

Single-cell genomics reveals complex microbial and viral associations in ciliates and testate amoebae

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Protists play important roles in nutrient cycling across ecosystems, yet the composition and function of their associated microbiomes remain poorly studied. Here, we use cultivation-independent single-cell isolation and genome-resolved metagenomics to investigate the microbiomes and viromes of more than 100 uncultivated ciliates and amoebae from diverse environments. Our findings reveal unique microbiome structures and complex associations with bacterial symbionts and viruses, with stark differences between ciliates and amoebae. We recover 117 microbial genomes affiliated with known eukaryotic endosymbionts, including *Holosporales*, *Rickettsiales*, *Legionellales*, *Chlamydiae*, and *Babelota*, and 258 genomes linked to host-associated *Patescibacteriota*. Many show genome reduction and genes related to toxin-antitoxin systems and nucleotide parasitism, indicating adaptation to intracellular lifestyles. We also identify more than 80 giant viruses from diverse lineages, some actively expressing genes in single-cell transcriptomes, along with other viruses predicted to infect eukaryotes or symbiotic bacteria. The frequent co-occurrence of giant viruses and microbial symbionts, especially in amoebae, suggests multipartite interactions. Together, our study highlights protists as hubs of microbial and viral associations and provides a broad view of the diversity, activity, and ecological importance of their hidden partners.

Protists, microeukaryotes that are not fungi, plants or animals¹, play instrumental roles in global ecosystems where they contribute to nutrient cycling and shape the structure and function of microbial communities^{2,3}. Heterotrophic protists are well known for their role in grazing, a process in which other microbes are being taken up through phagocytosis and digested⁴. This turnover of microbial biomass ultimately makes nutrients available to higher trophic levels². Grazing isn't the sole process that protists are involved in. Both heterotrophs

and autotrophs (i.e., unicellular algae) play central roles in biomineralization⁵, while autotrophs and mixotrophs contribute to organic carbon fixation^{6,7}.

Recent studies suggest that protists may harbor complex microbiomes⁸. For example, some amoebae have evolved strategies to maintain a regime of bacteria as part of their microbiome as food to ensure a constant nutrient supply⁹. Some of the protist-associated bacteria are resistant to digestion¹⁰ and may even be able to replicate

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inside their eukaryotic host cells¹¹. This ability might lead to host dependency¹² or even protection against pathogens¹³. Further, amoebae represent natural reservoirs for a wider range of human pathogens¹⁴. This complexity could have far-reaching implications, not only for the surrounding microbial communities and nutrient cycling but also for ecosystem, animal, and plant health.

Despite their importance, our insights into the roles of protists in ecosystems and particularly their interactions with associated microbes and viruses remain limited to a few well-studied groups such as *Acanthamoeba* and *Paramecium*^{15–17}. These have received attention due to their medical relevance and their ability to be cultivated under axenic or monoxenic conditions. Only recently, evidence emerged on the presence of complex and often species-specific microbiomes of protists^{8,18–20}. However, the vast diversity of unexplored eukaryotes leaves broad gaps in our understanding of diversity, function and associations of less well-studied protist lineages^{11,21–23}.

To address these limitations, we collected over 100 individual cells of diverse microbial eukaryotes directly from the environment, including ciliates and testate (i.e., shell-building) amoebae. Most of these organisms are understudied and have not been successfully maintained in culture. Using cultivation-independent single cell isolation, whole genome amplification and genome-resolved metagenomics, as well as single cell transcriptomics, we provide insights into the unique composition of the protist microbiome and virome. We uncover associations with a wide range of putative pathogens, including both microbial and eukaryotic symbionts and viruses known to infect amoebae and ciliates. Our findings underscore the role of these studied ciliates and amoebae as important players in complex microbial and viral associations in the environment and shed light on the intricate roles that these organisms play within ecological networks.

Results and discussion

Protist microbiome composition and diversity

The metagenomic binning of sequences from 104 single amplified genomes (SAGs) belonging to three testate amoeba and eight ciliate species (Fig. 1a and Supplementary Data 1) yielded a total of 724 prokaryotic metagenome-assembled genomes (MAGs; Supplementary Data 1, 2). According to MIMAG standards²⁴, 439 were of low, 209 of medium and 76 of high quality. Sequencing depth varied between samples, but the greatest number of MAGs per gigabase (Gb) of reads was recovered from the amoeba *Hyalosphenia elegans* and the ciliate *Loxodes* sp., the latter of which was deeply sequenced (Fig. 1a). *Hyalosphenia elegans* showed a much higher recovery rate of MAGs per Gb of sequence data compared to its sister species, *Hyalosphenia papilio*. This trend was also visible in the overall microbiome diversity; *Hyalosphenia elegans* was associated with a greater number of detected bacterial and archaeal phyla ($n=16$), orders ($n=52$) and genera ($n=52$), whereas despite higher sequencing depth *Hyalosphenia papilio* plateaued at 15 phyla, 35 orders and 80 genera (Fig. 1b). This may reflect the relative sizes of these organisms as more of the genomic material may be host for the larger *H. papilio* (100–150 μm) compared to *H. elegans* (80–120 μm) and also the presence of additional microalgal symbionts in *H. papilio*²⁵. Overall, microbiome diversity varied strongly among the sampled lineages; for example, MAGs recovered from *Spirostomum* sp. belonged to just two different bacterial phyla, while those from *Loxodes* sp., isolated from the same low pH environments as the testate amoebae, belonged to up to 16 different bacterial phyla, which corresponded to at least 60 orders and 145 genera (Fig. 1). When comparing the protist microbiomes, it became apparent that ciliates and amoebae have distinct microbiome compositions (Fig. 1a). Distribution of bacterial classes was more similar between amoebal lineages and more dissimilar between

