



Divergence time estimates and the evolution of major lineages in the florideophyte red algae

Citation

Yang, Eun Chan, Sung Min Boo, Debashish Bhattacharya, Gary W. Saunders, Andrew H. Knoll, Suzanne Fredericq, Louis Graf, and Hwan Su Yoon. 2016. "Divergence Time Estimates and the Evolution of Major Lineages in the Florideophyte Red Algae." Scientific Reports 6 (1) (February 19). doi:10.1038/srep21361.

Published Version

doi:10.1038/srep21361

Permanent link

http://nrs.harvard.edu/urn-3:HUL.InstRepos:33983356

Terms of Use

This article was downloaded from Harvard University's DASH repository, and is made available under the terms and conditions applicable to Open Access Policy Articles, as set forth at http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#OAP

Share Your Story

The Harvard community has made this article openly available. Please share how this access benefits you. <u>Submit a story</u>.

Accessibility

Divergence time estimates and the evolution of major lineages in the florideophyte red algae

Eun Chan Yang^{1,2}, Sung Min Boo³, Debashish Bhattacharya⁴, Gary W Saunders⁵, Andrew H Knoll⁶, Suzanne Fredericq⁷, Louis Graf⁸, & Hwan Su Yoon^{8,*}

¹Marine Ecosystem Research Division, Korea Institute of Ocean Science &

Technology, Ansan 15627, Korea

²Department of Marine Biology, Korea University of Science and Technology,

Daejeon 34113, Korea

³Department of Biology, Chungnam National University, Daejeon 305-764, Korea

⁴Department of Ecology, Evolution and Natural Resources, Rutgers University, New Brunswick, NJ 08901, USA

⁵Department of Biology, University of New Brunswick, Fredericton, NB E3B 5A3 Canada

⁶Department of Organismic and Evolutionary Biology, Harvard University,

Cambridge, MA 02138, USA

⁷Department of Biology, University of Louisiana at Lafayette, Lafayette, LA 70504-3602, USA

⁸Department of Biological Sciences, Sungkyunkwan University, Suwon 16419, Korea

*Corresponding author: Hwan Su Yoon (email: <u>hsyoon2011@skku.edu</u>)

The Florideophyceae is the most abundant and taxonomically diverse class of red algae (Rhodophyta). However, many aspects of the systematics and divergence times of the group remain unresolved. Using a seven-gene concatenated dataset (nuclear EF2, LSU and SSU rRNAs, mitochondrial *cox*1, and plastid *rbc*L, *psa*A and *psb*A genes), we generated a robust phylogeny of red algae to provide an evolutionary timeline for florideophyte diversification. Our relaxed molecular clock analysis suggests that the Florideophyceae diverged approximately 943 (817-1,049) million years ago (Ma). The major divergences in this class involved the emergence of Hildenbrandiophycidae [ca. 781 (681-879) Ma], Nemaliophycidae [ca. 661 (597-736) Ma], Corallinophycidae [ca. 579 (543-617) Ma], and the split of Ahnfeltiophycidae and Rhodymeniophycidae [ca. 508 (442-580) Ma]. Within these clades, extant diversity reflects largely Phanerozoic diversification. Divergences within Florideophyceae were accompanied by evolutionary changes in the carposporophyte stage, leading to a successful strategy for maximizing spore production from each fertilization event. Our research provides robust estimates for the divergence times of major lineages within the Florideophyceae. This timeline was used to interpret the emergence of key morphological innovations that characterize these multicellular red algae.

Introduction

The Florideophyceae is the most taxon-rich red algal class, comprising 95% (6,752) of currently described species of Rhodophyta¹ and possibly containing many more cryptic taxa². Florideophytes are relatively well known to the public because they are common along many shorelines and provide economically important cell wall-derived compounds such as agar and carrageenan. The Florideophyceae is equally well known to biologists, in part because of its characteristic (mostly): i) triphasic life cycle consisting of a carposporophyte, gametophyte and tetrasporophyte, ii) the presence of pit-plugs between adjacent cells, and iii) postfertilization cell-cell fusion mechanisms³. Carposporophytes may produce thousands of diploid carpospores from a single syngamy taking advantage of cell-to-cell fusions that occur between the fertilized carpogonium and auxiliary cells. Each carpospore develops into a tetrasporophyte that produces four haploid tetraspores per diploid cell through meiotic division. These additional carposporophyte and tetrasporophyte phases are thought to be evolutionary innovations that explain the success of the red algae. There are, however, a few exceptions to a triphasic life cycle including the 'asexual tetrasporophytes' of the Hildenbrandiales, the derived diphasic life cycle in Palmariales, and abbreviated life cycles (i.e., lacking a free-living tetrasporophyte) in some Acrochaetiales⁴, Nemaliales, and Gigartinales⁵.

Pit connections linking neighboring cells are one of the diagnostic features characterizing red algal orders. Diverse combinations of pit connection compartments (i.e., plug core with different number of cap layers and membranes) with molecular data have been used to define the ordinal boundaries of the Florideophyceae. In addition, comparison of the developmental pathway of reproductive structures in male and female gametophytes and tetrasporophytes have also been used to study the diversifictaion of red algal genera and families. Recent phylogenetic studies based on molecular data have resulted in a revised classification system that recognizes 29 orders in five subclasses: Ahnfeltiophycidae, Corallinophycidae,

Hildenbrandiophycidae, Nemaliophycidae, and Rhodymeniophycidae^{1,6-8}. Although previous molecular studies have produced robust phylogenies for relationships among some Florideophyceae⁶, deep relationships among and within other subclasses remain poorly resolved⁹, and the evolutionary timeline for florideophyte divergence has rarely been studied^{8,10}.

Two important sets of Proterozoic fossils constrain the early evolutionary history of the red algae. The first is commonly considered to be the oldest known taxonomically resolved eukaryotic fossil, the ca. 1,250-1,100 million years ago (Ma) *Bangiomorpha pubescens* from the Hunting Formation, Arctic Canada¹¹ (see review of *Bangiomorpha pubescens* age constraints¹²). The second consists of anatomically preserved florideophyte fossils from the ca. 580 (635-551) Ma, Doushantuo Formation, southern China, that exhibit growth forms and features closely resembling reproductive structures of modern corallines¹³⁻¹⁵. Based on Doushantuo fossils¹³, the split between the Bangiophyceae and Florideophyceae must have occurred during the Neoproterozoic Era, or earlier. Using these two fossils as calibration points in a multigene phylogenetic analysis. Yoon et al.¹⁰ suggested that the first red alga originated approximately 1,500 Ma, and the Florideophyceae evolved approximately 800 Ma. In another study considering red algal fossil data, Saunders and Hommersand⁶ summarized available data that were consistent with the major lineages of florideophytes diverging before 600–550 Ma, at the end of the Proterozoic Eon. Both studies, however, relied on a limited sampling of florideophyte taxa, and divergence times remain uncertain for the major lineages within the class. Aguirre et

al.¹⁶ studied coralline red algal phylogeny and divergence times based on fossil records; well-preserved coralline skeletons in Mesozoic and Cenozoic sedimentary rocks include species placed within the Sporolithaceae (136–130 Ma), Hapalidiaceae (115–112 Ma), and Lithophylloideae (65.5–61.7 Ma), providing additional calibration points for molecular clock analysis^{16,17}.

Estimating divergence times using molecular data and fossil constraints can considerably advance the evolutionary study of florideophytes and algae in general¹⁰. To estimate divergence times associated with the Florideophyceae and its constituent subclasses, we performed phylogenetic and molecular clock analyses of combined data (nucleotide sequence of rRNA + amino acid sequence of coding DNA sequence, CDS) from three nuclear (EF2, LSU, and SSU), one mitochondrial (*cox*1), and three plastid (*psa*A, *psb*A, and *rbc*L) genes from 27 florideophycean orders (missing only 2 orders: Entwisleiales and Pihiellales). We generated 180 new sequences consisting primarily of three plastid genes and subsequently compiled a dataset with previously published sequences (mainly EF2, LSU, SSU, and *cox*1) from GenBank.

In order to estimate the divergence time of the Florideophyceae from other red algae, we used seven constraints (Fig. 1, see the methods for details): three red algal fossil dates, (a) 1,222-1,174 Ma for stem taxon; i.e. the filamentous and spore-bearing red alga *Bangiomorpha*^{11,12}, (b) 633-551 Ma for Doushantuo fossil-coralline¹³⁻¹⁵ algae, and (c) Cenozoic corallines^{16,17} (c1-c2). Four published divergence dates for land plants¹⁸ were also used as constraints, including (d) 471-480 Ma for the divergence between liverworts (*Marchantia*) and vascular plants , (e) 410-422 Ma for the divergence time between ferns (*Psilotum*) and seed plants , (f) 313-351 Ma for the divergence time between gymnosperms (*Pinus*) and angiosperms, and (g) 138-162 Ma for the monocot-eudicot split (*Zea* and *Arabidopsis*). Divergence times were

estimated using Bayesian relaxed-clock methods^{19,20}. The results are discussed in light of key morphological transitions, such as the origin of the triphasic life cycle and of a diversity of fertilization and diploidization modes in the Florideophyceae.

Results and Discussion

Phylogeny of the Florideophyceae. The maximum likelihood (ML) phylogeny inferred from the seven-gene concatenated dataset is shown in Fig. 1 (see also Supplementary Fig. S1). The ML topology was congruent with the Bayesian tree. The phylogeny resolved a monophyletic lineage including the Florideophyceae and Bangiophyceae (node '1' in Fig. 1 and Supplementary Fig. S1) with 60% ML bootstrap (MLB) and 1.0 Bayesian posterior probability (BPP) supports. Within the Florideophyceae, five strongly supported (100% MLB and 1.0 BPP in Supplementary Fig. S1) groups were recovered, equivalent to the five subclasses of Florideophyceae: Ahnfeltiophycidae, Corallinophycidae, Hildenbrandiophycidae, Nemaliophycidae, and Rhodymeniophycidae. These five lineages were recognized in previous studies on the basis of ultrastructural attributes and multigene phylogenetic analyses^{7,9,21}. Therefore, it is highly likely that they reflect accurately the subgroups of Florideophyceae.

The subclass Hildenbrandiophycidae diverged first within the Florideophyceae (node '2' in Fig. 1; 100% MLB and 1.0 BPP in Supplementary Fig. S1) with its deep position consistent with previous studies based on ultrastructural and molecular data^{7,9,22-24}. The sole order in this subclass is the Hildenbrandiales, characterized by pit plugs with a single cap layer covered by a membrane²⁵. In contrast, the earlier diverging Bangiales (Bangiophyceae) have pit plugs with a cap but no membrane, and the pit plugs of Compsopogonophyceae represent the ancestral

type consisting simply of a plug core, but lacking both a cap and membrane²⁶. The Hildenbrandiales includes two genera, *Hildenbrandia* and *Apophlaea*, which form crustose thalli or extensive crusts with upright portions. This order is characterized by the vegetative phase having abundant secondary pit connections that link neighbouring cells. This is atypical in being established without conjunctor cell formation²⁷. Zonately and irregularly dividing tetrasporangia have been reported²⁴. but it has not been established whether they form meiotically or mitotically 27 . There are no reports of recognizable gametophytic reproductive structures (e.g., eggs, referred to as carpogonia, or spermatangia) or a sexual life history for hildenbrandialean species. Based on the current phylogenetic reconstructions (Supplementary Fig. S1) it is equally likely that the ancestor of Hildenbrandiophycidae was either of the biphasic type that characterizes other earlier diverging clades of red algae (e.g., Bangiophyceae and Rhodellophyceae) or the triphasic type characteristic of the remaining florideophyte subclasses. The only certainty is that the triphasic pattern evolved somewhere between the divergence of the Bangiophyceae and Florideophyceae (node '1' in Fig. 1), and the divergence of the Nemaliophycidae and Ahnfeltiophycidae-Corallinophycidae-Rhodymeniophycidae clade (node '3' in Fig. 1).

The monophyly of the Nemaliophycidae is strongly supported (node '4' in Fig. S1; 100% MLB and 1.0 BPP in Supplementary Fig. S1), consistent with previous studies^{7,9,21,28,29}. The Nemaliophycidae is characterized by the presence of pit plugs with two cap layers²⁵, which likely evolved from pit plugs with a single cap layer (e.g., Bangiales and Hildenbrandiales)^{25,26,29}. Interordinal relationships within the Nemaliophycidae, however, were not resolved. The Batrachospermales and Thoreales were positioned deep in the lineage with moderate support (68% MLB and 1.0 BPP in

Supplementary Fig. S1); in contrast, phylogenetic relationships among the Acrochaetiales, Balbianiales, Balliales, Colaconematales, Nemaliales, Palmariales, and Rhodachlyales were weakly supported. We were unable to add representatives of the Entwisleiales to this study, therefore broader taxon sampling with additional sequence data may resolve these relationships.

The Corallinophycidae was positioned between the Nemaliophycidae and Ahnfeltiophycidae-Rhodymeniophycidae clade. This subclass is characterized by pit plugs with a domed outer cap layer and thalli that are mineralized due to calcite precipitation^{6,21,30}. The distinctiveness of the Corallinophycidae is consistent with previous molecular and morphological studies^{7,9,29,31,32}. This group includes four orders: the Corallinales, Hapalidiales, Rhodogorgonales, and Sporolithales. The Rhodogorgonales is positioned deepest in this lineage (100% MLB and 1.0 BPP in Supplementary Fig. S1) followed by the divergence between the distinctive Sporolithales, Corallinales and Hapalidiales (node 'C1, and C2, respectively' in Supplementary Fig. S1; 100% MLB and 1.0 BPP). Relationships among orders and families within the subclass were congruent with previous studies^{16,17,31,32}.

The Ahnfeltiophycidae is sister to the Rhodymeniophycidae at node 5 (Fig. 1; 100% MLB and 1.0 BPP in Supplementary Fig. S1). The sister relationship of these two subclasses was strongly supported in our results and is congruent with previous studies based on ultrastructure and multigene phylogenies^{7,9}. The Ahnfeltiophycidae have naked pit plugs, lacking caps and membranes³³, and include the Ahnfeltiales and Pihiellales³⁴. The latter was not included in this study.

The Rhodymeniophycidae is strongly supported as a monophyletic group with 100% MLB and 1.0 BPP (node '6' in Fig. 1), including 12 orders (out of 13 extant orders²) comprising the Acrosymphytales, Bonnemaisoniales, Ceramiales, Gelidiales,

Gigartinales, Gracilariales, Halymeniales, Nemastomatales, Peyssonneliales, Plocamiales, Rhodymeniales, and Sebdeniales. All have pit plugs covered by a membrane only; the Gelidiales is an exception in having a thin inner cap under the membrane^{21,25}. In the Gelidiales, and to a lesser extent the Gracilariales²¹, pit plugs have a striated plug core for which the taxonomic utility has yet to be established. Interordinal relationships of this diverse subclass were not fully resolved in the current study. Nine orders, including Acrosymphytales, Ceramiales, Gelidiales, Gracilariales, Halymeniales, Nemastomatales, Plocamiales, Rhodymeniales, and Sebdeniales form a clade (92% MLB and 0.87 BPP in Supplementary Fig. S1), whereas relationships among the more deeply diverging Bonnemaisoniales, Gigartinales, and Peyssonneliales were largely unresolved (Supplementary Fig. S1). This conflicts with previous multigene phylogenetic analyses that weakly resolved the Ceramiales as an early-diverging group within the Rhodymeniophycidae⁷. However, interrelationships among the orders remain largely unresolved, in particular those subsequent to the divergence of the Peyssonneliales, except for the monophyletic assemblage of the three orders Halymeniales, Rhodymeniales, and Sebdeniales (99% MLB and 0.99 BPP in Supplementary Fig. S1). As indicated in Verbruggen et al.⁹. this is one region of the red algal phylogenetic tree that is in need of considerably more investigation.

