A chromosome-level genome assembly of *Pyropia haitanensis* (Bangiales, Rhodophyta)

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Abstract
*Pyropia haitanensis* (Bangiales, Rhodophyta), a major economically important marine crop, is also considered as an ideal research model of Rhodophyta to address several major biological questions such as sexual reproduction and adaptation to intertidal abiotic stresses. However, comparative genomic analysis to decipher the underlying molecular mechanisms is hindered by the lack of high-quality genome information. Therefore, we integrated sequencing data from Illumina short-read sequencing, PacBio single-molecule sequencing and BioNano optical genome mapping. The assembled genome was approximately 53.3 Mb with an average GC% of 67.9%. The contig N50 and scaffold N50 were 510.3 kb and 5.8 Mb, respectively. Additionally, 10 superscaffolds representing 80.9% of the total assembly (42.7 Mb) were anchored and orientated to the 5 linkage groups based on markers and genetic distance; this outcome is consistent with the karyotype of five chromosomes (n = 5) based on cytological observation in *P. haitanensis*. Approximately 9.6% and 14.6% of the genomic region were interspersed repeat and tandem repeat elements, respectively. Based on full-length transcriptome data generated by PacBio, 10,903 protein-coding genes were identified. The construction of a genome-wide phylogenetic tree demonstrated that the divergence time of *P. haitanensis* and *Porphyra umbilicalis* was ~204.4 Ma. Interspecies comparison revealed that 493 gene families were expanded and that 449 were contracted in the *P. haitanensis* genome compared with those in the *P. umbilicalis* genome. The genome identified is of great value for further research on the genome evolution of red algae and genetic adaptation to intertidal stresses.

KEYWORDS
comparative genomic analysis, genome annotation, genome assembly, *Pyropia haitanensis*, repeat annotation, whole-genome sequencing
1 | INTRODUCTION

Red algae (Rhodophyta) are an ancient eukaryotic group that extended back to 1.6–1.0 billion years ago according to the observation of the cellular and subcellular structures of multicellular rhodophytes Rafatazmania and Ramathalys in fossils using synchrotron radiation X-ray tomographic microscopy (Bengston, Sallstedt, Belivanova, & Whitehouse, 2017). Red algae comprise a monophyletic lineage of ~7,200 photosynthetic species, which belong to the Archaeplastida (Plantae) derived from primary endosymbiosis (Yoon, Müller, Sheath, Ott, & Bhattacharya, 2006). The secondary and tertiary endosymbioses of red algae have given rise to the most abundant, species-rich and ecologically significant groups of algae and other eukaryotes present on Earth today, such as cryptophytes, haptophytes, apicomplexans, stramenopiles and dinoflagellates (Archibald, 2012; Hoek, Mann, Jahns, & Jahns, 1995; Reyes-Prieto, Weber, & Bhattacharya, 2007). Genomic studies on red algae will provide valuable information on the evolution of oxygenic photosynthesis. Unfortunately, only a limited number of whole-genome data sets for red algae have been reported, including those for the hot-spring alga Cyanidioschyzon merolae, the mesophilic alga Porphyridium purpureum, the extremophilic alga Galderia sulphuraria, as well as the multicellular red seaweeds Chondrus crispus, Gracilariopsis chorda and Porphyra umbilicalis (Bhattacharya et al., 2013; Brawley et al., 2017; Collen et al., 2013; Lee et al., 2018; Nozaki et al., 2007). The genomic information of Pyropia haitanensis would help to reveal the adaptation mechanisms of intertidal seaweeds and help to reconstruct the evolutionary history of red algae.

In Rhodophyta, several species of the genus Pyropia (previously named Porphyra, and commonly called "nori") are well known for their economic value in the seaweed industry, such as P. haitanensis, P. yezoensis and P. tenera (Sutherland et al., 2011). According to the FAO’s statistics, nori production in the year 2016 was ~1.8 million tons in fresh weight with a commercial value over 1.5 billion USD (http://www.fao.org/fishery/factsheets/en). Pyropia haitanensis is a native species distributed along the coastline of south China. This species is cultivated at a large scale with the highest annual production among all the nori species. The current total annual harvest of P. haitanensis is ~88,000 tons (dry weight), which accounts for approximately 75% and more than 50% of the total nori production in China and the world, respectively (Guo et al., 2018). With the aid of a high-quality genome of P. haitanensis, modern molecular genetic techniques such as QTL mapping and GWAS will be used to identify the key loci of the important economic traits such as productivity, taste and colour, which undoubtedly will enhance the efficiency of molecular breeding of this economically important marine crop.

