**RESEARCH ARTICLE**

**Tissue-Specificity of Gene Expression Diverges Slowly between Orthologs, and Rapidly between Paralogs**

*Nadezda Kryuchkova-Mostacci¹,², Marc Robinson-Rechavi¹,²*  

¹ Department of Ecology and Evolution, University of Lausanne, Lausanne, Switzerland, ² Swiss Institute of Bioinformatics, Lausanne, Switzerland  

*marc.robinson-rechavi@unil.ch*

**Abstract**

The ortholog conjecture implies that functional similarity between orthologous genes is higher than between paralogs. It has been supported using levels of expression and Gene Ontology term analysis, although the evidence was rather weak and there were also conflicting reports. In this study on 12 species we provide strong evidence of high conservation in tissue-specificity between orthologs, in contrast to low conservation between within-species paralogs. This allows us to shed a new light on the evolution of gene expression patterns. While there have been several studies of the correlation of expression between species, little is known about the evolution of tissue-specificity itself. Ortholog tissue-specificity is strongly conserved between all tetrapod species, with the lowest Pearson correlation between mouse and frog at \( r = 0.66 \). Tissue-specificity correlation decreases strongly with divergence time. Paralogs in human show much lower conservation, even for recent Primate-specific paralogs. When both paralogs from ancient whole genome duplication tissue-specific paralogs are tissue-specific, it is often to different tissues, while other tissue-specific paralogs are mostly specific to the same tissue. The same patterns are observed using human or mouse as focal species, and are robust to choices of datasets and of thresholds. Our results support the following model of evolution: in the absence of duplication, tissue-specificity evolves slowly, and tissue-specific genes do not change their main tissue of expression; after small-scale duplication the less expressed paralog loses the ancestral specificity, leading to an immediate difference between paralogs; over time, both paralogs become more broadly expressed, but remain poorly correlated. Finally, there is a small number of paralog pairs which stay tissue-specific with the same main tissue of expression, for at least 300 million years.

**Author Summary**

From specific examples, it has been assumed by comparative biologists that the same gene in different species has the same function, whereas duplication of a gene inside one species to create several copies allows them to acquire different functions. Yet this model was little...
tested until recently, and then has proven harder than expected to confirm. One of the problems is defining "function" in a way which can be easily studied. We introduce a new way of considering function: how specific is the activity ("expression") of a gene? Genes which are specific to certain tissues have functions related to these tissues, whereas genes which are broadly active over many or all tissues have more general functions for the organism. We find that this "tissue-specificity" evolves very slowly in the absence of duplication, while immediately after duplication the new gene copy differs. This shows that indeed duplication leads to a strong increase in the evolution of new functions.

Introduction

The ortholog conjecture is widely used to transfer annotation among genes, for example in newly sequenced genomes. But has been difficult to establish whether and how much orthologs share more similar functions than paralogs [1,2]. The most widely accepted model is that orthologs diverge slower, and that the generation of paralogs through duplication leads to strong divergence and even change of function. It is also expected that in general homologs diverge functionally with time. The test of these hypotheses poses fundamental questions of molecular evolution, about the rate of functional evolution and the role of duplications, and is essential to the use of homologs in genome annotations.

Surprisingly, there are several studies which have reported no difference between orthologs and paralogs, or even the opposite, that paralogs would be more functionally similar than orthologs. Tests of the ortholog conjecture using sequence evolution found no difference after speciation or duplication in positive selection [3], nor in amino acid shifts [4]. The debate was truly launched by Nehrt et al. [5] who reported in a large scale study, based on expression levels similarity and Gene Ontology (GO) analysis in human and mouse, that paralogs are better predictors of function than orthologs. Of note, methodological aspects of the GO analysis of that study were criticized by several other authors [6,7]. Using a very similar GO analysis but correcting biases in the data, from 13 bacterial and eukaryotic species, Altenhoff et al. [8] found more functional similarity between orthologs than between paralogs based on GO annotation analysis, but the differences were very slight.

An early comparison of expression profiles of orthologs in human and mouse reported that they were very different, close to paralogs and even to random pairs [9]. Further studies, following Nehrt et al. [5], found little or no evidence for the ortholog conjecture in expression data. Rogozin et al. [10] reported that orthologs are more similar than between species paralogs but less similar than within-species paralogs based on correlations between RNA-seq expression profiles in human and mouse. Wu et al. [11] found only a small difference between orthologs and paralogs. Paralogs were significantly more functionally similar than orthologs, but by classifying in subtypes they reported that one-to-one orthologs are the most functionally similar. The analysis was done on the level of function by looking at expression network similarities in human, mouse, fly and worm.