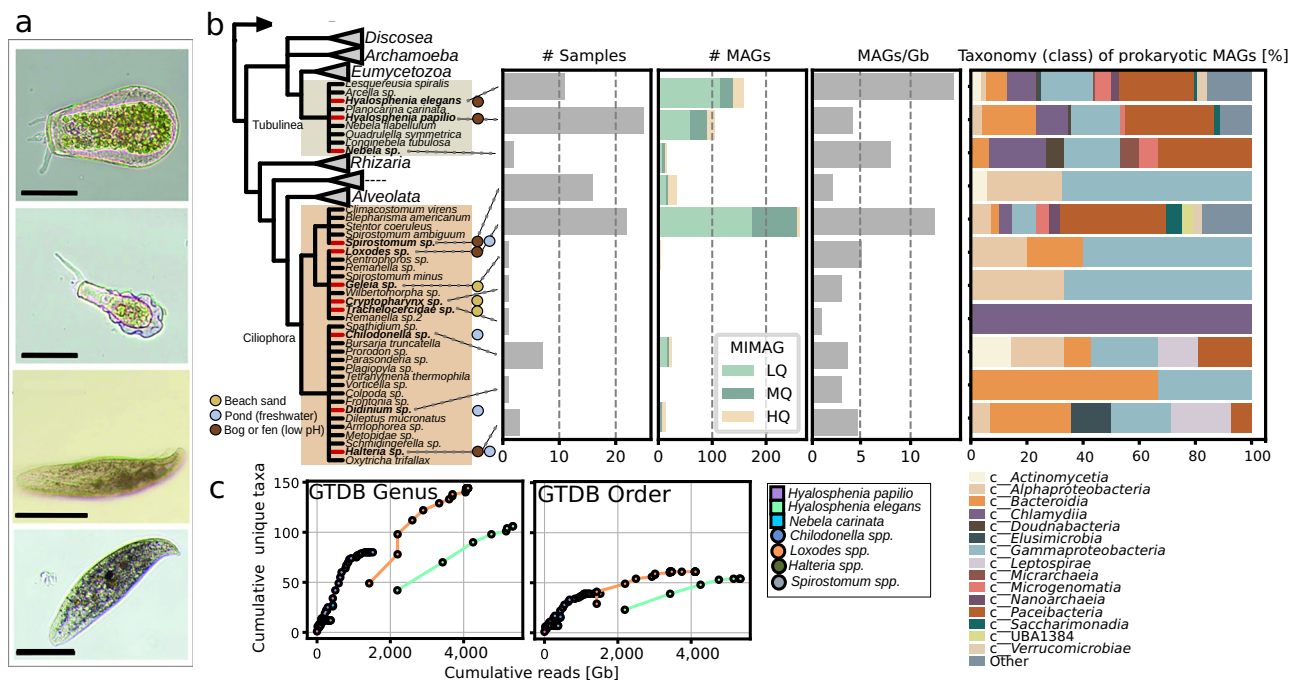


Fig. 1 | Amoeba (*Tubulinea*) and ciliate (*Ciliophora*) lineages sampled in this study. a From top to bottom: *Hyalosphenia papilio* (50 μm scale bar), *Hyalosphenia elegans* (50 μm scale bar), “small” *Loxodes* sp. (100 μm scale bar), “large” *Loxodes* sp. (100 μm scale bar). Note that the images are taken quickly prior to single-cell work, as the aim is to minimize stress to the cells. The micrographs shown are representative of the multiple cells independently isolated for each lineage, the numbers of which are indicated in panel (b). **b** Simplified phylogenetic

tree indicates protist lineages sampled in this study (in bold with red branches), bars indicate the number of samples sequenced, the total number of MAGs per protist lineage and the number of MAGs per Gb sequenced. The right panel shows the taxonomic distribution of bacterial and archaeal MAGs associated with sampled protists. **c** Collector's curves comparing the overall richness of bacterial and archaeal MAGs at different taxonomic levels for each protist lineage.

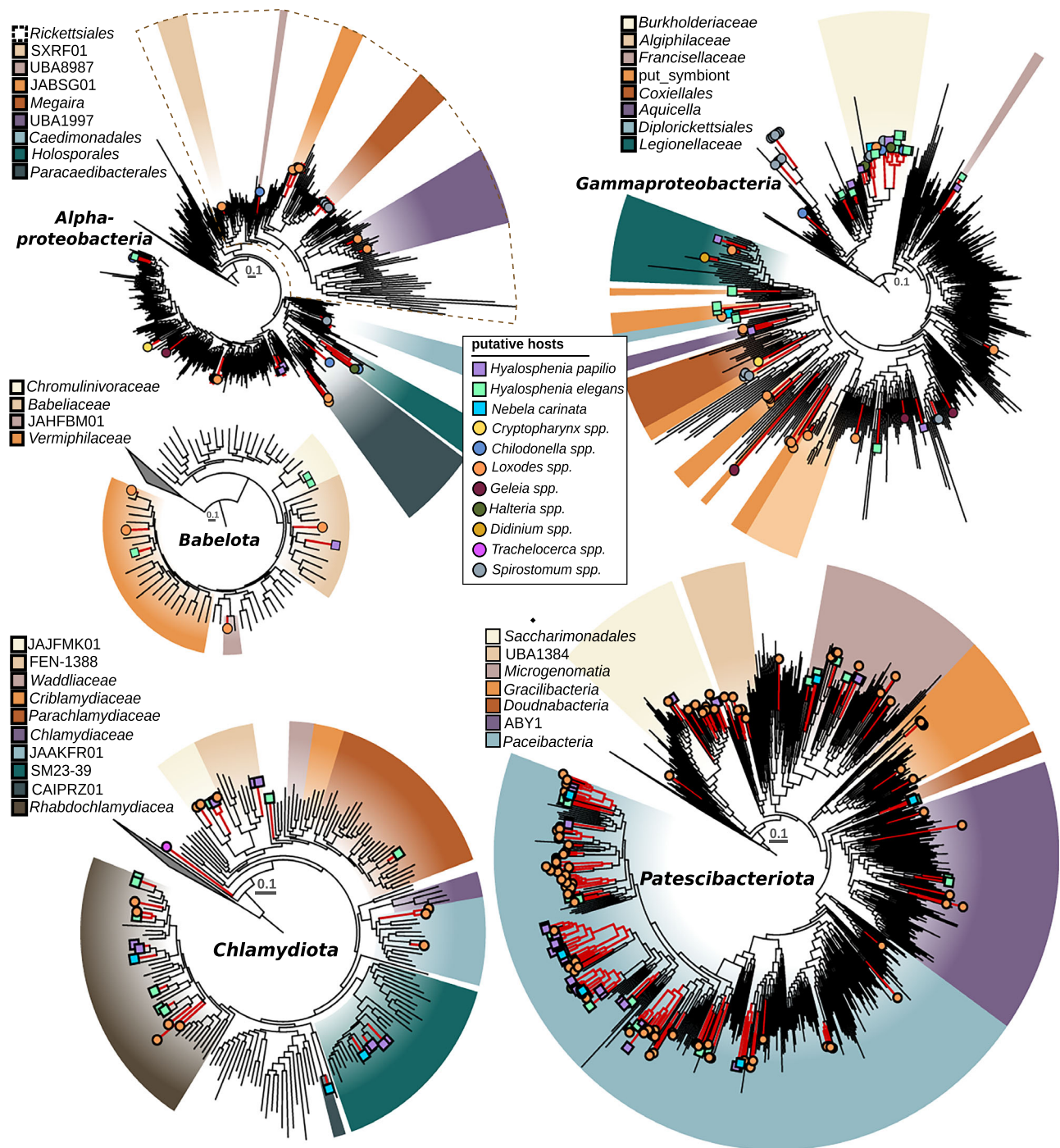


Fig. 2 | Amoeba and ciliate associated bacteria affiliated with known and suggested bacterial endosymbiont clades. Phylogenetic trees indicate the position of putative symbiont MAGs (red branches) that were recovered from protist microbiomes generated in this study. Colored shapes at the terminal branches show the protist lineages the MAGs were associated with. Each tree represents a

