Old genes in new places: A taxon-rich analysis of interdomain lateral gene transfer events

Auden Cote-L’Heureux\textsuperscript{1,}a, Xyrus X. Maurer-Alcalá\textsuperscript{2,}b, Laura A. Katz\textsuperscript{1,3,*}

\textsuperscript{1} Department of Biological Sciences, Smith College, Northampton, Massachusetts, United States of America, \textsuperscript{2} Institute of Cell Biology, University of Bern, Bern, Switzerland, \textsuperscript{3} Program in Organismic Biology and Evolution, University of Massachusetts Amherst, Amherst, Massachusetts, United States of America

\textsuperscript{a} Current address: Division of Invertebrate Zoology and Sackler Institute for Comparative Genomics, American Museum of Natural History, New York city, New York, United States of America

\textsuperscript{b} Current address: Northeastern University, Department of Biology, Boston, Massachusetts, United States of America

\textsuperscript{*} lkatz@smith.edu

Abstract

Vertical inheritance is foundational to Darwinian evolution, but fails to explain major innovations such as the rapid spread of antibiotic resistance among bacteria and the origin of photosynthesis in eukaryotes. While lateral gene transfer (LGT) is recognized as an evolutionary force in prokaryotes, the role of LGT in eukaryotic evolution is less clear. With the exception of the transfer of genes from organelles to the nucleus, a process termed endosymbiotic gene transfer (EGT), the extent of interdomain transfer from prokaryotes to eukaryotes is highly debated. A common critique of studies of interdomain LGT is the reliance on the topology of single-gene trees that attempt to estimate more than one billion years of evolution. We take a more conservative approach by identifying cases in which a single clade of eukaryotes is found in an otherwise prokaryotic gene tree (i.e. exclusive presence). Starting with a taxon-rich dataset of over 13,600 gene families and passing data through several rounds of curation, we identify and categorize the function of 306 interdomain LGT events into diverse eukaryotes, including 189 putative EGTs, 52 LGTs into Opisthokonta (i.e. animals, fungi and their microbial relatives), and 42 LGTs nearly exclusive to anaerobic eukaryotes. To assess differential gene loss as an explanation for exclusive presence, we compare branch lengths within each LGT tree to a set of vertically-inherited genes subsampled to mimic gene loss (i.e. with the same taxonomic sampling) and consistently find shorter relative distance between eukaryotes and prokaryotes in LGT trees, a pattern inconsistent with gene loss. Our methods provide a framework for future studies of interdomain LGT and move the field closer to an understanding of how best to model the evolutionary history of eukaryotes.
another kind of inheritance that is not so clear-cut but which has important effects on many groups of organisms. The "lateral" transfer of genetic material, transfer of genetic material between individuals that is not from parent to offspring, profoundly impacts the evolution of bacteria and archaea (i.e. prokaryotes). Rapid resistance to antibiotics is largely driven by this well-accepted phenomenon, and the rise of broad-scale genomic datasets has allowed detailed exploration of lateral gene transfer events in both prokaryotes and eukaryotes. However, there are few widely accepted standards for detecting genes that have been laterally transferred in eukaryotes, where many lineages remain under sampled. Here, we use a taxon-rich phylogenomic approach to develop and deploy a suite of methods for analyzing lateral gene transfer in eukaryotes. We first identify several hundred genes that have likely been transferred laterally from prokaryotes to eukaryotes, then we characterize their functions to detect broad-scale patterns, and finally, for most of these events, we reject the alternative hypothesis: gene loss explains the topology of laterally-transferred gene trees. Together, these data and approaches will be of use to future studies of LGT events in lineages across the eukaryotic tree of life.

Introduction

Lateral gene transfer (LGT), the transfer of genetic material that is not from parent to offspring, is often neglected in models of eukaryotic evolution. This is problematic given the potential innovations enabled by such events and in light of existing data on the ubiquitous nature of LGT in bacteria and archaea. In some archaeal lineages, for instance, LGT is so pervasive that linkage disequilibrium is near that of sexual eukaryotes [1], and LGT is a major driver behind phenomena such as antibiotic resistance in bacteria [2,3]. Although first discovered and most widely studied in bacteria, LGT is not strictly limited to prokaryotes; endosymbiotic gene transfer (EGT), the transfer of genes from mitochondrial or plastid genomes (or endosymbiont nuclei/nucleomorphs in photosynthetic eukaryotes that acquired plastids from other eukaryotes) to the nucleus, is also well-documented [4–6]. Outside of EGT, however, the extent to which LGT affects eukaryotic evolution is debated [7,8]. Some have argued that, while EGTs are relatively common, other interdomain transfer events are very rare and have little effect on eukaryotic genomes [4]. Other studies suggest LGTs are not so uncommon [8–15], though the preponderance of gene loss has likely obscured these events [16].

Past attempts to identify laterally transferred genes (LTGs) in eukaryotes have relied on detecting deviations from eukaryotic monophyly in single-gene trees and, to a lesser extent, discoveries of aberrant composition to identify very recent (i.e. not yet ameliorated) LTGs, though due to rapid amelioration many past studies have found no significant difference in GC content between putative LGTs and other protein-coding genes [17–22]. We apply filtration by composition in a phylogenomic context to mitigate the effect of contamination in transcriptomic data. Sequence-similarity and BLAST-based metrics such as the alienicity index have been used to detect LTGs in eukaryotes, though some have argued that these methods are best as a starting place for selecting candidate genes before proceeding with more detailed analyses. The finding of numerous spurious LGTs in the human genome is a notable example of problems with relying too heavily on approaches such as these [22–27].

In many situations, detecting deviations from eukaryotic monophyly (i.e. a subset of eukaryotes appear to fall nested among prokaryotes, though other eukaryotes possess the gene) is challenging in that it requires estimating single gene trees at large time scales, and such topologies are often also consistent with other evolutionary scenarios such as ancient
paralogy and subsequent differential gene loss [4,28]. Furthermore, assessment of tree topologies using likelihood-ratio tests frequently fail to reject eukaryotic monophyly, demonstrating the lack of support in these single-gene trees [29]. Indeed, some LTGs identified based on aberrant gene-tree topologies have turned out to be spurious—results of contamination or lack of data from other eukaryotic lineages [23,30]. In recent years, a number of studies have developed and applied methods to combat these pitfalls, including detailed analysis of sequence divergence between putative donors and recipients relative to outgroup eukaryotes [31–36].

Here, we present an approach for detecting LGTs from prokaryotes to eukaryotes that captures the evolutionary history and functional landscape of these events among diverse lineages. We rely on PhyToL, a phylogenomic pipeline developed by our group, coupled with a taxon-rich dataset that prioritizes whole genome sequences and then transcriptomic datasets to encompass the diversity of eukaryotes [16,37]. Then, rather than attempt to interpret the topology of gene trees that contain disparate groups of eukaryotes, we focus on gene families (GFs) present only in prokaryotes and a specific “recipient” eukaryotic group to the exclusion of other eukaryotes (i.e. exclusive presence). We identify candidate LTGs only after intensive data curation (i.e. assessment of contamination in transcriptomic data, and the evaluation of individual scaffolds of whole genome taxa) and thorough exploration of alternative hypotheses, and show the efficacy of using rigorous criteria for identifying candidate LTGs.

In total, we analyzed whole genome and transcriptome data from 1,531 species (genomic data from 688 bacteria, 114 archaea, and 189 eukaryotes, plus an additional 540 eukaryotes with transcriptomic data) accessed from GenBank or generated by our lab (S1 Table). We mitigated contamination by removing low quality and highly contaminated transcriptomes, as well as sequences with aberrant composition as compared to a set of conserved gene families (see methods). For taxa with genome sequence data, we further curated data by mapping sequences of interest to genomic scaffolds and analyzing the nearby protein-coding regions (CDS), only accepting LTGs located on scaffolds longer than 10 kb for which we could identify nearby CDSs with BLAST hits to closely-related eukaryotes. To mitigate the possibility of scaffolding errors due to incorporation of bacterial contamination (especially in genomes assembled from short-read data), we carefully analyzed coding regions in the vicinity of each gene of interest, giving special manual attention (e.g. by extensive sequence similarity searching against prokaryote genomes) to regions with several nearby putative LTGs (S9 and S10 Tables). Through a combined analysis of Gene Ontology [38] (GO) terms and PFam [39] domains, functional characterization of the LTGs revealed trends in the functional distributions both within and between recipient categories.

**Results and discussion**

**Analysis of interdomain LGTs in eukaryotes**

We identified 306 gene transfer events from 295 GFs into a variety of eukaryotic clades (S2 and S3 Tables). Many of these are instances of putative endosymbiotic gene transfer (EGT), which we consider separately from other LGT events. Using the phylogenomic pipeline PhyToL [40], the initial selection of GFs based on the presence of potential recipients yielded over 1,700 candidate LTGs (Fig 1A). We then generated multi-sequence alignments for these GFs using Guidance v. 2.02 [41], a tool that allows rigorous homology assessment, and then constructed gene trees using RaxML v. 8.0 [42] as incorporated into PhyToL (Fig 1B). After extensive data curation, including visual inspection of all alignments and trees, we retained only LTGs for which all or nearly all of the eukaryotes belong to a targeted recipient clade or group (Fig 1C). This approach is conservative in that it will exclude ancient events, cases...
where a vertically transmitted gene was lost and then re-acquired from a prokaryote, and most cases of intradomain (i.e. eukaryote-to-eukaryote) transfer (e.g. [43]).

We find certain prokaryotic groups overrepresented in distinct categories of LTGs, though discerning donor lineages in individual LGT events is confounded by a variety of factors including rampant gene transfer among prokaryotes [3,29,44–46]. The case of Cyanobacteria as the endosymbiont ancestor in EGT trees is the most prominent (Figs 1E and S4). Our approach selected against EGTs from the mitochondrial genome as these transfer events tend to be very ancient [47] and therefore generally do not show exclusive presence. Regardless, we do not see a disproportionate amount of Alpha-proteobacteria (the mitochondrial endosymbiotic ancestor) relative to other bacterial groups in putative LTG trees (Figs 1E and S4). In contrast, a recent study found enriched proteobacterial presence in putative LTGs to phytoplankton, which was explained by the ecological similarities of the putative donor and recipient lineages [48].

While differential gene loss remains a valid explanation for how exclusive presence can arise [4,28], this would lead to the expectation of relatively large divergence between eukaryotes and prokaryotes in LTG trees. To assess whether LTGs were more recently in prokaryotic
ancestors than vertically-transmitted genes (VTGs; Fig 1D), we conducted a test based on branch-lengths to identify systematic biases, similar to analyses that have been proposed and/or conducted in recent studies of LGT, though these studies were conducted without the explicit intent of comparing to simulated instances of gene loss [31,33,49]. We first selected a group of putative VTGs based on their presence in all five major clades of eukaryotes (see methods). Next, we mimicked gene loss by subsampling these VTG trees to match the taxonomic distribution of both eukaryotes and prokaryotes for a given LTG. We then generated an alignment and gene tree for each subsampled VTG to compare the ratios of the average branch length within the eukaryotic clades to the distance between the eukaryotic clade and the last common ancestor of the prokaryotes.