On the other hand, the ortholog conjecture has been supported by several studies of gene expression. Contra Yanai et al. [9], several studies have reported good correlations between expression levels of orthologs, between human and mouse [12], or among amniotes [13]. Moreover, some studies have reported changes of expression following duplication, although without explicitly testing for the ortholog conjecture: duplicated genes are more likely to show changes in expression profiles than single-copy genes [14,15]. Chung et al. [16] reported through network analysis in human that duplicated genes diverge rapidly in their expression...
profile. Recently Assis and Bachtrog [17] reported that paralog function diverges rapidly in mammals. They analysed among other things difference in tissue-specificity between a pair of paralogs and their single copy ortholog in closely related species. They conclude that divergence of paralogs results in increased tissue-specificity, and that there are differences between tissues. Finally, several explicit tests of the ortholog conjecture have also found support using expression data. Huerta-Cepas et al. [18] reported that paralogs have higher levels of expression divergence than orthologs of the similar age, using microarray data with calls of expressed/not expressed in human and mouse. They also claimed that a significant part of this divergence was acquired shortly after the duplication event. Chen and Zhang [7] re-analysed the RNA-seq dataset of Bra-wand et al. [13] and reported that expression profiles of orthologs are significantly more similar than within-species paralogs.

Thus while the balance of evidence appears to weigh towards confirmation of the ortholog conjecture, functional data has failed so far to strongly support or invalidate it. Even results which support the ortholog conjecture often do so with quite slight differences between orthologs and paralogs [8,10]. Yet expression data especially should have the potential to solve this issue, since it provides functional evidence for many genes in the same way across species, without the ascertainment biases of GO annotations or other collections of small scale data. Part of the problem is that the relation between levels of expression and gene function is not direct, making it unclear what biological signal is being compared in correlations of these levels. Another problem is that the comparison of different transcriptome datasets between species suffers from biases introduced by ubiquitous genes [19] or batch effects [20].

In our analysis we have concentrated on the tissue-specificity of expression. Tissue-specificity indicates in how many tissues a gene is expressed, and whether it has large differences of expression level between them. It reflects the functionality of the gene: if the gene is expressed in many tissues then it is "house keeping" and has a function needed in many organs and cell types; tissue-specific genes have more specific roles, and tissue adjusted functions. Recent results indicate that tissue-specificity is conserved between human and mouse orthologs, and that it is functionally informative [21]. Moreover, tissue-specificity can be computed in a comparable manner in different animal datasets without notable biases, as long as at least 6 tissues are represented, including preferably testis, nervous system, and proportionally not too many parts of the same organ (e.g. not many parts of the brain).

Are there major differences between the evolution of tissue-specificity after duplication (paralogs) or without duplication (orthologs)? We analyse the conservation of one-to-one orthologs and within-species paralogs with evolutionary time, using RNA-seq datasets from 12 species.

**Results**

We compared orthologs between 12 species: human, chimpanzee, gorilla, macaque, mouse, rat, cow, opossum, platypus, chicken, frog, and fruit fly. Overall 7 different RNA-seq datasets were used, including 6 to 27 tissues (see Materials and Methods). Three comparisons were performed with the largest sets as focal data: 27 human tissues from Fagerberg et al., 16 human tissues from Bodymap, and 22 tissues from mouse ENCODE [22–24]. For all analyses we used tissue-specificity of expression as described in Materials and Methods.

The first notable result is that tissue-specificity is strongly correlated between one-to-one orthologs. The correlations between human and four other species are presented in Fig 1A for illustration. This confirms and extends our previous observation [21], which was based on one human and one mouse datasets. Correlation of tissue-specificity varies between 0.74 and 0.89 among tetrapods, and is still 0.43 between human and fly, 0.38 between mouse and fly. The
latter is despite the very large differences in anatomy and tissue sampling between the species compared, showing how conserved tissue-specificity can be in evolution.

The correlation between orthologs decreases with divergence time (Fig 2). The decline is linear. An exponential model is not significantly better: ANOVA was not significantly better for the model with \( \log_{10} \) of time than for untransformed time for any dataset \((p > 0.0137, q > 1\%)\). The trend is not caused by the outlier fly data point: removing it there is still a significant decrease of correlation for orthologs (see S1 Fig). Results are also robust to the use of Spearman instead of Pearson correlation between tissue-specificity values.

