Large Differences in the Haptophyte Phaeocystis globosa Mitochondrial Genomes Driven by Repeat Amplifications

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The haptophyte Phaeocystis globosa is a well-known species for its pivotal role in global carbon and sulfur cycles and for its capability of forming harmful algal blooms (HABs) with serious ecological consequences. Its mitochondrial genome (mtDNA) sequence has been reported in 2014 but it remains incomplete due to its long repeat sequences. In this study, we constructed the first full-length mtDNA of P. globosa, which was a circular genome with a size of 43,585 bp by applying the PacBio single molecular sequencing method. The mtDNA of this P. globosa strain (CNS00066), which was isolated from the Beibu Gulf, China, encoded 19 protein-coding genes (PCGs), 25 tRNA genes, and two rRNA genes. It contained two large repeat regions of 6.7 kb and ∼14.0 kb in length, respectively. The combined length of these two repeat regions, which were missing from the previous mtDNA assembly, accounted for almost half of the entire mtDNA and represented the longest repeat region among all sequenced haptophyte mtDNAs. In this study, we tested the hypothesis that repeat unit amplification is a driving force for different mtDNA sizes. Comparative analysis of mtDNAs of five additional P. globosa strains (four strains obtained in this study, and one strain previously published) revealed that all six mtDNAs shared identical numbers of genes but with dramatically different repeat regions. A homologous repeat unit was identified but with hugely different numbers of copies in all P. globosa strains. Thus, repeat amplification may represent an important driving force of mtDNA evolution in P. globosa.

Keywords: mtDNA sizes, mitochondrial genomes, Phaeocystis globosa, repeat region, Haptophyta

INTRODUCTION

The haptophyte Phaeocystis globosa is a cosmopolitan phytoplankton that plays a pivotal role in global carbon and sulfur cycles by releasing substantial quantities of dimethylsulfide propionate (DMSP) (Schoemann et al., 2005). It is also an important HAB species that causes blooms in various ocean regions including the Arabian sea, the Southern North Sea, and the coastal waters of East Asia and Southeast Asia (Madhupratap et al., 2006; Chen et al., 2002; Schoemann et al., 2005;
Hai et al., 2010; Rousseau et al., 2013; Shen and Qi, 2021). *P. globosa* blooms outbreak frequently, bringing serious ecological hazards, and causing great economic losses of aquaculture industry and nuclear power facility threats (Shen et al., 2000; Qi et al., 2004; Hai et al., 2010; Cao et al., 2017).

Accumulating evidence suggested the *P. globosa* has high level of genetic diversity. Various physiological and ecological characteristics were observed among different *P. globosa* strains. For example, *P. globosa* strains collected in different ocean regions (e.g., Hong Kong and Shantou, China) have different acute toxic effects to *Artemia sinica* (Wei and Jiang, 2005). Sizes of *P. globosa* colonies found in the Chinese coastal waters can reach 3 cm in diameters, which are substantially larger than those found in European waters (Qi et al., 2004; Rousseau et al., 2007). The temperatures at which *P. globosa* blooms occurred were different for *P. globosa* blooms recorded in different geographical regions. For example, most *P. globosa* blooms occurred in Chinese coastal waters when temperatures were 16–22°C (Shen et al., 2000; Qi et al., 2004), which were higher than that in European waters (Seuront et al., 2006; Blauw et al., 2010). Cell surface structures of *P. globosa* strains isolated in Europe and Beibu Gulf, China were different. For example, small circular bulges were observed in small flagellate cells of China strains, while such characteristics were not observed in European strains (Hu et al., 2019).

Much effort has been made to ascertain *P. globosa* genetic diversity as a way to understand phenotypical differences. Common molecular markers (18S rDNA, 28S rDNA, ITS, and *rbcL*) have been applied to distinguish *P. globosa* strains with limited success (Hu et al., 2019; Song et al., 2020). The ITS region has been demonstrated to be a poor molecular marker because it has been found to have high intra-genome variation, while 18S rDNA, 28S rDNA and *rbcL* have been found to be over conservative that they cannot be used to effectively distinguish different strains (Hu et al., 2019; Song et al., 2020). Thus, these common molecular markers could not be adequately used to probe *P. globosa* genetic or phenotypic diversity. Molecular markers with high resolution are urgently needed to resolve *P. globosa* genetic diversity in environment.

Comparative analysis of the organelle genomes has been fruitful for identifying molecular markers with improved resolution. The complete chloroplast genome (cpDNA) of the first *P. globosa* strain was published in 2014, which was 107,461 bp in length (Smith et al., 2014). One molecular marker *rbcS*-rpL27 based chloroplast sequence intergenic spacer was found to be able to separate phylogeographic populations of *P. globosa* (Zhang et al., 2021). Comparative analysis of the cpDNAs of *P. globosa* strains facilitated the development of a new molecular marker *pgcp1* with even higher resolution for tracking *P. globosa* genetic diversity (Song et al., 2020).

Relative mutation rate of mtDNAs has been found to be higher than that of cpDNAs in *P. globosa* (Smith et al., 2014). The mtDNAs of *P. globosa* and *P. antarctica* have been published but they are still incomplete, due to the complex repeat regions (Smith et al., 2014). Repeat regions are ubiquitous in algal mtDNAs, some of which can be very large in length. For example, mtDNA of the cryptophyte *Rhodomonas salina* contains a 4.7 kb long repeat region, 11 repeat motifs ∼40–700 bp long occurring up to 31 times, forming a complex repeat structure. Tandem repeats are the major arrangement but the region also includes a large, 3 kb, inverted repeat and several potentially stable 40–80 bp long hairpin structures (Hauth et al., 2005). The cryptophyte *Hemiselmis andersonii* has a 20 kb repeat region, comprised of numerous tandem and dispersed repeat units of between 22 and 336 bp, and occurring up to 100 times. Adjacent to these repeats are 27 copies of palindromic sequences predicted to form stable DNA stem-loop structures (Kim et al., 2008).

The mtDNA of *Phaeodactylum tricornutum* has an even larger repeat region of 35 kb, consisting almost entirely of various combinations of several kinds of tandem repeat, including five elements 33-404 bp (Oudot-Le Secq and Green, 2011). The chlorophyte *Pedinomonas* mtDNA contains a 9 kb-repeated sequence, 13 repeat elements range in size from 6 to 389 bp (ele-01 to ele-13) were characterized. Although most members within an element family are identical, some are not (Turmel et al., 1999). We hypothesize that change in repeat regions may be, at least partially, due to differential repeat unit amplification, which is an important driving force for *P. globosa* mtDNA differences in evolution.

