



Bolidophyceae, a Sister Picoplanktonic Group of Diatoms – A Review

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Pico- and nano-phytoplankton (respectively, 0.2–2 and 2–20 μm in cell size) play a key role in many marine ecosystems. In this size range, Bolidophyceae is a group of eukaryotes that contains species with cells surrounded by 5 or 8 silica plates (Parmales) as well as naked flagellated species (formerly Bolidomonadales). Bolidophyceae share a common ancestor with diatoms, one of the most successful groups of phytoplankton. This review summarizes the current information on taxonomy, phylogeny, ecology, and physiology obtained by recent studies using a range of approaches including metabarcoding. Despite their rather small contribution to the phytoplankton communities (on average less than 0.1%), Bolidophyceae are very widespread throughout marine systems from the tropics to the pole. This review concludes by discussing similarities and differences between Bolidophyceae and diatoms.

Keywords: bolidophyceae, parmales, diatoms, genetic diversity, mitosis, geographical distribution, seasonal dynamics and silicification

INTRODUCTION

Following the appearance of oxygenic photosynthesis in the ancestors of cyanobacteria, this complex process was distributed across all eukaryotic lineages via permanent primary, secondary, and tertiary endosymbioses (Not et al., 2012). Ocean photosynthesis is dominated by phytoplankton, a functional group of single cell organisms including prokaryotes and eukaryotes. In the late 70's, early 80's the work of Waterbury et al. (1979) and Johnson and Sieburth (1982) revealed the importance of very small cells, some below one micron in size, for primary productivity, which importance was formalized with the concept of the microbial loop by Azam et al. (1983). However, it was only in the mid 90's, when researchers began to investigate the eukaryotic compartment of picophytoplankton, and realized that while cyanobacteria are

very little diversified, at the least at the genus level with a couple of taxa (*Prochlorococcus*, *Synechococcus*) dominating, eukaryotes turned out to be very diverse with picoplankton taxa distributed widely across several branches of eukaryotic tree of life (Vaulot et al., 2008).

Microphytoplankton such as diatoms, dinoflagellates or coccolithophorids that produce short lived blooms such as *Emiliania huxleyi*, have been extensively investigated, in contrast to other members of the picoplanktonic community. A large number of photosynthetic picoeukaryotes species (and clades) described to date belong to Stramenopiles (also called Heterokonts), which are characterized by flagellated cells, produced at least at some point of their life cycles, with two unequal flagella (heterokont), one being ornamented with hair-like structures called mastigonemes. Their plastids are thought to have been acquired through secondary endosymbiosis and typically contain chlorophylls *a* and *c*. Stramenopiles besides include diatoms, a very successful lineage which cells are encased in ornamented silica structures, Dictyochophyceae, also called silicoflagellates, and Pelagophyceae, well known because of the toxic algal blooms of the brown tide species *Aureococcus anophagefferens* and *Aureoumbra lagunensis* (Gobler and Sunda, 2012) or *Pelagomonas calceolata*, frequently isolated from sea water. Although often seen as less diverse, some of these groups play important ecological roles in marine ecosystems are fundamental for our understanding of the evolution of algae.

Bolidophyceae, a class created by Guillou et al. (1999a) is the Stramenopiles group phylogenetically nearest to the diatoms. They are often detected in molecular surveys, although in low abundance. We now know that they can occur as two distinct forms, either silicified pico-sized (2–5 μm) or non-silicified flagellated (1–1.7 μm) cells (Guillou et al., 1999a; Ichinomiya et al., 2011, 2016) and cultures from both forms have been isolated from the marine environment. This review summarizes the discovery of Bolidophyceae and current information of phylogeny, ecology, and physiology obtained by recent studies using a range of approaches. We discuss similarities and differences between Bolidophyceae and diatoms to explore the evolutionary link between these silicified algal groups.

DISCOVERY AND TAXONOMY

Well before the creation of the class Bolidophyceae, their silicified forms (Figure 1) were first reported from scanning electron microscopy (SEM) images in oceanic samples from the North Pacific (Iwai and Nishida, 1976). Initially, they were thought to be resting cysts of silicified loricate choanoflagellates (Silver et al., 1980). However, the observation of red auto fluorescence indicating the presence of chlorophyll and the existence of a chloroplast in sectioned cells observed by transmission electron microscopy revealed that they were active phytoplankton cells

