Developmental Constraints on Vertebrate Genome Evolution

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Abstract

Constraints in embryonic development are thought to bias the direction of evolution by making some changes less likely, and others more likely, depending on their consequences on ontogeny. Here, we characterize the constraints acting on genome evolution in vertebrates. We used gene expression data from two vertebrates: zebrafish, using a microarray experiment spanning 14 stages of development, and mouse, using EST counts for 26 stages of development. We show that, in both species, genes expressed early in development (1) have a more dramatic effect of knock-out or mutation and (2) are more likely to revert to single copy after whole genome duplication, relative to genes expressed late. This supports high constraints on early stages of vertebrate development, making them less open to innovations (gene gain or gene loss). Results are robust to different sources of data—gene expression from microarrays, ESTs, or in situ hybridizations; and mutants from directed KO, transgenic insertions, point mutations, or morpholinos. We determine the pattern of these constraints, which differs from the model used to describe vertebrate morphological conservation (“hourglass” model). While morphological constraints reach a maximum at mid-development (the “phylotypic” stage), genomic constraints appear to decrease in a monotonous manner over developmental time.

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Introduction

To what extent do the processes of embryonic development constrain genome evolution? Correlations between developmental timing and morphological divergence have long been observed, but the mechanisms and molecular basis of such patterns are poorly understood. The most commonly used measure of selective pressure on the genome, the ratio of non-synonymous to synonymous substitutions (dN/dS) in protein coding genes, has been of limited help in this case. Stronger constraints have been found on genes expressed in late embryonic stages in Drosophila [1], but most other studies have failed to report robust evidence for a lower dN/dS ratio in genes expressed at constrained developmental stages [2–5]. A different approach has been to characterize which genes are duplicated, and which are not: studies of C. elegans [2] and Drosophila [6] have found less duplication of genes expressed in early development. These results show that it is possible to identify developmental constraints at the genomic level. They have a few limitations though. One is that the data available has limited the characterization of developmental time to broad categories such as “early” and “late”. A second is the difficulty of relating results from two derived invertebrate species, to morphological evolution models in vertebrates [7].

Indeed it is in vertebrates that the fundamental models of developmental constraint on evolution have been established, starting in the nineteenth century with the “laws” of von Baer [8], claiming a progressive divergence of morphological similarities between vertebrate embryos, with the formation of more general characters before species-specific characters. Integration of these observations within evolutionary biology has not always been straight-forward [9–11]. More recently, an “hourglass” model was proposed to describe morphological evolution across development [12,13]: in the earliest stages of development (cleavage, blastula) there is in fact a great variety of forms in vertebrate embryos. Later in development, a “phylotypic” or conserved stage is observed, where many morphological characteristics are shared among vertebrates. This stage is usually presumed to be around the pharyngula stage. After this bottleneck, a “von Baer-like” progressive divergence is again observed. The conserved phyloptypic stage has been explained by assuming higher developmental constraints [13–15]. The limits on morphological evolution would be placed by the structure of animal development, making some changes unlikely or impossible. How such limitations are encoded in the genome, or impact its evolution, is still an open question.

In this work, we investigate the existence and timing of constraints on genes expressed in vertebrate development. We use representatives of the two main lineages of vertebrates, a teleost fish and a tetrapode, and we explore the impact of experimental gene loss, and of gain of gene copies in evolution. We find that timing of development has a strong impact in both cases, but that the pattern of constraints on genome evolution does not follow the morphological hourglass model. High constraints are present in early stages of development and relax progressively over time.