Using this simple branch length comparison method, we found that the distance between the eukaryotes and prokaryotes in most LGTs was shorter than for the corresponding subsampled VTG, with the distance commonly falling below the first quartile of their corresponding VTG distribution (Figs 1D, 2C, 3E and 4C, and S11 and S12 Tables). Correct interpretation of the results is contingent on the homogeneity of substitution rates before and after transfer, and it is possible that some of the cases in which our LGTs match VGTs that mimic loss are due to elevated rates of evolution immediately following gene transfer. The low relative branch length ratios of some putative LGTs are driven by both a decreased distance between eukaryotes and prokaryotes (consistent with a hypothesis of LGT) and an increase in branch length among recipient eukaryotes (S1–S3 Figs), consistent with an accelerated rate of evolution post-transfer. These patterns are also consistent with elevated rates of evolution following gene loss in all but the remaining clade. Additionally, it is possible that some of these genes were transferred from eukaryotes into prokaryotes, but this is unlikely due to our selection criteria of exclusive presence; in our candidate LTGs, the relative diversity of prokaryotes is almost always much greater than that of eukaryotes.

We identified putative LTGs in either monophyletic eukaryotic groups or into groups of organisms that share emergent functional properties (i.e. photosynthetic or anaerobic lineages). In total, we found 189 putative EGTs, 52 LGT events in Opisthokonta, 19 in Amoebozoa, 16 within Excavata, and 17 among SAR (Stramenopila, Alveolata, and Rhizaria); we also identified 14 LTGs unique to anaerobic eukaryotes belonging to two or more major eukaryotic groups, which we hypothesize may involve intra-domain transfer. We created a separate pipeline to assess the potential function of relatively ancient (and, by our methods of detection, widely retained) LTGs in eukaryotes. Numerous studies have explored the functions of individual putative LTGs on a case-by-case basis. For example, an analysis of the genome of the choanoflagellate Monosiga brevicollis revealed hundreds of putative LTGs, the majority of which are involved in carbohydrate and amino acid metabolism or stress responses [50], and another study of bacterial transfer into Ochrophyta emphasized the possible role of LGT in the evolution of secondary metabolic pathways in eukaryotes [51]. Our methods of LTG identification and curation allowed us analyze the high-level functional trends of LTGs in each taxonomic group; below we discuss these cases in three sections: 1) EGTs; 2) LGTs into non-photosynthetic eukaryotes; and 3) the special case of LGT into and between anaerobic eukaryotes.

**EGTs into photosynthetic eukaryotes**

Endosymbiotic gene transfer is distinct from other forms of non-vertical gene transfer in that its mechanism is relatively well understood, and it results from transfer of genes from a permanent intracellular symbiont [5,6]. As transfer of genes from endosymbionts to the nucleus is well-documented in photosynthetic eukaryotes [4–6,52–55], we use EGTs as a control for our
assessment of other interdomain gene transfers and as a pilot for our deployment of a functional analysis pipeline. We characterize gene families as EGTs when they are exclusively or nearly-exclusively found in prokaryotes and photosynthetic eukaryotes. Photosynthetic lineages in our pipeline include Archaeplastida (green algae, red algae, glaucocystophytes) whose ancestor acquired a plastid from a cyanobacterium [56], and clades that acquired plastids from other eukaryotes through secondary or tertiary endosymbiosis (eg. many stramenopiles, dinoflagellates, haptophytes, and cryptophytes) [52,57]. We consider genes present in only these clades as EGTs even if some lineages are thought to have lost the ability to photosynthesize, as there is a strong possibility that genes were transferred into the nuclear genome before ancestral plastid loss. Another alternative is that some of these events are transfers from the nucleus (or nucleomorph) of a photosynthetic eukaryote, with only some of these genes being originally of cyanobacterial origin.

Given the complex evolution of genes involved in photosynthesis, for which numerous hypotheses have been proposed [57,58], we recognize that our simplistic approach may categorize some LGTs as EGTs; any genes transferred laterally from a free-living prokaryote (i.e. not an endosymbiont) into a photosynthetic lineage will be incorrectly categorized by our study.

---

**Fig 2. The taxonomic and functional distributions of putative EGTs.** (A) The distribution of 189 GFs, categorized by putative function, subject to transfer events (columns) across all photosynthetic eukaryotes included in the study (Dino: Dinoflagellates; Rhiz: Chlorarachniophytes (Rhizaria); Hapto: Haptophytes; Crypto: Cryptophytes; Strepto: Streptophytes; Rhodo: Rhodophytes; Gla: Glaucophytes). (B) Functional categories based on the co-occurrence of Gene Ontology terms designate 52 EGTs as plastid-related (dark green, lefthand panel), 10 as metabolic, catabolic or biosynthetic (medium green, central panel), and 61 as other organelar or transport related (light green, right panel). (C) In comparisons of relative branch lengths, putative EGTs into Archaeplastida (red dots) consistently fall below the relative distance between eukaryotes and prokaryotes in VTG trees (box plots), inconsistent with the alternative hypothesis of gene loss and consistent with the results of similar analyses in other, non-photosynthetic taxonomic groups such as Opisthokonta (Fig 3).

https://doi.org/10.1371/journal.pgen.1010239.g002
While this is a possibility, it would be difficult to confidently rule out EGT partly because frequent LGT and gene loss among prokaryotes can obscure inferences on possible donors. As EGTs appear to be more common and stable than other forms of non-vertical transfer, we chose EGT as the most parsimonious hypothesis. Another scenario that we consider EGT is if a gene were to be transferred from a free-living bacterium into a eukaryote that later became an endosymbiont, and in turn transferred from the degrading nucleomorph into the recipient genome.

As expected, more Cyanobacteria appear in EGT trees than in any other category of LTGs (Figs 1E and S4), consistent with the cyanobacterial ancestry of the plastid [56,59]. We also find topologies consistent with the secondary and tertiary transfers of plastids, as lineages of photosynthetic eukaryotes frequently nest among archaeplastida; our trees also include lineages that have likely lost their plastid but retained genes of plastid-origin in their nuclear genome (e.g. Apicomplexa and Perkinsozoa, which appear in many of our EGT trees; S6 Fig) [52,58,60–62].

Deploying a pipeline that automates assessment of function through analysis of GO terms and PFam domains (see methods), we find that EGTs are frequently assigned plastid-related functions (Fig 2), consistent with previous literature [4,5,63]. Most of the EGTs exclusive to photosynthetic eukaryotes that acquired their plastid secondarily were assigned metabolic, catabolic or other biosynthetic functions (Fig 2). These candidate EGTs have elevated levels of Alpha-proteobacterial presence relative to other EGTs and are less likely to have a plastid-related function than trees containing Archaeplastida (Fig 2), consistent with plastid-related genes of non-cyanobacterial origin, a phenomenon which has been documented [57,58,60,64].

Our approach to the functional analysis of EGTs contrasts with the more ‘piecemeal’ approach of previous studies in which researchers focus on exploration of one or a few candidate genes or lineages [53,65–69]. Also, many studies focus either on functional differences between genes retained in the plastid as compared to those that are transferred to the nucleus, or on EGTs in lineages that have subsequently lost the ability to photosynthesize [55,70]. When the more general functional distribution of putative EGTs has been thoroughly analyzed and compared to other groups of gene families, it has been with a relatively small number of photosynthetic eukaryotes [4]. In contrast, we use EGTs to exemplify a functional analysis pipeline that, combined with intensive manual data curation, takes full advantage of our taxon-rich dataset to make inferences on the function of genes in the recipients of EGT (Fig 2A and 2B).

We also assess the alternative hypothesis of gene loss as an explanation for the 12 putative ETGs that are only present in Archaeplastida and bacteria (and sometimes only cyanobacteria), and that have taxonomic distributions that match (subsampled) conserved gene trees. Notably, the relative distance between eukaryotes and prokaryotes was below the first quartile of that of the VTGs in 8 out of 12 cases, and above the median in only two (Fig 2C), a trend inconsistent with gene loss and consistent with EGT. Moreover, the trends in the distance between eukaryotes and prokaryotes in these EGTs, the average branch length within eukaryotes, and their ratio, are similar to those of LTGs found exclusively in non-photosynthetic eukaryotes (S1–S3 Figs).

**LTGs unique to non-photosynthetic eukaryotes**

Using the requirement of exclusive presence, we identify a total of 52 interdomain LGT events into Opisthokonta: four into a common ancestor of all Opisthokonta, 41 specific to fungi, and seven specific to metazoa (Fig 2A). Within fungi, we found 28 LGTs specific to Dikarya, with 9 transfers unique to ascomycetes and no transfers solely into Basidiomycetes despite the presence of 10 Basidiomycetes with completed genomes in our database. While Chytridiomycetes
and Mucoromycetes are recipients of several LGTs, we found no LTGs retained in the genomes of microsporidia (a causative agent of wasting diarrhea in patients with AIDS), which likely reflects both elevated rates of gene loss in these highly-streamlined genomes [68,71] as well as our limited sampling of the group and focus on ‘older’ events, as LGTs have previously been documented in this lineage [72]. We did discover several putatively recent transfer events unique to single fungal species when assessing candidate LTG into other lineages. For example, we detect a transfer of an EPSP synthase (OG5_131267) gene from Proteobacteria into Aspergillus oryzae in a gene tree that we first identified for the presence of green algae nested among Cyanobacteria. As with other such cases, our curation mapped this gene to a robust genomic scaffold, found no other BLAST hit to eukaryotes, and rejected the monophyly of the eukaryotes on this tree by AU test, indicating multiple putative transfer events for this gene family (S2 Table).

The majority of the GFs putatively transferred into Opisthokonta have functions related to either: 1) intracellular transport, structure, and communication; or 2) metabolic activity (Fig 3C and 3D). Though the GFs belonging to the first category are found in both metazoa and fungi, LTGs involved in metabolic and biosynthetic function are unique to fungi. Additionally, four of the five ‘intracellular transport’ LTGs transferred into the last common ancestor of fungi contain the major facilitator superfamily (MFS_1) domain (Fig 3A), expanding on Fig 3. A summary of 52 interdomain LGT events in Opisthokonta, most of which are unique to fungi. (A) The presence of GFs (columns) in opisthokonta species, categorized by function (colors). Nodes with fewer than five and five or more inferred events are represented by open and closed circles, respectively. The presence of the MFS_1 domain is indicated by triangles. (B) Gene families categorized by function using Gene Ontology terms fall into two major categories: intracellular transport, structure and communication (dark blue) and metabolic activity (light blue). (C) Exemplar trees showing LGT into fungi and (D) Metazoa with bootstrap values greater than 50% denoted by the gray circles. Blue boxes indicate all eukaryotes (Opisthokonta) in the tree. (E) In relative branch length comparisons, Opisthokonta LTGs (red dots) consistently fall below the first quartile of the relative distance between eukaryotes and prokaryotes in VTG trees (box plots), inconsistent with the alternative hypothesis of gene loss.

https://doi.org/10.1371/journal.pgen.1010239.g003
previous work suggesting that some MFS subfamilies originated from a limited number of transfer events into fungi [73].

Across the 39 GFs putatively transferred into Opisthokonta that met our criteria for branch-length comparison, only six are consistent with a scenario of gene loss: one LTG lies above the median of its corresponding VTG distribution, and five fall above the first quartile. These six LTGs that fall above the first quartile represent either cases of gene loss that give rise to trees with exclusive presence or cases in which changes in functional constraints altered relative branch lengths (i.e. a rapid period of protein evolution as a transferred gene is first incorporated into the recipient genome). For the remaining 33 GFs that lie below the first quartile, 12 fall entirely outside of the estimates for vertical trees that mimic gene loss (Fig 2E). This pattern, which is consistent with LGT and not gene loss, is as or more pronounced as in cases of putative EGT (Fig 2C).

