplication hypothesis: one tree shows a sister sequence relationship for the zebrafish genes but differs slightly from the expected organismal tree and in eight trees, one zebrafish gene is the sister sequence to a clade which includes the second zebrafish gene and orthologues from Xenopus, chicken, mouse and man. For these nine gene trees, deviations from the predictions of the fish-specific genome duplication hypothesis are poorly supported. The two zebrafish orthologues for each of the three remaining genes are tightly linked and are, therefore, unlikely to have been formed during a genome duplication event. We estimated that the unlinked duplicated zebrafish genes are between 300 and 450 Myr . Thus, genome duplication could have provided the genetic raw material for teleost radiation. Alternatively, the loss of different duplicates in different populations (i.e. 'divergent resolution') may have promoted speciation in ancient teleost populations.


Keywords: genome duplication; speciation; phylogenetics; zebrafish (Danio rerio); comparative genomics

## 1. INTRODUCTION

Major transitions, including the evolution of eukaryotes, metazoans, Bilateria and Vertebrata, may have required the genetic raw material provided by gene and/or genome duplications (Ohno 1970; Lundin 1993, 1999; Sidow 1996; Holland 1999; Patel \& Prince 2000). Ohno (1970) presented comparative data on genome size and chromosome numbers to support his hypothesis that one or more genome duplications preceded the evolution of vertebrates. Ohno further proposed that the new redundant genes produced by genome duplication evolved new functions that were necessary for vertebrate evolution. The apparent functional connection between duplicate genes and the evolution of vertebrates was more fully asserted by Holland (1992). In mice, paralogues Hox-1.5 and Hox1.6 (renamed $H_{o x A 3}$ and $H_{o x A 1}$ respectively-De Robertis 1994) have overlapping expression domains and are at least partially functionally redundant. Holland proposed that overlapping expression domains among paralogous genes (Fitch 1970) delimit the expression domain of their single ancestral gene and that non-overlapping expression domains represent postduplication gains of function. Holland (1992) also
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suggested that post-duplication gains of function, particularly in Hox genes, facilitated the evolution of vertebratespecific features such as the control of neural crest cell fate and organogenesis, hindbrain differentiation and otic morphogenesis. It is clear that duplicated genes can evolve previously non-existent functions. Expansion of repetitive regions in one copy of a duplicated pancreatic trypsinogen-like gene produced a gene for antifreeze glycoproteins in Antarctic fish (Cheng \& Chen 1999) and mutations in duplicated opsin genes led to the evolution of trichromatic vision in New and Old World primates (Dulai et al. 1999). However, the causal link between gene duplication and major evolutionary transitions remains a matter of speculation.

Ohno's hypothesis that big leaps in evolution required the creation of new gene loci with previously non-existent functions emphasized genome duplication via tetraploidy as the mechanism for the production of new genes. Gene number comparisons support this model. Spring (1997) uncovered an average of three orthologous genes in humans for each of 52 Drosophila genes and proposed that the additional human genes were produced during two genome duplications. However, Spring's hypothesis, which has recently been referred to as the 'one to four rule' (Ohno 1999) and the '2R' hypothesis (Hughes 1999a), remains highly controversial (Hughes 1999a; Wang \& Gu 2000).

Genome duplication in Actinopterygii (ray-finned fishes) is the focus of this study. The recent discovery of 'extra' Hox gene clusters in zebrafish (Danio rerio) and pufferfish (Takifugu rubripes) led Amores et al. (1998) to the conclusion that a chromosome doubling event, probably by whole genome duplication, occurred after the divergence of ray-finned and lobe-finned fishes. Hox genes encode DNA-binding proteins and occur in one or more clusters of up to 13 genes per cluster. In Sarcopterygii (a monophyletic group including lobe-finned fishes, amphibians, reptiles, and mammals) there appear to be four Hox clusters labelled $\mathrm{A}, \mathrm{B}, \mathrm{C}$ and D with each cluster occurring on a different chromosome. In contrast, zebrafish possess at least seven Hox clusters and the pufferfish has two 'Hox A' clusters (Amores et al. 1998; Aparicio 2000). As in sarcopterygians, fish Hox clusters occur on different chromosomes. Following Amores et al.'s (1998) conclusion that genome duplication was the explanation for the 'extra' Hox clusters in fish, Meyer \& Schartl (1999) expanded the 'one to four rule' to the 'one to four to eight rule' to account for this additional genome duplication. Teleostei is the most diverse of all vertebrate groups and includes approximately 25000 species (Nelson 1994). Major teleost lineages are believed to have arisen between $c a .100$ and 200 Myr ago (Carroll 1997; Lydeard \& Roe 1997) and Amores et al. (1998) and Meyer \& Schartl (1999) proposed that genome duplication facilitated this radiation.

Stellwag (1999) suggested that, with respect to Hox cluster number, the zebrafish is not representative of actinopterygians and that the genome duplication proposed by Amores et al. (1998) might be limited to only a few derived fish or even the zebrafish lineage alone. This argument was weakened when it was discovered that medaka (Oryzias latipes), which is placed in a different teleost superorder than zebrafish, also possess seven Hox clusters (Naruse et al. 2000). Other criticisms of the teleost genome duplication hypothesis have focused on the fact that Hox genes reveal the history of only a small portion of the entire genome. Most fishes have smaller genomes than humans (Ohno 1970; Hinegardner \& Rosen 1972). The zebrafish genome is approximately half the size of the human genome (Hinegardner \& Rosen 1972). Morizot et al. (1991) estimated that the genome of the platyfish (Xiphophorus) is five times smaller than the human genome and Elgar et al. (1999) estimated that the pufferfish genome is eight times smaller than the human genome. Although genome size and gene content may not be correlated, Elgar et al. (1999) suggested that the duplication of Hox clusters by regional duplication is easier to reconcile with fish genome size data than genome duplication.

The goal of our study was to use a phylogenetic approach to evaluate the hypothesis that the 'extra' Hox genes and the rest of the genome in fishes were produced during a genome duplication in a teleost ancestor rather than by a series of regional duplications. The genome duplication hypothesis makes clear predictions about the number of genes in fishes compared with humans and about the topology of gene trees: a gene tree should match the expected organismal tree but have two zebrafish orthologues for each human gene and the zebrafish orthologues should be sister sequences in a phylogenetic


Figure 1. Phylogenetic topology predicted assuming the ancestor of actinopterygian fishes experienced a genome duplication. This topology, referred to as the 'duplication topology', also assumes that no genes have been lost in the taxa surveyed. Supplements to the term homology are described in the figure: 'or thology' (Fitch 1970) describes the relationship between homologous genes (i.e. genes descended from a common ancestral gene) that occur in different species; 'paralogy' (Fitch 1970) describes the relationship between homologous genes that occur within an individual (e.g. genes produced by genome or by tandem duplication). Duplicated zebrafish genes are 'co-orthologues' of their human orthologues (Gates et al. 1999).
analysis (figure 1). We refer to this predicted topology as the 'duplication topology'. Furthermore, pairs of zebrafish orthologues from different genes should have been formed at the same time and should be unlinked.
Human and zebrafish protein sequences were obtained from the non-redundant (NR) protein database at the National Center for Biotechnology Information (NCBI, Bethesda, MD, USA) to determine whether gene numbers and gene phylogenies support the fish-specific duplication hypothesis. We also collected sequences from Mus musculus, Gallus gallus and Xenopus laevis so that we could reconstruct the reliable phylogenies necessary to identify orthologues among the sequences retrieved in our basic local alignment search tool (BLAST) searches. Map data are available for most of the zebrafish genes in our survey and we used these data to determine whether anciently duplicated genes are distributed throughout the zebrafish genome.

