Highly Conserved Mitochondrial Genomes among Multicellular Red Algae of the Florideophyceae

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Data deposition: Newly determined and annotated 11 mitochondrial sequences of red algae have been deposited at GenBank under the accessions KF290995, KF649303–5, and KJ398158–64.

Abstract

Two red algal classes, the Florideophyceae (approximately 7,100 spp.) and Bangiophyceae (approximately 193 spp.), comprise 98% of red algal diversity in marine and freshwater habitats. These two classes form well-supported monophyletic groups in most phylogenetic analyses. Nonetheless, the interordinal relationships remain largely unresolved, in particular in the largest subclass Rhodymeniophycidae that includes 70% of all species. To elucidate red algal phylogenetic relationships and study organelle evolution, we determined the sequence of 11 mitochondrial genomes (mtDNA) from 5 florideophycean subclasses. These mtDNAs were combined with existing data, resulting in a database of 25 florideophytes and 12 bangiophytes (including cyanidiophycean species). A concatenated alignment of mt proteins was used to resolve ordinal relationships in the Rhodymeniophycidae. Red algal mtDNA genome comparisons showed 47 instances of gene rearrangement including 12 that distinguish Bangiophyceae from Hildenbrandiophycidae, and 5 that distinguish Hildenbrandiophycidae from Nemaliophycidae. These organelle data support a rapid radiation and surprisingly high conservation of mtDNA gene syntheny among the morphologically divergent multicellular lineages of Rhodymeniophycidae. In contrast, we find extensive mitochondrial gene rearrangements when comparing Bangiophyceae and Florideophyceae and multiple examples of gene loss among the different red algal lineages.

Key words: Florideophyceae, mitochondrial genome, ordinal relationship, red algal evolution, rhodophytes, Rhodymeniophycidae.

Introduction

Mitochondria (mt) supply chemical energy to cells and have a long evolutionary history that extends to the origin of eukaryotes (Gray et al. 1999; Burger et al. 2003; Koonin 2010). Most mt genomes are characterized by a compact structure (exceptions include plants that contain exceptionally large mt genomes) and maternal inheritance. Due to their ubiquitious distribution and conserved functions, complete mtDNA data

© The Author(s) 2015. Published by Oxford University Press on behalf of the Society for Molecular Biology and Evolution. This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com have been used in diverse studies including population genetics, phylogenetics, and organellar genome evolution (Unseld et al. 1997; Turmel et al. 2002; Ogihara et al. 2005; Ratnasingham and Hebert 2007; Zink and Barrowclough 2008; Morin et al. 2010). For example, analysis of mtDNA data from land plants revealed gene replacements through horizontal gene transfer and endosymbiotic gene transfer to the nucleus (Adams and Palmer 2003; Bergthorsson et al. 2003, 2004). Similar to plants, Rhodophyta (red algal) mtDNA data have been used in systematic and population studies as molecular markers (Saunders 2005; Yang et al. 2008; Freshwater et al. 2010).

Red algae belong to an anciently diverged group (Supergroup Plantae or Archaeplastida) whose ancestor hosted the plastid primary endosymbiosis (Bhattacharya et al. 2004). This phylum is composed of seven classes (i.e., Bangiophyceae, Compsopogonophyceae, Cyanidiophyceae, Florideophyceae, Porphyridiophyceae, Rhodellophyceae, and Stylonematophyceae) with the most species-rich class Florideophyceae the containing subclasses Hildenbrandiophycidae, Nemaliophycidae, Corallinophycidae, Ahnfeltiophycidae, and Rhodymeniophycidae (Saunders and Hommersand 2004; Yoon et al. 2006; Le Gall and Saunders 2007; Le Gall et al. 2010). A putative member of the Bangiophyceae played a central role in the eukaryote tree of life as a donor through a single or more likely multiple secondary plastid endosymbioses that gave rise to chlorophyll-c containing groups, such as diatoms, dinoflagellates, haptophytes, and cryptophytes (Yoon et al. 2004; Chan et al. 2011; Bhattacharya et al. 2013). Despite the importance of red algae in eukaryote evolution, only a few representative genomes (nuclear and organellar) have been characterized thus far (Ohta et al. 2003; Hancock et al. 2010; Smith et al. 2012; Janouškovec et al. 2013). Although the complete mtDNA from 26 red algal species has been determined (two Cyanidiophyceae [Leblanc et al. 1995; Jain et al. 2015], ten Bangiophyceae [Burger et al. 1999; Mao et al. 2012; Smith et al. 2012; Hwang et al. 2013; Hughey et al. 2014; Kong et al. 2014; Silva and Hughey 2014; Bangia fuscopurpurea, Bi G, Cao M, Mao Y, unpublished data, NC_026905], and 14 Florideophyceae species [Leblanc et al. 1995; Hancock et al. 2010; Zhang et al. 2012; Campbell et al. 2014; DePriest et al. 2014; Kim, Yang, Yi, et al. 2014; Kim, Yang, Boo, et al. 2014; Tablizo and Lluisma 2014; Yang, Kim, Boo, et al. 2014; Yang, Kim KM, Kim SY, et al. 2014; Kim et al. 2015; Lee et al. 2015]), the broader evolutionary history of this genome has been challenging to reconstruct because of insufficient taxon sampling and the high level of structural variation seen in the available mt genomes. Apart from a recent report of the corallinophycidean (nongeniculate) species Sporolithon durum (Kim et al. 2015), the remaining mt genome data are primarily from the Rhodymeniophycidae.