The presence of Actinobacteria is substantially greater in GFs putatively transferred into fungi (Figs 1E and S4), consistent with literature proposing early transfer events from Actinobacteria into the ancestor of fungi [74]. The striking similarities between fungi and Actinobacteria, from morphology to shared environment, have been linked to the potentially important role of ecology in determining patterns of transfer [75,76]. The potential role of shared ecology is further demonstrated by evidence for gene transfer between fungi and oomycetes [77]; though we focus on interdomain events, we recover a putative decarboxylase with a gene tree topology and presence/absence pattern consistent with lateral transfer from bacteria to fungi followed by transfer from fungi to Phytophthora ramorum, the sudden oak death pathogen.

Lateral transfer has been extensively studied in Opisthokonta, especially in fungi, where intimate symbiotic relationships (i.e. in lichens and mycorrhizal species) that often involve prokaryotes have predisposed some lineages to higher rates of LGT [35,78–82]. In fact, interdomain LGT has been implicated in important fungal innovations from gravity-sensing organs to pathogenic mechanisms and toxin-encoding genes [34,35,83,84]. In addition, some fungal species and genera seem to be especially prone to receiving LGTs, including the genera Aspergillus and Fusarium, for which genomic data are available [35,85] and in which we also observe frequently in putative LGT trees. LGT into other opisthokonts has been a point of interest to many, especially since the early spurious claims of hundreds of LTGs in the human genome [25,26], but specific mechanisms underlying individual transfers remain unknown. In fungi, in vitro experiments have been able to introduce foreign genetic material from prokaryotes into a variety of fungi [86]. Proposed mechanisms for LGT in vivo include transfer mediated by mobile genetic elements (e.g. viruses or transposable elements) and/or during long-term associations with parasites or symbionts [15,87].

In addition to Opisthokonta, we identified LTGs specific to one of three other major eukaryotic clades: 19 to Amoebozoa, 16 within Excavata, and 17 to SAR (Stramenopila, Alveolata, and Rhizaria). Many of the LTGs in the former two groups are found in anaerobic obligate parasites (e.g. Entamoeba (Amoebozoa), Trichomonas and Giardia (Excavata); S3 Table), which likely reflects both bias in the available data and also possibly gene transfers that support a transition to a strictly anaerobic lifestyle (see "LGT into anaerobic eukaryotes" below). Within Amoebozoa, we recover multiple putative LTGs in the slime mold Dictyostelium, such as a previously reported [88] siderophore transport-related protein (important in iron scavenging), plus other undocumented transfers into Discosea, Mycetozoa, and Tubulinea (S2 Table). The LTGs exclusive to members of SAR are found primarily in oomycetes and other stramenopiles (N = 11), apicomplexans (N = 4), and ciliates (N = 3). Apicomplexans such as Plasmodium also occasionally appear sister to photosynthetic eukaryotes in EGT trees, consistent with the presence of the plastid-derived apicoplast in these lineages (S6 Fig). Though our findings are generally consistent with the numerous studies that report interdomain LTGs into
the predominantly microbial lineages of Excavata, Amoebozoa, and SAR, our stringent methods contrast with analyses that focus on specific taxa and/or rely on topologies of single gene trees [22,70,77–79].

LGT into anaerobic eukaryotes

Multiple transitions from aerobic to strictly anaerobic/microaerophilic life strategies required eukaryotes to alter fundamental features of their metabolism, and previous analyses suggest that this involved the acquisition of genes from prokaryotes [10,12,89–93]. Such transfers are analogous to the emergence of photosynthesis in eukaryotes after the acquisition of the plastid through endosymbiosis, which led to a combination of inter- and intradomain EGTs. Endosymbioses and other intimate relationships among anaerobic lineages may also provide opportunities for transfers; consistent with the hypothesis is the presence of the anaerobic endosymbiont *Perkinsella* (Excavata) within the amoebozoan *Paramoeba* [94].

Given that the last eukaryotic common ancestor (LECA) likely contained pathways for both aerobic and anaerobic metabolism [95,96], interpreting the evolutionary history of ‘anaerobic’ genes must be done with caution. Indeed, some past findings of putative LTGs specific to anaerobes have been revised with additional data from homologous pathways in aerobic lineages [95]. For example, the sparse distributions among paraphyletic eukaryotes of genes

Fig 4. Using strict exclusive-presence criteria and extensive curation, we found evidence for both inter- and intradomain transfer involving anaerobic eukaryotes. (A) A glycy1-radical enzyme-activating enzyme (GRE-AE) is found exclusively in anaerobic eukaryotes (colored boxes) and prokaryotes. (B) Curation of genomic sequences (green dots in Fig 3A) kept only those that mapped to scaffolds longer than 10 kb and with robust nearby CDS as represented by the three scaffold sections: LTGs are shown in color and the presence/absence of nearby GFs in other eukaryotic and prokaryotic groups are shown by filled and empty squares, respectively (Ba: Bacteria; Arc: Archaea; Orphan: Eukaryotic orphan lineages; Ex: Excavata; Plt: Archaeplastida; Am: Amoebozoa; Op: Opisthokonta); sliding window GC content shows amelioration of these LTGs; CDSs (black boxes) with no presence/absence data are not in the PhyloToL database (e.g. lineage specific genes). (C) In most branch length comparisons, LTGs (red dots) fell below the first quartile of the branch-length ratios of the corresponding set of conserved genes (box plots) subsampled to mimic gene loss. This includes the distance between SAR and Excavata in the GRE-AE gene tree (A); the number of VTG trees is given for LTGs with fewer than 20 VTGs in their corresponding distribution.
involved in the anaerobic conversion of pyruvate to Acetyl-CoA, such as pyruvate-formate lyase (PFL), its activating enzyme (PFL-AE), and pyruvate:ferredoxin oxidoreductase (PFO) have led to the suggestion of interdomain followed by intradomain transfer [10,89,97]. In contrast to this hypothesis, the presence of these genes in various aerobic lineages (e.g. green algae) suggests that they were present in LECA and underwent extensive differential loss [95].

We identified 42 GFs found exclusively or nearly exclusively in eukaryotes sharing anaerobic functional and/or ecological contexts (e.g. Archamoeba, Parabasalids, Trypanosomatids, and Apicomplexa); in many of these cases, the eukaryotes are sister to anaerobic and/or pathogenic bacteria (S3 Table). Several previously reported anaerobic LTGs did not pass our conservative criteria due to their widespread presence in other eukaryotes, including PFL, PFL-AE, and PFO [10,89,97,98]. However, we did recover a glycyl-radical enzyme-activating enzyme (GRE-AE) highly similar to PFL-AE that was previously documented in Giardia [12], and later Entamoeba and Trichomonas [99]; we additionally find it in other Archamoebae and anaerobic members of SAR (Fig 4A). We expand the taxonomic scope of other previously-reported GFs, such as the alcohol dehydrogenase EhADH3B originally reported as an LTG unique to the human parasite Entamoeba histolytica [100] where it is associated with pathogenicity [101]; we recovered this GF in other Archamoebae, Trichomonas, and Blastocystis. Similarly, we find the nitroreductase Fd-NR2 in Entamoeba, expanding its presence from a previous report as an LTG only in Giardia [102] (S3 Table).

To further assess candidate anaerobe LTGs, we carefully curated each sequence by mapping it to the genome to account for contamination (Fig 4B and S9 and S10 Tables) and tested the alternative hypothesis of gene loss using relative branch length comparisons (Fig 4C). We only retained sequences that mapped to scaffolds longer than 10 kb and with clearly eukaryotic coding domains on the same scaffold as the LTG (Fig 4C). We excluded from our study several putative LTGs in the excavate Trimastix marina and in the breviates Pygsua biforma and Leniisia limosa, as these transcriptomes contained evidence of contamination by other eukaryotes in our analyses of control groups of genes conserved across eukaryotes (S9 and S10 Tables). Of the 14 interdomain LGTs that met our criteria for relative branch-length comparison, the majority lay below the first quartile and four are above the median (i.e. consistent with gene loss; Fig 4C), inconsistent with gene loss as an explanation for the pattern of inheritance of most of these putative LTGs. Fourteen anaerobe-specific LTGs are exclusive to paraphyletic eukaryotic lineages (i.e. contain anaerobic Amoebozoa, Excavata and/or SAR). Here we used the branch-length comparison approach to assess the seven cases of putative intradomain transfer that had large enough corresponding distributions of VTGs: three LTGs lay below the first quartile of the corresponding conserved-gene distribution and two lay above the median (Fig 3C), a signal consistent with LGT but with less support than the interdomain cases.

**Synthesis**

Non-vertical inheritance of genetic material clearly confounds attempts to reconstruct the tree of life [3,103–106]. However, the extent to which this is a problem is difficult to discern with such varying standards as those currently used for discovering LGT, many of which leave differential gene loss as a possible alternative explanation [4,23,26,95]. Here we present rigorous and conservative methodology for identifying interdomain LGT events in eukaryotes: we require exclusive presence of GFs in taxon-rich analyses and apply multiple rounds of curation including analyzing compositional bias, mapping genomic cases to robust scaffolds, and carefully interpreting transcriptomes that often include sequences from contaminants (Fig 1A–1D). We identified 306 LGT events (Fig 1E) and then assessed the alternative hypothesis of differential gene loss to find that our candidate LTGs are consistently outliers (Figs 2C, 3E and...
4C), though this varies between recipient groups. Using a definition based on exclusive presence and testing these hypotheses by branch-length ratio comparisons provides an important step towards clarifying criteria for robustly identifying LGT events and thus determining how pervasive LGT is as a phenomenon in eukaryotes.

Methods

Taxon selection

Analyses here rely on PhyloToL [40], which includes 540 eukaryotic species, 688 bacteria and 114 archaea, and represents a combination of whole genome taxa (all prokaryotes and 189 eukaryotes) plus numerous lineages represented by transcriptomes (i.e. ‘transcriptomic taxa’; Tables 1 and S1). Our intention in sampling was to create a relatively even set of taxa from across the eukaryotic tree of life, given data availability; to this end, we have undersampled plants, animals and fungi. Data are largely from GenBank (S1 Table), representing mostly lineages that can be cultivated. We also included 111 single cell transcriptomes from diverse microeukaryotes characterized in our lab that survived the rigorous data curation described below (S9 and S10 Tables). As part of data management, all taxa/cells are named with ten-digit codes that represent their major clade (e.g. Op = Opisthokonta), “minor” clade (e.g. Op_me = metazoa) and species (Op_me_Hsap = Homo sapiens). Though controversial, we include the major clade Excavata (Ex_), excepting the genus Malawimonas, and we note that the LGTs involving Excavata all include only a subset of the clade (i.e. do not rely on the monophyly of the group).

Initial curation of transcriptomic taxa

We evaluated all transcriptomic data prior to analysis of candidate LTG multi-sequence alignments (MSAs) and trees using PhyloToL [40] to assign transcripts to gene families. To mitigate both quality and contamination issues frequent with transcriptome data from microeukaryotes, we assessed the number of transcripts initially assigned to gene families (GFs) and the proportion of transcripts determined to be likely bacterial (i.e. contamination or potential LGTs), defined by PhyloToL as those that return a top BLAST hit to bacteria with an e-value at least 10^3 times less than that of the top eukaryotic hit. We excluded samples with high ratios of putative contaminating bacterial sequences to non-bacterial sequences, and samples likely contaminated by food sources; this approach removed a total of 55 transcriptomes. We took a similar approach to identify eukaryotic transcriptomes contaminated by other eukaryotes based on a pilot analysis of 35 gene trees constructed from highly conserved gene families (i.e. those present in a large number of species across all seven major clades). We identified 253 transcriptomic samples (from our lab and GenBank) that either failed to appear in these trees (i.e. experimental failure) or that had high levels of contamination (e.g. by food source, commonly co-cultured lineages such as Bodo) assessed using our knowledge of the organisms’ ecology and data source. In a second round of taxon curation, we removed additional taxa and sequences (i.e. contaminants and/or sequences from food sources) based on their performance in sets of conserved genes (i.e. present across diverse eukaryotic lineages), including the 408 MSAs generated to explore compositional bias and codon usage (see below).