The correlation between within-species paralogs is significantly lower than between orthologs (ANOVA \( p < 0.0137, q < 1\% \) for all datasets) (Fig 2). Moreover, there is no significant decline in correlation with evolutionary time (neither linear nor exponential) for paralogs. This may indicate almost immediate divergence of paralogs upon duplication, although other scenarios are possible (see Discussion).

The results are consistent using human or mouse as focal species (Fig 2A and 2B). Results are also consistent using a different human RNA-seq dataset (Fig A in S1 Fig).

This main analysis is based on the correlation of tissue-specificity for orthologs called pairwise between species. The number of orthologs used in the analysis is thus variable (available in Table B in S1 Table). An additional analysis was also performed using the same orthologs for all tetrapods, 4785 genes (Fig B-D in S1 Fig). Correlations of these "conserved orthologs" are not significantly different from those observed over all orthologs.

The analysis was also performed on all the datasets with tissue-specificity calculated without testis (Fig E-G in S1 Fig). The correlation between orthologs becomes significantly lower (ANOVA \( p = 0.000178 \)), while between paralogs it does not change significantly (ANOVA \( p = 0.846 \)). Even though the correlation between orthologs becomes weaker there is still a
significant difference between orthologs and paralogs (ANOVA \( p = 1.299 \times 10^{-7} \)). The same analysis was also performed removing 4 other main tissues (brain, heart, kidney and liver) (Fig H-K in S1 Fig). For the brain the correlation between orthologs becomes significantly lower (ANOVA \( p = 0.000289 \)), but stays higher than for paralogs; for other tissues there is no significant difference. For paralogs the correlation never changes significantly.

We also performed the analysis removing genes on sex chromosomes (Fig L-N in S1 Fig). This analysis was done without frog, as sex chromosome information is not available. This does not change significantly the correlations between either orthologs (ANOVA \( p = 0.856 \)) or paralogs (ANOVA \( p = 0.755 \)).

In general paralogs have lower expression and are more tissue-specific than orthologs (Fig O in S1 Fig), which is consistent with the dosage-sharing model [25,26]. Young paralogs are very tissue-specific, and get more ubiquitous with divergence time (Fig 1B and Fig P in S1 Fig); this is true for all datasets, and for \( \tau \) calculated with or without testis. We also tested for asymmetry by comparing paralog pairs to the closed possible non duplicated outgroup; e.g., we compared each Eutheria specific paralog to the non-duplicated opossum outgroup (one-to-two ortholog; Fig 3). We observe that the higher expressed paralog has a stronger correlation with the outgroup, thus appears to keep more the ancestral tissue-specificity, while the lower expressed paralog has a lower correlation and appears to become more tissue-specific (Fig 3), which is consistent with a form of neo-functionalization.

When both orthologs of a pair are tissue-specific (\( \tau > 0.8 \)), they are most often expressed in the same tissue (Fig 4). The same is observed when both paralogs are tissue-specific and are younger than the divergence of tetrapods. But for Euteleostomi and Vertebrata paralogs, if both are tissue-specific then they are as likely to be expressed in the different as in same tissues; most of these are expected to be ohnologs, i.e. due to whole genome duplication. This analysis was performed on the Brawand et al. (2011) dataset, because it has the most organisms with the same 6 tissues. This result does not change after removing testis (Fig Q in S1 Fig), nor changing the \( \tau \) threshold from 0.8 to 0.3 (Fig R-S in S1 Fig). Also after removing all tissue-specific genes (\( \tau > 0.8 \)), the difference between orthologs and paralogs is smaller but stay significant (ANOVA \( p = 0.001 \)) (Fig T in S1 Fig).

**Discussion**

Our results show that most genes have their tissue-specificity conserved between species. This provides strong new evidence for the evolutionary conservation of expression patterns. Using tissue-specificity instead of expression values allows easy comparison between species, as bias of normalisation or use of different datasets has little effect on results [21]. All of our results were confirmed using three different focus datasets, from human or mouse, and thus appear to be quite robust.

The conservation of expression tissue-specificity of protein coding genes that we find is high even for quite distant one-to-one orthologs: the Pearson correlation between \( \tau \) in human or mouse and \( \tau \) in frog is \( R = 0.74 \) (respectively \( R = 0.66 \)) over 361 My of divergence. Even between fly and mammals it is more than 0.38. Moreover, this tissue-specificity can be easily compared over large datasets without picking a restricted set of homologous tissues (e.g. in
The correlation between orthologs is strongest for recent speciations, and decreases linearly with divergence time. This decrease shows that we are able to detect a strong evolutionary signal in tissue-specificity, which has not always been obvious in functional comparisons of orthologs (e.g. [5, 8]).