In this study, we constructed the first complete mtDNA of *P. globosa*, which contained two large repeat regions of 6.7 kb and ∼14.0 kb in length, respectively. We identified several repeat units with various numbers of copies. Comparative analysis of six *P. globosa* mtDNAs revealed that the copies numbers varied substantially. Our results revealed that change in repeat units is a driver of *P. globosa* mtDNA diversity.

**MATERIALS AND METHODS**

*Phaeocystis globosa* Strain Isolation, Culture and Characterization

The five *P. globosa* strains (CNS000066, CNS00069, CNS00087, CNS00262, and CNS00266) described herein were isolated from samples collected in 2019 from *P. globosa* bloom waters of the Beibu Gulf, China and Vietnam sea area (Table 1). Single-cell isolation was achieved by selecting individual cells using a micropipette, followed by multiple washes before transferring each single cell to 24-well culture dishes for growth and characterization. They grew in L1 seawater culture medium (without Na₂SiO₃) (Guillard and Hargreaves, 1993). Cultures were maintained at 20°C under a 12:12 h light: dark cycle with approximately 30 µM s⁻¹ m⁻² of cool white fluorescent illumination.

Identification of the cultured algae was done based on microscopic morphological characters and phylogenetic analysis based on 18S rDNA. ZEISS Axio Imager Z2 light microscope (Zeiss, Siederecht, Netherlands) were used for morphological observation. 18S rDNA was assembled basing Illumina reads with GetOrganelle (Jin et al., 2020), with publicly available 18S rDNA sequences of *P. globosa* as reference sequences. The assembled sequences were further validated by aligning sequencing reads against the assembled result using BWA (Li and Durbin, 2009) and SAMtools (1.9) (Li et al., 2009) and visualized using IGV (Thorvaldsdottir et al., 2013). For phylogenetic analysis, 40 18S rDNA sequences were used. The sequences were aligned using MAFFT (Katoh and Standley, 2013). The ambiguously
TABLE 1 | Phaeocystis globosa strains information used in this study.

| Strains          | Geographic origin | 18S rDNA Accession number | Sampling time | Longitude (E°) | Latitude (N°) |
|------------------|-------------------|---------------------------|---------------|----------------|--------------|
| CNS00066         | Beibu Gulf, China | MN927483                  | 2019.02       | 109.00         | 20.50        |
| CNS00069         | Beibu Gulf, China | MN927485                  | 2019.02       | 109.00         | 21.17        |
| CNS00087         | Vietnam sea area  | MN927500                  | 2019.08       | 108.13         | 10.60        |
| CNS00262         | Vietnam sea area  | MW675291                  | 2019.08       | 108.13         | 10.60        |
| CNS00266         | Vietnam sea area  | MW675290                  | 2019.08       | 108.13         | 10.60        |

aligned regions were further manually edited and adjusted using MEGA7.0 (Kumar et al., 2016). There were a total of 1562 positions in the final dataset. The maximum likelihood (ML) tree and aBayes (BI) tree based on 18S rDNA were inferred using W-IQ-TREE, with best substitution model TNe + I + G4 was selected using default parameters of W-IQ-TREE (Trifinopoulos et al., 2016). The support for nodes was assessed by performing 1000 bootstrap replicates. For analysis of sequence variability between Phaeocystis strains, 1633 positions in the final 18S rDNA sequence matrix, 420 positions in the 18S rDNA V4 sequence matrix were obtained, 171 positions in the 18S rDNA V9 sequence matrix were obtained. Numbers of pair-wise variations sites among P. globosa strains were calculated using BioEdit (Hall, 1999).

DNA Preparation and Genome Shotgun Sequencing

Cultures at the exponential growth phase were harvested and concentrated by centrifugation. Two sequencing methods were used for the genomes in this study. All five strains obtained in this study were sequenced using Illumina sequencing, while the strain CNS00066 was also sequenced using PacBio Sequel II sequencing technology.

For Illumina sequencing, total nucleic acids were extracted using the OMEGA HP Plant DNA Mini Kit (OMEGA Biotek, Inc., United States) and quantified using a NanoDrop One spectrophotometer (Labtech International Ltd., Uckfield, United Kingdom). DNA samples of five P. globosa strains were prepared for whole genome sequencing. DNA degradation and contamination were monitored on 1% agarose gels. DNA concentration was measured using Qubit® DNA Assay Kit in Qubit® 2.0 Fluorometer (Life Technologies, CA, United States). Sequencing libraries were generated using NEBNext® DNA Library Prep Kit following manufacturer’s recommendations and indices were added to each sample. The genomic DNA was randomly fragmented to a size of 350 bp by shearing, then DNA fragments were end polished, A-tailed, and ligated with the NEBNext adapter for Illumina sequencing, and further PCR enriched by generic adapter P5 and P7 oligos. The PCR products were purified (AMPure XP system) and resulted in libraries were analyzed for size distribution by Agilent 2100 Bioanalyzer system and quantified using real-time PCR. Qualified libraries were sequenced on an Illumina platform according to effective concentration and data volume at Novogene (Beijing, China). Sequencing was done using NovaSeq PE150 (Illumina, San Diego, CA, United States).

For strain CNS00066, the long-read DNA sequencing technology PacBio Sequel II was used for constructed the complete mitochondrion genome. For PacBio Sequel II sequencing, high quality genomic DNA was extracted using a modified CTAB method. RNase A was used to remove RNA contaminants. The quality of the DNA was checked by agarose gel electrophoresis. One long insert library (15 kb) was prepared, which was sequenced using the PacBio Sequel II platform (FraserGen, Wuhan, China). We generated 6.44 Gb HiFi reads.

Quality Control and Assembly

For Illumina sequencing data, methods of quality control and assembly were similar with Song et al. (2020). Raw reads in FASTQ format were first processed through a series of quality control (QC) procedures with in-house C scripts by (1) removing reads with ≥10% unidentified nucleotides (N); (2) removing reads with >50% bases having Phred quality <5; (3) removing reads with >10 bp aligned to the adapters, allowing ≤10% mismatches; (4) removing putative PCR duplicates generated by PCR amplification in the library construction process (read 1 and read 2 of two paired-end reads that were completely identical). The filtered reads were assembled with SPAdes (Bankevich et al., 2012) and GetOrganelle (Jin et al., 2020).

