Selection in the evolution of gene duplications
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Abstract

Background: Gene duplications have a major role in the evolution of new biological functions. Theoretical studies often assume that a duplication per se is selectively neutral and that, following a duplication, one of the gene copies is freed from purifying (stabilizing) selection, which creates the potential for evolution of a new function.

Results: In search of systematic evidence of accelerated evolution after duplication, we used data from 26 bacterial, six archaeal, and seven eukaryotic genomes to compare the mode and strength of selection acting on recently duplicated genes (paralogs) and on similarly diverged, unduplicated orthologous genes in different species. We find that the ratio of nonsynonymous to synonymous substitutions (\(K_s/K_a\)) in most paralogous pairs is <<1 and that paralogs typically evolve at similar rates, without significant asymmetry, indicating that both paralogs produced by a duplication are subject to purifying selection. This selection is, however, substantially weaker than the purifying selection affecting unduplicated orthologs that have diverged to the same extent as the analyzed paralogs. Most of the recently duplicated genes appear to be involved in various forms of environmental response; in particular, many of them encode membrane and secreted proteins.

Conclusions: The results of this analysis indicate that recently duplicated paralogs evolve faster than orthologs with the same level of divergence and similar functions, but apparently do not experience a phase of neutral evolution. We hypothesize that gene duplications that persist in an evolving lineage are beneficial from the time of their origin, due primarily to a protein dosage effect in response to variable environmental conditions; duplications are likely to give rise to new functions at a later phase of their evolution once a higher level of divergence is reached.

Background

Gene duplications are traditionally considered to be a major evolutionary source of new protein functions. The conventional view, pioneered by Susumu Ohno, holds that a gene duplication produces two functionally redundant, paralogous genes and thereby frees one of them from selective constraints. This unconstrained paralog is then free to accumulate neutral mutations that would have been deleterious in a unique gene [1]. Although the most likely outcome of such neutral evolution is for one of the paralogs to fix a null mutation and become a pseudogene, there is also the possibility of fixation of mutations that lead to a new function [2-6].

One of the predictions of the conventional model of evolution of duplicated genes is the rapid loss of paralogs due to null mutations [2,3,5,7]. However, this prediction was not supported by studies on isozyme spectra of polyploids in a number of organisms (reviewed in [8]). Furthermore, a study of 17 pairs of duplicated genes in the tetraploid frog *Xenopus laevis* has shown that both copies were subject to
purifying selection [9], contrary to the notion of neutrality of one of the copies [1]. The failure of empirical research to support Ohno’s model has led to the proposal of two alternative hypotheses.

The ‘subfunctionalization’ hypothesis is based on the same assumption as Ohno’s model, namely that duplicated genes are redundant in function and, accordingly, a duplication event is selectively neutral [6,10]. However, it was argued that, as natural selection does not ‘know’ which duplicated gene should be under selection and which is free of selective constraint, both paralogs experience a period of relaxed selection and accelerated evolution. During this period, both genes might accumulate mutations that impair different functions of the ancestral gene, so that, after a certain point, none of the paralogs is capable of substituting for the ancestor [6,10].

The second hypothesis holds that a gene duplication that leads to a new function is preceded by a period of ‘gene sharing’, such that the original, unduplicated gene encodes two distinct functions. When two paralogs of such a bifunctional gene are produced by duplication, each may be driven by positive selection to specialize in one of these functions, which it performs more efficiently than the ancestor gene, leading to the creation of two indispensable genes with distinct functions [11].

The strong interest in the evolution of gene duplications stems from the notion that duplication leads to new functions [1,12,13]. With the increasing availability of genomic data, it became clear that numerous gene families have diverged in function through series of duplications [14,15], including many lineage-specific expansions identified in each of the genomes sequenced [16-19]. Such creation of novel gene functions obviously provides a long-term, but not a short-term [20] advantage for gene duplication.

Duplicated genes have also been observed to affect fitness immediately after duplication (see Discussion), providing a short-term advantage for duplication. Although these observations remained largely unnoticed by evolutionary biologists investigating the evolution of gene duplications, such short-term advantage is crucial to the long-term fate of a duplication as group selection favoring a duplication because of its long-term advantage cannot overcome the individual selection that depends exclusively on short-term effects [20].

