An Extremely Streamlined Macronuclear Genome in the Free-Living Protozoan *Fabrea salina*

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**Abstract**

Ciliated protists are among the oldest unicellular organisms with a heterotrophic lifestyle and share a common ancestor with *Plantae*. Unlike any other eukaryotes, there are two distinct nuclei in ciliates with separate germline and somatic cell functions. Here, we assembled a near-complete macronuclear genome of *Fabrea salina*, which belongs to one of the oldest clades of ciliates. Its extremely minimized genome (18.35 Mb) is the smallest among all free-living heterotrophic eukaryotes and exhibits typical streamlined genomic features, including high gene density, tiny introns, and shrinkage of gene paralogs. Gene families involved in hypersaline stress resistance, DNA replication proteins, and mitochondrial biogenesis are expanded, and the accumulation of phosphatidic acid may play an important role in resistance to high osmotic pressure. We further investigated the morphological and transcriptomic changes in the macronucleus during sexual reproduction and highlighted the potential contribution of macronuclear residuals to this process. We believe that the minimized genome generated in this study provides novel insights into the genome streamlining theory and will be an ideal model to study the evolution of eukaryotic heterotrophs.

**Key words:** ciliate, genome streamlining, genome evolution.

**Introduction**

With the evolution of prokaryotes into eukaryotes, the complexity of organisms has increased dramatically, but phenotypic complexity is not proportional to genome size. For example, the prokaryotic genome *Sorangium cel lulosum* (14.78 Mb) (Han et al. 2013) is even larger than that of the unicellular eukaryotic green alga *Ostreococcus tauri* (12.56 Mb) (Derelle et al. 2006). Genome size evolution is deemed to be shaped by distinct directions, that is, expansion or contraction (Koonin 2009). Studies on the genomes of *Prochlorococcus* (Garcia-Fernández et al. 2004), SAR11 (Grote et al. 2012), and SAR86 (Molloy 2012) have contributed to the genome streamlining theory, which explains the observed small, compact, and relatively AT-rich genomes. The characteristics of a typical streamlined genome include a small genome size, a low ratio of noncoding DNA and low numbers of paralogs (Giovannoni et al. 2014). In theory, a large effective population size is essential for the efficiency of evolutionary selection, and an oligotrophic marine environment is the base of selective pressure (Giovannoni et al. 2014). Additional studies further revealed the ubiquity of streamlined genomes in bacteria and archaea in the ocean (Getz et al. 2018). However, the status of genome streamlining in eukaryotes is not clear.

Ciliates are among the most diverse clades of unicellular eukaryotes, with 4,500 recorded free-living species and an estimated number of 27,000–40,000 species (Foissner et al. 2007). Ciliates originated approximately 1,100 million years ago (Fernandes and Schrago 2019), and they disperse virtually overall freshwater, marine, and terrestrial environments. An important feature of ciliates is that each cell harbors two types of nuclei, the diploid germline micronucleus (MIC) and the polyploid somatic macronucleus (MAC). High species and population diversity, as well as unusual genome characteristics, make ciliates good model organisms for studies of genome evolution and environmental adaptation. Previous studies on the ciliate *Paramecium tetraurelia* revealed that it has experienced three whole-genome duplication events, resulting in a somatic genome of 72.09 Mb (Aury et al. 2006). Additionally, the largest somatic genome (117 Mb) among the sequenced ciliate species was reported in *Entodinium caudatum* (Park et al. 2021). However, the minimum number of genes in the somatic genome required to maintain heterotrophic free-living life remains unknown.
**Fabrea salina** is a heterotrichous ciliate that dominates in hypersaline environments such as salt marshes, hypersaline lakes, and solar salterns. It can grow and reproduce at almost all salinities in the range of 35–180 ppt (Hotos 2021), and it is highly adaptable to various environmental conditions, with a short generation period and significant resistance to harsh environments (Guermazi et al. 2008). The body length along the long axis of *F. salina* is approximately 100–350 µm (fig. 1A). The major morphological characteristic of *F. salina* is that its oral cavity forms a vortex-like S-shaped feature. Under a suitable growth environment, cells proliferate stably by asexual reproduction. In vegetative cells, most of them contain one ribbon-like MAC, while a few may have multiple MACs (supplementary fig. S1A and B, Supplementary Material online). In this study, we employed both Illumina and Nanopore sequencing technologies to assemble the somatic genome of *F. salina* and described the characteristics of its extremely streamlined genome. By combining both genome and transcriptome data, we analyzed transcriptional changes in macronuclear residuals during sexual reproduction. In addition, through comparative genomic and transcriptomic analyses, we further systematically explored its genetic basis for adaptation to hypersaline environments.

**Results and Discussion**

**Shotgun Sequencing and Assembly of the Macronuclear Genome of *F. salina***

To address the assembly challenge arising from highly fragmented nanochromosomes of ciliates, we employed both Illumina and Nanopore sequencing technologies to generate short and long reads for *F. salina*, respectively. Then, we assembled an 18.35 Mb (18,352,030 bp, N50 = 265.05 kb) draft genome based on 185.8 million Illumina paired-end reads (27.86 Gb) and 0.33 million Nanopore long-reads (27.86 Gb) by using a combination of the Flye (Kolmogorov et al. 2019) – Racon (Vašer et al. 2017) – Medaka – Pilon (Walker et al. 2014) pipeline (fig. 18 and C). We found that 97.98 and 94.56% of the Illumina and Nanopore DNA-seq reads, respectively, and 95.82% of 60.5 million RNA-seq reads could be properly aligned to the assembled genome (fig. 1D), indicating its high completeness. Notably, 86 of 106 assembled chromosomes (15.85 Mb) were capped with telomeres. We further used three different approaches (BUSCO [Seppey et al. 2019], EUKCC [Saary et al. 2020] and CEGMA [Parra et al. 2007]) to evaluate its completeness and potential contamination. As shown in figure 1D, the completeness of the assembled *F. salina* genome was comparable with that of well-studied ciliate model species (supplementary table S1, Supplementary Material online). The GC content distribution of assembled contigs did not show any algal or bacterial contaminants with discordant GC content (supplementary fig. S1C, Supplementary Material online).