Candidate gene-family selection

We started with 13,630 gene families (GFs) as defined by OrthoMCL [107] release 5.0 and incorporated them into PhyloToL, based on their presence in diverse eukaryotes (Table 1). For
transcriptomes, PhyloToL translates nucleotide sequences ≥200bp after determining the appropriate genetic code and assigns amino acid sequences to GFs based on similarity using USEARCH [108] with a maximum e-value of 1e-10, and then combines these with sequences from OrthoMCL. We identified potential LGTs in predefined sets of potential recipient groups (Opisthokonta, Amoebozoa, Excavata, SAR, photosynthetic eukaryotes, and anaerobic eukaryotes) based on two relatively lenient criteria: 1) GFs where the proportion of the eukaryotes of the potential recipient clade were greater than the mean plus the standard deviation across all 13,630 GFs, and 2) the number of prokaryotes was greater than the 65th percentile across all GFs (Fig 1A). For cases where we had large numbers of single-cell transcriptome sequences (e.g. ciliates, Arcellinida), we applied a similar approach to sequences that PhyloToL determined to be possible bacterial contaminants (i.e. they had a BLASTx hit to a bacterial sequence in the OrthoMCL database that was at least 10^3 times less than its best BLASTx hit to a eukaryote); here we chose GFs for which the proportion of taxa in a potential recipient clade fell significantly above the general distribution (identified using Mahalanobis distance; p-value < .001), with a minimum of at least 67% of the eukaryotes belonging to the recipient clade.

**Refinement of putative EGT/LGT gene families**

We assessed homology for all candidate LTGs using up to five iterations of Guidance [41] version 2.02, a tool that removes sequences below user-specified thresholds (in our case, seqCut-off = 0.3 and colCutoff = 0.4) after multisequence alignment (MSA) reconstruction with MAFFT [109]. We masked gaps at 95% using trimAl [110] version 1.2 and constructed preliminary gene trees using RAxML [42] version 8.0 as implemented in PhyloToL (model PROTGAMMALG). We curated the resulting gene trees to focus on those that had nearly-exclusive presence of putative recipient clades (i.e. we allowed singletons or small second clades only if they survived our strict curation as described below) and then we went on to use several rounds of sequence curation described in the “Gene tree curation” section below. In other words, the gene trees were a tool for discovery of transferred gene families and we did not rely on single gene tree topologies as we finalized our list of LGTs.

We also curated all EGTs to avoid inclusion of plastid sequences present in transcriptomes. Using custom Python and R scripts, we compared putative EGT sequences against a set of 408 highly-conserved nuclear-encoded gene families (i.e. present in all five eukaryotic major clades) by plotting G+C content at third-position four-fold degenerate sites (GC3) against the effective number of codons (ENC) [111]. We also ran a correspondence analysis of the relative synonymous codon usage (RSCU) to distinguish nuclear and plastid genes. These analyses combined led to the removal of 29 putatively-plastid sequences that were significantly outlying (Mahalanobis distance; p-value < .001) in either of these plots (S5 Fig).

**Sequence-level curation of candidate LGTs**

In addition to initial curation of taxa, we refined candidate LTGs to meet a set of conservative criteria. Because of bacterial by-catch in eukaryotic transcriptome studies, we removed clades of transcriptomic sequences with insufficient taxonomic representation. These clades are defined as containing only sequences designated by PhyloToL as likely bacterial (see above) that contained either <4 species or <33% of the single-cell transcriptomes generated in our lab for a given taxon. We also curated every sequence in gene trees that fell into any of four categories: 1) sequences sister to one or no other closely-related eukaryotes (e.g. an Amoebozoa sequence in an otherwise all fungal clade); 2) non-recipient eukaryotic sequences in trees with at least five genera of recipient eukaryotes; 3) sequences in trees with fewer than five genera of recipient eukaryotes; and 4) sequences from trees containing only anaerobes that are from
multiple major clades of eukaryotes. The resulting collection of transcriptome and genome sequences were compared against the “nr” database using the “qblast” function in the Biopython library[112], and only sequences with robust hits to eukaryotes in the “nr” database (i.e. to taxa not included in PhyloToL) were retained (S10 Table).

We further inspected all transcriptome sequences that fell into any of the four categories and removed those that lay outside either the distribution of GC3s plotted against the ENc, or a correspondence analysis plot of the RSCU of sequences from 408 conserved gene trees (Mahalanobis distance; p-value < .001). We kept sequences that robustly hit multiple closely-related species in the nr database, and removed those that did not. We also removed some clades of very closely-related samples for which we had large numbers of transcriptomes (e.g. multiple individuals of the genus *Hyalosphenia*) if they did not hit other closely-related species and if the inferred LGT tree had a topology within eukaryotes inconsistent with vertical inheritance (i.e. interdigitation of species between genera; S10 Table). In other words, we took considerable care to rule out contamination and misidentification as we made inferences about LGTs from taxa represented only by transcriptomic data.

We took a separate approach for the curation of sequences from taxa with a whole genome assembly available (S9 Table) by evaluating whether these sequences mapped to genuine eukaryotic contigs or instead represented contaminating bacterial sequences (S9 Table). We removed all sequences that mapped to contigs shorter than 10 kb, as well as those on contigs containing no other annotated protein-coding regions (CDS). For genomic sequences on contigs longer than 10 kb that hit no closely-related eukaryotes by BLASTp against the “nr” database, we analyzed nearby CDSs and CDSs at the ends of the contig using BLASTp; here our goal was to look for bacterial material in these assemblies. If very few of these robustly hit closely-related eukaryotes and/or many robustly hit only prokaryotes, we removed the sequences. Aberrance in compositional bias was not used as a criterion in determining robustness of genomic sequences.

### Additional gene-tree curation

As a final curation step, we looked for cases in which GF designations in OrthoMCL split homologs into multiple gene trees, leading to an overestimation of LGTs. To this end, we choose representative sequences from each candidate LTG tree (i.e. sequences representing clusters of $\geq$75% identity generated using the VSEARCH [113]—cluster_smallmem command-line tool) and used the BLASTp tool in the BLAST+ executables package [114] to identify homologs among our original 13,630 GFs. We combined GFs for all sequences of candidate LTGs that hit an alternative gene family(s) with an e-value of an order of magnitude
at least half of that of its best hit, and assessed homology using Guidance as described above. We manually inspected gene trees generated for all resulting alignments and removed eight candidate LGTs in which “recipient” eukaryotes interdigitated with other eukaryotes introduced by the alternative GF. We combined an additional 17 candidate LTGs either with other LTGs or with GFs not initially selected by our methods. Twenty-five gene families combined with alternative gene trees exhibited near-reciprocal monophyly of GFs (i.e. represented ancient gene duplication), and therefore were not changed except for the removal of contaminating sequences that interdigitated among other eukaryotes in the alternative gene family.

We evaluated the few candidate LTG trees with polyphyletic eukaryotes by AU testing, constraining topologies to have eukaryotic monophyly; putative EGTs were not tested except for those containing non-photosynthetic taxa not sister to the putative recipient clade. Constraint-tree construction and AU testing was conducted using IQ-Tree [115] through the CIPRES Science Gateway [116] REST API. The final trees available in the supplementary materials were constructed using IQ-Tree version 2.1.2 through the CIPRES Science Gateway REST API using the LG model, gamma site rate distribution (-m LG+G) and 1,000 ultrafast bootstraps (-bb 1000), and we include the most likely constrained trees for cases in which eukaryotic monophyly was accepted.

Functional analysis
To assess the function of LGTs, we analyzed Gene Ontology [38] (GO) terms returned for each sequence using the EggNOG-mapper tool [117], implementing the Diamond model [118] under default parameters. We obtained additional GO terms using the InterPro2GO online database [119], accessed in April 2021, and PFam domains [39] identified by HMMer [120] for each sequence with a maximum domain overlap of 5 amino acids and an e-value of 1x10^{-5}. GO terms were slimmed using the generic GO-Slim database as accessed on the GO website (http://current.geneontology.org/ontology/subsets/goslim_generic.obo). We mapped functional descriptions onto single gene trees and manually evaluated the results as finalized parameters. We summarized functional categories (Fig 3A and S6–S8 Tables) by analyzing the overlap of GO term presence in GFs in each recipient category using UpSet plots (Fig 3B; created using the UpSetR package [121]; each combination of GO terms was manually assigned to each broad category) and custom Python scripts.

Branch length ratio calculation & comparison
To assess the alternative hypothesis that exclusive presence in LTG trees is due to differential loss of genes that were present in the last eukaryotic common ancestor, we compared relative branch lengths between putative recipient and donor lineages in LTG trees to those in a corresponding set of highly conserved gene families sampled to mimic gene loss. The basis of this assessment is that the divergence between eukaryotes and prokaryotes in trees that mimic gene loss should be much greater compared to LTG trees, even in comparisons of highly conserved genes (i.e. our assessment is conservative). Though comparing the ratio of branch lengths within eukaryotic recipients to the last common ancestor with prokaryotes accounts for varying functional constraint between GFs, the analysis is based on the assumption of homogeneous substitution rates (i.e. constant functional constraint) across lineages within a given tree.

For all gene trees containing a single putative interdomain transfer event, we selected a set of 20 to 50 conserved trees (i.e. present in all five eukaryotic major clades, as well as in diverse prokaryotes) that contained all the eukaryotes and prokaryotes found within the LTG tree. Branch length comparisons were not conducted for LTGs with fewer than 20 matching conserved genes, except for the three intradomain transfers analyzed and marked in Fig 4C. We
subsampled the conserved trees to match the taxonomic distribution of both eukaryotes and prokaryotes in the LTG tree, plus or minus one taxon (mimicking the gene loss necessary to explain the taxonomic distribution). In cases where there are multiple recipient clades within an LTG tree and eukaryotic monophyly could be rejected by AU testing, we either selected the eukaryotes from the largest clade (when only one recipient major clade present) or we tested both clades (when multiple major eukaryotic groups present). When paralogs were present in the eukaryotic clade in LTG trees, paralogous sequences on branches closest to the eukaryotic root were selected and all others were removed when rebuilding the tree. In VTG trees, paralogous sequences on branches furthest to the eukaryotic root were selected and all others were removed when rebuilding the tree. This served to mitigate any potential heterogeneity in substitution rates and biased towards the null hypothesis of equal branch length ratios in LTG and VTG trees exhibiting gene loss. For the seven gene families where anaerobes of two different eukaryotic major clades are sister to each other in an LGT tree, we focused only on assessment of the intradomain LGT by sampling conserved gene trees to match the taxonomic distribution of the eukaryotic clade under consideration only (i.e. no prokaryotes were included in the subsampled tree).

