Characterization of complete chloroplast genome of *Ulva torta* (Mertens) Trevisan, 1841

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**ABSTRACT**

*Ulva torta* (Mertens) Trevisan, 1841 was a global temperate widespread species. Green tide blooms caused by the green algae of the *Ulva* species occurred frequently in China. As a newly discovered species in the green tide bloom area, it was necessary to explore the relationship between *U. torta* and other green algae of the *Ulva* species. The complete chloroplast genome of *U. torta* was 105,423 bp in size. A total of 100 genes were annotated in the genome, containing 70 protein-coding genes, 27 transfer RNA (tRNA) genes, and three rRNA genes. The chloroplast genome had high AT content (74.76%). Phylogenetic analysis showed *U. torta* was clustered with *Ulva meridionalis*. This work could be useful for studying the evolution and genetic diversity of *U. torta*.

*U. torta* (gametophyte) was collected from the sea area of Rudong, Jiangsu, China in November, 2020 (32°25′52″N, 121°24′35″E). The specimen was deposited at the herbarium of Shanghai Ocean University Museum (collected by Qinlin Wen, wenq587@163.com) under the voucher number SHOU2020RDA21121. The specimen was sent to Sangon Biotech (Shanghai) Co., Ltd. for DNA extracting and high-throughput sequencing. DNA was extracted from this sample using the company’s Dzup (Plant) Genomic DNA Isolation Reagent. TruSeq DNA Sample Prep Kit was used to prepare genomic shotgun library (Illumina, USA), and then paired-end sequences were obtained by Illumina HiSeq 2500 platform. We obtained 10,626,405 raw read pairs and 1.59 Gbp data, with a single read length of 150 bp.

The chloroplast genome of *Ulva compressa* (MT916929) (Xia et al. 2021), *Ulva meridionalis* (MN889540) (Liu et al. 2020), *U. flexuosa* (KX579943) (Cai et al. 2017), *Ulva prolifera* (KX342867) (Jiang et al. 2019) and *U. linza* (KX058323) (Wang et al. 2017) had been studied by our laboratory before. Chloroplast genome of *U. prolifera* was taken as seed sequences for sequence splicing about the complete chloroplast genome of *U. torta* using NOVOPlasty software (Sedanza et al. 2020). The extended chloroplast genome contigs was obtained, and then the extended contigs were compared with the chloroplast genome sequence of *U. prolifera* to obtain the correct contig sequence. Then the complete chloroplast genome sequence was perfectly assembled.

Complete chloroplast genome of *U. torta* was 105,423 bp in size (GenBank accession number MZ703011). The chloroplast genome composition was biased toward AT content, at 74.76%, which was significantly higher than the GC content. When we annotated the whole chloroplast genome, *Ulva ohnoi* was mainly used for reference. A total of 100 functional genes were encoded in the genome, including 70 protein-coding genes, three rRNA genes (rrn 16 s gene, rrn 23 s gene, rrn 5 s gene, respectively), and 27 tRNA genes. In order to verify the phylogenetic position of the newly obtained species within *Ulva* species and further clarify the evolutionary relationship, phylogenetic analysis was carried out with 12 species (*Pseudendoclonium akinetum* as an outgroup taxonomically belongs to Chlorophyta, Ulvophyceae, Ulvalales, Krommanniaceae, *Pseudendoclonium*). All chloroplast genomes sequences were aligned with the BioEdit sequence software (Hall 1999). Clustal X software was used to perform multiple

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alignment analysis on qualified sequences. A Maximum Likelihood (ML) tree was constructed by Mega 7.0 (Kumar et al. 2016), and the accuracy of the phylogenetic tree was verified by the Bootstrap test repeated 1000 times, omitting less than 50% of the phylogenetic tree branch values. The result showed Ulva torta was related to Ulva meridionalis (Figure 1).

In this study, we analyzed complete chloroplast genome of Ulva torta. Currently, Ulva macroalgal blooms occur frequently in the Southern Yellow Sea of China (Zhang et al. 2014; Zhang et al. 2017; Zhao et al. 2019; Liu et al. 2020; Xiao et al. 2020; Liu et al. 2021). This study about chloroplast genomes of Ulva species will be useful for studying their genetic diversity.

Acknowledgment
Thanks to Sangon Biotech (Shanghai) Co., Ltd. for DNA extracting and high-throughput sequence.

Ethics statement
Research materials (macroalgae) involved in this research were all carried out under the existing legal framework in China, and there was no ethics involved.

Authors’ contributions
Jianheng Zhang, Peimin He and Jinlin Liu were involved in the conception and design. Shuang Zhao and Song Gao were involved in the data analysis of this work. Qinlin Wen, Weiming Yang and Jingshi Li were involved in the material acquisition, data interpretation and paper drafting. All authors critically revised this paper for intellectual content and approved the final version to be published. All authors have agreed to be accountable for all aspects of the work.

Disclosure statement
No potential conflict of interest was reported by the author(s).

Figure 1. A ML phylogenetic tree for Ulva torta based on the whole chloroplast genomes of 10 closely related species (U. compressa is 94,226 bp (MT916929) (Xia et al. 2021), Ulva meridionalis is 88,653 bp (MN889540) (Liu et al. 2020), U. flexuosa is 89,414 bp (KX579943) (Cai et al. 2017), U. prolifera is 90,066 bp (KX342867) (Jiang et al. 2019), U. linza is 1,251 bp (XX058323) (Wang et al. 2017), U. ohnoi is 103,313 bp (AP018696) (Suzuki et al. 2018), Ulva sp. is 99,983 bp (KP720616) (Melton et al. 2015), Ulva fasciata is 96,005 bp (KT882614) (Melton and Lopez-Bautista 2016), Ulva lactuca is 95,997 bp (MH730972) (Hughey et al. 2019) and Ulva pertusa is 104,380 bp (MN853875) (Han et al. 2020)) and one outgroup (Pseudendoclonium akinetum is 195,867 bp (AY835431) (Pombert et al. 2006)).

Funding
This work was financially supported by National Key R&D Program of China [grant numbers 2016YFC1402103], Natural Science Foundation of Shanghai [grant numbers 21ZR1427400], Project of Key Laboratory of Marine Ecological Monitoring and Restoration Technologies, MNR [grant numbers 202008], Project of Technology Innovation Center for Land Spatial Eco-restoration in Metropolitan Area, MNR [grant numbers CXZX202006].

Data availability statement
The support genome sequences data of this study are openly available in GenBank of NCBI (https://www.ncbi.nlm.nih.gov/) under the accession no. MT970311. The associated BioProject, Bio-Sample, and SRA accession numbers are PRJNA768934, SAMN22561212, and SRR16629507, respectively.

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