Pyropia haitanensis naturally inhabits a niche in the upper region of the intertidal zone (Sahoo, Tang, & Yarish, 2002). Routine tidal turning periodically exposes it to the air, and it inevitably experiences the drastic changes in environmental factors such as osmotic pressure, temperature, light and UV radiation (Blouin, Brodie, Grossman, Xu, & Brawley, 2011). P. haitanensis can survive even after losing 85%–95% of its cellular water (Wang, Mao, Kong, Cao, & Sun, 2015). The thriving nature of P. haitanensis suggested that long-term evolutionary selection has made this species highly adaptable to the combined harsh stresses of the intertidal region. Thereby, this species is considered a model of intertidal red seaweed for physiology and genetic research on stress tolerance. Due to its distinctive evolutionary position in the red algal clade, P. haitanensis might harbour different genetic mechanisms of stress tolerance from those of high plants, which are probably derived from green algae. The genome information of P. haitanensis is a valuable source for the identification of unique genetic signatures involved in environmental adaptation.

Furthermore, genome sequences of P. haitanensis with relatively higher integrity and completeness are unavailable, which has been one of the major constraints to improve research on the physiology, cytology, genetics and genomics of Pyropia. Currently, the development of high-throughput sequencing technologies for sequencing DNA, RNA and proteins has reduced sequencing time and cost, etc. Hitherto, there are already four generations. Every sequencing generation and its relevant sequencing platforms have advantages and disadvantages. Thus, it is necessary to assess their limitations and applications. Second-generation sequencing is currently the most common because of its higher throughput, but the short-read lengths and amplification biases have become disadvantages (Ari & Arikan, 2016). Single-molecule real-time (SMRT) is another sequencing technology that is currently in use, which can overcome the short-read lengths and biases without any amplification step (Roberts, Carneiro, & Schatz, 2013). The appearance of an optical map can further place short reads on genomic fragments, even those totalling several millions of bases (Neely, Deen, & Hofkens, 2011). Hence, in this study, the combined techniques of Illumina short-read sequencing, PacBio single-molecule sequencing and BioNano optical mapping were used to assemble the genome of P. haitanensis. Subsequently, gene prediction, repeat annotation, functional annotation, gene family expansion and contraction, and phylogenetic relationship were determined according to standard procedures to elucidate the gene repertoire of P. haitanensis.

2 | MATERIALS AND METHODS

2.1 Sample information

A laboratory-cultured genetically pure line, Pyropia haitanensis PH40 (♀), was used in this study to eliminate the interference caused by genotypic differences. The original thallus was collected from a nori farm in Putian, Fujian Province, China. The material was first identified by amplification of its 18S rRNA gene as described in a previous study (Müller, Sheath, Vis, Crease, & Cole, 1998), as well as by its morphologies. Single somatic cells were enzymatically isolated from the thallus, and the allele homozygous sporophytes (concho-cels) were obtained after the haploid doubling spontaneously. The genetically homogenous gametophytes were then developed from the homozygous sporophytes and cultured for DNA and RNA sample collection. Another strain PH37 (♂) used in this study was also harvested from Putian, Fujian Province, China, and purified with
the same method mentioned above. The gametophytes were cultured in a light incubator under the following conditions: 20 ± 1°C with 50–60 μmol photons-m⁻²·s⁻¹ illumination during a 12 h:12 h light-dark cycle. The culture medium of Provasoli’s enriched seawater (PES) (Starr, 1987) was refreshed every five days. To remove surface bacteria from *P. haitanensis* gametophytes, the thalli were harvested and mixed with quartz sands. Physical vibration was carried out in a homogenizer (Precellys 24), followed by several rounds of washing to remove the polysaccharides and bacteria from the surface of gametophytes. Subsequently, the samples were collected and immediately frozen in liquid nitrogen for total genomic DNA extraction using the CTAB method (Yang, Wang, Liu, & An, 1999).

### 2.2 Libraries construction

Five micrograms and 10 μg of genomic DNA were used to construct Illumina TruSeq paired-end sequencing libraries (500-bp insert sizes) and mate pair libraries (5 kb in size), respectively, according to the manufacturer's instructions. Meanwhile, a total of 10 μg of DNA was used to construct a 20-kb library using the PacBio Pacific Biosciences SMRT Bell Template Kit 1.0. To further carry out optical map construction, 2 μg of purified high molecular weight (HMW) genomic DNA was isolated and labelled according to standard BioNano protocols with the single-stranded nicking endonuclease BspQI. To assist in the genome annotation of *P. haitanensis*, total RNAs isolated from various stressful conditions (osmotic pressure, temperature, illumination, etc.) were equally mixed together to prepare the transcriptome sequencing libraries for SMRT platforms following the manufacturer’s instructions. For SMRT sequencing, full-length RNA libraries were constructed according to the manufacturer's instructions with minor modifications. To avoid overamplification of small fragments, we optimized the amplification cycle at 14 in a preliminary test. Then, three gel fractions, containing fragments >3, 2–3 and 1–2 kb, were collected and purified using the QIAquick Gel Extraction Kit. The extracted products were amplified using the 5′ Primer IIA and purified using 0.5 × AMPure beads (#A63880; Beckman, http://www.beckmancoulter.com) for subsequent sequencing.

