Combined Genome and Transcriptome Analyses of the Ciliate *Schmidingerella arcuata* (Spirotrichea) Reveal Patterns of DNA Elimination, Scrambling, and Inversion

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**Recommended Citation**

Smith, Susan A.; Maurer-Alcalá, Xyrus X.; Yan, Ying; Katz, Laura A.; Santoferrara, Luciana F.; and McManus, George B., "Combined Genome and Transcriptome Analyses of the Ciliate *Schmidingerella arcuata* (Spirotrichea) Reveal Patterns of DNA Elimination, Scrambling, and Inversion" (2020). Biological Sciences: Faculty Publications, Smith College, Northampton, MA.  
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Combined Genome and Transcriptome Analyses of the Ciliate *Schmidingerella arcuata* (Spirotrichea) Reveal Patterns of DNA Elimination, Scrambling, and Inversion

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Accepted: 24 August 2020

Data deposition: This project has been deposited at NCBI under the Bioproject ID: PRJNA626068 and the accession numbers SAMN14615117 (MIC genome), SAMN14615214 (MAC transcriptome), and SRR11933493 (raw reads); transcriptome annotation is available on figshare at the link: https://doi.org/10.6084/m9.figshare.12686621.

Abstract

*Schmidingerella arcuata* is an ecologically important tintinnid ciliate that has long served as a model species in plankton trophic ecology. We present a partial micronuclear genome and macronuclear transcriptome resource for *S. arcuata*, acquired using single-cell techniques, and we report on pilot analyses including functional annotation and genome architecture. Our analysis shows major fragmentation, elimination, and scrambling in the micronuclear genome of *S. arcuata*. This work introduces a new nonmodel genome resource for the study of ciliate ecology and genomic biology and provides a detailed functional counterpart to ecological research on *S. arcuata*.

Key words: genome architecture, single-cell ‘omics, tintinnid, micronucleus, macronucleus, Ciliophora.

Significance

Our understanding of genome organization in nonmodel ciliates is limited because 1) most species are uncultivable and 2) it requires the separate amplification of a ciliate’s two nuclei (the germline micronucleus and the somatic macronucleus), which is technically difficult. By using single-cell ‘omics, we were able to separately sequence both genomes of the ciliate *Schmidingerella arcuata*, which revealed patterns of extensive genome rearrangement and fragmentation, and also allowed us to analyze functional details of its somatic genome. This research contributes information on the genomic architecture and functional of an ecologically important ciliate and expands our understanding of ciliate genomics beyond model species.

Introduction

Ciliates are an ancient and diverse clade of microbial eukaryotes that inhabit nearly every environment on Earth. They have long served as models in the research of cell biology and recently have become an ideal system for the study of genome fragmentation and organization (Greider and Blackburn 1985; Lynn 2008; Parfrey et al. 2011). Although genome rearrangements have been discovered throughout the eukaryotes, the phenomenon appears to be especially...
elaborate within ciliates, which can exhibit the mass elimination, fragmentation, and scrambling of loci (Landweber et al. 2000; Prescott 2000; Chalker 2008; Bracht et al. 2013; Chen et al. 2014; Gao et al. 2015; Maurer-Alcalá, Knight, et al. 2018).

The complexity of ciliate genome architecture results from their nuclear dimorphism, a unique segregation of germline and somatic functions into separate nuclei (Lynn 2008). Chromosomal rearrangements can occur between the germline-limited micronuclear (MIC) and the transcriptionally active macronuclear (MAC). Following conjugation, the zygotic nucleus divides to form a new MIC and a new MAC. In creating the new MAC, a series of rearrangements and deletions can occur, including the extensive elimination of MIC segments called internally eliminated sequences (IESs) (Gratias and Bétermin 2001; Riley and Katz 2001; Katz et al. 2003; Fass et al. 2011). Regions that are not eliminated (i.e., present in both MIC and MAC) are called macronuclear-destined sequences (MDSSs) and are cut and pieced together to form MAC loci (Swart et al. 2013) (fig. 1A). In some ciliates, MDSS regions are arranged out of order in the MIC, a complex pattern of genomic architecture called “scrambling.” Scrambled loci can additionally be “inverted” if they are transcribed on opposing strands of the MIC scaffolds (fig. 1A).