A recent genome-wide analysis of duplicated genes of eukaryotes has suggested that they evolve under purifying selection, with an apparent early phase of relaxed constraint or even near-neutrality [21]. Here we show that the same pattern of purifying selection is observed among bacterial and archaeal paralogs and demonstrate that amino-acid substitution rate is greater in paralogs than in orthologs with the same level of divergence at synonymous sites. We also present evidence in support of genome-wide short-term advantage of duplicated genes, and discuss its relevance to the evolution of new gene functions.

**Results**

We identified recent gene duplications in 26 bacterial, six archaeal and three eukaryotic complete genomes, two completely sequenced chromosomes from *Arabidopsis*, and among the available sequences from three mammalian species - human, mouse and rat (Table 1). Because the duplications were identified through the similarity of full-length protein sequences (see Materials and methods), this procedure primarily, if not exclusively, detected functional genes rather than pseudogenes. For comparison, orthologs with no apparent recent duplications were identified in pairs of closely related organisms, namely the bacteria *Chlamydia trachomatis* and *C. muridarum* (603 orthologous pairs) and two strains of *Helicobacter pylori* (1,202 orthologous pairs), and the three mammals (3,428 orthologous pairs).

Selection was measured in terms of the ratio of the rate of nonsynonymous substitutions (Kn), which are usually subject to selective pressure, to the rate of synonymous substitutions (Ks), which are assumed to be (nearly) neutral. A Kn/Ks ratio of 1 is assumed to indicate neutrality, Kn/Ks > 1 is a signature of positive selection at the amino-acid level, and Kn/Ks < 1 is indicative of purifying selection, which is the most common mode of selection [14,15]. Such analysis is applicable to homologs with a relatively low level of divergence, in which synonymous substitutions have not yet reached saturation.

In all genomes analyzed, the nonsynonymous substitution rate was significantly lower than the synonymous substitution rate, such that Kn/Ks < 1 (Figure 1a,b and Table 1), in agreement with the recent genome-wide study of paralogs.

### Table 1

|                      | Archaea | Bacteria | S. cerevisiae | A. thaliana | C. elegans | D. melanogaster | Mammalia |
|----------------------|---------|----------|---------------|-------------|------------|-----------------|----------|
| Total number of      | 43      | 330      | 166           | 325         | 318        | 103             | 2,948    |
| paralogous clusters  |         |          |               |             |            |                 |          |
| Average ± standard   | 0.266 ± | 0.346 ±  | 0.104 ±       | 0.293 ±     | 0.244 ±    | 0.233 ±         | 0.451 ±  |
| error (standard      | 0.062   | 0.033    | 0.020         | 0.013       | 0.015      | 0.041           | 0.016    |
| deviation)           | (0.231) | (0.301)  | (0.171)       | (0.149)     | (0.162)    | (0.180)         | (0.257)  |

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Pairwise comparisons of the $K_n/K_s$ values for different clades revealed that the strength of selection was generally similar in bacterial, archaeal and eukaryotic sets of paralogous genes, although apparently stronger selection was observed among the yeast paralogs and weaker selection was seen in mammalian paralogs (Tables 1,2).

The observation that $K_n \ll K_s$ for most of the recently diverged paralogs is, in itself, not sufficient to conclude that both paralogs in each pair are subject to purifying selection. The same result would have been obtained if, in accordance with Ohno’s original model, one of the paralogs was free from selective constraints, whereas the other one was maintained by purifying selection. Therefore, we carried out a relative rate test on amino-acid substitutions [15] on those pairs of paralogs, for which an ortholog was available that was more distant from either of the paralogs than the paralogs were from each other (orthologs that were too distant to be used for the relative rate test were discarded; see Materials and methods). A comparison of the evolutionary distances between each of the paralogs and the ortholog used as an outgroup showed that, of a total of 101 analyzed pairs, only five evolve at significantly ($p < 0.05$) different rate. The proportions of paralogous pairs that evolved at significantly different rates in each of the analyzed clades are shown in Table 3. Thus, in the majority of cases at least, both paralogs evolved under similar levels of purifying selection, in agreement with the results of an early analysis of 17 pairs of paralogs from *X. laevis* and a more recent analysis of 19 pairs of paralogs in human and teleost lineages [9,22].