Examination of the average branch length within eukaryotic clades in LGT trees as compared to subsampled vertical trees revealed potential discrepancies in substitution rate, inconsistent with the assumption of homogenous substitutions (S1–S3 Figs). This measure in LTG trees frequently fell above the distribution of that in the corresponding VTG trees, hence lowering the relative distance between eukaryotes and prokaryotes. This observation holds for both LGT and EGT GFs (S1–S3 Figs), and is consistent with both differential loss of rapidly evolving genes and gene transfer, as accelerated evolutionary rates of a gene following transfer is possible. Analysis of the absolute distance between eukaryotes and prokaryotes revealed strong bias of LGT trees to lie below their corresponding VTG distributions, which is inconsistent with gene loss and consistent with LGT (S1–S3 Figs).

For all relative branch length comparisons, we used MAFFT version 7.407 to align the amino-acid sequences of all LTG and subsampled VTG trees. We then constructed single-gene trees for both VTGs and LTGs using IQ-TREE version 2.12 through the CIPRES Science Gateway REST API, using the LG model and gamma site rate distribution (-m LG+G). To calculate the relative branch lengths, each resulting tree was rooted on the eukaryotic clade (trees that mimicked gene loss where the eukaryotes were non-monophyletic were not considered) and the distance from the base of the eukaryotic clade to the base of the prokaryotes was divided by the average branch length within the recipient eukaryote clade.

In the eukaryote-only trees generated to emulate intradomain LGT among anaerobic eukaryotes, the distance between the clades of each eukaryotic major group was compared to average branch lengths within them. Each LTG tree ratio was then compared to its distribution of conserved GFs (Fig 4C and S11 and S12 Tables). Conserved gene families were not included in the control set if AU testing rejected reciprocal monophyly of the two eukaryotic groups.

**Supporting information**

**S1 Table. List of all taxa used in the study that have putative LTGs.** Columns include: major clade (Amoebozoa (Am), Archaea (Za), Archaeplastida (Pl), Bacteria (Ba), Excavata (Ex), Opisthokonta (Op), SAR (Stramenopila, Alveolata, Rhizaria), and orphan lineages (EE)); taxon code and name, the number of putative LTGs, data type (genomic, EST or Illumina), data source (i.e. the database, institution or project from which the data was accessed), the accession numbers for the data used where applicable and taxonomy as designated by NCBI. (XLSX)
S2 Table. All of the GFs inferred to be transferred into non-anaerobic organisms and their putative recipient eukaryotic clades. Gene families with two or more OG5 numbers from OrthoMCL (column A) were determined to be homologs as described in the methods section. LTGs found in only one recipient clade (column B) and that returned sufficient functional information (Gene Ontology terms; column C) are assigned broad functional categories as illustrated in Figs 2 and S1. Additional functional information is given in the form of EC numbers, Pfam IDs and PFam domain names as returned by the EggNOG mapper tool and through a search with HMMer, respectively. (XLSX)

S3 Table. All of the putative LTGs found exclusively or nearly-exclusively in anaerobic eukaryotes. The predicted function for each gene family is reported if there is consensus among the annotations of the genomic eukaryotic sequences in OrthoMCL version 5.0. We also list putative recipients and donors, the latter defined as the most coherent/broadest grouping of prokaryotes with the LTG. For each GF, we include EC numbers, Pfam IDs, and PFam domain names as returned by the EggNOG mapper tool and through a search with HMMer, respectively. (XLSX)

S4 Table. Presence/absence data for all putative LTGs in every taxon listed in S1 Table. Presence of an LTG is denoted with a “1” regardless of the number of copies of the gene found in the genome (i.e. paralogs), and absence is denoted with a “0”. (XLSX)

S5 Table. Presence/absence data for conserved GFs in every taxon listed in S1 Table. Presence of a GF is denoted with a “1” regardless of the number of copies of the gene found in the genome (i.e. paralogs), and absence is denoted with a “0”. We use these conserved GFs to assess the quality of transcriptomic data, and for subsampling to generate trees that mimic gene loss in branch-length comparisons (S11 and S12 Tables). (XLSX)

S6 Table. The presence of each Gene Ontology (GO) term as returned by EggNOG and PFam across recipient groups (All Go Terms; columns B-F), those returned just by EggNog (G-K) and by Pfam only (L-P); GO terms were slimmed using the generic GO-Slim database as accessed on the GO website (http://current.geneontology.org/ontology/subsets/goslim_generic.obo) in April of 2021 and then counted. We also give the number of GFs with eukaryotic sequences that return each GO term through either source. (XLSX)

S7 Table. The presence of all un-”slimmed” Gene Ontology (GO) terms as returned by EggNOG and PFam across all putative LTGs unique to Opisthokonta or photosynthetic eukaryotes. A “1” denotes presence regardless of the number of eukaryotic sequences that returned the GO term, and a “0” denotes absence in eukaryotes. (XLSX)

S8 Table. The presence of all Pfam domains as returned by HMMer across all putative LTGs. A “1” denotes presence regardless of the number of eukaryotic sequences that returned the domain, and a “0” denotes absence in eukaryotes. (XLSX)

S9 Table. All curated genomic sequences and the criteria used for determining their inclusion or exclusion. For each species, we provide the unique sequence identifier (column D),
the reason for inclusion or exclusion (columns H-J), the sequence length, the length of the contig on which the putative LTG is placed (column H), the accession of the genomic data, and whether the sequence hit closely-related species or multiple other CDS on the contig hit closely-related species (columns I,J). Any notable literature pertaining to the sequence or GF is noted in the “other” column.

S10 Table. All curated transcriptomic sequences and the criteria used for determining their inclusion or exclusion. For each species, we provide the unique sequence identifier (column D), the reason for inclusion or exclusion (columns G-J) and whether the sequence hit closely-related species.

S11 Table. The branch-length distributions for all putative interdomain LGTs. For each LGT (column A) we provide distance for all comparison trees (Column B) for both LGT and VGT trees (column C). All subsampled VGT measurements are given for each corresponding LTG, and the recipient category of the LTGs (and their corresponding subsampled VTGs) match those in figures (Figs 2E and 3C and S1–S3).

S12 Table. The branch-length distributions for all putative intradomain LGTs. For each LGT (column A) we provide distance for all comparison trees (Column B) for both LGT and VGT trees (column C). All subsampled VGT measurements are given for each corresponding LTG; these LTGs match those at the bottom of Fig 3C.

S1 Fig. Measurements of relative branch length in twelve EGT trees containing only archaeplastida are consistent with those in non-photosynthetic recipient groups (Fig 2C). (A) For the majority of the EGTs, the ratio of the average branch length within the putative recipient eukaryote clade (EE) to the distance between the eukaryote clade and the prokaryotes (EP; red dots) lies outside estimates of their corresponding subsampled-VTG distributions (box plots; S11 Table). Either most or all of the prokaryotes in these trees are Cyanobacteria, consistent with plastid ancestry of these GFs. (B) Average branch length within the eukaryotic clade of the same 12 LTGs (red dots, GFs in the same order) are variable relative to their corresponding subsampled-VTG distributions (box plots), consistent with variable functional constraints on these GFs following transfer. (C) For the majority of GFs, distance between the eukaryote and prokaryote clades in the same 12 LTGs (red dots) are shorter than their corresponding subsampled-VTG distributions (box plots).

S2 Fig. Measurements of relative branch length in 21 LGT trees containing only Amoebozoa (blue boxes, 7 GFs), Excavata (yellow boxes, 8 GFs) and SAR (red boxes, 6 GFs). (A) In the majority of cases, the ratio between the average branch length within the putative recipient eukaryote clade (EE) and the distance (branch length) between the eukaryote clade and the prokaryotes (EP; red dots) are outside the range of their corresponding subsampled-VTG distributions (box plots). (B) The average branch length within the eukaryotic clade of the same LGTs (in the same order, red dots) are variable compared to their corresponding subsampled-VTG distributions (box plots). (C) The distance between the eukaryotic and prokaryotic clades in the same LGTs (red dots) tend to be smaller than in the same subsampled-VTG distributions.
S3 Fig. The average branch length within eukaryotic clades and between eukaryotes and prokaryotes of LTGs in Opisthokonta following order of GFs in Fig 2C. (A) The average branch length within clades of Opisthokonta in LTG trees tends to be longer than in their corresponding subsampled-VTG distributions. (B) The distance (branch length) between the Opisthokont clade and the prokaryote clade in LTG trees tends to be shorter than in their corresponding subsampled-VTG distributions. (TIFF)

S4 Fig. The distributions of possible donor lineages in LTG trees, with the caveat that both gene loss and prokaryote-prokaryote LGT after interdomain gene transfer events can obscure inferences. For each archaeal (top) and bacterial (bottom) clade that appeared abundantly in LTG trees unique to anaerobes (left panel), photosynthetic eukaryotes (central panel), or fungi (right panel), we measured the proportion of the taxa in that clade in the PhyloToL databases that appear in each tree (X-axis). For putative transfer events into anaerobic eukaryotes, there is greater representation of Methanobacteria, Thermotogae, Fusobacteria, and the low numbers of Proteobacteria and Cyanobacteria; Cyanobacteria are overrepresented in the EGT trees, as expected for genes involved in photosynthesis; and with LTGs in fungi there are more Alpha-proteobacteria, gamma-proteobacteria and Actinobacteria. (TIFF)

S5 Fig. Examples of data curation, including assessment of contamination by plastid-encoded (a) and other sequences in transcriptomes (b). (A) GC content at third-position four-fold degenerate sites plotted against the effective number of codons (ENc) shows that the majority of sequences in the diatom Extubocellulus spinifer in EGT trees (green) match patterns of sequences from conserved gene trees (orange); significantly outlying points (red; Mahalanobis distance; \( p < .001 \)) may be plastid encoded and these GFs were removed. Inset is a violin plot of the GC content of conserved sequences. (B) The same set of sequences plotted in a correspondence analysis showing that relative synonymous codon usage is significantly different for the same GFs (red box; assessed by Mahalanobis distance; \( p < .001 \)). (C) We also assessed the composition of transcriptomic sequences that appeared in clades lacking robust taxonomic representation (red lineages in tree; in this case, the stramenopile Devalopayella elegans) and removed those with compositional patterns distinct from highly conserved GFs (orange). Points in gray belong to the highly-conserved gene families, but returned top BLAST hits to bacteria with an e-value 10^3 times lower than to eukaryotes. (TIFF)

S6 Fig. An example EGT tree. Genes subject to EGT were identified by exclusive presence in photosynthetic eukaryotes, including the Archaeplastida that acquired their plastid from a cyanobacterial ancestor and lineages that acquired plastids secondarily (e.g. photosynthetic members of SAR). Apicomplexa and Perkinsozoa occasionally appear in EGT trees, consistent with photosynthetic ancestry in these non-photosynthetic organisms. (TIFF)

S7 Fig. The distribution of prokaryotic presence in putative LGT & EGT trees. (A) the number of gene trees in which each pair of bacterial/archaeal clades co-occur. (B) the number of gene families in which each prokaryotic clade is present (bars are in the same order as in (A)). (C) The number of prokaryotic clades in each gene tree based on the data in S4 Table; the “Cyano only” label highlights trees with a single bacterial clade, all but one of which are EGTs and contain only cyanobacteria. (TIFF)
S8 Fig. A decarboxylase showing putative LGT from fungi to Phytophthora ramorum. Fungi are in purple, P. ramorum in red. All other tips are bacteria or archaea. (TIFF)

S9 Fig. Distribution of pairwise identities between prokaryotes and eukaryotes among LGTs (blue), EGTs (green) and the VGT trees that we generated to mimic gene loss (red), the latter of which were sampled from conserved gene families. Across all gene trees, we calculated the pairwise identities between each eukaryotic and prokaryotic sequence after aligning each pair of sequences separately. Few pairs exceed 70% identity (vertical dashed line), an observation consistent with the “70% rule” defined by Ku and Martin (2017), who argued that interdomain comparisons with >70% identity are likely contaminants. The right skew of the VTG distribution is consistent with the conservative nature of these gene families, which we selected based on their wide distribution among eukaryotes and prokaryotes. (TIFF)

Acknowledgments
We are grateful to Rabindra Thakur (Smith College/UMass Amherst) for contributions in launching this study, and to Jane Carlton (New York University), George Zhang (University of Michigan), Jaime Huertas-Cepas (EMBL), Carlos Cantalapiedra (EMBL), and Sussanah Porter (UCSB) for helpful conversations. We also thank Mark Miller, Wayne Pfeiffer and others at CIPRES, and Tony Caldanaro and Steve Davis at Smith College for their technological help. Finally, we thank members of the Katz lab for feedback on drafts and figures.