Results

Constraints on Gene Loss-of-Function in Zebrafish

First, we used the phenotypes of gene loss-of-function as an indicator of selective pressure on genes. We extracted genes
essential for the viability of the zebrafish, giving a lethal phenotype when non functional [16]. We expect that the loss of a gene should be more deleterious if this gene is expressed at a developmental stage with strong constraints. Thus we estimated whether genes were expressed or not at each stage, and computed the ratio of expressed essential genes to expressed reference genes (no reported loss of function phenotype). We then plotted the variation across development of this ratio. We used two different types of data to evaluate the presence of gene expression: (i) expression patterns from in situ hybridizations (Figure 1A), and (ii) “present” or “absent” calls from an Affymetrix microarray experiment (Figure 1B). Results are consistent for both data types: the proportion of essential genes is higher among genes expressed in early development, with a significant negative correlation. For the in situ hybridizations (Figure 1A), a linear regression is significant, but a parabola is not. The parabola has been suggested as the quantitative expectation of an hourglass-like model [3,17]. These results indicate a continuous trend over developmental time, with stronger constraints on early development.

Constraints on Gene Loss-of-Function in Mouse

We performed a similar analysis in mouse, with some differences of methodology due to the data available. For

![Figure 1. Variation across zebrafish development of the expression of essential genes compared to non-annotated genes.](https://www.plosgenetics.org/doi/10.1371/journal.pgen.1000311.g001)

**A** In-situ

| Time (hours) | Ratio Essential/Non-Annnotated |
|-------------|-------------------------------|
| 0           | 0.12                          |
| 11          | 0.08                          |
| 12          | 0.06                          |
| 13          | 0.04                          |
| 14          | 0.02                          |
| 15          | 0.01                          |
| 16          | 0.00                          |

**B** Microarray

| Time (hours) | Ratio Essential/Non-Annnotated |
|-------------|-------------------------------|
| 0           | 0.08                          |
| 11          | 0.06                          |
| 12          | 0.04                          |
| 13          | 0.02                          |
| 14          | 0.01                          |
| 15          | 0.00                          |

*Figure 1. Variation across zebrafish development of the expression of essential genes compared to non-annotated genes.* At each time point, the ratio of the number of essential genes expressed on the number of non-annotated genes expressed is plotted. A gray box on the x-axis indicates the phylotypic period. (A) Gene expression as reported using in situ hybridization data. The x-axis is proportional to time. A weighted linear regression was fitted to the data and the regression line plotted. (B) Gene expression as reported by “present” or “absent” calls from Affymetrix array data. The x-axis is in logarithmic scale. A Spearman correlation was computed (coefficient ρ). doi:10.1371/journal.pgen.1000311.g001
expression, we used of a large amount of EST (Expressed Sequence Tags) data from libraries spanning development, from which we deduced presence or absence of expression (see Methods). Only phenotypes obtained by the targeted knock-out technique were used. As knock-out experiments with no observable phenotype are reported in mouse, we can use these as a reference set, instead of non annotated genes as in zebrafish. The ratio of expressed essential genes to expressed reference genes is significantly negatively correlated with developmental time (Figure 4A), as in zebrafish (Figure 1).

Repeating the same approach with genes inducing a phenotype reported as “abnormal” when they are not functional, no significant trend is detected compared to genes inducing no phenotype, after multiple testing correction (Figure 4B). Moreover, these genes can be used as a reference for essential genes (Figure 4C), with results very similar to the use of genes inducing no phenotype after loss of function (Figure 4A). Thus in mouse, genes inducing abnormal phenotypes when non-functional have a behavior more similar to the reference set of “non essential” genes.

**Constraints on Gene Duplication**

The fish specific whole genome duplication [19] provides us with a natural experiment on constraints on gene doubling: after this event approximately 85% of duplicated genes lost one copy, and the subset which retained both copies is known to be biased relative to function and selective pressure [20]. Thus we tested if duplicate gene expression pattern in zebrafish development was biased compared to singletons. We plotted the median expression profiles of duplicates originating from the fish specific whole genome duplication, and of singletons, genes whose duplicate copy has been lost after the genome duplication (Figure 5). Duplicates are less expressed in early stages of development. The difference of median expression decreases progressively, similar to the observations for essential or abnormal phenotype genes. Larval time points show a maximum expression of duplicates relative to singletons.