Correlation between within-species paralogs is much lower than between orthologs. Whereas the expression of young paralogs has been recently reported to be highly conserved [17], we find a large difference between even very young paralogs in tissue-specificity. In Assis and Bachtrog [17], the measure of tissue-specificity is not clearly defined, but it seems to be TSI [27], which performed poorly as an evolutionarily relevant measure in our recent benchmark [21]; they also treated female and male samples as different “tissues”, confounding two potentially different effects. The low correlation that we observed for young paralogs does not decrease significantly with divergence time. It is possible that on the one hand paralogs do

Fig 3. Distribution of tissue-specificity in paralogs compared to an outgroup ortholog. For each graph, paralogs of a given phylogenetic age are compared to the closest outgroup un-duplicated ortholog; thus these paralogs are “in-paralogs” relative to the speciation node, and are both “co-orthologs” to the outgroup. X-axis, τ of unduplicated ortholog. Y-axis, τ of paralogs. Blue points are values for the paralog with highest maximal expression of the pair of paralogs, orange points are values for the other.

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diverge in tissue-specificity with time, and that on the other hand this trend is compensated by biased loss of the most divergent paralogs. It is also possible that we lack statistical power to detect a slight decrease in correlation of paralogs, due to low numbers of paralogs for many branches of the phylogeny. The most likely interpretation is that for small-scale paralogs (defined as not from whole genome duplication [28]) there is an asymmetry, with a daughter gene which lacks regulatory elements of the parent gene upon birth; further independent changes in tissue-specificity in each paralog would preserve the original lack of correlation. In any case, we do not find support for a progressive divergence of tissue-specificity for paralogs.

The overall conservation of tissue-specificity could be due to a subset of genes, and most notably sex-related genes. Indeed, the largest set of tissue-specific genes are testis-specific [21]. To verify the influence of sex-related genes, we performed all analyses without testis expression data, or without genes mapped to sex chromosomes. After removing testis expression from all datasets the correlation between paralogs does not change significantly, while between orthologs is gets significantly weaker. The lower correlation of orthologs suggests that testis specific genes are conserved between species, and as they constitute a high proportion of tissue-specific genes, they contribute strongly to the correlation. Removing sex chromosome located genes

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Fig 4. Difference of tissue-specificity between orthologs and paralogs. Each bar represents the number of gene pairs of a given type for a given phylogenetic age, for which both genes of the pair are tissue-specific ($\tau > 0.8$). In dark color, the number of gene pairs specific to the same tissue; in light color, the number of gene pairs specific to different tissues. Orthologs are in red, in the left panel, paralogs are in blue, on the right panel; notice that the scales are different for orthologs and for paralogs. Orthologs are one-to-one orthologs to human and paralogs are within-species paralogs in human. The overall proportions of pairs in the same or different tissues are indicated for orthologs and paralogs; in addition, for paralogs the proportion for pairs younger than the divergence of tetrapods (whole genome duplication) is also indicated.
does not change results significantly. After removing testis expression the differences of con-
servation of tissue-specificity between orthologs and paralogs stay significant. Overall, it
appears that tissue-specificity calculated with testis represents a true biological signal, and
given its large effect it is important to include this tissue in analyses.

In general paralogs are more tissue-specific and have lower expression levels. This could be
explained if ubiquitous genes are less prone to duplication or duplicate retention. Yet we do
not observe any bias in the orthologs of duplicates towards more tissue-specific genes (Fig 3;
see also S1 Fig). With time both paralogs get more broadly expressed (Fig 1 and Fig P in S1
Fig). In the rare case where both paralogs are tissue-specific, small-scale young paralogs are
expressed in the same tissue, while genome-wide old paralogs (ohnologs) are expressed in dif-
ferent tissues (Fig 4). With the data available, we cannot distinguish the effects of paralog age
and of duplication mechanism, since many old paralogs are due to whole genome duplication
in vertebrates, whereas that is not the case for the young paralogs. In many cases the higher
expressed paralog has a similar tissue-specificity to the ancestral state, while the lower
expressed paralog is more tissue-specific (Fig 3).

We have studied gene specificity without taking into account alternative splicing, or the possi-
ability that different transcripts are expressed in different tissues, because it is still difficult to
call transcript level expression reliably [29]. This would probably not change our main obser-
vations, that tissue-specificity is conserved among orthologs, diverges with evolutionary time,
and follows the ortholog conjecture. Of note, recent results have not supported an important
role of alternative splicing for differences in transcription between tissues [30,31].