Author Contributions
Conceptualization: Auden Cote-L’Heureux, Xyrus X. Maurer-Alcalá, Laura A. Katz.
Data curation: Auden Cote-L’Heureux, Laura A. Katz.
Formal analysis: Auden Cote-L’Heureux, Xyrus X. Maurer-Alcalá, Laura A. Katz.
Funding acquisition: Laura A. Katz.
Investigation: Auden Cote-L’Heureux.
Methodology: Auden Cote-L’Heureux, Laura A. Katz.
Project administration: Laura A. Katz.
Resources: Laura A. Katz.
Software: Auden Cote-L’Heureux.
Supervision: Laura A. Katz.
Visualization: Auden Cote-L’Heureux.
Writing – original draft: Auden Cote-L’Heureux, Laura A. Katz.
Writing – review & editing: Auden Cote-L’Heureux, Xyrus X. Maurer-Alcalá, Laura A. Katz.

References
1. Papke RT, Koenig JE, Rodriguez-Valera F, Doolittle WF. Frequent recombination in a saltern population of Halorubrum. Science. 2004; 306: 1928–1929. https://doi.org/10.1126/science.1103289 PMID: 15591201
2. Munita JM, Arias CA. Mechanisms of Antibiotic Resistance. Microbiol Spectr. 2016; 4. https://doi.org/10.1128/microbiolspec.VMBF-0016-2015 PMID: 27227291
3. Bapteste E, O’Malley MA, Beiko RG, Ereshefsky M, Gogarten JP, Franklin-Hall L, et al. Prokaryotic evolution and the tree of life are two different things. Biol Direct. 2009; 4: 34. https://doi.org/10.1186/1745-6150-4-34 PMID: 19788731

4. Ku C, Nelson-Sathi S, Roettger M, Sousa FL, Lockhart PJ, Bryant D, et al. Endosymbiotic origin and differential loss of eukaryotic genes. Nature. 2015; 524: 427–432. https://doi.org/10.1038/nature14963 PMID: 26287458

5. Ponce-Toledo RI, López-García P, Moreira D. Horizontal and endosymbiotic gene transfer in early plastid evolution. New Phytol. 2019; 224: 618–624. https://doi.org/10.1111/nph.15965 PMID: 31135958

6. Timmis JN, Ayliffe MA, Huang CY, Martin W. Endosymbiotic gene transfer: Organelle genomes forge eukaryotic chromosomes. Nat Rev Genet. 2004; 5: 123–U16. https://doi.org/10.1038/nrg1271 PMID: 14735123

7. Martin WF. Too Much Eukaryote LGT. BioEssays News Rev Mol Cell Dev Biol. 2017;39. https://doi.org/10.1002/bies.201700115 PMID: 29068466

8. Leger MM, Eme L, Stairs CW, Roger AJ. Demystifying Eukaryote Lateral Gene Transfer (Response to Martin 2017 10.1002/bies.201700115). BioEssays News Rev Mol Cell Dev Biol. 2018; 40: e1700242. https://doi.org/10.1002/bies.201700242 PMID: 29543982

9. Andersson JO. Lateral gene transfer in eukaryotes. Cell Mol Life Sci. 2005; 62: 1182–1197. https://doi.org/10.1007/s00018-005-4539-z PMID: 15761667

10. Hampl V, Stairs CW, Roger AJ. The tangled past of eukaryotic enzymes involved in anaerobic metabolism. Mob Genet Elem. 2011; 1: 71–74. https://doi.org/10.4161/mge.1.1.15588 PMID: 22016847

11. Allen JF, Raven JA, Doolittle WF, Boucher Y, Nesbo CL, Douady CJ, et al. How big is the iceberg of which organellar genes in nuclear genomes are but the tip? Philos Trans R Soc Lond B Biol Sci. 2003; 358: 39–58. https://doi.org/10.1098/rstb.2002.1185 PMID: 12594917

12. Andersson JO, Sigjoren AM, Davis L a. M, Embley TM, Roger AJ. Phylogenetic analyses of diplomonad genes reveal frequent lateral gene transfers affecting eukaryotes. Curr Biol. 2003; 13: 94–104. https://doi.org/10.1016/s0096-9822(03)00003-4 PMID: 12546782

13. Keeling PJ, Burger G, Durnford DG, Lang BF, Lee RW, Pearlman RE, et al. The tree of eukaryotes. Trends Ecol Evol. 2005; 20: 670–676. https://doi.org/10.1016/j.tree.2005.09.005 PMID: 16701456

14. Keeling PJ, Palmer JD. Horizontal gene transfer in eukaryotic evolution. Nat Rev Genet. 2008; 9: 605–618. https://doi.org/10.1038/nrg2386 PMID: 18591983

15. Sibbald SJ, Eme L, Archibald JM, Roger AJ. Lateral Gene Transfer Mechanisms and Pan-genomes in Eukaryotes. Trends Parasitol. 2020; 36: 927–941. https://doi.org/10.1016/j.pt.2020.07.014 PMID: 32828660

16. Katz LA, Grant JR. Taxon-rich phylogenomic analyses resolve the eukaryotic tree of life and reveal the power of subsampling by sites. Syst Biol. 2015; 64: 406–415. https://doi.org/10.1093/sysbio/syu126 PMID: 25540455

17. Bemm F, Weiß CL, Schultz J, Förster F. Genome of a tardigrade: Horizontal gene transfer or bacterial contamination? Proc Natl Acad Sci. 2016; 113: E3054–E3056. https://doi.org/10.1073/pnas.1525116113 PMID: 27173902

18. Francois CM, Durand F, Figuet E, Galtier N. Prevalence and Implications of Contamination in Public Genomic Resources: A Case Study of 43 Reference Arthropod Assemblies. G3 GenesGenomeGenetics. 2020; 10: 721–730. https://doi.org/10.1534/g3.119.400758 PMID: 31867277

19. Lupo V, Van Vlierberghe M, Vanderschuren H, Kerff F, Baurain D, Cornel T. Contamination in Reference Sequence Databases: Time for Divide-and-Rule Tactics. Front Microbiol. 2021; 12: 3233. https://doi.org/10.3389/fmicb.2021.755101 PMID: 34745061

20. Jaramillo VDA, Sukno SA, Thon MR. Identification of horizontally transferred genes in the genus Colletotrichum reveals a steady tempo of bacterial to fungal gene transfer. BMC Genomics. 2015; 16: 2. https://doi.org/10.1186/1471-2164-16-2 PMID: 25555398

21. Azad RK, Lawrence JG. Detecting Laterally Transferred Genes. In: Anisimova M, editor. Evolutionary Genomics: Statistical and Computational Methods, Volume 1. Totowa, NJ: Human Press; 2012. pp. 281–308. https://doi.org/10.1007/978-1-61779-582-4_10

22. Gladyshev EA, Meselson M, Arkhipova IR. Massive Horizontal Gene Transfer in Bdelloid Rotifers. Science. 2008; 320: 1210–1213. https://doi.org/10.1126/science.1156407 PMID: 18511688

23. Kurland CG, Canback B, Berg OG. Horizontal gene transfer: a critical view. Proc Natl Acad Sci U S A. 2003; 100: 9658–9662. https://doi.org/10.1073/pnas.1632870100 PMID: 12902542

24. Matriano DM, Alegado RA, Conaco C. Detection of horizontal gene transfer in the genome of the choanoflagellate Salpingoeca rosetta. Sci Rep. 2021; 11: 5993. https://doi.org/10.1038/s41598-021-85259-5 PMID: 33727612
25. Lander ES, Linton LM, Birren B, Nusbaum C, Zody MC, Baldwin J, et al. Initial sequencing and analysis of the human genome. Nature. 2001; 409: 860–921. https://doi.org/10.1038/35057062 PMID: 11237011

26. Salzberg SL, White O, Peterson J, Eisen JA. Microbial Genes in the Human Genome: Lateral Transfer or Gene Loss? Science. 2001; 292: 1903–1906. https://doi.org/10.1126/science.1061036 PMID: 11358996

27. Huynen M, Snel B, Bork P. Lateral Gene Transfer, Genome Surveys, and the Phylogeny of Prokaryotes. Science. 1999; 286: 1443–1443. https://doi.org/10.1126/science.286.5444.1443

28. Wolf YI, Koonin EV. Genome reduction as the dominant mode of evolution. BioEssays News Rev Mol Cell Dev Biol. 2013; 35: 829–837. https://doi.org/10.1002/bies.201300037 PMID: 23801028

29. Naranjo-Ortiz MA, Gabaldón T. Fungal evolution: cellular, genomic and metabolic complexity. Biol Rev. 2020; 95: 1198–1232. https://doi.org/10.1111/brv.12605 PMID: 32301582

30. Richards TA, Monier A. A tale of two tardigrades. Proc Natl Acad Sci. 2016; 113: 4892–4894. https://doi.org/10.1073/pnas.1603862113 PMID: 27064885

31. Almeida FC, Leszczyniecka M, Fisher PB, Desalle R. Examining ancient inter-domain horizontal gene transfer. Evol Bioinforma Online. 2008; 4: 109–119. PMID: 19204812

32. Adato O, Ninyo N, Gophna U, Snir S. Detecting Horizontal Gene Transfer between Closely Related Taxa. PLoS Comput Biol. 2015; 11: e1004408. https://doi.org/10.1371/journal.pcbi.1004408 PMID: 26439115

33. Cummings TFM, Gori K, Sanchez-Pulido L, Gavriliidis G, Moi D, Wilson AR, et al. Citrullination Was Introduced into Animals by Horizontal Gene Transfer from Cyanobacteria. Mol Biol Evol. 2022; 39: msab317. https://doi.org/10.1093/molbev/msab317 PMID: 34730808

34. Mistry J, Chapman S, Williams L, Qureshi M, Salazar GA, Sonnhammer ELL, et al. Pfam: The protein families database in 2021. Nucleic Acids Res. 2021; 49: D412–D419. https://doi.org/10.1093/nar/gkaa913 PMID: 33125078

35. Ceron-Romero MA, Maurer-Alcala XX, Grattepanche J-D, Yan Y, Fonseca MM, Katz LA. PhylotoL: A Taxon/Gene-Rich Phylogenomic Pipeline to Explore Genome Evolution of Diverse Eukaryotes. Mol Biol Evol. 2019; 36: 1831–1842. https://doi.org/10.1093/molbev/msz103 PMID: 31062861

