

Article

The morphology, genetic diversity, and distribution of *Ulva meridionalis* in Chinese seas

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Abstract: Green tides originate from the rapid growth of green macroalgae and their large accumulation. In the past few decades, the severity and frequency of green tides have increased and the range of their geographical distribution has widened. In recent years, *Ulva meridionalis* Horimoto et Shimada, ~~sp. nov., a new *Ulva* species belonging to the Ulvaceae family,~~ has been reported in many countries. This species has stable morphological characteristics and its length can reach 3 m in indoor cultures. Its cells contain starch granules, and the sporangium and gametangium of each cell contain eight spores and 16 gametes, respectively, which confer a high proliferation potential. The phylogenetic tree constructed in this study showed that the Internal Transcribed Spacer sequence identified *U. meridionalis* with a high identification reliability, and the genetic relationship between *U. meridionalis* and *Ulva pertusa* in the ITS sequence was close. The haplotype network analysis clarified the relationship of the *U. meridionalis* samples collected from four different sea areas in China and indicated that they are closely related. Five haplotypes were identified: Hap_2 and Hap_1 were the most frequent, and they were also the haplotypes shared among the three groups. The degree of subspecies formation was not reached among these *U. meridionalis* samples collected from the Chinese sea. Up to 20 years ago, *U. meridionalis* had only been recorded in Japan. After 2007, it was found to be widely distributed in the United States, China, French New Caledonia, French Polynesia, and Australia, where it proliferated. It has spread as a new kind of green tide-forming macroalga. The present study found that *U. meridionalis* is widely distributed in the Chinese seas; specifically, there have been small-scale blooms in the Bohai Sea, Yellow Sea, and South China Sea. Further investigations should focus on establishing whether *U. meridionalis* will cause large-scale green tide events in the future.

Keywords: *Ulva meridionalis*; Ulvaceae; algal bloom; Yellow Sea; intertidal zone

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1. Introduction

A green tide is an ecological phenomenon caused by the rapid proliferation and aggregation of green macroalgae after they detach from specific substrata and reach a free-floating state. Most green tides occur in semi-closed sea areas, such as estuaries or inner bays. If a green tide cannot be removed in time, it will cause serious damage to the coastal tourism, aquaculture industry, offshore ecosystem, as well as to human health [1]. There is a higher risk of biological invasion associated with green algae than with red and brown algae [2]. *Ulva* species are common to the global oceans and estuaries, and they are usually

dominant in green tides due to their rapid growth, diverse reproduction modes, and strong adaptability to the environment [3,4].

Ulva spp. can spread to different marine areas through ballast water, seafood trade, or attached to ships and other floating objects [5-8], and they tend to bloom in new habitats as invasive species [9]. At the same time, as marine eutrophication continues to intensify worldwide, green tides are becoming more frequent at a global scale [10]. At present, more than 70 *Ulva* species are known in the world [11,12], some of which are the main dominant species causing green tide outbreaks [13,14]. Approximately 20 *Ulva* species have been recorded in China [15], and among those responsible for the outbreaks are: *Ulva prolifera*, *Ulva compressa*, *Ulva intestinalis*, *Ulva linza*, *Ulva flexuosa*, and *Ulva pertusa* [16-20].

However, over the last 10 years a new species causing green tides has emerged: *Ulva meridionalis* Horimoto et Shimada, sp. nov. (*U. meridionalis*). *U. meridionalis* is a green alga belonging to the order Ulvales, family Ulvaceae, and *Ulva* genus [21]. It is native to the tropical and subtropical areas of the Indian and Pacific Oceans [22], but can also grow in temperate areas and is commonly found on the gravel or reefs of intertidal zones, bays, and estuaries. In 2000, samples of *U. meridionalis* were collected for the first time in the estuary of the Todoroki River, Japan, and the first small-scale outbreak occurred in its estuary area [23]. In 2015, a large amount *U. meridionalis* was found in the intertidal zone of Townsville, Queensland, Australia, accounting for 94% of the *Ulva* specimens sampled. [24]. In 2020, a large biomass of green macroalgae with high species diversity was found in the intertidal zone of locations along the eastern coast of the United States and Gulf of Mexico. The study reported the existence of *U. meridionalis* in North America for the first time, but the species did not cause green tides in the region [8]. In 2022, for the first time, researchers detected a considerable biomass of *U. meridionalis* in Oceania. This species may have emerged in this region due to natural diffusion or maritime traffic [22]. At present, there is an increasing trend of green tides at the regional scale in Oceania.

Over the past five years, *U. meridionalis* has also been found in China [25-26]. At present, the species has mainly bloomed in the Bohai Sea (Fig. 1A), Yellow Sea (Fig. 1B), and South China Sea (Fig. 1C), but no outbreaks have been recorded in the East China Sea. Current observations show that *U. meridionalis* mainly grows near intertidal mudflats, dikes, offshore plastic waste, and sewage outfalls, where it becomes the dominant species. In November 2018, a *U. meridionalis* outbreak occurred in the South China Sea (Xie et al., 2020), and this species accounts for about 92% of the biomass of algal blooms. Similarly, in 2021, a green tide consisting primarily of *U. meridionalis* occurred for the first time in Yingkou, Dalian, Liaoning Province. In addition, *U. meridionalis* drops from the attached state and floats in the local sea area (e.g., the Bohai Sea area), and a large quantity of *U. meridionalis* has been found in offshore mariculture ponds [27]. The aim of the present study was to investigate *U. meridionalis* using basic biology, molecular identification, and phylogenetic methods for analysis. This study may contribute to further understanding the regularity of green tides caused by *U. meridionalis* and providing a dataset and reference to control future outbreaks.