In Rhodymeniophycidae, all orders were strongly supported as monophyletic groups (98-100% MLB and 1.0 BPP in Supplementary Fig. S1) with the exception of the Gigartinales (61% MLB and 1.0 BPP in Supplementary Fig. S1) and the Plocamiales (52% MLB and 1.0 BPP in Supplementary Fig. S1), which is consistent with published data^{7,9,35}. Once again, relationships in this part of the tree need additional study including improved taxon sampling (e.g., *Hummbrella hydra*, the

lone member of the Pseudoanemoniaceae), more sequence data and further exploration of analyses options such as data partitions and evolutionary model selections (see Le Gall et al.⁷ and Verbruggen et al.⁹). The Ceramiales, the largest florideophycean order, was recovered as monophyletic (98% MLB and 1.0 BPP in Supplementary Fig. S1), and resolved as sister to the Acrosymphytales ('node 7' in Fig. 1; 82% MLB, 1.0 BPP in Supplementary Fig. S1). Previous studies^{9,36}, however, have indicated that the Inkyuleeaceae may not join the remainder of the Ceramiales as a monophyletic group, which when resolved may require further taxonomic revision.

Divergence time estimation, fossils, and the evolution of florideophyte algae.

Divergence time estimation using fossil constraints usually entails a large degree of uncertainty. This is because taxonomic assignment and age determination may be uncertain, and for poorly sampled lineages, the oldest recognized fossils may singificantly postdate the the origin of the group^{37,38}. For these reasons, we tested the impact of calibration constraints on the estimated divergence times of red algae using the parametric prior distributions available in BEAST. We compared the posterior mean estimates of nodes between the uniform (*uni*) and normal (*nor*) prior distributions. Regression analysis suggests that use of *uni* results in markedly older divergence time estimates than under *nor* (Supplementary Table S1). The correlation coefficient (slope *b*) of estimates (Y) to C7 *nor* (X) showed that *uni* ages (e.g., *b*=1.4936 for C7 *uni*) were significantly older than *nor* ages (*b*=0.9794 for C7 *nor*). For instance, the largest differences were found in the node 'r' of C7 *uni* age (2,816 Ma 95% High Posterior Density [HPD]: 1,415-5,663 Ma) to C7 *nor* age (1,694 Ma, 95% HPD: 1,484-1,925 Ma), i.e., Δ node 'r' mean = 1,122 Ma. The age of node '1' of C7 *uni* (1,661 Ma, 95% HPD: 883-3,443 Ma) was 1.8-fold older than that of C7 *nor*

(943 Ma, 95% HPD: 817-1,049 Ma), i.e., \triangle node '1' mean = 718 Ma. In the C7 *uni* age, 1,661 Ma for node '1' (the first appearance of the Florideophyceae) was much older than the taxonomically defined oldest eukaryote fossil¹² (*Bangiomorpha*, node 'a'). Therefore, we used *nor* estimates for the inferences described below.

Under the nor approach, age estimates that removed one constraint 'b, d-g' at each time (C6-b nor, C6-d nor, C6-e nor, C6-f nor, and C6-g nor) and using only the outgroup (C5-*bc nor*), showed a negligible effect (b > 0.96). However, removal of constraint 'a' (C6-a nor, b=0.8491) and using constraint 'b' only (C1b nor, b=0.8566) led to an underestimation of the divergence time. For example, the age of node 'r' in C6-a nor (1,362 Ma, ca. 20% younger) and C1b nor (1,609 Ma, 5% younger) were younger than that of C7 nor. The analyses with red algal fossils only (C2ab nor) and the *Bangiomorpha* fossil (C1*a nor*) showed little difference for the age of node 'r', i.e., \triangle node 'r' mean to C7 nor were 4 Ma (0.2%) for C2ab nor and 20 Ma (1.2%) for Cla nor. These results indicate that constraint 'a' (early stem group of red algae, *Bangiomorpha*) is more critical for divergence time estimates regarding red algae than the coralline fossils deposited in the Doushantuo formation. Removal of constraint 'c' (C6-c nor) led to a little overestimation (b=1.1162). For example, the age of 'node r' was overestimated in C6-c nor age (2,287 Ma). Regardless of distribution priors, the mean age of all nodes using only secondary calibration (C1c *nor* and C1*c uni*) derived from a previous study¹⁶ resulted in drastcally younger ages for all nodes (Supplementary Table S1). In general, the divergence time estimates and 95% credibility intervals varied among constraint scenarios under both uniform (uni) and normal (nor) prior distributions. However, florideophycean time estimates from independent analyses overlapped with each other within the 95% HPD.

To to test the robustness of the time estimates, we compared results from different speciation tree priors including i) Yule, ii) Birth-Death and iii) Birth-Death Incomplete (see Supplementary Table S2). The date estimations, however, were largely congruent with each other (b=1.0175 with r²=0.9986 for the Birth-Death and b=1.0163 with r²=0.9976 for the Birth-Death Incomplete priors to the Yule prior), suggesting highly robust results. For example, the ages of node '1' were 943 Ma, 917 Ma, and 949 Ma with Yule, Birth-Death and Birth-Death Incomplete tree priors, respectively. Based on these comparisons, the normal calibration priors with the Yule speciation process (C7 *nor*) were chosen for the split time estimates for the major florideophycean lineages (Fig. 1).

The divergence time of the Florideophyceae from a common ancestor with the Bangiophyceae was calculated as 943 Ma (95% HPD: 817-1,049 Ma) for node '1' (Fig. 1), a late Mesoproterozoic to early Neoproterozoic³⁹ estimate that is consistent with previous results. Lim et al.⁴⁰ suggested an origin of red algae in early eukaryotic evolution at 1,300-1,400 Ma, followed by a florideophycean (7 genera) split from *Porphyra* (Bangiophyceae) without any detailed phylogenetic analysis. Yoon et al.¹⁰ estimated the divergence time of florideophytes (ca. 800 Ma) by using the relaxed molecular clock analysis. Although they used multigene data (16S rRNA, *psaA*, *psaB*, *psbA*, *rbcL*, and *tufA*) as well as multiple fossil constraints including two red algal fossils, only two florideophycean species (*Chondrus* and *Palmaria*) were included in the analysis. Here, we included representatives of almost all florideophycean orders (27 out of 29 orders) with seven fossil constraint data and a seven-gene dataset for the divergence time estimation.

The oldest convincing geological evidence for red algae comes from the erect filamentous microfossils of *Bangiomorpha* preserved by early diagenetic silicification

in tidal flat/lagoonal carbonates from Arctic Canada¹¹. Radiometric dates on volcanic rock constrain the age of these fossils to be younger than 1.267 ± 2 Ma and older than 723 ± 3 Ma, but an U-Th-Pb whole rock age of $1,092 \pm 59$ Ma for black shale that underlies the fossiliferous horizon⁴¹, an unpublished Pb-Pb date on correlative carbonates and arguments from sequence stratigraphy¹⁵ suggest that the true age lies closer to the lower radiometric boundary. Because the gross morphology of Bangiomorpha is similar to that of extant Bangia species, it might be recognized as a taxon in crown Bangiophyceae. If correct, this suggests that the divergence of the Bangiophyceae and Florideophyceae occurred prior to 1,200-1100 Ma, ca. 200-300 Ma earlier than our molecular clock estimate. However, several *Bangiomorpha*-like, simple filamentous species occur among the deeply diverging Compsopogonophyceae (i.e., Compsopogon, Compopogoniopsis, Erythrotrichia, Rhodochaete) and Stylonematophyceae (i.e., Bangiopsis, Purpureofilum, Stylonema) (see Supplementary Fig. S1). In fact the original description of *Bangiomorpha* noted that the multicellular holdfast had greater similarity to *Erythrotrichia* than *Bangia*¹¹. The diagnostic packet-formation during sexual reproduction in the Bangiales, and as reasonably posited for *Bangiomorpha*, has subsequently been reported in three species of the Compsopogonophyceae indicating that this feature has evolved at least twice within the red algae. It is thus possible that *Bangiomorpha* associates with any one of a number of the deep red algal lineages, possibly even an extinct lineage that evolved characters in parallel to the Bangiophyceae and Compsopogonophyceae. Therefore, it would not be unreasonable to place the *Bangiomorpha* constraint as a stem taxon to the early branching lineages of red algae (node 'a' in Fig. 1). Our estimate for the time of the initial red algal divergence casts doubt on the

interpretation of budding coccoidal microfossils from the ca. 1,850 Ma Gunflint Formation, Canada, as red algae⁴².

The major divergences within the Florideophyceae (nodes '2–5' in Fig. 1) occurred during the mid-Neoproterozoic to early-Paleozoic eras beginning with the Hildenbrandiophycidae, with an estimated divergence time of 781 (95% HPD: 681-879) Ma (node '2' in Fig. 1). During this time, somewhere between 'nodes 1 and 3' the so-called triphasic (gonimoblast development on the female gametophyte) life cycle evolved in red algae. Did this happen in the ancestor of all Florideophyceae (between nodes '1 and 2') with subsequent loss in the Hildenbrandiophycideae, or was this subclass ancestrally biphasic with gonimoblast development evolving between nodes '2 and 3'? Only through elucidation of the sexual pattern for the Hildenbrandiophycidae will this question be resolved. Whenever it originated, the triphasic pattern is characterized by free-living haploid male and female gametophytes, which produce the gametes. The second phase, a sporophyte called the carposporophyte, involves postfertilization development of diploid gonimoblast filaments on the female gametophyte and produces carpospores. The third phase is a second diploid sporophyte, termed a tetrasporophyte in Florideophyceae because meiosis typically results in four-spored meiosporangia (tetrasporangia) with each haploid tetraspore germinating into a gametophyte. Under the assumption that a successful life history tends to maximize the potential for genetic recombination and genetic diversity from the union of a single pair of gametes. Searles⁴³ concluded that selection has favored the evolution of a gonimoblast stage in red algae as compensation for the presumed inefficient fertilization attributed to the absence of motile gametes; however, more recent research has shown that fertilization may not be as inefficient as previously thought (e.g. see Maggs et al.⁴⁴).

The oldest known florideophyte fossils occur in Ediacaran rocks from southern China¹³. Half a dozen taxa of thalloid algae preserved in three-dimensional cellular detail by early diagenetic phosphate precipitation reveal features that ally them to florideophyte algae. Whereas several of the preserved populations have been interpreted as early branching florideophytes – perhaps stem groups or early crown representatives of extant taxa – the presence of filamentous "cell fountains," cortexmedulla differentiation, conceptacles, possible cell fusions, carposporangia, and tetrasporangia (cruciate and stalked) suggest that stem corallines are present in this assemblage as well¹³. Radiometric dates¹⁵ constrain these fossils to be younger than 632.5 ± 0.5 Ma and older than 551.1 ± 0.7 Ma, and a sequence boundary beneath the fossils might correlate with 580 Ma glaciations in the northern hemisphere. These age constraints are within the 95% credibility intervals of our molecular clock estimates (i.e., estimate without constraint 'b') for initial florideophyte diversification. Independent evidence of Ediacaran red algae comes from organic matter containing high abundances of C^{27} steranes⁴⁵ – red algae are unusual among algae for the predominance of C^{27} molecules in their sterol profiles⁴⁶.

The split of the Nemaliophycidae (node '3' in Fig. 1) from the Ahnfeltiophycidae-Corallinophycidae-Rhodymeniophycidae (ACR lineage) occurred 661 (95% HPD: 597-736) Ma. The Corallinophycidae diverged at 579 (95% HPD: 543-617) Ma followed by the split of Ahnfeltiophycidae and Rhodymeniophycidae about 508 (95% HPD: 442-580) Ma (node '5'). Diversification within Nemaliophycidae began at 331 (95% HPD: 202-458) Ma (node '4'); Nemaliophycidae have cruciate tetrasporangia and mono- and bi-sporangia. Carposporophytes produce spore-bearing gonimoblasts directly from the fertilized carpogonium in this group, and the fertilization nucleus is thus not transferred to separate generative auxiliary cells for production of the gonimoblasts. Cell-to-cell fusions in nemaliophycidaen gonimoblast development are restricted to cells of the carpogonial branch resulting in a secondarily formed carpogonial fusion cell. In this case, as gonimoblast development proceeds, the cytoplasm of the carpogonial branch cells extends and fuses around the cells' pit plugs with the result that the pit plugs become dislodged and the carpogonial branch cells form open connections with one another, forming a fusion cell. These cell-to-cell fusions do not involve postfertilization connecting cells or connecting filaments. In the Ahnfeltiales (Ahnfeltiophycideae)³³, as well as the Gelidiales⁴⁷ and Gracilariales⁴⁸ (Rhodymeniophycidae), generative auxiliary cells are not present and the fertilization nucleus remains in the carpogonium, which facultatively fuses with neighbouring vegetative cells following fertilization resulting in another type of carpogonial fusion cell that cuts off gonimoblast initials. In the remainder of the Rhodymeniophycidae, fusion cells may incorporate a generative auxiliary cell, supporting cell, adjacent sterile filaments, and nutritive vegetative gametophyte cells acting as auxiliary cells⁴⁹. In the Rhodogorgonales, auxiliary cells and connecting filaments are absent, and gonimoblast filaments cut off directly from the fertilized carpogonia elongate, with portions of the intercalary gonimoblast cells expanding in size and consecutively initiating secondary gonimoblasts at the point of fusion with terminal cells of specialized vegetative filaments⁵⁰; it is likely that such independent, unique cell-cell fusion mechanisms not involving the production of auxiliary cells was ancestral in corallinophycidaen diversification (asterisk in Fig. 1).

Various types of pre- and post-fertilizational cell-to-cell fusion mechanisms form the basis for classifying the florideophyte red algae and can be found in the remainder of the Rhodymeniophycidae^{49,51,52}, i.e. the Acrosymphytales,

Bonnemaisoniales, Ceramiales, Gigartinales, Halymeniales, Nemastomatales, Peyssonneliales, Plocamiales, Rhodymeniales and Sebdeniales. The great diversity in pre- and postfertilization strategies in the Rhodymeniophycidae suggests that the ancestors of these taxa "experimented" on multiple occasions (i.e., at all taxonomic levels) on how to enhance carpospore production from a single fertilization event. Some taxa did it by the direct production of carposporophytes from the fertilized carpogonium, whereas other taxa produced generative auxiliary cells along with a suite of accompanying diploidization strategies. Because many strategies evolved in parallel, along with accompanying reversals, in all of these lineages, considerably better phylogenetic resolution at all taxonomic levels is needed before the evolutionary pathways for these complex postfertilization patterns can be resolved for Rhodymeniophycidae.

The Rhodymeniophycidae is the largest subclass (5,017 spp) of the Florideophycidae, having begun to diversify about 412 (95% HPD: 359-477) Ma (node '6' in Fig. 1). Its members are diverse with respect to many morphological characters in addition to the postfertilization richness discussed above. For example, all three major types of tetrasporangia division are present; viz., zonate, cruciate, and tetrahedral. However, this feature was also gained and lost on numerous occasions, appearing in some, but not all, Acrosymphytales, Ceramiales and Rhodymeniales. Thus, resolving evolutionary pathways of tetrasporangial diversity in this subclass also awaits improved phylogenetic resolution. Lack of interordinal phylogenetic resolution within the Rhodymeniophycidae is a common theme in the literature^{7,9,31,36,50,52} and remains consistent with our analyses (Supplementary Fig. S1). Indeed, the only relationship consistently resolved is a group^{33,51,52} including the Halymeniales, Rhodymeniales and Sebdeniales (99% MLB and 0.99 BPP in Supplementary Fig. S1). Clearly better phylogenetic resolution is needed within Rhodymeniophycidae before the evolutionary patterns of the intricate anatomical features characterizing the many species of this subclass can be explored.