Understanding mtDNA evolution in Rhodophyta, in particular as it relates to the sister taxa Bangiophyceae and

Florideophyceae that comprise greater than 98% of red algal diversity (Guiry MD and Guiry GM 2015) is of great interest to understand mitochondrial evolution in general. The Bangiophyceae contains approximately 193 reported species, all with simple thalli, including the economically important Porphyra and Pyropia (source of sushi wrap, kim or nori) from which mtDNA from ten species has been determined (Mao et al. 2012; Smith et al. 2012; Hwang et al. 2013; Hughey et al. 2014). The Florideophyceae is the largest class of red algae containing greater than 95% (6,760 spp.) of reported species diversity in the phylum Rhodophyta [approximately 7,100 species (Guiry MD and Guiry GM 2015)]. Phylogenetic relationships of the five subclasses within the Florideophyceae are well-resolved based on molecular and ultrastructural data (e.g., Saunders and Hommersand 2004; Le Gall and Saunders 2007). The Hildenbrandiophycidae, the earliest diverging subclass (which likely lacks a gametophyte stage) has a sister relationship to the remaining subclasses (Pueschel and Cole 1982; Le Gall and Saunders 2007). Of the remaining well-supported subclasses, the Rhodymeniophycidae is the largest, containing >5,000 species (70% of red algae). In spite of the strongly supported monophyly of the subclass, relationships among the 12 rhodymeniophycidan orders remain unresolved (Saunders et al. 2004; Le Gall and Saunders 2007; Verbruggen et al. 2010).

Here, we report 11 mtDNA sequences, 3 from Ahnfeltia plicata (Ahnfeltiophycidae), Hildenbrandia rubra (Hildenbrandiophycidae), Palmaria palmata (Nemaliophycidae), and the remaining 8 from the following rhodymeniophycidan species: Asparagopsis taxiformis (Bonnemaisoniales), Ceramium japonicum (Ceramiales), Gelidium elegans (Gelidiales), Plocamium cartilagineum (Plocamiales), Riquetophycus sp. (Peyssonneliales), Schizymenia dubyi (Nemastomatales), Schimmelmannia schousboei (Acrosymphytales), and Sebdenia flabellata (Sebdeniales).

The aims of the present work were 1) to elucidate phylogenetic relationships at higher taxonomic levels and identify major evolutionary transitions in the Rhodophyta using a database of 37 mt genomes; 2) to compare the coding sequences (CDSs) content and synteny between the Bangiophyceae and Florideophyceae, and among the five florideophycean subclasses; and 3) to determine the extent of mtDNA gene synteny and loss within the Rhodymeniophycidae.

Materials and Methods

Taxon Sampling, DNA Isolation, Genome Sequencing, and Annotation

The red algal species used in this study were chosen to sample all 5 subclasses of the Florideophyceae, as well as 12 orders of the Rhodymeniophycidae (table 1). We collected nine red algal species from the field and two species from culture collections

Table 1

Red Algal Taxonomy and General Features of the Mitochondrial Genome Used in the Present Study

Taxa	Size (bp)	GC Content (%)	Number of Gene					Accession
			CDS ^a	tRNA	rRNA	Intron	Intronic ORF	
Cyanidiophyceae								
Cyanidioschyzon merolae	32,211	36.5	35	25	3	_	_	NC_000887
Galdieria sulphuraria	21,428	44.0	19	7	2	_	_	NC_024666
Bangiophyceae								
Bangia fuscopurpurea	43,517	33.0	31	23	2	5	6	NC_026905
Porphyra purpurea	36,753	33.5	29	24	2	2	2	NC_002007
Porphyra umbilicalis	29,123	31.9	25	24	2	1	1	NC_018544
Pyropia fucicola	35,035	32.5	31	23	2	4	4	NC_024288
Pyropia haitanensis	37,023	30.7	24	24	2	4	5	NC_01775
Pyropia kanakaensis	39,300	30.0	30	25	2	5	3	NC_024289
Pyropia perforata	40,042	31.8	24	23	2	5	5	KF_515974
Pyropia tenera	42,269	32.8	25	23	2	5	6	NC_021475
Pyropia yezoensis	41,688	32.7	27	27	2	5	5	NC_017837
Wildemania schizophylla	29,156	33.2	26	23	2	2	2	NC_024579
Average:	37,391	32.2	27.2	23.9	2	3.8	3.9	
Florideophyceae								
Hildenbrandiophycidae								
Hildenbrandia rubra ^b	33,066	36.4	25	18	2	2	2	NC_026055
Nemaliophycidae								
Palmaria palmata ^b	29,735	32.2	26	22	2	2	_	NC_026056
Corallinophycidae	-							_
Sporolithon durum	26,202	28.4	23	20	2	1	_	NC_023454
Ahnfeltiophycidae	-							_
Ahnfeltia plicata ^b	32,878	33.4	24	24	2	3	2	NC_026054
, Rhodymeniophycidae								_
Asparagopsis taxiformis ^b	26,097	26.7	25	24	2	1	_	NC_024843
Ceramium japonicum ^b	26,110	28.5	22	24	2	_	_	KJ398159
Chondrus crispus	25,836	27.9	29	24	4	1	_	NC_001677
Gelidium elegans ^b	24,922	29.5	23	19	2	1	_	NC_023053
Gelidium vagum	24,901	30.5	23	18	2	1	_	NC_023077
Gracilaria chilensis	26,898	27.6	25	25	2	1	_	NC_02683
Gracilaria salicornia	25,727	28.4	25	20	2	1	_	
Gracilariophila oryzoides	25,161	28.1	23	20	2	1	_	NC 01477
Gracilariopsis andersonii	27,036	28.0	27	19	2	1	_	NC_014772
Gracilariopsis chorda	26,543	27.6	26	24	2	1	_	NC_02325
Gracilariopsis lemaneiformis	25,883	27.5	26	21	2	1	_	JQ071938
Grateloupia angusta	27,943	30.2	25	19	2	2	1	NC_023094
Grateloupia taiwanensis	28,906	31.4	25	24	2	2	1	KM99923
Kappaphycus striatus	25,242	30.1	24	24	2	1	_	NC_024265
Plocamiocolax pulvinata	25,894	26.0	25	20	2	2	_	NC_014773
Plocamium cartilagineum ^b	26,187	23.6	24	25	2	2	_	KJ398160
Rhodymenia pseudopalmata	26,166	29.5	24	21	2	-	_	NC_023252
Riquetophycus sp. ^b	26,261	25.7	24	22	2	1	_	KJ39816
Schimmelmannia schousboei ^b	25,816	26.7	23	21	2	1	_	KJ398162
Schizymenia dubyi ^b	26,347	25.8	25	24	2	1	_	KJ398163
Sebdenia flabellata ^b	25,887	28.5	23	21	2	1	_	KJ398164
Average:	26,866	28.7	24.6	21.7	2.1	1.3	0.08	

 $^{\rm a}{\rm Number}$ of protein-coding genes exclude intronic-orf. $^{\rm b}{\rm Newly}$ determined mitochondrial genomes.