2. Materials and Methods

2.1 Sample collection and molecular identification

From May to November 2021, green macroalgae were collected in the intertidal zones of the Bohai Sea, Yellow Sea, East China Sea, and South China Sea. The samples were washed with sterilized seawater and all epiphytes on the surface of algae were removed with a sterilized brush. Then, the samples were numbered for subsequent molecular identification.

The Ezup Column Plant Genomic DNA Purification Kit was used to extract DNA from the samples. The ITS primer sequences were ITS-F (5'-TCTTTGAAACCG-TATCGTGA-3') and ITS-R (5'-GCTTATTGATATGCTTAAGTTCA GCGGGT-3') [28]. The

ITS region was amplified via polymerase chain reaction (PCR). The PCR reaction mix contained 2 µL of sample DNA, 2 µL of upstream primer, 2 µL of downstream primer, 25 µL of PCR-Mix, and 19 µL of dd-H₂O. The PCR amplification procedure was carried out at the following conditions: initial denaturation at 94°C for 5 min; 30 cycles of denaturation at 94°C for 1 min, primer annealing at 60°C for 1 min, extension at 60°C for 2 min, and holding at 60°C for 10 min. The samples were stored at 4°C. All the high-quality PCR products were sent to Sangong Biotechnology Co. Ltd (Shanghai, China) for DNA sequencing, and all the samples identified as *U. meridionalis* were used for subsequent research (Table 1).

2.2 Morphological and microscopic observations

Through observation, it was found that the morphological characteristics of *U. meridionalis* in the four sampled areas were consistent, and the samples collected from Liaoning Province in the Bohai Sea area were selected (Station number: CN9) (Table 1) to conduct further basic biological observations. The optical microscope (E200, Nikon, Tokyo, Japan) was used to observe the branching structure of the algae in detail, and the length and width of cells in the base, middle, and upper regions were measured. The starch granules and morphology of algal cells were photographed. Using the punching method to induce algal release [29], then the sporangium, tetraflagellate meiospore, and biflagellate gametes produced by promoting gamete release were observed and photographed under an optical microscope. Then, the single cell culture of *U. meridionalis* was carried out at 20°C, 62.5–75 µmol m⁻² s⁻¹, and a 12 h light: 12 h dark photoperiod (12L: 12D) for 65 days; the culture medium was changed every 3 days. Photographs of *U. meridionalis* were taken with a Sony camera (7M3, SONY, Tokyo, Japan) as morphological records.

2.2 Morphological and microscopic observations

The *U. meridionalis* ITS sequences were obtained for the following samples collected from the four sea areas object of this study: (1) Bohai Sea (Sample number: CN9-1-China; CN9-2-China; CN9-3-China); (2) Yellow Sea (Sample number: CN1-1-China; CN2-1-China; CN2-2-China; CN3-1-China; CN3-2-China; CN4-1-China; CN4-2-China; CN5-1-China; CN6-1-China; CN6-2-China; CN7-1-China; CN7-2-China; CN7-3-China; CN7-4-China); (3) East China Sea (Sample number: CN8-1-China); and (4) South China Sea (Sample number: CN10-1-China; CN10-2-China; CN10-3-China) (Table 1).

The following sequences, which were similar to that of *U. meridionalis*, were downloaded from the National Biotechnology Center (National Center for Biotechnology Information, NCBI) database: MK426969 *U. meridionalis* (Zhanjiang, Guangdong, South China Sea), MK426968 *U. meridionalis* (Zhanjiang, Guangdong, South China Sea), MK426970 *U. meridionalis* (Yancheng, Jiangsu, Yellow Sea), MK426972 *U. meridionalis* (Rizhao, Shandong, Yellow Sea), MK426971 *U. meridionalis* (Yancheng, Jiangsu, Yellow Sea), and other related species (HQ902008 *U. pertusa*, MG017465 *U. prolifera*, AJ000203 *U. linza*, EU933981 *U. compressa*, HM031176 *U. flexuosa*, KC661326 *U. erecta*, KC661346 *U. fasciata*, MF139299 *U. intestinalis*, MZ596117 *U. tepida*). Then, a phylogenetic tree was constructed in MEGA7.0.14 using the Maximum Likelihood (ML) method. In addition, DNASP5 software [30,31] was used to count and analyze the haplotypes of 26 samples (21 samples from the four sea areas in this study and five samples from the NCBI), and Network 5.0 was used to perform the haplotype network analysis and examine the evolutionary relationship among haplotypes in each sample.

Table 1. Geographic location of *U. meridionalis* sample stations.