phylum (*Chlamydiota*, *Babelota*, *Patescibacteriota*) or class (*Alphaproteobacteria*, *Gammaproteobacteria*) and subclades that were previously shown (in *Chlamydiota*, *Babelota*, *Alphaproteobacteria*, *Gammaproteobacteria*) or suggested (in *Patescibacteriota*) to be host-associated are highlighted with colored wedges.

ciliates, with the freshwater *Loxodes* and the marine *Trachelocercidae* having the least proportion of shared taxa with other ciliates.

Protist microbiomes harbor microbes across known endosymbiont clades

A large proportion of recovered taxa belong to groups of known host-associated bacteria with a facultative or even obligate intracellular lifestyle. Specifically, 117 prokaryotic MAGs grouped with known

bacterial endosymbionts and 258 with putative symbionts (*Patescibacteriota*²⁶) (Fig. 2). Alphaproteobacterial endosymbionts ($n = 28$) were exclusively found in association with ciliates, particularly *Megaira* and *Caedimonadales* in *Spirostomum*, and *Paracaedibacterales* with *Loxodes*, *Chilodonella* and *Halteria* (Fig. 2). In addition, several novel and currently uncharacterized lineages within the order *Rickettsiales* were detected within *Loxodes* and, to a lesser extent, *Chilodonella*. Three of the sampled *Loxodes* cells contained bacteria that

grouped in the gammaproteobacterial family UBA6186 together with *Candidatus Azoamicus ciliatocola*, a bacterial endosymbiont of ciliates with cosmopolitan distribution²⁷ that has been shown to generate energy for its host by denitrification²⁸. Members of *Holospirales* and *Megaira* have previously been associated with different ciliates²⁹, but none of the other lineages have been identified as ciliate endosymbionts thus far. In both species of *Hyalosphenia* sampled in this study, likely host-associated *Gammaproteobacteria* of the family *Francisellaceae* were found. Further, *Diplorickettsia* were present in *Hyalosphenia elegans*, along with *Coxiellales*-related bacteria in the ciliate *Cryptopharynx*, and members of *Legionellales* were present in the ciliates *Didinium* and *Loxodes*, as well as the testate amoeba *Hyalosphenia papilio*. For *Didinium* and *Cryptopharynx*, gammaproteobacteria were the only associated putative symbionts. Another group of protist symbionts, the phylum *Babelota* (former *Dependentiae*), had 12 members from four different families associated with amoebae and ciliates sampled in this study; *Chromulinavoraceae* were exclusively found with *Hyalosphenia papilio* (an amoeba that harbors green algal symbionts), while other families were mixed between *Hyalosphenia* species and *Loxodes*. Our findings suggest that some *Babelota* may represent ciliate symbionts.

One of the best studied symbiont clades is the phylum *Chlamydia*, known to infect a wide diversity of eukaryotic hosts¹⁷. Here, we identified 56 chlamydial MAGs associated with diverse ciliates and amoebae. Four family-level lineages that consist solely of metagenome-assembled genomes were associated with either *Hyalosphenia* (f_FEN-1388), *Loxodes* (f_JAAKFR01), *Hyalosphenia* and *Loxodes* (f_JAJFMA01), or *Hyalosphenia* and *Nebela* (f_SM23-39). Further, the only bacterial symbiont which was found associated with *Tracheohercya* was a highly divergent member of the *Chlamydiota*, potentially representing a novel family or even order-level lineage without any closely related relatives (Fig. 2). The amoeba *Hyalosphenia* was found to be associated with *Parachlamydiaceae*, a group previously shown to infect different amoebae, particularly *Acanthamoeba castellanii*. In *Acanthamoeba* it confers protection against giant virus infection³⁰, but it is also associated with disease in humans and other animals³¹. Most chlamydial MAGs recovered in this study were associated with four different ciliates (*Loxodes*) and amoebae (*Hyalosphenia* and *Nebela*). These were affiliated with *Rhabdochlamydiaceae*, a group previously shown to be predominantly associated with insects and other metazoa³².

Diverse Patescibacteria make up a large fraction of the protist microbiome

In addition to members of well-known intracellular bacteria, we recovered 258 MAGs (25 high quality, 79 medium quality and 154 low quality) representing members from all major groups of the phylum Patescibacteria, mainly associated with *Loxodes*, *Nebela*, and *Hyalosphenia papilio* and, to a lesser extent, with *Hyalosphenia elegans*, *Halteria*, and *Chilodonella* (Fig. 2). Given the reduced genomes of *Patescibacteriota* and features such as lytic transglycosylases and phospholipases and a large number of transporters that may underlie host interaction^{26,33}, it's plausible that some or all of these might be closely associated with amoebae and ciliates. This aligns with a previous study that provided experimental evidence of an uncharacterized *Parcubacterium* as an intracellular bacterium in the ciliate *Paramecium* sp.³⁴. However, reports also exist of association with other bacteria^{35,36} and of a potential free-living lifestyle for *Patescibacteriota*^{26,37,38}. Our results indicate some patterns of co-occurrence, such as clades of *Patescibacteria* composed of MAGs derived from *Loxodes* (Fig. 2). However, the high overall diversity of *Patescibacteriota* MAGs and the absence of a clear host-specificity pattern hampered any predictions in regards to endosymbiosis, as distinguishing true symbiosis from co-picked or ingested cells is particularly challenging for this bacterial phylum and its putative host

(which lacks a robust reference genome) without additional lines of experimental evidence.