Then, the MAKER (Holt and Yandell 2011) pipeline was used to annotate the MAC genome. By combining transcriptome-based, homology-based, and ab initio approaches, we finally obtained 9,918 protein-coding gene models, 89 tRNAs, and 76 rRNAs. Among the predicted protein-coding genes, almost all of them (99.9%) were supported by RNA-seq data, and 88.5% had significant hits in the InterPro database by InterProScan (Jones et al. 2014) (fig. 1D). The gene number in each contig correlates with contig length. The median gene length was 1,145 bp, and each chromosome had an average of 95.3 genes (fig. 1F and G). Most genes (68.2%) contained only one exon, while multiexon genes had 5,156 introns in total (fig. 1H). Most introns are 15 bp (fig. 1I), which is similar to other heterotrich ciliates (e.g., *Stentor coeruleus* [Slabodnick et al. 2017] and *Condylostoma magnum* [Swart et al. 2016]). We also found that the *F. salina* genome contained canonical introns with a highly conserved branch point adenosine (fig. 1J), similar to that in *S. coeruleus* (Slabodnick et al. 2017). We have scanned the genome and confirmed that the *F. salina* genome was translated with a standard codon (supplementary table S2, Supplementary Material online).

In addition, we uncovered a high consistency of contig copy number in three *F. salina* individuals based on their sequencing depth distribution (cor = 0.96) (fig. 1K). We observed that the assembled contig with the greatest copy number variation was the mitochondrial genome (supplementary fig. S1D, Supplementary Material online), which was 42,610 bp and contained 2 rRNA genes, 5 tRNA genes, 26 known protein-coding genes, and 20 unclassified open reading frames. Similar to other sequenced ciliate mitochondrial genomes (Huang et al. 2021), the *F. salina* mitochondrial genome is a linear molecule.

**General Characteristics of the Smallest Genome in Free-Living Protozoa**

To characterize the structure of the *F. salina* MAC genome, nine well-studied ciliate genomes were compared with *F. salina*. The phylogenetic tree showed that *S. coeruleus* was a closely related species of *F. salina* (fig. 2A). The length of the basal branch leading to *Heterotricha* was shorter than that of the other classes, suggesting that *Heterotricha* was a class of ancient ciliates that experienced more conservative evolution (Fernandes and Schrago 2019). The number of chromosomes in *F. salina* is similar to that of *Oligohymenophorea* but fewer than that of *Spirotricha* characterized by fragmented genomes termed nanochromosomes. Meanwhile, 96.7% of the *F. salina* genome was covered by genes (fig. 2A), indicating the greatest nucleotide utilization efficiency and the highest gene density. The gene number of *F. salina* was much lower than that of other free-living ciliates but comparable with that of the parasitic ciliate *Ichthyophthirius multifiliis* (fig. 2A). Notably, *F. salina* has the smallest genome, even smaller than *I. multifiliis* (48.7 Mb) and the symbiotic ciliate *Paramecium bursaria* (29.2 Mb). Next, we compared the genome size with all published protozoan genomes and found that *F. salina* has the smallest genome among all
free-living protozoa described thus far (fig. 28). Its size is even smaller than Tiegheostelium lacteum (23.4 Mb) (Narita et al. 2020), parasitic Plasmodium falciparum (23.5 Mbp) (Moser et al. 2020), and parasitic Entamoeba histolytica (20.8 Mb) (Al-Ardi 2021). Considering that parasites and symbionts are dominated by reductive evolution with extensive losses of genes and introns (Wolf and Koonin 2013), their genomes are usually smaller than their free-living relatives. Unexpectedly, free-living F. salina has the smallest genome, suggesting that it may have undergone unusual genome streamlining.

A typical streamlined genome is usually characterized by a smaller genome size, a lower ratio of noncoding DNA, fewer paralogs, and nonessential genes (Giovannoni et al. 2014). To investigate genome streamlined traits in F. salina, orthogroups were determined using OrthoFinder (Emms and Kelly 2019) in nine representative ciliates as well as F. salina. A total of 207,761 genes were assigned to 33,032 orthogroups (fig. 2C). A total of 2,556 orthogroups were found in more than seven ciliates and thus were considered ciliate core orthogroups (CCOs). A total of 4,037 of 9,918 coding genes in F. salina could be
classified into CCOs. The ratio of CCOs in \textit{F. salina} (40.7\%) was only lower than that of \textit{I. multifiliis} (53.5\%) but significantly higher than that of its relative \textit{S. coeruleus} (30.0\%) (\(\chi^2\) test, \(P < 2.2 \times 10^{-16}\)) (fig. 2D). The proportion of species-specific genes (nonessential genes) showed an opposite trend, in which \textit{F. salina} was significantly fewer than \textit{S. coeruleus} (\(\chi^2\) test, \(P < 2.2 \times 10^{-16}\)) (fig. 2E). In addition, the gene number of \textit{F. salina} in each orthogroup was much lower than that of \textit{S. coeruleus} (fig. 2F), indicating its highly compact genome with limited paralogs.
Next, nonsynonymous/synonymous substitution ratio analysis (Ka/Ks) revealed that gained and expanded gene families showed higher Ka/Ks values than contracted gene families (Mann–Whitney test, \( P = 2.7 \times 10^{-13} \) and \( P = 1.3 \times 10^{-8} \), respectively) (fig. 2G), suggesting that the gained and expanded gene families may be subjected to more relaxed selective pressure. We performed Gene Ontology (GO) functional enrichment analysis on the contracted genes and found that among 317 contracted orthogroups, most of them were involved in organelle part (32.8%), catalytic activity (49.8%), or metabolic process (50.8%) (fig. 2H). GO:0010766 (negative regulation of sodium ion transport) and GO:1903288 (positive regulation of potassium ion transmembrane transport) were only present in one expanded orthogroup, where \( \text{KO} \) retention of sodium ion transport) and GO:1903288 (positive regulation of potassium ion transmembrane transport) were only present in one expanded orthogroup, where \( \text{KO} \) had a comparable number of genes. We found that even compared with the parasitic ciliate, \( \text{F. salina} \) exhibited a significantly lower gene content in the signaling and cellular processes and metabolic pathway (Fisher’s exact test, \( P < 0.01 \)) (supplementary fig. S3A, Supplementary Material online). Notably, the number of genes related to genetic information processing in \( \text{F. salina} \) was significantly higher than those in other ciliates (21.43–42.80%) but comparable with that in JCVI-syn3A (Fisher’s exact test, \( P = 0.698 \)), an artificially designed bacteria with the fewest genes necessary for life (Breuer et al. 2019). This finding suggests that genes in \( \text{F. salina} \) should be functionally optimized during evolution, with only the most viable and essential genes being retained.