We next sought to compare the strength of purifying selection acting on paralogs to that acting on orthologs with the same level of sequence divergence, which were identified as bidirectional best hits [23] between two pairs of bacterial genomes (*C. trachomatis-C. muridarum* and two strains of *...* [21].

**Figure 1**

Synonymous and nonsynonymous substitution rates for (a,b) paralogs and (c,d) orthologs from (a,c) bacteria and (b,d) mammals. All points with $K_s > 0.5$ (approaching saturation) were assigned a $K_s$ value of 0.5. In (b), the blue circles represent human duplicated genes and the red circles show duplicated genes from mouse and rat. In (d), the blue circles are human-rodent orthologous comparisons, and the red circles are the mouse-rat comparisons.
H. pylori) and three pairs of mammalian species (human-mouse, human-rat and mouse-rat). The results of this comparison indicated that paralogs were subject to significantly weaker purifying selection than similarly diverged orthologs (compare Figure 1a and c; b and d). Specifically, the Kn/Ks ratio for orthologs was approximately twice the ratio for paralogs in bacteria, and approximately three times the ratio for paralogs in mammals (Table 4).

Because we restricted our gene comparisons to cases in which the synonymous divergence rate (Ks) did not exceed 0.5 substitutions per site, a substantial part of the human-rodent orthologous comparisons was excluded from the analysis (Figure 1d). The use of the orthologous comparisons with a lower Ks, and thus with potentially stronger selection in the synonymous sites, could, in principle, bias our comparison of the Kn/Ks ratio. To rule out this possibility, we calculated the Kn/Ks ratio for the center of the rodent orthologous comparisons (0.15 < Ks < 0.25) and compared it to the Kn/Ks ratio of paralogs within the same range of Ks. The difference in the Kn/Ks ratio in this range of Ks (0.448 in paralogs versus 0.175 in orthologs) was not substantially different from the difference between the Kn/Ks ratios for 0.05 < Ks < 0.5 (Table 4). Extrapolating this observation to the human-rodent comparisons, it appears most likely that the exclusion of the human-rodent orthologous pairs with Ks > 0.5 did not substantially bias the results.

Weak selection acting on synonymous sites or differences in mutation bias among related species might potentially result in homogeneity of codon usage in genes within one genome, which would translate into a much greater heterogeneity of codon vocabulary in orthologous genes compared to paralogous genes. Under this scenario, synonymous divergence could be substantially underestimated for paralogs, but not for orthologous genes in our dataset, falsely suggesting an acceleration of evolution of amino-acid substitutions (larger Kn/Ks values) in paralogous genes. If there was such homogeneity of codon usage in paralogous genes, paralogs should show a greater correlation between codon bias and Ks values than orthologs. This prediction is not supported by linear least-square regressions, which fail to detect any substantial differences in the correlation between Ks and codon bias in paralogous versus orthologous genes (data not shown), suggesting that the observation of a higher Kn/Ks ratio in paralogous genes cannot be explained by an underestimate of synonymous divergence in paralogs.

### Table 2

|                                   | Bacteria | S. cerevisiae | A. thaliana | C. elegans | D. melanogaster | Mammalia |
|-----------------------------------|----------|---------------|-------------|------------|-----------------|----------|
| Archaea                           | 0.359    | 0.0706        | 0.746       | 0.895      | 0.708            | 0.0456   |
| Bacteria                          | 3.65e-10 | 0.128         | 0.00538     | 0.0370     | 0.0513           |          |
| S. cerevisiae                     | 1.47e-12 | 1.07e-7       | 0.00915     | 5.74e-28   |                 |          |
| A. thaliana                       |          | 0.0141        | 0.181       |            | 7.57e-14         |          |
| C. elegans                        |          | 0.789         |             |            | 4.81e-19         |          |
| D. melanogaster                   |          |               |             |            | 5.71e-5          |          |

Significant p-values (< 0.05) are in bold.

### Table 3

**Results of the relative rate test for recently diverged paralogs**

|                  | Archaea | Bacteria | S. cerevisiae | C. elegans | D. melanogaster | Mammalia |
|------------------|---------|----------|---------------|------------|-----------------|----------|
| N_{diff}/N_{ident} | 2/7     | 1/10     | 0/15          | 0/2        | 2/11            | 2/49     |

N_{diff}, number of paralogous genes pairs that evolved at significantly (p < 0.05) different rates; N_{ident}, number of paralogous pairs that evolved at (approximately) the same rate.