(UTEX; http://web.biosci.utexas.edu/utex/ and CCAP; http:// www.ccap.ac.uk) (supplementary table S1, Supplementary Material online). Prior to processing, fresh samples or silica gel-preserved vouchers were transported to the lab and cleaned several times using sterilized seawater under the dissecting microscope.

Total genomic DNA was extracted using the DNeasy Plant Mini Kit (Qiagen, Hilden, Germany). We used the Ion Torrent Personal Genome Machine (PGM; Life Technologies, San Francisco, CA) to do shotgun DNA sequencing. About 100 ng of total DNA was used for library preparation (200 bp read) based on the manufacturer's instructions. One or two independent libraries of each species were used for sequencing (supplementary table S2, Supplementary Material online). The high quality (Q20) data obtained from each species were assembled using MIRA (Chevreux et al. 1999) under default settings that were incorporated into the Ion Torrent PGM and the CLC de novo assembler under default settings in the CLC Genomics Workbench v.5.5.1 (CLC Bio, Aarhus, Denmark). The Riguetophycus sp. sample differed from the other samples in that the library was constructed and subsequently sequenced using the Illumina platform by Eureka Genomics (Hercules, CA). In this case, one paired-end, nonindexed library was constructed from the DNA of one vegetative individual (2 µg). Sequencing consisted of one lane, with a paired end run of 51 cycles. The resulting reads were assembled as above using the CLC de novo assembler. Contigs were scanned for organelle sequences using local Basic Local Alignment Search Tool (BLAST) (Altschul et al. 1997) with red algal mitochondrial genomes as search gueries. The mitochondrial contigs were then used for scaffolding using Geneious Pro v.5.3.6 (http://www.geneious.com), followed by "read mapping to reference" implemented in the CLC Genomics Workbench. The average coverage of mitochondrial consensus sequences was higher than $74 \times$ for all samples (supplementary table S2, Supplementary Material online). Sanger sequencing was used for gap-filling with each mtDNA sample to produce complete genomes (supplementary fig. S1, Supplementary Material online).

Open-reading frame (ORF) finding and gene prediction were done using BLAST (BLASTn and BLASTx), GeneMarkS (Besemer et al. 2001), and Geneious Pro. Large and small subunits of ribosomal RNA (rRNA) were identified using BLASTn with published red algal rRNAs as queries, whereas transfer RNAs (tRNA) were identified using the tRNAscan-SE Search Server (http://lowelab.ucsc.edu/ tRNAscan-SE/; Schattner et al. 2005). All protein-coding, rRNA, and tRNA genes, and intron boundaries within genes in the novel mitochondrial genomes were confirmed by multiple sequence alignment with published red algal data using the T-Coffee package (http://www. tcoffee.org; Di Tommaso et al. 2011). Novel mt genome data were deposited in GenBank (table 1).

Phylogenetic Analyses

To reduce tree artifacts due to high DNA divergence, amino acid sequences were used to reconstruct phylogenies. The inferred amino acid sequences of every protein-coding gene from 37 mtDNAs were aligned using the T-Coffee package (under the default setting of -mode fmcoffee) and then refined manually. This concatenated alignment comprised 6,436 amino acids (data set c24p, 24 CDS; gene no. 1–24 in supplementary table S3, Supplementary Material online). The alignment and tree files are available from TreeBase (http://treebase.org/treebase-web/search/study/summary.html? id=15150&x-access-code=7bdd870b7e5e820b3737098d36e 3cec4&agreement=ok), the RedToL website (dbdata.rutgers.edu/redseq/), and from the corresponding author upon request.

The phylogeny and statistical support for monophyletic groups were inferred using the maximum-likelihood (ML and ML bootstrap analysis) method with RAxML v.8.1.17 (Stamatakis 2006) under the specified models. We used the best fit independent amino acid model (amino acid substitution matrix with empirical frequencies [F] + gamma distributed rate heterogeneity [G]) for each gene partition (24 genes) of the concatenated data set to reduce the heterogeneity effect among genes on the phylogeny. The specific amino acid substitution matrix for each partition was selected by using the protein model selection procedure provided in the RAxML program website (http://sco.h-its.org/exelixis/resource/down load/software/ProteinModelSelection.pl). This script automatically determines the best-scoring model from all available (18 amino acid substitution matrices with/without amino acid freguencies estimation) to use with RAxML. The best-fit models for the genes were as follows: MTZOA for atp6 (ATP synthase F0 subunit 6), atp9, cob (apocytochrome b), cox2 (cytochrome c oxidase subunit 2), cox3, nad1 (NADH dehydrogenase subunit 1), nad2, nad3, nad4, nad5, JTTDCMUT for atp8, rpl16 (ribosomal protein subunit L16), rpl20, rps3 (ribosomal protein subunit S3), rps11, rps12, CPREV for cox1, nad4L, nad6, sdh3 (succinate:cytochrome c oxidoreductase subunit 3), FLU for sdh2, and JTT for sdh4, secY, ymf39. The independent evolutionary models for the partitions of each gene were unlinked by "-g" options. The "-f a" option was used in RAxML for the simultaneous search for the best likelihood tree and rapid bootstrap analysis with "-# 1,000" (1,000 bootstrap replications) with default options of "-i" (automatically optimized SPR branch rearrangement for heuristic search) and "-c" (25 distinct rate categories).