The resulting assembly scaffolds from the mtDNA were determined using BLAST (Johnson et al., 2008) of the complete mtDNA of the strain CNS00066 and publicly available partial mtDNA of P. globosa strain Pg-G(A) (NC_021637) (Smith et al., 2014). One mtDNA scaffold which contained one N region was obtained for each strain, respectively. Finally, clean reads were mapped to the assembled mtDNA to validate and to correct the wrong bases, using BWA (Li and Durbin, 2009) and SAMtools (1.9) (Li et al., 2009) and visualized using IGV (Thorvaldsdottir et al., 2013). We successfully completed the repeat region 1 in the strains CNS00069, CNS00087, CNS00262, and CNS00266, all of which were validated using Sanger DNA sequencing. The sequences were further validated by aligning sequencing reads against the assembled sequences using BWA (Li and Durbin, 2009) and SAMtools (1.9) (Li et al., 2009) and visualized using IGV (Thorvaldsdottir et al., 2013).

For PacBio Sequel II sequencing data, data were mapped to partial P. globosa mtDNA (KC967226) with minimap2 (version 2.5) (Li et al., 2018) using the default parameters, and a total 7.03Mb of potential mitochondrial HiFi reads were obtained. Reads depth of coverage was >160×. The mtDNA HiFi reads were locally assembled with MECAT2 software (default parameter) (Xiao et al., 2017) to obtain circular mitochondrial config 43,585 bp. The mtDNA was validated through an
additional round of alignment using BWA (Li and Durbin, 2009) and SAMTools (1.9) (Li et al., 2009) and visualized using IGV (Thorvaldsdottir et al., 2013).

In order to facilitate comparative analysis, the starting sites of mtDNAs were all adjusted to cox1.

Annotation of *P. globosa* mtDNA and Phylogenetic Analysis of Haptophyta

The mtDNAs was annotated with GeSeq (Tillich et al., 2017), ORF-FINDER (Rombel et al., 2002), and BLAST homology searches. Transfer RNAs (tRNAs) and ribosomal RNAs (rRNAs) were identified using tRNAscan-SE 2.0 (Lowe and Chan, 2016) and RNAmmer (Lagesen et al., 2007), respectively. The annotated data were deposited into NCBI GenBank database under the accession number of MW435860-MW435863, and MW435865. The graphical gene map was designed with the OrganellarGenomeDRAW program (Greiner et al., 2019). The phylogenetic tree of Haptophyta was constructed and the analysis involved 18 haptophyte taxa with two cryptophyte members as an outgroup. The dataset was composed of 3959 sites and assembled from the following 13 proteins: ATP6, ATP9, COX1, COX2, COX3, COB, NAD1, NAD2, NAD3, NAD4L, NAD4, NAD5, and NAD6. All amino acid sequences were aligned using MAFFT (Katoh and Standley, 2013). Regions showing poor alignment were trimmed with trimAl v1.2 (Capella-Gutierrez et al., 2009) using the parameter -automated1, and total amino acid sequence were concatenated using ALTER (Glez-Pena et al., 2010). There was a total of 3959 positions in the final dataset. Maximum likelihood (ML) tree and aBayes (BI) tree were inferred using W-IQ-TREE. The dataset was partitioned by protein, and best substitution model of each partition was estimated using default parameters of W-IQ-TREE (Trifonovopoulos et al., 2016), the mtZOA + G4 substitution model for ATP6, COX2, COX3,COB, NAD1, NAD3, NAD4L proteins, mtZOA + I substitution model for ATP9 protein, mtZOA + I + G4 substitution model for COX1 protein, mtInv + F + G4 substitution model for NAD2 protein, mtZOA + F + G4 substitution model for NAD4 protein, mtZOA + F + I + G4 substitution model for NAD5 protein, cpREV + F + I + G4 substitution model for NAD6 protein. Bootstrap analysis was performed using the ultrafast bootstrap approximation with 1000 replicates.

Synteny Analysis

The synteny comparison of seven mtDNAs of *Phaeocystis* was visualized using Mauve ver. 2.3.1 under the progressive mode (Darling et al., 2010). The gene content and synteny of strain CNS00069, strain Pg-G(A) and *P. antarctica* strain CCMP1374 compared to that of strain CNS00066, respectively, was performed using circos-0.69 (Krzywinski et al., 2009).

Repeat Analysis

A dot-plot similarity matrix of *P. globosa* strain CNS00066 mtDNA against itself was done with Dotter (Barson and Griffiths, 2016), in order to show repeat regions. The repeat regions were analyzed using Tandem Repeats Finder (Benson, 1999) and local BLAST using this mtDNA as the database. For stem–loop structure prediction and calculation of minimum Gibbs free energies in DNA, we utilized RNAstructure used default parameters (Bellaousov et al., 2013). Here, we considered only stem–loop structures which were max expect results predicted, although numerous smaller and less stable structures. The repeat units sequence matrix was aligned by MAFFT (Katoh and Standley, 2013), and the ambiguously aligned regions were further manually edited and adjusted by eye. The sequence similarity was analyzed using BioEdit (Hall, 1999).

RESULTS

Structural Features of the mtDNAs and Base Composition

Five *P. globosa* strains (CNS00066, CNS00069, CNS00087, CNS000262, and CNS00266) obtained in this study were identified based on comparative analysis of 18S rDNA sequences. The five *P. globosa* strains were nested within *P. globosa* monophyletic clade with robust support (Figure 1). The numbers of nucleotide differences between *P. globosa* strains were 0-16 nucleotides, which were much smaller than that of other congeneric species (19-74 nucleotides). V4 region of 18S rDNA was also analyzed, and the numbers of nucleotide differences between *P. globosa* strains were 0-6 nucleotides, which were much smaller than that of other congeneric species (11-20 nucleotides). 18 of 25 *Phaeocystis* 18S rDNA sequences have complete V9 region, and we analyzed the V9 region of these 18 strains. The number of V9 region nucleotide differences between *P. globosa* strains were 0-2 nucleotides, which was indistinguishable from other congeneric species (0-20 nucleotides). This indicated that V9 region does not provide enough resolution for species identification in *Phaeocystis*.

Of these five *P. globosa* strains, CNS00066 was a solitary-celled (Figure 1), which never formed colonies under laboratory culture condition, although it was originally isolated from a *P. globosa* colony. CNS00066 cells were 3-4.5 µm in diameter with two flagella and one haptonema emerging from a depression in the cell body. Strains CNS00069, CNS00087, CNS000262, and CNS00266 had both solitary cells and colonies, with cells being 4-8 µm in diameter, and colonies being up to 0.3 cm in diameter. Some cells in colonies had flagella and haptonema, while other did not.