As most ciliates are not amenable to culture, research on ciliate genomics and nuclear architecture has mostly been limited to a few model species (e.g., Paramecium and Tetrahymena) (Hamilton et al. 2016). In addition, traditional sequencing methods are biased toward highly amplified MAC regions, which has made it challenging to isolate MIC regions. However, the process of multiple-displacement amplification used in single-cell genomics is biochemically biased for long-template DNA (2–70 kb), which allows for the selection of MIC chromosomes (Spits et al. 2006). Combined with single-cell transcriptomics, we are now able to elucidate the patterns of genome rearrangement, elimination, and scrambling, all from a single cell (Maurer-Alcalá, Knight, et al. 2018).

Here, we use single-cell ’omics (genomics and transcriptomics) to study the MIC genome and the transcriptome (a proxy for the gene-sized chromosomes of the macronuclear genome) of S. arcuata, a marine ciliate (class Spirotrichea, order Tintinnida). Long used as an ecological model in planktonic food web studies, S. arcuata is ubiquitous in coastal waters, where it periodically dominates the ciliate community (Agatha and Strüder-Kypke 2012; Dolan and Pierce 2013; Santoferrara et al. 2018). Schmidingerella arcuata is also one of the few marine ciliates amenable to culture and thus represents a ciliate that is both ecologically relevant and cultivable (Dolan 2012; Montagnes 2013; Echevarria et al. 2016; Jung et al. 2016; Cobb 2017; Gruber et al. 2019). Although tintinnids have a long history of taxonomic study (Müller 1779; Haeckel 1866), there exists no published data on their MIC or genomic architecture, and only limited transcriptome data exist for Schmidingerella (Keeling et al. 2014), the only tintinnid genus with transcriptome data. Here, we present a genome and transcriptome resource for S. arcuata, acquired using single-cell techniques, and we report on pilot analyses of its genome architecture and transcriptome. This represents a new resource for the study of ciliate genomic architecture and provides a detailed genomic counterpart to ecological research on this model microzooplankton.

Materials and Methods

Culturing

Schmidingerella arcuata was collected from the surface waters of northeastern Long Island Sound, CT (41.31°N, 72.06°W), using a 20-μm-mesh plankton net. Single cells were isolated with drawn capillaries and moved to six-well culture plates with 0.2-μm-filtered sample water. Clonal cultures of S. arcuata were fed saturating concentrations (~3 × 10^3 cells/ml) of the dinoflagellate Heterocapsa triquetra and the prymnesiophyte Isochrysis galbana (strain TISO). Cultures were kept at 18°C under a 12:12 h light:dark cycle. Morphology and 18S rDNA sequences (see Santoferrara et al. 2013) confirmed the taxonomic identification of S. arcuata (Agatha and Strüder-Kypke 2012).

Isolation of Single Cells

Individuals were transferred from growing cultures to autoclaved, 0.2-μm-filtered seawater and starved for 12 h to ensure the clearance and digestion of prey. The cells were then picked and rinsed a minimum of five times in autoclaved, 0.2-μm-filtered seawater using drawn capillaries under a stereo microscope. Each cell was then transferred into the appropriate buffer for transcriptome or genome sequencing and brought to volume with nuclease-free water (as specified in the kits detailed below).