The Ceramiales constitutes the most diverse florideophyte order (2,654 spp) with a divergence time estimated as 335 (95% HPD: 284-395) Ma (node '7' in Fig. 1). The Ceramiales is distinguished from other florideophycean species by the formation of auxiliary cells after fertilization. Our relaxed clock analysis suggests that this unifying feature evolved during the Devonian (419-359 Ma) to Carboniferous (359-299 Ma) periods³⁹ of the Paleozoic Era.

With the exception of the Conchocelis stage of Porphyra-like algae, preserved as endoliths in Paleozoic carbonates ^{53,54}, red algae are represented in Phanerozoic rocks largely as calcareous skeletons. Several extant florideophyte clades are known to precipitate CaCO₃: as aragonite in a few members of the Peyssonneliales and Nemaliales, and as calcite in the Corallinales (Ca(Mg)CO₃), Sporolithales, Hapalidiales, and Rhodogorgonales. With some uncertainty, representatives of the first two groups have been reported from Carboniferous and Permian (>350 Ma) carbonates^{55,56}, consistent with our molecular clock inference of major florideophyte diversification during the late Paleozoic Era. Remnants of stem group corallines that contain features of biocalcification, partitions, calcified sporangial compartments and trichocytes, occur as well, and can be traced back to the Ordovician Period (485 - 445)Ma)^{57,58}, after the emergence of non-calcified corallines preserved in the Doushantuo rocks (635 - 551 Ma). The evolution of skeletal biomineralization within this clade occurred within the context of an ecosystem-wide increase in carbonate skeletonization that characterizes the Ordovician marine radiation⁵⁹, reflecting an increase in predation pressure, a change in seawater chemistry, or both⁶⁰.

In contrast, crown group diversification of coralline algae appears to be restricted to mid-Mesozoic and younger oceans, as indicated by both fossils and molecular clock estimates^{16,17}. This places coralline diversification within the context of the Mesozoic marine revolution during which many skeleton-forming clades evolved protective responses to the radiation of shell-crushing predators^{61,62}. We note however that the single-gene based molecular clock analysis of Aguirre et al.¹⁶, which does not take into account stem group corallines in Ediacaran to Ordovician rocks¹³⁻ ^{15,54,57,58}, yields estimates of family level diversification within the Corallinales that are about two times younger than our estimates. This apporach also yields an age for the divergence of corallines from nemalialeans (338.26 Ma) far younger than our estimate (579 Ma). If we accept that the Doushantuo Formation contains stem group corallines, then the estimate of Aguirre et al.¹⁶ for the coralline-nemalialean divergence must be too young. One might relax the interpretation that the Doushantuo fossils are stem group corallines, but any clock that accepts them as at least stem group florideophytes is unlikely to yield a Carboniferous date for the nemalialeancoralline divergence. Brooke and Riding⁵⁶ interpret Ordovician fossils as corallines and perhaps even stem group Sporolithaceae; this would also require a divergence within the Corallinales older than that estimated by Aguirre et al.¹⁶. For now, it may be most judicious to note the difference in estimates governed by different calibration strategies, and look to continuing improved sampling of living florideophytes as well as better paleontological constraints, in particular for Paleozoic and older fossils.

Furthermore, when we compared time estimates with/without the two constraints of Aguirre et al. ('c1' for the split of the Sporolithales of 133 Ma, and 'c2' for the split of Corallinaceae and Hapalidiaceae of 117 Ma based on the Cenozoic corallines¹⁷), all divergence times were within the 95% credibility interval (see

Supplementary Table S1 and Fig. S2). It may be that the crown group divergence among the Corallinaceae and Sporolithaceae is accurately captured by the fossil record¹⁷ but that divergence between the two clades was much earlier. Our molecular clock based on seven-genes with broad taxon sampling is consistent with fossils in suggesting that the coralline lineage diverged from other rhodophytes long before the crown group radiation of corallines. This, in turn, is consistent with other evidence that shows the time interval between total group divergence and crown group diversification can be long in eukaryotic clades⁶³.

Few molecular clock analyses have estimated the divergence times within the red algae. In the analysis of Parfrey et al.⁶⁴, which included 88 taxa (4 of them rhodophytes) and 15 genes, divergence estimates depended on molecular model and choice of paleontological constraints. In general however, time estimates for radiation within the Rhodymeniophycidae, the Bangiophyceae-Florideophyceae split, and the initial divergence of red algae are similar to or slightly younger than those reported here. More broadly, both fossils⁶⁵ and several recent molecular clock analyses^{66,67} suggest a mid-Proterozoic origin of photosynthetic eukaryotes but Neoproterozoic and later diversification of taxa within the major clades of Archaeplastida. In contrast, the molecular clock estimates of Berney and Pawlowski⁶⁷ suggest that red and green algae diverged only about 900 million years ago.

Conclusion

We studied the major diversification events within red algae using a multigene dataset (concatenated genes of nuclear EF2, LSU and SSU rRNAs, plastid encoded *psa*A, *psb*A, and *rbc*L, and mitochondrial *cox*1). Our ML phylogeny supports a sister group relationship between the Bangiophyceae and Florideophyceae, and resolves

relationships among the five subclasses of the Florideophyceae. The multigene relaxed clock estimation using multiple fossil constraints suggests that florideophytes arose near the beginning of the Neoproterozoic Era. The major evolutionary divergences within the class occurred in mid-Neoproterozoic to early Paleozoic oceans, beginning with the split of the Hildenbrandiophycidae followed by the appearance of the Nemaliophycidae, Corallinophycidae, Ahnfeltiophycidae and Rhodymeniophycidae. Radiation of the rhodymeniophycidaen algae is thought to have occurred during the mid Paleozoic Era.

These major divergences were accompanied by evolutionary innovations in the carposporophyte stage, maximizing spore production from each fertilization event. The Nemaliophycidae and Ahnfeltiophycidae and Corallinophycidae (in part) did not evolve generative auxiliary cells and the site of fertilization and diploidization in these taxa thus remain restricted to the carpogonium. Nonetheless, in most Rhodymeniophycidae and Corallinophycidae (in part), these two processes (likely independently) became decoupled with this division of labor resulting in the great diversity of carposporophyte types that characterize each of the subclasses. It is precisely the sequence of events starting with the establishment of the female reproductive system in relation to vegetative growth, and leading to the postfertilization carposporophyte that has traditionally formed the basis of red algal classification.

Our research provides the first comprehensive estimation of divergence dates within the Florideophyceae using molecular and fossil data. Although these results are clearly working hypotheses, we are buoyed by the observation that there was considerable congruence between the current results with both previous time

estimates and the fossil record. Future studies that incorporate additional red algal diversity should be used to test the ideas put forth in our study.

Methods

Taxon sampling and sequencing strategy. To establish a well-resolved phylogeny for the red algae, we selected 91 red algal taxa representing 34 orders, 27 of them florideophycean and seven non-florideophycean^{6,7,68}. We generated 180 new sequences for *rbc*L (n = 49), *psa*A (61), *psb*A (58), EF2 (2), SSU (1), LSU (2), and *cox1* (7) genes from florideophycean taxa to reduce missing data in the multigene alignment. Publicly available sequences of red algae, eight green algal representatives (including land plants), and three cyanobacterial species were downloaded from GenBank (Supplementary Table S3). The cyanobacteria *Nostoc* sp. PPC 7120, *Synechocystis* sp. PCC 6803, and *Thermosynechococccus elongatus* BP-1 were used as an outgroup to root the tree.

Genomic DNA was extracted from approximately 5 mg of algal biomass that had been pulverized in liquid nitrogen. The DNeasy Plant Mini Kit (Qiagen GmbH, Hilden, Germany) or Invisorb Spin Plant Mini Kit (Invitek, Berlin-Buch, Germany) was used for DNA extraction following the manufacturers' instructions. PCR and sequencing reactions were conducted using specific primers for each of three plastid genes⁶⁹.: psaA130F, psaA971F, psaA1110R, psaA1530F, psaA1760R, and psaA-3 for *psa*A; psbA-F, psbA-R1, psbA-500F, psbA-600R, and psbA-R2 for *psb*A; and F7, R753, F645, and RrbcS start for *rbc*L. PCR amplification was performed in a total volume of 25 μ l that contained 0.5 U *TaKaRa Ex Taq*TM DNA polymerase (Takara Shuzo, Shiga, Japan), 2.5 mM of each dNTP, 2.5 ul of the 10X *Ex Taq*TM Buffer (Mg²⁺ free), 2 mM MgCl₂, 10 pmol of each primer, and 1-10 ng template DNA. The reaction was carried out with an initial denaturation at 94°C for 10 min, followed by 35 cycles of amplification (denaturation at 94°C for 30 sec, annealing at 50°C for 30 sec, and extension at 72°C for 2 min), with a final extension at 72°C for 10 min. The PCR products were purified using a High Pure PCR Product Purification Kit (Roche Diagnostics GmbH, Mannheim, Germany), in accordance with the manufacturers' instructions. The sequences of the forward and reverse strands were determined for all taxa using commercial sequencing services. The electropherogram output for each specimen was edited using the program Chromas Lite v.2.1.1.

(http://www.technelysium.com.au/chromas.html).

Alignment and phylogeny. Alignments were generated manually using Se-Al v.2.0a11 (http://tree.bio.ed.ac.uk/software/seal/). Because *rbc*L sequences of green algae and land plants are derived from the cyanobacterial primary endosymbiont rather than from a proteobacterium, as in red algae, they were coded as missing data. We used a mixed model of amino acid and DNA sequences for phylogenetic analysis. Translated amino acid sequences for protein-coding genes were used to reduce the possibly misleading effects of nucleotide bias or mutational saturation in our data sets. In addition, only conserved regions of the rDNA alignment were used for phylogenetic analysis. The final dataset contained 102 taxa, comprising a total of 6,966 characters, including 2,053 amino acid positions (500 *psa*A, 301 *psb*A, 463 *rbc*L, 568 EF-2, and 221 *cox*1) and 4,913 DNA positions (3,070 LSU and 1,843 SSU). The final alignment is available upon request from HSY and the Supplementary Information of the journal website at http://www.nature.com/naturecommunications/.

Appropriate evolutionary models were selected separately for each protein and DNA alignment. For the amino acid dataset, the best-fit model was chosen using ModelGenerator v.0.85 (<u>http://bioinf.nuim.ie/modelgenerator/</u>) and preliminary Bayesian analyses using MrBayes v.3.2⁷⁰. The LG model was selected by ModelGenerator as a best fit from available amino acid models under the both AIC (Akaike information criterion) and BIC (Bayesian information criterion). We used the LG model for Maximum Likelihood (ML) analyses using RAxML v.8.1⁷¹. Because MrBayes does not support the LG model, the CPREV substitution model was selected for the Bayesian inference (BI). For the DNA alignment, the GTR substitution model was used.

ML and bootstrap analyses of protein + DNA combined data were conducted under the LG and GTR with independent rate heterogeneity (LG + F + G and GTR + G mixed model) using RAxML. We used 100 independent tree inferences using the default option of which automatically optimized SPR rearrangement and 25 distinct rate categories for this program to identify the best tree. Bootstrap values were calculated with 1,000 replicates using the same substitution model.

Bayesian posterior probabilities from the combined data were estimated under the CPREV + G and GTR + G mixed model using MrBayes. Two independent Metropolis-coupled Markov chain Monte Carlo (MCMCMC) runs with 20 million generations with four chains were run simultaneously. Every 200th generation tree was sampled and compared to determine the burn-in point. Based on the average standard deviation of split frequencies between two independent runs, 5,000,000 generations were determined to appropriate burn-in point. The remaining 150,002 trees from two runs sampled from the putative stationary distribution were used to infer Bayesian posterior probabilities (BPP) at the nodes. The ML and Bayesian tree topologies were congruent with each other.

Molecular clock analysis. The ML tree from RAxML was used to estimate divergence times. The hypothesis of rate constancy among taxa was tested by comparing likelihoods using the likelihood ratio test, given the best ML tree topology with and without the constraint of a molecular clock. As expected from the heterogenous branch lengths, a clock-like evolution of substitution rates was rejected for our data set (data not shown; see also Yoon et al.¹⁰). Thus, the multigene relaxed clock methods were used to estimate divergence times. The Bayesian relaxed dating was conducted using the BEAST v.1.8.1 software package²⁰. This method accommodates rate variation both among lineages and among genes. Branch lengths were estimated using only combined protein data under a LG model. We used combined data (DNA + protein; two partitions) with the separate GTR + G and LG +F + G models for each part. Topologies and branch lengths were linked for partitions and other parameters were unlinked, i.e., substitution matrixes, base or amino acid frequencies, and gamma parameters. The Yule process prior for speciation and uncorrelated lognormal relaxed clock models were used. The prior constraint time of selected nodes were set with upper and lower limits by using normal distribution of mean and standard deviation. The MCMC analyses were run for 20 million generations with sampling every 1,000th generation. Convergence of the MCMC algorithm was assessed by plotting of likelihood value. Initial 5 million generations were discarded as burn-in. In addition, the Birth-Death and Birth-Death Incomplete speciation priors were used to test the robustness of the time estimation (see

Supplementary Table S2). TreeAnnotator (part of BEAST package) was used to visualize the maximum clade credibility phylogeny and time estimation.

The following seven constraints were used for divergence time estimation (see Supplementary Fig. S1). The first node 'a' was constrained at 1,174-1,222 ($1,198\pm12$) Ma based on a Pb-Pb date for carbonates correlative with well-preserved fossils of the multicellular filamentous red alga *Bangiomorpha*^{11,12}. Because there are simple filamentous species within the Compsopogonophyceae and Stylonematophyceae (i.e., Erythrotrichia, Compsopogon, Rhodochaete, Bangiopsis, and Stylonema), we used node 'a' for the constraint point for the *Bangiomorpha* fossil data, instead of the node '1' ancestral to the Bangiophyceae and Florideophyceae (see Supplementary Fig. S2). Coralline algae (Corallinophycidae) provide an excellent context to stratigraphy¹³⁻¹⁵ and molecular clock analysis because of their calified cell wall structure^{16,17}. We constrained node 'b' to a date of $635 - 551 (593 \pm 21)$ Ma based on an early stem group of corallinalean red algae from the Doushantuo Formation, China¹³⁻¹⁵. The Doushantuo Formation contains the stem group of corallines, i.e., *Thallophyca* and Paramecia. Light microscopy and SEM data revealed fossilized features of the coralline algae such as complex pseudoparenchymatous thalli, specialized reproductive structures (carposporophytes, tetrasporangia and tetraspore-like cells) and cell to cell fusion¹⁴. The U-Pb zircon dates of volcanic ash beds within the Doushantuo Formation, animal fossils, and synchronous deglaciation records support the age of the formation¹⁵. Other coralline data from Aguirre et al.^{16,17} were available for calibration points, i.e., 130-136 (133 \pm 1.5) Ma for node 'c1' for the Sporolithales split, and 114-120 (117 ± 1.5) Ma for 'c2' for the split of Corallinales and Hapalidiales. The four nodes marked 'd-g' were designated constraints of the Viridiplantae (green algae and plants) used in a recent study¹⁸, including three fossil constraints, i.e., 471480 (475.5 ±2.25) Ma for the age of land plants (node 'd'), 401-422 (411.5 ±5.25) Ma for the age of the euphyllophytes (node 'e'), 313-351 (332 ±9.5) Ma for the age of the seed plants (node 'f'), and 138-163 (150.5 ±6.25) Ma for the split of Eudicotyledoneae (node 'g').