## 2. METHODS

## (a) Database searches

Protein sequences of zebrafish (Danio rerio), human (Homo sapiens), mouse (Mus musculus), chicken (Gallus gallus) and the African clawed frog (Xenopus laevis) were obtained by Blastp (Altschul et al. 1990). For all searches we selected the NR search option (see http:/www.ncbi.nlm.nih.gov/blast/html/blastcgihelp. html\#nucleotide_databases). With a few exceptions, human 'reference sequences' (Maglott et al. 2000) were used as BLASTp query sequences. Most genes surveyed were those used in a gene number comparison between Drosophila and humans (Spring 1997), but the mammalian genes that Gates et al. (1999) describe as having two zebrafish orthologues were also included. Species were surveyed one at a time to improve the identification of a drop in sequence similarity, which was used as a 'cut-off'. Sequences above the cut-off value were pasted to NCBI clipboards and then downloaded in FASTA format, a format that includes the sequence definition line and sequence characters.


Figure 2. Neighbour-joining tree of the retinoic acid receptor genes retrieved using BLASTp (gene identification numbers shown). Sequences that varied only in length or by very few amino-acid substitutions were removed prior to analysis (see § 2 ). The tree shows paralogous clades of $R A R \alpha, R A R \beta$, and $R A R \gamma$ genes. Bootstrap values (Felsenstein 1985) are shown ( 500 bootstrap reiterations).

## (b) Sequence alignment and phylogeny reconstruction

When BLASTp identified one or more putative zebrafish orthologues, protein sequences from all species were aligned using ClustalX (Thompson et al. 1997). For each alignment, a preliminary tree was drawn from the Clustal dendrogram file using TreeView v. 1.6.0 (Page 1996). This tree facilitated the identification of identical sequences, sequences that varied only in length, and sequences within species that differed by few amino acids, all of which were removed from the alignment. Very similar sequences could be alleles at one locus or evidence of recent tandem duplications. In either case they were not likely to be important for our study of genome duplication in the teleost ancestor.

Phylogenies were reconstructed from the remaining sequences using Poisson-corrected genetic distances and the neighbourjoining (NJ) algorithm (Saitou \& Nei 1987) in Treecon (Van de Peer \& De Wachter 1994). These first NJ phylogenies included many clades of orthologous and paralogous genes (e.g. figure 2). From these large trees we identified sets of orthologous genes (i.e. genes which occurred in monophyletic groups that matched the expected organismal topology). Sequences of orthologous genes were realigned and edited using BioEDiT (http://www. mbio.ncsu.edu/RNaseP/info/programs/BIOEDIT/bioedit.html). Regions where the alignment was unambiguous were retained and reanalysed using NJ and maximum likelihood (ML) methods. For these last phylogenetic analyses the most closely related human paralogues (identified from the first NJ analyses) were used as outgroups. Support for nodes was evaluated by 500 bootstrap reiterations (Felsenstein 1985). Tree-PuZZLE v. 5.0 (Strimmer \& Von Haeseler 1996) was used to reconstruct ML
trees (substitution models were selected for each analysis automatically by the program).

## (c) Dating duplication events

In order to estimate the age of zebrafish paralogues, the number of nucleotide substitutions at third codon positions was plotted against divergence dates for different taxa (Nei \& Kumar 2000). Since most third-codon position substitutions do not result in amino-acid replacements, the rate of fixation of these substitutions is expected to be relatively constant in different protein-coding genes (e.g. Nei et al. 2000) and to reflect the overall mutation rate (Hughes 1999b). Alternatively, one can use the number of synonymous substitutions per synonymous sites to estimate divergence times (Nei \& Kumar 2000; Nei et al. 2000). However, for the genes surveyed here, there is an approximately linear relationship between the number of thirdposition substitutions and the number of synonymous substitutions and therefore both approaches are expected to give similar results. Estimation of the number of substitutions at third-codon positions, corrected for multiple events per site according to Tajima \& Nei (1984), was done for 26 pairs of genes (no DNA sequence was available for the two zebrafish GDF6 genes). All computations were done with the software package Mega2 (Nei \& Kumar 2000).

Divergence dates between different taxa were taken from literature and were as follows: genome duplication in Xenopus, 30 Myr ago (Hughes \& Hughes 1993); divergence between human and mouse, 100 Myr ago (Li et al. 1990; Kumar \& Hedges 1998); divergence between reptiles (represented by the bird Gallus gallus) and mammals, 310 Myr ago (Kumar \& Hedges 1998); divergence between amphibians and amniotes, 360 Myr ago (Kumar \&

Hedges 1998); and divergence between ray-finned fish and Sarcopterygii, 450 Myr ago (Kumar \& Hedges 1998).

## 3. RESULTS

## (a) Gene numbers and phylogenetic analyses

BLASTp searches uncovered a large number of sequences for each species, many of which differed only in length or by very few amino-acid replacements. Neighbour-joining analyses of the longest sequences often identified many (up to 15) different monophyletic groups of orthologous genes (e.g. figure 2). Groups of orthologous and paralogous genes analysed together are listed together in different blocks in table 1. Groups of orthologous genes within these clades are presented on separate rows within blocks in table 1 .

Variation in the length of sequences in different species meant that for some genes a large proportion of the available data could not be used for phylogenetic analyses. Furthermore, sequence variation among taxa meant that large portions of some sequences could not be unambiguously aligned.

For 27 genes, NJ analyses produced a well-supported clade with two zebrafish genes, one human, mouse and/or chicken gene and one or two Xenopus genes. Eighteen of these 27 trees had the 'duplication topology' (figure 3a). In one tree (EN2) zebrafish genes are sister sequences but, unexpectedly, they cluster with the two Xenopus genes (figure $3 a$ ). For eight trees (figure $3 b$ ) one of the two zebrafish genes was the sister sequence to a monophyletic group that included the second zebrafish gene and orthologous genes from Xenopus, chicken, mouse and human. Phylogenies of the eight genes shown in figure $3 b$ have the 'outgroup topology'. Eighteen of the 19 genes with zebrafish orthologues as sister sequences using NJ methods also had this sister sequence relationship when ML methods were used (for ISL2, ML analyses produce the 'outgroup topology'). Among the eight genes in figure $3 b$, ML analysis produced the 'duplication topology' for FKD5, HOXC6 and SOX11. Maximum likelihood analyses of SNAP25 data supported the hypothesis that the two zebrafish genes (snap25,1 and snap25,2) were sister sequences, but the zebrafish, mouse and human SNAP25 sequence did not form a monophyletic group when ML methods were used. Both phylogenetic methods produced the 'outgroup topology' for four genes (DLX2, $\mathcal{F} A K 2$, $\mathcal{N T N 1}$ and OTX1).

Bootstrap support for the duplication topology or the outgroup topology was low for some trees in figure 3, even when the same topology was produced by both phylogenetic methods. To test whether the tree topologies shown in figure 3 were significantly better than the alternative topology, we performed a Kishino-Hasegawa test (Kishino \& Hasegawa 1989) as implemented in TreePuZZLE (Strimmer \& Von Haeseler 1996). As already might have been expected on the basis of the bootstrap analysis, user-defined trees where the two zebrafish genes are sister sequences were not found to be significantly worse than the $D L X 2, \mathcal{F A K} 2, \mathcal{N T N 1}$ and SOX11 trees shown in figure $3 b$. However, our application of the Kishino-Hasegawa test also produced unexpected results. The Kishino-Hasegawa test failed to reject the 'outgroup topology' in many cases even when NJ and ML analyses
produced the 'duplication topology' with high bootstrap support. For these genes the likelihood of a sister sequence relationship between zebrafish paralogues (i.e. the 'duplication topology') was always the highest, but the 'outgroup topology' was not significantly worse. The KishinoHasagawa test appears to have low resolving power for our datasets, which may be too conserved and include too few samples (A. von Haeseler, personal communication).

## (b) The age of the duplicated genes

To estimate the date of the fish-specific duplication, we plotted known divergence dates between different taxa against the number of nucleotide substitutions at thirdcodon positions (see § 2). Although we initially included the split between ray-finned fish (Actinopterygii) and Sarcopterygii, this divergence and the corresponding number of substitutions between zebrafish and the other vertebrates were omitted from the final analysis since the nucleotide substitutions at third codon positions were clearly saturated (not shown). This is probably also true for the amphibian-amniote divergences (as shown by the large differences in number of substitutions; figure 4) and to some extent for the divergence between the chicken and mammals (Nei \& Kumar 2000). However, based on the plot of figure 4, complete saturation probably does not occur much earlier.