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Sample numbers	Continent	Country and region	Longitude	Latitude	Habitats	Reference
CN1	Asia	Yancheng, Jiangsu Province, China	120°01'06.372"E	34°25'14.940"N	Attached to the intertidal zone	This study
CN2	Asia	Yancheng, Jiangsu Province, China	120°29'12.498"E	33°49'29.484"N	Attached to the pond	This study
CN3	Asia	Yancheng, Jiangsu Province, China	120°31'26.526"E	33°46'42.630"N	Floating in the pond	This study
CN4	Asia	Nantong, Jiangsu Province, China	121°17'22.158"E	32°27'37.794"N	Attached to the seawall	This study
CN5	Asia	Nantong, Jiangsu Province, China	121°56'11.893"E	31°42'51.779"N	Attached to the wastewater outlet	This study
CN6	Asia	Nantong, Jiangsu Province, China	121°39'14.670"E	32°04'28.422"N	Attached to the intertidal mud-flat	This study
S964	Asia	Yancheng, Jiangsu Province, China	120°45'E	33°15'N	Attached to the pond	[60]
U246-8	Asia	Yancheng, Jiangsu Province, China	120°47'E	33°15'N	Attached to the pond	[60]
S023	Asia	Lianyungang, Jiangsu Province, China	119°12'E	34°56'N	Attached to the intertidal zone	[60]
CN8	Asia	Ningbo, Zhejiang Province, China	121°47'03.120"E	29°32'47.274"N	Attached to the pier	This study
S199	Asia	Ningbo, Zhejiang Province, China	121°31'E	29°32'N	Attached to the intertidal zone	[60]
N001	Asia	Ningbo, Zhejiang Province, China	121°33'E	29°26'N	Attached to the intertidal zone	[60]
CN9	Asia	Dalian, Liaoning Province, China	122°05'01.198"E	40°14'54.776"N	Floating in the intertidal zone	This study
CN10	Asia	Wenchang, Hainan Province, China	110°44'39.000"E	19°24'59.000"N	Attached to the intertidal zone	This study
S654	Asia	Haikou, Hainan Province, China	110°32'E	20°01'N	Attached to the intertidal zone	[60]
S203	Asia	Haikou, Hainan Province, China	110°16'E	20°01'N	Attached to the intertidal zone	[60]
S981-1a	Asia	Qinhuangdao, Hebei Province, China	119°37'E	39°55'N	Attached to the intertidal zone	[60]
CN7	Asia	Qingdao, Shandong Province, China	120°06'37.756"E	36°12'26.316"N	Floating in the pond	[25,37]

H940	Asia	Qingdao, Shandong Province, China	120°20'E	36°03'N	Attached to the intertidal zone	[60]
S115	Asia	Qingdao, Shandong Province, China	120°21'E	36°02'N	Attached to the intertidal zone	[60]
S027	Asia	Qingdao, Shandong Province, China	120°41'E	36°14'N	Attached to the intertidal zone	[60]
S149	Asia	Qingdao, Shandong Province, China	120°10'E	35°53'N	Attached to the intertidal zone	[60]
S079	Asia	Qingdao, Shandong Province, China	120°20'E	36°03'N	Attached to the intertidal zone	[60]
U261-3	Asia	Rizhao, Shandong Province, China	119°36'E	35°29'N	Attached to the intertidal zone	[60]
N177-1	Asia	Weihai, Shandong Province, China	122°09'E	37°31'N	Attached to the intertidal zone	[60]
N186-1	Asia	Weihai, Shandong Province, China	122°11'E	37°30'N	Attached to the intertidal zone	[60]
U225a	Asia	Zhanjiang, Guangdong Province, China	110°04'E	20°17'N	Attached to the intertidal zone	[60]
U277-1a	Asia	Zhanjiang, Guangdong Province, China	110°25'E	21°13'N	Attached to the seawall	[60]
S209	Asia	Beihai, Guangxi Zhuang Autonomous Region, China	109°09'E	21°24'N	Attached to the intertidal zone	[60]
S217	Asia	Shenzhen, Guangdong Province, China	114°01'E	22°31'N	Attached to the intertidal zone	[60]
S599	Asia	Xiamen, Fujian Province, China	118°05'E	24°34'N	Attached to the pond	[60]
RH008	Asia	Ishigaki Island, Okinawa Prefecture, Japan	124°15'E	24°22'N	Attached to the estuary	[23]
RH001-007	Asia	Ishigaki Island, Okinawa Prefecture, Japan	124°15'E	24°22'N	Attached to the estuary	[23]
RH009-037	Asia	Ishigaki Island, Okinawa Prefecture, Japan	124°15'E	24°22'N	Attached to the estuary	[23]
RH043-047	Asia	Ishigaki Island, Okinawa Prefecture, Japan	124°15'E	24°22'N	Attached to the estuary	[23]