KO-based functional enrichment analysis revealed that genes involved in DNA replication, mitochondrial biogenesis, ubiquitination, transcription, G protein—coupled receptors, and lectins were enriched in \( \text{F. salina} \) compared with \( \text{I. multifilisi} \) (Fisher’s exact test, \( P < 0.05 \)) (fig. 3C). In addition, the gene content related to DNA replication and mitochondrial biogenesis was the highest in \( \text{F. salina} \) compared with any other ciliate (supplementary fig. S3B, Supplementary Material online), which suggests that a higher proportion of energy and replication-related genes retained in streamlined genomes may have a selective advantage in extreme environments.

Limited Chromosomal Drift During Asexual Reproduction

One of the characteristics of ciliates is that they can reproduce asexually via less accurate amitosis, which may lead to uneven distribution of MAC polyploid chromosomes in progeny cells, known as chromosomal copy drift (CCD) (Zheng et al. 2021). The occurrence of CCD results in a deviation of 1:1 in the allele frequency of the heterozygous sites on the genome. Given that CCD is thought to reduce cell fitness and thus many ciliates cannot be maintained for a long period under laboratory conditions (Bell and Graham 1988; Zheng et al. 2021), we investigated the CCD in the \( \text{F. salina} \) MAC genome by analyzing the minimum allele frequencies (MAFs) of heterozygous loci on each chromosome. We detected a total of 37,653 heterozygous sites under a strict threshold (minimum frequency >0.05, total coverage of each site >5x, and coverage of the minor allele >5x). The MAF of 15.77% heterozygous loci was <0.4 (referring to unbalanced sites, with a large deviation from 1:1), which was the lowest proportion among the 10 ciliate species (fig. 3D and supplementary fig. S4A, Supplementary Material online), indicating a limited influence of CCD in \( \text{F. salina} \). Interestingly, we observed that the overall proportion of balanced sites (MAF > 0.4) in the genome was negatively correlated with chromosome number (supplementary fig. S4B, Supplementary Material online).

Next, we determined the proportion of balanced sites on each chromosome to explore potential factors affecting CCD. The proportion of balanced sites showed a significant positive correlation with the copy number and chromosome length (cor = 0.28 and 0.32, respectively) (fig. 3E and F). Likewise, chromosome length and copy number were positively correlated (cor = 0.37, \( P = 0.0004 \)) (supplementary fig. S4C, Supplementary Material online).
online), showing that longer contigs tended to have more copies. We hypothesized that more functionally important genes tend to be located on the chromosomes where CCD is less likely to occur to reduce the impact of unequal MAC division. As expected, we found that the CCO genes in *F. salina* were significantly enriched in the top 50% of contigs with higher copies (Fisher’s exact test, *P* = 0.01) but not in the top 50% of contigs with longer lengths (Fisher’s exact test, *P* = 0.13) (fig. 3G). Chromosomes with more copies are more likely to avoid the CCD effect, and similar observations were also made in *Oxytricha trifallax* (Swart et al. 2013) and *Halteria grandinella* (Zheng et al. 2021).

Cryptically Excised Internally Eliminated Sequences Increase Transcriptional Diversity During Sexual Reproduction

During sexual reproduction in ciliates, the MAC is diminished and forms a variable number of MAC residuals. Generally, most parental MAC residuals are completely
degnerated, while a few of them may be retained and eventually fuse with the new MAC (Gong et al. 2020). However, it is still not clear whether MAC residuals play a role in sexual reproduction. During the formation of new MACs, DNA segments of MICs named internally eliminated sequences (IESs) are spliced (Zheng et al. 2021). Therefore, detecting whether the IESs are absent in the sequence can be used as a signal to identify the origin of the transcriptionally expressed genes.

Given that some IESs may be alternatively retained in MACs, we identified 461 candidates alternatively spliced (AS) IESs with pointer structures. The median AS ratio was 1.39%, which is comparable with observations in P. tetraurelia (Duret et al. 2008) and H. grandinella (Zheng et al. 2021), indicating that cryptically excised IESs are retained in most copies of the polyploid MAC genome, with only a small fraction of the polyploid chromosomes undergoing complete excision of IESs. The median lengths of AS-IESs and pointers were 14 and 4 nt, respectively (figs. 3H and I). However, the pointer sequence did not show any obvious motif and may have undergone a different excision mechanism compared with the typically fully excised IES. We found that 346 IESs were located in the exonic region (exonic IESs), which was longer than those located in other regions (Mann–Whitney test, \( P = 0.0003 \)) (fig. 3J), with median lengths of 16 and 12 nt, respectively. The AS ratio of exonic IESs was significantly lower than that in other regions (Mann–Whitney test, \( P < 0.001 \)) (fig. 3K). Among all the IESs, 133 were found in the transcriptome data, and the AS ratio at the transcriptional level was higher than that at the genomic level (Mann–Whitney test, \( P < 0.001 \)) (fig. 3L). Genes containing exonic IESs have multiple spliced isoforms in the genome, which may have different functions. These results suggest that the IES in the exonic regions greatly increases protein diversity, which may represent a fine regulation of gene content and their related functions in ciliates.