### 2.3 Genome sequencing and assembly

To estimate the genome size of *P. haitanensis*, the low-quality reads and sequences aligning to the chloroplast (Accession no: KC464603) and mitochondrion (NC_017751) genomes of *P. haitanensis* were removed using the NGS QC Toolkit and Bowtie 2 (parameters: -v very-sensitive; version: 2.0.2) (Langmead, Trapnell, Pop, & Salzberg, 2009). Different K-mer frequencies were calculated by Jellyfish and genome size (Luo et al., 2012). For genome assembly, subreads from PacBio were used to assemble the nuclear genome of *P. haitanensis* using the RS_HGAP_Assembly3 protocol in SMRT ANALYSIS v2.3.0 with default parameters (Chin et al., 2013). Then, mate pair data sets were aligned to the above-assembled contigs using SSPACE (Boetzer, Henkel, Jansen, Butler, & Pirovano, 2010). Meanwhile, PacBio long reads were mapped to the scaffold sequences using BLASR, and the gaps that resulted from the scaffolds were filled using PBJelly2 with default parameters (English et al., 2012). Finally, Quiver was run again to polish the accurate consensus at the base level.

To improve the assembly, optical maps of the BioNano system were further used for scaffolding. A labelled DNA sample was loaded onto the Saphyr Chip nanochannel array, and the stretched DNA molecules were then imaged with the BioNano Saphyr system. Raw image data were converted into bnx files, and AutoDetect (BioNano Genomics) software generated basic labelling and DNA length information. Access (BioNano Genomics) software was used to filter and remove <150 Kb low-quality reads, and then, IrySolve (BioNano Genomics) was used to carry out the assembly of BioNano's genome maps and the 'Hybrid Scaffold' between genome maps from BioNano and sequence maps. Further gap filling using the reads that not used in the last step was achieved by RefAligner (BioNano Genomics). To remove the potential contamination of bacterial sequences in the current assembly, we applied a postprocessing step. We cut each scaffold into 100 bp overlapping 1-Kb windows and blasted them against the NT database using BLASTn. The blast results were further analysed using MEGAN to search for bacterial hits. scaffolds that met the following three criteria were considered to be bacterial contamination and removed from the final genome: (a) over 60% of windows in the scaffold had best hits as bacterial sequences with identity >70%; (b) the sequencing depth was <5; and (c) there was no cDNA support in these ‘bacterial windows.’ To assess the quality of the assembled genome, K-mer frequency distribution, the full-length transcriptome sequencing data map rate and Benchmarking Universal Single-Copy Orthologs (BUSCO) analysis were used.

### 2.4 Genetic map construction and scaffold anchoring

To construct a genetic map of *P. haitanensis*, the gametophytic blades of PH40 (♀) and PH37 (♂) were selected as parents for crossing experiments. The blades from these two pure lines were cocultured in a flask until carposporangia appeared. Then, the fertilized female blade was selected and cultured until reproductive cells were released. Subsequently, the fertilized carpospores were cultured to generate heterozygous conchocelis. The heterozygote was then confirmed using two SSR markers in our laboratory. After confirmation, the heterozygous gametophytes (F1) were then developed from the homozygous conchocelis and used to establish double haploid populations (DH). Each individual F1 gametophyte was digested with 50–60 μg of purified high molecular weight (HMW) genomic DNA were used to construct a 20-kb library using the PacBio Pacific Biosciences SMRT Bell Template Kit 1.0. To further carry out optical map construction, 2 μg of purified high molecular weight (HMW) genomic DNA was isolated and labelled according to standard BioNano protocols with the single-stranded nicking endonuclease BspQI. To assist in the genome annotation of *P. haitanensis*, total RNAs isolated from various stressful conditions (osmotic pressure, temperature, illumination, etc.) were equally mixed together to prepare the transcriptome sequencing libraries for SMRT platforms following the manufacturer’s instructions. For SMRT sequencing, full-length RNA libraries were constructed according to the manufacturer’s instructions with minor modifications. To avoid overamplification of small fragments, we optimized the amplification cycle at 14 in a preliminary test. Then, three gel fractions, containing fragments >3, 2–3 and 1–2 kb, were collected and purified using the QIAquick Gel Extraction Kit. The extracted products were amplified using the 5′ Primer IIA and purified using 0.5 × AMPure beads (#A63880; Beckman, http://www.beckmancoulter.com) for subsequent sequencing.

