Phylogenomic approaches underestimate eukaryotic gene transfer

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Phylogenomic approaches have shown that eukaryotes acquire genes via gene transfer. However, there are two fundamental problems for most of these analyses; only transfers from prokaryotes are analyzed and the screening procedures applied assume that gene transfer is rare for eukaryotes. Directed studies of the impact of gene transfer on diverse eukaryotic lineages produce a much more complex picture. Many gene families are affected by multiple transfer events from prokaryotes to eukaryotes, and transfers between eukaryotic lineages are routinely detected. This suggests that the assumptions applied in traditional phylogenomic approaches are too naïve and result in many false negatives. This issue was recently addressed by identifying and analyzing the evolutionary history of 49 patchily distributed proteins shared between Dictyostelium and bacteria. The vast majority of these gene families showed strong indications of gene transfers, both between and within the three domains of life. However, only one of these was previously reported as a gene transfer candidate using a traditional phylogenomic approach. This clearly illustrates that more realistic assumptions are urgently needed in genome-wide studies of eukaryotic gene transfer.

Transfer of genetic material between different organismal lineages is important in prokaryote evolution. Studies of single gene families as well as phylogenomic studies in the last decade have shown that also eukaryotes are affected by this evolutionary mechanism. However, the importance of the process is still uncertain; only modest numbers of gene transfer candidates are typically reported from eukaryotic genome projects, whereas directed studies suggest gene transfer to be important in the adaptation process of eukaryotes. This is an intriguing incongruity. To understand these differences it is useful to consider the assumptions applied in the screening procedures in the phylogenomic approaches used in genome projects (referred to as ‘traditional phylogenomic approaches’ herein) and how well they match the knowledge we currently have from more directed studies of eukaryote gene transfer. Here I will argue that the match is really poor leading to a high number of false negatives.

Studies of gene transfers in eukaryotes using phylogenomic methods typically identify eukaryotic proteins with high sequence similarity to a prokaryotic protein, but with no or significantly weaker similarity to any eukaryotic protein. This is indeed a strong indication of a gene transfer event. The problem is that these traditional phylogenomic approaches only identifies protein families in which a single transfer has occurred between a prokaryote and a eukaryote, which probably is a rarity. Comparative genomics studies of prokaryotes have shown that most protein families are patchily distributed; they are absent from a small or large fraction of the genomes (Fig. 1). These genes are distributed via gene transfer and provide diversifying functions and niche adaptation to the recipients. Eukaryote genome evolution, on the other hand, has been viewed as mainly influenced by genome expansion and a few major endosymbiotic events. However, there are data suggesting that gene transfer of patchily distributed proteins is important for the diversification process also for eukaryotes.
Here I will review some recent findings of adaptation by gene acquisition in eukaryotes obtained by the usage of phylogenomic approaches for the study of gene transfers between specific eukaryotic groups, and one attempt to use a novel approach to study patchily distributed proteins. With an increased understanding of the evolutionary dynamics of all classes of gene families (Fig. 1) we can apply realistic assumptions to large-scale studies of eukaryotic gene transfer.

**Directed Studies Identify Gene Sharing Leading to Adaptation**

Many of the most devastating diseases in plants are caused by fungi or oomycetes. These are two distantly related groups of eukaryotes that have similar lifestyles. Both feed by osmotrophy. The cells secrete enzymes that decompose organic material and the metabolites are imported into the cell. The similarity in lifestyle between the groups is an example of convergent evolution. Fungi are more closely related to animals than to oomycetes, whereas diatoms, a group of photosynthetic algae, are a sister group to oomycetes. Absence of phagotrophy has been assumed to be a barrier to gene transfer. Indeed, the oomycete and fungi genomes are not among the genomes for which traditional phylogenomic studies have indicated a significant role of gene transfer in eukaryotes. Nevertheless, targeted evolutionary studies have suggested that gene transfer contributed to the similarities between the groups.