Next, we induced conjugation and sequenced the transcriptome at five consecutive time points (supplementary figs. S5–S8, Supplementary Material online). Eighteen RNA-seq libraries were built and sequenced to generate 3,655 million paired-end reads (54.83 Gb). As shown in Supplementary figure S8, the MAC residual becomes regular and rounded. The MAC residual was still present 72 h after conjugation (CP) separation (supplementary figs. S7 and S8, Supplementary Material online). Interestingly, all 77 exonic IESs could be detected at all five time points, and samples from different conjugation periods were clearly separated (supplementary fig. S5A, Supplementary Material online), indicating that MAC residuals retained transcriptional activity during sexual reproduction. Next, 33 exonic IESs present in more than five samples were selected to describe the expression pattern in detail (supplementary fig. S6, Supplementary Material online). The results showed that despite being located on the same chromosome, different genes showed opposite directions (supplementary fig. S6, Supplementary Material online), suggesting a dynamic regulation of the ratio between different versions of the chromosome during conjugation. Next, six genes that have a highly variable exonic IES (SD > 2) were identified (supplementary fig. S5B, Supplementary Material online). Fabrea_09599, Fabrea_02192, and Fabrea_00615 have a WD40 repeat domain (Pfam ID: PF00400) that is associated with multiple functions, including RNA processing and control of cell division (Suganuma et al. 2008). Fabrea_00480 was annotated as DIAPH2 (diaphanous 2, K05741) involved in cytoskeletal proteins. Fabrea_09158 was annotated as FRAS1 (extracellular matrix protein FRAS1, K23379), which is associated with a variety of physiological processes, including shape change, proliferation, and motility (Wu 2004). These findings indicate that the cryptically excised IESs in the MAC may play important functions during sexual reproduction.

Genetic Basis for Salinity Tolerance

To better understand its broad salinity adaptation, we analyzed the gene expression profiles of F. salina at different salinity levels: 80‰, 130‰, and 180‰ (fig. 4A). A total of 1,129 differentially expressed genes (DEGs) were found in the DEG analysis of the high and optimum salinity groups, and 1,353 DEGs were detected between optimum and low salinity (fig. 4B). However, only 23.4% of the DEGs were shared in both comparisons, and clustering analysis showed a different expression signature (fig. 4C and F), suggesting that F. salina may have distinct mechanisms to adapt to low and high osmotic pressures. Further analysis confirmed that DEGs of high salinity group were significantly enriched (\( \chi^2 \) test, \( P = 5.86 \times 10^{-6} \)) in the expanded gene families (fig. 4D and F), most of which were involved in G protein-coupled receptors and chaperones and folding catalysts (fig. 4G). These findings mirror the results of previous studies, which showed that G protein receptors play an important role in the response to abiotic stresses, especially high salinity (Liu et al. 2018). Activation of chaperones and folding catalysts is a key way for organisms to deal with misfolded proteins resulting from the accumulation of reactive oxygen species owing to salt stress (Zhang et al. 2021). Next, we found that the most significant changed DEGs were enriched in the genes specific to the high salinity group (Fisher’s exact test, \( P = 1.4 \times 10^{-7} \)), rather than those shared with the low salinity group (fig. 4H and I). Functional enrichment analysis showed that the GO terms such as biosynthetic process, regulation of cellular process, cellular component organization or biogenesis, were up-regulated in high salinity environments, whereas the GO terms related to carbohydrate derivative binding, ion binding, microtubule-based process were down-regulated (fig. 4I).

Microorganisms counteract salt stress by the accumulation of compatible solutes that act as osmoprotectants to help cells resist osmotic pressure without altering intracellular enzyme activity (Harding et al. 2016). A large number of
osmoprotectants, including proline, glycine betaine (GB), ectoine, and glycerol, have been reported in halophilic and halotolerant organisms (Weinisch et al. 2019). Weinisch et al. (2019) found that GB and ectoine increased linearly with salinity stress in _F. salina_. Here, we observed an incomplete biosynthetic pathway for GB, with the absence of Ect/5-hydroxyectoine-related genes (ectABCSD) in the MAC genome. For betaine aldehyde dehydrogenase (BADH), one of the important gene involved in the biosynthetic pathway of GB, its abundance was negatively correlated with salinity (cor = −0.74, P = 0.023). Interestingly, we found that the expression of enzymes related to glycerol biosynthesis decreased with increasing salinity (fig. 4K and L). The abundance of ACP6 was significantly negatively correlated with salinity (cor = −0.685, P = 0.042). The expression level of GK in the 80% salinity group was lower than that in the 180% group, although this difference was not statistically significant.

*Fig. 4. *Genetic adaptation for salinity tolerance. (A) Flow chart of the experimental design and transcriptome analysis of _F. salina_ under salinity stress. (B) The number of differentially expressed genes in the high and low salinity groups. Both groups were compared against the optimum salinity group. (C) Venn diagram of differentially expressed genes in the two groups. (D) The bar graph indicates the enrichment of DEGs in expanded and contracted gene families. (E) Heatmap for differentially expressed genes. (F) Kn/Ks of differentially expressed genes. P value was determined based on Mann–Whitney test. (G) KEGG enrichment analysis of DEGs in different categories. The color of the dots represents the P value. (H) Differential significance distribution of the DEGs in the high salinity group. (I) Significant GO terms (P < 0.05) in the high salinity-specific DEGs. The P value in I and J was determined based on Fisher’s exact test. (K) Distribution of gene expression at different salinities. The Spearman correlations and their significance are marked at the bottom of the image. The adjusted P values resulted from DEG analysis and are marked at the top of the image. ****Adj. P < 0.0001, **Adj. P < 0.001, *Adj. P < 0.01, *Adj. P < 0.05. (L) Schematic diagram of the pathways associated with glycerol and PA synthesis. Orange and blue colors represent up-regulated and down-regulated steps under hypersalinity stress, respectively. GPDH, glycerol 3-phosphate dehydrogenase; GPAT, glycerol-3-phosphate acyltransferase; LPAAT, lysophosphatidic acid acyltransferase; DGK, diacylglycerol kinase; GK, glycerol kinase; GPDH, glycerol 3-phosphate dehydrogenase; GPAT, glycerol-3-phosphate acyltransferase; LPAAT, lysophosphatidic acid acyltransferase; DGK, diacylglycerol kinase; GK, glycerol kinase; MGLL, monoglyceroide lipase; ACP6, lysophosphatidic acid phosphatase type 6 precursor; DGK, diacylglycerol kinase; DGAT, diacylglycerol acyltransferase; DHAP, dihydroxyacetone-phosphate; G3P, glycerol-3-phosphate dehydrogenase; MGA, monoacylglycerol; LPA, lysophosphatidic acid; PA, phospholipid acid; DAG, diacylglycerol; TAG, triacylglycerol.
(adj. \( P = 0.064 \)). These results suggest that the compatible solutes in \( F. \) *salina* may not be fully self-synthesized and confirmed by findings in the heterotrophic ciliate *Schmidendorfithrix salinarum*, which could directly absorb GB and ectoine from the environment (Weinisch et al. 2018).

