The first two mitochondrial genomes for the genus Ramaria reveal mitochondrial genome evolution of Ramaria and phylogeny of Basidiomycota

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Abstract
In the present study, we assembled and analyzed the mitogenomes of two Ramaria species. The assembled mitogenomes of Ramaria cfr. rubripermanens and R. rubella were circularized, with sizes of 126,497 bp and 143,271 bp, respectively. Comparative mitogenome analysis showed that intron region contributed the most (contribution rate, 43.74%) to the size variations of Ramaria mitogenomes. The genetic contents, gene length, tRNAs, and codon usages of the two Ramaria mitogenomes varied greatly. In addition, the evolutionary rates of different core protein coding genes (PCGs) in Phallomycetidae mitogenomes varied. We detected large-scale gene rearrangements between Phallomycetidae mitogenomes, including gene displacement and tRNA doubling. A total of 4499 bp and 7746 bp aligned fragments were detected between the mitochondrial and nuclear genomes of R. cfr. rubripermanens and R. rubella, respectively, indicating possible gene transferring events. We further found frequent intron loss/gain and potential intron transfer events in Phallomycetidae mitogenomes during the evolution, and the mitogenomes of R. rubella contained a novel intron P44. Phylogenetic analyses using both Bayesian inference (BI) and Maximum Likelihood (ML) methods based on a combined mitochondrial gene dataset obtained an identical and well-supported phylogenetic tree for Basidiomycota, wherein R. cfr. rubripermanens and Turbinellus floccosus are sister species. This study served as the first report on mitogenomes from the genus Ramaria, which provides a basis for understanding the evolution, genetics, and taxonomy of this important fungal group.

Keywords: Phallomycetidae, Mitochondrial genome, Intron, Gene rearrangement, Evolution, Phylogenetic analysis

INTRODUCTION
Species from the genus Ramaria are usually known as coral fungi because of their colorful and finely branched sporocarps. Ramaria species are conspicuous ectomycorrhizal fungi associated with conifers, which are widely distributed in the northern hemisphere (Nouhra et al. 2005). The micro and macro morphological features of Ramaria are quite diverse. Some Ramaria fruiting bodies are considered to be poisonous, including Ramaria flavobrunnescens and Ramaria rufescens (Huang et al. 2009; Perez-Moreno and Ferrera-Cerrato 1995; Scheid et al. 1995).
2022), while other Ramaria species are edible, such as Ramaria botrytis, Ramaria madagascariensis, Ramaria largetii, Ramaria flava, and Ramaria formosa (Aprotosoaie et al. 2017; Khaund and Joshi 2014; Liu et al. 2014, 2015). Polysaccharide, sesquiterpenes, and glucan extracted from the basidiocarps of Ramaria exhibited antioxidant, antigenotoxic, antitumor, antimicrobial, and immunoenhancing activities (Barros et al. 2008; Bhanja et al. 2013; Centko et al. 2012; Dong et al. 2020; Fu et al. 2022; Li 2017; Liu et al. 2013; Zou et al. 2021). All in all, the accurate identification and classification of Ramaria species is very important for the development and utilization of Ramaria species.

Ramaria species belong to the order Gomphales and the subclass Phallomycetidae, which occupy a unique phylogenetic position in the higher Basidiomycetes (Nouhra et al. 2005). Taxonomy of the Gomphales has traditionally relied upon morphological features now known to be subject to parallel evolution and phenotypic plasticity. Inaccurate classification limits the discovery and utilization of Ramaria species. Subsequently, morphological and molecular data (mit-atp6-DNA, mit-12S-rDNA, and nuc-25S-rDNA) have been used to infer inter- and intra-specific relationships among Ramaria and closely related genera, which reveal that Gomphales species are closely related to taxa in the Geastrales, Phallales, and Hysterangiales (Giachini et al. 2010). Mitochondrial genomes (mitogenomes) have been reported to be an effective tool for analyzing the phylogeny of basidiomycetes. Up to now, we have reported four complete mitogenomes from Phallomycetidae, including Turbinellus floccosus (Cheng et al. 2021), Sphaerobolus stellatus (Ye et al. 2020), Dictyophora indusiata, and Phallus echi-novalvatus (Chen et al. 2020a). However, the characteristics of mitogenomes from the genus Ramaria are still unknown, which limits our comprehensive understanding of mitochondrial evolutionary pattern and genetics of Ramaria species.