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Meyer, McKay, & Matz, 2012). These libraries were sequenced on an Illumina HiSeq system to generate single-end reads with a length of 50bp. Subsequently, reads were trimmed to remove sequences with adapters, those without restriction sites and those containing ambiguous bases and of low-quality value. Meanwhile, sequence reads from putative plastid and mitochondrial origins of P. haitanensis were also removed. The remaining reads were analysed using the RADtyping program v1.0 with default parameters (Fu et al., 2013) for genotyping. The markers that could be genotyped in at least 80% of offspring were used to calculate the genetic distance and draw linkage maps using JoinMap 4.0 at LOD 7.0 (Van Ooijen, 2006). The linkage group numbers were selected at a LOD threshold of more than 4.0. Meanwhile, genetic distances between markers and marker sequences were used to anchor scaffolds to the linkage groups using the R package.

2.5 | Repeat elements

Repeat elements occupy a major proportion of the nuclear DNA in most eukaryotic genomes and have been demonstrated to have structural and functional roles (Bischoff, Olmo, & Heslop-Harrison, 2015). REPEATMODELER (version: 1.0.8) was used to analyse consensus sequences of interspersed repeats in genomes of P. haitanensis (Smit & Hubley, 2008). Consensus sequences that were shorter than 80 bp were discarded (Wicker et al., 2007). The remaining consensus sequences were used as the library in REPEATMASKER (version: open-4-0-7) to predict interspersed repeat elements in the whole genome (Chen, 2004). Meanwhile, Tandem Repeats Finder (Benson, 1999) was used to identify tandem repeat sequences in P. haitanensis genome.

2.6 | Gene prediction and functional annotation

After repeats’ masking, we used a combination of de novo prediction, homology searches and transcript isoform based methods to predict gene structures of P. haitanensis. De novo prediction was performed using AUGUSTUS (Stanke et al., 2006). For homologous annotation, we queried the P. haitanensis genome scaffolds against a database containing protein sequences from five organisms (Chondrus crispus, Gracilariopsis chorda, Cyanidioschyzon merolae, Po. umbilicals and Porphyridium purpureum). At the same time, transcript isoforms of P. haitanensis were mapped to the genome using BLAST and then assembled by PASA (Haas et al., 2008). Finally, EVM was used to integrate these gene models from the above methods. To further detect the function of the protein-coding genes in P. haitanensis, the predicted protein sequences were aligned against several public databases (NR, InterPro, GO, KOG, KEGG, CAZyme and Conserved Domains Database [CDD]).

2.7 | Gene family expansion and contraction

To further examine the genome divergence and conservation among red algae, we carried out a phylogenetic analysis based on single-copy orthologous groups using the P. haitanensis genome and other five red algal genomes to build orthologous genes using ORTHOMCL (Li, Stoeckert, & Roos, 2003), with Cyanophora paradoxa as the out-group species. Genome sequences were aligned using the program MAFFT version 5 (Katoh, Kuma, Toh, & Miyata, 2005) and were further trimmed using trimAl with the option “automated1” (Capella-Gutiérrez, Silla-Martínez, & Gabaldón, 2009). Maximum likelihood (ML) analyses were conducted using RAxML-8.2.4 (Stamatakis, 2014). The best model and parameter settings were chosen according to the Akaike information criterion using PROTEST 3.0 (Abascal, Zardoya, and Posada 2005). A Bayesian phylogenetic tree was constructed using MRBAYES 3.2 under the same model (Huelsenbeck & Ronquist, 2001). Four incrementally heated Metropolis-coupled Monte Carlo Markov chains were run for 10,000,000 generations for the concatenated data set, and runs were sampled every 1000th generation. Convergence and stationarity of the log-likelihood and parameter values were assessed using TRACER v1.5 (Rambaut, Drummond, Xie, Baele, & Suchard, 2018). The initial 10% were discarded as burn-in. A time-calibrated phylogeny was inferred using a relaxed molecular clock method as implemented in BEAST v1.8.3 (Drummond, Suchard, Xie, & Rambaut, 2012). We set the most recent common ancestor with a lognormal prior, an offset of 950 Ma, and a standard deviation of 25.0 based on the divergence of Florideophyceae and Bangiophyceae (Herron, Hackett, Aylward, & Michod, 2009; Yang et al., 2016).