To test concordance of monophyletic groups between different data sets, we prepared 23 sub-data sets from the c24p (table S3) based on three criteria. First, based on the mean amino acid (AA) P-distance of each gene alignment that was calculated with PAUP* v4.0b10 (Swofford 2002), 24 CDSs were sorted by mean values (ranging from 0.046 [*atp9*] to 0.621 [*rps3*]), and then the genes with high mean P-distances

were incrementally removed for the sub-data sets by 0.05 intervals (PV1–8). Second, 24 CDSs were ordered by AA length of gene (ranging from 156 [*atp8*] to 684 [*nad5*]), and the genes were incrementally removed for the sub-data sets by 50 AA length intervals (PL1-8). Third, 24 CDSs were sorted by translation products as listed in supplementary table S3, Supplementary Material online. The electron transport, oxidative phosphorylation protein (PE), and ribosomal protein (PR) genes were grouped as sub-data sets that include PE (*atp-cob-cox-nad-sdh-secY-ymf39*), PE1-5 (*atp6-9*, *cob-cox1-3*, *nad1-6*, *sdh2-4*, and *secY-ymf39*), and PR (*rpI-rps*). We compared the ML bootstrap values among the full c24p and 23 sub-data sets (100 resampling for bootstrap analysis; supplementary table S2 and fig. S2, Supplementary Material online).

Bayesian Inference (BI) was done using MrBayes v.3.2.4 (Ronguist and Huelsenbeck 2003) using the best-fit model for the data set of the c24p. Preliminary searches under the mixed aamodel (prset aamodel=mixed;) suggested the JONES amino acid substitution model as the best-fit for BI. The JONES+G model was used as a fixed model [prset aamodel=fixed(jones)] for all partitions. All model parameters were unlinked for each partition by using the program settings [prset applyto=(all) ratepr=variable; lset applyto=(all) rates=gamma ngammacat=4; unlink revmet=(all) statefreq=(all) shape=(all);]. Twenty million Metropolis-coupled Markov Chain Monte Carlo (MC³) simulations were completed with the following parameters: Two independent runs with different random start points, one cold chain and five heated chains for each run, and tree sampling at every 1,000th generation. The Bayesian Posterior Probabilities (BPP) of the monophyletic nodes were calculated using trees (total 2,002) post burn-in; that is, 10 million generations.

Alternative tree topologies were evaluated using the Approximately Unbiased (AU) and Shimodaira Hasegawa (SH) tests (Shimodaira 2002) implemented in the CONSEL v.1.20 package (Shimodaira and Hasegawa 2001). All 24-CDS combined data (c24p) were used for the paired-site test with the best phylogeny and alternative phylogenies. We selected five alternative relationships of rhodymeniophycidan orders based on sub-data set analyses that had more than 50% of nodes supported with greater than 80% maximum-likelihood bootstrap (MLB) (supplementary fig. S2, Supplementary Material online). The test was performed with 100,000 bootstrap replicates under the same amino acid model settings and partitions as used in the best ML search. We repeated the test three times to confirm the reproducibility of the analysis.

Results and Discussion

Characteristics of 11 Novel mt Genomes

We determined 11 mt genomes from representatives of the 4 florideophycean subclasses; that is, Ahnfeltiophycidae,

Hildenbrandiophycidae, Nemaliophycidae, and Rhodymeniophycidae (table 1 and supplementary fig. S1, Supplementary Material online). These data were combined with existing mt genomes from 10 bangiophycean, 2 cyanidiophycean, and 14 florideophycean taxa, including 6 genomes recently determined by our group (Kim, Yang, Yi, et al. 2014; Kim, Yang, Boo, et al. 2014; Yang, Kim, Boo, et al. 2014; Kim, Yang, Boo, et al. 2014; Yang, Kim, Boo, et al. 2014; Yang, Kim KM, Kim SY, et al. 2014; Kim et al. 2015; Lee et al. 2015). This resulted in a database of 37 mt genomes from red algae for the phylogenetic and comparative analyses.

The newly determined mt genomes have a compact gene arrangement with 44–52 predicted genes (i.e., 22–26 proteincoding, 2 rRNAs, and 18–25 tRNAs; supplementary fig. S1. Supplementary Material online), which are shared among most red algae (Hancock et al. 2010; Zhang et al. 2012; Yang, Kim KM, Kim SY, et al. 2014). Differences were found in the absence of a few genes including atp8, nad4L, rpl20, sdh3, sdh4, secY, and ymf39, for example, the lack of atp8 in Ahnfeltia, Hildenbrandia, and Plocamium (see supplementary table S3, Supplementary Material online). The rrl (large subunit rRNA) of Ahnfeltia, Hildenbrandia, and *Palmaria* includes an intron (with/without an intronic ORF); the cox1 gene contains an intron with an ORF in Ahnfeltia and Hildenbrandia and without an ORF in Plocamium; and trnl in all species includes an intron except in Ceramium, Gelidium, and Hildenbrandia (supplementary fig. S1 and table S3, Supplementary Material online). The mt genomes of the Bangiophyceae were larger in size (i.e., min. = 29.1 kb, max. = 43.5 kb, average = 37.4 kb) and had a higher GC-content (min. = 30%, max. = 33.5%, average = 32.2%) than those of the Florideophyceae (size: min. = 24.9 kb, max. = 33.1 kb, average = 26.9 kb; GC-content: min. = 23.6%, max. = 36.4%, average = 28.7%). Within the Florideophyceae, H. rubra contained the largest mt genome and the highest GC content (33.1 kb, 36.4% GC; see table 1).

Mitochondrial Genome-Based Red Algal Phylogeny

Phylogenetic analysis of concatenated mitochondrial data (c24p) from 37 red algal species strongly supports the currently accepted relationships of Rhodophyta (fig. 1). These include a close relationship of Bangiophyceae and Florideophyceae (Müller et al. 2001; Saunders and Hommersand 2004), the recognition of each of the five subclasses, and intersubclass relationships within the Florideophyceae (Le Gall and Saunders 2007; Verbruggen et al. 2010).