E16	Asia	Tokushima, To- kushima Prefec- ture, Japan	124°15'E	24°22'N	Attached to the estuary	[23]
UNA0007 1829	North America	Cedar Point, Ala- bama, America	88°08'13.459"W	30°18'37.865"N	Floating in the intertidal zone	[8]
UNA0007 1885	North America	Coden, Alabama, America	88°15'30.200"W	30°22'54.894"N	Floating in the intertidal zone	[8]
UNA0007 2126	North America	Choctawatchee Bay Bridge, Flor- ida, America	86°09'20.999"W	30°25'41.401"N	Floating in the intertidal zone	[8]
UNA0007 2127	North America	Choctawatchee Bay Bridge, Flor- ida, America	86°09'21.038"W	30°25'41.402"N	Floating in the intertidal zone	[8]
UNA0007 2079	North America	Copano Fishing Pier, Texa, Amer- ica	97°01'32.624"W	28°06'47.833"N	Floating in the intertidal zone	[8]
UNA0007 2312	North America	Charlotte Harbor, Florida, America	82°04'17.893"W	26°57'22.950"N	Floating in the intertidal zone	[8]
NOU2187 15	Oceania	NA, Moindou, French New Cal- edonia	NA	NA	Attached to the intertidal zone	[22]
NOU2187 42	Oceania	NA, Poe, French New Caledonia	NA	NA	Attached to the intertidal zone	[22]
NOU2187 43	Oceania	NA, Poe, French New Caledonia	NA	NA	Attached to the intertidal zone	[22]
NOU2187 55	Oceania	NA, Bourail, French New Cal- edonia	NA	NA	Attached to the intertidal zone	[22]
NOU2188 03	Oceania	NA, islet Double, French New Cal- edonia	NA	NA	Attached to the intertidal zone	[22]
NOU2188 47	Oceania	NA, Poe, French New Caledonia	NA	NA	Attached to the intertidal zone	[22]
NOU2188 26	Oceania	NA, Cap Goule- vain, French Pol- ynesia	NA	NA	Attached to the intertidal zone	[22]
NOU2188 32	Oceania	NA, Cap Goule- vain, French Pol- ynesia	NA	NA	Attached to the intertidal zone	[22]
NOU2187 70	Oceania	NA, Tahiti, French Polynesia	NA	NA	Attached to the intertidal zone	[22]
NOU2153 08	Oceania	NA, Marquises, French Polynesia	NA	NA	Attached to the intertidal zone	[22]
NOU2153 09	Oceania	NA, Marquises, French Polynesia	NA	NA	Attached to the intertidal zone	[22]
NOU2188 67	Oceania	NA, Moorea, French Polynesia	NA	NA	Attached to the intertidal zone	[22]
TSV31	Oceania	Townsville, Queensland, Australia	146°48'39.999"E	19°14'42"S	Attached to the intertidal zone	[24]

3. Results

3.1 Basic biological characteristics of *U. meridionalis*

In a natural environment and under favorable growth conditions, the length and width of *U. meridionalis* can reach 45 cm and 0.8 cm, respectively. The alga is light green or yellow green, it has a smooth surface, and is easy to break. Under indoor culture conditions, it can grow to a length of 3 m (Fig. 2A). It has a tubular structure and an obvious main branch structure (Fig. 2B), especially folds in the wider part of the algae. In addition, it is also characterized by a monolayered tubular structure and a double-layer cell structure. The width of the transverse branch, which basically appears in the area near the base, decreases toward the branch node (Fig. 2C). For the collected samples, the length of algal cells in the base region was 9–30 μm , the width was 7–17 μm (Fig. 2D), and the cells contained starch granules (Fig. 2E). In the middle region, the length and width of algal cells was 8–23 μm and 6–16 μm , respectively (Fig. 2F), and in the upper region, the same parameters measured 8–20 μm and 5–16 μm , also respectively (Fig. 2G). Both the length and width of *U. meridionalis* cells decreased moving from the base to the upper region. The chloroplasts of each cell contained 1–11 starch granules; specifically, the chloroplasts of the base and middle algal cells mainly had 4–5 and 4–6 starch granules, respectively, while there were 3–4 starch granules in the chloroplasts of the upper algal cells. The surface cells of *U. meridionalis* had a rounded triangle, rectangle, or polygon shape. Generally, after induced dispersal, *U. meridionalis* forms spores in about 3.5 days, which gradually disperse, and sporophytes and gametophytes can form a complete individual. The gametangium of *U. meridionalis* (Fig. 2H), tetraflagellate meiospore are ovate and have a reddish eyespot (Fig. 2I), it is manifested as negative phototaxis. The biflagellate gametes are oval (Fig. 2J–K), it shows positive phototaxis with reddish eyespot.