Functional and comparative genomics reveal symbiont-specific traits

We performed a comparative genomic analysis of the protist-associated bacteria focusing on metabolic potential and traits relevant to symbiosis (Fig. 3). First, we assessed the relationship between genome completeness and metabolic potential, which revealed a strong positive correlation between genome completeness and the mean completeness of KEGG metabolic modules for all MAG categories (Fig. 3a) and clearly distinguished free-living microbes with the highest module completeness, from predicted symbionts and *Patescibacteriota*, that had the least complete metabolic capabilities, indicative of genome reduction (Fig. 3a). Next, we screened for Toxin-Antitoxin (TA) systems, which can play a role in symbiosis maintenance. We found that TA systems were prevalent across most symbiont groups, particularly in the *Gammaproteobacteria* (including *Legionellales* and UBA6186) and *Patescibacteriota*, suggesting these systems are a common feature of the here protist-associated bacteria, with exception of members of the phylum *Babelota* which were entirely devoid of TA systems (Fig. 3b). Finally, we performed a comparative genomics analysis of the four *Loxodes*-derived MAGs from the order UBA6186 and their close relative '*Candidatus Azoamicus ciliatocola*', which is a known endosymbiont of anaerobic ciliates (Fig. 3c). Our results indicate the shared absence of key metabolic functions, such as genes involved in the tricarboxylic acid (TCA) cycle, or the presence of ATP/ADP translocases, which have a key function in nucleotide parasitism³⁹. However, in contrast to *Azoamicus*, our MAGs do not encode for genes linked to denitrification (nitrate and nitrite reductase), which was a hallmark feature of '*Candidatus Azoamicus ciliatocola*' that utilizes the denitrification pathway to generate energy for its host^{27,39}. Instead, the UBA6186 members found in *Loxodes* encode for cytochrome bd and cytochrome o oxidases, possibly enabling limited respiration at the trace oxygen levels characteristic of *Loxodes*' microaerophilic habitat. Further, the presence of genes coding for malic enzyme and genes involved in proline degradation suggests alternative strategies for energy generation and metabolic flexibility. This likely allows these putative endosymbionts to utilize diverse carbon sources and maintain redox balance through NADPH production, which may be particularly advantageous in the nutrient-limited intracellular environment where primary substrates are scarce.

Giant viruses and virophages are frequently found in protist microbiomes and genes transcribed in situ

The microbiomes of the ciliates and amoebae sampled in our study did not only contain sequences of various host-associated bacteria but 81 giant viruses metagenome assembled genomes (GVMAGs) (Supplementary Data 3). Taxonomic identification with *gvcass*⁴⁰ and phylogenomic analysis revealed that these GVMAGs belonged to diverse lineages within the viral phylum Nucleocytoviricota¹⁶ (Fig. 4a). Specifically, those associated with *Hyalosphenia elegans* belonged to several orders, including *Asfuvirales*, *Pandoravirales*, *Algavirales*, and *Imitervirales*. Viruses associated with *Loxodes* were highly diverse; whereas, those linked to *Hyalosphenia papilio* and *Chilodonella* were confined to a few clades within *Imitervirales*. In previous studies, giant viruses have not been found to directly infect ciliates⁴¹. However, the frequent presence of diverse giant viruses in the ciliates *Loxodes* and *Chilodonella* sampled here suggests members of *Ciliophora* as underappreciated potential hosts for these viruses. For samples where sequences from more than one eukaryote were found, inferring sequence-based putative associations is challenging. For example, members of *Algavirales* might more likely infect the green algae⁴² that are symbiotic to *Hyalosphenia papilio* and were also detected in the