To test how the calibrations affected the age estimation of red algae, we devised 14 distinct calibration scenarios based on the constraint category³⁷ (Supplementary Table S1). The Bangiomorpha fossil (constraint 'a') and green plant lineage (constraints 'd-g') were considerd "outgroup" constraints. Coralline fossils of the Doushantuo formation (constraint 'b') and estimated times of coralline classes (constraints 'c1 and c2') were consided "ingroup" constraints of the Florideophyceae. The constraint 'b' was "safe" because it showed high level of taxonomic resolution and accurate age^{14,15}. However, constraints 'c1 and c2' were considered "risky" because these were "secondary" calibrations derived from a previous estimation^{16,17}. In the first calibration scenario, we used all seven age constraints (C7) for age estimation at the nodes. In the second, we used six constraints excluding one constraint (C6-*i*, i = a-g) in order to evaluate the impact of each constraint. The constraint 'c' is only a "secondary" constraint in the present study, in order to provide an empirical test for the impact of the probability range in the normal prior distribution (mean and S.D.). We also applied additional analyses with two and ten times the standard deviation (S.D.), i.e., C6-c 2SD and C6-c 10SD. In the third calibration scenario we compared two analyses with "outgroup" only (C5-bc) and "green" lineage only (C4-*abc*). In the fourth, we used a red algal constraint only, i.e., C2ab (constrains 'a' and 'b') and C1a, C1b, and C1c (constraint 'a', 'b', and 'c', respectively).

The parametric prior distributions were compared using Bayesian inference implemented in the BEAST package. In the uniform prior distribution, all calibrations were specified with minimum boundary and arbitrarily large maximum boundary (1.0e100). In the normal prior distribution, all calibrations were specified with minimum and maximum boundaries described above. We compared the ages of all estimates (Y; dependent variable) and those under the C7 scenario with normal prior distribution (C7 *nor*, X; independent variable) by regression analysis (Y = a + bX; Supplementary Table S1).

References

- Guiry, M. D. & Guiry, G. M. AlgaeBase. (2015) Available at: http://www.algaebase.org (Accessed: 9th June 2015).
- Saunders, G. W. Applying DNA barcoding to red macroalgae: a preliminary appraisal holds promise for future applications. *Phil. Trans. R. Soc. London. B* 360, 1879-1888 (2005).
- Garbary, D. J. & Gabrielson, P. W. Taxonomy and evolution. in *Biology of the Red Algae.* (eds Cole, K. M. & Sheath, R. G.) Ch. 18, 477-498 (Cambridge University Press, 1990).
- Stegenga, H. The life histories of *Rhodochorton purpureum* and *Rhodochorton floridulum* (Rhodophyta, Nemaliales) in culture. *Br. Phycol. J.* 13, 279-289 (1978).
- 5. Fredericq, S. & Lopez-Bautista, J. Characterization and phylogenetic position of the red alga *Besa papillaeformis* Setchell: an example of progenetic

heterochrony? *Constancea* 83.2 (2002). Available at: <u>http://ucjepsberkeleyedu/constancea/83/indexhtml (Accessed: 9th June 2015,</u> The online continuation of *University of California Publications in Botany,* 1902-2002. Special Festschrift to honor PC Silva)

- Saunders, G. W. & Hommersand, M. H. Assessing red algal supraordinal diversity and taxonomy in the context of contemporary systematic data. *Am. J. Bot.* 91, 1494-1507 (2004).
- Le Gall, L. & Saunders, G. W. A nuclear phylogeny of the Florideophyceae (Rhodophyta) inferred from combined EF2, small subunit and large subunit ribosomal DNA: Establishing the new red algal subclass Corallinophycidae. *Mol. Phylogenet. Evol.* 43, 1118-1130 (2007).
- Yoon, H. S., Zuccarello, G. C. & Bhattacharya, D. in Cellular Origin, Life in Extreme Habitats and Astrobiology, Vol.13. Red Algae in the Genomic Age (eds Seckbach, J. & Chapman, D. J.) Evolutionary history and taxonomy of red algae, 25-42 (Springer, 2010).
- Verbruggen, H. et al. Data mining approach identifies research priorities and data requirements for resolving the red algal tree of life. *BMC Evol. Biol.* 10, 16 (2010).
- Yoon, H. S. et al. A molecular timeline for the origin of photosynthetic eukaryotes. *Mol. Biol. Evol.* 21, 809-818 (2004).
- Butterfield, N. J. *Bangiomorpha pubescens* n. gen., n. sp.: implications for the evolution of sex, multicellularity, and the Mesoproterozoic/Neoproterozoic radiation of eukaryotes. *Paleobiology* 26, 386-404 (2000).
- Knoll, A. H. The multiple origins of complex multicellularity. *Annu. Rev. Earth Planet Sci.* 39, 217–239 (2011).

- Xiao, S., Knoll, A. H., Yuan, X. & Pueschel C. M. Phosphatized multicellular algae in the Neoproterozoic Doushantuo formation, China, and the early evolution of florideophyte red algae. *Am. J. Bot.* **91**, 214-227 (2004).
- Xiao, S., Zhang, Y. & Knoll, A. H. Three-dimensional preservation of algae and animal embryos in a Neoproterozoic phosphorite. *Nature* 391, 553-558 (1998).
- Condon, D. et al. U-Pb ages from the Neoproterozoic Doushantuo Formation, China. *Science* 308, 95-98 (2005).
- Aguirre, J., Perfecti, F. & Braga, J. C. Integrating phylogeny, molecular clocks, and the fossil record in the evolution of coralline algae (Corallinales and Sporolithales, Rhodophyta). *Paleobiology* 36, 519-533 (2010).
- Aguirre, J., Riding, R. & Braga, J. C. Diversity of coralline red algae: origination and extinction patterns from the early Cretaceous to the Pleistocene. *Paleobiology* 26, 651-667 (2000).
- Magallón, S., Hilu, K. W. & Quandt, D. Land plant evolutionary timeline: gene effects are secondary to fossil constraints in relaxed clock estimation of age and substitution rates. *Am. J. Bot.* **100**, 556-573 (2013).
- Thorne, J. L. & Kishino, H. Divergence time and evolutionary rate estimation with multilocus data. *Syst. Biol.* 51, 689-702 (2002).
- Drummond, A. J., Suchard, M. A., Xie, D. & Rambaut, A. Bayesian phylogenetics with BEAUti and BEAST 1.7. *Mol. Biol. Evol.* 29, 1969-1973 (2012).
- Pueschel, C. M. An expanded survey of the ultrastructure of red algal pit plugs.
 J. Phycol. 25, 625-636 (1989).
- 22. Choi, H.-G., Kraft, G. T. & Saunders, G. W. Nuclear small-subunit rDNA sequences from *Ballia* spp (Rhodophyta): proposal of the Balliales ord. nov.,

Balliaceae fam. Nov., *Ballia nana* sp. nov. and *Inkyuleea* gen. nov. (Ceramiales). *Phycologia* **39**, 272-287 (2000).

- Pueschel, C. M. Secondary pit connections in *Hildenbrandia* (Rhodophyta, Hildenbrandiales). *Br. Phycol. J.* 23, 25-32 (1988).
- Nichols, H. W. Culture and development of *Hildenbrandia rivularis* from Denmark and North America. *Am. J. Bot.* 52, 9-15 (1965).
- Pueschel, C. M. & Cole, K. M. Rhodophycean pit plugs: an ultrastructural survey with taxonomic implications. *Am. J. Bot.* 69, 703-720 (1982).
- Scott, J., Thomas. J. & Saunders, B. Primary pit connections in *Compsopogon coeruleus* (Balbis) Montagne (Compsopogonales, Rhodophyta). *Phycologia* 27, 327-333 (1988).
- Pueschel, C. M. Ultrastructural observations of tetrasporangia and conceptacles in *Hildenbrandia* (Rhodophyta: Hildenbrandiales). *Br. Phycol. J.* 17, 333-341 (1982).
- Saunders, G. W. & Bailey, J. C. Phylogenesis of pit-plug associated features in the Rhodophyta: inferences from molecular systematic data. *Can. J. Bot.* 75, 1436-1447 (1997).
- Harper, J. T. & Saunders, G. W. A re-classification of the Acrochaetiales based on molecular and morphological data, and establishment of the Colaconematales ord. nov. (Florideophyceae, Rhodophyta). *Eur. J. Phycol.* 37, 463-476 (2002).
- Pueschel, C. M., Trick, H. N. & Norris, J. N. Fine structure of the phylogenetically important marine alga *Rhodogorgon carriebowensis* (Rhodophyta, Batrachospermales?). *Protoplasma* 166, 78-88 (1992).

- Le Gall, L., Payri, C., Bittner, L. & Saunders, G. W. Multigene phylogenetic analyses support recognition of the Sporolithales ord. nov. *Mol. Phylogenet. Evol.* 54, 302-305 (2010).
- Nelson W. A. et al. Multi-gene phylogenetic analyses of New Zealand coralline algae: *Corallinapetra nvaezelandiae* gen. et sp. nov. and recognition of the Hapalidiales ord. nov. *J. Phycol.* 51, 454-468 (2015).
- Maggs, C. A. & Pueschel, C. M. Morphology and development of *Ahnfeltia* plicata (Rhodophyta); Proposal of Ahnfeltiales ord. nov. J. Phycol. 25, 333-251 (1989).
- Huisman, J. M., Sherwood, A. R. & Abbott, I. A. Morphology, reproduction, and the 18S rRNA gene sequence of *Pihiella liagoraciphila* gen. et sp. nov. (Rhodophyta), the so-called 'monosporangial discs' associated with members of the Liagoraceae (Rhodophyta), and proposal of the Pihiellales ord. nov. *J. Phycol.* **39**, 978-987 (2003).
- 35. Withall, R. D. & Saunders, G. W. Combining small and large subunit ribosomal DNA genes to resolve relationships among orders of Rhodymeniophycidae (Rhodophyta): recognition of the Acrosymphytales ord. nov. and Sebdeniales ord. nov. *Eur. J. Phycol.* **41**, 379-394 (2006).
- Choi, H.-G. et al. Phylogenetic relationships among lineages of the Ceramiaceae (Ceramiales, Rhodophyta) based on nuclear small subunit rDNA sequence data. J. Phycol. 44, 1033-1048 (2008).
- Sauquet H. et al. Testing the impact of calibration on molecular divergence times using a fossil-rich group: the case of *Nothofagus* (Fagales). *Syst. Biol.* 61, 289-313 (2012).

- Ho, S. Y. W. & Phillips, M. J. Accounting for calibration uncertainty in phylogenetic estimation of evolutionary divergence times. *Syst. Biol.* 58, 367-380 (2009).
- Walker J. D., Geissman J. W., Bowring, S. A. & Babcock, L. E. The Geological Society of America Geologic Time Scale. *Geol. Soc. Am. Bull.* 125, 259-272 (2013).
- Lim, B. L., Kawai, H., Hori, H. & Osawa, S. Molecular evolution of 5S ribosomal RNA from red and brown algae. *Jpn. J. Gen.* 61, 169-176 (1986).
- Turner E. C. & Kamber, B. S. Arctic Bay Formation, Borden Basin, Nunavut (Canada): Basin evolution, black shale, and dissolved metal systematics in the Mesoproterozoic ocean. *Precambrian Res.* 208-211, 1-18 (2012).
- 42. Tappan, H. Possible eukaryotic algae (Bangiophycidae) among early Proterozoic microfossils. *Bull. Geol. Soc. Am.* **87**, 633-639 (1976).
- 43. Searles, R. B. The strategy of the red algal life history. *Am. Nat.* 115, 113-120 (1980).
- Maggs, C. A. et al. Speciation in red algae: members of the Ceramiales as model organisms. *Integr. Comp. Biol.* 51, 492-504 (2011).
- Elie, M. et al. A red algal bloom in the aftermath of the Marinoan Snowball Earth. *Terra Nova* 19, 303–308 (2007).
- Kodner, R. B., Pearson, A., Summons, R. E. & Knoll, A. H. Sterols in red and green algae: quantification, phylogeny, and relevance for the interpretation of geologic steranes. *Geobiology* 6, 411-420 (2008).
- Hommersand, M. H. & Fredericq, S. An investigation of cystocarp development in *Gelidium pteridifolium* with a revised description of the Gelidiales (Rhodophyta). *Phycologia* 27, 254-272 (1988).

- Fredericq, S. & Hommersand, M. H. Proposal of the Gracilariales ord. nov. (Rhodophyta) based on an analysis of the reproductive development of *Gracilaria verrucosa. J. Phycol.* 25, 213-227 (1989).
- 49. Hommersand, M. H. & Fredericq, S. in Biology of the Red Algae (eds Cole, K. M. & Sheath, R. G.) Sexual reproduction and cystocarp development, 305-345 (Cambridge University Press, 1990).
- Fredericq, S. & Norris J. N. A new order (Rhodogorgonales) and family (Rhodogorgonaceae) of red algae composed of tropical calciferous genera, *Renouxia* sp. nov. and *Rhodogorgon. Crypt. Bot.* 5, 316-331 (1995).
- 51. Withall, R. D. & Saunders, G. W. Combining small and large subunit ribosomal DNA genes to resolve relationships among orders of Rhodymeniophycidae (Rhodophyta): recognition of the Acrosymphytales ord. nov. and Sebdeniales n ord. nov. *Eur. J. Phycol.* **41**, 379-394 (2006).
- Krayesky, D. M. et al. A new order of crustose red algae based on the Peyssonneliaceae with an evaluation of the ordinal classification of the Florideophyceae (Rhodophyta). *Proc. Biol. Soc. Washington* 123, 364-391 (2009).
- 53. Campbell, S. E. *Palaeoconchocelis starmachii*, a carbonate boring microfossil from the upper Silurian of Poland (425 Million Years Old): Implications for the evolution of the Bangiaceae (Rhodophyta). *Phycologia* 19, 25-36 (1980).
- Vogel, K. & Brett, C. E. Record of microendoliths in different facies of the Upper Ordovician in the Cincinnati Arch region USA: The early history of light-related microendolithic zonation. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 281, 1–24 (2009).
- 55. Wray, J. L. Calcareous Algae. Amsterdam: Elsevier. (1977).

- James, N. P., Wray, J. L. & Ginsburg, R. N. Calcification of encrusting aragonitic algae (Peyssonneliaceae): implications for the origin of late Paleozoic reefs and cements. *J. Sed. Res.* 58, 291-303 (1988).
- 57. Brooke, C. & Riding, R. Ordovician and Silurian coralline red algae. *Lethaia*31, 185-195 (1998).
- Riding, R. & Braga, J. C. *Halysis* Høeg, 1932 An Ordovician coralline red alga? *J. Paleo.* 79, 835-841 (2005).
- Harper, D. A. T. The Ordovician biodiversification: Setting an agenda for marine life. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 232, 148-166 (2006).
- Pruss, S., Finnegan, S., Fischer, W. W. & Knoll, A. H. Carbonates in skeletonpoor seas: New insights from Cambrian and Ordovician strata of Laurentia. *Palaios* 25, 73-84 (2010).
- Vermeij, G. J. The Mesozoic marine revolution; evidence from snails, predators and grazers. *Paleobiology* 3, 245-258 (1977).
- Steneck, R. A., Hacker, S. D. & Dethier, M. N. Mechanisms of competitive dominance between crustose coralline algae – An herbivore-mediated competitive reversal. *Ecology* 72, 938-950 (1991).
- Knoll, A. H., Wörndle, S. & Kah, L. Covariance of microfossil assemblages and microbialite textures across a late Mesoproterozoic carbonate platform. *Palaios* 28, 453-470 (2013).
- Parfrey, L., Lahr, D., Knoll, A. H. & Katz, L. A. Estimating the timing of early eukaryotic diversification with multigene molecular clocks. *Proc. Natl. Acad. Sci. USA.* 108, 13624–13629 (2011).
- Knoll, A. H. Paleobiological perspectives on early eukaryotic evolution. *Cold* Spring Harb. Perspect. Biol. Doi: 101101/cshperspecta016121 (2014).