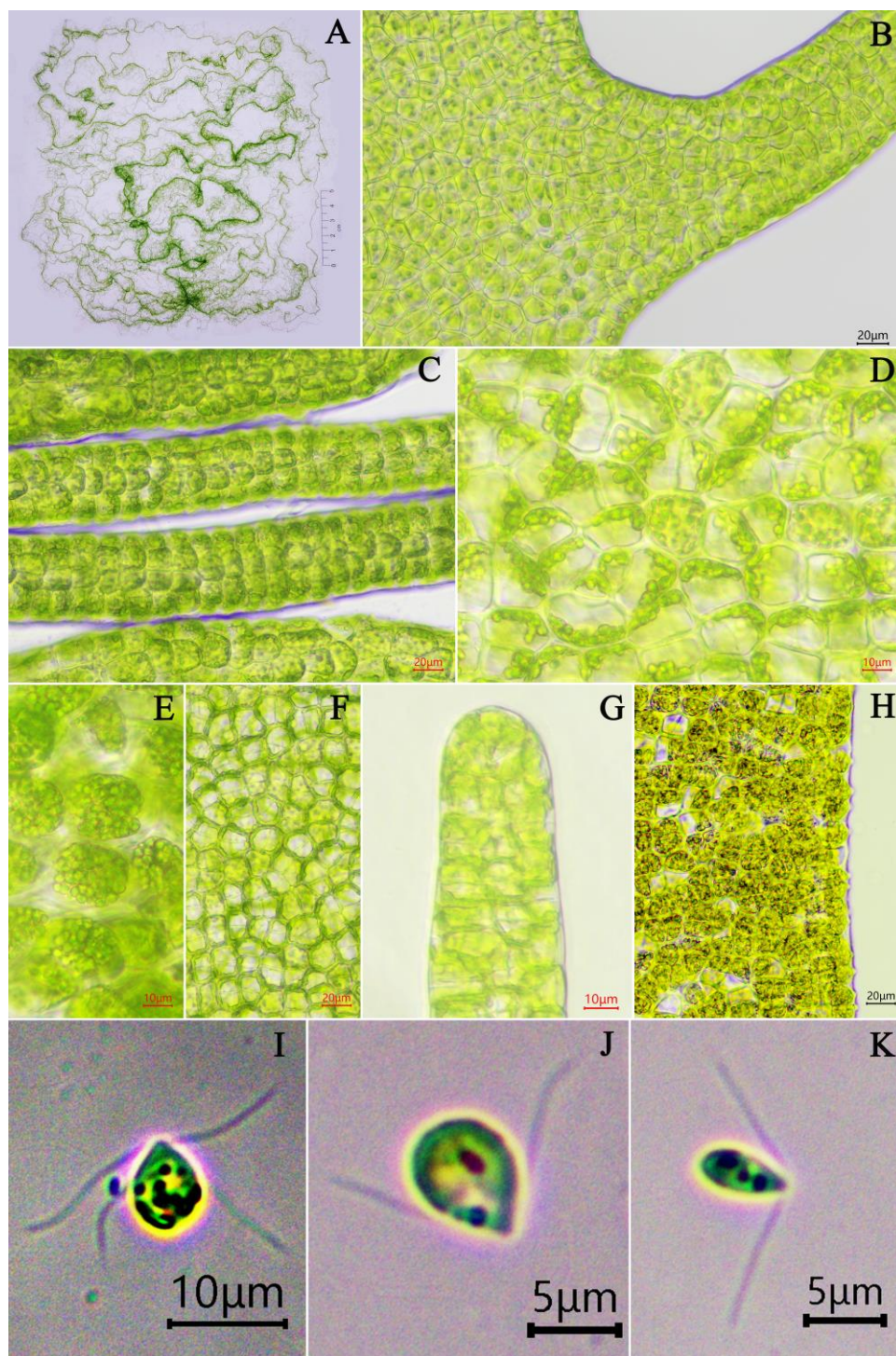


Figure 2. Photographs of *U. meridionalis* depicting: (A) morphology of the macroalga under indoor culture conditions; (B) main branching structure; (C) proximity of the horizontal branch to the base area; (D) base area; (E) cells containing more starch granules; (F) middle region; (G) upper region; (H) gametangium; (I) tetraflagellate meiospore; (J) biflagellate female gamete; (K) biflagellate male gamete.

3.2. Genetic relationship among *U. meridionalis* samples

Based on the ITS-ML phylogenetic tree (Fig. 3), all the samples examined in this study fell into seven evolutionary clusters, and HM031176 *U. flexuosa*, HQ902008 *U. pertusa*, and EU933981 *U. compressa* were grouped into separate evolutionary clusters.

The following sequences were grouped into single clusters: MF139299 *U. intestinalis* and MZ596117 *U. tepida*, (with a 100% sequence similarity); KC661326 *U. erecta* and KC661346 *U. fasciata* (sequence similarity, 99%); MG017465 *U. prolifera* and AJ000203 *U. linza* (sequence similarity, 100%) (Fig 3). There are two reasons for this: on one hand, Although the ITS sequence as a bar code has a strong universality, it is not sufficient to distinguish few species under certain circumstances; on the other hand, it is also possible that the same species have different names.

All the *U. meridionalis* sequences were grouped into a single cluster (Sample numbers: CN9-1-China; CN9-2-China; CN9-3-China; CN1-1-China; CN2-1-China; CN2-2-China; CN3-1-China; CN3-2-China; CN4-1-China; CN4-2-China; CN5-1-China; CN6-1-China; CN6-2-China; CN7-1-China; CN7-2-China; CN7-3-China; CN7-4-China; CN8-1-China; CN10-1-China; CN10-2-China; CN10-3-China; MK426969 *U. meridionalis*; MK426970 *U. meridionalis*; MK426972 *U. meridionalis*; MK426971 *U. meridionalis*; MK426968 *U. meridionalis*), showing that the ITS bar code can identify *U. meridionalis* with a high accuracy. Among these sequences, the relationship between genetic distance and proximity was determined as follows: *U. meridionalis* samples collected from Wenchang City, Hainan Province in the South China Sea (CN10-3-China; CN10-2-China) $\Rightarrow$ *U. meridionalis* samples collected from Nantong City, Jiangsu Province in the Yellow Sea (CN6-1-China) $\Rightarrow$ *U. meridionalis* samples collected from Yancheng City, Jiangsu Province in the Yellow Sea (CN5-1-China; CN4-1-China; CN4-2-China; CN3-1-China; CN3-2-China; CN2-2-China; CN2-1-China; CN1-1-China) $\Rightarrow$ *U. meridionalis* samples collected from in Dalian City, Liaoning Province in the Bohai Sea (CN9-2-China; CN9-1-China; CN9-3-China) $\Rightarrow$ *U. meridionalis* samples collected from Nantong City, Jiangsu Province in the Yellow Sea (CN6-2-China) $\Rightarrow$ *U. meridionalis* samples collected from Wenchang City, Hainan Province in the South China Sea (CN10-1-China) $\Rightarrow$ *U. meridionalis* samples collected from Qingdao City, Shandong Province in the Yellow Sea (CN7-1-China; CN7-2-China; CN7-3-China; CN7-4-China) $\Rightarrow$ *U. meridionalis* samples collected from Ningbo City, Zhejiang Province in the East China Sea (CN8-1-China). Individual, basic group differences were detected between the above *U. meridionalis* samples, but the degree of subspecies differentiation was not reached (Fig. 3). Also, the ITS sequence indicated a close genetic relationship between *U. meridionalis* and *U. pertusa*, which is inconsistent with the results of the phylogeny of *U. meridionalis* based on organelle genome analysis.