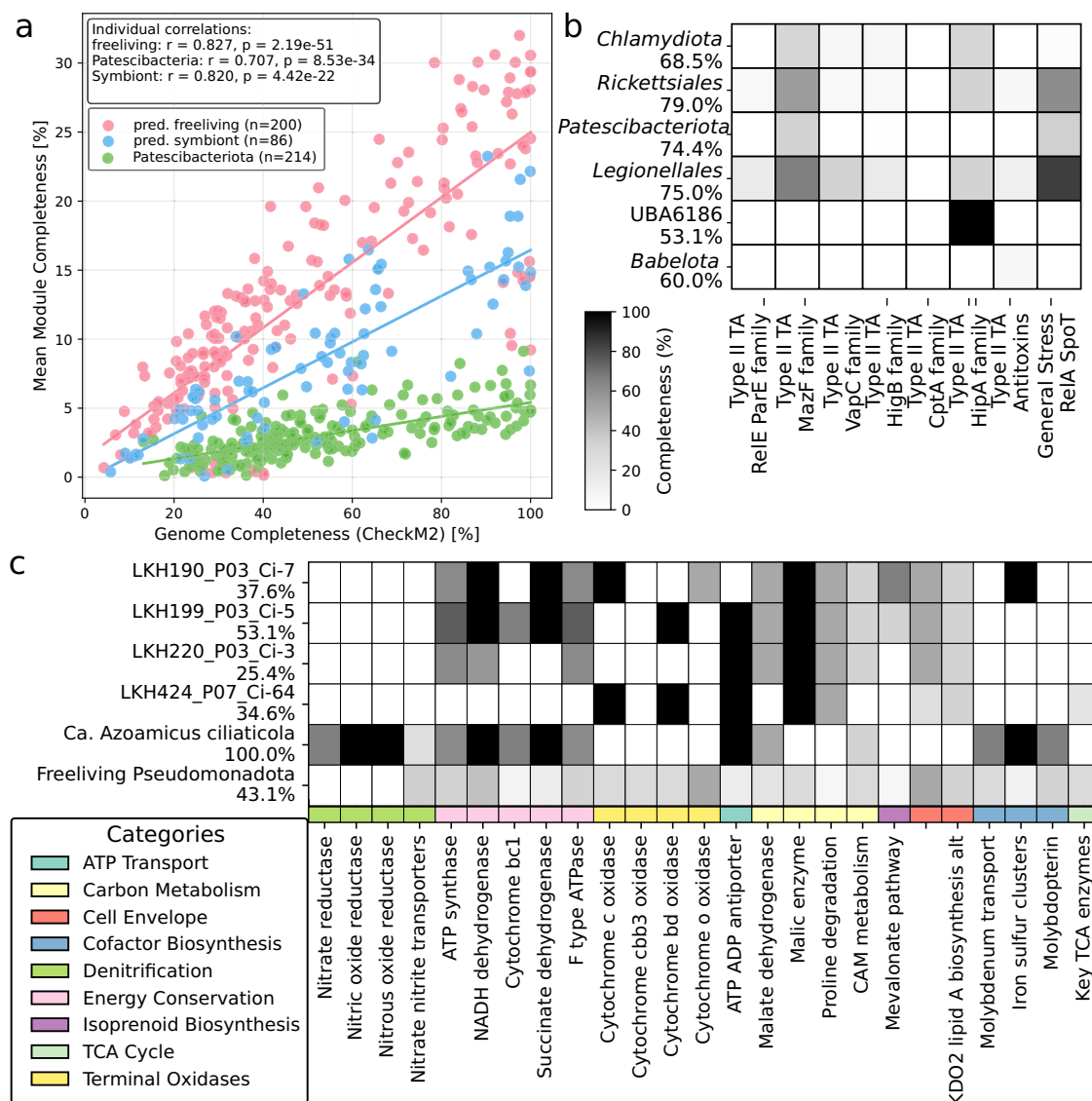


Fig. 3 | Functional and comparative genomic analyses of protist-associated bacteria. **a** Correlation between genome completeness (CheckM2) and mean metabolic module completeness for MAGs affiliated with known free-living bacteria ($n = 200$), symbiont clades ($n = 86$), and *Patescibacteriota* ($n = 214$). Each data point represents an individual MAG, with regression lines, Pearson correlation coefficients (r), and p -values (two-sided) shown for each category. **b** Heatmap showing the prevalence of Toxin-Antitoxin (TA) system gene families across major members of known and putative symbiont clades. Percentages indicate the group average of

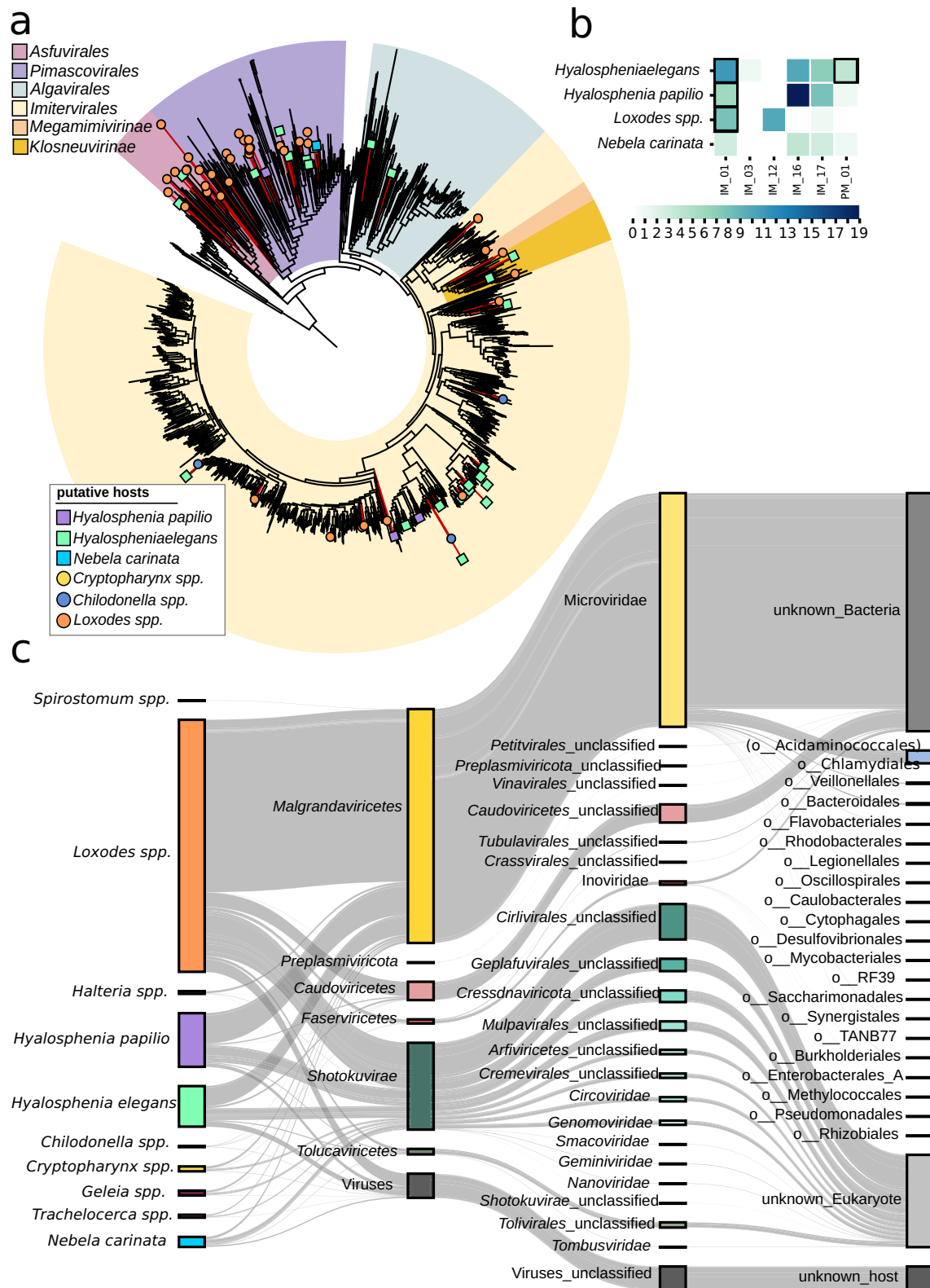
estimated genome completeness (CheckM2). Columns represent different TA gene families or related stress response genes. **c** Comparative heatmap showing the completeness of selected metabolic modules for four *Azoamicus*-related MAGs (order UBA6186) from this study, the reference genome of '*Candidatus Azoamicus ciliaticola*', and an aggregate of closest related free-living bacteria in *Pseudomonadales* and *Enterobacteriales* (freeliving *Gammaproteobacteria*). Rows represent the individual MAGs with their estimated genome completeness (CheckM2).

same samples, rather than the *Hyalosphenia papilio* itself. Further, it has been shown that giant viruses are frequently ingested as food⁴³. Such uptake may not lead to an infection in amoebae and ciliates, and viruses may accumulate in the cytoplasm, or in some cases, multiple highly similar viruses are taken up at the same time^{44,45}.

To better understand if some of the detected giant virus lineages are actively infecting the protists, we analyzed single-cell transcriptomes from similar amoebae and ciliates directly isolated from our sampling sites (Supplementary Data 1). Using these data, we were able to confirm gene expression for viruses of the *Imiterviales* family IM_01 (*Mesomimiviridae*) in several *Hyalosphenia elegans*, *Hyalosphenia papilio* and *Loxodes* cells. Further, we found genes of members of the Pimascovirales family PM_01 expressed in *Hyalosphenia elegans*. In-depth experimental assessment of the protist lineages sampled here, which, however, are challenging to maintain in the lab, will be

required to fully establish a direct connection between giant viruses and a particular protist host.