- Eme, L., Sharpe, S. C., Brown, M. W. & Roger, A. J. On the age of the eukaryotes: evaluating evidence from fossils and molecular clocks. *Cold Spring Harb. Perspect. Biol.* Doi: 101101/cshperspecta016139 (2014).
- 67. Berney, C. & Pawlowski, J. A molecular time-scale for eukaryote evolution recalibrated with the continuous microfossil record. *P. Roy. Soc. Lond. B Bio.*273, 1867-1872 (2006).
- Yoon, H.S. et al. Defining the major lineages of red algae (Rhodophyta). J. *Phycol.* 42, 482-492 (2006).
- Yoon, H. S., Hackett, J. D. & Bhattacharya, D. A single origin of the peridininand fucoxanthin-containing plastids in dinoflagellates through tertiary endosymbiosis. *Proc. Natl. Acad. Sci. USA*. 99, 11724-11729 (2002).
- Ronquist, F. et al. MrBayes 3.2: Efficient Bayesian phylogenetic inference and model choice across a large model space. *Syst. Biol.* 61, 539-542 (2012).
- 71. Stamatakis, A. RAxML Version 8: A tool for phylogenetic analysis and postanalysis of large phylogenies. Bioinformatics Doi: 10.1093/bioinformatics/btu033 (2014).

Acknowledgements

We thank Drs Ga Youn Cho, Myung Sook Kim, and Junxue Mei for providing several red algal samples and Tae-Wook Kang for assistance in data analysis. This work was partly supported by the Korean Rural Development Administration Nextgeneration BioGreen21 (PJ011121), the National Research Foundation of Korea (MEST: 2014R1A2A2A01003588), and Marine Biotechnology Program (PJT200620) funded by Ministry of Oceans and Fisheries, Korea to HSY, the Basic Science Research Program through the National Research Foundation of Korea (NRF) founded by the Ministry of Science, ICT and Future Planning (2014R1A1A1006512) to ECY, and by a grant from the U.S. National Science Foundation (0936884, 1317114, 0937978) to HSY, DB, and SF for work on algal evolution. AHK's research is sponsored, in part, by the NASA Astrobiology Institute.

Authors contributions

ECY, SMB and HSY designed the project. ECY and LG performed experiments, data analysis. ECY, SMB and HSY drafted the manustript. DB, GWS, AHK and SF contributed data interpretation and manuscript revision. All authors read and approved the final manuscript.

Additional information

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications/

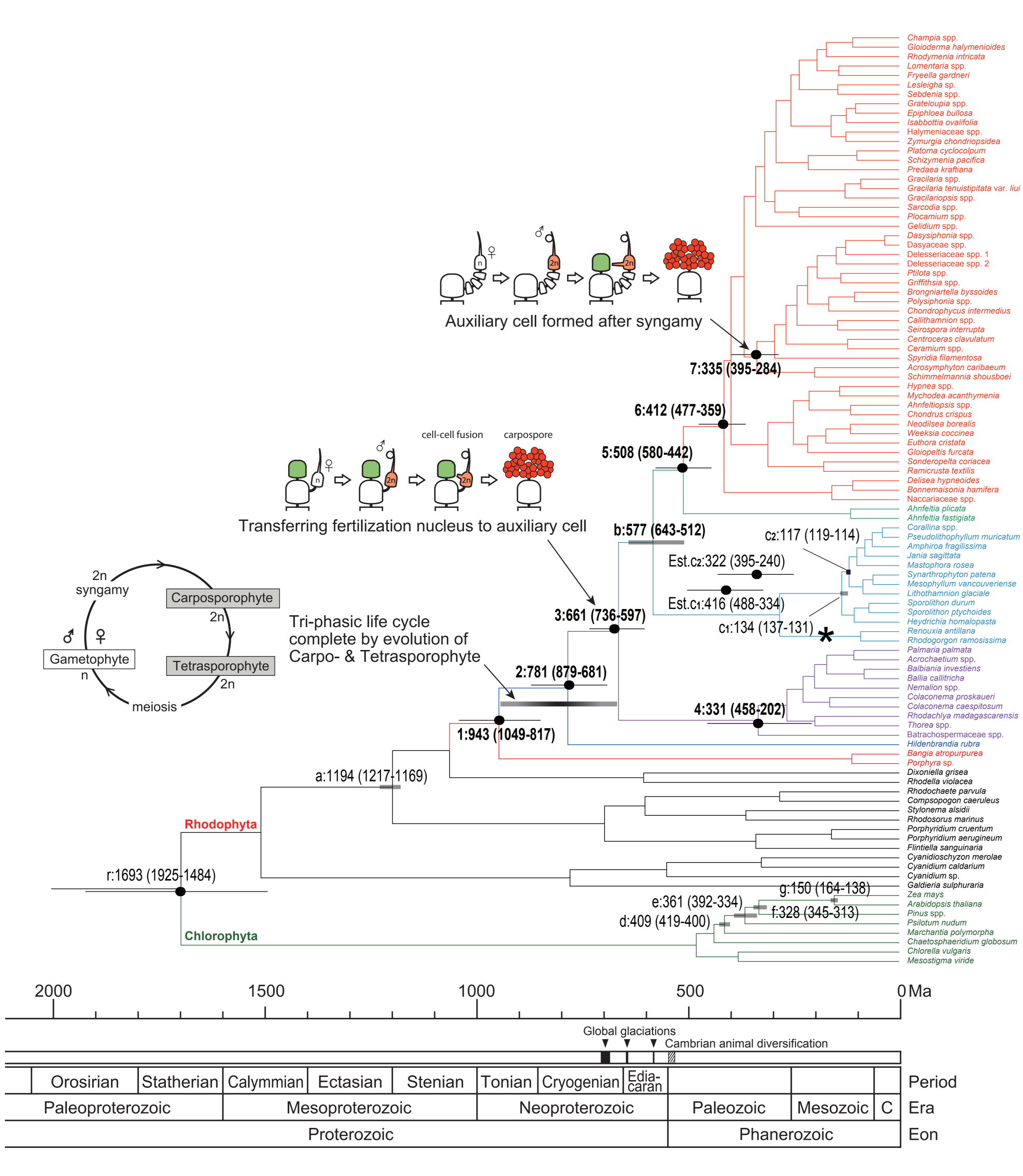
Competing financial interests

The authos declare no competing financial interests.

Figure legends

Figure 1 | Divergence time and evolution of the Florideophyceae.

Estimated times of major divergences based on multigene relaxed clock analysis using the best RAxML tree. Branch lengths are proportional to divergence times (i.e., millions of years ago, Ma). Labels on the node refer to the same splits shown in Supplementary Fig. S1. Seven major divergence times (nodes '1-7') and seven constraints (nodes 'a-g') are indicated by mean and 95% HPD (horizontal bar) in parenthesis. The node 'r' refers to the root time of the Plantae (i.e., red and green algae). The 'Est.c1' and 'Est.c2' are estimated times of nodes 'c1 and c2' without constraint age, respectively. The three key evolutionary events are indicated with diagrams at the best estimate time frame (arrows). Complete triphasic life cycle of red algae accomplished by the evolution of carposporophyte and tetrasporophyte between time frame of nodes '1' and '3'. On the female gametophyte, the fertilization nucleus in the carpogonium (fertilized egg) moved to an auxiliary cell by various so-called 'cell-to-cell fusion' mechanisms followed by carposporophyte development and sporic meiosis. This feature is commonly found in most of the florideophycean red algae (except the Hidenbrandiophycidae and Corallinophycidae) and may have evolved at the time of node '3'. Asterisk (*) indicates loss of cell-to-cell fusion in the Corallinophycidae. Formation of an auxiliary cell after fertilization (syngamy) is a unique feature of the Ceramiales shown as node '7'. The geologic timeline is given under the chronological timeline in million years scale. Three global glaciations were hypothesized to have occurred 716-670, 645-635, and 581-579 Ma ago (three arrowheads in the Neoproterozoic Era). The Cambrian animal diversification occurred approximately 520–543 Ma, at the beginning of the Paleozoic Era.



		Rhodymeniales	
		Sebdeniales	
		Halymeniales	
		Nemastomatales	
		Gracilariales	i
		Plocamiales Gelidiales	
YCEAE	Rhodymeniophycidae	Ceramiales	
FLORIDEOPHY		Acrosymphytales	
DEC		Gigartinales	
OR		Peyssonneliales	
		Bonnemaisoniales	
L LL	Ahnfeltiophycidae	Ahnfeltiales	
		Corallinales	ז
	Corallinophycidae	Hapalidiales	
		Sporolithales	
		Rhodogorgonales	
	Nemaliophycidae Hildenbrandiophycidae	Palmariales Acrochaetiales Balbianiales Balliales Nemaliales Colaconematales Rhodachlyales Thoreales Batrachospermales	
	BANGIOPH		
YCEAE	RHODELLOPH COMPSOPOGONOPH		
	STYLONEMATOPH		
ICEAE	PORPHYRIDIOPH		
YCEAE	CYANIDIOPH		

Supplementary Information

Divergence time estimates and evolution of major lineages in the florideophyte red algae

Eun Chan Yang^{1,2}, Sung Min Boo³, Debashish Bhattacharya⁴, Gary W Saunders⁵, Andrew H Knoll⁶, Suzanne Fredericq⁷, Louis Graf⁸, & Hwan Su Yoon⁸*

¹Marine Ecosystem Research Division, Korea Institute of Ocean Science &

Technology, Ansan 15627, Korea

²Department of Marine Biology, Korea University of Science and Technology,

Daejeon 34113, Korea

³Department of Biology, Chungnam National University, Daejeon 305-764, Korea

⁴Department of Ecology, Evolution and Natural Resources, Rutgers University, New Brunswick, NJ 08901, USA

⁵Department of Biology, University of New Brunswick, Fredericton, NB E3B 5A3 Canada

⁶Department of Organismic and Evolutionary Biology, Harvard University,

Cambridge, MA 02138, USA

⁷Department of Biology, University of Louisiana at Lafayette, Lafayette, LA 70504-2451, USA

⁸Department of Biological Sciences, Sungkyunkwan University, Suwon 16419, Korea

List of files

Supplementary Figure S1. Phylogenetic relationships of the Florideophyceae.

Tree of the Florideophyceae with other red algal classes and green plants inferred by a likelihood analysis using the concatenated EF2, *cox*1, *psa*A, *psb*A, *rbc*L (protein), and LSU, SSU (DNA) supermatrix under the LG+F+G and GTR+G mixed model. Maximum likelihood bootstrap (MLB) support value and Bayesian posterior probabilities (BPP) are shown near the branches. Red algal systematic ranks (order, subclass, and class) are indicated by the taxon name. The characters 'a-g' in dark circle indicates that fossil or molecular based constraint points for relaxed molecular clock estimation. Numbers '1-7' in dark circle indicates major divergence points in the Florideophyceae.

Supplementary Figure S2. The best phylogeny of red algae with selected 31 nodes including seven constraints.

Supplementary Table S1. The constraint scenarios, age estimates (in Ba) for selected nodes, and regression analysis results with the normal and uniform distribution priors.

Supplementary Table S2. Time estimations comparison resulted in the BEAST analyses under different priors, i) Yule Process, ii) Birth-Death and iii) Birth-Death Incomplete process.

Supplementary Table S3. Species examined in the present study.

The sequence determined in this study are shown in **bold** text.

Supplementary Data1. Seven gene combined alignment.

The data set used in the present study. Nexus format alignment included partition and the best tree topology which have been inferred from the maximum likelihood search and used in molecular clock estimations.