Mitochondrial genome is called the ‘second genome’ of eukaryotes, which is considered to be obtained from bacteria through endosymbiosis (Gray et al. 2001). Independent origin, single parent inheritance and several available molecular markers make the mitogenome an effective tool to infer the phylogenetic relationship of species (Abuduaini et al. 2021; Basse 2010). In addition, mitochondrial gene arrangement, variation of conserved genes, dynamic changes of introns, structural variation of tRNA, codon usage and so on also provide rich genetic information for the analysis of species variation and evolution (Fonseca et al. 2021; Ren et al. 2021; Wu et al. 2021a; Zhang et al. 2021a). The mitogenome of fungi has been less studied than that of animals. Mitogenome mutations in animals are considered to be closely related to the occurrence of diseases (Du et al. 2022; Pickett et al. 2022; Zhong et al. 2022). In addition, mitogenomes have also become a powerful tool for analyzing animal systematics and population genetics (Garcia-Souto et al. 2022; Wolsberger et al. 2022; Zhang et al. 2021b). However, the relationship between fungal mitogenome variation and fungal growth, development and stress response are still unknown. The purpose of obtaining the complete sequence of fungal mitogenome is to further analyze the relationship between fungal mitochondrial sequence mutation and fungal traits. It is reported that the mitogenome of fungi varies greatly in gene arrangement, repeat sequence content, gene number and intron species, even among closely related species (Araujo et al. 2021; Fonseca et al. 2021; Li et al. 2020b; Li et al. 2021b). While most basidiomycetes have been reported to contain a series of core protein coding genes, including atp6, atp8, atp9, cob, cox1, cox2, cox3, nad1, nad2, nad3, nad4, nad4l, nad5, nad6, and rps3 (Li et al. 2021a), the large variations in the content and structure of fungal mitogenomes make it very difficult to obtain a complete fungal mitogenome. Up to now, less than 150 basidiomycete mitogenomes have been published (https://www.ncbi.nlm.nih.gov/ genotype/browse#!/organelles/), while hundreds of thousands of basidiomycetes have been described in nature. The limited number of available mitogenomes limits our understanding of the genetics and evolution of this diverse fungal group (Basidiomycota).

In the present study, the mitogenomes of two Ramaria species, including R. cfr. rubripermanens and R. rubella, were assembled and annotated. We also compared with the two Ramaria mitogenomes with other mitogenomes from subclass Phallomycetidae. The aims of this study are: 1) to reveal the characterizations of the two Ramaria mitogenomes; 2) to reveal the variations or conservativeness between Ramaria and other Phallomycetidae mitogenomes by comparative mitogenomic analysis; 3) to reveal the intron dynamics of Phallomycetidae mitogenomes; 4) to understand the phylogenetic status of Ramaria in the phylum Basidiomycota based on the combined mitochondrial gene set. This study served as the first report on the mitogenome from the genus Ramaria, which will promote our understanding of the origin, evolution, and genetics of Ramaria species and closely related genera.

**MATERIALS AND METHODS**

**Mitogenome assembly and annotation**

We assembled the complete mitogenomes of R. cfr. rubripermanens and R. rubella using the raw sequencing data from the Sequence Read Archive (SRA) database under the accession numbers of SRR5801920 and SRR3747475, respectively, which are publicly available.
(Li et al. 2018a, Miyauchi et al. 2020). The specimen of R. cfr. rubripermanens was that collected in Yunnan used in our previous study (Li et al. 2019a), deposited in the Yunnan Edible Mushroom Research Initiative, Yunnan Agricultural University (number MG17). That of R. rubella was collected by the US Joint Genome Institute (JGI; https://mycocosm.jgi.doe.gov/Ramac1/Ramac1.home.html) for the 1000 Fungal Genome project, and is stored in the USDA Forest Products Laboratory collection, Madison, WI (number UT-36052-T). Quality control steps were conducted to obtained clean reads from the raw sequencing data, which included removing adapter reads using AdapterRemoval v2 (Schubert et al. 2016) and filtering low-quality sequences using ngsShoRT 2.2 (Chen et al. 2014). The two mitogenomes of Ramaria were assembled using NOVOPlasty v4.2.1 based on the K-mer size of 29 (Dierckxsens et al. 2017). Gaps between contigs were complemented based on polymerase chain reaction and pyrosequencing. We further obtained the two circularized assembled Ramaria mitogenomes. The two complete Ramaria mitogenomes were annotated according to previously described methods (Li et al. 2018b). First, the protein-coding genes (PCGs), open reading frames (ORFs), rRNAs, tRNAs, and introns of the two Ramaria mitogenomes were annotated according to the results of MFannot (Valach et al. 2014) and MITOS (Bernt et al. 2013) based on the mitochondrial genetic code 4. ORFs longer than 100 aa were selected for further modification or annotation based on the NCBI Open Reading Frame Finder (Coordinators 2017) and BLASTP searches against the NCBI non-redundant protein sequence database (Bleasby and Wootton 1990). Intron and exon boundaries were detected by using exonate v2.2 (Slater and Birney 2005). The previously annotated tRNA genes in the Ramaria mitogenomes were further verified by tRNAscan-SE v1.3.1 software (Lowe and Chan 2016). We drew the physical maps of the two Ramaria mitogenomes using OGDraw v1.2 (Lohse et al. 2013).

**Sequence analysis**

We calculated base compositions of the two Ramaria mitogenomes and other mitogenomes from subclass Phallomycetidae using DNASTAR Lasergene v7.1 software (http://www.dnastar.com/). Strand asymmetries of the tested Phallomycetidae mitogenomes were calculated according to the following formulas: AT skew = [A − T] / [A + T], and GC skew = [G − C] / [G + C]. We determined codon usages of the two Ramaria mitogenomes based on Sequence Manipulation Suite (Stothard 2000). The nonsynonymous (Ka) and synonymous (Ks) substitution rates for the 15 core PCGs (atp6, atp8, atp9, cob, cox1, cox2, cox3, nad1, nad2, nad3, nad4, nad4L, nad5, nad6, and rps3) in the 6 Phallomycetidae mitogenomes were calculated using the DnaSP v6.10.01 (Rozas et al. 2017). The genetic distances between each pair of the 15 core PCGs were calculated using MEGA v6.06 based on the Kimura-2-parameter (K2P) substitution model (Caspersen 2016). In addition, we conducted BlastN searches of the two Ramaria mitogenomes against themselves (Chen et al. 2015) to find intra-genomic duplications or interspersed repeats in the two Ramaria mitogenomes (>50 bp). Tandem repeats in the two Ramaria mitogenomes were determined by using Tandem Repeats Finder (Benson 1999) with default parameters. We also conducted BlastN searches of the two Ramaria mitogenomes against their nuclear genomes to identify any gene fragments that transferred between their nuclear and mitochondrial genomes.