Two scenarios can explain this result. First, retention of two copies may be more likely after the whole genome duplication for genes less expressed in early development. Second, the retention of genes may be unbiased relative to development, but duplicate genes may evolve secondarily lower expression in early development. To get a proxy of the ancestral state before whole genome duplication, we used again mouse data, which has diverged from zebrafish before the fish specific duplication. We compared mouse orthologs of zebrafish duplicates to mouse orthologs of zebrafish singletons, regarding their expression in development (Figure 6). Mouse orthologs of duplicates are significantly less expressed in early development compared to orthologs of singletons. This result in mouse is consistent with the observations in zebrafish, and the most parsimonious explanation is that expression was similar in the ancestor of the two lineages. Therefore we can accept the first hypothesis: after the fish specific whole genome duplication, there was preferential retention of duplicates less expressed in early development.

To check if this phenomenon is particular to the fish specific genome duplication, we repeated this analysis with the two ancient rounds of genome duplication (“2R”), which occurred in the ancestor of vertebrates [21]. It is difficult to distinguish between the two whole genome duplications since no model species diverged from the vertebrate lineage between them. Therefore we looked at the median expression profiles of genes with any duplication at the origin of vertebrates, compared to singletons, whose duplicates were lost after both whole genome duplications. For zebrafish, we restricted this analysis to genes which are singletons regarding the fish specific whole genome duplication. Similarly to fish specific duplicates, duplicates from 2R are significantly less expressed than singletons in the early development of zebrafish (Figure S1) and mouse (Figure S2). Thus
mechanisms of retention after whole genome duplication seem to be conserved during vertebrate evolution (see also Text S1).

**Constraints on Gene Sequence**

To check if sequences of genes expressed at different stages in development are experiencing different selective pressure, we used the non synonymous to synonymous substitution ratios ($d_{NS}/d_S$). In zebrafish, we used an approach similar to Davis et al. [1]: at each stage we performed the correlation between $d_{NS}/d_S$ and gene expression from microarray data (Figure S3). It has been shown that genes retained in duplicate tend to evolve slowly [20,22]. To control for that factor, we kept only strict singletons in the analysis (genes whose duplicate was lost after 2R and fish-specific genome duplications). At all stages the correlation is negative, confirming that genes with higher expression levels are under stronger purifying selection [23,24]. We note that correlation at the “adult” stage (90 days) is weaker (Figure S3): the link between expression and selective constraints on sequences appears stronger in development than in adult. But there is not a significant trend over time (Spearman $r = 0.08; p = 0.68$).

In mouse, we considered only singletons after 2R genome duplication, and we compared the slowest evolving genes (25% lower $d_{NS}/d_S$) with the fastest evolving genes (25% higher $d_{NS}/d_S$). There is a significant correlation with time of expression (Figure 7). Genes with strong sequence constraints (low $d_{NS}/d_S$) tend to be expressed early in development.

Figure 3. Expression of four groups of genes, clustered according to their expression in zebrafish development.
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What is the function of the genes whose evolution is constrained by expression in early development? We analyzed enrichment or depletion in Gene Ontology [25] categories for the clusters based on gene expression (Figure 3). Using the Molecular Function ontology, genes whose expression is highest in early development are significantly enriched in fundamental processes of the cell, such as RNA processing, transcription, and DNA replication (Table S1). This is very similar to the categories observed to be enriched in housekeeping genes [26]. It is also consistent with the categories depleted in fish specific duplicates [20]. Conversely, genes highly expressed in early development are depleted in receptor or channel activity, while these activities are enriched in genes highly expressed in late development. Fewer terms are significant for the Biological Process ontology, and results are essentially consistent with the Molecular Function. Overall, the genes expressed in early development, which appear constrained against gene duplication or loss of function, seem to be housekeeping genes involved in basic cellular processes.

Discussion

Recent discussion of the evolution of ontogeny [27] has allowed the clarification of several important points. The first is that models must be explicitly defined, to allow testing. Poe and Wake [17] distinguish three models for the evolution of ontogeny: the
Constraints on Genome Evolution

Figure 6. Variation across mouse development of the ratio of expressed orthologs of zebrafish singletons after the fish specific genome duplication (FSGD) relative to orthologs of zebrafish duplicates. Legend as in Figure 4.