Single-Cell Transcriptome and Genome Amplification

The SMART-Seq2 v4 Ultra Low input RNA kit (Cat: 634889; Takara, Mountain View, CA) was used for whole-transcriptome amplification (WTA) following manufacturer’s protocols, with the exception that we quartered the reaction volumes. For whole-genome amplification (WGA), the Repli-g single-cell kit (Cat: 150343; Qiagen, Hilden, Germany) was used following manufacturer’s protocols. The products (cDNA for WTA, gDNA for WGA) were quantified with the dsDNA Qubit assay (Invitrogen, Waltham, MA) and polymerase chain reaction-checked with eukaryotic 18S rDNA (Medlin et al. 1988) and genus-specific ITS (Costas et al. 2007) primers. Minimum bacterial contamination was confirmed by polymerase chain reaction with 16S rDNA primers (Lane et al. 2000; Prescott 2000; Chalker 2008; Bracht et al. 2013; Chen et al. 2014; Gao et al. 2015; Maurer-Alcalá, Knight, et al. 2018).
Fig. 1—Genome architecture in *Schmidingerella arcuata*. (A) Patterns of nuclear architecture in the germline MIC and somatic MAC. The MIC contains the required MDSs for the generation of functional genes during development of the MAC. MDSs can be interrupted by IESs. Development of the MAC requires the precise excision of IESs and the correct rearrangement of MDS regions. MDS loci may be scrambled (e.g., MDS 1–4), inverted (e.g., MDS 4), or...
1991). Sequencing libraries were prepared with the Illumina Nextera XT kit (Cat: FC1311096; Illumina, San Diego, CA) then processed with Illumina HiSeq 2500 at Macrogen Sequencing (Geumcheon-gu, Seoul, South Korea).

Transcriptome and Genome Assembly
Raw reads from WTA and WGA sequencing were trimmed for quality and size (Q28 and minimum length of 200 and 1,500 bp, respectively) using BBMap (V38.39; Bushnell 2014) After trimming, two single-cell WTAs were assembled together using maSPades (V3.13.1, Bankevich et al. 2012), and seven single-cell WGAs were assembled together using both SPAdes (V3.13.1) and MEGAHIT (V1.2.9; Li et al. 2015). The MEGAHIT genome assembly was used for the final analysis because it yielded a higher mapping continuity (i.e., the amount of transcripts mapped to the WGA assembly per kilobase). Assemblies were processed through custom python scripts (http://github.com/maurerax/KatzLab/tree/HTS-Processing-PhyloGenPipeline) for the removal of rDNA and prokaryotic transcripts, and for the identification of orthologous gene families using USEARCH (V9.2; Edgar 2010) with OrthoMCL databases (V2.0.9; Fischer et al. 2011) (Maurer-Alcalá, Knight, et al. 2018). Additional steps included the prediction of open reading frames with AUGUSTUS (Hoff and Stanke 2019) using an Escherichia coli model to eliminate bacterial contaminants. Stop-codon usage was determined using a custom Python script, which quantified the frequency of in-frame occurrences of TAG/TGA/ATAA when each codon was used as a termination site. The completeness of the MIC genome assembly was analyzed using Benchmarking Universal Single-Copy Ortholog (BUSCO; Waterhouse et al. 2018) (E-value < 10^{-3}, alveolate lineage database). OmicsBox (V5.2.5; Götz et al. 2008) was used with InterProScan (V5.42; Hunter et al. 2009) and BlastX (NCBI nonredundant database, E-value < 10^{-4}; V2.8.1; Altschul et al. 1990) for functional annotation and for the identification of gene ontology (GO) terms involved in Kyoto Encyclopedia of Genes and Genomes (Kanehisa and Goto 2000) pathways.

Genome Architecture Analysis
Custom Python scripts were used to identify and organize the genome architecture of S. arcuata (Maurer-Alcalá, Knight, et al. 2018). Putative MIC loci for MAC MDSs were identified via mapping of the MAC transcriptome to the MIC genome sequences using BLAST. For transcripts to be considered “MIC mapped,” at least 60% of their length was required to be mapped to the MIC (length threshold as suggested in Maurer-Alcalá, Knight, et al. 2018). Loci were classified into categories of “unmapped,” “nonscrambled,” and “scrambled.” MDS–IES borders were required to be flanked by pairs of short (2–10 bp) tandem repeats called pointer regions to discriminate them from possible intron-exon boundaries (Bracht et al. 2013) (fig. 1B and C). The GC content of MDS–IES boundaries was determined by evaluating the 40 bp located at both ends (i.e., the 5’ and 3’) of an MDS in the MIC.