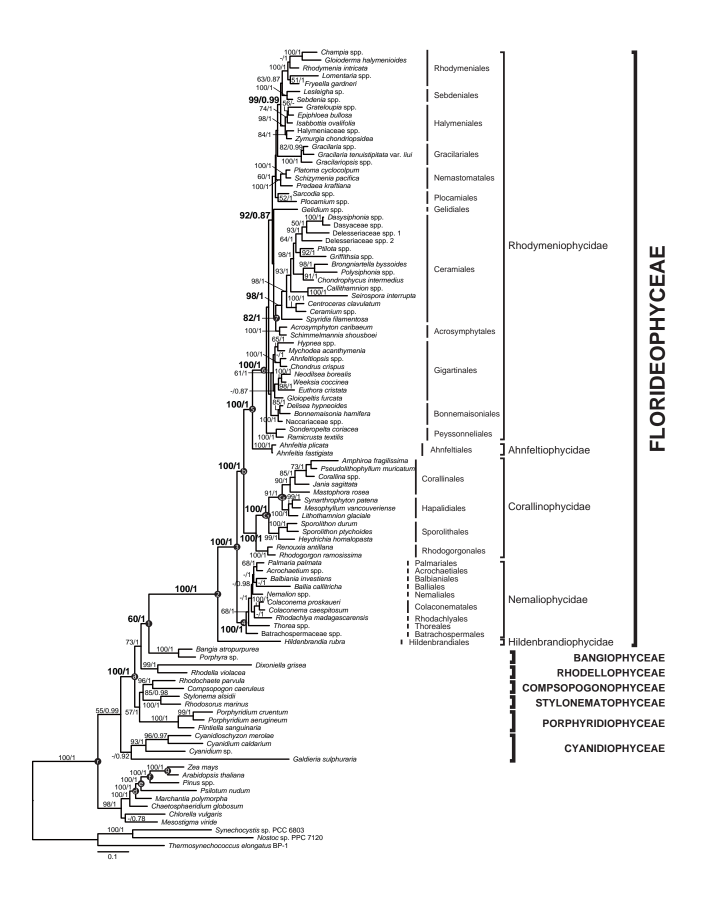


Figure S1

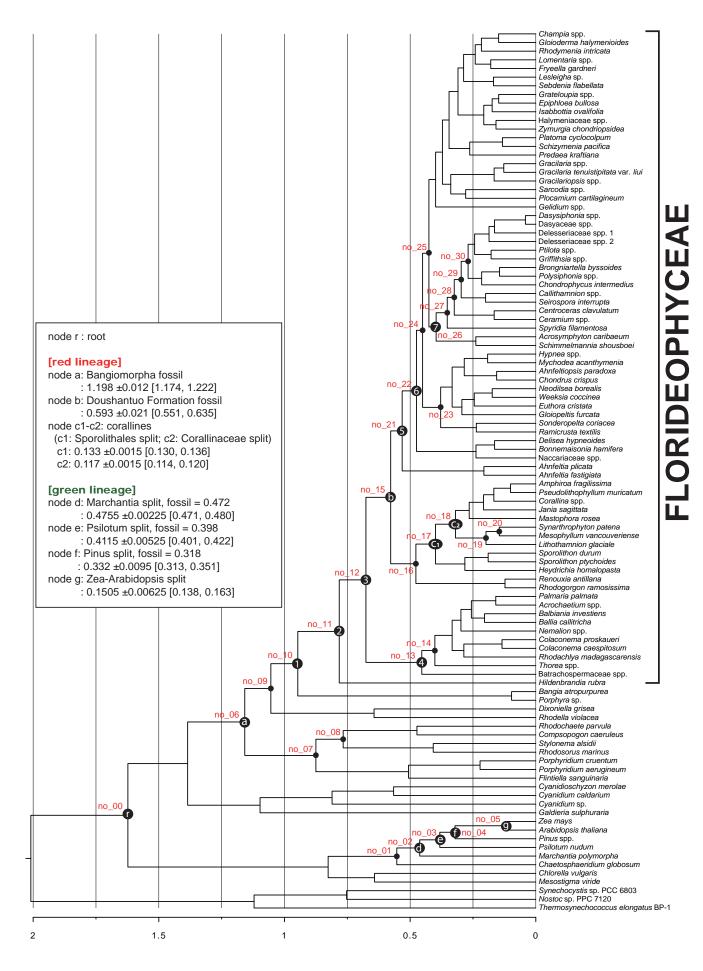


Figure S2

a	+	+	+		+	+	+	+	+	+	+	+	+		+	+			+		+	+	+	+	+	+	+		+	+			 Bangiomorpha fossil Doushantuo Formation fossil 	red algae red algae
c1, c2		-	+	+		+	+	+			+	+			+		+					+	+	÷.	+	+			+		+		c. Corallines estimation	red algae
d	+	+	+	+	+	+	+	+	+	+	+	+	+	+				+	+	+	+	+	Ŧ	+	+	+	+	+				Ŧ	d. Land plants age	green algae
e	+	+	+	+	+	+	+	+	+		+	+	+	+					+	+	+	+	+		+	+	+	+					e. Euphyllophytes split	green algae
f	+	+	+	+	+	+	+	+	+	+		+	+	+					+	+	+	+	+	+		+	+	+					f. Seed plants split	green algae
g	+	+	+	+	+	+	+	+	+	+	+		+	+					+	+	+	+	+	+	+		+	+					g. Eudicotyledoneae split	green algae
note:		x2	x10				x2	x10					out	green	red												out	green	red					
		S.D.	S.D.				S.D.	S.D.					only	only	fossils												only	only	fossils				e de la companya de la	≦u⊡
																		_																
Prior for constraint:	Normal d																		Uniform														H	ШЧ
Node id	C7	C7 (2SD)	C7 (105D)	C6 -a	C6 -b	C6 -c	C6 -c (2SD)	C6 -c (10SD)	C6 -d	С6 -е	C6 -f	C6 -g	C5 -bc	C4 -abc	C2 ab	C1 a	C1 b	C1 c	C7	C6 -a	C6 -b	C6 -c	C6 -d	С6 -е	C6 -f	C6 -g	C5 -bc	C4 -abc	C2 ab	C1 a	C1 b	C1 c	<u>6</u>	
r — node 00	1.6026	1.7008		1 2610	1.6740	2,2860		1.3448	1 7 2 6 0	1.6900	1.6761	1 7 6 1 9	10057	n.a.	1 (000	1.7144	1 6090	0.5221	2.0160	2 2045	2 6 9 4 9	2 2742	2.4236	2 7 2 0 4	2.5042	2.4966	2 6 1 1 0	2 2000	2 2041	2.2675	2.0170	0.7602	Le Le	≣O
r — node_00 node 01			0.3794																	0.8440			0.8751											≦≻
d - node 02			0.3357					0.3567												0.7067										0.6634			18	≡_
			0.2730																											0.5337			18	≣⊥
e — node_03 f — node 04	0.3611			0.3659				0.2931										0.1205		0.5755					0.5156						0.4590			E T
																							0.2584										1	
g — node_05			0.1072																														-th	
a — node_06			0.7443																				1.7063										102	≣ш
node_07			0.4694																				1.2983											≣o ∣ι
node_08			0.4092																				1.1435											ē 🗆 🛛
node_09			0.6751																				1.5794										I L	=
(1) — node_10			0.6063																	1.3790			1.4411										IN 48	
(2) — node_11			0.4416							0.7607													1.1966						1.1282		0.9826			≣O I.
(3) — node_12			0.3777																				1.0437										B	
(4) — node_13			0.2440							0.3464										0.6586					0.7162		0.7485		0.6335		0.5106		미니션	≘ I`
node_14			0.1973																	0.5770					0.6307				0.5366		0.4244			┋╙╴╽╭
b — node_15			0.3256					0.5018										0.1941		0.9226		0.8864	0.9118						0.8723		0.7627			Ē 4
node_16			0.2220																				0.7785										∽」└Ր─═	≣ L
c1 — node_17			0.1617																				0.6834							0.5723				
c2 — node_18			0.1329																	0.5208					0.5669					0.4637				
node_19	0.0839	0.0856	0.0917	0.0835														0.0748					0.3604				0.3801	0.3096	0.3068	0.2975	0.2574	0.1054		GREE
node_20	0.0602			0.0603				0.1283										0.0573					0.2770				0.2874			0.2306				=
(5) — node_21			0.2962																				0.8362							0.7645		0.2601		
(6) — node_22		0.3958		0.3968				0.4138												0.7413					0.7978		0.8201	n.a.		0.6706	n.a.	0.2316		
node_23			0.2122					0.3221							0.3930	n.a.	n.a.	0.1291		0.5784			n.a.		0.6305			n.a.	n.a.	n.a.	n.a.	0.1808		
node_24		0.3696		0.3685						0.3890					0.4736			0.1519		0.7054			n.a.		0.7379			n.a.	0.6711	n.a.	n.a.	0.2198		
node_25			0.2424																				0.6792							0.6029	n.a.	0.2068		
(7) — node_26	0.3345	0.3151	0.2262	0.3163	0.3143	0.3848	0.3991	0.3448	0.3439	0.3231	0.3238	0.3499	0.3845	0.5057	0.4246	0.4574	0.4171	0.1335	0.7285	0.6171	0.7092	0.5954	0.6364	0.7103	0.6702	0.6449	0.6954	0.5842	0.6325	0.5613	n.a.	0.1928		
node_27	0.2927	0.2788	0.2016	0.2801	0.2789	0.3402	0.3590	0.3093	0.3091	0.2860	0.2880	0.3118	0.3447	0.4517	0.3788	0.4073	0.3706	0.1196	0.6498	0.5505	0.6341	0.5317	0.5628	0.6334	0.6000	0.5767	0.6205	0.5200	0.5631	0.4952	0.5680	0.1715		
node_28	0.2655	0.2542	0.1856	0.2536	0.2538	0.3131	0.3299	0.2846	0.2845	0.2601	0.2616	0.2820	0.3157	0.4145	0.3478	0.3755	0.3414	0.1099	0.5995	0.5055	0.5834	0.4907	0.5194	0.5809	0.5512	0.5325	0.5736	0.4791	0.5167	0.4544	0.5157	0.1586		
node_29			0.1691																				0.4744											
node_30	0.2147	0.2061	0.1511	0.2048	0.2080	0.2571	0.2736	0.2408	0.2348	0.2075	0.2123	0.2300	0.2590	0.3379	0.2880	0.3125	0.2837	0.0912	0.4994	0.4183	0.4856	0.4085	0.4293	0.4830	0.4499	0.4376	0.4741	0.3999	0.4306	0.3734	0.4195	0.1305		
gression slope (b)	1	0.9794	0.6438	0.8491	0.9645	1.1162	1.0116	0.7353	1.0014	0.9832	0.9990	1.0131	1.0046	0.9349	0.8990	0.9035	0.8566	0.2723	1.4936	1.1825	1.4632	1.2388	1.2695	1.4066	1.3544	1.3159	1.3978	1.2212	1.2054	1.2215	1.0799	0.4168	Regression function	
						0.0000	0.0500	0.0000	0.0050	0.0074	0.0040	0.9975	0.0555	0.0505	0.0628	0.0576	0.05.40	0.0504	0.0591	0.9549	0.0505	0.0007	0.0546	0.0400	0.0521	0.0526	0.05.27	0.0576	0.0575	0.005.4	0.0559	0.0506	Y = a + bX (Y, C7; X, CN)	

Table S1. The constraint scenarios, age estimates (in Ba) for selected nodes, and regression analysis results with the normal and uniform distribution priors.

2.5 2.0 1.5 1.0 0.5 c node_00 node_01 node_02 node_19 node_20 node. node

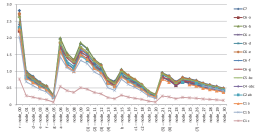
2.5

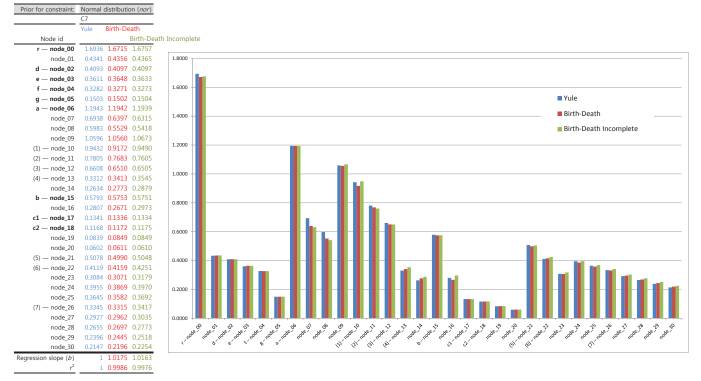
2.0

1.5

1.0

0.5





Supplementary Table S2. Time estimations comparison resulted in the BEAST analyses under different priors, i) Yule Process, ii) Birth-Death and iii) Birth-Death Incomplete process.

Table S3. Species examined in the present study. The sequences determined in this study are shown in bold text. Taxa name used in tree highlighted.

Taxa, collection deta	ils, and voucher m	umber				
Plastid			Nuclear			Mitochondria
psaA	psbA	rbcL	EF-2	SSU	LSU	cox1
RHODOPHYTA						
BANGIOPHYCEAE						
Bangiales Nägeli						
Bangia atropurpurea	(Roth) C. Agardh					
AY119698	AY119734	AY119770	EF033517	D88387	AF419107	DQ442887
Porphyra purpurea (R	toth) C. Agardh					
NC_000925	NC_000925	NC_000925	EF033519	L26201	EF033596	NC_002007
COMPSOPOGONOPHYCI	EAE					
Compsopogonales Skuja						
	eus (Balbis ex C. Agardh					
AY119701	AY119737	AF087116	-	AF342748	-	-
Rhodochaetales Bessey						
Rhodochaete parvula	Thuret					
AY119707	AY119743	AY119777	-	AF139462	FJ973376	-
CYANIDIOPHYCEAE						
Cyanidiales T. Christensen						
Cyanidioschyzon mere	elae Luca, Taddei et Var	ano				
AY119693	AY119693	AY119765	-	AB158485	AB158485	NC_000887
Cyanidium caldarium	(Tilden) Geitler					
NC_001840	NC_001840	NC_001840	-	AB090833	-	-
Cyanidium sp. Sybil C						
AY391363	AY391366	AY391369	-	-	-	-
Galdieria sulphuraria	(Galdieri) Merola					
AY119695	AY119731	AY119767	-	AB091230	-	-
PORPHYRIDIOPHYCEAE	3					
Porphyridiales Kylin ex Sku						
Flintiella sanguinaria	Ott					
AY119704	AY119740	AY119774	-	AY342749	-	-
Porphyridium aerugin	eum Geitler					
AY119705	AY119741	AY119775	-	AF168623	-	-