36. Sela I, Ashkenazy H, Katoh K, Pupko T. GUIDANCE2: accurate detection of unreliable alignment regions accounting for the uncertainty of multiple parameters. Nucleic Acids Res. 2015; 43: W7–14. https://doi.org/10.1093/nar/gkv318 PMID: 25883146

37. Stamatakis A. RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. Bioinformatics. 2014; 30: 1312–1313. https://doi.org/10.1093/bioinformatics/btu333 PMID: 24451623

38. Xia J, Guo Z, Yang Z, Han H, Wang S, Xu H, et al. Whitefly hijacks a plant detoxification gene that neutralizes plant toxins. Cell. 2021; 184: 1693–1705.e17. https://doi.org/10.1016/j.cell.2021.02.014 PMID: 37370502

39. Gogarten JP, Townsend JP. Horizontal gene transfer, genome innovation and evolution. Nat Rev Microbiol. 2005; 3: 679–687. https://doi.org/10.1038/nrmicro1204 PMID: 16138096

40. Gogarten JP, Doolittle WF, Lawrence JG. Prokaryotic evolution in light of gene transfer. Mol Biol Evol. 2002; 19: 2226–2238. https://doi.org/10.1093/oxfordjournals.molbev.a004046 PMID: 12446813

41. Ku C, Nelson-Sathi S, Roettger M, Garg S, Hazkani-Covo E, Martin WF. Endosymbiotic gene transfer from prokaryotic pangenomes: Inherited chimerism in eukaryotes. Proc Natl Acad Sci. 2015; 112: 10139–10146. https://doi.org/10.1073/pnas.1421385112 PMID: 25733873

42. Roger AJ, Muñoz-Gómez SA, Kamikawa R. The Origin and Diversification of Mitochondria. Curr Biol. 2017; 27: R1177–R1192. https://doi.org/10.1016/j.cub.2017.09.015 PMID: 29112874
48. Fan X, Qiu H, Han W, Wang Y, Xu D, Zhang X, et al. Phytoplankton pangenome reveals extensive prokaryotic horizontal gene transfer of diverse functions. Sci Adv. 2020; 6: eaba0111. https://doi.org/10.1126/sciadv.aba0111 PMID: 32494685

49. Susko E, Steel M, Roger AJ. Conditions under which distributions of edge length ratios on phylogenetic trees can be used to order evolutionary events. J Theor Biol. 2021; 526: 110788. https://doi.org/10.1016/j.jtbi.2021.110788 PMID: 34097914

50. Yue J, Sun G, Hu X, Huang J. The scale and evolutionary significance of horizontal gene transfer in the choanoflagellate Monosiga brevicollis. BMC Genomics. 2013; 14: 729. https://doi.org/10.1186/1471-2164-14-729 PMID: 24156600

51. Dorrell RG, Villain A, Perez-Lamarque B, Audren de Kerdrel G, McCallum G, Watson AK, et al. Phylogenomic fingerprinting of tempo and functions of horizontal gene transfer within ochrophytes. Proc Natl Acad Sci. 2021; 118: e2009974118. https://doi.org/10.1073/pnas.2009974118 PMID: 33419955

52. Keeling PJ. The endosymbiotic origin, diversification and fate of plastids. Philos Trans R Soc B-Biol Sci. 2010; 365: 729–748. https://doi.org/10.1098/rstb.2009.0103 PMID: 20124341

53. Nowack ECM, Vogel H, Groth M, Grossman AR, Melkonian M, Glockner G. Endosymbiotic Gene Transfer and Transcriptional Regulation of Transferred Genes in Paulinella chromatophora. Mol Biol Evol. 2011; 28: 407–422. https://doi.org/10.1093/molbev/msq209 PMID: 20702568

54. Keeling PJ. The Number, Speed, and Impact of Plastid Endosymboses in Eukaryotic Evolution. In: Merchant SS, editor. Annual Review of Plant Biology, Vol 64. Palo Alto: Annual Reviews; 2013. pp. 583–607. Available: https://gateway.webofknowledge.com/gateway/Gateway.cgi?GWVersion=2&SrcAuth=DOISource&SrcApp=WOS&KeyID=10.1146%2Fannurev-plant-050312-120144&DestApp=DOISource&SrcAppID=5EWyTzfOX9hQBl3U7u&SrcTitle=ANNUAL+REVIEW+OF+PLANT+BIOLOGY%2C+VOL+64&DestDOIRegistrantName=Annual+Reviews

55. Szafranski P. Intercompartmental Piecewise Gene Transfer. Genes. 2017; 8: 260. https://doi.org/10.3390/genes8100260 PMID: 28984842

56. Gould SB, Waller RR, McFadden GI. Plastid evolution. Annu Rev Plant Biol. 2008; 59: 491–517. https://doi.org/10.1146/annurev.arplant.59.032607.092915 PMID: 18315522

57. Larkum AWD, Lockhart PJ, Howe CJ. Shopping for plastids. Trends Plant Sci. 2007; 12: 189–195. https://doi.org/10.1016/j.tplants.2007.03.011 PMID: 17416546

58. McFadden GI. Primary and secondary endosymbiosis and the origin of plastids. J Phycol. 2001; 37: 951–959. https://doi.org/10.1046/j.1529-8817.2001.00433.x

59. Zimorski V, Ku C, Martin WF, Gould SB. Endosymbiotic theory for organelle origins. Curr Opin Microbiol. 2014; 22: 38–46. https://doi.org/10.1016/j.mib.2014.09.008 PMID: 25306530

60. Dorrell RG, Bowler C. Secondary Plastids of Stramenopiles. In: Hirakawa Y, editor. Secondary Endosymbioses. London: Academic Press Ltd-Elsevier Science Ltd; 2017. pp. 57–103. Available: https://www.webofscience.com/wos/woscc/full-record/WOS:000414512900004

61. McFadden GI, Waller RE, Reith ME, Lang-Umnasch N. Plastids in apicomplexan parasites. Plant Syst Evol. 1997; 261–287.

62. Fernández Robledo JA, Caler E, Matsuoka M, Keeling PJ, Shanmugam D, Roos DS, et al. The search for the missing link: A relic plastid in Perkinsus? Int J Parasitol. 2011; 41: 1217–1229. https://doi.org/10.1016/j.ijpara.2011.07.008 PMID: 21889509

63. Reyes-Prieto A, Hackett JD, Soares MB, Bonaldo MF, Bhattacharya D. Cyanobacterial Contribution to Algal Nuclear Genomes Is Primarily Limited to Plastid Functions. Curr Biol. 2006; 16: 2320–2325. https://doi.org/10.1016/j.cub.2006.09.063 PMID: 17141613

64. Llorente B, de Souza FSJ, Soto G, Meyer C, Alonso GD, Flawia MM, et al. Selective pressure against horizontally acquired prokaryotic genes as a driving force of plastid evolution. Sci Rep. 2016; 6: 19036. https://doi.org/10.1038/srep19036 PMID: 26750147

65. Chen X, Schreiber K, Appel J, Makowka A, Fahrbrich B, Roettger M, et al. The Entner–Doudoroff pathway is an overlooked glycolytic route in cyanobacteria and plants. Proc Natl Acad Sci. 2016; 113: 5441–5446. https://doi.org/10.1073/pnas.1521916113 PMID: 27114545

66. Pyrh J, Žásky V, Fellows JD, Grosche C, Włoga D, Striepen B, et al. The iron-sulfur scaffold protein HCF101 unveils the complexity of organelar evolution in SAR, Haptista and Cryptista. BMC Ecol Evol. 2021; 21: 46. https://doi.org/10.1186/s12862-021-01777-x PMID: 33740894

67. Bhattacharya D, Nosenko T. Endosymbiotic and horizontal gene transfer in chromalveolates. J Phycol. 2008; 44: 7–10. https://doi.org/10.1111/j.1529-8817.2007.00433.x PMID: 27041032

68. Carter DR. Plastocyanin-ferredoxin oxidoreduction and endosymbiotic gene transfer. Photosynth Res. 2008; 97: 245–253. https://doi.org/10.1007/s11120-008-9333-y PMID: 18661249
69. Burki F, Imanian B, Hehenberger E, Hirakawa Y, Maruyama S, Keeling PJ. Endosymbiotic Gene Transfer in Tertiary Plastid-Containing Dinoflagellates. Eukaryot Cell. 2014; 13: 246–255. https://doi.org/10.1128/EC.00299-13 PMID: 24297445

70. Suzuki S, Endoh R, Manabe R, Ohkuma M, Hirakawa Y. Multiple losses of photosynthesis and convergent reductive genome evolution in the colourless green algae Protococca. Sci Rep. 2018; 8: 940. https://doi.org/10.1038/s41598-017-18378-8 PMID: 29343788

71. Keeling PJ, Fast NM. Microsporidia: Biology and evolution of highly reduced intracellular parasites. Annu Rev Microbiol. 2002; 56: 93–116. https://doi.org/10.1146/annurev.micro.56.012302.160854 PMID: 12142484

72. Dean P, Sendra KM, Williams TA, Watson AK, Major P, Nakjang S, et al. Transporter gene acquisition and innovation in the evolution of Microsporidia intracellular parasites. Nat Commun. 2018; 9: 1709. https://doi.org/10.1038/s41467-018-03923-4 PMID: 29703975

73. Gonçalves C, Coelho MA, Salema-Ooom M, Gonçalves P. Stepwise Functional Evolution in a Fungal Sugar Transporter Family. Mol Biol Evol. 2016; 33: 352–366. https://doi.org/10.1093/molbev/msv220 PMID: 26474848

74. Schmitt I, Lumbsch HT. Ancient Horizontal Gene Transfer from Bacteria Enhances Biosynthetic Capabilities of Fungi. Plos One. 2009; 4: e4437. https://doi.org/10.1371/journal.pone.0004437 PMID: 19212443

75. Seipke RF, Kaltenpoth M, Hutchings MI. Streptomycetes as symbionts: an emerging and widespread theme? Fems Microbiol Rev. 2012; 36: 862–876. https://doi.org/10.1111/j.1574-6976.2011.00313.x PMID: 22091965

76. Jain R, Rivera MC, Moore JE, Lake JA. Horizontal gene transfer accelerates genome innovation and evolution. Mol Biol Evol. 2003; 20: 1598–1602. https://doi.org/10.1093/molbev/msg154 PMID: 12775714

77. Richards TA, Soanes DM, Jones MDM, Vasieva O, Leonard G, Paszkiewicz K, et al. Horizontal gene transfer facilitated the evolution of plant parasitic mechanisms in the oomycetes. Proc Natl Acad Sci. 2011; 108: 15258–15263. https://doi.org/10.1073/pnas.1105100108 PMID: 21878562

78. Qiu H, Cai G, Luo J, Bhattacharya D, Zhang N. Extensive horizontal gene transfers between plant pathogenic fungi. BMC Biol. 2016; 14: 41. https://doi.org/10.1186/s12915-016-0264-3 PMID: 27215567

79. Torres-Cortés G, Ghignone S, Bonfante P, Schüßler A. Mosaic genome of endobacteria in arbuscular mycorrhizal fungi: Transkingdom gene transfer in an ancient mycoplasma-fungus association. Proc Natl Acad Sci. 2015; 112: 7785–7790. https://doi.org/10.1073/pnas.1501540112 PMID: 25964335

80. Feurtey A, Stukentsekh EH. Interspecific Gene Exchange as a Driver of Adaptive Evolution in Fungi. Annu Rev Microbiol. 2018; 72: 377–398. https://doi.org/10.1146/annurev-micro-090817-062753 PMID: 29927707

