New gene origin and deep taxon phylogenomics -

opportunities and challenges

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Abstract

The immense morphological and phenotypic diversity within eukaryotes coincides with large-scale differences in genic repertoires including the presence of thousands of new genes in every genome. New genes arise through duplication and divergence of existing coding sequences or *de novo* from non-coding sequences. These processes together cause individual genomes to contain up to one-third of orphan genes without any detectable homology in other lineages. Recently, deep taxon phylogenomics, the genome comparisons of extremely closely related species, provided novel insight into the evolutionary dynamics of such rapidly evolving genes. This review focuses on deep taxon phylogenomics and its importance in studying the evolution of new genes, and discusses challenges and opportunities.

New genes can arise from ancestrally coding and non-coding regions

Genetic studies in multiple species have demonstrated that biological innovation is frequently coupled with the evolution of new genes [1–5]. But how new genes arise in the first place remains controversial and is becoming a major research topic in evolutionary biology. Controversy results from the fact that there are multiple potential mechanisms of origin of new genes. Proper definitions of "new," "taxonomically-restricted," "orphan," and "de novo" genes are therefore crucial, even if some of these terms can overlap (see Figure I, Box 1, and Glossary). New genes are genes that emerged recently in a given lineage (Box 1, Figure I). In principle, each genome can be divided into regions that encode for proteins and regions that do not. Therefore, the primary distinction between different types of new genes is whether they originate from ancestrally protein-coding or non-coding genomic sequences [6,7] (see Glossary). While duplication of protein-coding genes is certainly one of the major forces generating new genes, the origin of orphan genes which lack traceable homologs in other lineages, is much less understood. It was previously considered that duplication-divergence would be the only mechanism to produce orphan genes out of previously protein-coding material [8], but multiple studies have shown that divergence alone, exaptation of transposable elements, strand-switching, and resurrection of pseudogenes can create orphan genes without the need of duplication [9-12] In prokaryotes, new genes can also arise frequently by horizontal gene transfer (HGT) [13]. HGT can be considered as a special case of gene duplication where a copy of a foreign gene is integrated into the host genome. In contrast, HGT in eukaryotic genomes can only explain a meager fraction of new genes despite numerous well-documented reports [14–16].

De novo genes make up the most controversial class of new genes (see Glossary). Even though the idea of gene emergence through *de novo* creation from non-coding sequence was raised in the first half of the 20th century [17], duplication was considered the only viable

mechanism for gene birth until the end of the last decade. The chief proponent for the exclusivity of the duplication mechanism, Susumu Ohno, claimed in his renowned book *"Evolution by gene duplication"* that due to the relentless pressure of natural selection only a redundant cistron is unfettered to emerge as a new gene locus [18]. The stance of stalwarts such as Ohno and Francois Jacob against the *de novo* gene origin [19], coupled with the enormous difficulty of finding evidence supporting such mechanisms had stifled investigation into gene origin apart from duplication. It took decades until the sheer abundance of orphan genes (see Glossary and Box 1 for definition) in eukaryotic genomes and the first evidence of *de novo* gene birth has been intensively discussed [8,22–25]. In this review, we focus on the importance of deep taxon phylogenomics in studying new gene origin and their evolutionary dynamics and we will discuss associated opportunities and challenges.

Deep taxon phylogenomics is indispensable for understanding new gene origin

Since the first sequencing projects, eukaryotic genomes were known to contain substantial fractions of orphan genes, but whether orphan genes are the result of strong divergence or *de novo* origin is still debated. Historically, two complementary approaches have been used to identify the origin of orphan genes: bioinformatic methods for remote homology detection [26,27] and phylogenomic approaches including phylostratigraphy (see Glossary) [28– 32]. While both approaches are widely used, it is important to note that they have distinct capabilities to investigate different classes of new genes. Methods like PSI-blast [26] or HHsearch [27] can detect distantly related proteins, which are missed by BLASTP thus leading to initial classification as orphan genes. Such approaches work best for large orphan gene families, where many sequences are available to construct generalized sequence profiles. However, methods for remote homology detection are better suited to identify extensive

divergence rather than *de novo* origin, as orthologous non-coding nucleotide sequences get scrambled beyond recognition at deeper evolutionary time scales.