The most notable finding of the phylogenetic analysis was the resolution of ordinal relationships among 21 species of Rhodymeniophycidae (fig. 1). The subclass Rhodymeniophycidae is the most morphologically diverse of the red algae and includes greater than 5,015 reported species under the current 12-order taxonomic system (Guiry MD and Guiry GM 2015). The lineages in this subclass likely underwent a rapid radiation as indicated by the mostly short and dense

intermediate nodes among the orders, with only a few ordinal relationships resolved with weak to moderate statistical support (Le Gall and Saunders 2007; Verbruggen et al. 2010). Our mtDNA phylogeny nonetheless provides novel insights into the relationships of some of the rhodymeniophycidan groups. Within the subclass, the Bonnemaisoniales (A. taxiformis) diverged first and is sister to the remaining 11 orders with high support (i.e., 100% MLB and 1.0 BPP) in the c24p data set (node-a) and in the sub-data sets (15 sub-data sets support node-a with >91% MLB; supplementary fig. S2, Supplementary Material online). The Peyssonneliales (Riquetophycus sp.) and Gigartinales (Chondrus and Kappaphycus) grouped together with strong support (nodeb in fig. 1, 100% MLB and 1.0 BPP) using c24p and 15 subdata sets (>91% MLB in supplementary fig. S2, Supplementary Material online) in this restricted taxa data set of Gigartinales sensu lato. However, the monophyly of node-c (11 orders) was less well supported, that is, 75% MLB and 1.0 BPP in the c24p. The topology of sub-data sets (i.e., the sister relationship of Asparagopsis and Kappaphycus as shown in fig. 1 as alt-1) was not rejected in both AU and SH tests (supplementary fig. S3C, Supplementary Material online). A larger number of taxa are likely required to resolve the phylogenetic relationships of the Bonnemaisoniales, Gigartinales, and Peyssonneliales (Krayesky et al. 2009).

After the divergence of the Gelidiales (Gelidium spp.; 94% MLB and 1.0 BPP; node-d), two monophyletic clades (node-e and node-h) were found, that is, node-e: Plocamiales (Plocamiocolax pulvinata and P. cartilagineum) + Ceramiales (C. japonicum) (node-f) + Acrosymphytales (S. schousboei) clade (58% MLB and 0.59 BPP) and node-h: Rhodymeniales (Rhodymenia pseudopalmata) + Sebdeniales (Se. flabellata) (node-k) + Halymeniales (Grateloupia spp.) (node-i) +Gracilariales (Gracilaria spp., Gracilariopsis spp., and Gracilariophila oryzoides) + Nemastomatales (Schiz. dubyi) (node-i) clade (67% MLB and 0.99 BPP). Interordinal relationships within each clade are mostly congruent with results from prior studies (Le Gall and Saunders 2007; Le Gall et al. 2010; Verbruggen et al. 2010). However, interordinal relationships of the Acrosymphytales, Ceramiales, and Plocamiales were unresolved and an alternative topology (alt-2 in fig. 1) was not rejected by the tree topology tests (supplementary fig. S3B, Supplementary Material online). Because the Ceramiales is the largest group of red algae including greater than 2,655 species (Guiry MD and Guiry GM 2015), additional mt genome data from representative taxa of the order (such as an early branched Spyridia) may provide better resolution of this node.

Mt genome data included in this analysis resolved the relationship of 16 of the 25 florideophycean orders (Guiry MD and Guiry GM 2015). It has generally been difficult to resolve class-level relationships among red algae (i.e., Compsopogonophyceae, Porphyridiophyceae, and Stylonematophyceae) and intraclass relationships (i.e.,

Nemaliophycidae and Rhodymeniophycidae) using multigene data (Saunders and Hommersand 2004; Yoon et al. 2006; Le Gall and Saunders 2007; Verbruggen et al. 2010; Scott et al. 2011). However, our analysis using mtDNA data (data set c24p and 23 sub-data sets) provides moderate to strong support for intraclass relationships in the Rhodymeniophycidae. Based on these results, we expect that mt genome data from major representatives could be a promising tool to further resolve the relationships of the red algae.

Mitochondrial Genome Evolution in Red Algae

To infer mt genome evolution in the red algae, we compared gene synteny from 35 genomes (excluding the highly divergent extremophiles Cyanidioschyzon merolae and Galdieria sulphuraria) that represent two classes (Bangiophyceae and Florideophyceae), all 5 of the florideophycean subclasses, and all 12 rhodymeniophycidan orders (fig. 2). These data (supplementary fig. S1, Supplementary Material online) encompass the phylogenetic breadth of the Florideophyceae. The overall structure of mt genomes was mapped onto the best phylogeny (fig. 1) to infer the ancestral genome content, patterns of gene acquisition and loss, and types of rearrangements (i.e., numbers on fig. 2). Because of a conserved linear arrangement of three adjacent genes around mt rRNAs (rrs [small subunit rRNA]-nad4L-rrl, approximately 4,300 bp in length excluding intron regions), we used it as 5'- and 3'anchor points of the mtDNA maps (NR segment in fig. 2). We infer that a total of 51 mitochondrial genes were present in the hypothetical last common ancestor of Bangiophyceae and Florideophyceae (Ra in fig. 2) including 24 protein-coding genes, 2 rRNAs, and 25 tRNAs (supplementary table S4, Supplementary Material online).

gene content of Bangiophyceae The and Florideophyceae is nearly identical in terms of proteincoding genes (supplementary table S3, Supplementary Material online). However, two major syntenic differences involving inversion/translocation events were found between Bangiophyceae (e.g., Porphyra) and Hildenbrandiophycidae (Hildenbrandia) (see no. 1-12 in fig. 2) and between Hildenbrandiophycidae and Nemaliophycidae (Palmaria) (see no. 14, 15, 17–19). The majority of rearrangements occurred through gene inversions (12 inversions of 17 rearrangements, fig. 2) between Bangiophyceae and Hildenbrandiophycidae, and between Hildenbrandiophycidae and Nemaliophycidae. Rearrangements of a single gene were more frequent than simultaneous multigene rearrangements (i.e., more than two genes involved; i.e., genes no. 5, 7, 10, and 14).