The mtDNA of *P. globosa* strain CNS00066 was a circular molecule of 43,585 bp (Figure 2A). It had two large blocks of repeat regions. The repeat regions segregated the coding region into two parts and the genes were encoded on two strands, respectively, suggesting that the genome may be transcribed in two units. Aside from the repeat region, the mtDNA was relatively compact with 92.8% of sequence specifying genes and structural RNAs, with only 7.2% non-coding regions. The overall AT content of the mtDNA of CNS00066 was 70.6%, which was similar to mtDNAs of other species in the

1 http://blast.ncbi.nlm.nih.gov/
2 http://ogdraw.mpimp-golm.mpg.de/
3 http://rna.urmc.rochester.edu/RNAstructureWeb/
FIGURE 1 | The phylogenetic analysis of the haptophyte species based on 18S rDNA sequences. The evolutionary history was inferred by using the Maximum Likelihood method and aBayes (BI) method based on TN93 + I + G4 substitution model. The analysis involved 40 18S rDNA sequences. There were a total of 1562 positions in the final dataset. Evolutionary analyses were conducted using W-IQ-TREE. Numbers at the branches represent aBayes support/ultrafast bootstrap support (%), respectively. Only bootstrap values above 0.50 aBayes support and 50% ML bootstrap support are shown. The strains obtained in this study were bold. Morphology of strain CNS00066 was shown.
FIGURE 2 | Circular map of the *P. globosa* mtDNA and phylogenetic analysis of haptophyte species. (A) Circular map of the mtDNA of *P. globosa* strain CNS00066. Genes were colored according to the functional categories listed in the index below the map. The GC content is indicated on the inner circle. The repeat regions (repeat 1 and repeat 2) are in gray. Genes showed inside and outside the circle are transcribed clockwise and counter-clockwise, respectively. (B) Phylogenetic tree of haptophyte species based on 13 mitochondrial proteins (ATP6, ATP9, COX1, COX2, COX3, COB, NAD1, NAD2, NAD3, NAD4L, NAD4, NAD5, and NAD6). Phylogenetic tree was reconstructed on the IQ-TREE web server (Trifinopoulos et al., 2016). Numbers at the branches represent aBayes support/ultrafast bootstrap support (%), respectively. Our strains were shown in bold.
Coccolithophyceae (also known as Prymnesiophyceae, consisting of Phaeocystales, Isochrysidales, and Prymnesiales) (66.7-71.7%), which was higher than that of species in Pavlovophyceae (Pavlovala) (60.8-62.7%) (Table 2). The longest intergenic spacer was located between trnR-ACG and trnS-UGA, which was 379 bp.

**Phylogenetic Analysis of Haptophyte Species**

To explore the evolutionary relationship between *P. globosa* and other haptophyte members, we constructed a phylogenetic tree using protein sequences encoded by 13 PCGs in the mtDNAs that were shared by all haptophyte members (Figure 2B). The mtDNAs of five strains obtained in this study clustered with other *P. globosa* strains in a monophyletic clade with robust support. The mtDNAs of strain CNS00066 (MW435860) clustered with *P. globosa* (KC967226) with moderate support. The mtDNAs of four strains CNS000669 (MW435863), CNS00067 (MW435865), CNS000262 (MW435861), and CNS000266 (MW435862), which were isolated from the Beibu Gulf in China and Vietnam sea area, clustered together in a different clade. The species in Coccolithophyceae, Pavlovophyceae, and Rapppephyceae all formed well supported branches, and Rapppephyceae was sister to the Pavlovophyceae. The phylogenetic relationship was generally agreed with the 18S rDNA-based phylogenetic relationship in this study and previous results basing mitochondrial and plastid dataset (Kawachi et al., 2021) with minor differences, which Rapppephyceae was sister to the Coccolithophyceae.

**Gene Content and Synteny Analysis of Phaeocystis Species**

The six mtDNAs of *P. globosa* (five obtained in this study and KC967226) shared an identical number (19) of PCGs (Figures 3A-C). Four of these genes (*atp4, atp6, atp8* and *atp9*), which encode subunits of the ATP synthase complex, were identified. Seven genes encoding NADH dehydrogenase subunits, *nad1*-6 and *nad4L*, were detected. Three genes encoding complex IV (cytochrome c oxidase subunits), *cox1, cox2* and *cox3* were identified. Four genes encoding ribosomal proteins, *rpl16, rps12, rps14* and *rps3* were detected. One gene encoding complex III (cytochrome c reductase) *cob* was identified. A total of 25 tRNA genes and two rRNA genes (*rrnL* and *rrnS*) are present in the *P. globosa* mtDNA. 5S rRNA gene was not detected. None of the genes contained intron except *rrnL* (23S rDNA gene) in mtDNA of *P. globosa* strain Pg-G(A), which contained a single intron.

We performed a comparative analysis of seven mtDNAs of the genus *Phaeocystis*. Comparison of genome organization and gene order were shown in Figure 3A and Supplementary Figure 1. Five conserved gene blocks (blocks a-e) were identified among mtDNAs of *Phaeocystis*. Within *P. globosa* strains, two gene arrangement events were discovered. The first was mtDNAs of strain CNS000666 and strain Pg-G(A), which showed identical gene orders (blocks a-b-c-d-e), and the second were mtDNAs of strains CNS00069, CNS00087, CNS00262, and CNS00266, which showed another identical gene order, and blocks order was a-b-d-e-c. Compared with the first genome organization, the block b was inverted and reversed, and the block c and the blocks d-e were reciprocally translocated in the second genome organization. A high degree of syntenic conservation was identified between *P. globosa* strains CNS000666, Pg-G(A) and *P. antarctica*, except that block e (*nad5*) was inverted and reversed in *P. antarctica*. Compared to *P. globosa*, another additional tRNA gene *trnE-UUC* was found in *P. antarctica*.