The overall picture that we obtain for the evolution of tissue-specificity is the following. In
the absence of duplication, tissue-specificity evolves slowly, thus is mostly conserved, and tis-
sue-specific genes do not change their main tissue of expression (Figs 2 and 4). After small-
scale duplication (i.e., not whole genome) paralogs diverge rapidly in tissue-specificity, or
already differ at birth. This difference is mostly due to the less expressed paralog losing the
ancestral specificity, while the most expressed paralog keeps at first closer to the ancestral state,
as estimated from a non-duplicated outgroup ortholog (Fig 3). But over time, even the most
expressed paralog diverges much more strongly than a non-duplicated ortholog. While paralog
divergence is rapid, in the small number of genes which stay tissue-specific for both paralogs
the main tissue of expression is mostly conserved, for several hundred million years (i.e. origin
of tetrapods, Fig 4). With increasing age of the paralogs, they both tend to become more
broadly expressed (Fig 1 and Fig P in S1 Fig) while keeping a low correlation. For whole
gene duplicates we have less information, because of the age of the event in vertebrates and
the lack of good outgroup data. The main difference is that when two genome duplication
paralogs are both tissue-specific, they are often expressed in different tissues (Fig 4).

We have used tissue-specificity to estimate the conservation of function, rather than Gene
Ontology annotations or expression levels. We believe that this metric is less prone to system-
atic errors, whether annotation biases for the Gene Ontology, or proper normalisation
between datasets and choice of few tissues for expression levels. Our results confirm the Ortho-
log Conjecture on data which is genome-wide and functionally relevant: orthologs are more
similar than within-species paralogs. Moreover, orthologs diverge monotonically with time, as
expected. On the contrary, even young paralogs show large differences.

**Material and Methods**

RNA-seq data from 12 species (human, gorilla, chimpanzee, macaque, mouse, platypus, opos-
sum, chicken, gorilla, cow, frog, rat and fruit fly) were used for the analysis. We recovered all
animal RNA-seq data sets which cover at least 6 adult tissues, and were either pre-processed in
Bgee [32], or provided pre-processed data from the publication, as of June 2015. For human, mouse and chicken we used several datasets. All the datasets with the corresponding number of tissues are summarized in Table 1. The numbers of genes used for the analysis are in Table A and B in S1 Table.

The orthology and paralogy calls and their phylogenetic dating for paralogs were taken from Ensembl Compara (Version 75) [33]. Phylogenetic dating was converted to absolute dates using the TimeTree data base [34].

For the human dataset from Fagerberg et al. [22] and the fly dataset [36], FPKM values were downloaded from the respective papers Supplementary Materials; the mouse ENCODE project dataset was processed by an in house script (TopHat and Cufflinks [40]); all other data were processed by the Bgee pipeline [32]. For all analyses gene models from Ensembl version 75 were used [41]. Only protein-coding genes were used for analysis. For the analysis of paralogs the youngest couple was taken (Fig U in S1 Fig), and sorted according to the maximal expression, i.e. the reference paralog (called ”gene” in our R scripts) is always the one with the highest maximal expression. This choice gives the highest correlation compared to a random sorting (Fig V in S1 Fig).

Analyses were performed in R version 3.2.1 [42] using Lattice [43], plyr [44], gplots [45] and qvalue [46,47] libraries.

As a measure for tissue-specificity we used $\tau$ (Tau) [48]:

$$\tau = \frac{\sum_{i=1}^{n} (1 - \bar{x}_i)}{n - 1}; \quad \bar{x}_i = \frac{x_i}{\max_{1 \leq i \leq n}(x_i)}$$

$\tau$ is calculated on the log RNA-seq expression data. The values of $\tau$ vary from 0 to 1, where 0 means ubiquitous expressed genes and 1 specific genes. We have recently shown that $\tau$ is the best choice for calculating tissue specificity among existing methods [21]. For comparing tissue-specific genes, they were called with $\tau \geq 0.8$, and assigned to the tissue with the highest expression.

A special case is testis-specificity, as many more genes are expressed in testis than other tissues. For control analysis, all genes with maximal expression in testis were called ”testis specific”, independently of $\tau$ value.