The *trnl* gene (without intron; no. 6) exists between two large conserved regions, that is, *nad2-sdh4-nad4-nad5-atp8-atp6* (ANS segment; approximately 6,500 bp) and *ymf39-cox3-cox2-cox1* (no. 7 CY segment; approximately 4,000 bp

GBE

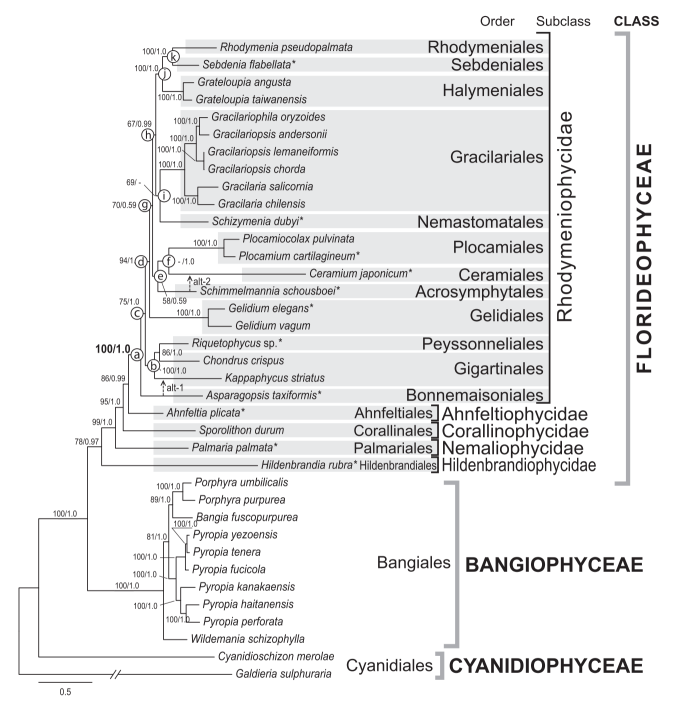
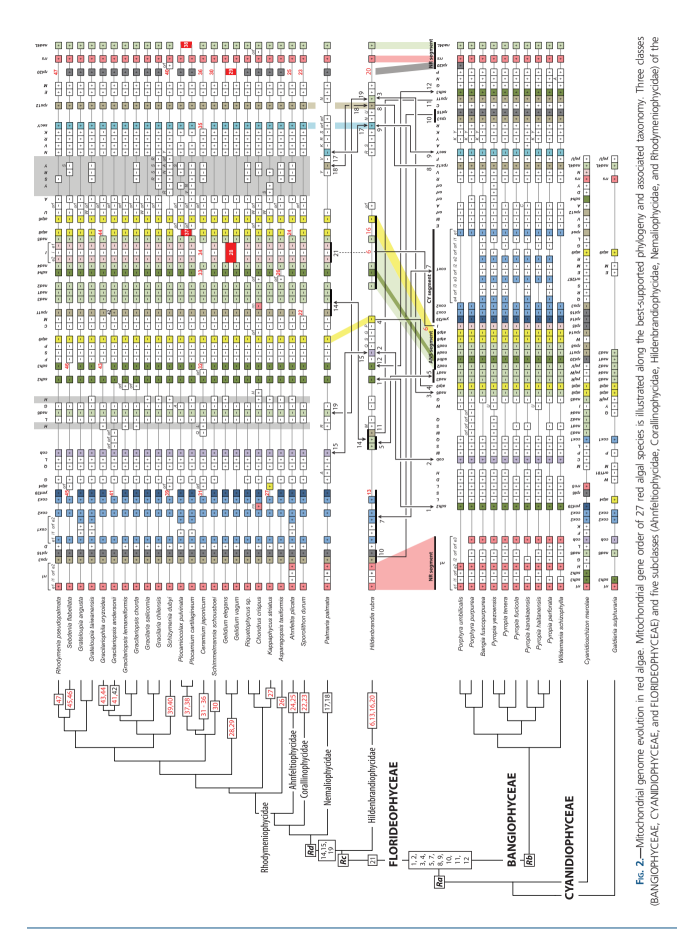


Fig. 1.—Phylogeny of the red algae (Rhodophyta) based on mt genome data. Tree constructed using the ML method based on 24 concatenated genes (6,345 amino acids from 24 protein coding). The support values for each node are calculated from MLB and BPP. Asterisks after species names indicate newly determined mt genomes, followed by red algal order, subclass, and class system. Alternative tree branch position of species (alt) indicated by dot line with arrow, that is, alt-1 for *Asparagopsis* and alt-2 for *Schimmelmannia*. Alternative topologies are available in supplementary figure S3, Supplementary Material online.

in length when the intron and intronic ORF are excluded) in the bangiophycean lineage. In the Florideophyceae, the CY segment is inverted (5'-cox1-cox2-cox3-ymf39-3'), whereas the ANS segment is retained with the *trnl-intron* in all species (no. 21) but not in *Hildenbrandia* (no. 6; 5 bp intergenic region between *nad4* and *nad5*), *Gelidium* (no. 28), and *Ceramium* (no. 34). This indicates that the *trnl* gene was present in the common ancestor of Bangiophyceae and Florideophyceae



(Ra in fig. 2, supplementary table S4, Supplementary Material online). In the Bangiophyceae, the ANS and CY segments were present in all bangiophycean species with/without intron and intronic-ORF in cox1. However, in the Florideophyceae, the ANS segments acquired an intron in trnl and a reduced intron and intronic-ORF in cox1 of the CY segment. The trnl has gained an intron and it is inserted in the ANS segment (i.e., between nad4 and nad5) in the florideophycean ancestor (Rc). Thus, the trnl-intron was lost independently in Hildenbrandia, Gelidium, and Ceramium. This trnl-mediated intron acquisition is the first report of such an event in red algae; however, acquisition of heterogeneous DNA (protein-coding genes and introns) has been reported in the Viridiplantae, that is, in green algae and land plants (Palmer et al. 2000; Bergthorsson et al. 2004; Smith et al. 2013).