**Comparative Analysis of mtDNAs**

Comparison of 18 mtDNAs of species in the phylum Haptophyta (Phaeocystales, Isochrysidales, Prymnesiales, Pavlovales, and Pavlovulinae) revealed interesting features in gene content (Figure 4). They shared 36 core genes, including 14 PCGs (*atp4, atp6, atp9*, *cob, cox1, cox2, cox3, nad1, nad2, nad3, nad4, nad4L, nad5, and nad6*), two rRNAs (*rrnL* and *rrnS*), and 20 tRNAs genes (*trnA-UAG, trnD-GUC, trnE-UUC, trnF-GAA, trnH-GUG, trnI-GAU, trnK-UUU, trnL-AAA, trnL-UAG, trnM-CAU, trnM-CAU, trnQ-UAG, trnR, trnS-UCU, trnS-GCU, trnT-UUC, trnV-UAC, trnW, and trnY-GUA*). These mtDNAs also showed important differences. For example, *rpl14* and *rps19* were only found in Pavolaves (Pavlovophyceae). The *rps14* was absent from *Chrysochromulina* sp. NIES-1333 (Prymnesiales) and *Pavulumina ranunculiformis* NIES-3900, the *rps3* was absent from *Chrysochromulina* sp. NIES-1333 (Prymnesiales), Pavolaves, Haptophyta sp. isolate H2 and *P. ranunculiformis* NIES-3900. The *nad9* was found only in *P. ranunculiformis* NIES-3900. The *atp8* gene was absent only in Isochrysidales, while the *dam* was found only in the *Emiliania huxleyi* (Isochrysidales). The *dam* gene, which encodes a DNA adenine methyltransferase, is common in bacterial genomes. Using tblastn search, we found that *dam* gene was also present in a closely related species *Gephyrocapsa oceanica* (Isochrysidales) and distantly related *Aureococcus anophagefferens* (Ochrophyta) mtDNAs, but absent from other studied mtDNAs. It may be gained via horizontal gene transfer. Further investigation is needed to test this possibility. Compared with mtDNAs of other Haptophyte species, the *Chrysochromulina* sp. NIES-1333 mtDNA encoded three extra *trnM-CAU* genes and one extra *trnS-ACU* gene, leading to a total of six *trnM-CAU* genes and three *trnS* genes. The six *trnM-CAU* genes were dispersed in genome, with sequence identities ranging from 36.4% to 100%. In contrast, no obvious similarity was found among three *trnS* genes. The *rps8* gene was only found in Isochrysidales and Prymnesiales. *P. antarctica* (Phaeocystales) had an extra *trnE-UUC* gene. The SS rRNA gene was found only in Pavolaves (Pavlovophyceae). The *trnG-UGC, trnP-UGG* and *trnS-UGA* genes were only absent in Haptophyta sp. isolate H2. The *trnW* gene contained UCA anticodon in Coccolithophyceae and CCA anticodon in Pavlovophyceae. The *trnR* gene contained AGC anticodon in Coccolithophyceae and Pavlovophyceae, a UCG anticodon in Rapppephyceae. Alternative start codons were used in different haptophyte mtDNAs. AUG was the primary start codon in haptophyte mtDNAs and only known start codon in mtDNAs of *Phaeocystis*. In addition, GUG may serve as the translation initiation codon for *atp4, rps8* in *E. huxleyi*, *atp9*, *nad4*, and *rpl14* in *D. viridis, atp4* in Haptophyta sp. isolate H2, *atp9, rpl14*, and *rps14* in *Pavlova*. The UUG may serve as the translation initiation codon.
TABLE 2 | The known mitochondrial genomes from 19 taxa in the haptophyte members.

| Class                | Order       | Species                     | Accession number | The known repeat length (> 250 bp) | Size (bp) A+T (%) | References               |
|---------------------|-------------|-----------------------------|------------------|-----------------------------------|-------------------|--------------------------|
| Coccothrahisiales   | Phaeocystales | Phaeocystis globosa CNS00066 | MW435860         | two, 6.7 kb + 14 kb               | 43,585            | 70.6 This study          |
|                     |             | Phaeocystis globosa CNS00069 | MW435863         | ~343 bp                           | 24,330 (partial)  | 66.7 This study          |
|                     |             | Phaeocystis globosa CNS00087 | MW435865         | ~1.4 kb                           | 25,390 (partial)  | 67.2 This study          |
|                     |             | Phaeocystis globosa CNS00262 | MW435861         | ~705 bp                           | 24,764 (partial)  | 66.9 This study          |
|                     |             | Phaeocystis globosa CNS00266 | MW435862         | ~334 bp                           | 24,328 (partial)  | 66.7 This study          |
|                     |             | Phaeocystis globosa Pg-G(A)  | KC067226          | ~709 bp                           | 24,477 (partial)  | 69.5 Smith et al., 2014  |
|                     |             | Phaeocystis antarctica      | JN131834 & JN131835 | ~763 bp                           | 19,177 + 8,370 (partial) | 70.3 Smith et al., 2014 |
| Isochrysidales      |             | Emiliana huxleyi COMP1516   | JN022704          | ~1.9 kb                           | 28660             | 71.5 Smith and Keeling, 2012 |
|                     |             | Emiliana huxleyi COMP373     | AY342361          | ~2.2 kb                           | 29,013 (partial)  | 71.7 Puerta et al., 2004 |
| Prymnesiales        |             | Chrysochromulina tobinii COMP291 | NC_036938         | none                              | 24,009 (partial)  | 67.5 Hovde et al., 2014  |
|                     |             | Chrysochromulina parva UW 1161 | AB990144         | two, ~1.6 kb + 1.6 kb             | 34,291            | 70.0 Nishimura et al., 2014 |
|                     |             | Haptophyta sp. isolate H2    | MK18944           | none                              | 25,180            | 77.9 Wideman et al., 2020 |
| Pavlovophyceae      | Pavloales   | Pavlova lutheri COMP1325     | HQ908424          | four, ~6 kb                       | 34,086 (partial)  | 62.7 –                  |
|                     |             | Pavlova sp. NNA-492         | MN564259          | three, ~7.3 kb                    | 36,202            | 62.5 Hulatt et al., 2020 |
|                     |             | Diacronema viridis voucher KMMCC 0113 | MW044630         | ~3.8 kb                           | 29,282            | 60.8 –                  |
|                     |             | Diacronema viridis culture CCMP620 | MW044629         | ~3.8 kb                           | 29,282            | 60.8 –                  |
| Rappephyceae        | Pavlovulinales | Pavlovulina ranunculiformis NIES-3900 | LC564891 & LC564892 | none                            | 11,199 + 9,863 (partial) | 54.1 Kawachi et al., 2021 |

*–* represent no references.

for ATP6, ATP9, NAD4, NAD4L, NAD9 in P. ranunculiformis NIES-3900, ATP9, COX1, COX2, COX3, COB, NAD1, NAD4, RPL16, and RPS3 in Chrysochromulina sp. NIES-1333, COX3 in C. tobinii CCMP291, COX2 and NAD2 in Haptophyta sp. isolate H2, and RPS14 in genus Pavlova.

We also compared PCGs in mtDNAs of Haptophyte species with that of Ancoracysta twista, Rhodomonas salina (Cryptophyceae), Aureococcus anophagefferens (SAR clade, Pelagophyceae), Phaeodactylum tricornutum (SAR clade, Bacillariophyceae), Lotharella oceanica (SAR clade, Chlorarachniophyceae) (Figure 5). These six mtDNAs shared 16 genes, ATP6 and ATP9 genes encoded subunits of the ATP synthase complex, COX1, COX2 and COX3 genes encoded complex IV (cytochrome c oxidase subunits), COB gene encoded encoding complex III (cytochrome c reductase), NAD1, NAD2, NAD3, NAD4, NAD5 and NAD6 genes encoded NADH dehydrogenase subunits. RPS3, RPS12, and RPS14 genes encoded ribosomal proteins. These genes are probably the most essentially important core genes and perform basic functions. Interestingly, P. globosa mtDNA contained the lowest number (19) of PCGs, compared to other five mtDNAs.