Divergence dates for different vertebrate lineages are controversial and may differ considerably whether based on palaeontological or molecular calibration (Kumar \& Hedges 1998; Gu 1998; Lee 1999). Nevertheless, if we consider the dates used as reliable, and using 1.02 (s.d. $=0.24$ ) as the average number of substitutions per site between the 23 pairs of unlinked zebrafish coorthologues (see below), the fish-specific genome duplication occurred ca. 350 Myr ago. Since the third codon positions have probably reached saturation, as indicated by the high number of estimated substitutions per site when both zebrafish genes are compared, this calculation is at the limit of our ability to estimate dates. In conclusion, the fish-specific genome duplication is probably older than 300 million years, if we assume that thirdcodon positions are not completely saturated at the time of the reptilian-mammalian divergence. Furthermore, assuming that the genome duplication is not older than the divergence of the Actinopterygii and Sarcopterygii, the duplication probably occurred between 300 and 450 Myr ago.

## (c) Map positions

Zebrafish co-orthologues shown in figure 3 are distributed among 16 of the 25 zebrafish linkage groups (table 2). For DLL and MSX3, one co-orthologue occurs on linkage group (LG) 1 and the other on LG13, and for DLX2 and EN1, one zebrafish co-orthologue occurs on LGl and the other on LG9. For $E N 2$ and $S H H$, one zebrafish coorthologue occurs on LG2 and the other on LG7. For BMP2, SNAP25 and SOX11 one co-orthologue occurs on LG17 and the other occurs on LG20. Lastly, for three genes (HOXB5, HOXB6 and RAR $\alpha$ ) one co-orthologue occurs on LG3 and the other on LG12. Thus, portions of LG1 and LG13, LG1 and LG9, LG2 and LG7, LG17 and LG20, and LG3 and LG12 appear to be paralogous (table 2).

Table 1. Surveyed genes.
(Blocks separated by blank lines identify families of genes uncovered in BLAST searches and used for tree reconstruction. Rows (some comprised of more than one line) identify genes that are orthologous to a single human gene according to our phylogenetic analyses. Genes with topologies that support the fish-specific genome duplication hypothesis are shaded. '-', no orthologous genes found in databases.)

| human gene name | Homo sapiens | Danio rerio | Mus musculus | Gallus gallus | Xenopus laevis |
| :---: | :---: | :---: | :---: | :---: | :---: |
| ABL1 | 4885045 | - | 125137 | - | - |
| ABL 2 | 6382060 | - | - | - | 7248894 |
| ALDOA | 4557305 | - | 7548322 | - | 1944025 |
| ALDOB | 4557307 | - | - | 113610 | - |
| ALDOC | 113613 | - | 113614 | 226855 | 3928511 |
| APP | 4502167 | 8050809 | 6680708 | 6465892 | 320195 |
| APLP1 | 4885065 | - | 6680700 | - | - |
| APLP2 | 4502147 | - | 1086521 | - | - |
| ANK1 | 4502089 | - | 1168457 | 1245423 | - |
| ANK2 | 4502091 | - | - | 1245425 | - |
| ANK3 | 4502093 | - | 710549 | 1245427 | - |
| BMP2 | 4557369 | $\begin{aligned} & 2804175 \\ & 2149148 \end{aligned}$ | 6680794 | 2501173 | 115070 |
| BMP4 | 4502423 | 2149144 | 461633 | 2501175 | $\begin{aligned} & 399122 \\ & 477512 \end{aligned}$ |
| BMP5 | 339560 | - | 6671642 | 1881823 | - |
| BMP6 | 4502425 | - | 6680798 | - | - |
| BMP7 | 4502427 | 6573121 | - | 6970053 | 4096790 |
| BMP8 | 4502429 | - | 6671644 | - | - |
| BRNI (POU3-tf2) | 5453936 | $\begin{aligned} & 1730449 \\ & 2495310 \end{aligned}$ | 6679425 | - | - |
| POU3-tf3 (outgroup) |  | 5031983 |  |  |  |
| В ¢ $^{\prime}$ | 4557377 | - | 2507603 | - | - |
| ITK | 7949058 | 2353318 | - | - | - |
| TEC | 4507429 | - | 420220 | - | - |
| ТХК | 4507743 | - | 1174826 | - | - |
| CDH 1/3/14 | 4757960 | - | - | 115417 | 13432108 |
|  | 4502721 | - | - | 416739 | 13432110 |
| CDH2 | 14589889 | 2133885 | - | 115422 | $\begin{aligned} & 416743 \\ & 115425 \end{aligned}$ |
| CDH12 ${ }^{\text {a }}$ | 2119627 | - | 6680904 | 3023428 | - |
|  |  | - |  | 2134302 | - |
| cad7 | - | - | 7549750 | 2134303 | 2119628 |
| cad11 | - | 1345125 | 6753372 | 3511021 | 3377485 |
| $C A L M ~^{\text {b }}$ | 5901912 | - | 6680832 | 3415119 | 6137739 |
| CALM2 ${ }^{\text {b }}$ | 4502549 | - |  |  |  |
| CALM ${ }^{\text {b }}$ | 4885109 | - |  |  |  |
| CD X1 | 4502763 | - | 1170313 | 1170316 | 435578 |
| CD X2 | 4502765 | - | 1170314 | 1737445 | - |
| CD X4 | 4885127 | 283775 | 1083362 | 547650 | 2134077 |
| COL4A 1 | 7656985 | - | 115312 | 7271901 | - |
| COL4A 3 | 177894 | - | 6680968 | - | - |
| COL4A 5 | 4502955 | - | 2119170 | - | - |
| CTS H | 4758096 | - | 7106279 | - | - |
| CTS K | 4503151 | - | 6681085 | 1017831 | - |
| CTS L | 4503155 | 1752664 | 6753558 | 2144502 | 2706547 |
| CTSS | 4758098 | - | 3850787 | - | - |
|  |  |  | 2961621 |  |  |
| Catlop-p | - | - | 5306071 | - | - |
| Catm | - | - | 7715970 | - | - |