### Table 1. Datasets used in the paper.

| Organisms/ datasets     | Fagerberg | Brawand | Bodymap | ENCODE | Necsulea | Merkin | Keane |
|-------------------------|-----------|---------|---------|---------|----------|--------|-------|
| Dataset ID              | E-MTAB-1733 | GSE30352 | GSE30611 | GSE36025 (mouse) | GSE43520 | GSE41637 | GSE30617 |
| RPKM/FPKM source        | Supp. mat. | Bgee    | Bgee    | Supp. mat.; [35] | Bgee    | Bgee    | Bgee    |
| Human Homo sapiens      | 27        | 8       | 16      |         |          |        |       |
| Gorilla Gorilla gorilla | 6         |         |         |         |          |        |       |
| Chimpanzee Pan troglodytes | 6       |         |         |         |          |        |       |
| Macaque Macaca mulatta  | 6         |         |         |         |          |        |       |
| Mouse Mus musculus      | 6         | 22      | 9       | 6       |          |        |       |
| Rat Rattus norvegicus   |           |         |         |         |          |        |       |
| Bos taurus              |           |         |         |         | 9        |        |       |
| Opossum Monodelphis domestica | 6   |         |         |         |          |        |       |
| Platypus Ornithorhynchus anatinus | 6 |         |         |         |          |        |       |
| Chicken Gallus gallus   | 6         |         |         |         |          |        |       |
| Frog Xenopus tropicalis |           |         |         |         |          |        |       |
| Fly Drosophila melanogaster | 6   |         |         |         |          |        |       |
| Citations               | [22]      | [13]    | [23]    | [24,36] | [37]     | [38]   | [39]  |

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Over all ANOVA tests performed (112 tests), we used a q-value threshold of 1% of false positives, corresponding to a p-value threshold of 0.066.

Supplementary Materials

Additional Supplementary files are available on Figshare: https://dx.doi.org/10.6084/m9.figshare.3493010.v2

Supporting Information

S1 Fig. Additional figures, including alternative versions of Figs 2 and 3 with different parameters or datasets.

S1 Table. Numbers of genes used in the analyses, per species and dataset.

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Author Contributions

Conceptualization: NKM MRR.
Data curation: NKM.
Funding acquisition: MRR.
Investigation: NKM MRR.
Methodology: NKM MRR.
Software: NKM.
Supervision: MRR.
Validation: NKM MRR.
Visualization: NKM.
Writing – original draft: NKM MRR.
Writing – review & editing: NKM MRR.

References

1. Studer RA, Robinson-Rechavi M. How confident can we be that orthologs are similar, but paralogs differ? Trends Genet. 2009; 25:210–6. doi: 10.1016/j.tig.2009.03.004 PMID: 19368988
2. Gabaldón T, Koonin E V. Functional and evolutionary implications of gene orthology. Nat. Rev. Genet. Nature Publishing Group; 2013; 14:360–6. doi: 10.1038/nrg3456 PMID: 23552219
3. Studer R, Penel S, Duret L, Robinson-Rechavi M. Pervasive positive selection on duplicated and non-duplicated vertebrate protein coding genes. Genome Res. 2008; 18:1393–402. doi: 10.1101/gr.076992.108 PMID: 18562677
4. Studer RA, Robinson-Rechavi M. Large-scale analysis of orthologs and paralogs under covarion-like and constant-but-different models of amino acid evolution. Mol. Biol. Evol. 2010; 27:2618–27. doi: 10.1093/molbev/msq148 PMID: 20551039
5. Nehrt NL, Clark WT, Radivojac P, Hahn MW. Testing the ortholog conjecture with comparative functional genomic data from mammals. PLoS Comput. Biol. 2011; 7:e1002073. doi: 10.1371/journal.pcbi.1002073 PMID: 21695233

6. Thomas PD, Wood V, Mungall CJ, Lewis SE, Blake JA. On the use of gene ontology annotations to assess functional similarity among orthologs and paralogs: A short report. PLoS Comput. Biol. 2012; 8:1–7.

7. Chen X, Zhang J. The ortholog conjecture is untestable by the current gene ontology but is supported by RNA sequencing data. PLoS Comput. Biol. 2012; 8:e1002784. doi: 10.1371/journal.pcbi.1002784 PMID: 23209392

8. Altenhoff AM, Studer RA, Robinson-Rechavi M, Dessimoz C. Resolving the ortholog conjecture: orthologs tend to be weakly, but significantly, more similar in function than paralogs. PLoS Comput. Biol. 2012; 8:e1002514. doi: 10.1371/journal.pcbi.1002514 PMID: 22615551

9. Yanai I, Graur D, Ophir R. Incongruent expression profiles between human and mouse orthologous genes suggest widespread neutral evolution of transcription control. OMICS. 2004; 8:15–24. doi: 10.1089/153623104773547462 PMID: 15107234

10. Rogozin IB, Managadze D, Shabalina SA, Koonin EV. Gene family level comparative analysis of gene expression in mammals validates the ortholog conjecture. Genome Biol. Evol. 2014; 6:754–62. doi: 10.1093/gbe/evu051 PMID: 24610837