Phylogenomic approaches allow for studying the ages and origins of new genes, in particular they enable the characterization of orphan and taxonomically-restricted genes [28,32]. However, the power of such approaches lies with their phylogenetic resolution (Fig. 1). While genomes of different insect orders are too divergent for studying the evolution and origin of new genes, comparisons between closely related members of the same genus, such as Drosophila, can be used to reliably measure evolutionary constraints as well as to make statements about gene origin [28,33–35]. Studies in the last decade began to use closely related species that are separated only for short time periods (Figure 1A). Such deep taxon phylogenomics can capture recent evolutionary origins and dynamics. In principle, the use of a robust phylogenomic framework allows the identification of enabling mutations, i.e. mutations that allow the transition from non-coding to coding sequences, which is considered to be the strongest indicator of a de novo origin. Indeed, some recent studies have elucidated the mechanisms of origin and evolutionary dynamics of new genes, i.e. studies in primates [11,20,37–39], yeast [29,40,41], mouse [36,42] and Drosophila [34,35,43] spearheaded this research. Such established model organisms for genome evolution can rely on well curated genomic resources, which are important to distinguish true orphan genes from annotation artifacts. In addition, recent studies also employed deep taxon sampling also in non-classical model organisms, such as Drosophila pseudobscura [28], the yeast genus Lachancea [29,44], rice [30,45], and the nematode Pristionchus pacificus [9,31], the latter of which will be discussed in more detail in the following section.

Finally, it is important to note that many aspects of genome architecture can have a strong influence on the frequency and patterns of new gene origin [46]. Specifically, differences in genomic composition, *i.e.*, fraction of coding sequence, intron size, and transposable elements can lead to largely different contributions of origin mechanisms. For example, the

frequency of *de novo* origin would be expected to be much higher in a mammalian genome with 3% coding sequences as opposed to a nematode genome with 20-30% coding sequences. Therefore, it is very likely that in order to reveal the full complement of mechanisms of origin of new genes, genomic analysis must be coupled with both broad taxonomic coverage and deep taxon phylogenomics.

Deep taxon phylogenomics of *Pristionchus* and the origin and evolution of new genes in nematodes

Fungi, insects, mammals and plants have traditionally provided important insight into genome evolution and the occurrence of new genes. Recently deep taxon phylogenomics was applied to study genome evolution and the emergence of new genes in nematodes of the genus *Pristionchus*. The hermaphroditic species *P. pacificus* is a satellite model to *Caenorhabditis elegans* with an established functional toolkit [47–49]. This species is a soil nematode that is reliably found in association with scarab beetles in world-wide samplings, which enabled the systematic search for related species with similar associations [50–54]. As a result, a collection of more than 40 *Pristionchus* species is currently available, some of which form sterile F1 hybrids indicating that these species are extremely closely related. A phylogenetic framework of the *Pristionchus* genus was established by transcriptome sequencing of all species and revealed striking patterns of gene loss [55].

Initial sequencing of the *P. pacificus* genome classified roughly one third of all genes as orphan genes when *P. pacificus* was compared with *C. elegans* [56–58]. These two nematodes belong to distinct families, the Diplogastridae and Rhabditidae, respectively, and their protein sequence divergence is five times larger than the divergence between human and mouse [56]. Given that until recently the *P. pacificus* genome was the only diplogastrid genome available, the high number of orphan genes is likely due to sparse taxon sampling. Nevertheless, as initial gene annotations were completely based on automated gene predictions with the guidance of

limited transcriptome and proteome data [56,57], it was important to investigate if *P. pacificus* orphan genes are real. Indeed, RNA-seq data as well as comparative and population genomic data revealed that the vast majority of orphan genes is expressed under standard laboratory conditions and is under evolutionary constraint [59].

Utilizing the high phylogenetic resolution of *Pristionchus* nematodes, a more detailed phylogenomic analysis was performed by sequencing the genomes of eight species with a ladder-like phylogeny and two outgroups to determine the age of new genes and to study their evolutionary dynamics [31]. This analysis revealed a number of genome-wide features. First, deep taxon sampling and the selection of species with a ladder-like phylogeny allowed for the classification of new genes into age classes and to contrast their evolutionary dynamics at various time scales. Second, genes of new age classes are localized at the chromosomal periphery, whereas they are rare in the chromosomal centers. This observation also reflects general trends of nematode chromosome architecture such as higher recombination rate and more genetic diversity at the chromosome arms as well as distinct epigenetic profiles between chromosome arms and centers [58,60]. Third, many new genes show weaker expression as opposed to old genes, thus confirming previous findings. This suggests that expression either increases or becomes broader over time [31] and is consistent with the "out of Testis" hypothesis which states that either sexual selection or special cellular environments may create favorable conditions for the expression of gene-like sequences [25]. Interestingly, a parallel study of *P. pacificus* epigenetic profiles found evidence that new genes arise in the vicinity of enhancers of older genes [60]. This is consistent with previous finding of abundant transcriptional activity around cis-regulatory regions [61,62] and suggests that enhancers may function as promoters for new genes [60]. Fourth, new genes have a higher propensity of being lost than older and conserved genes. And finally, new genes exhibit only weak evolutionary constraints.