**Comparative mitogenomic analysis and intron analysis**

We conducted comparative mitogenomic analysis of the 6 Phallomycetidae mitogenomes reported to assess variations and conservations between different mitogenomes, which included genome sizes, GC contents, base compositions, codon usage, gene, and intron numbers. We calculated the contribution rate of different regions to the size variation of the two Ramaria mitogenomes, which is calculated by the following formula: size difference of region / size different of the entire mitogenome *100%. Introns in cox1 genes of the 6 Phallomycetidae mitogenomes were classified into different position classes (Pcls) according to previously described methods (Ferandon et al. 2010). Cox1 genes of the 6 Phallomycetidae mitogenomes were first aligned with the cox1 gene of Ganoderma calidophilum (Li et al. 2019b), which was used as reference in previous studies (Ye et al. 2020), by using Clustal W (Thompson et al. 1994). The same Pcl was judged by the same insert site in the corresponding reference gene and conservation of sequences, which was named according to their insert site (nt) in the reference gene. Introns belonging to the same Pcl were considered as orthologous intron and usually had a high sequence similarity (Ferandon et al. 2010).

**Phylogenetic analysis**

We constructed a phylogenetic tree of 84 Basidiomycota mitogenomes using two phylogenetic inference methods based on the combined mitochondrial gene set (14 core PCGs), which included Bayesian inference (BI) method and Maximum Likelihood (ML) methods. The phylogenetic status of Ramaria species in the phylum Basidiomycota was investigated according to the phylogenetic tree. The phylogenetic tree was constructed based on previously described methods (Huang et al. 2021). Tubera calosporum from the phylum Ascomycota was used as
individual mitochondrial genes were first aligned using MAFFT v7.037 (Katoh et al. 2019), and then concatenated into a combined mitochondrial gene dataset using the SequenceMatrix v1.7.8 (Vaidya et al. 2011). Ambiguous regions of these sequences were trimmed. In this dataset, the alignment sequence length of each species used to construct the evolutionary tree reached 21,990 bp. We conducted preliminary partition homogeneity test to detect if there were potential phylogenetic conflicts between different mitochondrial genes. Best-fit models of partitioning schemes and evolution for the combined mitochondrial gene dataset were detected by using PartitionFinder 2.1.1 software (Lanfear et al. 2017). MrBayes v3.2.6 (Ronquist et al. 2012) was used to conduct the BI analysis. Two independent runs with four chains (three heated and one cold) each were conducted simultaneously for $1.0 \times 10^7$ generations. Each run was sampled every 1000 generations. We assumed that stationarity had been reached when the estimated sample size (ESS) was greater than 100, and the potential scale reduction factor (PSRF) approached 1.0. The first 50% samples were discarded as burn-in, and the remaining trees were used to calculate Bayesian posterior probabilities (BPP) in a 50% majority-rule consensus tree (Wang et al. 2020b). RAxML v 8.0.0 (Stamatakis 2014) was used to conduct the ML analysis. We assessed Bootstrap values (BS) through an ultrafast bootstrap approach with 1,000 replicates.

Data availability
The complete mitogenomes of R. cfr. rubripermanens and R. rubella were deposited in the GenBank database (Benson et al. 2018) under the accession numbers OM272988 and OM272989, respectively.

RESULTS
PCGs in the two Ramaria mitogenomes
In the present study, we assembled the two Ramaria mitogenomes. The assembled mitogenomes of R. cfr. rubripermanens and R. rubella were circularized, with sizes of 126,497 bp and 143,271 bp, respectively (Fig. 1). The GC content of the R. cfr. rubripermanens and R. rubella mitogenomes were 28.92% and 31.69%, respectively (Additional file 1: Table S1). Both the Ramaria mitogenomes had negative AT skews and positive GC skews. A total of 18 and 20 free-stranding PCGs were

![Fig. 1](image_url) Circular maps of the two Ramaria mitochondrial genomes. Genes are represented by different colored blocks. Colored blocks outside each ring indicate that the genes are on the direct strand, while colored blocks within the ring indicates that the genes are located on the reverse strand. The inner grayscale bar graph shows the GC content of the mitochondrial sequences. The circle inside the GC content graph marks the 50% threshold.
detected in the mitogenomes of *R. cfr. rubripersimans* and *R. rubella*, respectively. Both the *Ramaria* mitogenomes contained a whole set of core PCGs, including *atp6*, *atp8*, *atp9*, *cob*, *cox1*, *cox2*, *cox3*, *nad1*, *nad2*, *nad3*, *nad4*, *nad4L*, *nad5*, *nad6*, and *rps3* (Additional file 1: Table S2). In addition, the mitogenome of *R. cfr. rubripersimans* contained 2 non-conserved PCGs encoding proteins with unknown functions and 1 non-conserved PCG encoding a putative GIY-YIG endonuclease. The *R. rubella* mitogenome contained 2 non-conserved PCGs encoding proteins with unknown functions, 1 non-conserved PCG encoding an RNA polymerase and 2 non-conserved PCGs encoding DNA polymerases. The mitogenomes of *R. cfr. rubripersimans* and *R. rubella* contained 38 and 37 introns, respectively, which were distributed in *atp6*, *cob*, *cox1*, *cox2*, *cox3*, *nad1*, *nad2*, *nad4*, *nad4L*, *nad5*, *rns*, and *rnl* genes. Two of the 38 introns from *R. cfr. rubripersimans* mitogenome belonged to the group II, 32 belonged to the group I, and 4 were of unknown types. The mitogenome of *R. rubella* contained 3 group II introns, 30 group I introns, and 4 unknown introns. The mitogenomes of *R. cfr. rubripersimans* and *R. rubella* also contained 40 and 38 ORFs in introns, which encoded putative LAGLIDADG homing endonucleases, GIY-YIG homing endonucleases, reverse transcriptase/maturase, and proteins with unknown functions.