### Table 4

**A comparison of the Kn/Ks ratios for paralogs and orthologs**

|                                | Bacteria | Mammals |
|--------------------------------|----------|---------|
| All proteins                   |          |         |
| Paralogs                       | 0.346 ± 0.033 (0.301) | 0.451 ± 0.016 (0.257) |
| Orthologs                      | 0.164 ± 0.004 (0.128) | 0.131 ± 0.003 (0.131) |
| Proteins with predicted signal peptides (probably secreted) |          |         |
| Paralogs                       | 0.352 ± 0.079 (0.137) | 0.476 ± 0.038 (0.239) |
| Orthologs                      | 0.184 ± 0.014 (0.118) | 0.197 ± 0.014 (0.174) |
| Predicted membrane proteins    |          |         |
| Paralogs                       | 0.342 ± 0.036 (0.185) | 0.448 ± 0.030 (0.246) |
| Orthologs                      | 0.162 ± 0.004 (0.107) | 0.142 ± 0.004 (0.132) |

The numbers are the average Kn/Ks ratio ± standard error (standard deviation), 0.05 < Ks < 0.5.
Because genes with different functions evolve at different rates [24,25], the observed relaxation of selection in paralogs could potentially be due to an over-representation of fast-evolving genes in the paralogous gene set. In an attempt to isolate the effect of duplication per se on purifying selection, we analyzed paralogs and orthologs that appeared to encode proteins with similar functions, namely secreted and membrane proteins that were identified by signal peptide and transmembrane helix prediction. The difference in the $K_s/K_a$ ratios for paralogous versus orthologous genes was approximately the same for the predicted secreted and membrane proteins as it was for the complete analyzed gene sets (Table 4), indicating that the acceleration of evolution probably is an inherent feature of duplicated genes.

Discussion
The present analysis confirms the earlier observations that paralogs evolve under purifying selection [21], which typically acts with similar strength on both duplicated copies of genes [9,22]. Additionally, however, we found that paralogs evolve significantly faster than unduplicated genes with a similar level of divergence, which is compatible with the notion that gene duplications are a source of new protein functions.

The inconsistency of empirical evidence with Ohno’s model prompts questions on the validity of some of the assumptions underlying this model. One major assumption, which was inherited by the subfunctionalization model, is that one gene copy is sufficient to perform the respective function, so that a gene duplication is redundant and has no effect on fitness [1,10]. This notion has been widely accepted, and often becomes one of the central postulates of models of duplicated gene evolution [3,7,26,27]. Should this be the case, however, a duplication event would only very rarely achieve fixation [28,29]; moreover, in the event that a duplication is slightly deleterious, it would be effectively prevented from achieving fixation [30].

Although the notion of duplication producing redundant genes is central to current theories of duplicated gene evolution, the short-term benefits of gene duplications are well known. This is illustrated by the numerous observations of adaptive gene amplifications in response to antibiotics [31-33], anticancer drug treatments and exposure to various toxins [34-39] or heavy metals [40-44], nutrient limitations [32,33,45-50], pesticide treatments [51-53], extreme temperatures [54,55] and symbiotic and parasitic interactions [56,57]. Combining this information with the observations that recently duplicated genes evolve under purifying selection (21) and our present work), it seems reasonable to hypothesize that a majority of duplicated genes that achieve fixation in a population increase fitness when present in two or more copies in a genome and thus are subject to purifying selection from the moment of duplication.

Recently duplicated paralogs appear to be a nonrandom group enriched in genes coding for proteins involved in different aspects of the organisms’ interaction with the environment (see Additional data files). In particular, a substantial fraction of these paralogs encode (predicted) membrane or secreted proteins. The prevalent functions of duplicated genes varied among different organisms. In bacteria, the majority of these genes encoded different types of surface molecules, which, in pathogens, are involved in interaction with substrate cells. In yeast, there was an emphasis on membrane transporters as well as on genes involved in stress response (for example, heat shock). In multicellular eukaryotes, receptors (for example, olfactory receptors) and (predicted) secreted signaling molecules were predominant. Proteins with a predicted signal peptide were more prevalent in Arabidopsis thaliana (35% of all paralogs versus 17% of all unduplicated genes from the proteome), Caenorhabditis elegans (41% among paralogs versus 31% among unduplicated genes) and Drosophila melanogaster (39% in paralogs versus 23% in orthologs) and predicted membrane proteins were more numerous in mammals (56% in paralogs versus 49% in unduplicated genes). Each of these differences in the functional composition of the sets of paralogs and unduplicated genes were statistically highly significant ($p < 0.0001$, $\chi^2$ test).