3 | RESULTS AND DISCUSSION

3.1 | Material identification and Genome assembly

The material used in this study was identified as Pyropia haitanensis according to its morphology, life history, as well as its reproductive structure, etc (Figure 1). The blade was 15–16 cm in length and 2-3 cm in width, with a red to brown colour. Additionally, it had an umbilicate base, which can help the blade attach to substratum. The molecular marker and alignment results also supported identification of the specimen as P. haitanensis (Figure S1). Scanning electron microscopy showed that bacteria had been removed from the surface of the algae (Figure S2). And a total of ~22.1 Gb of raw sequence data were obtained using the Illumina platform for P. haitanensis. Based on calculation of the K-mer frequency by Jellyfish, the estimated genome size of P. haitanensis was approximately 38.5 Mb (Table S1). For genome assembly, ~5.0 Gb of subreads from the PacBio RSII platform with a mean length of 5.7 kb were used to assemble the nuclear genome of P. haitanensis. A 59.7 Mb assembly was produced consisting of 1,839 contigs with an N50 of 510.3 kb. Then, the number of scaffolds built based on ~1.8 Gb of Illumina mate pair sequencing data was reduced to 1,168 and the length of N50 increased to 913.7 kb. Scaffolding using PacBio long reads allowed us to improve the assembly to 663 scaffolds (totalling 59.2 Mb) with a scaffold N50 of 912.3 kb. For optical map construction, a total of 93.8 Gb of molecular data were obtained (Table 1). Combined with optical mapping data, we finally yielded a P. haitanensis genome with a size of 53.3 Mb. Among the 195 scaffolds, 11 pseudomolecules had lengths larger
than 0.4 Mb and covered 88.4% of the genome region. The contig N50 and scaffold N50 were 510.3 kb and 5.8 Mb, respectively, and the length of the longest scaffold was 7.6 Mb (Table 2). The average GC content of this genome was as high as 67.9%, which is the highest among all the published algal genomes. The phenomenon of high GC content was also found in the Bangiophyceae species \textit{Po. umbilicalis} (65.8%) (Brawley et al., 2017) and green algae \textit{Chlamydomonas reinhardtii} (64%) (Merchant et al. 2007). Compared with the assembly results of the published macroalgae, including \textit{Chondrus crispus} (scaffold N50 = 240.0 kb), \textit{Po. umbilicalis} (scaffold N50 = 202.0 kb) and \textit{Saccharina japonica} (scaffold N50 = 252.0 kb), the assembly of \textit{P. haitanensis} genome had the fewest scaffolds and the longest N50 and the highest contiguity and coverage (Ye et al., 2015).

### TABLE 1 Genome and transcriptome sequencing information of *Pyropia haitanensis*

| Sequencing platforms | Library size | Data size (Gb) | Depth |
|----------------------|-------------|----------------|-------|
| DNA library          |             |                |       |
| Illumina             | 500 bp      | 22.1           | 220   |
| Illumina             | 5 kb        | 1.8            | 47    |
| PacBio               | 20 kb       | 6.4            | 99    |
| BioNano              | ~           | 93             | 1,860 |
| RNA library          |             |                |       |
| PacBio               | 1–2 kb      | 1.5            | 12    |
| 2–3 kb               | 1.3         | 12             |       |
| >3 kb                | 1.5         | 12             |       |

#### 3.2 Anchor scaffolds by genetic maps

The genome sequencing of male and female parents and their offspring produced 32,327,297, 35,177,866 and 1,031,682,186 reads, respectively. These reads then were mapped to the genome for subsequent genotyping. The results showed that 1,367 SNPs were shared between the two parents. One hundred and twenty-nine loci that met the linkage requirement were used to construct the genetic map. Finally, five linkage groups were constructed using these markers, with a number of markers ranging from 9 to 45. The length per group ranged from 88.6 cM to 284.0 cM, with an average of 171.4 cM. Based on the markers and genetic distance, 10 pseudomolecules representing 80.9% of the total assembly (42.7 Mb)
were anchored and orientated to the 5 linkage groups (Figure 2). Among them, pseudomolecules 12, 26, 32 and 110 were anchored to one chromosome, and pseudomolecule 9 was mapped to one chromosome. Meanwhile, pseudomolecules 13 and 27 and pseudomolecules 80 and 201 were placed on two different chromosomes, respectively, based on the markers and their distance. The remaining pseudomolecule 140 was anchored to one chromosome. The number of linkage groups established in this study is consistent with the cytological observations (Tseng & Sun, 1989; Yan et al., 2008).

### 3.3 Genome evaluation

To assess the quality of the assembled genome, three approaches were used. First, the final assembled genome size of this species (53.3 Mb) was similar to the size calculated based on the K-mer frequency distribution (46.5 Mb). Second, we obtained a total of 17,383 unigenes from the PacBio system. Then, these transcriptome sequencing data were mapped to the current assembly by BLAT (Kent, 2002), and >87.2% of PacBio isoforms could be successfully aligned. Third, we performed Benchmarking Universal Single-Copy Orthologs (BUSCO) analysis, and 85.5% of the eukaryotic single-copy genes were detected in the *P. haitanensis* genome. This number is higher than the values in *C. crispus* (84.5%) and *Po. umbilicalis* (74.3%) (Figure S3). Interestingly, we also noticed that the ‘complete’ percentage of BUSCO in red algae was generally lower than those in other species. The reason for this possibly lies in independent evolution after primary endosymbiosis, leading to great genome diversity in red algae (e.g. reduction of the genome contents of the red algae (Qiu, Price, Yang, Yoon, & Bhattacharya, 2015)). The relative lack of red algal genome information in public databases might be another reason.