As a lipid second messenger, phosphatidic acid (PA) was reported to be involved in intracellular salt stress signaling in eukaryotes (Shen et al. 2019). In *F. salina*, we found that the expression of GPAT, which acylates G3P to produce LPA, was significantly positively correlated with salinity (cor = 0.84, \( P = 0.004 \)). The same expression trend was also observed in GPDH (cor = 0.84 and \( P = 0.004 \)). The expression of another important gene, lysophosphatidic acid acyltransferase (LPAAT), which produces PA with lysophosphatidic acid (LPA) and acyl-CoA as substrates, was significantly higher in the 180‰ and 130‰ groups than in the 80‰ group (adj. \( P = 0.049 \) and adj. \( P = 0.001 \), respectively) (fig. 4K and L). Such increased expression of the enzymes associated with PA synthesis was also observed in the alga *Parachlorella kessleri* grown under salt stress conditions (You et al. 2019). We demonstrated that the accumulation of PA may play an important role in resistance to high osmotic pressure.

### Conclusion

In this study, we used both Illumina and Nanopore sequencing technologies to assemble a high-quality macronuclear genome of *F. salina* characterized by an unusually streamlined and compacted genome in free-living protozoa. We revealed the minimal gene content that allows eukaryotes to live freely and heterotrophically, which suggests that genome streamlining is in fact not unique to prokaryotes but has also been adopted by some eukaryotes. Given their dual nuclei system, ciliates represent an ideal model to study genome expansion or contraction events. Our study provides a unique and valuable resource for understanding the genomic conservation and evolutionary history of eukaryotic organisms.

### Methods and Materials

#### Culture of *F. salina*

*Fabrea salina* was collected from the BeiDaGang wetland along the Bohai Sea coast at Tianjin (38°57’N, 117°37’E), China, by the Tianjin Academy of Agricultural Sciences and was identified by its morphological features using differential interference contrast microscopy. Cells were incubated in automatically filtered sterile seawater (pH 7.8) at 30°C and 80 pSU (actual units of salinity) under lighting. *Dunaliella salina* was supplied as food source. After 7–10 days of culture, the population density reached 100 cells/1 ml of medium. The status and cell numbers of *F. salina* were checked using an anatomical lens.

#### Nucleic Acid Isolation and Sequencing

As described previously (He et al. 2019), *F. salina* cells were isolated through a 15-μm-pore-size nylon filter membrane and washed three times. Then, the cells were starved for 1 day, and 1 × penicillin–streptomycin antibiotics (Invitrogen, Carlsbad, USA) were added to eliminate bacterial contamination. Before extracting nucleic acids, the cells were washed five times with sterile water to remove any bacteria that may be attached to the cell surface. For single-cell genome sequencing, 12 cells were placed in three sterile 0.2-ml tubes as biological replicates, with 3, 4, and 5 cells in each tube. DNA extraction was carried out following the Repli-g single-cell kit manufacturer’s instructions (Qiagen). Then, approximately 12,000 cells were harvested and generated 1 μg DNA for Nanopore long-read sequencing. To obtain high-quality genomes, we constructed two types of libraries. The native library was constructed following with native barcoding genomic DNA protocol (with EXP-NBD104, EXP-NBD114, SQK-LSK109, ONT, UK). The polymerase chain reaction (PCR) library was constructed according to PCR barcoding genomic DNA protocol (with SQK-LSK109, EXP-PBC001, ONT, UK) with 12 PCR cycles. The library mean length was 8 kb and was sequenced on R9.4 FlowCells using the MinION sequencer (ONT, UK) for 24 h. ONT Guppy software (v1.8) was used for base calling on fast5 files, and the “passed filter” reads were used for downstream analysis.

In addition, the Illumina paired-end sequencing DNA library was constructed by shearing genomic DNA to 300 bp fragments on a Covaris S220 system (Covaris, Woburn, MA, USA). A Nextera DNA Flex Library Prep kit (Illumina, San Diego, CA, USA) was used following the manufacturer’s protocol. Three DNA libraries were sequenced using the Illumina HiSeq 2500 platform (Illumina, San Diego, CA, USA) for 2 × 150 bp reads and generated 27.86 Gb of data.

For the single-cell transcriptome, whole-transcriptome amplification was performed using a SMART-Seq2 v4 Ultra Low Input RNA kit (NEB, USA) following the manufacturer’s protocols. Sequencing libraries were prepared with the Illumina Nextera XT kit and sequenced using an Illumina HiSeq 2500 platform. High-throughput sequencing for three samples produced 54.83 Gb of 2 × 150 pair-end data in total.

#### Genome Assembly

First, Nanopore long reads were trimmed with porechop (version 0.2.4) (https://github.com/rrwick/Porechop) to remove adapters from read ends and split sequences with internal adapters. The processed reads were assembled to draft the genome by Flye (version 2.8.1) with the default parameters (Kolmogorov et al. 2019). Subsequently, a combination of polishing methods was used, including Racon (Vaser et al. 2017), Medaka (version 1.3) (https://github.com/nanoporetech/medaka) and Pilon (version 1.23) polishing with Illumina sequencing reads (Walker et al. 2014). A total of 185,798,514 Illumina paired-end reads were used to correct the draft genome. Finally, we obtained a high-quality *F. salina* MAC genome (18.3 Mb) with an N50 of 265.05 kb and a
complete mitochondrial genome (42,610 bp) (see the Supplementary Materials for more details).