11. Wu Y-C, Bansal MS, Rasmussen MD, Herrero J, Kellis M. Phylogenetic identification and functional characterization of orthologs and paralogs across human, mouse, fly, and worm. bioRxiv. 2014;

12. Liao B-Y, Zhang J. Evolutionary conservation of expression profiles between human and mouse orthologous genes. Mol. Biol. Evol. 2006; 23:530–40. doi: 10.1093/molbev/msj054 PMID: 16280543

13. Brawand D, Soumillon M, Necsulea A, Julien P, Csárdi G, Harrigan P, et al. The evolution of gene expression levels in mammalian organs. Nature. 2011; 478:343–8. doi: 10.1038/nature10532 PMID: 22012392

14. Gu Z, Rifkin SA, White KP, Li W-H. Duplicated genes increase gene expression diversity within and between species. Nat. Genet. 2004; 36:577–9. doi: 10.1038/ng1355 PMID: 15122255

15. Hummiecki L, Wolfe KH. Divergence of spatial gene expression profiles following species-specific gene duplications in human and mouse. Genome Res. 2004; 14:1870–9. doi: 10.1101/gr.2705204 PMID: 15466287

16. Chung W-Y, Albert R, Albert I, Nekrutenko A, Makova KD. Rapid and asymmetric divergence of duplicate genes in the human gene coexpression network. BMC Bioinformatics. 2006; 7:1–14.

17. Assis R, Bachtrog D. Rapid divergence and diversification of mammalian duplicate gene functions. BMC Evol. Biol. BMC Evolutionary Biology; 2015; 15:1–7.

18. Huerta-Cepas J, Dopazo J, Huynen MA, Gabaldón T. Evidence for short-time divergence and long-time conservation of tissue-specific expression after gene duplication. Brief. Bioinform. 2011; 12:442–8. doi: 10.1093/bib/bbr022 PMID: 21515902

19. Piasecka B, Robinson-Rechavi M, Bergmann S. Correcting for the bias due to expression specificity improves the estimation of constrained evolution of expression between mouse and human. Bioinformatics. 2012; 28:1865–72. doi: 10.1093/bioinformatics/bts266 PMID: 22576178

20. Gilad Y, Mizrahi-Man O. A reanalysis of mouse ENCODE comparative gene expression data. F1000Research. 2015; 4:121. doi: 10.12688/f1000research.6536.1 PMID: 26236466

21. Kryuchkova-Mostacci N, Robinson-Rechavi M. A benchmark of gene expression tissue-specificity metrics. Brief. Bioinform. 2016; 1–10.

22. Fagerberg L, Hallstrom BM, Oksvold P, Kampf C, Djureinic D, Odeberg J, et al. Analysis of the human tissue-specific expression by genome-wide integration of transcriptomics and antibody-based proteomics. Mol. Cell. Proteomics. 2014; 13:397–406. doi: 10.1074/mcp.M113.039605 PMID: 24309898

23. Farrell CM, O’Leary NA, Harte RA, Loveland JE, Wilming LG, Wallin C, et al. Current status and new features of the Consensus Coding Sequence database. Nucleic Acids Res. 2014; 42:D865–72. doi: 10.1093/nar/gkt1059 PMID: 24217909

24. The ENCODE Project Consortium. A user’s guide to the encyclopedia of DNA elements (ENCODE). PLoS Biol. 2011; 9:e1001046. doi: 10.1371/journal.pbio.1001046 PMID: 21526222

25. Lan X, Pritchard JK. Coregulation of tandem duplicate genes slows evolution of subfunctionalization in mammals. Science. 2016; 352:1009–13. doi: 10.1126/science.aad8411 PMID: 27199432

26. Gout J-F, Lynch M. Maintenance and loss of duplicated genes by dosage subfunctionalization. Mol. Biol. Evol. 2015; 32:2141–8. doi: 10.1093/molbev/msv095 PMID: 25908670
27. Julien P, Brawand D, Soumillon M, Necseula A, Liechti A, Schütz F, et al. Mechanisms and evolutionary patterns of mammalian and avian dosage compensation. PLoS Biol. 2012; 10:e1001328. doi: 10.1371/journal.pbio.1001328 PMID: 22615540

28. Davis JC, Petrov D a. Do disparate mechanisms of duplication add similar genes to the genome? Trends Genet. 2005; 21:548–51. doi: 10.1016/j.tig.2005.07.008 PMID: 16098632