Since the split of the hildenbrandiophycidan ancestor (i.e., node Rd in fig. 2), gene synteny has been highly conserved, with changes only occurring in tRNAs and in intronic regions. Lineage-specific CDS gene losses were found in florideophycean species (red numbers in fig. 2). For example, there were losses of atp8 (no. 16), rpl20 (no. 20), trnl (no. 6), and ymf39 (no. 13) in Hildenbrandia, whereas losses of rps11 (no. 22) and rpl20 (no. 23) occurred in Sporolithon. In contrast to the conserved CDS gene content (except the rps11 inversion in Gracilariopsis andersonii; no. 42), there are several types of variation in tRNAs (i.e., gene duplication and insertion/deletion, see fig. 2 and supplementary table S3, Supplementary Material online) not only between the subclasses but also within the Rhodymeniophycidae in trnH (H—H gray regions around trnL-nad6-trnG) and/or trnY and one of each trnR and trnS (Y-S-R-Y gray region between trnA and trnN).

In this study, the new florideophycean mt genome data surprisingly show that overall gene synteny is conserved among subclasses, except for the Hildenbrandiophycidae. Based on the fossil record and molecular clock studies, the florideophycean ancestor (*Rc*) originated approximately 900 Ma and the nemaliophycidan ancestor (*Rd*) appeared at least 550 Ma (Corallinophycidae fossils) (Butterfield 2001;

Xiao et al. 2004; Yoon et al. 2004). Despite the long evolutionary history of the florideophycean red algae, most are characterized by a highly conserved mtDNA synteny (fig. 2).

Reduction of Protein-Coding Genes

Based on our current collection of mt genomes, at least 51 genes (excluding hypothetical proteins) were present in the common ancestor (Ra) of Bangiophyceae and Florideophyceae (supplementary table S4, Supplementary Material online). We inferred a total of 29 lineage-specific gene losses during red algal evolution (no. 6, 13, 16, 20, 22-41, 43-47 in fig. 2). We marked these gene losses on the map of percent AT-content versus gene length, which shows a negative correlation between these values (fig. 3). Surprisingly, genes that were absent in some species were grouped in the top-left corner. These genes are small in size (<800 bp in mean length) and have a high AT content (>73% mean AT content). The rpl20 (relatively short with the highest AT content) was lost in 18 species among 35 investigated red algae (except Cyanidiophyceae), and this was followed by gene losses of ymf39 (6 species), atp8 (5 species), sdh3 (3 species), sdh4 and nad4L (2 species), and rps11 and secY (1 species). Gene loss appears however to be uncorrelated with phylogenetic relationships (see, red number on fig. 2). For example, within the strongly supported clade of the Halymeniales, Sebdeniales, and Rhodymeniales (100% MLB), rpl20 was lost only in R. pseudopalmata (no. 47), whereas it was retained in only three species. Interestingly, 9/10 bangiophycean species lost the rpl20 gene, except for Porphyra umbilicalis. It is reasonable to infer the rpl20 data as representing four independent losses in Porphyra purpurea, B. fuscopurpurea, the ancestor of six Pyropia species, and Wildemania schizophylla, respectively, as opposed to one loss in the ancestor of the Bangiophyceae (Rb) with a regain of rpl20 because it is extremely improbable to regain a gene in the correct reading frame after its loss.

There are three genes located near the boundary of the gene loss; that is, *nad6*, *rps3* and *rpl16*, which are small in size (612, 691, and 419 bp mean length, respectively) and have a

Fig. 2.—Continued

red algal taxonomic system are indicated on the branches. Gene synteny is anchored by *rrl* (mt-rRNA large subunit) at the start and end *rrs* (mt-rRNA small subunit) at the end followed by *nad4L*, which shared an arrangement in all species (NR segment). Protein-coding and rRNA genes are abbreviated as shown in supplementary figure S1, Supplementary Material online. The tRNA genes are labeled using the one-letter code of their corresponding amino acid. The box indicates only gene existence, and box size does not correspond to gene size. The color code for genes corresponds to supplementary figure S1 and table S1, Supplementary Material online. Plus (+, forward) and minus (-, reverse) refer to the direction of gene transcription. The one letter code on the tRNA gene box indicates when alternative tRNA genes exist. Light-gray vertical boxes (three positions) point to variable tRNA gene positions in the Ahnfeltiophycidae, Corallinophycidae, and Rhodymeniophycidae. Evolutionary changes in the position of protein-coding genes are indicated with numbers (1–46) and mapped on the phylogeny; black numbers with blunt lines between genomes indicate gene rearrangement (e.g., no. 4); gene rearrangements are marked with a thick horizontal line, that is, no. 5, 7, 10, and 14; and the *trnl-intron* region event (no. 21) is mapped on *Palmaria* and on the remaining phylogeny and corresponds to acquisition time. Strictly conserved regions are denoted with colored shadows between clades; that is, conserved ANS (*nad2-sdh4-nad4-nad5-atp8-atp6*), CY (*ymf39-cox3-cox2-cox1*), and NR (*rpl20-rrs-nad4L-rrl*) segments between Bangiophyceae and Florideophyceae, *atp9* between *Hildenbrandia* and *Palmaria*, and *rps12-secY* between *Hildenbrandia* and *Sporolithon*.