A number of genes for which amplification has been shown to have a role in adaptation to various conditions were among the recent duplicates in our dataset. Examples include genes for the yeast hexose transporter, whose amplification has been shown to increase fitness in a low-glucose environment [49], the nematode P-glycoprotein gene, the classical eukaryotic multidrug-resistance gene [36], and the fruit fly copper-binding metallothionein gene whose duplication is implicated in copper tolerance [40,41]. Many genes that are known to amplify in response to selection, primarily during anticancer drug treatments, were found to be present in multiple copies in mammalian genomes as well, in particular, aminoacyl-tRNA synthetases, glutamine synthetase and other anabolic enzymes, adenosine deaminase, ornithine decarboxylase, AMP deaminase, and N-acetyl glucosaminyltransferase [34,46]. Permeases, transporters, synthetases and various detoxification enzymes, which, in general, tend to amplify as a response to stressful conditions [32-34,46,48,51], also showed a substantial presence among the recently duplicated genes in all species (see Additional data files). Duplication of genes belonging to these functional groups is thought to be adaptive because increase in the production of the corresponding proteins may either increase the efficiency of transport of a nutrient into the cell or of toxins out of the cell, increase the rate of catabolism of toxic substances, or allow a greater rate of synthesis of metabolites, for example amino acids, that are required in increased amounts under the given conditions [32-34,46,48,51].

Thus, the observation that purifying selection appears to act on all recent duplicates and examination of the functions of
recently duplicated genes do not support the notion that gene duplication results in true functional redundancy and duplications may achieve fixation despite being redundant [26]. The alternative hypothesis - that gene duplications are fixed in a population by positive selection in all organisms - is supported by a combination of evidence of adaptive duplications from many types of living organisms: prokaryotes [31,33,45,46,48,50,55,56], protists [35,58,59], plants [39,44], fungi [43,49], invertebrates [40,41,51-53], non-mammalian vertebrates [54], as well as mammalian somatic tissues [34,36-38]. Combining these observations with the suggestion that gene duplication may be a general mechanism of adaptation to various conditions of environmental stress [32,33,46,48-50,52,53,55,60], we suggest that, in both prokaryotes and eukaryotes, most paralogs that are fixed in a population have a direct effect on fitness from the moment of duplication, and aid in the adaptation to various environmental conditions, primarily through a protein dosage effect.

That the short-term benefit of a gene duplication is a direct effect on protein dosage also stems from a variety of experimental observations in a number of organisms, prokaryotic and eukaryotic. Gene duplication may be a temporary mechanism to increase protein or RNA dosage, as in the case of rRNA genes in amphibian oocytes and ciliate macronuclei, the chorion genes in some dipterans, actin genes in chicken as well as drug transporters in somatic tissues (see [34,37] for reviews). Protein dosage effects have also been demonstrated in a number of other studies of inheritable adaptive gene duplications [32,34,35,43,44,46,49,51,53,61]. Furthermore, there is evidence from the analysis of the yeast genome that duplicated genes tend to be from those sets of functions that are more highly expressed [62], supporting a general role for selection on protein dosage in duplicated genes.

We show here a substantial acceleration of evolution in recently diverged paralogs compared to orthologs with the same level of synonymous sequence divergence. This acceleration may be explained by positive selection or by a relaxation of purifying selection or by a combination of the two. A few cases of differentiation of paralogs driven by positive selection have been described [63-66]; however, such cases are notoriously difficult to uncover. Thus, we make no attempt here to examine in greater detail the selective forces that act on duplicated genes. In general, however, only a small fraction of sites in some genes appears to evolve under positive selection [15], and relaxation of purifying selection is likely to be the main mechanism behind the acceleration of evolution after duplication.