Assessment of Genome Assembly
First, the integrity of core eukaryotic genes in the MAC genome was assessed using CEGMA (version 2.5) (Parra et al. 2007) with default parameters. Two hundred of 248 core eukaryotic orthologous groups (KOGs) were identified. Then, a relaxed restriction of CEGMA search was performed on the remaining genes following the published approach of S. coeruleus (Slabodnick et al. 2017). Hmmscan (HMMER, version 3.3.2, http://hmmmer.org/) was used to assign the Pfam domain in KOGs to the predicted genes in *F. salina*. With this analysis, 46 additional core eukaryotic genes were determined under the full-sequence e-value $< 1 \times 10^{-3}$. Finally, the updated HMMs for two undetected KOGs (KOG0563 and KOG2531) in the EggNOG database were downloaded and reassigned to genes by hmmscan. Three genes were assigned to them. Above all, 248 core eukaryotic genes were found in our assembled genome. This result is similar to the completeness rate of fully well-studied ciliate genomes, such as *S. coeruleus*, *Tetrahymena thermophila*, *P. bursaria*, *F. salina*, *S. coeruleus*, *T. thermophila*, *E. octocarinatus*, and *O. trifallax*, which commonly contain 220–243 KOGs.

Additionally, EukCC (version 4.0.6) (Saary et al. 2020) and BUSCO (version 0.2) (Seppey et al. 2019) were used to assess the completeness and contamination based on the predicted gene amino acid sequence in *F. salina* with default parameters. The results showed that the completeness was 75.76% and 83.6% in BUSCO and EukCC, respectively. The contamination was 3.03% and 6.4% in BUSCO and EukCC, respectively. The other nine published ciliate genomes were assessed using CEGMA (version 2.5) (Parra et al. 2007) with default parameters. Two hundred of 20,356 transcripts were assembled using Trinity (version 2.11.0) with default parameters. A total of 20,356 transcripts were obtained. Cd-hit (version 4.8.1) (Li and Godzik 2006) was used to eliminate redundancy at a threshold of 0.95 similarity (~0.95). For homology evidence, amino acid sequences in nine published ciliates were collected as cross-species protein homology evidence.

InterProScan (version 5.48-83.0) (Jones et al. 2014) was used to identify conserved functional motifs and protein domains. The predicted proteins were aligned with parameters (-iprlookup -goterms) to the following databases: PROSITE, HAMAP, PFAM, PRINTS, PRODOM, SMART, TIGRFAMS, PIRSF, SUPERFAMILY, CATH-Gene3D, and PANTHER. The results showed that 8,781 of 9,918 genes contained motifs or protein domains in the *F. salina* assembled genomes. Additionally, eggNOG-mapper (Huerta-Cepas et al. 2019) was used based on the webserver http://eggnog-mapper.embly.de/. A total of 5,995 genes were assigned to GO terms, KEGG KO, or EC numbers. For noncoding RNA prediction, rRNA and tRNA were predicted by cmscan based on the Pfam (version 14.4) (Kalvari et al. 2018) database. tRNAscan-SE (version 2.0.7) (Chan and Lowe 2019) was used to identify tRNA with default parameters.

Codetta (version 2.0) was used to predict the genetic code from nucleotide sequence data (Shulgina and Eddy 2021).

Detection of Telomeres
To identify the telomere in the MAC genome of *F. salina*, 200 bp sequences from both ends of assembled contigs were extracted and analyzed by a custom R script that recognized the telomere repeat 5-CACCCTAA-3. Briefly, the sequences of contig ends were cut into 8 nt kmers, and kmers with significantly higher frequencies were assembled and manually validated.

Gene Prediction and Annotation
The MAKER pipeline (version 2.31.11) (Holt and Yandell 2011) was used to predict gene models. The pipeline includes Augustus (Nachtweide and Stanke 2019), SNAP (Korf 2004), and GeneMark-ES (Borodovsky and Lomsadze 2011) to identify repeats, align ESTs and proteins to the genome, produce ab initio gene predictions, and then combine these data into gene annotations with evidence-based quality values. Considering the short intron length (15–16 nt), the source code (filename: intronmodel.cc, types.cc, extrinsic.cc, G1.pm, protein.pm, alttest.pm, est.pm) in Augustus was modified to change the minimum length limit to 9 nt for introns (the default is 39 or 20). Then, the source code was recompiled using these updated settings.

The transcriptome evidence for gene prediction came from two sources. The first was a published *F. salina* transcriptome assembly (MMETSP1345) containing 10,034 assembled transcripts produced by the Marine Microbial Eukaryotic Transcriptome Sequencing Project (MMETSP) (Keeling et al. 2014). The second was generated in this study, which contained 9.07 Gb of PE150 transcriptome sequencing data. After quality filtration and adapter removal, processed reads were assembled using Trinity (version 2.11.0) with default parameters. A total of 20,356 transcripts were obtained. Cd-hit (version 4.8.1) (Li and Godzik 2006) was used to eliminate redundancy at a threshold of 0.95 similarity (~0.95). For homology evidence, amino acid sequences in nine published ciliates were collected as cross-species protein homology evidence.

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Codetta (version 2.0) was used to predict the genetic code from nucleotide sequence data (Shulgina and Eddy 2021).

Phylogenetic Analysis
The *F. salina* 18S rRNA gene was identified by Pfam. 18S rRNA genes of the following nine ciliates and an outgroup were downloaded from NCBI: *H. grandinella* (MG002432.1), *S. lemniae* (AM233915.1), *O. trifallax* (FJ545743.1), *E. octocarinatus* (LT623905.1), *T. thermophila* (M10932.1), *I. multifiliis* (U17354.1), *P. tetraurelia* (AB252009.1), *S. coeruleus* (JQ282899.1), *P. bursaria* (MG589318.1), and the outgroup *Plasmodium falciparum* (MF155937.1). Then, multiple sequence alignment and maximum likelihood tree construction were performed in MEGAX (Kumar et al. 2018). Gene density was defined as the ratio of the total length of protein-coding genes to the genome length.