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early conservation model à la von Baer [8]; the hourglass model, characterized by a conserved phylotypic stage [12,28]; and the adaptive penetrance model (an inverted hourglass). The second point is that quantitative testing is important to distinguish between these models. At the morphological level, several studies have used heterochrony data from vertebrates to quantify the amount of change at each stage of development [17,29]. Surprisingly, this led to rejection of both the early conservation and the hourglass models, although which model is favoured remains disputed [27]. The third point that should be clarified is the distinction between constraints at the level of patterns, and constraints at the level of processes [29]. The studies of heterochrony in vertebrates are typically concerned with the pattern.

In this framework, our results clearly provide a quantitative test which supports the early conservation model. By studying not morphological structures but features of the genome and its expression, this test concerns the level of processes, not patterns. Thus an important point to be made is that our results should be taken neither in contradiction nor in support of any specific model at the level of patterns, given our still limited knowledge of causal relationships between process and patterns in ontogeny [30]. On the other hand, our results do appear to be in contradiction with previous reports of a maximum of constraints on processes around the phylotypic stage of vertebrates [3,4,31].

We use two simple measures of constraint on the expression of a gene at a developmental stage: if expression of one copy is needed, then (i) removing it may be deleterious, and (ii) increasing the number of copies may also be deleterious. This view is consistent with a recent study in yeast which suggests that constraints influencing the ability to lose certain genes or to maintain them in duplicate may be similar [32]. We expect gain or loss of genes highly expressed at more constrained developmental stages to be counter-selected. And indeed, we find a clear and significant trend: early development is strongly constrained, then constraints diminish during development in a continuous manner. Genes highly expressed in early development are more frequently essential, and less frequently preserved in double copy after genome duplication. Thus early development is less robust against gene loss and against gene doubling. Trends are conserved between mouse and zebrafish, representatives of the two main lineages of bony vertebrates, and between 2R and fish specific genome duplications. An indication of how strong these constraints are is our capacity to predict which genes were kept in duplicate in zebrafish based on expression pattern in mouse. Despite more than 400 MY of independent evolution, and the use of relatively noisy data (mix of EST libraries), more than a quarter of the variance in gene retention is explained (Figure 6; $r^2 = 0.27$). There is also some signal for early conservation at the level of coding sequences, at least in mouse (Figure 7). What we do not see is any genomic evidence for specific constraints at a phylotypic stage.

Both in zebrafish and in mouse, the pharyngula stage appears to be part of the general trend from stronger genomic constraints in early development, towards weaker genomic constraints at later stages. We believe that our data are sufficiently detailed, and exhibit sufficiently strong signal, that a maximum of genomic constraints at the phylotypic stage would be visible. So where does the contradiction with previous studies come from?

An early quantitative study [31] found that when screens were done in rodents for the induction of teratogenesis, most abnormalities were obtained by applying teratogens during the phylotypic stage. This was interpreted [31] as supporting strong constraints at the phylotypic stage, due to inductive interactions. But these screens aimed not to test developmental robustness, but to obtain abnormal embryos for experimental work. As remarked by Bininda-Emonds et al. [29], Galis and Metz [31] define the phylotypic stage broadly as including most organogenesis. If application of teratogens in early development resulted in lethality before organogenesis, it would not be of interest to the researchers performing the screens. Thus it seems that what Galis and Metz [31] measured was the potential for a stage to produce morphological abnormalities, not the overall constraints on ontogeny at each stage. There seems to be little reason to suppose that such data provide “an accurate model of natural selection” [33], unlike e.g. the retention of duplicate genes over long evolutionary periods.