81. Dal Grande F, Widmer I, Wagner HH, Scheidegger C. Vertical and horizontal photobiont transmission within populations of a lichen symbiosis. Mol Ecol. 2012; 21: 3159–3172. https://doi.org/10.1111/j.1365-294X.2012.05482.x PMID: 22384938

82. Tunji M, Kora P. Vertical and horizontal gene transfer in lichens. Period Biol. 115: 9.

83. Nguyen TA, Greig J, Khan A, Goh C, Jedd G. Evolutionary novelty in gravity sensing through horizontal gene transfer and high-order protein assembly. PLOS Biol. 2018; 16: e2004920. https://doi.org/10.1371/journal.pbio.2004920 PMID: 29689046

84. Lacroix B, Citovsky V. Transfer of DNA from Bacteria to Eukaryotes. mBio. 2014; 7: e00863–16. https://doi.org/10.1128/mBio.00863-16 PMID: 27406565

85. Wisecaver JH, Slot JC, Rokas A. The Evolution of Fungal Metabolic Pathways. PLOS Genet. 2014; 10: e1004816. https://doi.org/10.1371/journal.pgen.1004816 PMID: 25474404

86. Richards TA, Leonard G, Soanes DM, Talbot NJ. Gene transfer into the fungi. Fungal Biol Rev. 2011; 25: 98–110. https://doi.org/10.1016/j.fbr.2011.01.028 PMID: 21334300

87. Frost LS, Leplae R, Summers AO, Toussaint A. Mobile genetic elements: The agents of open source evolution. Nat Rev Microbiol. 2005; 3: 722–732. https://doi.org/10.1038/nrmicro1235 PMID: 16138100

88. Eichinger L, Pachebat JA, Giöckner G, Rajandream M-A, Sugcang R, Berriman M, et al. The genome of the social amoeba Dictyostelium discoideum. Nature. 2005; 435: 43–57. https://doi.org/10.1038/nature03481 PMID: 15875012

89. Hug LA, Stichmann A, Roger AJ. Phylogenetic distributions and histories of proteins involved in anaerobic pyruvate metabolism in eukaryotes. Mol Biol Evol. 2010; 27: 311–324. https://doi.org/10.1093/molbev/msp237 PMID: 19805439
90. Eme L, Gentekaki E, Curtis B, Archibald JM, Roger AJ. Lateral Gene Transfer in the Adaptation of the Anaerobic Parasite Blastocystis to the Gut. Curr Biol CB. 2017; 27: 807–820. https://doi.org/10.1016/j.cub.2017.02.003 PMID: 28262486

91. Alsmark C, Foster PG, Sicheritz-Ponten T, Nakjang S, Embley TM, Hirt RP. Patterns of prokaryotic lateral gene transfers affecting parasitic microbial eukaryotes. Genome Biol. 2013; 14: R19. https://doi.org/10.1186/gb-2013-14-2-r19 PMID: 23442822

92. Grant JR, Katz LA. Phylogenetic study indicates widespread lateral gene transfer in Entamoeba and suggests a past intimate relationship with parabasalids. Genome Biol Evol. 2014; 6: 2350–2360. https://doi.org/10.1093/gbe/evu179 PMID: 25146649

93. Alsmark UC, Sicheritz-Ponten T, Foster PG, Hirt RP, Embley TM. Horizontal gene transfer in eukaryotic parasites: a case study of Entamoeba histolytica and Trichomonas vaginalis. Methods Mol Biol Clifton NJ. 2009; 532: 489–500. https://doi.org/10.1007/978-1-60327-853-9_28 PMID: 19271203

94. Tanifuji G, Cenci U, Moog D, Dean S, Nakayama T, David V, et al. Genome sequencing reveals metabolic and cellular interdependence in an amoeba-kinetoplastid symbiosis. Sci Rep. 2017; 7: 11688. https://doi.org/10.1038/s41598-017-11866-x PMID: 28916813

95. Gould SB, Garg SG, Handrich M, Nelson-Sathi S, Gruenheit N, Tielens AGM, et al. Adaptation to life on land at high O$_2$ via transition from ferredoxin-to NADH-dependent redox balance. Proc R Soc B Biol Sci. 2019; 286: 20191491. https://doi.org/10.1098/rspb.2019.1491 PMID: 31431166

96. Zimorski V, Mentel M, Tielens AGM, Martin WF. Energy metabolism in anaerobic eukaryotes and Earth’s late oxygenation. Free Radic Biol Med. 2019; 140: 279–294. https://doi.org/10.1016/j.freeradbiomed.2019.03.030 PMID: 30935869

97. Rosenthal B, Mai Z, Caplivski D, Ghosh S, de la Vega H, Graf T, et al. Evidence for the bacterial origin of genes encoding fermentation enzymes of the amitochondriate protozoan parasite Entamoeba histolytica. J Bacteriol. 1997; 179: 3736–3745. https://doi.org/10.1128/jb.179.11.3736-3745.1997 PMID: 9171424

98. Horner DS, Hirt RP, Embley TM. A single eubacterial origin of eukaryotic pyruvate: ferredoxin oxidoreductase genes: implications for the evolution of anaerobic eukaryotes. Mol Biol Evol. 1999; 16: 1280–1291. https://doi.org/10.1093/oxfordjournals.molbev.a026218 PMID: 10486982

99. Gelius-Dietrich G, Henze K. Pyruvate Formate Lyase (PFL) and PFL Activating Enzyme in the Chytrid Fungus Neocallimastix frontalis: A Free-Radical Enzyme System Conserved Across Divergent Eukaryotic Lineages.1. J Eukaryot Microbiol. 2004; 51: 456–463. https://doi.org/10.1111/j.1550-7408.2004.tb00394.x PMID: 15352329

100. Clark CG, Alsmark UCM, Tazreiter M, Saito-Nakano Y, Ali V, Marion S, et al. Structure and Content of the Entamoeba histolytica Genome. Advances in Parasitology. Academic Press; 2007. pp. 51–190. https://doi.org/10.1016/S0065-308X(07)65002-7

101. König C, Meyer M, Lender C, Nehls S, Wallaschkowski T, Holm T, et al. An Alcohol Dehydrogenase 3 (ADH3) from Entamoeba histolytica Is Involved in the Detoxification of Toxic Aldehydes. Microorganisms. 2020; 8: E1608. https://doi.org/10.3390/microorganisms8101608 PMID: 33086693

102. Müller J, Müller N. Nitroreductases of bacterial origin in Giardia lamblia: Potential role in detoxification of xenobiotics. MicrobiologyOpen. 2019; 8: e904. https://doi.org/10.1002/mbo3.904 PMID: 31343119

103. Lapierre P, Lasek-Nesselquist E, Gogarten JP. The impact of HGT on phylogenomic reconstruction methods. Brief Bioinform. 2014; 15: 79–90. https://doi.org/10.1093/bib/bbt050 PMID: 22908214

104. Doolittle WF, Brunet TDP. What Is the Tree of Life? PLoS Genet. 2016; 12: e1005912. https://doi.org/10.1371/journal.pgen.1005912 PMID: 27078870

105. Doolittle WF. Phylogenetic Classification and the Universal Tree. Science. 1999; 284: 2124–2128. https://doi.org/10.1126/science.284.5423.2124 PMID: 10381871

106. Coleman GA, Davin AA, Mahendrarajah TA, Szántóh LL, Spang A, Hugenholz P, et al. A rooted phylogeny resolves early bacterial evolution. Science. 2021; 372: eabe0511. https://doi.org/10.1126/science.abe0511 PMID: 33958449

107. Chen F, Mackey AJ, Stockecket CJ Jr, Roos DS. OrthoMCL-DB: querying a comprehensive multi-species collection of ortholog groups. Nucleic Acids Res. 2006; 34: D363–D368. https://doi.org/10.1093/nar/gkl123 PMID: 16381887

108. Edgar RC. Search and clustering orders of magnitude faster than BLAST. Bioinformatics. 2010; 26: 2460–2461. https://doi.org/10.1093/bioinformatics/btq461 PMID: 20709691

109. Katoh K, Misawa K, Kuma KI, Miyata T. MAFFT: a novel method for rapid multiple sequence alignment based on fast Fourier transform. Nucleic Acids Res. 2002; 30: 3059–3066. https://doi.org/10.1093/nar/gkf436 PMID: 12136088
110. Capella-Gutiérrez S, Silla-Martínez JM, Gabaldón T. trimAl: a tool for automated alignment trimming in large-scale phylogenetic analyses. Bioinformatics. 2009; 25: 1972–1973. https://doi.org/10.1093/bioinformatics/btp348 PMID: 19505945

111. Wright F. The ‘effective number of codons’ used in a gene. Gene. 1990; 87: 23–29. https://doi.org/10.1016/0378-1119(90)90491-9 PMID: 2110097

112. Biopython: freely available Python tools for computational molecular biology and bioinformatics | Bioinformatics | Oxford Academic. [cited 5 Sep 2021]. Available: https://academic.oup.com/bioinformatics/article/25/11/1422/330687

113. Rognes T, Flouri T, Nichols B, Quince C, Mahé F. VSEARCH: a versatile open source tool for metagenomics. PeerJ. 2016; 4: e2584. https://doi.org/10.7717/peerj.2584 PMID: 27781170

114. Camacho C, Coulouris G, Avagyan V, Ma N, Papadopoulos J, Bealer K, et al. BLAST+: architecture and applications. BMC Bioinformatics. 2009; 10: 421. https://doi.org/10.1186/1471-2105-10-421 PMID: 20003500

115. Minh BQ, Schmidt HA, Chernomor O, Schrempf D, Woodhams MD, von Haeseler A, et al. IQ-TREE 2: New Models and Efficient Methods for Phylogenetic Inference in the Genomic Era. Mol Biol Evol. 2020/02/06 ed. 2020; 37: 1530–1534. https://doi.org/10.1093/molbev/msaa015 PMID: 32011700

116. Miller MA, Pfeiffer W, Schwartz T. Creating the CIPRES Science Gateway for inference of large phylogenetic trees. 2010 Gateway Computing Environments Workshop (GCE). 2010. pp. 1–8. https://doi.org/10.1109/GCE.2010.5676129

117. Huerta-Cepas J, Szklarczyk D, Heller D, Hernández-Plaza A, Forslund SK, Cook H, et al. eggNOG 5.0: a hierarchical, functionally and phylogenetically annotated orthology resource based on 5090 organisms and 2502 viruses. Nucleic Acids Res. 2019; 47: D309–D314. https://doi.org/10.1093/nar/gky1085 PMID: 30418610

118. Buchfink B, Xie C, Huson DH. Fast and sensitive protein alignment using DIAMOND. Nat Methods. 2015; 12: 59–60. https://doi.org/10.1038/nmeth.3176 PMID: 25402007

119. Mitchell A, Chang H-Y, Daugherty L, Fraser M, Hunter S, Lopez R, et al. The InterPro protein families database: the classification resource after 15 years. Nucleic Acids Res. 2015; 43: D213–221. https://doi.org/10.1093/nar/gku1243 PMID: 25428371

120. Eddy SR. Accelerated Profile HMM Searches. PLOS Comput Biol. 2011; 7: e1002195. https://doi.org/10.1371/journal.pcbi.1002195 PMID: 22039361

121. Conway JR, Lex A, Gehlenborg N. UpSetR: an R package for the visualization of intersecting sets and their properties. Bioinformatics. 2017; 33: 2938–2940. https://doi.org/10.1093/bioinformatics/btx364 PMID: 28645171