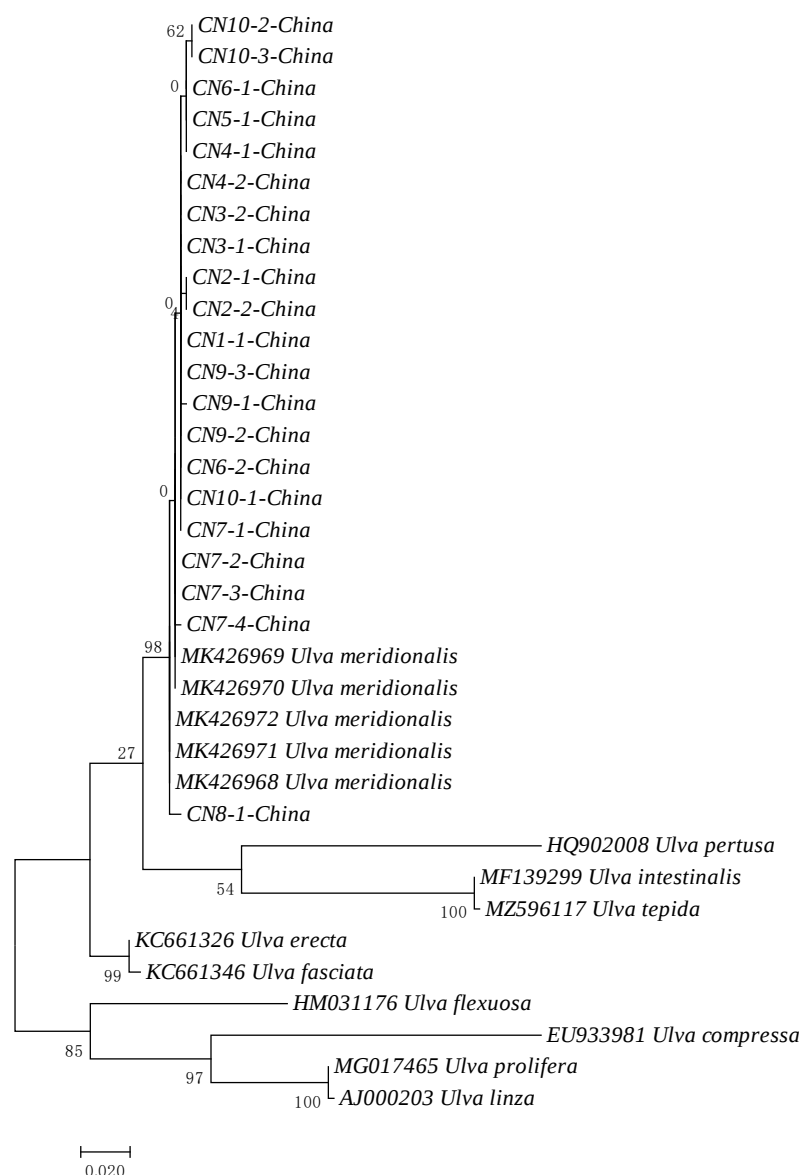


Figure 3. ML phylogenetic tree constructed based on the Kimura 2-parameter method.

The circle symbol represents one haplotype and its size reflects the number of different haplotypes. The different colors represent different sources from which the haplotypes were derived. The network evolution diagram (Fig. 4) shows that the *U. meridionalis* samples collected in the four sea areas in China are genetically close, and the haplotype evolution network showed a chain structure. A total of five haplotypes were detected: the samples from the East China Sea corresponded to the independent haplotype Hap_3; those from the South China Sea contained three haplotypes, Hap_1, Hap_2, and Hap_5; those from the Bohai sea presented two haplotypes, Hap_1 and Hap_2; and finally, the samples from the Yellow Sea contained three haplotypes, Hap_1, Hap_2, and Hap_4. Hap_2 and Hap_1 were the largest haplotypes and they were shared among the three groups (the South China Sea, the Bohai sea, the Yellow Sea), they are located in the center of the network evolution map and it is speculated that they are very likely the ancestral haplotypes of the group. Hap_3, which was detected in the *U. meridionalis* samples of the East China Sea, showed limited level of genetic isolation from the other haplotypes, however, it did not reach the degree of subspecies formation (Fig. 3). It is speculated that the samples with this haplotype may have experienced a short period of geographical isolation.

tion. At the same time, the haplotype network diagram supports the results of the phylogenetic tree. The *U. meridionalis* samples from the East China Sea may be local, but they may also be invasive, either derived from natural conditions or artificially introduced new strains. The *U. meridionalis* samples in the Yellow Sea, Bohai Sea, and South China Sea have expanded from haplotype Hap_2 and Hap_1 population in recent years, and no obvious systematic geographical pattern has been detected among them (Fig. 3–Fig. 4).