Results and Discussion
Genome and Transcriptome Resources
Assemblies for the MIC and MAC of S. arcuata are about 49 and 6 Mb in size, respectively (table 1). Of the 11,673 transcripts, which are a proxy for the gene-sized MAC chromosomes of this species, roughly 15% (1,712) were mapped to the MIC. BUSCO analyses estimate that the MIC genome resource is about 19% complete (Complete: 18.8%, Fragmented: 7.6%, Missing: 73.6%, n: 171). This indicates the need for deeper sequencing, although BUSCO analyses have been found to underestimate highly fragmented genomes (López-Escardó et al. 2017) and thus may not be a reliable indicator of completeness in S. arcuata. About 80% of MAC transcripts have significant BlastX hits (NCBI nonredundant database, E-value < 10^{-4}), and of those, 74% have a confident assignment of GO terms. The majority of MAC transcripts related to cellular components correspond to membrane and organelle activity, whereas catalytic and binding activities are the primary molecular functions, and localization and biological regulation comprise the main biological terms (supplementary fig. S1, Supplementary Material online). About 15% of MAC transcripts were placed in Kyoto Encyclopedia of Genes and Genomes pathways. The three primary pathways identified are thiamine metabolism, purine metabolism, and aminoaacyl-tRNA biosynthesis (supplementary fig. S1, Supplementary Material online). Annotation details regarding the MAC transcriptome can be found in supplementary data, Supplementary Material online, and full annotation.
files can be found at figshare at the link: https://doi.org/10.6084/m9.figshare.12686621.

We assessed stop-codon usage and found that the codons TGA and TAA were rarely found in-frame, and their usage in S. arcuata matched homologs in the Oxytricha trifallax transcriptome (GenBank BioSample SAMN02953822). The combination of TGA/TAA as stop codons is not observed in the few published transcriptomes for this genus (Keeling et al. 2014; Heaphy 2018) or for other Spirotrichs, with TAA/TAG reported for Euplotes crassus and TGA for Oxytricha trifallax and Stylonychia lemnæa (Kervestin et al. 2001; Lozupone et al. 2001; Swart et al. 2013; Heaphy 2018; Yan et al. 2019). However, ciliates have frequent stop-codon reassignments, and even context-dependent stop codons (Adachi and Cavalcanti 2009; Heaphy et al. 2016; Yan et al. 2019).

Patterns of Genome Architecture

Schmidingerella arcuata shows extensive genome fragmentation, including the unscrambling and inversion of loci during MAC formation (fig. 1B). We considered MIC loci as scrambled if their associated transcripts mapped to MDSs that were out of consecutive order in the MIC, and if those MDS–IES boundaries contained pointer regions (Maurer-Alcalá, Knight, et al. 2018). Scrambled loci were determined to be inverted if nonconsecutive MDSs appeared on both strands of germline scaffolds (fig. 1A and B). Of the MIC-mapped transcripts, roughly 36% were found to be scrambled. The average GC content at MDS–IES boundaries for S. arcuata (47.5%) was slightly higher than the overall GC for all germline-supported scaffolds (46.4%); this increase also occurs in other ciliate classes, although most report a sharper rise (10–14%) in %GC around these regions (Maurer-Alcalá, Knight, et al. 2018). Pointer sequence size was variable within and among MDS groups (fig. 1A), with a range of 2–10 bp. We found no evidence for alternative processing (more than one MAC sequence resulting from a single MIC region; Katz and Kovner 2010).