A more detailed study of orphan origins applied phylostratigraphy and additional quality filters to define a high confidence set of 29 species-specific genes, for which the origin could be traced based on manual inspection of syntenic regions in closely related genomes [9]. This revealed diverse divergence mechanisms including chimeric origin, alternative reading frame usage, and gene splitting with subsequent gain of *de novo* exons as well as cases of complete de novo origin (Figure 1 B-F). In addition, this study pointed out that technical problems such as annotation artifacts and heuristic failure of homology searches inflate the number of speciesspecific orphan genes. Together, this work and related studies of gene duplication established P. pacificus [62,63] as the primary system to study the evolution of new genes in nematodes. These studies also confirmed many known trends from vertebrates, insects, and plants, including the either low or spatiotemporally restricted expression of new genes [42,65,66] and the inverse relationship between evolutionary rate and age [28,30,67,68]. Importantly however, the availability of chromosome-scale assemblies revealed differences in the chromosomal distribution of new genes across phyla. For example, while novel genes cluster near the centromeric regions in rice genomes, nematodes have holocentric chromosomes and new genes preferentially cluster at the chromosome arms [30,31,58]. In Drosophila new genes seem to be enriched on certain sex chromosomal areas [28], whereas in humans an association with DNA replication timing has been reported [69]. These differences highlight the need to study the evolution and origin of new genes across different phyla.

What does it take to be a gene: from intra-species characterization to pervasive translation

Despite the power of deep taxon phylogenomics, it is not likely to provide a full understanding of the evolutionary dynamics of all new gene classes. Important challenges remain for multiple reasons. First, even at considerably high inter-species phylogenetic resolution, the detection of molecular fossils, such as enabling mutations, is often hampered by the rapid divergence of non-coding sequences. Importantly, apart from identifying the enabling mutations in a sister species, an additional non-coding sequence from an outgroup species is required to reliably infer that the enabling mutation is actually the derived state. With only one sequence from a sister species, a putative enabling mutation could actually represent a pseudogenization event of an ancestral gene of unknown origin. Given these difficulties, conclusive evidence for *de novo* origin is still quite rare [9,70,71]. The resulting large number of species-specific orphan genes, for which origin cannot be analyzed at the species level, either hints at rapid divergence of corresponding non-coding sequences in the sister species (e.g. complex structural variations, ORF switching) or indicates that the standard approaches for homology detection (e.g. BLAST) are inadequate to find traces of homology at the species level. Therefore, to achieve an optimal phylogenetic resolution, the inter-species comparison has to be complemented with intra-species studies to compare genomes of multiple diverging populations. Currently, corresponding resources have been developed in several species with the most comprehensive dataset available in humans [35,72].

Second, besides phylogenetic resolution additional conceptual problems exist. One important hurdle in the determination of the relative contribution of the two gene origin mechanisms is the ascertainment that a candidate gene is really protein-coding. As new genes tend to be expressed either weakly or in a very restricted manner [60,73], direct evidence of transcription and translation is often limited [59,71]. Similarly, indirect evidence based on evolutionary constraints tends to retain low statistical power for individual genes at the intraspecies level [59,74]. In addition, recent studies employing ribosome profiling proposed that large numbers of translated ORFs seem to be either non-functional or serve regulatory roles in the expression of downstream coding sequences while their actual peptide sequences hold little significance [75–78]. Thus, the ability to distinguish all bona fide protein-coding genes remains elusive even at a very narrow phylogenetic distance; measures of evolutionary constraint are

not powerful enough, and direct evidence of translation is either unavailable or inconclusive. Together with accumulating evidence that the lifespan of new genes may be associated with their mechanism of origin [8,28,79,80], the relative contribution between *de novo* and divergence will also depend on the definition of a gene in a given study [84]. If putative products of pervasive transcription and translation (see Glossary) with annotated ORFs are already considered as gene-like sequences, more genes can be classified as *de novo* genes, in contrast to a more conservative definition that only considers genes that survived long enough to have gained verifiable exon-intron boundaries [9] (Figure 2).