In addition to giant viruses, we were also able to recover sequences of 33 virophages from the sorted protists (Supplementary Data 4), of which 25 had a sufficient number of virophage hallmark genes to be placed into the *Lavidaviridae* taxonomic framework⁴⁶. 27 out of 33 virophage sequences were found in protist microbiomes, which also contained giant virus genomes. Virophages are known to integrate into their host genomes and become activated when the host encounters giant viruses, which they parasitize, potentially offering protection against giant virus infection⁴⁷. All virophages recovered here were on contigs with a length of 5–26 kb, which is the typical genome size range of known virophages^{48,49}, with none found on longer contigs or surrounded by protist genes. Virophage genes were not found actively expressed in metatranscriptome data from



independently sorted similar protists. Nevertheless, our findings suggest that the virophages are probably not integrated into the protist genome, but rather actively engaging in virus-virus interactions.

Protists are hot spots of DNA virus diversity

All sampled protists in our study were associated with a large number of other DNA viruses (Fig. 4c and Supplementary Data 5). Most of these

belonged to the subfamily *Gokushovirinae* from the family *Microviridae* in the order *Malgrandaviricetes*, which are ssDNA viruses that typically have small genomes and are known to infect host-associated bacteria⁵⁰. Host prediction based on iPHoP⁵¹ indicated a broad range of potential hosts, including *Legionellales*, *Coxiellales*, *Burkholderiales*, *Acidaminococcales*, and others (Fig. 4c). To a lesser extent, we found members of the order *Caudoviricetes*, which are diverse tailed dsDNA

Fig. 4 | Viral sequences found in ciliate and amoeba microbiomes.

a Phylogenetic tree of the *Nucleocytoviricota*. Red branches indicate giant virus genomes recovered in this study. Color of circles (ciliates) and squares (amoeba) at terminal branches corresponds to protist lineages. Colored wedges highlight order-level groups in the *Nucleocytoviricota*. **b** *Nucleocytoviricota* families for which genes were found to be expressed, as based on single-cell transcriptomes of four protist lineages. Numbers indicate the number of protist single cells associated with members of the same active *Nucleocytoviricota* family. Fields with a black outline

indicate pairs which were also identified in the genome data. In addition to these viral transcripts, there were others that could be assigned to viruses of the families IM_03, IM_12, IM_16 and IM17, but none of these were found in the SAG data. **(c)** Sankey diagram linking non-*Nucleocytoviricota* viral contigs of high completeness to protists and predicted hosts. Segments on the left are colored based on the protist lineages, segments in the two center columns represent viral lineages on the order and family level, segments on the right correspond to predicted hosts.



Fig. 5 | Putative multipartite association in ciliate and amoeba microbiomes, as based on SAG sequences. a The upper panel shows the distribution of different symbiont lineages, giant viruses and free-living bacteria in the individual assemblies of the sampled protists. The lower panel indicates other eukaryotes that were associated with the respective data sets. *Self-detection of protist hosts based on

18S rRNA gene screening. **b** Ranked impact of factors on uniqueness (upper heatmap) and richness (lower heatmap) of free-living bacteria, putative endosymbionts, *Patescibacteriota* and *Nucleocytoviricota*, with “6” (black) having the strongest impact and “1” (white) having the least impact on the respective factor.

viruses associated with free-living bacteria. We also identified other viruses that we could not taxonomically classify but that were predicted to infect intracellular bacteria in the order *Chlamydiales* (Fig. 4c). In addition, we found numerous ssDNA viruses from the Shotokuvirae, most of which belonged to the *Cressdnaviricota* orders *Arfiviricetes* and *Repensiviricetes*, all known to infect a wide range of eukaryotic hosts⁵². Host prediction for eukaryotic viruses is less advanced than for bacterial viruses, so it's not entirely clear which of the detected viruses may infect the protists or associated eukaryotes, or whether these viruses adhere to the protist surface or reside in the cytoplasm or food vacuoles prior to being degraded. Given the diversity and abundance of detected viruses, it's conceivable that some indeed infect protist hosts.

Complex multipartite associations in protists microbiomes

Our single cell study suggests that multipartite associations among protists, bacterial symbionts, giant viruses, and other viruses are prevalent (Fig. 5). However, for bacteria not commonly known to be obligate intracellular symbionts (e.g., many *Patescibacteriota*), the nature of the interaction, whether endosymbiotic, epibiotic, parasitic, recently ingested prey, or merely co-isolated contaminants can not be definitively determined from this dataset alone. Laboratory cultivation and imaging, such as fluorescence in situ hybridization, would provide validation. This is less of a concern for amoebae SAGs that contain sequences of giant viruses, *Chlamydia*, *Babelota* and *Gammaproteobacteria* affiliated with known intracellular bacteria. In ciliate SAGs, the co-occurrence pattern differed, and multipartite associations were mainly predicted in *Loxodes* sp., consisting of giant viruses and chlamydial and alphaproteobacterial symbionts, and to a lesser extent, *Babelota* or *Gammaproteobacteria*. There was only a single case (*Chilodonella* sp.) of a predicted multipartite association that involved three or more interaction partners. For other ciliate lineages we did not identify multiple interaction partners (Fig. 5). The overall lower

complexity of sequences from microbial symbionts and giant viruses in different ciliate samples can potentially be attributed to three factors: first, different grazing preferences compared to amoebae⁵³, second, the absence of other associated microeukaryotes in the same samples and finally the potential that some of the WGAs failed to amplify target organisms.

In contrast to the patterns for ciliates, most amoebae were associated with sequences from smaller, often flagellates such as kinetoplastids and chrysophytes. In the case of testate amoeba, washing of their shells is more difficult to achieve compared to ciliates or naked amoebae and may have hindered the complete removal of attached smaller eukaryotes. In addition, green algae were frequently detected, especially in *Hyalosphenia papilio*, which is known to be associated with endosymbiotic *Chlorella*²⁵. Notably, while chloroviruses are known to associate with *Chlorella*, we did not recover any giant viruses from *Hyalosphenia* samples containing *Chlorella*. In contrast, we sampled three *Hyalosphenia elegans* cells that were not associated with any other eukaryotes, but each associated with multiple giant viruses. Next, we tested how different factors shape the uniqueness and richness of distinct microbial and viral groups within protist microbiomes. Specifically, the sampling site was a key driver of uniqueness for free-living bacteria and giant viruses (*Nucleocytoviricota*), while host taxonomy most strongly influenced uniqueness in putative endosymbionts and *Patescibacteriota*. In contrast, microbiome richness was mainly linked to host taxonomy and sequencing depth. While it's plausible that host cell size may positively correlate with the capacity to harbor a greater diversity and abundance of associated microbes, in our analysis, morphology, cell size, and lifestyle had only moderate effects on the richness of associated taxa. These findings reinforce that both host traits (e.g., taxonomy, morphology) and environmental conditions (e.g., sampling site) collectively shape the complexity of protist-associated microbial and viral communities. Taken together, our analysis supports the notion that these protists are

sites of prevalent multipartite interactions and are tightly linked to the diversity, specificity, and ecological roles of their bacterial and viral partners.