Variations in Genome Architecture within and among Ciliate Classes

In general, the micronuclear arrangement of housekeeping genes in S. arcuata matched that of O. trifallax. However, we detected a major housekeeping gene with variable micronuclear organization among six ciliates with available data (fig. 1C). In this example, the beta-tubulin gene (paralog 1) is separated into two similar-sized MDSs in the MIC of S. arcuata, interrupted by a single IES and connected by an 8 bp TC-iterative pointer sequence (fig. 1C). In contrast, other ciliates of the class Spirotrichea (O. trifallax and S. lemnæa) separate the paralog into three or four MDSs, with comparatively shorter IES regions. Model ciliates of the Oligohymenophorea class (Paramecium caudatum and Tetrahymena thermophila) contain this paralog as either three MDS regions of variable size or an uninterrupted sequence in the MIC (Dupuis 1992; Libusová and Drábek 2006). The Phyllopharyngean ciliate C. unicinata divides the MIC gene into three consecutive MDSs of variable size (75–600 bp each) with two 6–7-bp pointer sequences (Harper and Jahn 1989; Zuñiga and Katz 2007; Katz and Kovner 2010).

Significance of S. arcuata-Omics Resources

This work contributes a MIC genome and MAC transcriptome resource for the ecologically important ciliate S. arcuata. Single-cell omics allowed selective amplification the MIC and MAC, which revealed genomic scrambling, elimination, and inversion in S. arcuata. This study provides a nonmodel genome and transcriptome resource to a field represented mostly by model ciliates. The included annotation details are a valuable resource for future ecological research on S. arcuata and closely related ciliates, which are currently underrepresented in detailed genome-scale analyses. Additionally, research on nonmodel ciliates are beginning to reveal the significance of genome architecture in molecular evolution (Maurer-Alcalá, Ying, et al. 2018; Maurer-Alcalá and Nowacki 2019; Yan et al. 2019). Recent models indicate that slight differences in IESs and specific architectural patterns (e.g., alternative processing and scrambling) among intraspecific ciliate populations can cause rapid incompatibility, potentially leading to incipient speciation (Katz and Kovner 2010; Goldman and Landweber 2012; Gao et al. 2015; Yan et al. 2019). These slight errors in the rearrangement of loci could theoretically accumulate more frequently than (non-neutral) point mutations, which may help to explain the large disparity between the molecular and morphological diversity in ciliates (Gao et al. 2015).
Supplementary Material

Supplementary data are available at Genome Biology and Evolution online.

Acknowledgments

This research was supported by the United States National Science Foundation (awards OCE1924572 to G.B.M. and L.F.S.; OCE-1924570 and DEB-1541511 to L.A.K.), Smith College, and the University of Connecticut’s Center for Genome Innovation and Computational Biology Core.