In summary, we argue that systematic approaches and individual case studies of intraand cross-species phylogenomic data sets are necessary to elucidate gene origin mechanisms and to quantify their contribution. Cross-species analyses at maximal phylogenetic resolution are needed to study evolutionary constrained and most likely functional new genes. At the same time, intra-species studies are well suited to characterize the raw material, out of which new genes are born. Finally, this has to be complemented by comprehensive transcriptomic data across life stages and tissues to test hypotheses about their regulation and developmental importance [81,82].

Linking mechanism of origin and evolutionary stability

Numerous new genes have been associated with phenotypic innovations, but most of them are taxonomically-restricted duplicates [1,4] or orphan genes of unknown origin [2,3]. This puts into doubt the importance of *de novo* genes for phenotypic evolution. In addition, the fast turnover and limited constraint acting on gene-like sequences raises the questions of how quickly new genes acquire functions at an organismic level, and which types of new genes live long enough to be integrated into the biology of their hosts. We hypothesize that the high numbers of reported proto-genes (see Glossary) and apparently non-functional peptides (as a result of pervasive transcription and translation), is just a byproduct of the cells[′] inability to

regulate the transcriptional and translational machinery tightly enough to only express those genes that are absolutely necessary at a given point in time. This exposes non-coding sequences to a low level of basal expression, and thus offers a playground out of which novel functions may evolve. The idea that random polypeptides might display many unfavorable properties, such as toxicity due to aggregation, has inspired the preadaptation hypothesis (see Glossary, [76,80]). This theory states that initial purging of weakly expressed but strongly deleterious peptides shifts the raw material towards more gene-like properties. In combination with loss of non-functional peptides due to genetic drift this leaves only a small pool of survivors [79] beyond a certain age. Even if these *de novo* genes have no particular function as of yet, they get better adjusted to the cellular environment due to their long exposure and may start interacting with other components of the cellular network. It is important to note here that new duplicates, unless they are immediately strongly selected against due to dosage imbalance [83], have at least a couple of advantages over *de novo* genes. First, they are less likely to exhibit toxicity as they derive from sequences that evolved over sufficiently long periods to remove their toxicity. Further, sequences arising from duplications can easily be longer than newly evolved ORFs, are more likely to have an optimized amino acid composition and codon usage, and have higher chances of containing secondary structures or functional motifs that readily allow them to interact with the existing cellular networks. Once the integration of a new gene into any cellular network provides a fitness advantage to the organism, selection will act to preserve such a gene leading to long-term survival. Thus, initial persistence of gene-like sequences may be the key to acquiring biological functions and evolutionary stability. The fundamental differences in starting conditions between different classes of new genes may explain why de novo genes are frequently generated but guickly lost, while at the same time duplicated genes dominate the population of new genes at larger evolutionary distances [28,79]. At smaller time scales, it was proposed that *de novo* gene birth may be more prevalent then gene duplication [40]. This initial observation from yeast was further supported by the finding that the amount of novel ORFs in

rodent genomes outnumbers species-specific duplications by at least an order of magnitude [42,78]. The proposed model (Figure 2) would indicate that quantifying the contribution of duplication vs. *de novo* formation to new gene emergence makes sense only if the variable lifespans of different classes of new genes are also considered. Thus, to fully understand the dynamics between gene emergence and loss, systematic studies are needed which measure the contribution of different origin mechanisms to new gene emergence across various time scales. Consequently, this requires deep intra-and inter-species taxon <u>phylogenomics</u>.

Concluding Remarks and Future Perspectives

The question of how molecular innovations facilitate phenotypic novelties is central to evolutionary biology, and understanding how new genes are formed and retained is the key to answering it. With the availability of deeply sampled phylogenomic data sets, it will soon be possible to construct models of genome evolution describing how new genes are formed and why they are retained. One of the most striking conceptual implications from recent studies is that the notion of a gene as a discrete molecular entity needs to be shifted to a continuum of sequence types with various degrees of biological activities. This spectrum on one end begins with products of pervasive transcription and translation, moves to very recent *de novo* genes or protogenes, includes pseudogenes and long non-coding RNAs, and finally ends with what we traditionally consider a functional gene (Figure 2). We hypothesize that persistence of gene-like sequences is crucial for acquiring biological functions and that different classes of new genes are not equally well equipped to survive this early phase. Thus, future studies at the interspecies and intra-species level (see Outstanding Questions) are required to group the origins and dynamics of new genes into a general framework that can be linked to phenotypic evolution.