**RNA genes in the *Ramaria* mitogenomes**

We detected two rRNA genes in both the *Ramaria* mitogenomes, which included the small subunit ribosomal RNA (*rns*) and the large subunit ribosomal RNA (*rnl*) (Additional file 1: Table S2). The average lengths of *rns* and *rnl* genes were 1687 bp and 3778 bp, respectively. The *rns* and *rnl* genes of *R. rubella* were 29 bp and 117 bp longer than that of *R. cfr. rubripersimans*. Both the *Ramaria* mitogenomes contained 25 tRNA genes, which were folded into classical cloverleaf structures (Fig. 2). The two mitogenomes contained 2 tRNAs with different anticodons coding for arginine, serine, and leucine, and 3 tRNAs with the same anticodons coding for methionine. The size of individual tRNAs ranged from 71 to 88 bp. tRNA genes of sizes >85 bp, including *trnS*(tga) and *trnL*(tag), contained larger extra arms than other tRNAs, indicating the size variations of tRNAs were mainly due to size variations of extra arms in the *Ramaria* mitogenomes. Of the 25 tRNA genes shared by the two *Ramaria* mitogenomes, 23 contained mutational sites that varied between the two mitogenomes. A total of 83 mutational sites were detected in the 25 tRNA genes between the two *Ramaria* mitogenomes, of which 24.10% variable sites were located on the extra arm, followed by the D arm.

**Mitogenome composition analysis**

The mitogenomes of *R. cfr. rubripersimans* and *R. rubella* both contained one overlapping nucleotide (Additional file 1: Table S2), which located across the neighboring genes *nad4L* and *nad5* (−1 bp). The length of intergenic sequences in the two *Ramaria* mitogenomes ranged from 1 to 4480 bp, and the longest intergenic sequence was located between *cox3* and *atp8* in the *R. cfr. rubripersimans* mitogenome.

The intronic regions accounted for the largest proportions of the *R. cfr. rubripersimans* and *R. rubella* mitogenomes, which accounted for 47.26% and 46.85% of their mitogenomes, respectively (Fig. 3). Intergenic regions accounted for the second largest proportions of the *R. cfr. rubripersimans* and *R. rubella* mitogenomes, accounting for 33.80% and 34.31% of the entire mitogenomes, respectively, which indicated that the two *Ramaria* mitogenomes have relatively relaxed structures. Protein coding regions occupied 13.20–13.67% of the two mitogenomes. RNA coding regions accounted for the smallest proportion of the two mitogenomes (5.17–5.74%). The *R. rubella* mitogenome was 16,774 bp larger than that of *R. cfr. rubripersimans*. Introncs regions contributed the most to the size expansion of the *R. rubella* mitogenome, with the contributing rate of 43.74%. Intergenic regions contributed to 38.17% of the size variation, and protein coding regions contributed to 17.23% of the *R. rubella* mitogenome expansion.

**Codon usage analysis**

Most of the core PCGs in the 6 *Phallomycetidae* mitogenomes used ATG as start codons, except that the *cox2* gene of *Sphaerobolus stellatus* used GTG, the *nad3* gene of *R. cfr. rubripersimans* used TTG, and the *nad6* gene of *S. stellatus* used TTG as start codons (Additional file 1: Table S3). The TAA was the most common used stop codon in the core PCGs of the 6 *Phallomycetidae* mitogenomes, followed by the TAG. The start and stop codons varied greatly between the two *Ramaria* species. The *atp8* and *cob* genes of *R. cfr. rubripersimans* used TAG as stop codons, while *R. rubella* used TAA as stop codons. The *nad4* gene of *R. rubella* used TAG as stop codon, while *R. cfr. rubripersimans* used TAA as stop codon.

Codon usage analysis indicated that the most frequently used codons in the two *Ramaria* mitogenomes were AAA (for lysine; Lys), TTT (for phenylalanine; Phe), TTA (for leucine; Leu), AAT (for asparagine; Asn), ATT (for isoleucine; Ile), and TAT (for tyrosine; Tyr) (Fig. 4)).
Fig. 2. Putative secondary structures of tRNA genes identified in the two *Ramaria* mitochondrial genomes. Residues conserved across the two mitochondrial genomes are shown in green, while variable sites are shown in red. All genes are shown in order of occurrence in the mitochondrial genome of *Ramaria cfr. rubripermanens*, starting from *trnW*.
The frequent use of A and T in codons contributed to a relative high AT content in the two *Ramaria* mitogenomes (average: 69.70%).