### 3.4 Repeat elements

For the repeat element analysis, the results showed that the repeat elements identified in *P. haitanensis* constituted 24.2% of the whole genome, including 14.6% as tandem repeat sequences and 9.6% as interspersed repeats. Among the tandem repeats, a total of 26,822

### Table 2

| Statistics of the final assembly of *Pyropia haitanensis* genome |
|---------------------------------------------------------------|
| **Contig** | **Scaffold** | **BioNano** |
|-------------|--------------|-------------|
| Total sequences | 1,497 | 230 | 195 |
| Total bases | 57,754,774 | 50,812,391 | 53,254,677 |
| Min sequence length | 504 | 740 | 60 |
| Max sequence length | 2,019,106 | 3,335,433 | 7,561,339 |
| Average sequence length | 38,580.3 | 220,923.4 | 273,100.9 |
| N50 length | 538,396 | 1,023,154 | 5,758,810 |
| N90 length | 14,603 | 143,036 | 158,429 |
| (G + C)s | 69.9% | 71.2% | 67.8% |

### Table 3

| Composition of repeat elements in genome of *Pyropia haitanensis* |
|---------------------------------------------------------------|
| **Class** | **Order** | **Superfamily** | **Number** | **Length (bp)** | **Percentage (%)** |
|-------------|----------|----------------|------------|-----------------|-------------------|
| Interspersed repeats | LTR | Gypsy | 413 | 1,327,093 | 2.49 |
| | | Copia | 544 | 278,151 | 0.52 |
| | | Caulimovirus | 83 | 114,994 | 0.22 |
| | | Other LTR | 76 | 85,379 | 0.16 |
| DNA | | CMC-EnSpm | 83 | 80,307 | 0.15 |
| | | PIF-Harbinger | 331 | 253,169 | 0.48 |
| | | PiggyBac | 228 | 69,767 | 0.13 |
| Unknown | 10,009 | 2,874,529 | 5.40 |
| Tandem repeats | Microsatellite | 26,822 | 1,695,878 | 3.18 |
| | Minisatellite | 60,360 | 4,290,390 | 8.06 |
| | Satellite | 3,586 | 1,776,700 | 3.34 |
microsatellites were identified, accounting for 3.2% of the genome. In addition, 60,360 (8.1%) minisatellite and 3,586 (3.3%) satellite DNAs were identified. LTR elements represented the majority of the confirmed interspersed repeats, occupying 3.4% of the genome, while the DNA elements comprised 0.8% (Table 3, Figure 3). Among LTRs, 1,040 full-length LTRs were predicted, 544 of which belonged to the Copia superfamily, 413 belonged to the Gypsy superfamily and 83 belonged to Caulimovirus superfamily. The remaining 76 LTRs were not full length and occupied 0.2% of the genome. When compared with closely related species, we noticed that the Po. umbilicalis genome had a substantial repeat element (43.9%) in its 87.7 Mb genome, including 17.7% DNA transposons (15.5 Mb) and 17.0% LTR elements (14.9 Mb) (Brawley et al., 2017). Comparison of the repeat landscape of the P. haitanensis genome and those in other species in red algae (Price et al., 2019) showed that the LTRs can be attributed to genome size variation.