Literature Cited

Adachi M, Cavalcanti AR. 2009. Tandem stop codons in ciliates that reassign stop codons. J Mol Evol. 68(4):424–431.
Agatha S, Strüder-Kypke MC. 2012. Reconciling cladistic and genetic analyses in chooreotrich ciliates (Ciliophora, Sprotrichia, Olgoirotieha). J Eukaryotic Microbiol. 59(4):325–350.
Altschul SF, et al. 1990. Basic local alignment search tool. J Mol Biol. 215(3):403–410.
Bankevich A, et al. 2012. SPAdes: a new genome assembly algorithm and its applications to single-cell sequencing. J Comput Biol. 19(5):455–477.
Bracht JR, et al. 2013. Genomes on the edge: programmed genome instability in ciliates. Cell 152(3):406–416.
Bushnell B. 2014. BBMap: a fast, accurate, splice-aware aligner. Lawrence Berkeley National Lab. (LBNL). Berkeley (CA): OSTI.
Chalker DL. 2008. Dynamic nuclear reorganization during genome remodeling of Tetrahymena. Biochim Biophys Acta Mol Cell Res. 1783(11):2130–2136.
Chen X, et al. 2014. The architecture of a scrambled genome reveals massive levels of genomic rearrangement during development. Cell 158(5):1187–1198.
Cobb S. 2017. Investigation of contact-based cues mediating food uptake in the marine tintinnid ciliate Favella sp. [master’s thesis]. [Chico (CA)]: California State University.
Costas BA, McManus G, Doherty M, Katz LA. 2007. Use of species-specific primers and PCR to measure the distributions of planktonic ciliates in coastal waters. Limnol Oceanogr Methods. 5(6):162–173.
Dolan JR. 2012. Tintinnid ciliates: an introduction and overview. In: Dolan JR, Agatha S, Coats DW, Montagnes DJ, Stoecker DK, editors. Biology and ecology of tintinnid ciliates: models for marine plankton. Oxford: Wiley-Blackwell. p. 1–16.
Dolan JR, Pierce RW. 2013. Diversity and distributions of tintinnids. In: Dolan JR, Agatha S, Coats DW, Montagnes DJ, Stoecker DK, editors. Biology and ecology of tintinnid ciliates: models for marine plankton. Oxford: Wiley-Blackwell. p. 214–243.
Dupuis P. 1992. The beta-tubulin genes of Paramecium are interrupted by two 27 bp introns. EMBO J. 11(10):3713–3719.
Echavarria ML, Wolfe G, Strom S, Taylor AR. 2014. Connecting alveolate cell biology with trophic ecology in the marine plankton using the ciliate Favella as a model. FEMS Microbiol Ecol. 90(1):18–38.
Echavarria ML, Wolfe GV, Taylor AR. 2016. Feast or flee: bioelectrical regulation of feeding and predator evasion behaviors in the planktonic alveolate Favella sp. (Sprotrichia). J Exp Biol. 219(3):445–456.
Edgar RC. 2010. Search and clustering orders of magnitude faster than BLAST. Bioinformatics 26(19):2460–2461.
Fass JN, et al. 2011. Genome-scale analysis of programmed DNA elimination sites in Tetrahymena thermophila. G3 (Bethesda) 1(6):515–522.
Fischer S, et al. 2011. Using OrthoMCL to assign proteins to OrthoMCL-D1 groups or to cluster proteomes into new ortholog groups. Curr Protoc Bioinf. 35(1):6–12.
Gao F, Roy SW, Katz LA. 2015. Analyses of alternatively processed genes in ciliates provide insights into the origins of scrambled genomes and may provide a mechanism for speciation. MBio 6(1):1–8.
Goldman AD, Landweber LF. 2012. Oxytricha as a modern analog of ancient genome evolution. Trends Genet. 28(8):382–388.
Götz S, et al. 2008. High-throughput functional annotation and data mining with the Blast2GO suite. Nucleic Acids Res. 36(10):3420–3435.