In previous studies, protists have been identified as hosts for diverse lineages of new endosymbionts, but such findings often relied on morphological descriptions following isolation^{11,54}. Here, we used cultivation-independent sequencing approaches to sample a diversity of uncultivated protists. The hand-picking method, while essential for targeting specific uncultured protists, can not completely eliminate the possibility of co-isolating transiently associated microbes and/or environmental DNA/virions. Yet, in our more than 100 datasets, the diversity of microbes affiliated with known microbial symbionts is unparalleled, as is the diversity of identified giant viruses and other smaller viruses, such as Arfiviricetes and Repensiviricetes that infect eukaryotes and Gokushovirinae that may infect associated symbionts. The frequent co-occurrence of sequences from diverse giant viruses, with genes found to be expressed in situ, and microbial symbionts likely representing multipartite associations, is particularly intriguing. Previously, *Acanthamoeba* and ciliate species have been highlighted as evolutionary melting pots and potential training grounds for pathogens of multicellular eukaryotes^{55–57}. Our findings provide strong support for this hypothesis and call for further experimental work to study the microeukaryotic microbiome and virome. Understanding their roles in shaping protist populations and surrounding microbial communities, not only from an evolutionary perspective, but also in the ecological context of contributing to ecosystem dynamics through nutritional symbioses and pathogenicity, is crucial.

Methods

Sample collection

Individual ciliates were isolated from environmental samples obtained from a range of different sampling sites (Supplementary Data 1). Amoeba were collected in low-pH bogs and fens and washed off the moss that they inhabit using prefiltered (2 µm filter) bog water after size-selecting over a 300 µm filter to discard large plant material. Arcellinida testate amoebae were then picked under an inverted microscope from the water samples using hand-held glass pipettes. They were transferred to a microscope slide with a drop of freshly filtered bog water in an attempt to wash off obvious contamination sticking to the outside of the shell. Each individual was photo documented and then transferred to a 0.2 ml tube for transcriptome/genome amplification. We sampled ciliates either from a small low pH (pH ~4.5) pond within a local fen or from the intertidal zone of a sandy beach. Samples were filtered over an 80 µm mesh (for sandy samples) or directly poured into small Petri dishes (for pond samples). Ciliates were then observed and hand-picked with glass pipettes under an inverted microscope. Cells were washed by passing through slides of in situ water 2–3 times on depression slides to remove obvious surrounding contaminants (e.g., other non-target micro-eukaryotes, sediment particles). Each individual was diluted with nuclease-free water or prefiltered (0.2 µm filter) in situ water preceding single-cell transcriptome/genome amplification.

Whole genome and transcriptome amplification and sequencing

Whole genome amplifications of individual cells were performed using the Repli-g Single-Cell Kit (Qiagen, cat. 150345) following the manufacturer's instructions. Most samples we incubated for the recommended 8 hours, whereas for a few small ciliates with low expected DNA content (e.g., *Cryptopharynx* spp. and *Wilbertomorpha* spp.), we extended the incubation time to 10 to 12 h. Single-cell whole transcriptome amplifications were carried out using the SMART-Seq v4 Ultra Low Input RNA Kit for Sequencing (Clontech, cat. 634895, 634896) according to the manufacturer's instructions. This kit utilizes oligo(dT) priming to selectively reverse transcribe polyadenylated

(poly-A) mRNA, thereby enriching eukaryotic mRNA transcripts. Products of both genome and transcriptome amplifications were purified using the Ampure XP for PCR Purification system (Beckman Colter). A Qubit 3.0 fluorometer (Invitrogen) was used to measure the DNA concentration. Sequencing libraries for the transcriptome samples were prepared using the Nextera XT DNA Library Preparation Kit (Illumina). Library preparation for the genomic samples as well as the high-throughput sequencing of all libraries (both for genomes and transcriptomes) was carried out at the genome sequencing center at the University of California, San Diego, or at the Institute for Genome Sciences at the University of Maryland, Baltimore. A HiSeq 4000 (Illumina) sequencing platform was used for the majority of samples. The only exceptions are seven genomic samples, six from *Hyalosphenia elegans* and one from *Loxodes* sp., that were sequenced on a NovaSeq 6000 sequencing system. Samples were sequenced at 10–80 million reads per sample (Supplementary Data 1).

Sequence data quality control and assembly

The 150 bp paired-end raw sequencing reads were quality checked using FastQC (v0.11.8)⁵⁸, and the BBDMap toolkit (v38.87)⁵⁹ was used to remove adapters and trim poor-quality reads. Given that our single-cell 'omics data from uncultivated protists tend to be complex and lack reference genomes/transcriptomes, we chose to apply more stringent trimming parameters. Transcriptome reads were trimmed with "trimq=24, minlen=100" and genome reads with "trimq=28, minlen=125". Read normalization was performed with bbnorm (BBDMap toolkit v38.87) prior to the assembly. De novo assemblies were then performed using SPAdes (v3.14.1)⁶⁰ using the option -sc. Further, CrossBlock from the BBTools software package (v38.87)⁵⁹ was used with default settings to minimize the effects of cross-talk between assemblies of multiplexed libraries (Supplementary Data 6).

Metagenomic binning, bin QC, gene calling, annotation and symbiont prediction

Contigs were filtered at 2 kb length and subsequently organized into genome bins based on tetranucleotide sequence composition with MetaBat2⁶¹, yielding 6510 MAGs with an assembly size of above 20 kb. Completeness and contamination was estimated with CheckM1 (v1.1.5)⁶², CheckM2 (v1.0.2)⁶³ and an hmmsearch (v3.1b2) using a set of 56 universal single copy panorthologs (UNI56)⁶⁴ and taxonomy inferred with GTDB-tk (v1.7.0)⁶⁵. MAGs were retained if they had a CheckM1 (v1.1.5) and CheckM2 (v1.0.2) contamination of below 10% and at least 5 of UNI56 markers⁶⁴ or were classified as Nucleocytoviricota with gvcass (<https://github.com/NeLLi-team/gvcass>). Gene calling was performed with prodigal⁶⁶ using the -p meta option. Genes were then annotated with moducomp (<https://github.com/NeLLi-team/moducomp>).