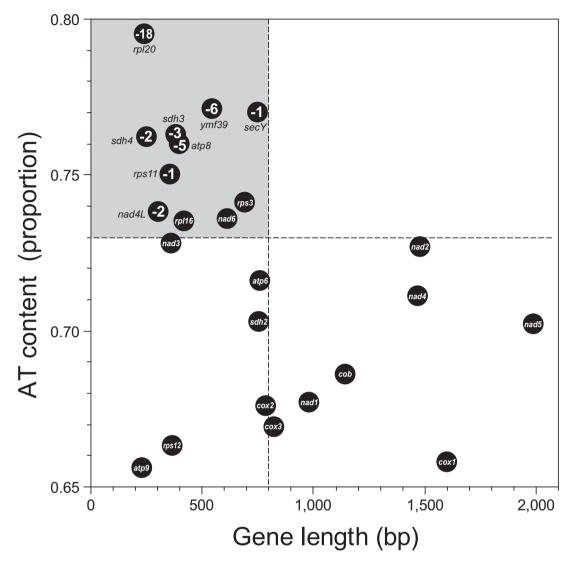


Fig. 3.—Protein-coding gene size reduction in red algal mitochondrial genomes. Correlation of AT contents (proportion) and sequence length (bp) of genes are illustrated with the minus number of species for each gene loss in bangiophycean and florideophycean red algae. Dark-gray circles with no number indicate that the gene is found in all taxa (35 species). The CDS of less than 800 bp in length and higher than 73% of AT content tend to be those lost.

high AT-content (73.6%, 74.1%, and 73.5%, respectively; fig. 3) that are retained in all 35 bangio-florideophycean red algal mt genomes. Based on the absence of the ribosomal and succinate dehydrogenease genes (rpl, rps, and sdh) in land plants (Adams et al. 2002; Notsu et al. 2002; Adams and Palmer 2003), and the trend of gene loss in red algae, rpl16 and rps3 may be candidates for future gene loss (or perhaps already lost in some yet to be studied rhodophytes) than nad6. Meanwhile, many lost genes (rpl20, rps11, sdh4, secY, and ymf39) and candidates for loss (rpl16, rps2, rps3, and sdh2) were lost in G. sulphuraria but retained in Cy. merolae (Cyanidiophyceae; fig. 2, supplementary table S3, Supplementary Material online). The pattern that high ATcontent genes are more frequently lost than low AT-content ones in red algae is an intriguing one, for which we currently have no explanation for the mechanisms. It may be a phenomenon restricted to this phylum, and further study is needed.

The lost genes have most likely been replaced by copies that were transferred to the nucleus (Lang et al. 1999). In red algae, *G. sulphuraria* is the only species with completed nuclear (Schönknecht et al. 2013) and mt genomes, and the mt genome has eight CDS losses (fig. 2, supplementary table S3, Supplementary Material online). We found one of the eight CDSs, mitochondrial-derived *sdh2* gene, in the nuclear genome of *G. sulphuraria* (XM_005707240.1|*G. sulphuraria* succinate dehydrogenase [ubiquinone] iron-sulfur protein [Gasu_20110] mRNA, complete CDS, 275 AA;

E value = 5e-105, 93% Query coverage, 59% identity) by using tBLASTn search with *sdh2* of *Cy. merolae* (gi|89543 61|ref|NP_059350.1; 265 AA) on the National Center for Biotechnology Information. However, other genes showed matches with corresponding plastid encoded genes (*rpl16*, *rps3*, *rps11*, *rps12*) or very short regions of similarity to nuclear genes (*rpl20*, *shd4*, *ymf39*). Additional genomes from Florideophyceae and mt-driven gene transfer to the nucleus are needed to test the genome reduction and gene transfer scenario presented here.

Among all reported mt genomes shown in figure 2, group Il introns are present in all species except Ceramium, Cyanidioschyzon, Galdieria, and Gelidium (Dai et al. 2003; Mao et al. 2012; Smith et al. 2012; Yang, Kim, Boo, et al. 2014). It is interesting, however, that only three genes (rrl, cox1, and trnl) contain introns with or without intronic-ORFs (Leblanc et al. 1995; Hancock et al. 2010; Hwang et al. 2013; Kim, Yang, Boo, et al. 2014; Yang, Kim, Boo, et al. 2014). In the bangiophycean species (Bangia, Porphyra, Pyropia, and Wildemania), one or two copies of rrl-introns are present (Smith et al. 2012; Hwang et al. 2013), whereas these tend to be lost in most florideophycean taxa (i.e., only present in Hildenbrandia, Palmaria, and Ahnfeltia). A similar pattern is found for the cox1 intron, where 1–3 copies are present in bangiophycean taxa but sporadically present in the Florideophyceae (supplementary table S3, Supplementary Material online). In contrast, the florideophycean lineage acquired the trnl-intron without the intronic-ORF after divergence of the Hildenbrandiophycidae.

The distribution of introns in red algae together with the 11 novel mt genomes suggests that the majority of red algal introns (*rrl* and *cox1*) originated prior to the split of the two major classes (*Ra* in fig. 2; the last common ancestor of Bangiophyceae and Florideophyceae). Subsequent intronic-ORF loss resulted in pseudogenization of intronic-ORFs, followed by intron loss. Intron gain, however, may occur with an ORF-lacking intron, as seen for the *trnl*-intron, which is an apparent case of a recent intron gain.

Conclusions

Our study elucidates mt genome evolution in the Florideophyceae through analysis of this genome in *Ahnfeltia*, *Hildenbrandia*, *Palmaria*, and diverse rhodymenio-phycidan species. The phylogeny based on concatenated amino acid mt genome data supports the current taxonomic system for Bangiophyceae and Florideophyceae. We demonstrate that all Florideophyceae mt genomes have compact and conserved architectures but they also contain several unique features, such as gene rearrangements that discriminate lineages and the gain of group II introns in *tml* genes. Comparative analyses suggest that the mtDNA in Florideophyceae (except the Hildenbrandiophycidae), although representing a wide variety of morphologically divergent taxa,

is surprisingly highly conserved with little variation in gene composition and with shared synteny across the class. This result is consistent with a rapid radiation of the class followed by strong constraints on mitochondrial gene content and genome evolution.

Supplementary Material

Supplementary figures S1–S3 and tables S1–S4 are available at *Genome Biology and Evolution* online (http://www.gbe. oxfordjournals.org/).

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