Our claim of the prevalence of the protein dosage effect as the short-term advantage of duplicated genes is not incompatible with the possibility of a long-term advantage in terms of new functions or the observation that duplicated genes evolve at a faster rate. Indeed, evolution of a new beneficial function in one of the duplicates, once a relatively high level of divergence (which will be different for different genes, depending on the specific function involved) is reached, appears to be common among duplicated genes as demonstrated for several paralogous gene families [33,52,54,67]. Inasmuch as this happens, positive selection might contribute to the observed acceleration of evolution in duplicated genes.

However, assuming that gene duplications primarily evolve under purifying selection, at least soon after duplication, the observed acceleration of evolution may be explained by epistatic interaction between gene copies. In the absence of epistasis, a fitness function is an exponential function that depends on a single additive variable called fitness potential [68-71]. Assuming that the acceleration of evolution observed in paralogs is primarily due to a relaxation of purifying selection, the fitness potential variable describing the fitness function of diverging gene copies can be thought of as the effective amount of functional protein, which depends both on gene duplication and point mutations. Assuming that all point mutations are equally deleterious, such that it takes, for example, 10 point mutations to eliminate the increase in the amount of functional protein resulting from one duplication, \( p = d - m/10 \), where \( p \) is the fitness potential, \( d \) is the number of gene copies and \( m \) is the number of deleterious point mutations. By definition, in the absence of epistasis, selection against a deleterious mutation is constant for all values of the fitness potential.

However, if a fitness function grows at a less than exponential rate, then a relaxation of selection against deleterious mutations is expected [68-71]. For example, consider a linear fitness function \( f(p) = 2p \). In this case, the fitness of an organism with one gene copy is \( f(p) = 2p = 2(1) = 2 \), and the fitness of the same organism with one point mutation is \( f(p) = 2p = 2(d - m/10) = 2(1 - 1/10) = 1.8 \); thus the mutation reduced fitness by 10% (1 - 1.8/2). The fitness of an organism with two gene copies is \( f(p) = 2p = 2(2) = 4 \), the fitness of the same organism with two gene copies and one point mutation is \( f(p) = 2(2 - 1/10) = 3.8 \); in this case, the same mutation will cause only a 5% reduction in fitness (1 - 3.8/4). Thus, the observed acceleration of evolution in paralogous genes implies that either their divergence occurs under positive selection or that the fitness function for gene duplications grows at a slower than exponential rate.

Experimental evidence from several organisms supports the notion that the increase of fitness due to gene duplication is far from being exponential; more specifically, under static environmental conditions, there is an optimal gene copy number [49,50,53]. As duplication events are known to incur a fitness cost [50,53], the fitness function of duplicated genes most probably resembles an inverted parabola, such that it contains a clearly defined maximum [28,53]. Numerous observations that gene duplications that increase protein dosage can be pathogenic [61,72-75] further support the model of optimum copy number for each gene.
Thus, it seems reasonable to hypothesize that, for each gene, there is an optimum number of copies per genome that may vary depending on environmental conditions. Assuming that the environmental condition that determines the optimum gene number fluctuates, positive selection would regulate the gene copy number in the genome accordingly. Under this model, if environmental conditions, and therefore the number of gene copies, fluctuate frequently, then most paralogs will be closely related, such that an excess of (nearly) identical genes will be found within a genome, as indeed has been observed for many genomes ([21] and data not shown).

The present observation that duplicated genes experience a substantial relaxation of selection compared to unduplicated genes is compatible with the traditional view that gene duplications make a major contribution to the evolution of new gene functions. Additionally, the repertoire of protein functions among recent duplicates suggests that many gene duplications contribute to adaptation of the organism to various forms of environmental stress. The results of the present analysis of recent duplications suggest a two-stage evolutionary model of gene duplication: in the first stage, immediately after duplication and during the early phase of their evolution, paralogs are retained and are subject to purifying selection because of the short-term advantage of protein dosage regulation; at a later stage in their evolution, gene duplications are likely to provide a long-term advantage by enabling the creation of new functions.

Materials and methods

Genome sequence data

The complete genome sequences of bacteria and archaea, the yeast Saccharomyces cerevisiae, the nematode C. elegans and the fruit fly D. melanogaster, and the sequences of chromosomes II and IV of the thale cress A. thaliana were extracted from the Genomes division of the Entrez retrieval system [76]. The complete sequences of cDNAs from mouse, rat and humans were extracted from GenBank.