Richards and coworkers studied this phenomenon further. They could identify dozens of gene transfers between the groups using a wide range of genomes from both groups together with clustering and phylogenetic methods. Interestingly, all transfers except one were reported to have occurred in the direction from fungi to oomycetes. Many of the transferred genes encode secreted decomposing enzymes and were specifically acquired by plant-tissue colonizing oomycetes. These results show that oomycete most likely are more recent plant pathogens than fungi and that transfer of genetic material from a distantly related eukaryotic group have played an important role in evolution of their pathogenic lifestyle. These fascinating results would not have been obtained with a traditional phylogenomic approach in which genes with strong sequence similarities to other eukaryotes would have been assumed to be present in the common eukaryotic ancestor.

Studies of gene transfer are indeed able to shed light on the diversification process of eukaryotes. Animals and fungi are both members of Opisthokonta. No photosynthetic member has been identified in this group. Chaoanoflagellates are a group of free-living microbial eukaryotes which are the closest relatives to animals. Sun and coworkers used a directed phylogenomic approach to search for genes of algal origin in the genome of *Monosiga brevicollis*, a phagotrophic unicellular choanoflagellate. They reconstructed phylogenetic trees for all genes in the genome. Using realistic filtering criteria they were able to identify 103 genes with strong support for algal origin, mostly from haptophytes, diatoms and green algae. This could be the result of repeated transfer of genes from food; choanoflagellates feed on bacteria and other eukaryotes. Alternatively, or rather in addition, the genes could have been introduced from a past algal endosymbiont in the lineage leading to *Monosiga*. Interestingly, a quarter of the identified genes appeared to first have been transferred from bacteria to a eukaryotic alga, and then secondarily to choanoflagellates. However, such a bacterial origin would not have been detected using a typical phylogenomic approach since the strongest sequence similarity would be to algal gene of bacterial origin. Functions in amino acid and carbohydrate metabolism dominated among the gene transfer candidates, indicating that these choanoflagellates have adapted by acquisition of algal genes that expand their metabolic repertoire.

**A Novel Approach to Study Patchily Distributed Proteins**

The two examples outline above test well-defined hypotheses about gene transfers by using directed phylogenomic methods.
Traditional Phylogenomic Studies have Drastically Underestimated the Amount of Gene Transfer

Only a single protein among the 49 identified as patchily distributed was among the 18 gene transfer candidates in the original *D. discoideum* genome publication, and very few were among the 184 lateral gene transfer candidates reported from *N. gruberi*. This may be surprising, but is logical if the details of the methods applied are considered. Dictyostelium genes with significant similarity to a bacterial-specific Pfam domain and only present in Dictyostelium among eukaryotes were considered as gene transfer candidates. This conservative approach is unlikely to pick up false positives, but will be very prone to false negatives. Genes acquired via gene transfer in two or more different eukaryotes are excluded, as are any genes without sufficient sampling among prokaryotes to be included in Pfam. Similarly, the *N. gruberi* gene set was screened with similarity searches, and genes with significant similarity only to prokaryotes were considered as gene transfer candidates.

Again, gene families with repeated transfers are missed in the screen and eukaryote-to-eukaryote transfers are not even considered. The true number of gene families in these microbial eukaryotes are likely much larger than has been reported. These discrepancies should not be surprising from a biological viewpoint. Microbes live in steadily changing environments. Ecosystems are inhabited by distantly related organisms which have adapted to its specific condition. The spread of patchily distributed genes are part of this adaptation process, and there is no reason to assume that microbial eukaryotes do not take part of this flux of genetic material (Fig. 1). For example, if a gene provide the ability to utilize a carbon compound present in the environment it is likely to spread to different microbes in the environment previously lacking this ability (provided that there are mechanisms in action). The assumption that a gene has a vertical eukaryotic history is violated as soon as two eukaryotic lineages inhabit a similar environment and acquire their copy of a particular gene family independently during the adaptation process. The traditional phylogenomic approaches will fail to identify members of such protein families as gene transfer candidates because they assume that vertical inheritance is the norm for all protein families with gene transfer events as very rare exceptions. However, this is probably only the case for universal core genes, and certainly not for patchily distributed proteins (Fig. 1). Traditional phylogenomic approaches probably only have scratched the surface of the gene transfer events and thereby drastically underestimated the impact of the process on eukaryotic genome evolution.