Pophynidium cruentan (S.F.Gray) Nägeli EST_contig1070 EST_contig13758 - EST_contig2709 RHODELLOPHYCEAE - Bitanicharya -<	D 1: 1:	(C E Carro) Në zali					
RHODELLOPHYCEAE Rhodellades H.S. Yoon, K.M Willer, R.G. sheath, F.D. Ott et D. Bhattacharya Disoniella grisse (Getifer) Scott, Broadwater, Saunders, Thomas et Gabrielson AY119702 AY119703 AY119703 AY119773 AY119773 AY119773 AY119776 AB045581 AB045581 AB045581 AB045581 AB045581			EST	ECT			EET
Rhodellales H.S. Yoon, K.M. Müller, R.G. sheath, F.D. Ott et D. Bhattacharya Dixoniella grisei (Geitter) Scott, Broadwater, Saunders, Thomas et Gabrielson AY119730 AY119773 AY119775 AY119776 - EU861395 - - AY119706 AY119774 AY119778 - AF342750 - - - Rhodsors marinus Geitler -		ESI_contig5624	ESI_contig10/0	ESI_contig13/58	-	-	ESI_contig2/09
Dissniella grisea (Geitler) Scott, Broadwater, Saunders, Thomas et Gabrielson API 19702 API 19738 API 19773 AB045581 - - Rhodella violacea (Kommann) Wehrneyer - EU861395 -							
AY119702 AY119738 AY119773 - AB045581 - - Rhodella violacea (Kommann) Wehrmeyer - - EU861395 - - AY119706 AY119742 AY119776 - EU861395 - - Stylonematales Drew - <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>							
Rhodella violacea (Kommann) Wehrmeyer							
AY119706 AY119742 AY119776 EU861395 - - STYLONEMATOPHYCEAE -			AY119773	-	AB045581	-	-
STYLONEMATOPHYCEAE Stylonematales Drew							
Stylonematales Drew Rhodosorus marinus -			AY119776	-	EU861395	-	-
Rhodosorus marinus Geitler AY119708 AY119744 AY119778 - AF342750 - - Stylonema alsidii (Zanardini) Drw -<							
AY119708AY119744AY119778AF342750Skylonema alsidii (Zanardini) Drew							
Stylonema alsidii (Zanardini) Drew AY119709 AY119745 AY119779 AF168633 - - AY119709 AY119745 AY119779 - AF168633 -							
AY119709AY119745AY119779-AF168633FLORIDEOPHYCEAE <td< td=""><td></td><td></td><td>AY119778</td><td>-</td><td>AF342750</td><td>-</td><td>-</td></td<>			AY119778	-	AF342750	-	-
FLORIDEOPHYCEAE Ahnfeltiophycidae Ahnfeltia fastigiata (Linnaeus) J.V. Lamouroux; US: Alaska: Sitka: Kruzof Island (13 vii 2006); A313 FJ195596 FJ195601 FJ195605 DQ343668 AF419104 GQ497301 Ahnfeltia fastigiata (Linnaeus) J.V. Lamouroux; US: Alaska: Sitka: Kruzof Island (13 vii 2006); A313 GQ497301 Ahnfeltia plicata (Hudson) Fries DQ343668 AF419104 GQ497301 Corallinophycidae EF033537 Z14139 AF419105 - Corallinales P.C. Silva et H.W. Johansen (1986, p.250) emendavit W.A. Nelson, J.E. Sutherl., T.J. Farr et H.S. Yoon Amphiroa fragilissima (Linnaeus) J.V. Lamouroux -							
Ahnfeltiophycidae Ahnfeltiales C.A. Maggs et C.M. Pueschel Ahnfeltia fastigiata (Linnaeus) J.V. Lamouroux; US: Alaska: Sitka: Kruzof Island (13 vii 2006); A313 FJ195596 FJ195601 FJ195605 FJ195605 Ahnfeltia plicata (Hudson) Fries EF033537 Z14139 AF419105 - Corallinales P.C. Silva et H.W. Johansen (1986, p.250) emendavit W.A. Nelson, J.E. Sutherl., T.J. Farr et H.S. Yoon Amphiroa fragilissima (Linnaeus) J.V. Lamouroux U04039 EF033529 U60744 EF033599 - Corallina spp. (C. officinalis + C. pilulifera) Corallina officinalis Linnaeus		AY119745	AY119779	-	AF168633	-	-
Ahnfeltiales C.A. Maggs et C.M. Pueschel Ahnfeltiales C.A. Maggs et C.M. Pueschel Ahnfeltia fastigiata (Linnaeus) J.V. Lamouroux; US: Alaska: Sitka: Kruzof Island (13 vii 2006); A313 FJ195596 FJ195601 FJ195605 - DQ343668 AF419104 GQ497301 Ahnfeltia plicata (Hudson) Fries - EF033537 Z14139 AF419105 - Corallinophycidae - EF033529 Voon - - Corallinales P.C. Silva et H.W. Johansen (1986, p.250) emendavit W.A. Nelson, J.E. Sutherl., T.J. Farr et H.S. Yoon - - - Amphiroa fragilissima (Linnaeus) J.V. Lamouroux -	FLORIDEOPHYCEAE						
Ahnfeltia fastigiata (Linnaeus) J.V. Lamouroux; US: Alaska: Sitka: Kruzof Island (13 vii 2006); A313 FJ195596 FJ195601 FJ195605 - DQ343668 AF419104 GQ497301 Ahnfeltia plicata (Hudson) Fries - - EF033537 Z14139 AF419105 - Corallinophycidae - - EF033537 Z14139 AF419105 - Corallinales P.C. Silva et H.W. Johansen (1986, p.250) emendavit W.A. Nelson, J.E. Sutherl., T.J. Farr et H.S. Yoon Amphiroa fragilissima EF033599 U60744 EF033599 -							
Ahnfeltia fastigiata (Linnaeus) J.V. Lamouroux; US: Alaska: Sitka: Kruzof Island (13 vii 2006); A313 FJ195596 FJ195601 FJ195605 - DQ343668 AF419104 GQ497301 Ahnfeltia plicata (Hudson) Fries - - EF033537 Z14139 AF419105 - Corallinophycidae - - EF033527 Z14139 AF419105 - Corallinales P.C. Silva et H.W. Johansen (1986, p.250) emendavit W.A. Nelson, J.E. Sutherl., T.J. Farr et H.S. Yoon Af419105 - - Amphiroa fragilissima (Linnaeus) J.V. Lamouroux EF033529 U60744 EF033599 -	Ahnfeltiales C.A. Maggs et C.	M. Pueschel					
Ahnfeltia plicata (Hudson) Fries EF033537 Z14139 AF419105 - Corallinophycidae EF033537 Z14139 AF419105 - Corallinales P.C. Silva et H.W. Johansen (1986, p.250) emendavit W.A. Nelson, J.E. Sutherl., T.J. Farr et H.S. Yoon - - - Amphiroa fragilissima (Linnaeus) J.V. Lamouroux - - U04039 EF033529 U60744 EF033599 - Corallina spp. (C. officinalis + C. pilulifera) - - EF033530 L26184 AF419116 FM180081 Corallina officinalis Linnaeus - - EF033530 L26184 AF419116 FM180081 Corallina pilulifera Postel et Ruprecht; Japan: Chiba: Choshi (01 viii 2004); C1003 -	Ahnfeltia fastigiata (Lini	naeus) J.V. Lamouroux;	US: Alaska: Sitka: Kruzof	Island (13 vii 2006); A313			
CorallinophycidaeEF033537Z14139AF419105-Corallinales P.C. Silva et H.W. Johansen (1986, p.250) emendavit W.A. Nelson, J.E. Sutherl., T.J. Farr et H.S. Yoon-Amphiroa fragilissima(Linnaeus) J.V. Lamouroux-U04039EF033529U60744EF033599Corallina spp. (C. officinalis + C. pilulifera)Corallina officinalis Linnaeus-EF033530L26184AF419116FM180081Corallina pilulifera Postel et Ruprecht; Japan: Chiba: Choshi (01 viii 2004); C1003DQ787594DQ787538Mastophora rosea(C. Agardh) Setchell; USA: Guam: Pago Bay; PAGO0003B; RedToL #538KP224295KP224296KP224298Iania sagiitata (Lamouroux) Blainville; Australia: Tasmania: Stanley Breakwater; GWS016470; RedToL #536KP224288KC130175KC157580KC157591KP224299Pseudolithophyllum muricatum (Fosile) Steneck et R.T. PaineAY294373	FJ195596	FJ195601	FJ195605	-	DQ343668	AF419104	GQ497301
Corallinophycidae Corallinales P.C. Silva et H.W. Johansen (1986, p.250) emendavit W.A. Nelson, J.E. Sutherl., T.J. Farr et H.S. Yoon Amphiroa fragilissima (Linnaeus) J.V. Lamouroux	Ahnfeltia plicata (Hudso	n) Fries					
Corallinales P.C. Silva et H.W. Johansen (1986, p.250) emendavit W.A. Nelson, J.E. Sutherl., T.J. Farr et H.S. Yoon Amphiroa fragilissima (Linnaeus) J.V. Lamouroux EF033529 U60744 EF033599 - Corallina spp. (C. officinalis + C. pilulifera) Corallina officinalis Linnaeus -	-	-	-	EF033537	Z14139	AF419105	-
Amphiroa fragilissima (Linnaeus) J.V. Lamouroux EF033529 U60744 EF033599 - Corallina spp. (C. officinalis + C. pilulifera) EF033529 U60744 EF033599 - Corallina spp. (C. officinalis + C. pilulifera) EF033529 U60744 EF033599 - Corallina officinalis Linnaeus EF033530 L26184 AF419116 FM180081 Corallina pilulifera Postel et Ruprecht; Japan: Chiba: Choshi (01 viii 2004); C1003 - - - DQ787594 DQ787634 DQ787588 - - - Mastophora rosea (C. Agardh) Setchell; USA: Guam: Pago Bay; PAGO0003B; RedToL #538 KP224281 KP224287 KP224278 KP224295 KP224296 KP224298 Jania sagittata (Lamouroux) Blainville; Australia: Tasmania: Stanley Breakwater; GWS016470; RedToL #536 KP224288 KC134331 KC130175 KC157580 KC157591 KP224299 Pseudolithophyllum muricatum (Fosile) Steneck et R. T. Paine - - - - Corollitophyllum muricatum (Fosile) Steneck et R. T. Paine - - - -	Corallinophycidae				•		
Image: Construction of the second system of the s	Corallinales P.C. Silva et H.W	. Johansen (1986, p.250) emendavit W.A. Nelson,	J.E. Sutherl., T.J. Farr et H.	S. Yoon		
Corallina spp. (C. officinalis + C. pilulifera) Corallina officinalis Linnaeus Corallina officinalis Linnaeus Corallina pilulifera Postel et Ruprecht; Japan: Chiba: Choshi (01 viii 2004); C1003 DQ787594 DQ787534 DQ787594 DQ787558 Mastophora rosea C. Agardh) Setchell; USA: Guam: Pago Bay; PAGO0003B; RedToL #538 KP224281 KP224287 KP224281 KP224287 KP224282 KP224288 KP224282 KP224288 KC130175 KC157580 KC157591 KP224299 Pseudolithophyllum muricatum (Foslie) Steneck et R. T. Paine AY294373 -	Amphiroa fragilissima (I	Linnaeus) J.V. Lamouro	ux				
Corallina officinalis Linnaeus EF033530 L26184 AF419116 FM180081 Corallina pilulifera Postel et Ruprecht; Japan: Chiba: Choshi (01 viii 2004); C1003 EF033530 L26184 AF419116 FM180081 DQ787594 DQ787634 DQ787558 - - - - Mastophora rosea (C. Agardh) Setchell; USA: Guam: Pago Bay; PAGO0003B; RedToL #538 KP224281 KP224287 KP224278 KP224279 KP224295 KP224296 KP224298 Jania sagiitata (Lamouroux) Blainville; Australia: Tasmania: Stanley Breakwater; GWS016470; RedToL #536 KP224282 KP224288 KC130175 KC157580 KC157591 KP224299 Pseudolithophyllum muricatum (Foslie) Steneck et R.T. Paine - - - -	-	-	U04039	EF033529	U60744	EF033599	-
Corallina officinalis Linnaeus EF033530 L26184 AF419116 FM180081 Corallina pilulifera Postel et Ruprecht; Japan: Choshi (01 viii 2004); C1003 L26184 AF419116 FM180081 DQ787594 DQ787634 DQ787558 - - - Mastophora rosea (C. Agardh) Setchell; USA: Guam: Pago Bay; PAGO0003B; RedToL #538 KP224281 KP224287 KP224278 KP224279 KP224295 KP224296 KP224298 Jania sagittata (Lamouroux) Blainville; Australia: Tasmania: Stanley Breakwater; GWS016470; RedToL #536 KC134331 KC130175 KC157580 KC157591 KP224299 Pseudolithophyllum muricatum (Foslie) Steneck et R. T. Paine AY294373 - - -	Corallina spp. (C. officin	alis + C. pilulifera)			•		
EF033530 L26184 AF419116 FM180081 Corallina pilulifera Postel et Ruprecht; Japan: Chiba: Choshi (01 viii 2004); C1003 DQ787594 DQ787634 DQ787558 -							
Corallina pilulifera Postel et Ruprecht; Japan: Chiba: Choshi (01 viii 2004); C1003 DQ787594 DQ787634 DQ787558 -	-	-	-	EF033530	L26184	AF419116	FM180081
DQ787594 DQ787634 DQ787558 -	Corallina nilulifera Post	el et Ruprecht: Japan: C	hiba: Choshi (01 viji 2004)	· C1003			
Mastophora rosea KP224281 KP224287 KP224278 KP224279 KP224295 KP224296 KP224298 Jania sagiitata (Lamouroux) Blainville; Australia: Tasmania: Stanley Breakwater; GWS016470; RedToL #536 KC124298 KC134331 KC130175 KC157580 KC157591 KP224299 Pseudolithophyllum muricatum (Foslie) Steneck et R.T. Paine AY294373 C C C				-	-	-	-
KP224281 KP224287 KP224278 KP224279 KP224295 KP224296 KP224298 Jania sagittata (Lamouroux) Blainville; Australia: Tasmania: Stanley Breakwater; GWS016470; RedToL #536 #		*****		3B. RedToL #538			
Jania sagittata (Lamouroux) Blainville; Australia: Tasmania: Stanley Breakwater; GWS016470; RedToL #536 KP224282 KP224288 KC134331 KC130175 KC157580 KC157591 KP224299 Pseudolithophyllum muricatum (Foslie) Steneck et R.T. Paine - <td></td> <td>·····</td> <td></td> <td></td> <td>KD224205</td> <td>KD224206</td> <td>1/1221/208</td>		·····			KD224205	KD224206	1/1221/208
KP224282 KP224288 KC134331 KC130175 KC157580 KC157591 KP224299 Pseudolithophyllum muricatum (Foslie) Steneck et R.T. Paine - <						NF 224270	NF 224270
Pseudolithophyllum muricatum (Foslie) Steneck et R.T. Paine						KC157501	KD224200
AY294373				KC130173	NC13/300	KU13/391	NF 224299
	rseuaounophyllum mur	icatum (Poslie) Steneck					
Hapanoiales w.A. Neison, J.E. Sutheri., I.J. Farr et H.S. Yoon	-	-		-	-	-	
	Hapalidiales W.A. Nelson, J.E	. Sutherl., T.J. Farr et H	.S. Yoon	-		-	-

Mesophyllum vancouv		t R.T. Paine; Canada: British C			140150500	17 544 4200
	KP224289	KC134326	KC130171	KC157577	KC157589	KP224300
Lithothamnion glacial		foundland and Labrador: Mae		RedToL #545		
-	KP224290	KC134336	KC130177	-	KC157593	HM918805
) R.A. Townsend; Australia: V		······		
KP224283	KP224291	KC134328	EF033531	KC157578	EF033600	KP224302
hodogorgonales S. Freder		schel				
<i>Renouxia antillana</i> Fr	edericq et J.N. Norris					
-	-	U04181	-	-	-	
-	-	-	EF033534	EF033584	EF033601	GQ497316
Rhodogorgon ramosis	sima J.N. Norris et Buche					
-	-	U04183	EF033535	AF006089	EF033602	-
oorolithales L. Le Gall et (
Heydrichia homalopas	ta R.A. Townsend et Bor	owitzka				
-	DQ167931	-	-	AF411629	-	-
Sporolithon durum (Fo	oslie) R.A. Townsend et W	Voelkerling				
-	KC963421	-	-	AF411626	-	NC_023454
Sporolithon ptychoides	Heydrich					
-	KC870926	-	GQ149067	GQ149066	GQ149068	HQ422711
ildenbrandiophycidae						
ildenbrandiales Pueschel	et Cole				-	-
Hildenbrandia rubra (Sommerfelt) Meneghini;	CCAP 1368/2				
DQ787593	DQ787633	DQ787557/AF208801	EF033522	AF076995	AF419136	GQ497309
emaliophycidae						
crochaetiales Feldmann						
	savianum + A. secundati					
Acrochaetium savianu	m (Meneghini) Nägeli; C	CAP 1350/2	•			
DQ787597	DQ787637	DQ787561	-	-	-	-
Acrochaetium secundo	utum (Lyngbye) Nägeli		-		-	-
-	-	-	EF033523	AF079784	AF528044	-
Acrochaetium sp.	-		-			-
-	-	-	-	-	-	GQ497300
albianiales R.G. Sheath et	K.M. Müller		-			
Balbiania investiens (Lenormand ex Kützing)		•			-
-	-	AF132293	EF033524	AF132294	AF421124	-
alliales HG. Choi, Kraft	et G.W. Saunders		-			-
		aland: Marlborough: Kaikoura	a (20 viii 2002); B10			-
	DQ787635	DO787559	EF033525	AF236790	AF419106	

Batrachospermales Puescl						
	e spp. (B. gelatinosum + H					
	elatinosum (Linnaeus) De	e Candolle; UTEX LB149.	3			
DQ787596	DQ787636	DQ787560	-	AF026045	DQ903115	EU636744
Petrohua bernabei (G.W. Saunders					
-	-	-	EF033526	-	-	-
Colaconematales J.T. Har						
				Lonsdale Lighthouse Reef;		
KP224284	KP224292	KC134354	EF033528	AF079787	AF528046	KP224303
				i; GWSC008; RedToL #559		
KP224285	KP224293	KC134327	KC130172	AF079791	AF528049	KF364496
Nemaliales Schmitz						
	elminthoides + Nemalion					
Nemalion helmintho	ides (Velley in Withering	g) Batters				
-	-	-	EF033532	L26196	AY570376	-
	nd: Devon: Wembury Be					
DQ787598	DQ787638	DQ787562	-	-	-	-
Nemalion sp.						_
-	-	-	-	-	-	GQ497310
Palmariales Guiry et D. Ir						
		ern Ireland: Noyle: Garron				
DQ787599	DQ787639	DQ787563	EF033533	Z14142	Y11506	GQ497313
		cott, K.A. West, U. Karste			•	
				aunders; Madagascar: Ifaty;		
KP224286	KP224294	KC134337	EU262262	EU262260	EU262261	KP233841
Thoreales Müller, Sheath,			-			_
	acea + Thoreaceae sp.)					
Thorea violacea Bo						
AY119712	AY119747	AF029160	_	-	-	-
Thoreaceae sp.						
-	-	-	EF033536	AF420253	AF419145	-
Rhodymeniophycidae						
Acrosymphytales R.D. W			.			
	baeum (J. Agardh) G. Sjo	·····				
KM359975	KM359999	KM360022	EF033539	DQ343661	DQ343684	-
	<i>housboei</i> (J. Agardh) J. A					
KM359976	KM360000	KM360023	EF033540	AY437681	AF419130	-
Bonnemaisoniales Feldma	ann et Feldmann					