Table 1. continued

| human gene name | Homo sapiens | Daniorerio | Mus musculus | Gallus gallus | Xenopus laevis |
| :---: | :---: | :---: | :---: | :---: | :---: |
| DLL1 | 10518497 | $\begin{aligned} & 2809389 \\ & 1888392 \end{aligned}$ | 6681197 | 2134296 | 807696 |
| DELTA4 (outgroup) | 8926615 |  |  |  |  |
| DLX1 | 2829447 | 2842747 | 6753644 | - | - |
| DLX2 | 4758168 | $\begin{aligned} & 2842748 \\ & 1708243 \end{aligned}$ | 6753646 | - | $\begin{aligned} & 1079297 \\ & 1708249 \end{aligned}$ |
| DLX3 | 4885185 | 1346299 | 2495277 | 5830236 | $\begin{aligned} & 2134092 \\ & 1708245 \end{aligned}$ |
| DLX4 | 4503343 | - | 6681201 | - | - |
| DLX5 | 4885187 | 1708248 | 2495278 | 1708250 | 2134167 |
| DLX6 | 4885189 | 2842749 | 6014979 | - | 1708242 |
| DLX7 | - | 2842750 | - | - | - |
| DLX8 | - | 2842751 | - | - | - |
| TCF3/E2a | 181906 | 2118448 | - | 506759 | 283796 |
| TCF4\|E2b | 4507399 | - | 7305551 | - | - |
| TCF12/E2c | 4507391 | - | 346644 | 416847 | - |
| E2F2 | 4758226 | - | - | - | - |
| E2F3 | 4503433 | - | 3122045 | - | - |
| EGF | 4503491 | - | 6753732 | - | - |
| TGFA | 4507461 | - | 1351229 | - | - |
| HGL | 4758526 | - | - | 9297019 | - |
| AREG | 4502199 | - | 6753100 | - | - |
| DTR | 4503413 | - | 6754178 | 4761593 | - |
| TDGF1 | 4507425 | 8132035 | - | - | - |
| EGFR | 4885199 | - | 1352359 | 1070476 | - |
| ERBB2 | 4758298 | - | - | - | - |
| ERBB3 | 4503597 | - | - | - | - |
| ERBB4 | 4885215 | - | - | 4884676 | - |
| EGR1 | 4503493 | 1352361 | 6681285 | - | 7673684 |
| EGR2 | 4557549 | 462005 | 2507546 | - | 1169500 |
| EGR3 | 4758252 | - | 9055212 | - | - |
| EGR4 | 4503495 | - | 4704780 | 6707678 | - |
| EMX1 | 31140 | 2133842 | 729412 | - | - |
| EMX2 | 31142 | 2133843 | 729414 | - | - |
| EN1 | 7710119 | $\begin{aligned} & 4322044 \\ & 417127 \end{aligned}$ | 7106305 | 483162 | $\begin{aligned} & 1708255 \\ & 399907 \end{aligned}$ |
| EN2 | 7710121 | $\begin{aligned} & 417128 \\ & 417129 \end{aligned}$ | 6753752 | 483259 | $\begin{aligned} & 1708257 \\ & 1708256 \end{aligned}$ |
| EPAI | 2827756 | - | - | - | - |
| EPA2 | 4758278 | 3005903 | 6753758 | - | 3861464 |
| EPA3 | 4885211 | - | 125338 | 125337 | - |
| EPA4 | 4758280 | 3005933 | 6679657 | 2833208 | $\begin{aligned} & 8134439 \\ & 8134440 \end{aligned}$ |
| EPA5 | 1706628 | - | 6679659 | 1706627 | - |
| EPA7 | 4758282 | 1754761 | 2497573 | 8134447 | - |
| EPA8 | 7263928 | 8134436 | 6679663 | - | - |
| EPB1 | 2739208 | - | - | 8134448 | $\begin{aligned} & 8134450 \\ & 8134449 \end{aligned}$ |
| EPB2 | 1706664 | - | 1706665 | 2827774 | 2739062 |
| EPB3 | 4758288 | 2198795 | 1708165 | 2134386 | 974710 |
| EPB4 | 4758290 | $\begin{aligned} & 3005901 \\ & 3163942 \end{aligned}$ | 6753760 | - | $\begin{aligned} & 6689570 \\ & 6689572 \end{aligned}$ |
| EPB6 | 4758292 | - | - | 2833209 | - |
| $\begin{aligned} & \text { EVX1 } \\ & \text { EVX2 } \\ & \text { ever }^{\text {c }} \end{aligned}$ | $\begin{aligned} & 4503615 \\ & 553284 \end{aligned}$ | 4322046 1617040 630922 | $\begin{aligned} & 6679711 \\ & 6679713 \end{aligned}$ | - | 1708342 |

Table 1. continued

| VIL2 | 4507893 | - | 6678571 | 4514720 | - |
| :--- | :--- | :--- | :--- | :--- | :--- |
| RDX | 4506467 | - | 6677699 | 6179570 | - |
| MSN | 4505257 | - | 462608 | - | 6648536 |
| FGFr1 | 182532 | - | 309240 | 120045 | 214900 |
| FGFr2 | 4503709 | - | 214423 | 116098 | 544293 |
| FGFr3 | 4503711 | 8886017 | 47723 | 116097 | 2425188 |
| FGFr4 | 4503713 | 773667 | 6679789 | - | 2541908 |
|  |  |  |  | 1213275 |  |
| FKD5 | 8134472 | 2982343 | 2494502 | - | 3695057 |
| FXL1 (outgroup) | 13638268 | 2982347 |  | - |  |
| FLOT1 |  |  | - | - |  |
| flotillin1 (outgroup) | 3115387 | 12751185 | 6679811 |  |  |
|  | (Dros.) |  |  | - |  |

$\left.\begin{array}{llllll}\text { gdf6 } & - & 914116 & 1707885 \\ \text { (bovine) }\end{array}\right]$

Table 1. continued

| human gene name | Homo sapiens | Danio rerio | Mus musculus | Gallus gallus | Xenopus laevis |
| :---: | :---: | :---: | :---: | :---: | :---: |
| HOXA11 ${ }^{\text {e }}$ | 5031759 | 4322049 | 6754226 | 399992 | 2995957 |
|  |  | 1707451 |  |  |  |
| HOXC11 ${ }^{\text {e }}$ | 7657166 | 4322084 | - | - | - |
|  |  | 4322086 |  |  |  |
| HOXD11 | 400021 | 974813 | 123292 | 400020 | - |
| HOXA13 ${ }^{\text {c }}$ | 4504457 | $\begin{aligned} & 4322051 \\ & 4322053 \end{aligned}$ | 6680245 | - | - |
|  |  |  |  |  |  |
| HOXC13 ${ }^{\text {c }}$ | 7689387 | $\begin{aligned} & 4322090 \\ & 4322092 \end{aligned}$ | 1708359 | - | - |
|  |  |  |  |  |  |
| ID1 | 4504569 | 2253424 | 2827752 | - | - |
| ID2 | 4504571 | - | 109791 | 2935461 | 2134185 2134043 4587148 |
|  |  |  |  |  |  |
|  |  |  |  |  |  |
| ID3 | 2135331 | - | 6680341 | - | - |
| ID4 | 4504573 | - | 729812 | - | - |
| INSR | 4557884 | - | 6754360 | 4588602 | 5420052 |
| INSRR | 186555 | - | 6754362 | - | - |
| IGF1R | 4557665 | - | 3025894 | 2808533 | $\begin{aligned} & 1150692 \\ & 3037089 \end{aligned}$ |
|  |  |  |  |  |  |
| ISL1 | 124927 | 1708559 | 4469284 | 1708560 | - |
| isl2 | - | $\begin{aligned} & 1708564 \\ & 1708561 \end{aligned}$ | $\begin{aligned} & 1708563 \\ & (\mathrm{rat}) \end{aligned}$ | 1708562 | - |
| ITGA2B | 4504745 | - | 7262859 | - | - |
| ITGA5 | 4504751 | - | 6754378 | - | 3183037 |
| ITGA4 | 4504749 | - | - | - | - |
| ITGB3/4 | 124968 | - | 7949057 | 631019 | 2119641 |
| ITGB6 | 9446402 | - | 4324977 | - | - |
| ITGB7 | 4504777 | - | - | - | - |
| ITGB1 | 4504767 | - | 124964 | 124962 | 124961 |
|  |  |  |  |  | 124965 |
| ITGB2 | 4557886 | - | - | - | - |
| ITGB5 | 4504773 | - | 3478697 | - | - |
| Э ${ }^{\text {K1 }}$ | 4504803 | 1938358 | 1708580 | 4558482 | - |
| T YK2 | 4507749 | - | 5733095 | - | - |
| ЭАК2 | 4826776 | $\begin{aligned} & 3687398 \\ & 3687400 \end{aligned}$ | 6680508 | - | - |
| Э АК3 | 4557681 | - | 2499670 | - | - |
| LI (CAM) | 4557707 | $\begin{aligned} & 1065714 \\ & 1065716 \end{aligned}$ | 6651057 | 104799 | - |

NRCAM (outgroup) 6651380

| LAMA1 | 34226 | - | 6678656 | 1246110 | - |
| :--- | :--- | :--- | :--- | :--- | :--- |
| LAMA2 | 4557709 | - | 2497588 | - | - |
| LAMA3 | 4557711 | - | 1922889 | - | - |
| LAMB1 |  |  | - | - |  |
| LAMB2 | 4504951 | - | 126367 | - |  |
| LAMB3 | 4504953 | - | 6678658 | 2708707 | - |
|  | 4557713 | - | 6678660 | - | 267419 |
| LHX1 |  |  |  |  |  |
| Lhx5(outgroup) |  | 2497670 | 6678688 |  | continued |