3.5 Gene prediction

After repeats’ masking, de novo prediction predicted 11,725 gene models for P. haitanensis. Based on the homologous protein database established from the five red algae mentioned above, 31,389 protein-coding sequences were obtained. At the same time, we predicted 11,871 gene models using PASA software. Finally, EVM was used to integrate these gene models from the above methods to obtain a gene data set with 10,930 protein-coding sequences (ORFs), which is comparable to the gene repertoire of other sequenced red algae genomes (Bhattacharya et al., 2013; Brawley et al., 2017; Collén et al., 2013; Lee et al., 2018; Nozaki et al., 2007). These protein-coding genes in P. haitanensis were further employed to analyse their functions using several public databases. We identified 7,356 and 10,374 genes that showed homology to proteins in the NR and InterPro databases, respectively (Figure S4). A total of 3,147 genes were assigned to GO
| Gene name | Gene function | P. haitanensis | Porphyra umbilicalis | Chondrus crispus | Porphyridium purpureum | Cyanidioschyzon merolae |
|-----------|---------------|---------------|---------------------|-----------------|------------------------|---------------------------|
| ROS production | | | | | | |
| RBOH | NADPH oxidase | ph10359.t1 | OSX70888.1 | ccri|XP_005718545.1 | ppurjevm.model.contig_2134.3 | Cm|XP_005535894.1 |
| | | ph07364.t1 | OSX75422.1 | ccri|XP_005719187.1 | ppurjevm.model.contig_2149.17 | Cm|XP_005538987.1 |
| | | ph06070.t1 | OSX74398.1 | ccri|XP_005718335.1 | ppurjevm.model.contig_2146.22 | |
| | | ph07507.t1 | OSX73467.1 | ccri|XP_005716000.1 | ppurjevm.model.contig_3670.1 | |
| | | ph08568.t1 | OSX75676.1 | | ppurjevm.model.contig_502.2 | |
| | | ph05196.t1 | OSX69054.1 | | | |
| | | ph03740.t1 | OSX69091.1 | | | |
| | | ph11172.t1 | OSX72018.1 | | | |
| | | ph06827.t1 | | | | |
| | | ph03938.t1 | | | | |
| AOX | in mitochondria | ph03278.t1 | OSX69369.1 | ccri|XP_005719100.1 | ppurjevm.model.contig_2288.11 | Cm|XP_005536259.1 |
| PTX | in plastid | ph07793.t1 | OSX69826.1 | ccri|XP_005712075.1 | ppurjevm.model.contig_4450.5 | Cm|XP_005536398.1 |
| ABA regulatory net | | | | | | |
| PYR1/PYL/PCAR | | | | | | |
| PP2C | type-2C protein phosphatase | ph10951.t1 | OSX76300.1 | ccri|XP_005719405.1 | ppurjevm.model.contig_3479.1 | Cm|XP_005536535.1 |
| | | ph09239.t1 | OSX79480.1 | ccri|XP_005711405.1 | ppurjevm.model.contig_510.16 | Cm|XP_005538832.1 |
| | | ph02078.t1 | OSX71532.1 | ccri|XP_005719125.1 | ppurjevm.model.contig_3807.1 | Cm|XP_00553984.1 |
| | | ph11536.t1 | OSX77048.1 | ccri|XP_005712925.1 | ppurjevm.model.contig_4456.15 | Cm|XP_00553913.1 |
| | | ph07863.t1 | OSX81030.1 | ccri|XP_005711323.1 | ppurjevm.model.contig_2501.2 | |
| | | ph10321.t1 | OSX77620.1 | | ppurjevm.model.contig_3441.20 | |
| | | ph02405.t1 | OSX69983.1 | | ppurjevm.model.contig_441.27 | |
| | | ph06642.t1 | OSX71152.1 | | ppurjevm.model.contig_3468.6 | |
| | | ph08933.t1 | | | ppurjevm.model.contig_528.2 | |
| OST1 | Protein OPEN STOMATA kinase | ph00419.t1 | OSX79527.1 | ccri|XP_005713434.1 | ppurjevm.model.contig_2031.6 | |
| | | ph03789.t12 | OSX79650.1 | ccri|XP_005713325.1 | ccri|XP_005716962.1 | ccri|XP_005718769.1 |
classifications. Based on KEGG analysis, we could annotate a total of 1,830 genes (Table S2) and a total of 317 KEGG metabolic pathways in the genome of *P. haitanensis* (Figure S5). Moreover, the CAZyme database annotation showed that a total of 303 genes in the *P. haitanensis* genome were associated with carbohydrate metabolism-related enzymes (Table S3). In addition, 7,041 genes in *P. haitanensis* were assigned to CDD 1,295 superfamilies (Table S4).

3.6 Gene family expansion and contraction

To estimate the gene family expansion and contraction, the genome of *P. haitanensis* combined with five available red algae and an out-group species was selected to define the orthologous genes. We identified 622 single-copy orthologous genes within *P. haitanensis* and the other six species, which were used in phylogenetic analyses in the following study. Analysis suggested the divergence time of *P. haitanensis* and *P. umbilicalis* was 204.4 Ma (95% highest posterior density (HPD)=164.6–249.7 Ma), indicating that *P. haitanensis* was a more recently diverged lineage in the red algae (Figure 4).

A total of 493 orthologous groups (containing 2,514 genes) harboured more *P. haitanensis* paralogs than *P. umbilicalis* and were therefore defined as the expanded gene families. They mainly encoded ATP hydrolysis, nucleic acid metabolism, purine metabolism, cytoskeleton-associated proteins, ion-transporting proteins as well as E3 ubiquitin ligase, etc., according to their Pfam annotation (Tables S5 and S6). Meanwhile, 294 groups (containing 1,218 genes) with fewer *P. haitanensis* paralogs were defined as contracted gene families. These encoded phytochelatin synthase, sucrose transporter, cytochrome c oxidase copper chaperone, etc. Although the two closely related species are similar in morphology and physiology, the existence of large amounts of expanded and contracted gene families among them suggests that different environmental pressures have shaped their specific genetic contents to adapt to their individual habitats since they diverged from each other.