FJ195594	FJ195600	FJ195604	AY010232	L26182	AF419112	-
Delisea hypneoides H	Harvey					
KM359977	-	KM360024	EF033541	EF033585	EF033603	-
Naccariaceae spp. (A	accaria wiggii + Reticulo	caulis mucosissimus)				
Naccaria wiggii (Tu	mer) Endlicher					
-	-	-	-	-	-	GQ497312
Reticulocaulis mucos	sissimus Abbott					
KM359978	KM360001	KM360025	EF033542	DQ343656	DQ343680	-
miales Oltmanns						
	C. collabens + C. onsangi					
Callithamnion collab	ens (Rudolphi) McIvor et	Maggs				
-	-	-	-	DQ022769	DQ022974	-
			ellington: Breaker Bay (11 i	x 2002); C309		
DQ787602	DQ787642	DQ787565	-	-	-	EU194962
Seirospora interrupte						
DQ787615	DQ787649	DQ110903	-	DQ022774	DQ022799	EU194970
	um (C. Agardh) Montagn					
AY295137	AY178488	AY295175	EF033543	DQ343657	AF419113	EU194971
	ecundatum + C. virgatum					
	um Lyngbye; France: Roso					
DQ787605	DQ787644	DQ110904	_	-	-	EU194972
Ceramium virgatum	Roth		_			
-	-	-	EF033544	AF236793	EF033604	-
	orallinoides + G. okiensis)	-			
	des (Linnaeus) Trevisan					
AY295126	AY295146	AY295164	_	EU718690	-	-
Griffithsia okiensis 🛛	Cajimura		-			
-	-	-	-	-	-	EU194973
Ptilota spp. (P. gunn					-	
			e: Cushendun (24 iiv 2003)			
DQ787613	AY865154	DQ787575	-	EU718700	-	EU194975
Ptilota serrata Kützi	ng					
-	-	-	EF033545	-	EF033605	-
		Pohang: Wolpo (16 xi 200	02); S16			
DQ787618	DQ787652 collabens + H. plumosa)	DQ787579	-	EU718707	AF458717	AF458719

DQ787619	DQ787653	DQ787580	-	AF488384	-	-
Heterosiphonia plumo	osa (Ellis) Batters					
-	-	-	EF033546	-	EF033606	
Dasysiphonia spp. (D	. chejuensis + D. okiensi	is)				
Dasysiphonia okiensi:	s Kajimura; Japan: Oki I	sland: Kamo Bay (06 v 200	03); D56			
DQ787620	DQ787654	DQ787581	-	-	-	GQ497304
Dasysiphonia chejuen	sis I.K. Lee et W.A. We	st				
-	-	-	-	AF488388	-	-
Delesseriaceae spp. 1	(Delesseria serrulata +	Grinnellia americana)				
Delesseria serrulata I	Harvey; Korea: Gangreu	ng: Anin (30 i 2002); D19				
DQ787621	DQ787655	DQ787582	-	AF488403	-	-
Grinnellia americana	(C. Agardh) Harvey					
-	-	-	EF033547	-	EF033607	-
Delesseriaceae spp. 2	(Phycodrys rubens + Se	orella repens + Hymenena	sp.)			
Phycodrys rubens (Li	nnaeus) Batters; Norway	: Møre og Romsdal: Ona I	sland (28 vii 2004); P422			
DQ787622	DQ787656	DQ787583	-	-	-	-
Sorella repens (Okam	ura) Hollenberg	-				-
-	-	-	EF033548	AF488406	EF033608	-
Hymenena sp.						
-	-	-	-	-	-	GQ497305
Brongniartella byssoi	des (Goodenough et Wo	odward) F. Schmitz: Norw	ay: Møre og Romsdal: Pinnø	oya (29 vii 2004); B19		
DQ787623	DQ787657	DQ787584	-	-	-	-
Chondrophycus interr	<i>nedius</i> (Yamada) Garba	ry et Harper; Korea: Gangv	von: Goseong: Ayajin (30 vi	i 2002); L2		
DQ787624	DQ787658	DQ787585	-	-	-	-
Polysihponia spp. (P.	stricta + Polysiphonia s	p.)				
Polysiphonia stricta (Dillwyn) Greville; Norw	vay; Møre og Romsdal: Pin	nøya (29 vii 2004); P421		•	
DQ787625	DQ787659	AY958166	-	AF427535	-	-
Polysiphonia sp.	_	_				
-	-	-	-	-	-	GU385828
idiales Kylin		-				
	gans + G. australe + G. j					-
		ouroux; Korea: Jejudo: Seo	ongsan (23 vii 2002); G2			-
DQ787626	DQ787660	DQ787586	-	-	-	-
Gelidium australe J. A	Agardh					
-	-	-	EF033549	DQ343660	DQ343682	-
Gelidium purpurascer	s N L. Gardner				-	
$r \cdots r$	is rulli ourditer					
- artinales Schmitz	-	-	-	-	-	GQ497307

Ahnfeltiopsis spp. (A. p						
		pan: Chiba: choshi (27 vii 2	002); A12			
DQ787627	DQ787661	DQ787587	-	-	-	-
Ahnfeltiopsis linearis (C. Agardh) P.C. Silva e	t DeCew (GWS002887)				
-	-	-	-	-	GQ338104	GQ380028
Chondrus crispus Stac	khouse; GWS009227					
KM359979	KM360002	KM360026	EF033554	Z14140	AF419120	NC_001677
Euthora cristata (C. A	gardh) J. Agardh; GWS	000026 (G0433)				
KM359980	KM360003	KM360027	EF033555	AY437684	AF419124	GU140142
Gloiopeltis furcata (Po	ostels et Ruprecht) J.G. A	Agardh; GWS002264				
KM359981	KM360004	KM360028	EF033553	U33130	EF033612	-
Hypnea spp. (H. japon	ica + H. charoides)					
Hypnea japonica Tana	ka; Korea: Ulreungdo: '	Гоnggumi (27 viii 2003); Н	41			
FJ195597	DQ095851	DA095829	-	-	-	EU345986
Hypnea charoides J.V.	. Lamouroux					
-	-	-	-	AY437682	-	-
Mychodea acanthymer	nia Kraft; GWS000960	•				
KM359982	KM360005	KM360029	EF033557	EF033589	EF033614	-
Neodilsea borealis (Al	bott) Lindstrom; G022	3				
KM359983	KM360006	KM360030	EF033550	AF317112	EF033610	EU189312
Weeksia coccinea (Ha	rvey) Lindstrom; GWS0	01705				
KM359984	KM360007	KM360031	EF033552	AF317120	EF033611	EU189325
racilariales S. Fredericq et	M.H. Hommersand				-	
Gracilaria tenuistipita	ta var. liui Zhang et Xia	L				
AY673996	AY673996	AY673996		D001 (005	······	
Gracilaria spp. (G. tex		A10/3//0	-	DQ316995	-	EF434924
	ctorii + G. salicornia)	A10/3//0	-	DQ316995	-	EF434924
Gracilaria textorii De		n: Daecheon (17 i 2000); G4	-	DQ316995	-	EF434924
Gracilaria textorii De FJ195595			-	-	-	EF434924 EF434925
FJ195595	Toni; Korea: Chungnan DQ095832	n: Daecheon (17 i 2000); G4 DQ095793			-	
FJ195595	Toni; Korea: Chungnan	n: Daecheon (17 i 2000); G4 DQ095793	- 45 - EF033559		- - EF033615	
FJ195595 Gracilaria salicornia (-	Toni; Korea: Chungnan DQ095832	1: Daecheon (17 i 2000); G- DQ095793 m	-	-	- - EF033615	EF434925
FJ195595 Gracilaria salicornia (- Gracilariopsis spp. (G	Toni; Korea: Chungnan DQ095832 (C. Agardh) E.Y. Dawso - - chorda + G. andersona	1: Daecheon (17 i 2000); G- DQ095793 m	- EF033559	-	- - EF033615	EF434925
FJ195595 Gracilaria salicornia (- Gracilariopsis spp. (G	Toni; Korea: Chungnan DQ095832 (C. Agardh) E.Y. Dawso - - chorda + G. andersona	n: Daecheon (17 i 2000); G- DQ095793 on - ii + G. lemaneiformis)	- EF033559	-	- - EF033615	EF434925
FJ195595 Gracilaria salicornia (- Gracilariopsis spp. (G Gracilariopsis chorda	Toni; Korea: Chungnan DQ095832 (C. Agardh) E.Y. Dawse - - . chorda + G. andersoni (Holmes) Ohmi; Korea DQ095825	n: Daecheon (17 i 2000); G- DQ095793 on <i>i + G. lemaneiformis</i>) Jindo: Hoidong (09 iii 200 DQ095785	- EF033559	-	- - EF033615 -	EF434925
FJ195595 Gracilaria salicornia (- Gracilariopsis spp. (G Gracilariopsis chorda	Toni; Korea: Chungnan DQ095832 (C. Agardh) E.Y. Dawso - - - chorda + G. andersoni (Holmes) Ohmi; Korea:	n: Daecheon (17 i 2000); G- DQ095793 on <i>i + G. lemaneiformis</i>) Jindo: Hoidong (09 iii 200 DQ095785	- EF033559	-	- EF033615 -	EF434925
FJ195595 Gracilaria salicornia (Gracilariopsis spp. (G Gracilariopsis chorda Gracilariopsis anderse	Toni; Korea: Chungnan DQ095832 (C. Agardh) E.Y. Dawse chorda + G. andersoni (Holmes) Ohmi; Korea: DQ095825 onii (Grunow) Dawson;	n: Daecheon (17 i 2000); G- DQ095793 on <i>i + G. lemaneiformis</i>) Jindo: Hoidong (09 iii 200 DQ095785	- EF033559 1); G39 - EF033560	-	- EF033615 - -	EF434925
FJ195595 Gracilaria salicornia (Gracilariopsis spp. (G Gracilariopsis chorda - Gracilariopsis anderse	Toni; Korea: Chungnan DQ095832 (C. Agardh) E.Y. Dawse chorda + G. andersoni (Holmes) Ohmi; Korea: DQ095825 onii (Grunow) Dawson;	n: Daecheon (17 i 2000); G- DQ095793 on <i>i + G. lemaneiformis</i>) Jindo: Hoidong (09 iii 200 DQ095785 GWS002277	- EF033559 1); G39 - EF033560	-	- EF033615 - - AF419132	EF434925

Grateloupia spp. (G. div	varicata + G. subpecti	nata + Grateloupia sp.)				
Grateloupia divaricata	Okamura; Korea: Tae	an: Hakampo (11 xi 2001); (G50			
DQ787630	DQ787664	DQ787590	-	-	-	-
Grateloupia subpectinat	ta Holmes					
-	-	-	-	U33132	-	-
Grateloupia sp. GWS00	3281					
	-	-	-	-	-	GQ497308
Epiphloea bullosa (Harv						
KM359985	KM360008	KM360032	EF033562	AY437701	DQ343693	-
Isabbottia ovaliforlia (K						
KM359986	KM360009	KM360033	EF033563	EF033590	EF033616	-
Halymeniaceae spp. (P.						
Pachymenia carnosa (J.						
KM359987	KM360010	KM360034	EF033564	AF515289	DQ343695	-
Halymenia pseudoflores	ii F.S. Collins et M.A	Howe; CL033701	-			
-	-	-	-	-	-	GQ862076
Zymurgia chondriopside			-			
KM359988	KM360011	KM360035	EF033566	AF515304	DQ343698	-
Nemastomatales Kylin						
Predaea kraftiana Milla		32				
KM359989	KM360012	-	EF033567	AF515297	EF033618	-
Platoma cyclocolpum (N						
KM359990	KM360013	KM360036	EF033568	AF515292	DQ343706	-
Schizymenia pacifica (K						
KM359991	KM360014	KM360037	EF033569	EF033592	AF419129	GQ497319
Peyssonneliales D.M. Krayes			.			
Ramicrusta textilis C.M						
KM359992	KM360015	KM360038	-	FJ848977	FJ848970	-
Sonderopelta coriacea						
KM359993	KM360016	KM360039	EF033558	AY437689	DQ343691	-
Plocamiales G.W. Saunders e						
Plocamium spp. (P. carl						
		kon; Korea: Pohang: Guryeo	ongpo (31 i 2002); P13			
DQ787629	DQ787663	DQ787589	-	-	-	
Plocamium maggsiae G	.W. Saunders et Lehr	ıkuhl				
-	-	-	EF033570	AY437708	AF419141	
Plocamium sp. LLG017	1					
-	-	-	-	-	-	GQ497314

Sarcodia ciliata Zanard	ini; GWS001027					
KM359994	KM360017	KM360040	EF033572	DQ343666	DQ343708	
Sarcodia sp. GWS0025	97					
-	-	-	-	-	-	FJ499623
odymeniales Schmitz						
Champia spp. (C. chath						
Champia chathamensis			ington: Lyall Bay (03 viii 2	001); C255		
FJ195598	FJ195602	FJ195606	-	-	-	-
Champia affinis (j.d. He	oker et Harvey) Harve	y				
-	-	-	-	U23951	-	-
Champia gigantea M.J.	Wynne					
-	-	-	EU624164	-	EU624159	-
Fryeella gardneri (Setc						
KM359996	KM360019	KM360042	EF033578	AF085273	EF033622	GQ497306
Gloioderma halymenio						
KM359995	KM360018	KM360041	EF033574	DQ873283	DQ873283	-
		Lomentaria sp. GWS00188				
		am: Sacheon (18 vii 2001)				
FJ195599	FJ105603	FJ195607	EU624178	-	EU624155	-
Lomentaria australis (H	lützing) Levring	-	-			
-	-	-	-	U33134	-	-
Lomentaria sp. GWS00	1885					
-	-	-	-	-	-	GQ497311
	Okamura) Okamura; K	orea: Ulreungdo: Dodong (26 viii 2003); R13			
DQ787631	DQ787665	DQ787591	EU624196	AB381929	EU624150	-
odeniales R.D. Withall et (
Lesleigha sp. 1LH; GW						
KM359997	KM360020	KM360043	EF033581	AY437707	DQ343700	-
Sebdenia spp. (S. flabel			-			
Sebdenia flabellata (J. 1	Agardh) P.G. Parkinsor	1				
	-	-	EF033579	U33138	AF419134	-
-	CWC002074					
- Sebdeniaceae sp. Unkn	5wn; Gw 5002074					

NC_004115	NC_004115	-	_	AJ250110	-	NP_689386
Mesostigma viride La						
NC_002186	NC_002186	-	EU812184/ EU812185	AJ250109	AY591912	NC_008240
CHLOROPHYTA						
TREBOUXIOPHYCEAE						
Chlorella vulgaris Be	eijerinck					
NC_001865	NC_001865	-	-	GQ122334	AB237642	AB011523
STREPTOPHYTA						
Arabidopsis thaliana	(L.) Heynhold					
NC_000932	NC_000932	-	AC009894	NR_022795	AC006837	NC_001284
Marchantia polymorp						
NC_001319	NC_001319	-	-	AY342318	AY342318	NC_001660
	rgii + P. wallichiana + P	. kwangtungensis)				
Pinus thunbergii Parl						
NC_001631	NC_001631	-	-	-	-	-
Pinus wallichiana A.	B. Jacks					
-	-	-	-	X75080	AJ271114	-
Pinus kwangtungensi.	s = Pinus fenzeliana Han	dMazz.				
-	-	-	_	_	-	EF114116
Psilotum nudum (L.)						
NC_003386	NC_003386	-	_	X81963	EU161326	EU161444
Zea mays L.	-					
NC_001666	NC_001666	-	EU970836	AF168884	AJ309824	AY506529
Outgroup						
CYANOBACTERIA						
CYANOPHYCEAE						
Nostoc sp. PPC 7120	-					
NC_003272	NC_003272	-	_	_	-	NC_003272
Synechocystis sp. PCO						
NC_000911	NC_000911	-	_	-	-	NC_000911
Thermosynechococcu			-			
NC_004113	NC_004113	-	-	-	-	NC_004113