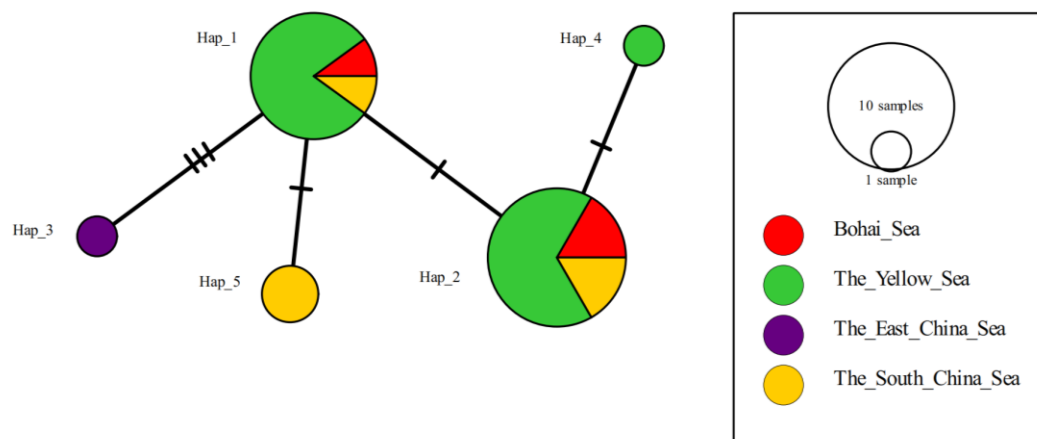


Figure 4. Network diagram of the *U. meridionalis* haplotypes derived from the ITS sequences.

4. Discussion

Since 1970s, the incidence of green tides has increased steadily [32] and, as the scale of these events continues to expand, prevention has become the focus of attention among researchers. In fact, with the increase in nitrogen supply, the growth rate of *Ulva* spp. also increases [33], and their distribution is expected to expand also due to global warming [34,35]. Therefore, controlling green tide outbreaks is a global challenge. For a long time, the species diversity and distribution of *Ulva* spp. has attracted considerable attention in terms of the investigation of algal resources, especially as species belonging to this genus easily cause green tide outbreaks. As more investigations in seaweed farms were conducted worldwide [36], new species or new records were successively reported. For example, *U. meridionalis*, which has a high growth rate with an average daily growth rate of 0.37 day⁻¹ and maximum daily growth rate of more than 1.12 day⁻¹ [21], was shown to be a new green tide-forming species [21,25,26,37]. At present, there are frequent outbreaks of harmful algal blooms around the world (such as red tides, green tides and golden tides) [38–40]. ~~The discovery of *U. meridionalis* not only enriches the species diversity of *Ulva* worldwide, but also increases the uncertainty of the future trend of green tide outbreaks in various countries.~~

As a eurythermal and salt-tolerant species, *U. meridionalis* is widely distributed. At present, it has been found in Asia, Oceania, and North America (Fig. 5; Table 1). Up to 20 years ago, this species had been recorded only in Japan. However, in recent years, it has also been reported in the United States, China (the Bohai Sea, the Yellow Sea, the East China Sea, and the South China Sea), Australia, French New Caledonia, and French Polynesia. Furthermore, massive proliferation of this macroalga has been reported in the Bohai Sea, Yellow Sea, and South China Sea. At present, *U. meridionalis* blooms as a dominant species in the tropical and subtropical coastal seawaters of China in spring and autumn (Fig. 1). In summer, large-scale outbreaks occur even in temperate regions, suggesting that the species has a strong diffusion ability and can gradually spread to higher northern latitudes, which may be due to global warming [34,35,41,42]. The distribution of *U. meridionalis* is increasingly extensive (Table 1); however, the routes through which it is spreading and the diffusion mechanism are unknown, and further studies should investigate these aspects.

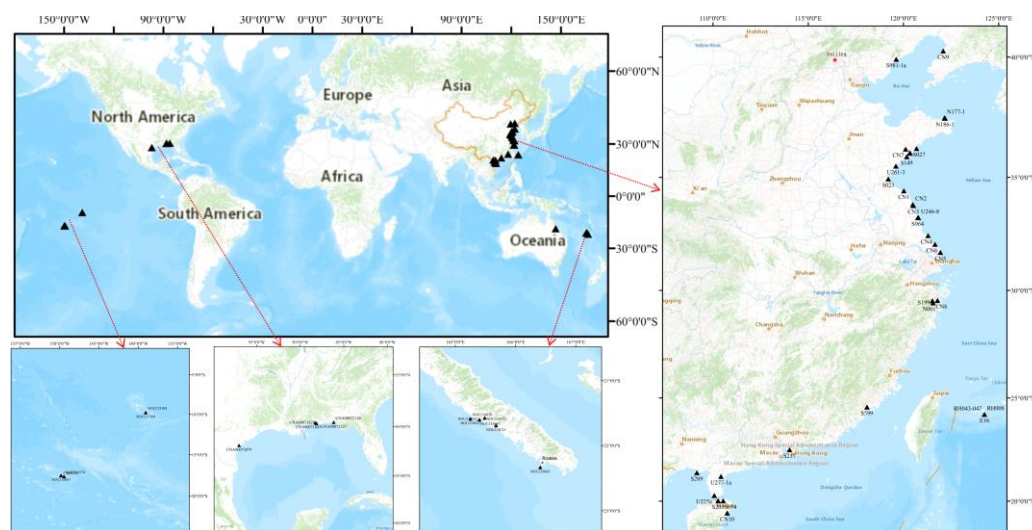


Figure 5. Global distribution of *U. meridionalis*. See Table 1 for details on the sampling stations.