It is worth noting that we observe a “peak” of constraints shortly after pharyngula (Figure 4B) for the expression profile of mouse genes which give an “abnormal” phenotype when knocked-out. The behavior of these genes is surprising, because in zebrafish the
trend for such genes was similar to that for essential genes. We suspect that the definition of abnormal phenotypes differs between databases and between investigators working in different species. Less severe phenotypes may be reported as “abnormal” in mouse, relative to zebrafish. Of note, data in ZFIN [16] come mainly from the reviewed literature, where minor abnormalities of phenotype are rarely reported, whereas data in the MGD [34] come also from genome wide mutagenesis, and thus include such minor abnormalities. Minor abnormalities in mouse phenotype may also be easier to detect because of the gross similarity with human in anatomy and physiology. In any case, these are the data in our study which most closely approximate the teratogenesis study, and the only data that do not support the early conservation model. Although this trend is statistically not significant, it is consistent with the observations of Galis and Metz [31]. This deserves to be further examined in future studies.

Two other studies which quantified a maximum of constraints at the phylotypic stage did use evolutionary measures of constraint. These studies [3,4] estimated constraints on the evolution of coding sequences, in relation to the timing of expression in mouse development from EST data. Despite similar experimental designs and data, we reached differing conclusions. First, we note that we did check for sequence conservation \( K_{R/N} \) trends over development. In zebrafish, we found no robust pattern (Figure S3), while in mouse we found support for the early conservation model (Figure 7). Second, in our analyses we found that small samples of ESTs could introduce important variability, which is why we used weighted regressions for all computations based on these data. For example, we see a very high ratio of mouse orthology of zebrafish singletons to duplicates for Thélier stage 5 (day 4) (Figure 6); but this is obtained based on only 628 genes with at least one EST at that stage (median over all stages: 3767). The weighted regression insures that such a point has a weak incidence on the statistical significance. Similar issues are visible in the data of Irie et al. [4], but are not addressed in their analysis. Indeed, the extreme points they use to support constraints at pharyngula are based on some of the smallest samples of their dataset. Finally, it should be noted that another study in mouse found an opposite pattern (relaxation of constraints near the phylotypic stage) using an alternative model [43]. This experiment uses an Affymetrix GeneChip Zebrafish Genome Array (A-AFFY-38). 15 stages were sampled, spanning from fertilization to adult stages (15 minutes, 6, 8, 9, 10, 11.7, 16, 24, 30 hours, 2, 4, 5, 14, 30, 90 days, covering zygote, segmentation, gastrula, pharyngula, hatching, larval, juvenile, adult). Two replicates were made per time point; we use both of them for computations, and the 2 values are plotted to give an order of the variability between replicates.

Raw CEL files were renormalized using the package gcRMA [44] of Bioconductor version 2.2 [45]. We used the “affinities” model of gcRMA, which uses mismatch probes as negative control probes to estimate the non-specific binding of probe sequences. The normalized values of expression are in \( \log_2 \) scale, which attenuates the effect of outliers. Mapping of \( D. \ rerio \) genes on Affymetrix probesets was made using Ensembl [46] annotation for zebrafish genome version Zv7 (unpublished).

We did not consider the first time point of the data (15 minutes, fertilization). Its behaviour was peculiar in many cases. We explain this by the presence of maternal transcripts in the embryo [47]. These transcripts are largely degraded by 6 hours of development [48], the second time point of the dataset. For the absolute detection of transcripts (presence or absence calls), the method we used [49] replaces all MM probe values by a threshold value which is based on the mean PM value (after gcRMA transformation) of probesets that are very likely to have absent target transcripts. This removes the influence of probe sequence affinity and results in better performance than the MAS 5 algorithm.

Significance of Trends in Zebrafish Development

For the zebrafish microarray data we first used a randomization approach to assess the significance of the difference between two curves of median expression across development (for example median expression of duplicates vs. singletons, or of essential genes vs. genes with no reported phenotype). If the two groups contain \( n_1 \) and \( n_2 \) genes, we pooled all these genes and randomly separated them into two new groups of same sizes \( n_1 \) and \( n_2 \). Then we calculated and recorded the difference between the two new curves of median expressions across development. After repeating this randomization 10,000 times, we could define 1%no and 1% confidence intervals.