**Repetitive sequences analysis**

We conducted BlastN searches of the two *Ramaria* mitogenomes against themselves, and identified 142 and 74 repeat elements in the mitogenomes of *R. cfr. rubripermanens* and *R. rubella*, respectively (Additional file 1: Table S4). The length of these repeat sequences ranged from 50 to 413 bp, with pair-wise nucleotide similarities ranging from 76.47 to 100%. The largest repeats were located in the intergenic region between *atp9* and *cox1* gene in the *R. cfr. rubripermanens* mitogenome. The largest repeats in the *R. rubella* mitogenome were located in the intergenic region between *trnM* and *trnE*, as well as in intergenic region between *trnE* and *trnT* genes, with each of 100 bp long. Repeat sequences accounted for 9.28% and 3.25% of the *R. cfr. rubripermanens* and *R. rubella* mitogenomes, respectively. A total of 37 and 58 tandem repeats were detected in the mitogenomes of *R. cfr. rubripermanens* and *R. rubella*, respectively (Additional file 1: Table S5). The longest tandem repeat sequence was detected in the mitogenome of *R. cfr. rubripermanens*, comprising 48 bp, which was located in the intergenic region between *cox3* and *atp8*. Tandem repeat sequences accounted for 1.49% and 1.98% of the *R. cfr. rubripermanens* and *R. rubella* mitogenomes, respectively.

We further conducted BLAST searches of the two *Ramaria* mitogenomes against their nuclear genomes to identify duplication events between nuclear and mitochondrial genomes. A total of 18 and 22 aligned fragments were detected in the mitogenome of *R. cfr. rubripermanens* and *R. rubella*, respectively (Additional file 1: Table S6). The length of these aligned fragments ranged from 100 to 1702 bp, with sequence identities ranging from 76.82 to 100%. The largest aligned fragment was found located in the third intron of *rnl* gene.
in the *R. rubella* mitogenome. The largest aligned fragment in the *R. cfr. rubripermanens* mitogenome (972 bp) was found located in the first exon of *nad2*.

A total of 4499 bp and 7746 bp aligned fragment were detected in the mitogenomes of *R. cfr. rubripermanens* and *R. rubella*, respectively.

**Fig. 4** Codon usage of the two *Ramaria* mitochondrial genomes. Frequency of codon usage is plotted on the y-axis. **a**, *Ramaria cfr. rubripermanens*; **b**, *Ramaria rubella*

**Fig. 5** Sequence information of core protein coding genes in the 6 Phallomycetidae species
Evolution and variation of core PCGs

Twelve of the 15 core PCGs showed sequence length variations, with the exceptions of \textit{atp9}, \textit{cox3} and \textit{nad4L}, which had identical gene length among the 6 \textit{Phallomycetidae} species (Fig. 5). The length variation of \textit{rps3} gene was the largest among the 15 core PCGs detected, up to 1545 bp. Among the 15 core PCGs detected, the average GC content of \textit{atp9} was the highest, reaching 39.04%, followed by \textit{cox1} gene, reaching 35.57%. The GC content of \textit{atp8} gene was the lowest, with an average of 22.29%. The GC contents of core PCGs varied among the 6 \textit{Phallomycetidae} species, indicating that there are frequent sequence mutations in core PCGs of \textit{Phallomycetidae} species. Fourteen of the 15 core PCGs contained negative AT skews, and only \textit{rps3} gene contained positive AT skew in the six \textit{Phallomycetidae} species, which showed that most core PCGs tend to evolve in the direction of T-rich rather than A-rich in the leading strand of core PCGs. The GC skews of core PCGs among different \textit{Phallomycetidae} species varied, indicating frequent G/C mutations in the core PCGs.

The \textit{rps3} gene was found had the largest average Kimura-2-parameter (K2P) genetic distance between the 6 \textit{Phallomycetidae} species among the 15 core PCGs detected, followed by the \textit{nad3} gene, which showed that these genes differentiated greatly in evolution (Fig. 6). The \textit{nad4L} gene had the smallest average K2P genetic distance between the 6 \textit{Phallomycetidae} species, indicating this gene was highly conserved. The \textit{rps3} gene had the largest non-synonymous substitutions rate (\textit{Ka}) among the 15 core PCGs, followed by the \textit{nad2} and \textit{nad3} genes. While the \textit{nad4L} gene had the smallest \textit{Ka} value. The average synonymous substitution rate (\textit{Ks}) of \textit{nad3} gene was the largest, while that of \textit{nad2} gene was the smallest among the 15 core PCGs. The \textit{Ka/Ks} values for all the 15 core PCGs were < 1, indicating that these genes were subjected to purifying selection.