Variable Repeat Lengths and Locations Among mtDNAs of Phaeocystis globosa Strains

To test our hypothesis that plasticity of repeat regions may be an important driver of P. globosa mtDNA diversity in evolution, we constructed mtDNAs of four additional P. globosa strains (CNS00069, CNS00087, CNS00262, and CNS00266). The mtDNAs of these strains were still incomplete, suggesting the presence of complex repeat sequences that remain to be resolved.
Nevertheless, all genes were successfully identified in these sequences, suggesting that the missing sequences were repeat regions most likely without containing any genes. Aside from the repeat regions, mtDNAs were also relatively compact which was similar to that of strain CNS00066. AT contents of these strains were 66.7-67.2%, which were a little lower than the complete mtDNA of CNS00066.

The mtDNAs of all *P. globosa* strains contained two noticeable repeat regions with varying lengths in different strains: repeat 1 and repeat 2 (Figure 6A). The length of complete repeat regions repeat 1 (larger than 250 bp were analyzed here) was rather variable among different *P. globosa* strains, which varied substantially between 334 bp-6.7 kb, with the longest repeat found in the strain CNS00066, and the shortest repeat found in strain CNS00266, suggesting that the repeat regions might be plastic. The gray rectangles in Figure 6A showed the end of the scaffold (represented using a segment of 1000N), and these regions were repeat 2 which are not yet constructed. The still unresolved repeat 2 within all *P. globosa* strains was located at the beginning of two transcription units, while repeat 1 was located at the ending of two transcription units. Within the mtDNA of the *P. antarctica* strain CCMP1374, three repeat regions were found, among which only one complete repeat region was obtained (763 bp in length), and two unfinished repeat regions were located at the ends of two transcription units. The current known repeats length of *Phaeocystis* strains was substantially shorter than the corresponding repeat in strain CNS00066.

The locations of the repeat regions were also variable among different strains. Within the strains CNS00066 and Pg-G(A), two repeat regions were located between block c and block d, block e and block a, respectively. Within strains (CNS00069, CNS00087, CNS00262, and CNS00266), two repeats were between block a and block b, block e and block c, respectively. Gene structure rearrangement was close to repeat regions, such as the block b in strains (CNS00069, CNS00087, CNS00262, and CNS00266) was inverted compared with that of CNS00066. This suggested that rearrangement in mtDNA of *Phaeocystis* may have occurred through repeat-mediated recombination.

**Characterization of Repeat Units in the *P. globosa* Strain CNS00066**

Two complex repeat regions were observed in mtDNA of strain CNS00066 (Figure 6B). Six repeat units (A-F) of different length (13 bp-203 bp) were identified in repeat regions in the strain CNS00066 (Figure 6C and Table 3). The number of copies of each repeat unit varied dramatically and the copies of each repeat unit were not identical. For example, repeat unit A (∼163 bp) had 21 copies with sequence identity ranging from 94.4 to 100%; Repeat unit B (∼51 bp) had 252 copies with sequence identity ranging from 63 to 100%; Repeat unit C (∼63 bp) had 48 copies with sequence identity ranging from 92 to 100%; Repeat unit D (34 bp) had 22 copies with sequence identity ranging from 91.1 to 100%; Repeat unit E (13 bp) had 18 copies with sequence identity ranging from 92.3 to 100%; Repeat unit F (∼203 bp) had only two copies with sequence identity of 99.0%. Interestingly, the 5′ 26 bp of the repeat unit C showed high similarity to repeat unit B, suggesting possible evolutionary relationship between these two repeat units.

Repeat region 1 can be divided into two segments repeat 1a and repeat 1b. A large inverted repeat p1 (which consisted of a palindrome sequence A-E-D-B- B-D-E-A) encompassing 573 bp spanning repeat 1a and repeat 1b. Repeat 1 was composed of repeat units A, B, D and E in various arrangements, mainly including B-D-E-A (∼260 bp) repeat units, D-E-A (∼210 bp) repeat units, and B-D-A (∼248 bp) repeat units.
Repeat 2 also consisted of two segments, repeat 2a and repeat 2b, and a large inverted repeat p2 (which is a palindrome arrangement of repeat units B-F-F-B, 588 bp) spanning repeat 2a and repeat 2b. Repeat 2 featured repeat units B, C and F. Maximal 106 consecutive tandem copies of repeat unit B were within repeat 2a. Repeat units B and C showed various arrangements with B-C (114 bp) repeat units being the common arrangement.

The palindromic sequences p1 and p2 which were adjacent to inversion points, exhibiting a strong potential to form stable
FIGURE 3 | Multicollinearity of seven mtDNAs of genus Phaeocystis members. (A) The genomes were aligned using Mauve 2.3.1. Each colored block indicates a region of synteny among the seven genomes. The block a contain cox1-trnG-UCC-rps3-trnR-UCU-trnI-GAU-atp9-trnC-GCA- atp6-trnM-CAU-trmD-GUC-nad4-trnL-UAA-nad2-cob-trnM-CAU gene cluster, block b contain trnT-UGU-nad3-trnM-CAU- atp8-trnR-ACG-trnS-UGA gene cluster, and the block c contain trnW-UCA-nad1-rps14-trnN-GUU-trnF-GAA-cox2-trnV-UAC-rps12-trnH-GUG-trnY-GUA-nad6-trnQ-UUG-trnL-UAG-rpl16-nad4L-trnE-UUC-trmP-UGG gene cluster, and the block d contain rnl-atp4-coc3-mts gene cluster, and the block e contain nad5 gene. (B) Comparison of genome organization and gene order among the genomes. The square brackets indicated the region of synteny among the genomes. Lines connecting blocks indicated putative homology. Compared with P. globosa, P. antarctica have the additional tRNA gene trnE-UUC, which is labeled by an asterisk (*). (C) The mitochondrion genomes content and synteny of strain CNS00069 and strain Pg-G(A) compared to that of strain CNS00066, respectively.

stem–loop structures (Supplementary Figure 2). The repeat units A, B, and C were also predicted to form stable stem-loop DNA structures (Figure 6D), and repeat unit B was predicted to fold into double-hairpin elements. Variation sites were circled in red. No stem-loop DNA structure was found in repeat units D, E and F.
Homologous Repeat Sequence Found in *P. globosa* mtDNAs

Because repeat regions were identified in all sequenced mtDNAs of *Phaeocystis* species, we hypothesized that the repeat units identified in different mtDNAs share homology. To test this hypothesis, we analyzed the repeats of mtDNAs of five *P. globosa* strains obtained in this study, in addition to the *P. globosa* strain Pg-G(A) and the *P. antarctica* strain. Results showed that homologous repeat unit A could be identified in all *P. globosa* mtDNAs (Figure 6E), but not in the *P. antarctica* mtDNA.