U. meridionalis has unique and stable morphological characteristics and it is clearly different from the *Ulva* spp. commonly found in China. At present, there is no obvious evidence that *U. meridionalis* is native to any particular country, therefore the possibility that this macroalga is a new alien invasive species entering China cannot be ruled out. Further molecular studies need to be conducted to trace its source. Previous comparative analyses based on the mitochondrial genome showed that the relatives of *U. meridionalis* and *U. flexuosa* are genetically close. However, comparative analyses of the chloroplast genome [37, 39] revealed that *U. meridionalis* is closely related to *U. linza* [25]. In the present study, a close genetic relationship was found between *U. meridionalis* and *U. pertusa* based on the ITS sequence. This is inconsistent with the results of previous phylogenetic studies of *U. meridionalis* based on organelle genomes [25, 37, 39]. In addition, there are relatively few studies on *Ulva* DNA barcode sequences and organelle genomes for the four sea areas in China sampled in the present study (For instance, there is a lack of research on the organelle genome of *U. meridionalis* and other *Ulva* spp. in different sites). At present, the number and type of gene sequences available from the NCBI are not sufficient to effectively determine the genetic relationship of *U. meridionalis* with other *Ulva* species, which remains to be further studied. Genome-wide research [43], plant organelle genome research [27], single nucleotide polymorphism technology, simple sequence repeat technology, DNA barcode sequence research, and other technologies could be adopted at their maximum capacity to clarify the origin of *U. meridionalis* and its outbreak mechanism. These technologies would also allow to explore whether the distribution and outbreaks of this macroalga are related to its own biological characteristics and environmental factors or they are affected by species variation, intraspecific hybridization, and other associated reasons.

At present, the area with the largest green tide outbreaks in the world is the Southern Yellow Sea. Green tides with *U. prolifera* as the dominant species have occurred in this sea in spring and summer for 15 consecutive years. These events occur on a large scale, they last long [44]. Whether *U. meridionalis* will bloom under suitable conditions in the Yellow Sea is uncertain. The possibility that this species may also cause large-scale outbreaks similarly to *U. prolifera*, harm the structure and function of the local ecosystem, and complicate the green tide dynamics in the Southern Yellow Sea should be reasons of major concern. In particular, it should be considered that at present *U. meridionalis* has appeared as a floating phenomenon and a large-scale outbreak has been reported in the Bohai Sea (Fig. 1A). In addition, this species is also widely distributed in the Yellow Sea, where its biomass is enormous, and large-scale floating has been reported in offshore ponds and intertidal areas as well (Fig. 1B).

U. meridionalis has a strong ecological adaptability. Studies have shown that at P and N concentrations of 0.125 mg/L and 4.02 mg/L, respectively, its maximum growth rate could reach 0.796 day⁻¹ [21]. Compared with the other *Ulva* spp. [33,45–51], *U. meridionalis* showed a very high growth rate. For example, the growth rate of *U. prolifera* in a wide salinity range between 5 and 30 and temperatures between 10 and 30°C was 0.37–0.89 day⁻¹. In comparison, at a 10–30 salinity range and temperature of 30°C, the growth rate of *U. meridionalis* could reach 1.4 day⁻¹ [51]. Under the best culture conditions, its daily growth rate exceeded 1 day⁻¹ [52]. Among the multicellular autotrophic plants, *U. meridionalis* showed the fastest growth rate, which makes it a high-quality algal biomass resource [51]. In addition, *U. meridionalis* is characterized by a high photosynthetic carbon sequestration rate, frequent cell proliferation, and rapid formation of algal bodies composed of cell walls in high temperature environments. The accumulated cell and cell wall components include polysaccharides and rare sugars [52–55]. The prospect of resource utilization is broad. At present, only few studies have been conducted on *U. meridionalis* globally. Just nine articles (two in Chinese and seven in English) have been published and the present study mainly reports the physiological and ecological characteristics of this macroalga [11,21,26,27,37,51,57–59]. In general, no systematic studies have been conducted on *U. meridionalis* fortunately, researchers are gradually beginning to focus on it. For instance, it has been shown that this species has a fast growth rate and a high ammonia nitrogen absorption rate [21,51]. It has a broad temperature and salinity resistance and may have several future applications, for example, in the advanced treatment of municipal sewage, in the sulfated polysaccharide extraction industry, and also as a potential pharmaceutical and food raw material [52]. It is suggested to further develop new ways to utilize *U. meridionalis* resources by transforming the biomass produced by this green tide-forming species into materials that are beneficial to human life. The use of *U. meridionalis* as a resource will contribute to the reduction of the algal biomass, and, consequently, to the control of potential large-scale outbreaks in the future.

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