3.7 ROS-ABA signalling pathway-related genes in *P. haitanensis*

ROS is an important secondary messenger that is poised at the core of signalling pathway in plants maintaining the normal metabolic fluxes and different cellular functions and responding to environment stresses (Quigley et al., 2009). The production of ROS in cell originated from NADPH oxidases (NOX) located different organelles (cell wall, chloroplast and mitochondria) (Bedard & Krause, 2007). The NOX in cell wall is also considered as ROS-generating respiratory burst oxidase (RBOH). In higher plants, RBOH is a family with more than ten members (Suzuki et al., 2011). We identified 10 members of RBOH in *P. haitanensis*, 8 in *P. umbilicalis*, 4 in *C. crispus* and 2 in *Cyanidioschyzon merolae* (Table 4). Compared to single-cell red algae, RBOH in *P. haitanensis* endured significant expanding during evolution. The numbers of AOX and PTX in *P. haitanensis* are 2, with no significant difference with other red algae species. Under the downstream signal pathway activated by ROS, MAPK cascade
is highly conserved and can be activated by phosphorylation (Xing, Ginty, & Greenberg, 1996). It plays major role in signal transduction of diverse stress responses even in combination of many stresses. The activation of MAPK cascade firstly is inhibited by MAPK repressor while induced by ROS (Son et al., 2011). The dual-specificity protein tyrosine phosphatase (DSPTP) is MAPK repressor in ROS pathway (Martell, Angelotti, & Ulrich, 1998). Only 1 was identified, *P. haitanensis*; however, 8 and 5 was identified in single-cell red algae species, *P. purpureum* and *C. merolae*, respectively. When the MAPK cascade was activated, the phosphorylation event can further activated many downstream factors, including transcript factors (TFs) etc. At present, MYB44, HSFA and ERF factors were identified to be activated by MAPK and involved in many stress and development process. We identified 16 MYB family TFs in *P. haitanensis*, including 12 MYB-like, respectively. Yet, only 1 HSFA was identified in *P. haitanensis*. There are no significant differences in the numbers of these two-type TFs in all red algae species studied. It was noting that ERF factor did not exist in either specie, which is an important TFs in ethylene signalling pathway.

ABA signalling pathway plays important in response to environmental stress, especially drought stress (Davies, Kudoyarova, & Hartung, 2005). The turning on of this pathway is dependent on the ABA receptor binding to ABA. Currently, the ABA receptor widely studied including PYR1/PYL/PCAR component. Its binding to ABA can inhibit PP2C, further inhibit OST1 kinase and activate MAPK. After that, the downstream response factors were activated. In addition, OST1 can activate the slow anion channel-associated (SLAC). We did not identify the presence of PYR1/PYL/PCAR type receptor in either red algae, but identified G protein receptor (GPCR), which is another receptor binding to ABA. The number of GPCR in *P. haitanensis* is 3. There are 10 PP2C in *P. haitanensis*, yet only 5 in *P. umbilicalis*, which indicated this gene family endured expanding in *P. haitanensis*. OST1 (1) and SLAC (2) were also identified in different red algae with no significant difference in numbers. Numerous reports highlight the importance of the ROS-ABA signalling pathway in responding to drought stress in higher plants (Cruz de Carvalho, 2008; Golldack, Li, Hartung, 2005). The turning on of this pathway is dependent on the phosphorylation event can further acti- vated many downstream factors, including transcript factors (TFs) etc. At present, MYB44, HSFA and ERF factors were identified to be activated by MAPK and involved in many stress and development process. We identified 16 MYB family TFs in *P. haitanensis*, including 12 MYB-like, respectively. Yet, only 1 HSFA was identified in *P. haitanensis*. There are no significant differences in the numbers of these two-type TFs in all red algae species studied. It was noting that ERF factor did not exist in either specie, which is an important TFs in ethylene signalling pathway.

4 | CONCLUSIONS

In this study, we reported a high-quality nuclear genome of *Pyropia haitanensis*, a red algal species of great economic, ecological and research value. We adopted multiple sequencing techniques to achieve an assembly with high contiguity and coverage. The investigation of genome characteristics and functional features yields further insights regarding the phylogenetic diversity of *P. haitanensis*. This genome will not only be a fundamental resource for deciphering the molecular mechanisms underlying the developmental processes of *P. haitanensis* and environmental adaptation mechanisms of intertidal seaweeds, but also help to reconstruct the evolutionary history of red algae.

AUTHOR CONTRIBUTIONS

Y.X.M. and D.M.W. conceived the study. C.M., X.Z.Y. and P.P.S. performed the experiments. K.P.X., G.Q.B., Y.L., F.N.K., X.H.T., Y.G. and G.Y.D. analysed and interpreted the assembly and annotations. K.P.X. and G.Q.B. performed the comparative genome analysis. C.M. and K.P.X. wrote the manuscript with input from all authors.

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DATA AVAILABILITY STATEMENT

The DNA sequencing data have been deposited into the NCBI Sequence Read Archive under the BioProject: PRJNA503796.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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