Intron dynamics in \textit{cox1} genes of \textit{Phallomycetidae}

A total of 156 introns were detected in the 6 \textit{Phallomycetidae} mitogenomes, which were distributed in \textit{atp6}, \textit{cob}, \textit{cox1}, \textit{cox2}, \textit{cox3}, \textit{nad1}, \textit{nad2}, \textit{nad4}, \textit{nad4L}, \textit{nad5}, \textit{rns}, and \textit{rnl} genes (Fig. 7). Six of the 156 introns belonged to the group II (Lambowitz and Zimmerly 2004, 2011), 10 were twintrons, and the rest belonged to the group I. We did not detect any putative ORF in 27 of the 156 introns, and about 82.69% introns contained putative ORFs, which encoded putative homing endonuclease, maturases, or reverse transcriptases (Michel et al. 1989). These introns are unevenly distributed in the host gene. \textit{Cox1} gene harbored the largest number of introns, accounting for 31.41% of the total introns, followed by \textit{cob} and \textit{rnl} genes, which harbored 16.03% and 13.46% of the total introns, respectively. We found that introns have a preference for particular host genes, for example, \textit{cox1} gene tends to be rich in introns, which is considered to be independent of the length of host genes (Megarioti and Kouvelis 2020;
Michel and Ferat 1995). The \textit{atp8}, \textit{atp9}, \textit{nad3}, and \textit{nad6} did not contain any intron in the 6 \textit{Phallomycetidae} mitogenomes. Intron dynamics in \textit{cox1} gene could significantly affect organization and size of \textit{Phallomycetidae} mitogenomes. We further classified introns in \textit{cox1} genes of the 6 \textit{Phallomycetidae} mitogenomes into different position classes (Pcls) using the \textit{cox1} gene of medical fungus \textit{Ganoderma calidophilum} (Li et al. 2019b) as the reference. Introns belonging to the same Pcl were considered as orthologous introns and had similar intron sequence or structures. The 49 introns in \textit{cox1} genes of the 6 \textit{Phallomycetidae} mitogenomes could be classified into 27 Pcls, which showed the rich diversity of \textit{Phallomycetidae} intron types. The class and number of introns in different \textit{Phallomycetidae} species varied, indicating potential intron loss/gain events in \textit{Phallomycetidae} evolution. A novel intron P44 was detected in \textit{R. rubella}, which belonged to the group II and contained a putative intronic ORF encoding reverse transcriptase. No homologous intron of P44 was detected in other \textit{Basidiomycota} species.

**Gene arrangement and comparative mitogenomic analysis**

We further compared mitochondrial gene arrangements of the 6 \textit{Phallomycetidae} species, including PCGs, rRNA genes, and tRNA genes. The results showed that five of the six \textit{Phallomycetidae} species have identical gene arrangement of PCGs and rRNA genes (Fig. 8). Only \textit{Sphaerobolus stellatus} had gene displacement, including \textit{nad2}, \textit{nad3}, \textit{rps3} and \textit{atp9} genes. As far as tRNA genes are concerned, \textit{S. stellatus} had the location transfer of four tRNA genes, including \textit{trnD}, \textit{trnR}, \textit{trnW}, and \textit{trnQ}. In addition, four tRNA doubling events have occurred in the \textit{S. stellatus} mitogenome, including \textit{trnM}, \textit{trnE}, \textit{trnT}, and \textit{trnS}. The mitogenome of \textit{R. cfr. rubripermanens} and \textit{R. rubella} each had one tRNA gene doubling event involving \textit{trnM} gene, respectively.

The size of the 6 \textit{Phallomycetidae} mitogenomes varied, ranging from 50,098 to 152,722 bp, with an average size of 112,345 bp (Additional file 1: Table S1). The two \textit{Ramaria} mitogenomes are larger than the average size of \textit{Phallomycetidae} mitogenomes. The mitogenome of \textit{R. rubella} was the second largest among the 6 \textit{Phallomycetidae} mitogenomes detected, which was only smaller than \textit{Sphaerobolus stellatus} (152,722 bp) from the order \textit{Gastrolatales}. The GC content of the 6 \textit{Phallomycetidae} mitogenomes ranged from 24.3 to 31.69%, with an average GC
content of 27.46%. The two *Ramaria* mitogenomes had a higher GC content than the average value. In addition, there is a large gap (2.77%) in GC content between the two *Ramaria* mitogenomes. Four mitogenomes from *Gomphales* and *Geastrales* had negative AT skews, while the other two mitogenomes from *Phallales* had positive AT skews. Four of the six mitogenomes had positive GC skews, including *R. cfr. rubripermanens*, *R. rubella*, *Dictyophora indusiata*, and *Phallus echinovolvatus*.

Each mitogenomes contained 16–58 PCGs, and the two *Ramaria* mitogenomes contained the greatest number of PCGs. There were 4–38 introns harboured in the mitogenomes of the 6 *Phallomycetidae* species, and the two *Ramaria* mitogenomes contained the greatest number of introns. All the 6 *Phallomycetidae* mitogenomes contained two rRNA genes. In addition, 24–26 tRNA genes were detected in the 6 *Phallomycetidae* mitogenomes.

**Phylogenetic analysis**

In the present study, the phylogenetic status of 84 Basidiomycete species was assessed based on combined mitochondrial gene dataset. An identical and well-supported phylogenetic tree was obtained by using both Bayesian inference (BI) and Maximum Likelihood (ML) methods (Fig. 9). All major clades within the phylogenetic tree had a high support value (BPP ≥ 0.96; BS ≥ 98). According to the phylogenetic tree, the 84 *Basidiomycota* species could be divided into 18 major clades, corresponding to the orders *Agaricales*, *Boletales*, *Cantharellales*, *Filobasidiales*, *Geastrales*, *Gomphales*, *Hymenochaetales*, *Microbotryales*, *Microstromatales*, *Phallales*, *Polyporales*, *Pucciniales*, *Russulales*, *Sporidiobolales*, *Tilletiales*, *Tramellales*, *Trichosporonales*, and *Ustilaginales* (Additional file 1: Table S7). Phylogenetic analysis indicated that *R. cfr. rubripermanens* and *T. floccosus* are sister species, which had a close relationship. The result is consistent with previous studies, indicating the mitogenome was an effective molecular marker for analyze the phylogenetic relationship of basidiomycetes.