Second, we calculated the Spearman correlation between developmental time and the difference between two curves of median expression across development. Bonferroni correction was applied to correct for multiple testing, considering the 9 tests computed with this microarray data (Figure 1; Figure 2; Figure 5; Figure S1; Figure S3; Figure S5A–D): \( \alpha = 0.05/9 = 0.0056 \).

Clustering of Microarray Data

In order to identity genes locally or highly expressed in early development, we used the Fuzzy C-Means soft clustering method implemented in the Bioconductor package Mfuzz [50]. After a
pre-filtering step (genes with sd <0.5 were removed), we ran the algorithm with the number of clusters set to c = 4. This gave one cluster of genes locally expressed across development (3641 probesets, 2261 Ensembl genes), one of genes highly expressed (2175 probesets, 1175 Ensembl genes), one of genes whose expression increased (1714 probesets, 1123 Ensembl genes) and one of genes whose expression decreased (3906 probesets, 2446 Ensembl genes) (Figure 3).

Mouse EST Data

EST (Expressed Sequence Tags) data were retrieved from BGEE (dataBase for Gene Expression Evolution, http://bgee.unil.ch/), a database comparing transcriptome data between species [51], including EST libraries from UniGene [52]. The mapping of UniGene clusters on Ensembl genes is taken from Ensembl (version 48) [46], where a percentage of identity of 90% is set as the minimum threshold to link an Ensembl gene with a UniGene cluster. Each library has been annotated manually to ontologies of anatomy and developmental stages, if it was obtained under non-pathological conditions, with no treatment (“normal” gene expression). We considered a gene expressed at one time point in development if at least one EST was mapped to this gene at this time point. Thus, we could retrieve the number of genes expressed at each time point of mouse (Mus musculus) development. From this set we extracted two groups to compare (for example essential/non-essential, or duplicates/singletons). As the total number of ESTs available at each time point is different, we use at each time point the ratio of the numbers of genes expressed in the two groups. We obtained similar results when we defined a gene as expressed if it had at least two ESTs mapped to it. Also, considering the ratio of the mean number of ESTs per gene at each stage, instead of the ratio of the number of genes expressed at each stage, gave similar results (not shown). We used data from 297 EST libraries, spanning 26 different developmental stages (from TS01 to TS27), corresponding to a total of 633,307 ESTs.

A weighted linear regression between developmental time and expression ratios was fit to the data, and a F-test was run to assess if the slope was significantly different from zero. Weights were the mean number of ESTs per gene at each stage, instead of the ratio of the number of genes expressed at each stage, gave similar results (not shown). We used data from 297 EST libraries, spanning 26 different developmental stages (from TS01 to TS27), corresponding to a total of 633,307 ESTs.

Zebrafish In Situ Data

In situ hybridization expression data from ZFIN [16] were retrieved using BGEE [51]. We considered only stages with more than 1000 genes expressed, starting when maternal genes are largely degraded (6 hours post-fertilization [48]). We retrieved all genes with at least one report of expression by in situ hybridization, at each time point of zebrafish development. From this set we extracted two groups (for example essential and non-annotated genes), and analyzed their ratio across development using the same methodology as with ESTs (see above).

Rate of Protein Evolution

The orthology relationships, and the values of $d_{N}$ (rate of nonsynonymous substitution per codon) and $d_{S}$ (rate of synonymous substitution per codon) were obtained from Ensembl version 48 [46]. We retrieved zebrafish genes with one-to-one orthologs in Tetraodon nigroviridis and Takifugu rubripes (divergence time is ~32 MYA between the two pufferfish species and ~130 MYA with Danio rerio [53]). We downloaded the pairwise $d_{N}$ and $d_{S}$ between Tetraodon and Takifugu, calculated with codeml from the PAML package in the Ensembl pipeline (model = 0, NSsites = 0) [54]. Ensembl considers that $d_{N}$ values are saturated when they reach a threshold which is $2*median(d_{S})$. See http://www.ensembl.org/info/about/docs/compara/homology_method.html for further details.