The repeat unit A showed variable copy numbers, different length and sequence similarities among different strains (Figure 6E). The lengths varied between 55 bp-161 bp. The repeat unit A appeared three times in strain CNS00069, 12 times in strain CNS00087, six times in strain CNS00262, three times in strain CNS00266, and five times in strain Pg-G(A). No homologous sequences were found within *P. antarctica*. The results indicated the repeat unit A may share a common ancestor within *P. globosa*.

DISCUSSION

The mtDNAs are generally compact with short intergenic regions (Gray et al., 1998; Boore, 1999). However, the sizes of mtDNAs...
FIGURE 6 | Continued
Song et al. Large Differences in *P. globosa* mtDNAs

FIGURE 6 | Repeat analysis of *P. globosa*. (A) Comparison of the repeat regions position and length among *P. globosa* strains. The black blocks represent the complete repeat regions, and their length was marked under blocks. The gray rectangle showed the end of the scaffold (marked with 1000N in the sequence), and this region is another repeat region which is unobtainable until now. The genome ends in a red vertical line. (B) Dot plot similarity matrix of *P. globosa* strain CNS00066 mtDNA against itself. Dots in the nucleotide similarity matrix represent regions of sequence similarity. The main diagonal represents the mtDNA on the x-axis matching against its partner of the y-axis, Dots adjacent to the main diagonal correspond to repeat DNA and the result indicated two large repeat regions. Dot plots were generated with Dotter under default parameters. (C) Repeat structures in *P. globosa* strain CNS00066 mtDNA. Six repeat units (A-F) of different length (13 bp-203 bp) were identified in repeat regions in the strain CNS00066. (D) Repeat units A, B, and C exhibit a strong potential to form hairpin secondary structures. (E) The repeat unit A showed variable length, sequence similarity and copy numbers among different strains. The yellow arrow boxes represent repeat unit A.
Hemiselmis andersenii (Kim et al., 2008), diatom of unit may have been amplified among different repeat regions and repeat 2 in strain CNS00066. This indicated that repeat in the same strains, such as repeat unit B both exist in repeat identified sequence homology among different repeat regions. Repeat unit could be ubiquitous mutational hotspots. We also the sequences among different repeat unit copies within strain regions were results of amplification of a few repeat units. The repeat regions have been reported between related chlorophyte species Haematococcus lacustris and Stephanosphaera pluvialis, suggesting recent common ancestor of these two species (Zhang et al., 2019; Smith, 2020).

All P. globosa strains shared same numbers genes, and these gene sequences were mostly conserved among all P. globosa strains, consistent with previous observations that mtDNAs are generally compact (Gray et al., 1998). Genes are encoded on both DNA strands and their orientation suggests the presence of two transcription units within P. globosa mtDNAs, e.g., starting from the repeat region 2 and ending in repeat region 1 in both transcription directions in strain CNS00066.

In addition to differences in repeat content among mtDNAs of different strains, genome rearrangement events were also identified. Repeat regions, especially palindromic sequences were adjacent to rearrangement of the mtDNA, suggesting that they may play an important role in genomic rearrangements, transcription and replication, as proposed for other algae including red alga Chondrus crispus (Leblanc et al., 1995) and cryptophyte algae Rhodomonas salina (Hauth et al., 2005). Microhomologies have also been associated with tandem duplications and structural variation in plant mtDNAs (Xia et al., 2020). Interesting, the palindromic sequence p2 are coincident with a change in the direction of “cumulative GC skew.” Additionally, the palindromic sequence p2 had minimum CG content (Supplementary Figure 3), suggesting that the palindromic sequence p2 may correspond to the origin of replication.

Among the known 18 mtDNAs of haptophyte species, nine were reported as complete sequences, and other nine were reported as partial sequences. Nine full-length mtDNAs all contain repeat regions of various lengths ranging from 1.9 to 20.7 kb (Puerta et al., 2004; Smith and Keeling, 2012; Hovde et al., 2014, 2019; Nishimura et al., 2014; Smith et al., 2014; Hulatt et al., 2020; Wideman et al., 2020). The mtDNA of the P. globosa strain CNS00066 had the longest repeat region among all constructed haptophyte mtDNAs. The repeat regions were usually composed of tandem repeat formed by multiple repeat units, such as Phaeocystis strains and Chrysochromulina tobinii CCMP291, Pavlova strains, and Diacronema viridis strains. Chrysochromulina tobinii CCMP291 has the second longest repeat regions in Haptophyta, which was

### Table 3: Repeat units within mtDNA of P. globosa strain CNS00066 and their sequence homology.

| Repeat unit | Copies number | Sequence length | Sequence matrix length | PID |
|-------------|---------------|-----------------|------------------------|-----|
| A           | 21            | 161-167 bp      | 162bp                  | 94.4-100% |
| B           | 252           | 46-54 bp        | 46bp                   | 63-100%   |
| C           | 48            | 63-64 bp        | 63bp                   | 92.0-100% |
| D           | 22            | 34bp            | 33bp                   | 91.1-100% |
| E           | 18            | 13bp            | 13bp                   | 92.3-100% |
| F           | 2             | 203-204 bp      | 204 bp                 | 99.0%     |

TABLE 3: Repeat units within mtDNA of P. globosa strain CNS00066 and their sequence homology.

Can vary considerably among related species and even between strains of same species, which have been found to be driven by differences in repeat content or the inclusion of introns (Liu et al., 2017). The mtDNAs sequences of two Ulva pertusa strains Up1 and Up2 were different by 4.7 kb different in length, which was due to two additional group II introns in two genes (cox1 and cox2) and tandem duplication mutations in non-coding regions in the larger mtDNA (Liu et al., 2017). A pair of 3.5 kb-long, cox1-harbouring repeats were found only in two Nannochloropsis strains CCMP527 and CCMP537, while intergenic regions of the six mtDNAs were generally conserved (Wei et al., 2013). Metschnikowia yeasts mtDNAs vary extensively in size and mainly contributed by different numbers and sizes of introns (Lee et al., 2020).