Impact of different factors on uniqueness and richness of the protist microbiome

We first grouped MAGs by library and calculated richness (unique genera per library) and uniqueness (taxa exclusive to each library within the dataset). Sequencing depth was categorized (<5 Gb = "low", 5–20 Gb = "medium", 20–100 Gb = "high", >100 Gb = "very high"), cell size was categorized (<100 micron = "small", 100–300 micron = "medium", >300 micron = "large"), lifestyle (mixotrophy, heterotrophy), other features ("shell", "microanaerobic", "shell and algae symbionts", "other") and used to normalize both richness and uniqueness. We then fit separate Ordinary Least Squares models for normalized richness and uniqueness, incorporating the factors host taxonomy, cell size, other features, sampling site, sequencing depth, and lifestyle as categorical predictors. Type II ANOVA was applied to each model, and partial eta-squared values were used to estimate each predictor's effect size. This approach allowed us to rank factors by their influence on microbial diversity metrics.

Screening for viruses and viral host prediction

To identify any viral sequences within the protist single-cell sequence data, geNomad (v1.6.0)⁶⁷ was used with default settings. Contigs that were predicted as viral were then subject to completeness and contamination estimate with CheckV (v1.02)⁶⁸. Host prediction was performed with iPhoP (v1.3.3)⁵¹ on contigs that were predicted as viral genomes with high completeness and no contamination. To identify giant virus metagenome assembled genomes (GVMAGs) and unbinned contigs that had a length of at least 50 kb were subject to classification with gvcass (v0.9) (<https://github.com/NeLLi-team/gvcass>)⁴⁰. In brief, 9 conserved giant virus orthologous groups (GVOGs)⁶⁹ were identified using hmmsearch, extracted, and used as a query for a diamond blastp search (v2.1.3)⁷⁰ against a database of the respective GVOG built from a representative set of bacteria, archaea, eukaryotes and viruses. The top 100 blastp hits were extracted, combined with the query sequence, aligned with mafft (v7.490; -linsi)⁷¹, trimmed with trimal (v1.4; -gt 0.1)⁷² and used to build a phylogenetic tree with IQtree (v2.3.0; LG4X)⁷³. The nearest neighbor in the tree was identified using branch length, and the existing taxonomic string for that reference genome was then taken into account for the final classification result. For a successful classification, the taxonomic strings from all identified nearest neighbors were compared at the different taxonomic levels (genus, family, order, class, phylum) to yield the final classification at the lowest taxonomic level on which all nearest neighbors were in agreement.

Eukaryotic phylogenomics

To reconstruct a phylogenetic tree showing the position of our focal taxa within the eukaryotic tree of life, we used our phylogenomic pipeline PhyloToL⁷⁴. Gene trees for 391 gene families that are highly conserved across eukaryotes and present in at least four out of five major eukaryotic clades were produced. We chose 250 taxa from all major eukaryotic clades as well as bacteria and archaea, with an even distribution of 25 taxa per major clade. In addition, one representative of each of our focal species was added. PhyloToL produces multi-sequence alignments using Guidance v2.0⁷⁵ and builds gene trees using RAXML (PROTGAMMALG)⁷⁶. In addition, we generated a supermatrix using the alignment concatenation option of PhyloToL and inferred a species tree with RAXML(v8) with PROTGAMMALG.

Bacterial and viral phylogenomics

An initial bacterial species tree was built with New Simple Genome Tree (nsgtree v.0.4.0, <https://github.com/NeLLi-team/nsgtree>) from all MAGs that had at least 5/56 single copy marker genes of the UNIS6 set of markers⁶⁴ together with a representative set of genomes from the GTDB database⁶⁹. GVOGs were identified with hmmsearch (v.3.3.2), aligned with mafft(v.7.508)⁷¹ and trimmed with trimal (v1.4)⁷². Clades of known symbionts were then selected and extracted from the tree, additional genomes (1 per family) were added to the selected clades and separate species trees were built.

To infer a species tree for the Nucleocytoviricota, GVMAGs and a reference dataset of published giant virus genomes⁶⁹ were combined, and nsgtree was employed with the set of phylogenetic markers GVOG8 (GVOGm0013, GVOGm0022, GVOGm0023, GVOGm0054, GVOGm0172, GVOGm0461, GVOGm0760, GVOGm0890)⁶⁹; genomes that had less than 4 out of 8 GVOGs were removed. GVOGs were identified with hmmsearch (v.3.3.2), aligned with mafft(v.7.508)⁷¹ and trimmed with trimal (v1.4)⁷². Only GVMAGs and reference giant virus genomes with at least four of the seven GVOGs and with no more than four copies of any of the GVOG8 were selected, totaling 82 GVMAGs with assembly sizes ranging from 28 kb to 1.43 Mb. The final tree was calculated with IQtree (v.2.1.11)⁷³ and visualized with iTOL (v.6)⁷⁷.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

Sequence data that support the findings of this study have been deposited in the Sequence Read Archive with the accession codes outlined in Supplementary Data 1. Metagenome assemblies, MAGs, annotations, alignments and phylogenetic trees that support the findings of this study can be accessed and directly downloaded at the National Energy Research Scientific Computing Center (NERSC) [https://portal.nersc.gov/cfs/nelli/ms_protistsymb/]. [https://portal.nersc.gov/cfs/nelli/ms_protistsymb/source_data.tar.gz]. Source data are provided in this paper.

Code availability

All code underlying the analyses for each figure has been deposited as custom scripts, Jupyter notebooks and a DuckDB with example queries to extract sequences or annotations, and can be downloaded as source data at NERSC Final code in Jupyter notebooks was organized and polished with Claude Code (Anthropic).

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Author contributions

Y.Y., A.W. and L.K. collected and processed the samples, identified protists and prepared them for sequencing. A.W. and L.K. provided a phylogenetic tree of sampled eukaryotes. F.S. and R.A. performed data analysis, and F.S. generated the figures. T.W. supervised research. F.S. and T.W. prepared the manuscript, with contributions from all authors. All authors read and approved the final manuscript.

Competing interests

F.S. serves as CEO of SampleX. This work was not funded by, nor does it benefit, SampleX. All other authors declare no competing interests.

Additional information

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