**DISCUSSION**

Mitogenomes have been reported to play an important role in eukaryotic growth and development, oxidative stress, and environmental response (Cai et al. 2021; Du et al. 2022; Guan et al. 2021). The mitogenome mutation of animals may lead to animal diseases and affect the metabolic level of animals (Chen et al. 2020b). However, as one of the most diverse eukaryotes on earth, the mitogenome structure and variation of basidiomycetes have been less studied (Li et al. 2022; Xu and Wang 2015). Up to now, the complete mitogenome of basidiomycetes available in public databases is less than 0.05% of the described basidiomycete species in nature. It is reported that the mitogenome of basidiomycetes varies greatly in gene content, repeat sequence, genome structure and genome size, which makes it difficult to obtain the complete mitogenome of fungi (Basse 2010; Fonseca et al. 2021; Mendoza et al. 2020). In the family *Phallomycetidae*, the largest mitogenome (152,722 bp) (Ye et al. 2020) is three times larger than the smallest one (50,098 bp) (Chen et al. 2020a). Great genome size variation has also occurred in *Ramaria* species. Previous studies have shown that the size variation of fungal mitogenome is closely related to plasmid-derived genes, repeat accumulation, dynamic changes of introns and variation of intergenic sequence (Boussau et al. 2011; Chen et al. 2021; Wang et al. 2020a; Zubaer et al. 2018). In the present study, we found that introns contribute the most to the mitogenome variation of *Ramaria* species, which is consistent with previous studies (Ye et al. 2020), which showed that introns play an important role in *Phallomycetidae* size variation.

It is reported that the ancestors of eukaryotes obtained mitochondria from bacteria through endosymbiosis.
In the long-term evolution process, most mitochondrial genes have been transferred to the nuclear genome, which is convenient for the overall regulation of cells (Adams and Palmer 2003; Barton and Jones 1983). However, some PCGs, tRNA genes and rRNA genes are still retained in the mitogenome (Burki 2016). Among them, a set of core PCGs play an important role in the energy metabolism of eukaryotes. We found that the length, codon usage and base composition of these core PCGs varied greatly between different Phallomycetidae species, even among closely related species. The effect of core PCG mutation on fungal growth needs to be further revealed. In addition, different core PCGs have different evolutionary rates and may undergo purifying selection. We also found that 23 of the 25 tRNAs in Ramaria species had site variation, and the frequency of mutation may have a certain impact on the growth or stress response of fungi. The mutation of tRNA was considered to be closely related to the efficiency of protein synthesis and may eventually affect the phenotype of eukaryotes (Hayashi et al. 2016; Lin et al. 2021; Povea-Cabello et al. 2020). In the present study, a number of non-conserved PCGs have been detected in Ramaria species, which encoded RNA polymerase, DNA polymerase, and proteins with unknown functions. RNA and DNA

Fig. 9 Molecular phylogeny of 84 Basidiomycota species based on Bayesian inference (BI) and Maximum Likelihood (ML) analysis of 14 protein coding genes. Support values are Bayesian posterior probabilities (before slash) and bootstrap (BS) values (after slash). The asterisk on the evolutionary clades indicates that the BPP value is 1 and the BS value is 100. Species and NCBI accession numbers for mitogenomes used in the phylogenetic analysis are provided in Additional file 1: Table S7.
DNA polymerase are plasmid-derived genes. In some species, they are integrated into the mitogenome, while in others, they exist independently (Wu et al. 2021a, 2021b). The dynamic change of plasmid-derived regions promoted the organization and size variations of fungal mitogenome, and finally makes the fungal mitogenome more diverse and complex (Himmelstrand et al. 2014).

In addition, some PCGs with unknown functions exist in Ramaria mitogenomes, indicating that the functions of Ramaria mitogenomes need to be further analyzed, so as to promote a comprehensive understanding of the function and origin of fungal mitogenome.

Introns are considered as potentially mobile genetic elements in fungal mitogenome, and can affect the organization and size of fungal mitogenomes (Mukhopadhyay and Hausner 2021). In the present study, we found that introns are closely related to the size variation of mitogenome in subclass Phallomycetidae. In basidiomycetes, most mitochondrial introns belong to the group I and only a few introns belong to the group II (Mullineux et al. 2010). Some introns in fungal mitogenome contain intronic ORFs, which can encode putative homing endonuclease, maturases, or reverse transcriptases (Belfort and Lambowitz 2019). Introns can be classified into different position classes according to their insertion sites on the mitochondrial PCGs. Introns belonging to the same Pcls are considered to be orthologous and have high sequence similarities (Cheng et al. 2021; Li et al. 2020a). In this study, we found that the number and types of introns in different Phallomycetidae species varied, even among closely related species, indicating that intron loss or gain events have occurred in the evolution of Phallomycetidae species. In addition, some rare introns in subclass Phallomycetidae were found to exist in a large number of species, indicating that there may be potential intron transfer events between them. We also detected a novel intron P44 in R. rubella species, which have not been detected in any basidiomycete species reported (Cheng et al. 2021). The origin and evolution of the novel intron P44 need to be further analyzed.