This study reported the first full-length mtDNA of P. globosa, which was 43,585 in size. This large mtDNA size was driven by two large complex repeat regions, whose combined length was almost half of the entire mtDNA size. Indeed, the length of repeat sequences in P. globosa was the longest repeat region among all revolved haptophyte mtDNAs. The repeat regions were results of amplification of a few repeat units. The sequences among different repeat unit copies within strain were not identical, and base substitutions and indels existed. Repeat unit could be ubiquitous mutational hotspots. We also identified sequence homology among different repeat regions in the same strains, such as repeat unit B both exist in repeat 1 and repeat 2 in strain CNS00066. This indicated that repeat unit may have been amplified among different repeat regions within mtDNA. The structural features of the repeat regions of P. globosa have some similarities compared to other algae such as cryptophyte Rhodomonas salina (Hauth et al., 2005) and Heniselmis andersenii (Kim et al., 2008), diatom Phaeodactylum tricornutum (Oudot-Le Secq and Green, 2011), and Chlorophyte Volvox carteri (Smith and Lee, 2010) which all were composed of several different repeat units, and usually include tandem repeat, reverse repeat and palindromic sequences.

Among P. globosa different strains, the length and location of repeat regions, number of repeat units were highly variable, but there is some regularity. For example, repeat unit A was identified in all P. globosa strains. It was not found in P. antarctica, suggesting that it may be P. globosa-specific. The repeat unit A retained obvious sequence similarity among different strains, and this indicated repeat unit A may have a shared origin, existing in the mtDNA of the most recent common ancestor of strains. The sequence similarity of repeat unit A among different strains was often lower than that among different copies within strains, suggesting that the repeat regions acquired mutations in different strains. The fast mutation rates of repeat regions have also been reported in species Ulva lactuca (Liu et al., 2020) and Ulva pertusa (Liu et al., 2017). Sequence similarity of repeat regions have been reported between related chlorophyte species Haematococcus lacustris and Stephanosphaera pluvialis, suggesting recent common ancestor of these two species (Zhang et al., 2019; Smith, 2020).
∼9.3 kb and consisted of four repeat units ranging from 85 bp to ∼1.5 kb. *Chrysochromulina* sp. NIES-1333 showed a very different mitochondrial structure from other haptophyte strains described above. The *Chrysochromulina* sp. NIES-1333 genome (34,291 bp) has two small intergenic repeats of 1624 bp and 1630 bp, and missing a large, complex repeated sequence array.

Seven of nine partial mtDNAs were obviously truncated in the repeat regions, including those of *Pavlova lutheri* CCMP 1325 and all *Phaeocystis* strains. No obvious repeat regions were found in *Chrysochromulina parva* UW 1161, Haptophyta sp. isolate H2 and *P. ranunculiformis* NIES-3900, and their mtDNAs are partial sequences and have not been successfully assembled. Homologous sequences of the repeat region of *P. globosa* have not been discovered in other haptophyte species. Length of repeat region of different *P. globosa* strains was variable. The repeat length polymorphism maybe used as molecular marker for specific strains.

Phylogenetic analysis using mtDNAs revealed that Rappephyceae was sister to the Pavlovophyceae, while Prymnesiophyceae split from their common ancestor earlier. However, the support was only moderate (Figure 2B), suggesting that the evolutionary relationships among these three classes (Rappephyceae, Pavlovophyceae, and Prymnesiophyceae) might be more complicated. Indeed, our 18S rDNA-based phylogenetic analysis and a recent study suggested that Rappephyceae was sister to the Prymnesiophyceae, while Pavlovophyceae split from their common ancestor earlier (Kawachi et al., 2021). Their exact evolutionary relationship will be resolved with more genomes for specific strains.

**CONCLUSION**

In this study, we successfully constructed the full-length mtDNA of *P. globosa* for the first time. It was a circular genome with a size of 43,585 bp, encoding 19 PCGs, 25 tRNA genes and two rRNA genes. Surprisingly it was found to contain two large-sized repeat regions of 6.7 kb and ∼14.0 kb in length, respectively. The combined length of these two repeat regions accounted for almost half of the entire mtDNA. The length and the repetitive nature of these two repeat regions made it challenging to construct the complete length of full-length mtDNA of *P. globosa*. Further analysis revealed that the repeat regions consisted of combinations of several of repeat units. To explore the nature of repeat sequences in other *P. globosa* strains, we have also constructed mtDNAs of four additional *P. globosa* strains (one from Beibu Gulf, China and three from Vietnam). Comparative analysis revealed that corresponding repeat regions were readily identified in all strains but with different lengths. A homologous repeat unit A was found to be prevalent among all *P. globosa* strains, but with different numbers of copies and length. In addition to differences in repeats, these mtDNAs also differed in gene structure rearrangement. Another surprising result identified in this project was the plasticity of the repeat regions of the mtDNAs of different *P. globosa* strains. Thus, repeats may be an important driving force of mtDNA evolution. In addition, the mtDNAs of different *P. globosa* strains showed variations, suggesting that the mtDNAs can be used as a super molecular marker for distinguishing different strains.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are publicly available. This data can be found here: https://www.ncbi.nlm.nih.gov/nuccore/MW435860,MW435863, MW435865,MW435861,MW435862; https://www.ncbi.nlm.nih. gov/nuccore/MN927483,MN927485,MN927500,MW575291,MW 575290, https://dataview.ncbi.nlm.nih.gov/object/SRR14711855, and http://www.ncbi.nlm.nih.gov/bioproject/732653.

**AUTHOR CONTRIBUTIONS**

NC conceived of the project. HS and YC organized the strain selection, cultivation, DNA preparation, and genome sequencing. HS organized the assembly, annotation, quality control, and analyzed the data with suggestions from others. HS and NC wrote the manuscript with inputs from others. FL revised the manuscript. All authors read and approved the final version of the manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2021. 676447/full#supplementary-material

Supplementary Figure 1 | The mitochondrion genomes content and synteny of *P. antarctica* strain CCMP1374 compared to that of strain CNS0066.
Supplementary Figure 2 | The partial palindromic (contains the repeat units near loop) sequences of p1 and p2. They exhibit a strong potential to form hairpin secondary structures.

Supplementary Figure 3 | GC content and GC skew of P. globosa strain CNS00066 mtDNA. GC content (black) and GC skew (green and purple). For GC skew, the center line indicates the average GC skew value for the genome. Green shading above the line denotes GC skew values greater than the genome average, whereas purple shading below the line denotes GC skew values less than the genome average. Major and minor tick marks on the outermost and innermost circles show nucleotide position on the genome in 1 kb increments. The arrow points to loop of the palindromic sequence p2 in repeat region 2.

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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