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Box 1 - What are new genes?

New genes are genes that based on sequence comparison appear to have emerged recently in a given lineage. By definition, they are taxonomically-restricted or even species-specific. Importantly, gene duplication events can also be taxonomically-restricted and thus, we also consider products of duplication events as new genes. Since taxonomically-restricted genes can be the result of gene loss, a comprehensive analysis of the phylogenetic context (e.g. phylostratigraphic analysis) is needed to distinguish losses from the emergence of new genes. New genes that lack detectable sequence homology (e.g. BLASTP e-value < 0.001) in other taxa are commonly referred to as orphan genes. The definition of orphan genes is always context-dependent and in sparsely sampled taxonomic clades, orphan genes make up to onethird of a gene set in a given genome [8]. Conversely, some orphan genes may be relatively old and the lack of detected homology may be a result of strong divergence. Such firm divergence can create entirely new sequences and such genes may act as the conduits of evolutionary innovation and arguably should be treated as new genes. Whether orphan genes are just a product of extensive divergence from ancestrally protein-coding genes or arose *de novo* from non-coding sequences is one of the most actively studied questions in evolutionary biology. While none of these different gene classes is absolutely identical to the other (Figure I), genes in all of these categories tend to show similar properties, such as spatiotemporally restricted expression, the high propensity of being lost, and relaxed evolutionary constraint; we consider all the different types of taxonomically-restricted genes (duplicated genes, orphan genes, and de novo genes) as new genes.

Figure I Different classes of new genes. New genes arose recently in an evolutionary lineage and are consequently taxonomically-restricted. Out of all taxonomically-restricted genes, we consider duplicated genes, orphan genes, and *de novo* genes as new genes.

Glossary

Deep taxon phylogenomics: Genome comparisons of closely related species with a well-defined molecular phylogeny.

De novo genes: Genes that emerged from an ancestrally non-coding sequence (intergenic, intronic, or non-coding RNAs).

Orphan genes: A gene without detectable protein homology outside a predefined taxonomic clade.

Pervasive translation: The idea that large parts of the genome are transcribed and translated at a background level.

Phylogenomics: Genomic analysis in a given phylogenetic framework.

Phylostratigraphic analysis: An approach to determine the age of a given sequence by tracing the founding member in a species tree based on BLAST searches in extant taxa.

Protein-coding sequence: Exonic sequences that are translated into peptides.

Proto-genes: Products of pervasive translation with some genic properties but without any biological function.

Pseudogenes: Genes that have lost functions due to mutations in their coding sequence (e.g. stop codons, frame shifts).

Figure legends

Figure 1 Deep taxon sampling and the origin of orphan genes. (A) Deep taxon sampling facilitates the investigation new gene formation in closely related genomes. Silhouette images of animals were taken from PhyloPic (www.phylopic.org) and are available for reuse under the Public Domain Dedication 1.0 license. (B-F) The diversity of orphan gene origin as identified in *Pristionchus* nematodes. (B) The schematic genomic locus shows a candidate orphan gene (red) with two neighboring syntenic anchor genes (blue, green). Closer investigation of the orthologous region in sister taxa may eventually reveal either *de novo* formation (B) or divergence (C) as the mechanism of origin. (D) Partial exonic duplications from different source genes can result in a chimeric orphan gene. Loss of detectable homology is caused by the small size of duplicated fragments combined with moderate divergence. (E) Gene split with subsequent gain of a completely new exon can result in a new orphan gene. In this case indels in the duplicated exon induce frameshift causing the loss of protein homology. (F) Partial duplication, intron gain, and transcription from the opposite strand generate an orphan gene. (G) Orphan genes can arise by duplication and actualization of an alternative ORF in a putatively overprinted gene.

Figure 2 Hypothetical model of birth rate and lifetime of new genes. The lifespans of new genes may depend on their mechanism of origin. Thus, in this model the relative contribution of different mechanisms will depend on the definition of a gene and the evolutionary time scale of the comparison.

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Orphan gene





Taxonomically Restricted Genes