Heterozygous Loci Detection
First, the genome sequencing data of nine published ciliates were downloaded from public databases (supplementary table S4, Supplementary Material online). Then, the sequence reads were aligned to the reference
Among the 5,173 orthogroups in F. salina nonymous substitutions (expansion events (involving 1129 genes). The rate of nonsy-317 contraction events (involving 580 genes) and 252 ex-
were considered CCOs, as they were detected in 207,761 (84.20%) of 246,731 genes were assigned to
MCL clustering algorithm with default parameters. A total of 60,623 genes (24.57%) were in 15,978 species-
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Orthology Group Analysis
The protein sequences of 10 species were downloaded from NCBI and other databases: H. graddinella (GCA_006369765), S. lemmnae (GCA_000751175), O. trifallax (http://oxy.ciliate.org/index.php/home/downloads), E. octocarinatus (http://ciliates.ihb.ac.cn/database/download/#eo), T. thermophila (http://ciliate.org), I. multiliis (GCA_000220395), P. tetraurelia (GCF_000165425), S. coeruleus (GCA_001970955), and P. bursaria (https://ngdc.cnbc.ac.cn/gwh/Assembly/138/show; GWHAA FB00000000). OrthoFinder (version 2.5.2) (Emms and Kelly 2019) was used to infer orthogroups based on the

Cells were selected before CP separation and 12, 24, 48, and 72 h after CP separation. For CP, one cell was stained with Hoechst 33342 to observe nuclear morphology, and the other cell was used for transcriptome sequencing.

Growth Rate at Different Salinity Levels
We set up 13 consecutive salinity gradients between 60‰ and 180‰, with three parallel experiments for each salinity. The volume of each cell culture flask was controlled at 50 ml, and the number of cells was controlled at 200. Dunaliella salina was provided as food and incubated for 2 weeks. During the incubation, sterile seawater or sterile distilled water of varying salinity was added daily to maintain the salinity of the culture. As a result, the maximal growth rate was at a salinity of 130‰. Therefore, we collected F. salina samples at three salinities (80‰, 130‰, and 180‰) with three replicates of each salinity for transcriptome sequencing. Gene expression was quantified using HISAT2 (version 2.0.5) and StringTie (version 1.3.4) (Pertea et al. 2016).

Alternatively Spliced IES Detection
The software ADFinder (Zheng et al. 2020) was used to detect the alternatively spliced IESs in the MAC genome without sequence data of the MIC genome. For the polyploid MAC genome, different versions of the same chromosome may exist with IESs deleted (excised) and IESs retained (nonexcised). As recommended, Bowtie2 (version 2.3.2) software (local, -k 5, -ma 3) was used to align our RNA-seq and DNA-seq reads to the MAC genome. Then, ADFinder was used to identify the position and pointer sequence of alternatively spliced IESs with default parameters.

Functional Enrichment Analysis
First, all protein-coding genes in ten ciliates were classified into KO by Kofamscan (version 1.3.0) (Aramaki et al. 2020) with the parameter “-E 0.05”. Second, the unassigned genes were further annotated using the most frequent KOs in their orthogroups. Then, all genes were classified into three protein families (metabolism, genetic information processing, and signaling and cellular processes) and 52 sublevel protein families based on the KEGG BRITE database (https://www.genome.jp/kegg/brite.html). As a result, 72.4% of 254,039 protein-coding genes were successfully annotated, and a median of 76.2% of genes was annotated in each ciliate. Then, the function enricher in the R package “clusterProfiler” (version 3.12.0) (https://github.com/YuLab-SMU/clusterProfiler) was used to perform enrichment analysis.

Then WEGO (Ye et al. 2018) (version 2.0) (https://wego.genomics.cn/) was used to perform GO classification and Fisher’s exact test was used to calculate P values. Cytoscape (https://cytoscape.org/) was used to visualize the network.

Salinity Stress-Related Genes
The following genes involved in salinity tolerance were selected: Fabrea_04266 (ACP6, lysophosphatidic acid
phosphatase type 6, K14395), Fabrea_06374 (BADH, beta-
in aldehyde dehydrogenase, K00130), Fabrea_06811
(DGK, diacylglycerol kinase [ATP], K00901), Fabrea_
09038 (GK, glycerol kinase, K00864), Fabrea_05028
(GPAT, glycerol-3-phosphate O-acetyltransferase, K13506),
Fabrea_08063 (GPDH, glycerol-3-phosphate dehydrogen-
ase, K00057), Fabrea_08457 (LPAAT, lysophosphatidic
acid acyltransferase/lysophosphatidylinositol acyltransfer-
ase, K13523), Fabrea_08078 (MGLL, acylglycerol lipase,
K01054), and Fabrea_02753 (DGAT, diacylglycerol
O-acetyltransferase, K11155). Then, we combined diffe-
rential expression analysis and Spearman correlation analysis
to assess the relationship between gene expression and sal-
inity. Differential expression analysis was performed in the
R package DESeq2 (version 1.24.0) (Love et al. 2014).
Correlation analysis of gene expression and salinity was
calculated using the function cor.test(method = “spear-
man”) in R.

Supplementary Material
Supplementary data are available at Molecular Biology and
Evolution online.

Author Contributions
F.Z. and M.M. conceived the project. B.Z., T.Z., and F.Z. per-
duced the manuscript with the contribution of all authors.
All authors read and approved the final manuscript.

Data availability
All data generated or analyzed during this study are in-
cluded in this published article, its supplementary informa-
tion files and publicly available repositories. Reads for the
genome assemblies have been deposited to NCBI GenBank (PRJNA822294). The gene training model for
Fabrea salina in Augustus and orthogroup results have been
deposited to GitHub (https://github.com/callAgene/Supplementary-Materials-for-Fabrea-salina).