Table 1. continued

| MEF2A | 5031907 | 1518141 | 7305265 | 4914481 | $\begin{aligned} & 913313 \\ & 913312 \end{aligned}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| MEF2C | 4505147 | 1518143 | 477011 | - |  |
| MEF2D | 5174545 | 1518145 | 2500877 | - | 2500878 |
| MSX1 | 123310 | - | 11177822 | 1708273 | 234375 |
| MSX2 | 1082306 | - | 547660 | 1170325 | 547691 |
| Msx 3 | - | $\begin{aligned} & 399912 \\ & 2506531 \end{aligned}$ | 6754756 | - | - |
| Msx ${ }^{\text {f }}$ |  | 399913 |  |  |  |
| $M s x A^{\text {f }}$ |  | 2506530 |  |  |  |
| MYOD1 | 4505309 | 3914105 | 6996932 | 3915780 | $\begin{aligned} & 127711 \\ & 127053 \end{aligned}$ |
| MYOG | 4505311 | - | - | - | - |
| MYOD5 | 5031929 | - | 6678982 | - | 127629 |
| MYH9 | 189030 | - | - | 127759 | 3660672 |
| MrH10 | 641958 | - | - | 212449 | 422615 |
| MYH11 | 2104553 | - | 7441402 | 3915778 | - |
| NFKB1 | 189180 | - | 6679044 | 222839 | - |
| NFKB2 | 4505383 | - | 5081604 | 2134380 | 3116208 |
| REL | 4506473 | - | 6677707 | 136185 | 1004330 |
| RELA | 307300 | - | 6677709 | 1729913 | 548721 |
| RELB | 5730007 | - | 6677711 | 5305228 | 1710086 |
| NOS1 | 987662 | - | 6724321 | - | - |
| NOS2A/B/C | 1228940 | - | 6754872 | 2498062 | - |
| NOS3 | 189212 | - | - | - | - |
| NTN1 | 4758840 | $\begin{aligned} & 2327065 \\ & 2394302 \end{aligned}$ | 4732097 | 2497605 | 2655297 |
| $\mathcal{N T N} 2$ (outgroup) | 5453810 |  |  |  |  |
| OTX1 | 417425 | $\begin{aligned} & 3024322 \\ & 3024327 \end{aligned}$ | 417426 | - | - |
| OTX2 | 417427 | 3024329 | 417428 | - | $\begin{aligned} & 644782 \\ & 3024328 \end{aligned}$ |
| OTX 5 | - | - | - | - | $\begin{aligned} & 6624755 \\ & 6252982 \end{aligned}$ |
| PAX2 | 4557821 | $\begin{aligned} & 3420031 \\ & 3024368 \end{aligned}$ | 417447 | 6683012 | $\begin{aligned} & 5815455 \\ & 2765055 \end{aligned}$ |
| PAX5 (outgroup) | 417449 |  |  |  |  |
| PBX1 | 4505623 | 7160792 | $\begin{aligned} & 2432009 \\ & 7110681 \end{aligned}$ | 8096555 <br> 8096557 | - |
| PBX2 | 4505625 | 7160798 | - | - | - |
| PBX3 | 5453852 | 7160796 | 2432017 | - | - |
| PBX4 | - | 5679283 | - | - | - |
| PTC1 | 4506247 | 4539024 | 6679519 | 6225890 | - |
| PTC2 | 4506245 | 6225889 | 6679517 | - | - |
| RAF1 | 4506401 | 534977 | - | 125489 | 125654 |
| ARAF1 | 4502193 | - | 125646 | - | - |
| BRAF | 4757868 | - | - | 464647 | - |
| RAN | 131845 | 2500061 | 6677677 | 1172839 | 6729160 |
| RASN(outgroup) | 6857182 (Dros.) | - | - | - | - |
| NRAS | 4505451 | 3334308 | 7242162 | - | 3334309 |
| HRAS | 4885425 | - | 6680271 | 31868 | - |
| KRAS2A | 131875 | - | 417590 | - | 2072749 |
| KRAS2B | 131879 | - | 131880 | - | $\begin{aligned} & 3599487 \\ & 464552 \end{aligned}$ |

Table 1. continued

| human gene name | Homo sapiens | Daniorerio | Mus musculus | Gallus gallus | Xenopus laevis |
| :---: | :---: | :---: | :---: | :---: | :---: |
| RALA | 4885569 | - | 131836 | - | - |
| RALB | 4506405 | - | - | - | 3955067 |
| RAR ${ }^{\text {a }}$ | 4160009 | 704370 | 133484 | 2119682 | 2119679 |
|  |  | 215026 |  |  | 2119680 |
|  |  |  |  |  | 283822 |
| RAR $\beta$ | 133492 | - | 91262 | 3041719 | - |
| RAR $\gamma$ | 133500 | 6094141 | 133501 | 623767 | 1839488 |
| RB1 | 4506435 | - | 6677679 | 459445 | - |
| RBL1 | 4506443 | - | 2498835 | - | - |
| RBL2 | 5032029 | - | 6685841 | - | - |
| RXRA | 4506755 | 1583309 | 6755384 | - | 283824 |
| RXRB | 1350911 | 1046299 | 1350912 | - | 1085220 |
|  |  | 1046297 |  |  | 840922 |
| RXRG | 5902068 | 8478106 | 1350914 | 133700 | 1710810 |
| SRC | 4885609 | - | 6678129 | 6175046 | 125705 |
| rest | 4885661 | - | 6678617 | 125869 | 321075 |
| FGR | 4885235 | - | 6753860 | - | - |
| FrN | 4503823 | - | 6679879 | 479367 | 125371 |
| LCK | 4885449 | - | 2117800 | 1170731 | - |
| LYN | 4505055 | - | - | - | 2114076 |
| НСК | 4504357 | - | 6754166 | - | - |
| BLK | 4502413 | - | 6680786 | - | - |
| SDC1 | 4506859 | - | 6755438 | - | 2547264 |
| SDC2 | 386787 | - | 6677891 | - | 2547266 |
| SDC4 | 4506861 | - | 6755442 | 1351051 | - |


| SNA 11 | 5729674 | 841424 | 6755586 | - | - |
| :--- | :--- | :--- | :--- | :--- | :--- |
| SLUG (outgroup) | 2832266 | - | - | - | - |
|  |  |  |  |  |  |
| SNAP25 | 134583 | 3703098 | 6755588 | 481202 | - |
| SNAP23 | -6685971 | - | 6678049 | - | - |


| SOX11 | 4507161 | 4099263 | 6678065 | 2982742 | 2522255 |
| :--- | :--- | :--- | :--- | :--- | :--- |
| SO X4 (outgroup) | 4507163 | 7572947 |  |  |  |


| STAT1 | 6274552 | 3687402 | 6678153 | - | - |
| :---: | :---: | :---: | :---: | :---: | :---: |
| STAT2 | 4885615 | - | 6561853 | - | - |
|  |  |  | 6014655 |  |  |
|  |  |  | 5051642 |  |  |
| STAT3 | 4507253 | 3687429 | 1711553 | - | 6177821 |
| STAT4 | 4507255 | - | 1174461 | - | - |
| STAT5a | 4507257 | - | 6755672 | 4960028 | - |
| STAT5b | 6912688 | - | 7242209 | - | - |
| TNC | 4504549 | 1065718 | 7106435 | 135584 | - |
| TNXB | 7671639 | - | 7441741 | 1419546 | - |
| TNR | 5730098 | - | - | 86419 | - |