Gratias A, Bètémier M. 2001. Developmentally programmed excision of internal DNA sequences in Paramecium aurelia. Biochimie 83(11–12):1009–1022.
Greider CW, Blackburn EH. 1985. Identification of a specific telomere terminal transferase activity in Tetrahymena extracts. Cell 43(2):405–413.
Gruber MS, Mühlthaler A, Agatha S. 2019. Ultrastructural studies on a model tintinnid—Schmidingerella meunieri (Kofoid and Campbell, 1929) Agatha and Strüder-Kypke, 2012 (Ciliophora). I. Somatic kineidits with unique ultrastructure. Acta Protozool. 57(3):195–214.
Haeckel E. 1866. VII. Character des Protistenreiches (VII. Character of the kingdom of Protists). In: Haeckel E, Reimer G, editors. Principen der generellen Morphologie der Organismen: wörtlicher Abdruck eines Teiles der 1866 erschienenen generellen Morphologie (Allgemeine Grundzüge der organischen Formen-Wissenschaft, mechanisch begründet durch die von Charles Darwin reformierte Deszendenz-Theorie). Berlin: Georg Reimer. p. 215.
Hamilton EP, et al. 2016. Structure of the germline genome of Tetrahymena thermophila and relationship to the massively rearranged somatic genome. Elife 5:e19090.
Harper DS, Jahn CL. 1989. Actin, tubulin and H4 histone genes in three species of hypotrichous ciliated protozoa. Gene 75(1):93–107.
Heaphy SM. 2018. Recoding and reassignment in protists [doctoral dissertation]. [Cork (United Kingdom)]: University College Cork.
Heaphy SM, et al. 2016. Novel ciliate genetic code variants including the reassignment of all three stop codons to sense codons in Condylostoma magnum. Mol Biol Evol. 33(11):2885–2889.
Hoff KJ, Stanke M. 2019. Predicting genes in single genomes with AUGUSTUS. Curr Protoc Bioinf. 65(1):e57.
Hunter S, et al. 2009. InterPro: the integrative protein signature database. Nucleic Acids Res. 37(Database):D211–D215.
Jahn CL, Klobutscker LA. 2002. Genome remodeling in ciliated protozoa. Annu Rev Microbiol. 56(1):489–520.
Jung JH, Choi JM, Coats DW, Kim YO. 2016. Eudubosquella costata n. sp. (Dinoflagellata, Syndinea), an intracellular parasite of the ciliate Schmidingerella arcuata: morphology, molecular phylogeny, life cycle, prevalence, and infection intensity. J Eukaryotic Microbiol. 63(3):3–15.
Kanehisa M, Goto S. 2000. KEGG: Kyoto Encyclopedia of Genes and Genomes. Nucleic Acids Res. 28(1):27–30.
Katz LA, Kovner AM. 2010. Alternative processing of scrambled genes generates protein diversity in the ciliate Chilodonella uncinata. J Exp Zool Part B 314B(6):480–488.
Katz LA, Lasek-Nesselquist E, Shoeybenos-West OL. 2003. Structure of the micronuclear alpha-tubulin gene in the phyllopharyngean ciliate Chilodonella uncinata: implications for the evolution of chromosomal processing. Gene 315:15–19.
Keehn PJ, et al. 2014. The Marine Microbial Eukaryote Transcriptome Sequencing Project (MMETSP): Illuminating the Functional Diversity of Eukaryotic Life in the Oceans through Transcriptome Sequencing. PLoS Biol. 12(6):e1001889.
Kervestin S, Frolova I, Kisselev L, Jean-Jean O. 2001. Stop codon recognition in ciliates: Euplotes release factor does not respond to reassigned UGA codon. EMBO Rep. 2(8):680–684.
Landweber LF, Kuo TC, Curtis EA. 2000. Evolution and assembly of an extremely scrambled gene. Proc Natl Acad Sci U S A. 97(7):3298–3303.
Lane DJ. 1991. 16S/23S rRNA sequencing. In: Stackebrandt E, Goodfellow M, editors. Nucleic acid techniques in bacterial systematics. New York: John Wiley & Sons. p. 115–175.

