

Mitochondrial DNA Part B



Resources

ISSN: (Print) 2380-2359 (Online) Journal homepage: https://www.tandfonline.com/loi/tmdn20

The complete plastid genome and phylogenetic analysis of *Gracilaria chilensis*

Guoliang Wang, Na Liu, Yue Li, Lei Zhang, Maria Dyah Nur Meinita, Weizhou Chen, Tao Liu & Shan Chi

To cite this article: Guoliang Wang, Na Liu, Yue Li, Lei Zhang, Maria Dyah Nur Meinita, Weizhou Chen, Tao Liu & Shan Chi (2020) The complete plastid genome and phylogenetic analysis of *Gracilaria chilensis*, Mitochondrial DNA Part B, 5:2, 1282-1283, DOI: 10.1080/23802359.2018.1431070

To link to this article: https://doi.org/10.1080/23802359.2018.1431070

9	© 2020 The Author(s). Published by Informa UK Limited, trading as Taylor & Francis Group.
	Published online: 02 Mar 2020.
	Submit your article to this journal 🗗
α	View related articles 🗗
CrossMark	View Crossmark data

Taylor & Francis Taylor & Francis Group

MITOGENOME ANNOUNCEMENT

OPEN ACCESS Check for updates



The complete plastid genome and phylogenetic analysis of *Gracilaria chilensis*

Guoliang Wang^{a,b,c*}, Na Liu^{a*}, Yue Li^a, Lei Zhang^a, Maria Dyah Nur Meinita^d, Weizhou Chen^e, Tao Liu^a and Shan Chi^{a,f}

^aCollege of Marine Life Sciences, Ocean University of China, Qingdao, China; ^bCAS Key Laboratory of Genome Sciences and Information, Beijing Institute of Genomics, Chinese Academy of Sciences, Beijing, China; University of Chinese Academy of Sciences, Beijing, China; ^dFaculty of Fisheries and Marine Science, Jenderal Soedirman University, Purwokerto, Indonesia; ^eMarine Biology Institute, Shantou University, Shantou, China; [†]Qingdao Haida BlueTek Biotechnology Co., Ltd, Qingdao, China

ABSTRACT

Gracilaria chilensis is an economically important species of macroalgae. The plastid genome sequence of G. chilensis is 185,640 bp with a GC content of 29.34%. A total of 236 genes were determined, containing 203 protein-encoding genes, three rRNA genes, 30 tRNA genes, and one intron (with intronic ORF) inserted into the trnM gene. The gene content and structure of Gracilariaceae species were relatively well conserved. The phylogenetic analysis, based on the red algal plastid genomes, suggested that G. chilensis had a closer relationship with Gracilaria tenuistipitata var. liui in Gracilaria.

ARTICLE HISTORY

Received 5 January 2018 Accepted 18 January 2018

KEYWORDS

Gracilaria chilensis; complete plastid genome; Gracilariaceae; phylogenetic analysis

Gracilaria chilensis is an economically important macroalga in the family Gracilariaceae. This family includes \sim 230 species in seven genera (Lyra et al. 2015). Currently, over 150 species of Gracilaria have been described. G. chilensis, distinguished by Bird et al. (1986), is a natural marine food with high fibre, high protein, low fat, low calories, and is rich in vitamins and minerals. Moreover, it is the most economically important alga, mainly used as a high-quality raw material for extracting agar and algin (Meinita et al. 2017). Besides, it is believed that G. chilensis can be used for bioremediation of ocean acidification, by absorbing excess CO2 from seawater (Gao and McKinley 1994). Both of these economic and environmental advantages can be achieved by integrating the cultivation of algae with fish farming.

In this study, we provide the red algal plastid genome by next-generation sequencing methods. The specimen was collected from Rongcheng, Shandong Province of China (37°09′52″N, 122°28′53″E), and stored in the Culture Collection of Seaweed at the Ocean University of China (sample accession number: 2016070050). Total DNA was extracted via the modified CTAB method (Doyle and Doyle 1990). The genome data of G. chilensis was obtained using an Illumina HiSeq system with paired-end, 150 bp sequencing. The complete plastid genome, using *Gracilaria salicornia* (NC 023785) as reference, was annotated with Geneious R10.1.3. The plastid genome map of G. chilensis was produced using OGDRAW software.