Table 1. continued

| WNT1 | 4885655 | 139740 | 139744 | - | 139748 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| WNT2a | 4507927 | 2501661 | 139751 | - | - |
| WNT2b | 13518017 | - | 6678591 | 5901876 | 3123031 |
| WNT3b | 6136371 | 263558 | 6678593 | 5821261 | 401416 |
| WNT3a | 6136340 | - | 7106447 | - | - |
| WNT118 | 4759320 | 7579033 | 6678589 | 1351423 | 1722841 |
|  |  | 3169687 |  |  |  |
| WNT10b | 5803223 | 263561 | 6756003 | - | - |
| WNT10a | - | 1175018 | 6678587 | 6141561 | - |
| WNT6 | - | - | 227508 | - | 401424 |
| WNT16 | 5732946 | - | 6249635 | - | - |
|  | 7706773 | - | - | - | - |
| WNT7a | 5509901 | - | 6678603 | - | 401418 |
| WNT7b | 6136361 | 263560 | 6678605 | 1245763 | 401419 |
| WNT7c | - | - | - | - | 401420 |
| WNT5a | 4507929 | - | 6678597 | 4512218 | 731158 |
| WNT5b | - | 2501662 | 6678599 | - | 465484 |
| WNT4 ${ }^{\text {a }}$ | - | 1351427 | 6678595 | 1351428 | 477511 |
|  |  | 4894948 |  |  |  |

${ }^{\text {a }}$ A well supported monophyletic group including human CDH12, Cad6 from M. musculus, and two divergent G. gallus sequences (cad10 and cad6b) did not show the expected organismal topology (CDH12 was the 'basal' sequence) and, therefore, may not be true orthologs.
${ }^{\mathrm{b}}$ CALM genes in the databases for human, mouse, chicken, and frog were identical. Thus, the placement of the mouse, chicken, and frog genes on the same row as CALM1 is arbitrary.
${ }^{\text {c }}$ BLastp turned up two zebrafish EVX genes. One was the sister sequence of the EVX1+EVX2 clade when Drosophila even-skipped (gi 123364) was used to root the tree.
${ }^{\mathrm{d}}$ GenBank included a short mouse sequence labelled Gdf6. The phylogenetic relationship between this gene and the GDF6 sequences included in table 1 was not resolved.
${ }^{\text {e }}$ For many Hox genes, only short conserved sequences that could not be placed within expected clades of orthologs were available (see § 4). Thus, in some cases, Hox genes are assigned to rows according to their names.
${ }^{\mathrm{f}}$ All $M S X$ genes shown formed a well-supported monophyletic group. However, the relationship between zebrafish msxD and msxA genes and the other MSX genes was not resolved.
${ }^{\mathrm{g}}$ WNT4 and WNT11 genes each form monophyletic groups with two zebrafish genes, but the tree topologies differ significantly from the expected organismal tree and may include two sets of orthologous genes as is the case for WNT2,WNT3, WNT5,WNT7 and WNT10 genes.

For ISL2, L1(CAM) and PAX2, zebrafish co-orthologues occur next to one another on the same chromosome (table 2). This observation suggests that duplicated ISL2, $L 1(C A M)$ and PAX2 genes in zebrafish were formed by tandem duplications. For this reason these three genes were not included in the estimate of the age of the fishspecific genome duplication reported above.

## 4. DISCUSSION

A genome duplication in the ancestor of teleost fishes is the most parsimonious explanation for the following observations: (i) many genes that occur once in chicken, mouse and man, and twice in Xenopus, a tetraploid frog, also occur twice in zebrafish; (ii) the phylogenetic analyses that were necessary to identify the two zebrafish coorthologues show, in most cases, that zebrafish genes are sister sequences as predicted by the genome duplication hypothesis; (iii) zebrafish co-orthologues are approximately the same age; and (iv) zebrafish co-orthologues are distributed throughout the zebrafish genome.

## (a) Gene number comparisons and gene tree topologies

The genome duplication hypothesis predicts that zebrafish will have more genes than humans. However, we found 140 cases among the 240 human genes included in our survey in which the database contained no zebrafish
orthologues. In a few cases (e.g. Hox genes) the shortage of zebrafish orthologues may be an artefact of our inability to assign some genes to specific clades. However, the shortage of fish genes is primarily due to the incomplete nature of the database: NCBI contains 1591 protein entries for zebrafish and 96009 protein entries for humans (23 November 2000).

Phylogenetic analyses identified 27 genes where orthologues that occur once in man, mouse and chicken, and often twice in Xenopus, also occur twice in zebrafish. For all of these genes, monophyly of the two zebrafish genes, plus orthologues from Xenopus, chicken, mouse and man, was well supported. For three of these genes, zebrafish coorthologues are closely linked. Therefore, despite our estimation that they are approximately the same age as the other duplicates, they are unlikely to have been produced by genome duplication. Although not all of the remaining 24 genes had the topology predicted by the fish-specific genome duplication hypothesis, most examples of the 'outgroup topology' are poorly supported by bootstrap reiterations and/or are not present when ML methods are used. A genome duplication event (or many gene duplications) prior to the Sarcopterygii-Actinopterygii divergence might explain the 'outgroup topologies' in figure $3 b$. However, if this is the case, then true orthologues of each of the 'basal' zebrafish genes must have been lost in Sarcopterygii. We believe it is more likely that some or all of the outgroup topologies shown in figure $3 b$ are tree


Figure 3. (See caption opposite.)


Figure 3. (Continued.) Phylogenies of duplicated fish genes. Trees were reconstructed using Poisson-corrected genetic distances and the neighbour-joining algorithm of Saitou \& Nei (1987) as implemented in Treecon (Van de Peer \& De Wachter 1994). Bootstrap values shown for nodes supported by more than $50 \%$ of 500 bootstrap reiterations (Felsenstein 1985). In all cases monophyly of the ingroup is well supported in an analysis that included other paralogues (see figure 2 ). The most closely related human paralogue was used to root the tree. (a) Phylogenies showing a sister sequence relationship for the zebrafish paralogues. Phylogenies of ISL2, L1 (CAM) and PAX2 genes had the same topologies as the genes shown here but the map positions of the zebrafish co-orthologues (table 2) suggest that they were not produced during genome duplication. (b) Phylogenies that include two zebrafish co-orthologues but not the expected sister sequence relationships. Maximum likelihood analyses (not shown) produce the duplication topology for FKD5, HOXC6 and SOX11.
reconstruction artefacts, perhaps caused by unequal rates of evolution in one of the zebrafish co-orthologues.

Synteny data indicate that zebrafish have two coorthologues for 10 human $H o x$ genes: B1, B5, B6, C6, B8, A9, A11, C11, A13, C13 (Amores et al. 1998). If these additional $H o x$ genes in zebrafish were produced by genome duplication, then we should have been able to reconstruct the 'duplication topology' for each of them. Instead, we found the topology predicted by the genome duplication hypothesis for only $H o x B 5$ and $H o x B 6$ genes (and for HoxC6 genes when ML methods were used). For Hox B1, HoxA11, HoxC11, HoxA13 and HoxC13, one or both of the zebrafish sequences in the database was 73 amino acids long or less and was comprised almost entirely of the highly conserved homeodomain, which is 60-63 amino acids long (Bürglin 1994). The lack of variation in
these short sequences precluded reliable tree reconstruction. For $H o x B 8$, only one zebrafish sequence ( $h o x B 8 b$ ) occurred in the database. For HoxA9 the two zebrafish genes, hoxA9a and hoxA9b, occurred within a wellsupported Hox9 clade and were sister sequences, but were not assigned to any of the four Hox9 clades.