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References
Al-Ardi MH. 2021. Illumination on the structure and characteristics
d of entamoeba histolytica genome. Preprints 2021030070.
Ames RM, Lovell SC. 2015. DupliPHY-Web: a web server for
DupliPHY and DupliPHY-ML. Bioinformatics 31:416–417.
Aramaki T, Blanc-Mathieu R, Endo H, Ohkubo K, Kanehisa M, Goto S,
Ogata H, Valencia A. 2020. KofamKOALA: KEGG Ortholog
assignment based on profile HMM and adaptive score threshold.
Bioinformatics 36:2251–2252.
Aury J-M, Jaillon O, Duret L, Noel B, Jubin C, Porcel BM, Ségurens B,
Daubin V, Anthouard V, Aich N, et al. 2006. Global trends of
whole-genome duplications revealed by the ciliate
Paramécium tetraurelia. Nature 444:171–178.
Bell G, Graham B. 1988. Sex and death in Prototaxa: the history of ob-
session. Cambridge, UK: Cambridge University Press.
Borodovsky M, Lomsadze A. 2011. Eukaryotic gene prediction using
GeneMark.hmm-E and GeneMark-ES. Curr Proto Bioinformatics
Chapter 4, Unit 4, 6–1–10.
Breuer M, Earnest TM, Mermrayn C, Wise KS, Sun L, Lynott MR,
Hutchison CA, Smith HO, Lapek JD, Gonzalez DJ, et al. 2019.
Essential metabolism for a minimal cell. Elife 8:e36842.
Chan PP, Lowe TM. 2019. tRNAscan-SE: searching for tRNA genes in
genomic sequences. Methods Mol Biol. 1962:1–14.
Danecek P, Bonfield JK, Liddle J, Marshall J, Ohan V, Pollard MO,
Whitwham A, Keane T, McCarthy SA, Davies RM, et al. 2021.
Twelve years of SAMtools and BCFtools. Gigascience 10:giab0088.
Derelle E, Ferraz C, Rombouts S, Rouzé P, Worden AZ, Robbens S,
Partensky F, Degroeve S, Eychenyl S, Cooke R, et al. 2006.
Nem gene analysis of the smallest free-living eukaryote
Ostreococcus tauri unveils many unique features. Proc Natl
Acad Sci U S A. 103:11647–11652.
Duret L, Cohen J, Jubin C, Dessen P, Goût J-F, Mousset S, Aury J-M,
Danecek P, Bonfield JK, Liddle J, Marshall J, Ohan V, Pollard MO,
Whitwham A, Keane T, McCarthy SA, Davies RM, et al. 2021.
Twelve years of SAMtools and BCFtools. Gigascience 10:giab0088.
Derelle E, Ferraz C, Rombouts S, Rouzé P, Worden AZ, Robbens S,
Partensky F, Degroeve S, Eychenyl S, Cooke R, et al. 2006.
Nem gene analysis of the smallest free-living eukaryote
Ostreococcus tauri unveils many unique features. Proc Natl
Acad Sci U S A. 103:11647–11652.
Duret L, Cohen J, Jubin C, Dessen P, Goût J-F, Mousset S, Aury J-M,
Jaillon O, Noël B, Arnaiz O, et al. 2008. Analysis of sequence vari-
bility in the macronuclear DNA of Paramecium tetraurelia: a
somatic view of the germline. Genome Res. 18:585–596.
Emms DM, Kelly S. 2019. OrthoFinder: phylogenetic orthology inher-
ence for comparative genomes. Genome Biol. 20:238.
Fernandes NM, Schrargo CG. 2019. A multigene timescale and divers-
fication dynamics of Ciliophora evolution. Mol Phylogenet Evol.
139:106521.
Foissner W, Chao A, Katz LA. 2007. Diversity and geographic distribu-
tion of ciliates (Protista: Ciliophora). In: Foissner W, ed. Protist
diversity and geographical distribution. Dordrecht: Springer. p. 111–129.
Garcia-Fernández JM, de Marsac NT, Dizel J. 2004. Streamlined regu-
lation and gene loss as adaptive mechanisms in Prachlorococcus
for optimized nitrogen utilization in oligotrophic environments.
Microbial Mol Biol Rev. 68:630–638.
Getz EW, Tithi SS, Zhang L, Aylward FO. 2018. Parallel evolution of
gene streamlining and cellular bioenergetics across the mar-
ine radiation of a bacterial phyllum. mbio 9:e01089-18.
Gionvannni SJ, Cameron Thrash J, Temperton B. 2014. Implications of
streamlining theory for microbial ecology. ISME J. 8:1553–1565.
Gong R, Jiang Y, Vallesé A, Gao Y, Gao F. 2020. Conjugation in
Euplotes raikovi (Protista, Ciliophora) reveals new insights into nucle-
ear events and macronuclear development from micronuclear and
macronuclear cells. Microorganisms 8:162.
Grote J, Thrash JC, Huggett MJ, Landry ZC, Carini P, Giovannini SJ,
Rappé MS. 2012. Streamlining and core genome conservation
among highly divergent members of the SAR11 clade. mbio 3:
e00252-12.
Guermazi W, Elloumi J, Ayahi H, Bouain A, Aleya L. 2008. Rearing of
Fabrea salina Henneuguy (Ciliophora, Heterotrichida) with three
unicellular feeds. C R Biol. 331:56–63.
Han K, Li Z-f, Peng R, Zhu L-p, Zhou T, Wang L-g, Li S-g, Zhang X-b, Hu
W, Wu Z-h, et al. 2013. Extraordinary expansion of a
Sorangium cellulosum genome from an alkaline milieu. Sci Rep.
3:2101.
Harding T, Brown MW, Simpson AGB, Roger AJ. 2016. Osmoadaptative
strategy and its molecular signature in obligately halophilic hetero-
 trophic protists. Genome Biol Evol. 8:2241–2258.
He M, Wang J, Fan X, Liu X, Shi W, Huang N, Zhao F, Miao M. 2019.
Genetic basis for the establishment of endosymbiosis in
Paramecium. ISME J. 13:1360–1369.
Holt C, Yandell M. 2011. MAKER2: an annotation pipeline and
genome-database management tool for second-generation gen-
ome projects. BMC Bioinformatics 12:491.