We selected a set of 4937 genes having $d_{S}$, $d_{N}$ and Affymetrix expression data. Among them 620 genes were strict singletons in fishes (loss of duplicates after 2R and after the fish-specific genome duplication). At each time point we performed the Spearman correlation between the $d_{S}/d_{N}$ ratio and expression, following Davis et al. [1]. A t-statistic was used to assess if the correlation coefficient was different from 0.

For the analysis in mouse we retrieved pairwise $d_{S}$ and $d_{N}$ between human and mouse, for genes with one-to-one orthologs (14,333 genes). We kept only the singletons for 2R genome duplication and separated the 25% with the highest $d_{S}/d_{N}$ and the 25% with the lowest $d_{S}/d_{N}$ (607 genes in each group). We then compared the expression across development of these two groups using EST data. Using the 10% highest and lowest $d_{S}/d_{N}$ gave similar results (not shown).

Genotypes and Phenotypes

Zebrafish mutants. Data on zebrafish mutants were retrieved from the Zebrafish Information Network (http://zfin.org/zf_info/downloads.html, April 2008) [16]. We selected mutant genotypes having a lethal or abnormal phenotype from the file “phenotype.txt”, paying attention that they were grown in normal conditions (ZDB-EXP-041102-1). These genotypes were mapped to ZFIN gene IDs using the file “genotype_features.txt” and then to Affymetrix probesets using Biomart [55]. This resulted in a dataset of 252 ZFIN IDs associated with a lethal phenotype (79 Affymetrix probesets), and 2870 ZFIN IDs associated with an abnormal phenotype (461 probesets). Annotated normal phenotype data are rare in ZFIN, due to a lack of report of such mutants in the literature, so we used non-annotated as a reference (7246 ZFIN gene IDs with expression data).

To be sure that the technique used in the phenotype screen did not bias our analysis, we separated the dataset of genotypes having an abnormal phenotype by technique (file “genotype_features.txt”: inversion, transgenic insertion, deficiency, point mutation, translocation, insertion, sequence variant or unspecified. Only transgenic insertions, point mutations and sequence variants provide enough data, with 343, 221 and 2424 ZFIN IDs respectively, corresponding to 309, 171 and 88 Affymetrix probesets respectively (Dataset S1 and Dataset S1).

Zebrafish morpholinos. The morpholinos knock-down phenotypes were downloaded from ZFIN (http://zfin.org/zf_info/downloads.html, April 2008) [16]. We selected morpholinos (file “pheno_environment.txt”) giving lethal or abnormal phenotypes (file “phenotype.txt”), paying attention that the genotypes were wild type (file “wildtypes.txt”). The probes were mapped to ZFIN gene IDs using the file “Morpholinos.txt” and then to Affymetrix probesets using Biomart [55]. Only “abnormal” phenotypes provided enough data, with 601 ZFIN IDs corresponding to 256 Affymetrix probesets (Dataset S1).

Mouse knock-outs. Data on mouse mutants were retrieved from the Mouse Genome Database (ftp://ftp.informatics.jax.org/
Identification of Duplicate Genes

Gene families were obtained from the HomolEns database version 3 (http://phil.univ-lyon1.fr/databases/homolens.html), which is based on Ensembl release 41 [46]. HomolEns is build on the same model as Hovergen [56], with genes organized in families, which include pre-calculated alignments and phylogenies. In HomolEns version 3, alignments are computed with MUSCLE [57] (with default parameters), and phylogenetic trees with PhyML [58]. Phylogenies are computed on conserved blocks of the alignments selected with GBLOCKS [59]. Using the TreePattern functionality of the FamFetch client for HomolEns, which allows scanning for gene tree topologies [60], we selected sets of genes with or without duplications on specific branches of the vertebrate phylogenetic tree.