Gates et al. (1999) and Barbazuk et al. (2000) included Hes5 among their list of genes with two zebrafish coorthologues. Both studies report that zebrafish genes her2 and her 4 are orthologous to mouse Hes5. However, our BLASTp searches turned up three additional zebrafish genes (her1, her3 and her7) that cluster with mouse Hes5 and the topology of the expanded tree (whether based upon NJ or ML methods) does not support the hypothesis that any pair of zebrafish genes are co-orthologues of mouse Hes5.
(b)


Figure 3. (Continued.)
(b) Age of co-orthologues

Since additional Hox clusters are present in both zebrafish and Takifugu (see § 1), the fish-specific genome duplication is believed to have happened before the divergence
of Cypriniformes (zebrafish) and Tetraodontiformes (Takifugu), at least 150 Myr ago (Nelson 1994; Cantatore et al. 1994). On the other hand, the duplication most probably took place after the divergence of ray-finned and


Figure 4. Substitutions at third-codon positions plotted against divergence dates (see § 2) for taxa included in this study. The divergence of Actinopterygii and Sarcopterygii (ca. 450 Myr ago) was excluded because third positions are saturated and the inclusion of these data would erroneously influence the regression. The average number of third-codon position substitutions between pairs of zebrafish co-orthologues is $1.02($ s.d. $=0.24)$.
lobe-finned fishes, ca. 450 Myr ago (Kumar \& Hedges 1998; Lee 1999), since all sarcopterygian species studied so far have four or fewer Hox gene clusters. This is consistent with our observations that for many phylogenetic trees, zebrafish paralogues appear to have been formed during the time interval between the divergence of amphibians and amniotes, and the divergence between reptiles (i.e. birds) and mammals (figure $3 a$ ).

A comparison of synonymous and non-synonymous substitutions in duplicated genes of varying ages and from a diversity of species suggests that genes experience a period of accelerated evolution shortly after gene duplication (Lynch \& Conery 2000). Acceleration in the rate of evolution of both zebrafish genes compared with frog, chicken, mouse and human genes might mean that the genome duplication is younger than it appears to be on our phylogenies (though an increase in non-synonymous mutations following a duplication event should not affect our genetic distance estimates based upon third-codon positions). Allotetraploidy might have also confounded our ability to date the fish genome duplication. Gene duplication (i.e. tetraploidy) occurs when cytokinesis fails during the first mitotic division of a fertilized egg (Sheppard et al. 1982). In autotetraploidy, 'duplicate' genes come from two individuals of the same species and are identical or are alleles at a given locus. With allotetraploidy the two genomes involved come from different species and may have diverged extensively at the faster-evolving loci before the tetraploidy, i.e. duplication event (Spring 1997). Thus, for genome duplication via allotetraploidy, divergence between coorthologues begins before the tetraploidy event (i.e. genome duplication).

Despite these possible sources of error in the estimation of the fish genome duplication, our estimate that the duplicated zebrafish genes are between 300 and 450 million years old indicates that genome duplication preceded the teleost radiation. Study of 'basal' actinopterygians (e.g. bichir, sturgeon, bowfin, gar) will help to
determine more accurately the date of the fish genome duplication.

## (c) Gene location

Comparative genomics has provided many new insights into the evolution of chromosomes. Radiation hybrid maps have shown that there are orthologous chromosome regions in human and mouse (Nadeau \& Sankoff 1998), in human and cat (Murphy et al. 2000), human and cattle (Band et al. 2000), and in human and zebrafish (Barbazuk et al. 2000). Genome duplication means that many species also possess paralogous chromosome regions (e.g. Morizot et al. 1991; Lundin 1993; Amores et al. 1998; Pébusque et al. 1998). Indeed, the term 'co-orthology' can be applied to regions of chromosomes as well as genes.

The duplicated zebrafish genes uncovered in this study occur on a large proportion of the 25 zebrafish linkage groups, but they do not appear to be randomly distributed in the zebrafish genome. Our phylogenetic data indicate that regions of zebrafish LG1 and LG9, LG2 and LG7, LG3 and LG12, LG11 and LG23, LG17 and LG20 are paralogous (table 2).

## (d) The retention and loss of duplicated genes

Several models have been proposed to explain the evolutionary persistence of duplicated genes in zebrafish. Gibson \& Spring (1998) argue that selection can prevent the loss of redundant genes (i.e. duplicates) if those genes code for components of multidomain proteins because mutant alleles disrupt multidomain proteins (i.e. are dominant negative mutations). Force et al. (1999) argue that when a gene with multiple functions is duplicated, the duplicates are redundant only for as long as each retains the ability to perform all ancestral roles. When one duplicate experiences a mutation that prevents it from carrying out one of its ancestral roles, the other duplicate is nolonger redundant. This is consistent with Sidow's (1996) proposition that a single unique function in an ocean of redundancy is enough to keep the gene afloat and prevent degenerative substitutions. According to Force et al.'s (1999) 'duplication degeneration-complementation' model, degenerative mutations preserve rather than destroy duplicated genes. Force et al. (1999) present EN1 as an example of their model. Zebrafish engla and englb appear to have divided the roles of their orthologues (e.g. human $E N 1$ ). It will be interesting to find out if the other coorthologues reported here have divided the roles of their sarcopterygian orthologues or are components of multidomain proteins. De Pinna (1996) provided a list of teleost synapomorphies. One convincing way to show that extra genes originating from genome duplication were responsible for the radiation of Teleostei would be to demonstrate that duplicated genes code for teleost-specific traits.

An alternative evolutionary link between the teleost radiation and genome duplication involves 'divergent resolution' (Lynch \& Conery 2000; Taylor et al. 2001). Lynch and Conery proposed that the loss of different duplicates in geographically isolated populations could reduce the fecundity of hybrids. They considered a young pair of functionally redundant, unlinked, duplicate genes in an ancestral species. One of the two duplicates is likely to be silenced (i.e. become a pseudogene) within the next one

Table 2. Genome location and genetic distance between pairs of co-orthologous genes.
(Map data were obtained from the Zebrafish Information Network: http://zfish. uoregon.edu/ZFIN/, Gates et al. (1999) and Barbazuk et al. (2000). Symbols denote possible paralogous chromosomes. 'Confidential' means that the gene has been mapped but data are not available. Genetic distances were computed using only third codon positions and corrected for multiple events per site according to Tajima \& Nei (1984). Estimated number of mutations per site are shown for $\operatorname{ISL2,} \operatorname{L1}(C A M)$ and PAX2 but these data are not included in the calculation of the mean because these zebrafish co-orthologues were probably produced by independent tandem duplications. Woods et al. (2000) recently reported that the two zebrafish ISL2 genes and the two zebrafish Pax2 genes do not occur on the same linkage groups (contrary to Barbazuk et al. 2000). Our phylogenies of ISL2 and Pax2 genes were consistent with the fish-specific genome duplication hypothesis (i.e. 'duplication topology' with high bootstrap support for all nodes) and the Tajima-Nei distance estimates for the ISL2 and Pax2 duplicates (table 2) are approximately the same as those for the other unlinked duplicates.)