In this study, we also compared the mitochondrial gene arrangement of the 6 Phallomycetidae species. Five of the six Phallomycetidae species had identical arrangement of PCGs and rRNA genes, which may inherit the gene arrangement from the ancestors of Phallomycetidae species. Compared with the gene arrangement of the ancestors of Phallomycetidae species, there was a large-scale gene rearrangement in PCGs and tRNA genes of Sphaerobolus stellatus (Ye et al. 2020). We also detected the doubling event of tRNAs in S. stellatus and the two Ramaria species. The mechanism of fungal mitochondrial gene rearrangement has not been fully analyzed. Previous studies have shown that fungal mitochondrial genome rearrangement may be related to the accumulation of repeat sequences (Aguileta et al. 2014). However, we only detected 2.50% repeat elements in the mitogenome of S. stellatus, which is lower than the repeat content of the two Ramaria species, indicating that the mitochondrial gene rearrangement of Phallomycetidae species may be affected by other factors other than repeats. In addition, we detected 4499 bp and 7746 bp aligned fragments between the mitochondrial and nuclear genomes of R. cfr. rubripermans and R. rubella, respectively, indicating the potential gene fragment transfer events between mitochondrial and nuclear genomes. The coevolution of mitochondrial genome and nuclear genome affects a series of ecological adaptation processes of animals and plants (Bar-Yaacov et al. 2012), which may also be happening in Ramaria.

Ramaria species is a diverse fungal group. Some Ramaria species are edible while others are considered poisonous (Barros et al. 2008; Liu et al. 2014). The misidentification of Ramaria species may lead to serious poisonous events. However, some morphological features of Ramaria species are easy to overlap, so it is not feasible to classify Ramaria species only by morphological features. Mitochondrial genome is considered to be a powerful tool to analyze the phylogenetic relationship of eukaryotes (Luchetti and Plazzi 2021). Compared with traditional multiple molecular markers, which requires multiple PCR and pyrosequencing, mitogenome can provide more genetic information and operate more conveniently. Usually, 15 core PCGs and 2 rRNA genes can be used as molecular markers for phylogeny in mitogenome, resulting in reliable and high support rate of eukaryotic phylogeny (Li et al. 2018b, 2021a; Nie et al. 2019). Although the phylogenetic analysis based on nuclear genome provides more abundant genetic information, the acquisition cost of nuclear genome is high and the amount of data is too large, which limits large-scale access to fungal nuclear genomes. Therefore, phylogeny based on mitochondrial genome is an important choice. Ramaria species occupies a unique phylogenetic position in basidiomycetes. While no mitogenome of Ramaria species has been reported. In this study, we obtained a highly supported phylogenetic tree based on the combined mitochondrial gene set through two phylogenetic inference methods. Phylogenetic analysis shows that the phylogenetic relationships of R. cfr. rubripermans and T. floccosus species is closer than that between the two Ramaria species, which is consistent with the previous studies based on multiple molecular markers (Giachini et al. 2010), indicating that the mitogenome is a reliable tool for analyzing the phylogenetic relationship of Phallomycetidae species. More mitogenomes of
Phallomycetidae species need to be obtained to understand the population genetics and phylogeny of Ramaria.

CONCLUSIONS

In this study, we obtained two novel mitogenomes from the Ramaria genus. The complete mitogenomes of R. cfr. rubripermanens and R. rubella showed large size variations, and the intron region contributed the most to the size variation of two mitogenomes (contribution rate, 43.74%). We further detected large variations in genetic contents, gene length, tRNAs, and codon usages of the two Ramaria mitogenomes. Comparative genomic analysis detected large-scale gene rearrangements between Phallomycetidae mitogenomes, including gene displacement and tRNA doubling. We first found possible gene transferring events between the mitochondrial and nuclear genomes of the two Ramaria species. The Phallomycetidae mitogenomes have been experienced frequent intron loss/gain and potential intron transfer events during the evolution, and a novel intron P44 was found in the R. rubella mitogenome. Phylogenetic analyses showed that phylogenetic relationships of R. cfr. rubripermanens and T. floccosus species is closer than that between the two Ramaria species. The present study provides the first record of Ramaria species is closer than that between the two Phallomycetidae species. The present study provides the first record of Ramaria species, thus providing a foundation for studying the evolution, genetics, and taxonomy of this important genus and related fungal group.

Abbreviations

Mitogenome: Mitochondrial genome; PCG: Protein-coding gene; Pcls: Position classes; Ks: Synonymous substitution rates; Ka: Nonsynonymous substitution rates; BI: Bayesian inference; ML: Maximum Likelihood.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s43008-022-00100-7.

Additional file 1 Table S1. Comparison on mitogenomes among six Phallomycetidae species. Table S2. Characterization and annotation of the two Ramaria mitogenomes. Table S3. Start and stop codons analysis of 6 species from Phallomycetidae. Table S4. Local BLAST analysis of the two Ramaria mitogenomes against themselves. Table S5. Tandem repeats detected in the two Ramaria mitogenomes. Table S6. Aligned fragments between mitogenomes and nuclear genomes of the two Ramaria species. Table S7. Species information used for phylogenetic analysis in this study.

Acknowledgements

Not applicable.

Author contributions

Conceived and designed experiments: QL, LZ, and CC. Analyzed the data: LL, QW, ZW, T2, PX, and WT. Wrote and reviewed the paper: QL and CC. All authors read and approved the final manuscript.

Funding

This study was funded by the open project of Key Laboratory of Integrated Pest Management on Crops in Southwest, Ministry of Agriculture (2021).

Availability of data and materials

All data generated or analyzed during this study are included in this published article [and its Additional files].

Declarations

Ethics approval and consent to participate

Not applicable.

Adherence to national and international regulations

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

Received: 24 April 2022 Accepted: 12 July 2022

Published online: 13 September 2022

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