Data analysis

Clusters of paralogs were identified by comparing the protein sequences encoded by all genes from each genome using the BLASTP program [77] followed by single-linkage clustering using the BLASTCLUST program (I. Dondoshansky, Y.I.W. and E.V.K., unpublished results; [78]). Each cluster included sequences that aligned over at least 95% of their lengths with a score density of 1.5 bits per position, which approximately corresponds to 75% identity. Probable orthologs were operationally defined as bidirectional best hits between proteins from two genomes [23]. A subset of probable orthologous pairs that met the criteria used for identification of paralogs (score density of 1.5 bit/position and aligned over at least 95% of their lengths) was selected. Orthologous pairs, in which one or both of the members was identified as part of a paralogous cluster, were excluded from further analysis to ensure a set of unique genes in the orthologous dataset.

Protein sequences were aligned using the CLUSTAL W program [79] and the corresponding nucleotide-sequence alignments were derived, by substituting the respective coding sequences for the protein sequences. The number of synonymous nucleotide substitutions per synonymous site (Ks), and the number of nonsynonymous nucleotide substitutions per nonsynonymous site (Kn) were estimated using the Pamilo-Bianchi-Li method [80,81]. Families of paralogs that included more than two genes were separated into all possible pairs, and each pair was subjected to the same analysis, after which an average of both Ks and Kn was taken and used as a single point. However, only paralogous clusters with two paralogs were used for the relative rate test. For the substitution rate calculations, the data from 26 bacterial genomes, six archaeal genomes and three mammalian genomes were combined into three kingdom-specific datasets; the remaining eukaryotic genomes were analyzed independently. The substitution rates for orthologous gene pairs were calculated using the same approach. Pairs of paralogs and orthologs with Ks > 0.5, which approach saturation in synonymous sites, and those with Ks < 0.05, which are subject to statistical and database-annotation errors, were excluded from all calculations of the Kn/Ks ratios.

For the relative rate test, the closest ortholog from the same taxon was identified for each pair of paralogs. To ensure correct identification, those ortholog candidates that were closer to either of the paralogs than the paralogs were to each other were discarded. In the remaining triplets, which consisted of an ortholog and two paralogs, p-distances P12, P13, and P23 for ortholog-paralog 1, ortholog-paralog 2, and paralog-paralog pairs, respectively, were calculated from the multiple alignment. The conversion between p-distances and corresponding evolutionary distances, D12, D13, and D23, was carried out using the gamma-distance correction [82,83] with an α parameter of 1.0. The significance of the difference between P12 and P13 was estimated using the normalized deviation \( |P_{12} - P_{13}| / \left( P_{12} P_{13} / L_{12} + P_{13} P_{12} / L_{13} \right)^{1/2} \), where \( L_{12} \) and \( L_{13} \) are the lengths of the paralogs [84]. To estimate the significance of the maximum possible difference between the paralogs, evolutionary rates, \( D_{OPmax} \) and \( D_{OPmin} \) were calculated as \( (D_{OP} \pm D_{PP})/2 \) (as follows from the triangle inequality, \( |D_{OP} - D_{OP2} - D_{PP}| \leq 0 \)). D12, D13, and D23 were converted to the respective p-distances \( P_{OPmax} \) and \( P_{OPmin} \); the significance of the maximum possible difference was calculated in the same manner as for the real \( P_{OP} \) and \( P_{OP2} \). Triplets that failed to yield significant difference between \( P_{OPmax} \) and \( P_{OPmin} \) (because of paralogs being too similar) were discarded because no amount of evolutionary pressure (resulting in different rates of evolution in paralogs 1 and 2) would be sufficient to produce a significant difference in \( P_{OP1} \) and \( P_{OP2} \) on such short evolutionary distances.
For prediction of protein function, detailed sequence analysis was carried out wherever needed using the PSI-BLAST program [78] and the SMART system for protein domain identification [85]. Signal peptides in protein sequences were predicted using the SignalP program [86] and membrane proteins using the MEMSAT program [87]. Codon bias was estimated using the effective number of codons measure [88].

### Additional data files
Annotated lists of all recently diverged paralogs and the list of orthologs analyzed in this study are available with the complete version of this paper, online, and at [89].

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