|  | symbol | symbol (zebrafish) | location (zebrafish) | Tajima-Nei distance |
| :---: | :---: | :---: | :---: | :---: |
| 1 | BMP2 | bmp $2 a$ | LG 17 • | 1.207 |
|  |  | bmp2b | LG 20 - |  |
| 2 | BRN1 | brn1.1 | LG 9 | 1.119 |
|  |  | brn1.2 | LG 6 |  |
| 3 | DLL1 | dla | LG 1 | 1.233 |
|  |  | dld | LG 13* |  |
| 4 | D LX2 | $d l x 2$ | LG 9 | 1.364 |
|  |  | dlx 5 | LG $1 \dagger$ |  |
| 5 | EN1 | engla | LG 9 | 0.931 |
|  |  | eng 16 | LG $1 \dagger$ |  |
| 6 | EN2 | eng2 | LG 7 | 1.199 |
|  |  | eng3 | LG $2 \Psi$ |  |
| 7 | EPB4 | rtk4 | unmapped | 0.975 |
|  |  | epa4 | unmapped |  |
| 8 | FKD5 | fkd3 | LG 25 | 1.027 |
|  |  | fkd5 | unmapped |  |
| 9 | FLOT1 | re2a | unmapped | 0.720 |
|  |  | re2b | unmapped |  |
| 10 | Hedgehog | shh | LG 7 | 1.389 |
|  |  | twh | LG $2 \Psi$ |  |
| 11 | HOXB5 | hoxb5a | LG 3 | 0.749 |
|  |  | hoxb5b | LG 12Ф |  |
| 12 | HOXB6 | hoxb6a | LG 3 | 0.876 |
|  |  | hoxb6b | LG 12Ф |  |
| 13 | HOXC6 | hoxC6a | LG 23 | 1.009 |
|  |  | hoxC6b | LG $11 \Theta$ |  |
| 14 | JАК2 | jak2a | confidential | 1.054 |
|  |  | jak2b | confidential |  |
| 15 | LHX1 | lhxl | LG 15 | 1.089 |
|  |  | lim6 |  |  |
| 16 | $m s x 3$ (mouse) | msxb | LG 1 | 1.590 |
|  |  | msxc | LG 13* |  |
| 17 | NTN1 | ntn1 | LG 3 | 0.863 |
|  |  | ntn1a | LG 6 |  |
| 18 | OTX1 | otx 1 | LG 17 | 1.047 |
|  |  | otx 3 | LG 1 |  |
| 19 | RARA | rara2a | LG 12 | 0.964 |
|  |  | rara2b | LG $3 \Phi$ |  |
| 20 | RXRB | rxre | LG 19 | 0.931 |
|  |  | rxrd | unmapped |  |
| 21 | SNA(il) | snail1 | LG 11 | 0.809 |
|  |  | snail2 | LG $23 \Theta$ |  |
| 22 | SNAP25 | snap 25,1 | LG 20 • | 0.594 |
|  |  | snap 25,2 | LG 17 • |  |
| 23 | SOX11 | sox11a | LG 17 • | 0.749 |
|  |  | sox11b | LG $20 \bullet$ |  |
| Mean (s.d.) |  |  |  | 1.02 (0.23) |
|  | gdf6 | dynamo | LG 19 | NA |
|  | (bovine) | radar | confidential |  |
|  | ISL2 | isl2 | LG 25 | 1.128 |
|  |  | isl3 | LG 25 |  |
|  | L1(CAM) | 11.1 | LG 23 | 1.187 |
|  |  | 11.2 | LG 23 |  |
|  | PAX2 | pax2 | LG 13 | 0.873 |

to two million years. If the ancestral species is divided into geographically isolated populations, then a different copy of the duplicated gene could become fixed in the two populations. If the two populations hybridize, the $\mathrm{F}_{1}$ progeny would be heterozygous in two respects. With respect to homologous chromosomes, one homologue would have a functional allele and the other a pseudogene. With respect to the entire genome, an $\mathrm{F}_{1}$ individual would have two functional alleles of the locus but those alleles would occur on different chromosomes. In the $\mathrm{F}_{2}$ generation, there is a $6.25 \%$ chance that an individual will receive only pseudogenes of a given duplicated and differentially resolved gene. If the gene in question is an essential gene, then $6.25 \%$ of the $\mathrm{F}_{2}$ generation would not survive. Furthermore, $25 \%$ of $\mathrm{F}_{2}$ individuals may also suffer reduced fitness because they would be haploid at this locus. Lynch \& Conery (2000) stated that with tens to hundreds of young unresolved gene duplicates present in most eukaryotic genomes, such genes could provide a common substrate for the passive origin of isolating barriers. However, genome duplication (e.g. in the ancestor of teleost fishes) provides many more than tens to hundreds of unlinked, duplicated genes. Divergent resolution of thousands of genes might be a very powerful isolating mechanism. One prediction of this model in which genome duplication leads to speciation is that tetraploid taxa should have more species than their diploid sister groups.

## (e) Terminology

In this paper we have adopted the term 'co-orthologue' (Gates et al. 1999). In our opinion, this term is useful because it conveys information about genome duplications that is not obvious from the term 'orthologue'. Supplements to orthology and paralogy have also been introduced by Holland (1999) and Sharman (1999): 'pro-orthologue' describes the relationship of a gene to one of the postduplication descendants of its orthologue. Human RARA is, for example, a pro-orthologue of the zebrafish genes rara $2 a$ and the rara $2 b$ (figure 2). 'Semi-orthologue' describes the relationship of one of a set of duplicated genes to a gene directly descended from the ancestor of the whole set (e.g. rara $2 a$ is semi-orthologous to $R A R A$ ). Because semi-orthologue implies 'half orthologue' it might be a more appropriate term than co-orthologue for comparisons between diploid fish genes and their human pro-orthologues. Such a naming approach could be extended to include other genic relationships. For example, genes in most actinopterygians might be considered 'octalogues' of their respective orthologous genes in invertebrates. However, attempts to describe such gene relationships numerically can become awkward. For example, how would the relationship between genes in tetraploid fish such as the goldfish (Carassius auratus) and genes in Drosophila be described? In this case a 1:16 gene ratio is expected, based upon the four genome duplications that probably separate these species. Even for a species between which a 1:2 or a 1:4 gene ratio is expected based upon genome duplication data, tandem duplications can disrupt the actual orthologue ratio. Therefore, we prefer the terms pro-orthologue and coorthologue to describe relationships between genes in taxa separated by any number of tandem or genome duplications.

## (f) Problems with gene nomenclature

Our conclusion that there was a genome duplication event in fish means that all genes in actinopterygian fish have co-orthologous relationships with their sarcoterygian (e.g. human) orthologues. Currently the names of many zebrafish genes reflect their co-orthologous relationship to orthologues or 'pro-orthologues' in sarcopterygians (e.g. $b m p 2 a$ and $b m p 2 b$; eng $1 a$ and eng $1 b$ ). However, in many cases the fact that a given zebrafish gene is one of two orthologues is not clear from its name. For example, the following pairs of genes were shown to be co-orthologues in our study: $d l a$ and $d l d, d l x 2$ and $d l x 5$, eng2 and eng $3, i s l 2$ and $i s l 3$, $r x r E$ and $r x r D$, shh and twhh, otx1 and otx 3 , $f k d 3$ and $f k d 5$, and dynamo and radar.

We propose all genes in diploid fish be given the same name as pro-orthologues in humans but that these names be appended with an ' $a$ ' or ' $b$ ' designation to reflect their co-orthologous relationships with human (and other sarcopterygian) genes. In cases where only one co-orthologue appears to have been retained, the ' $a$ ' designation serves as a reminder of the genes' duplication history.

Tiggy-winkle hedgehog (Ekker et al. 1995) highlights the potential confusion generated when the name of a gene lacks phylogenetic information. Tiggy-winkle hedgehog (twhh) and sonic hedgehog (shh) in zebrafish are equally orthologous (i.e. co-orthologous) to sonic hedgehog (SHH) in humans (present study; Zardoya et al. 1996). A PubMed search suggests that this fact is not widely appreciated: 29 references include the terms; shh+zebrafish and only five include twh + zebrafish. Furthermore, a gene named 'twhh' has been sequenced in goldfish. However, goldfish twhh cannot be orthologous to zebrafish twhh, as might be expected from its name, because goldfish are tetraploid (Zhang et al. 1999). That is, the goldfish twhh that has been sequenced can only be co-orthologous to zebrafish twh (i.e. one of two twhh co-orthologues).

Our phylogenetic study also turned up naming 'errors' in genes for which only one co-orthologue is currently known. Zebrafish rxra clusters with strong bootstrap support within the $R X R c$ clade. Conversely, zebrafish $r \times r c$ clusters with strong support within the $R X R a$ clade. As this list of confusing and erroneous names grows a complete review of fish gene nomenclature will become increasingly important just as it was for Hox genes in 1992 (De Robertis 1994).

Woods et al. (2000) recently reported that the two zebrafish $I s l 2$ genes and the two zebrafish Pax2 genes do not occur on the same linkage groups (contrary to Barbazuk et al. 2000). Our phylogenies of $I s l 2$ and Pax2 genes were consistent with the fish-specific genome duplication hypothesis (i.e., 'duplication topology' with high bootstrap support for all nodes), and the Tajima-Nei distance estimates for the $I s l 2$ and Pax2 duplicates (table 2) are approximately the same as those for the other unlinked duplicates.

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