The complete plastid genome of G. chilensis was mapped as a circular molecule of 185,640 bp with 29.34% GC content

(GenBank accession number MF401963), including 203 protein-encoding genes, three rRNA genes, 30 tRNA genes, and one intron interrupting the trnM gene. The nucleotide composition was 35.53% A, 14.66% C, 14.69% G, and 35.13% T. The length of the coding region was 145,521 bp, corresponding to 78.4% of the total length. The plastid genome of G. chilensis was compact, with 10 pairs of overlapping genes found with overlap lengths of 1–95 bp (carA-ycf53, psbD-psbC, ycf29-trnH, trnT-ilvB, rpl24-rpl14, rpl14-rps17, rps17-rpl29, rpl23rpl4, rps18-rpl33, and atpF-atpD). Synteny analysis of species published in the NCBI sequence database revealed that the size of the plastid genomes of Gracilariaceae was 180-185 kb, with small differences in GC content (average GC content, 28.41%). The gene numbers and structures were largely similar among Gracilariaceae species; their plastid genomes were relatively well conserved, with no gene rearrangement phenomena.

Phylogenetic analysis was conducted using 85 shared plastid protein sequences from 17 red algal plastid genomes, publicly available from the GenBank dataset, and using Cyanidioschyzon merolae as an outgroup. Our results indicated that all red algal taxa were clearly separated into different groups, according to their original class (Figure 1). The Florideophyceae species formed a large branch, in which five species from the Gracilariales order were grouped as a subbranch, in which G. chilensis showed a closer relationship with Gracilaria tenuistipitata var. liui in Gracilaria. This study also facilitated a novel interpretation of the evolutionary

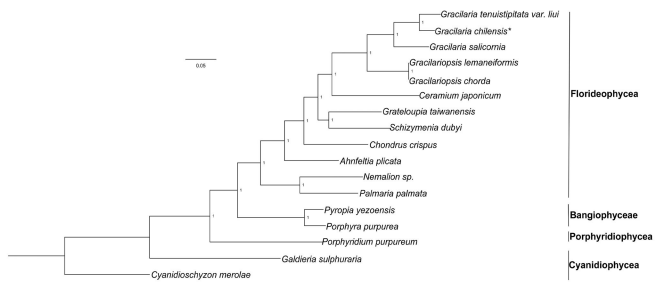


Figure 1. Phylogenetic analysis was performed using a Bayesian method, based on the plastid genomes of red algae as shown below: Cyanidioschyzon merolae (NC_004799), Galdieria sulphuraria (KJ700459), Porphyridium purpureum (NC_023133), Pyropia yezoensis (KC517072), Porphyra purpurea (U38804), Palmaria palmata (NC_031147), Nemalion sp. (LT622871), Ahnfeltia plicata (NC_031145), Chondrus crispus (NC_020795), Ceramium japonicum (NC_031174), Grateloupia taiwanensis (KC894740), Schizymenia dubyi (NC_031169), Gracilariopsis chorda (NC_031149), Gracilariopsis lemaneiformis (KP330491), Gracilaria salicornia (NC_023785), Gracilaria tenuistipitata var. liui (AY673996), and Gracilaria chilensis (MF401963). Asterisk indicates newly sequenced species in this study.

process from ancient unicellular algae to large multicellular complex algae.

Disclosure statement

No conflicts of interest for all the authors including the implementation of research experiments and writing this article were reported.

Funding

This work was supported by the China-ASEAN Maritime Cooperation Fund and the China Agriculture Research System-50.

References

Bird CJ, Mclachlan J, Oliveira ECD. 1986. Gracilaria chilensis sp.nov. (Rhodophyta, Gigartinales), from Pacific. Can J Bot. 64:2928-2934.

Doyle JJ, Doyle JL. 1990. Isolation of plant DNA from fresh tissue. Focus.

Gao K, Mckinley KR. 1994. Use of macroalgae for marine biomass production and CO₂, remediation: a review. J Appl Phycol. 6:45-60.

Lyra GM, Costa ED, de Jesus PB, de Matos JC, Caires TA, Oliveira MC, Oliveira EC, Xi ZX, de Nunes JM, Davis CC. 2015. Phylogeny of Gracilariaceae (Rhodophyta): evidence from plastid and mitochondrial nucleotide sequences. J Phycol. 51:356-366.

Meinita MDN, Marhaeni B, Hong YK, Jeong GT. 2017. Enzymatic saccharification of agar waste from Gracilaria verrucosa, and Gelidium latifolium, for bioethanol production. J Appl Phycol. 1-9.