![Fig. 1B](http://dictybase.org/)

Figure 1B shows an example of an individual gene tree. The exact details of the transfer events could in many cases not be traced, because the density of organismal sampling was too low. Nevertheless, there are no strong indications that any of the proteins have evolved solely via vertical inheritance and gene loss; gene transfer has likely affected all patchily distributed genes identified in the analysis to some extent. The soil-dwelling cellular slime mold *D. discoideum* was selected in the case study for two reasons: an active research community have produced a high quality annotation of the genome sequence (http://dictybase.org/), and only 18 potential gene transfers were reported in the original publication. I identified 49 protein families in the Dictyostelium genome which were shared with at least one prokaryotic species, but only a limited number of other eukaryotes and prokaryotes (Fig. 1). The evolutionary history of these patchily distributed families were analyzed further. For seven of the families there were no eukaryotic sequences except the Dictyostelium sequences. The remaining 42 families contained sequences from one or more eukaryotic species outside the Dictyostelium genus. The closest relative with a completely sequenced genome, the human parasite *E. histolytica*, was represented in only two families. In contrast, the amoebal flagellate *Naegleria gruberi* had a representative in 25 of the families. Dictyostelium and Naegleria are having somewhat overlapping lifestyles, they are both free-living heterotrophs that can be found in soil and they both undergo cell differentiation under certain conditions. However, they are distantly related eukaryotes classified within two different superfamilies: Amoebozoa and Excavata.

There exist at least two alternative plausible explanations for this striking gene-sharing pattern. These genes were present in the common ancestor of the Dictyostelium and Naegleria and distributed in eukaryotes strictly by vertical inheritance. In lineages that have different lifestyles (i.e., parasites) the genes have become obsolete and lost over evolutionary time. Alternatively, the genes have been distributed via gene transfer in more recent evolutionary timescales providing selective advantage to the recipient lineages. Phylogenetic analyses were performed on all protein families to distinguish between these alternatives. The results were striking. The vast majority of the phylogenetic trees showed strong indications of lateral gene transfer between prokaryotes and eukaryotes and within eukaryotes. Figure 1B shows an example of an individual gene tree. The exact details of the transfer events could in many cases not be traced, because the density of organismal sampling was too low. Nevertheless, there are no strong indications that any of the proteins have evolved solely via vertical inheritance and gene loss; gene transfer has likely affected all patchily distributed genes identified in the analysis to some extent. This conservative approach is unlikely to pick up false positives, but will be very prone to false negatives. Genes acquired via gene transfer in two or more different eukaryotes are excluded, as are any genes without sufficient sampling among prokaryotes to be included in Pfam. Similarly, the *N. gruberi* gene set was screened with similarity searches, and genes with significant similarity only to prokaryotes were considered as gene transfer candidates. Again, gene families with repeated transfers are missed in the screen and eukaryote-to-eukaryote transfers are not even considered. The true number of gene families in these microbial eukaryotes are likely much larger than has been reported. These discrepancies should not be surprising from a biological viewpoint. Microbes live in steadily changing environments. Ecosystems are inhabited by distantly related organisms which have adapted to its specific condition. The spread of patchily distributed genes are part of this adaptation process, and there is no reason to assume that microbial eukaryotes do not take part of this flux of genetic material (Fig. 1). For example, if a gene provide the ability to utilize a carbon compound present in the environment it is likely to spread to different microbes in the environment previously lacking this ability (provided that there are mechanisms in action). The assumption that a gene has a vertical eukaryotic history is violated as soon as two eukaryotic lineages inhabit a similar environment and acquire their copy of a particular gene family independently during the adaptation process. The traditional phylogenomic approaches will fail to identify members of such protein families as gene transfer candidates because they assume that vertical inheritance is the norm for all protein families with gene transfer events as very rare exceptions. However, this is probably only the case for universal core genes, and certainly not for patchily distributed proteins (Fig. 1). Traditional phylogenomic approaches probably only have scratched the surface of the gene transfer events and thereby drastically underestimated the impact of the process on eukaryotic genome evolution.
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