Regarding the fish-specific whole genome duplication, we found 1772 Ensembl IDs for duplicates in zebrafish, 8821 for singletons in zebrafish, 755 mouse orthologs of these duplicates, and 6043 mouse orthologs of these singletons. For the 2R whole genome duplications, we found 986 duplicates and 1266 singletons in zebrafish, and 2448 duplicates and 2705 singletons in mouse (Datasets S1 and S2).

Gene Ontology Analysis

Over and under representation of GO terms [25] was tested by means of a Fisher exact test, using the Bioconductor package topGO version 1.8.1 [61]. The reference set was all Ensembl genes mapped to a probeset of the zebrafish Affymetrix chip. The “elim” algorithm of topGO was used, allowing to decorrelate the graph structure of the gene ontology, reducing non-independence problems. A False Discovery Rate correction was applied, and gene ontology categories with a FDR <15% were reported.

Tools

R was used for statistical analysis and plotting (http://www.R-project.org/) [62], in conjunction with Bioconductor packages (http://www.bioconductor.org/, version 2.2)[45]. To retrieve genomic information we used the BioMart tool [55] or connected to the Ensembl MySQL public database [46].

Supporting Information

Figure S1 Expansion in zebrafish development of genes according to retention after vertebrate 2R whole genome duplications. Median expression profiles of vertebrate specific 2R duplicates in zebrafish in red dashed line and triangles, and of singletons in black solid line and circles. Legend as in Figure 2. Found at: doi:10.1371/journal.pgen.1000311.s001 (0.53 MB TIF)

Figure S2 Variation across mouse development of the ratio of expressed vertebrate 2R singletons, relative to duplicates. Legend as in Figure 4. Found at: doi:10.1371/journal.pgen.1000311.s002 (0.31 MB TIF)

Figure S3 Variation across zebrafish development of the Spearman correlation between gene sequence evolution and expression. Only singletons genes (for 2R and fish-specific genome duplications) were considered. We used the ratio of the rate of non-synonymous substitutions on the rate of synonymous substitutions (dN/dS) as a measure of selective pressure. Correlations below the dashed line are significantly different from 0 (p-value <0.05). The x-axis is in logarithmic scale. A gray box on the x-axis indicates the phylotypic period.

Found at: doi:10.1371/journal.pgen.1000311.s003 (0.41 MB TIF)

Figure S4 Expression in zebrafish development of genes with abnormal mutant phenotypes. Median expression profiles of zebrafish genes inducing abnormal phenotypes when non functional, for 4 different techniques, compared to non-annotated genes in black solid line and circles. The techniques are: morpholinos in purple dashed-dotted line and squares; transgenic insertions in green dashed line and triangles; point mutations in blue dashed line and diamonds; sequence variants in red dotted line and crosses. Points significantly different from the reference curve (non annotated genes) are filled. See Figure S5 for confidence intervals of the difference with the reference curve. The x-axis is in logarithmic scale. A gray box on the x-axis indicates the phylotypic period.

Found at: doi:10.1371/journal.pgen.1000311.s004 (0.41 MB TIF)

Figure S5 Significance of the expression difference between zebrafish genes inducing abnormal phenotypes when non-functional and non-annotated genes for 4 different techniques. These randomization plots refer to Figure S4. Legend as in Figure 2B.

Found at: doi:10.1371/journal.pgen.1000311.s005 (1.05 MB TIF)

Table S1 Gene Ontology analysis. The two groups analyzed are the genes experiencing an increase of expression along development (late expression, cluster 4) and the genes experiencing a decrease of expression (early expression, cluster 3) (Figure 3). Molecular Function and Biological process ontologies were analyzed with the “elim” algorithm of the Bioconductor package topGO (see Methods).

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Dataset S1 Details and characteristics of zebrafish gene sets used in this study. FSGD: Fish Specific whole Genome Duplication.

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Dataset S2 Details and characteristics of mouse gene sets used in this study. FSGD: Fish Specific whole Genome Duplication.

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Text S1 Supplementary text.

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Author Contributions
Conceived and designed the experiments: JR MRR. Performed the experiments: JR. Analyzed the data: JR MRR. Wrote the paper: JR MRR.

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