Li D, Liu CM, Luo R, Sadakane K, Lam TW. 2015. MEGAHIT: an ultra-fast single-node solution for large and complex metagenomics assembly via succinct de Bruijn graph. Bioinformatics 31(10):1674–1676.

Lisovský L, Dráber P. 2006. Multiple tubulin forms in ciliated protozoan Tetrahymena and Paramecium species. Protoplasma 227(2–4):65–76.

López-Escarló D, et al. 2017. Evaluation of single-cell genomics to address evolutionary questions using three SAGs of the choanoflagellate Monosiga brevicollis. Sci Rep. (1):711025.

Lozupone CA, Knight RD, Landweber LF. 2001. The molecular basis of nuclear genetic code change in ciliates. Curr Biol. 11(2):65–74.

Lynn DH. 2008. The ciliated protozoa. Springer. Orig. Published 1979. New York: Pergamon Press. p. 65–199.

Maurer-Alcalá XX, Knight R, Katz LA. 2018. Exploration of the germline genome of the ciliate Chilodonella uncinita through single-cell omics (transcriptomics and genomics). mBio. 9(1):1–9.

Maurer-Alcalá XX, Nowacki M. 2019. Evolutionary origins and impacts of genome architecture in ciliates. Ann N Y Acad Sci. 1447(1):110–118.

Maurer-Alcalá XX, Ying Y, et al. 2018. Twisted tales: insights into genome diversity of ciliates using single-cell ‘omics. Genome Biol Evol. 10(8):1927–1938.

Medlin L, Elwood HJ, Stickel S, Sogin ML. 1988. The characterization of enzymatically amplified eukaryotic 16S-like rRNA-coding regions. Gene 71(2):491–499.

Montagnes DJS. 2013. Ecophysiology and behaviour of tintinnids. In: Dolan JR, Agatha S, Coats DW, Montagnes DJS, Stoecker DK, editors. Biology and ecology of tintinnid ciliates: models for marine plankton. Oxford: Wiley-Blackwell. p. 86–122.

Montagnes DJS, et al. 2008. Selective feeding behaviour of key free-living protists: avenues for continued study. Aquat Microb Ecol. 53(1):83–98.

Müller OF. 1779. Zoologia Danica seu animalium Daniae et Norvegiae rarorum ac minus notorum descriptiones et historia 1. Weygandinis, Havniae et Lipsiae.

Parfrey LW, Lahr DJ, Knoll AH, Katz LA. 2011. Estimating the timing of early eukaryotic diversification with multigene molecular clocks. Proc Natl Acad Sci U S A. 108(33):13624–13629.

Pederson T. 2010. An Olympian protozoan. Nucleus 1(1):2–3.

Prescott DM. 2000. Genome gymnastics: unique modes of DNA evolution and processing in ciliates. Nat Rev Genet. 1(3):191–198.

Riley JL, Katz LA. 2001. Widespread distribution of extensive genome fragmentation in ciliates. Mol Biol Evol. 18(7):1372–1377.

Santoferrara LF, McManus GB, Alder VA. 2013. Utility of genetic markers and morphology for species discrimination within the order Tintinnida (Ciliiphora, Spirotrichea). Protist 164(1):24–36.

Santoferrara LF, Rubin E, McManus GB. 2018. Global and local DNA (metabarcoding reveal new biogeography patterns in tintinnid ciliates. J Plankton Res. 40(3):209–221.

Spits C, et al. 2006. Whole-genome multiple displacement amplification from single cells. Nat Protoc. 1(4):1965–1970.

Swart EC, et al. 2013. The Oxytricha trifallax macronuclear genome: a complex eukaryotic genome with 16,000 tiny chromosomes. PLoS Biol. 11(1):e1001473.

Wang Y, et al. 2017. A comparative study of genome organization and epigenetic mechanisms in model ciliates, with an emphasis on Tetrahymena, Paramecium and Oxytricha. Eur J Protistol. 61:376–387.

Waterhouse RM, et al. 2018. BUSCO applications from quality assessments to gene prediction and phylogenomics. Mol Biol Evol. 35(3):543–548.

Yan Y, Maurer-Alcalá XX, Knight R, Pond SLK, Katz LA. 2019. Single-cell transcriptomics reveal a correlation between genome architecture and gene family evolution in ciliates. mBio 10X6:1–13.

Zufall RA, Katz LA. 2007. Macronuclear and macronuclear forms of beta-tubulin genes in the ciliate Chilodonella uncinita reveal insights into genome processing and protein evolution. J Eukaryotic Microbiol. 54(3):275–282.

Zufall RA, Robinson T, Katz LA. 2005. Evolution of developmentally regulated genome rearrangements in eukaryotes. J Exp Zool Part B 304B(5):448–455.

Associate editor: Sujal Phadke