29. Pelechano V, Wei W, Jakob P, Steinmetz LM. Genome-wide identification of transcript start and end sites by transcript isoform sequencing. Nat. Protoc. 2014; 9:1740–59. doi: 10.1038/nprot.2014.121 PMID: 24967623

30. Ezkurdia I, Rodríguez JM, Carrillo-de Santa Pau E, Vázquez J, Valencia A, Tress ML. Most highly expressed protein-coding genes have a single dominant isoform. J. Proteome Res. 2015; 14:1880–7. doi: 10.1021/pr501286b PMID: 25732134

31. Tress ML, Abascal F, Valencia A. Alternative splicing may not be the key to proteome complexity. Trends Biochem. Sci. Elsevier Ltd; 2016; 0:1–13.

32. Bastian F, Parmentier G, Roux J, Moretti S, Lauder V, Robinson-Rechavi M. Bgee: integrating and comparing heterogeneous transcriptome data among species. Data Integr. Life Sci. Springer Berlin Heidelberg; 2008. p. 124–31.

33. Vilella AJ, Severin J, Ureta-Vidal A, Heng L, Durbin R, Birney E. EnsemblCompara GeneTrees: Complete, duplication-aware phylogenetic trees in vertebrates. Genome Res. 2009; 19:327–35. doi: 10.1101/gr.073585.107 PMID: 19029536

34. Hedges SB, Dudley J, Kumar S. TimeTree: A public knowledge-base of divergence times among organisms. Bioinformatics. 2006; 22:2971–2. doi: 10.1093/bioinformatics/bti505 PMID: 17021158

35. Kryuchkova-Mostacci N, Robinson-Rechavi M. Tissue-specific evolution of protein coding genes in human and mouse. PLoS One. 2015; 10:e0131673. doi: 10.1371/journal.pone.0131673 PMID: 26121354

36. Li JJ, Huang H, Bickel PJ, Brenner SE. Comparison of D. melanogaster and C. elegans developmental stages, tissues, and cells by modENCODE RNA-seq data. Genome Res. 2014; 24:1086–101. doi: 10.1101/gr.170100.113 PMID: 24985912

37. Necseula A, Kaessmann H. Evolutionary dynamics of coding and non-coding transcriptomes. Nat. Rev. Genet. Nature Publishing Group; 2014; 15:734–48. doi: 10.1038/nrg3802 PMID: 25297727

38. Merkin J, Russell C, Chen P, Burge CB. Evolutionary dynamics of gene and isoform regulation in mammalian tissues. Science. 2012; 338:1593–9. doi: 10.1126/science.1228186 PMID: 23258891

39. Keane TM, Goodstadt L, Danecek P, White M a, Wong K, Yalcin B, et al. Mouse genomic variation and its effect on phenotypes and gene regulation. Nature. 2011; 477:289–94. doi: 10.1038/nature10413 PMID: 21921910

40. Trapnell C, Roberts A, Goff L, Pertea G, Kim D, Kelley DR, et al. Differential gene and transcript expression analysis of RNA-seq experiments with TopHat and Cufflinks. Nat. Protoc. Nature Publishing Group; 2012; 7:562–78. doi: 10.1038/nprot.2012.016 PMID: 22383036

41. Flicek P, Ahmed I, Amode MR, Barrett D, Beal K, Brent S, et al. Ensembl 2013. Nucleic Acids Res. 2013; 41:D48–55. doi: 10.1093/nar/gks1236 PMID: 23203867

42. R Core Team. R: A language and environment for statistical computing [Internet]. Vienna, Austria; 2015. p. R Foundation for Statistical Computing, Vienna.

43. Sarcar D. Lattice: Multivariate data visualization with R [Internet]. New York: Springer; 2008.

44. Wickham H. The Split-Apply-Combine strategy for data analysis. J. Stat. Softw. 2011; 40:1–29.

45. Warnes G, Bolker B, Bonebakker L, Gentleman R, Huber W, Liaw A, et al. Gplots: Various R programming tools for plotting data [Internet]. 2016.

46. Storey J, Tibshirani R. Statistical significance for genomewide studies. Proc Natl Acad Sci U S A. 2003;2003.

47. Storey JD. Qvalue: Q-value estimation for false discovery rate control [Internet]. 2015.

48. Yanai I, Benjamin H, Shmoish M, Chalifa-Caspi V, Shklar M, Ophir R, et al. Genome-wide midrange transcription profiles reveal expression level relationships in human tissue specification. Bioinformatics. 2005; 21:650–9. doi: 10.1093/bioinformatics/bti042 PMID: 15388519