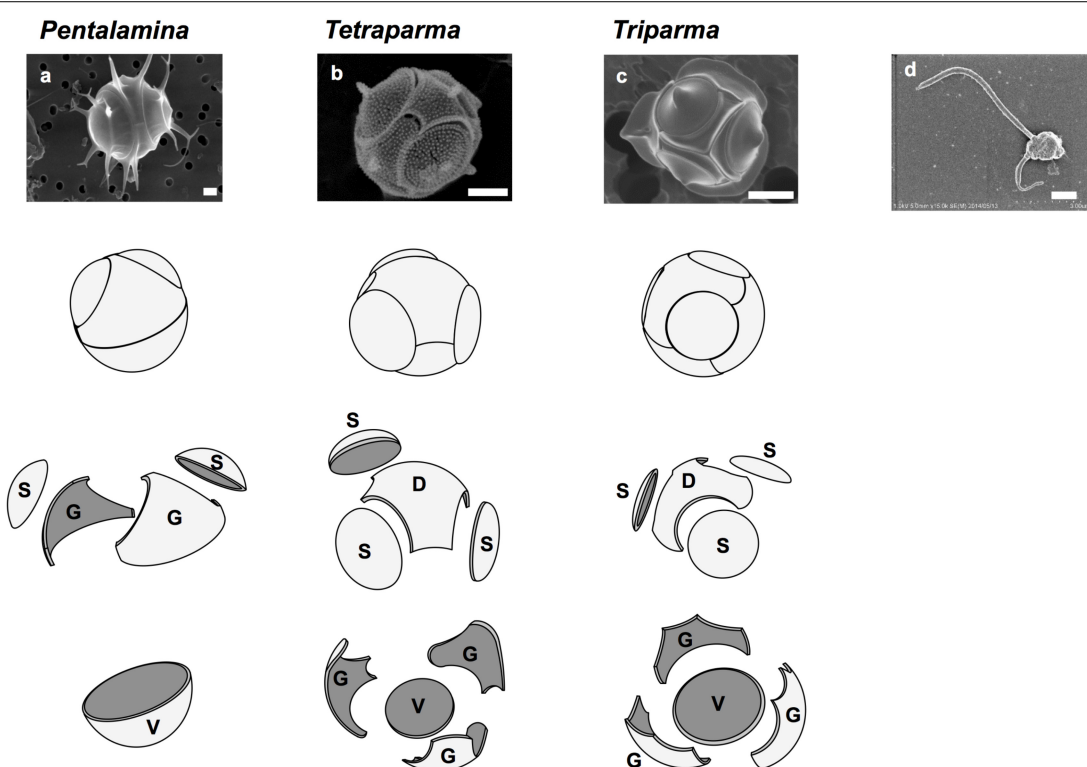


FIGURE 1 | Silicified and flagellated species of Bolidophyceae. **(a)** *Pentalamina corona*, **(b)** *Tetraparma pelagica*, **(c)** *Triparma laevis* f. *inornata*, and **(d)** *Triparma eleuthera*. Scale bars = 1 μm . Diagrams of silica plates in the three genera of silicified species (Parmales); D, dorsal plate; G, girdle plate; S, shield plate; and V, ventral plate. Redrawn from Booth and Marchant (1987) with permission.

(Marchant and McEldowney, 1986). Marchant and McEldowney (1986) could not establish their taxonomic position, although they suggested some morphological similarities with other algae groups such as Bacillariophyceae and Chrysophyceae.

Booth and Marchant (1987) tentatively established Parmales (Latin: small round shields) as a new order within Chrysophyceae. The taxonomy of Parmales was based on the morphological features of silica plates that can only be observed with SEM. Two families and three genera were established: Pentalaminaceae with one genus *Pentalamina* (Latin: five plates) and Triparmaceae with two genera, *Tetraparma* (Latin: four small round shields) and *Triparma* (Latin: three small round shields) (Figures 1a–c). *Pentalamina* has 2 circular shield plates of equal size, a larger ventral plate and 2 triradiate girdle plates. *Tetraparma* has 3 shield plates of equal size, a smaller ventral plate, a triradiate dorsal plate and 3 girdle plates. *Triparma* has 3 shield plates of equal size, a larger ventral plate, a triradiate dorsal plate and 3 girdle plates (Booth and Marchant, 1987, 1988; Kosman et al., 1993; Bravo-Sierra and Hernández-Becerril, 2003; Konno and Jordan, 2007; Konno et al., 2007).

The flagellated forms of Bolidophyceae (Figure 1d) were originally isolated from the Pacific Ocean and Mediterranean Sea, and described as two flagellated species, *Bolidomonas pacifica* and *B. mediterranea*, differing in the angle of the insertion of the two flagellum, swimming patterns as well as in 18S rRNA gene signatures (Guillou et al., 1999a). The name “*Bolidomonas*” refers to the rapid swimming behavior of the cells remembering a racing car. A variety, *B. pacifica* var. *eleuthera*, was later proposed based on both cultures and environmental sequences (Guillou et al., 1999b). Analyses of photosynthetic pigments as well as nuclear 18S rRNA and plastid RubisCO large subunit (*rbcL*) sequences (Guillou et al., 1999a; Daugbjerg and Guillou, 2001) demonstrated the sister relationship between *Bolidomonas* and diatoms, although *Bolidomonas* are flagellates and lack the siliceous frustule characteristic of diatoms. Bolidophyceae were thus proposed to be an intermediate group between diatoms and all other Stramenopiles (Guillou et al., 1999a).

For more than 24 years, Parmales escaped isolation. These silicified cells are small and difficult to distinguish them from other small phytoplankton in field samples under the light microscope. To overcome this problem, Ichinomiya et al. (2011) used the fluorescence dye PDMPO (2-(4-pyridyl)-5-((4-(2-dimethylaminoethylaminocarbamoyl) methoxy)phenyl) oxazole) (Shimizu et al., 2001), which is co-deposited with silicon into the solid silica matrix of the newly produced cell walls and fluoresces under UV excitation whenever silicic acid is polymerized forming biogenic silica. Using PDMPO staining and a serial dilution technique, the first Parmales strain was established from Oyashio region of the western North Pacific (Ichinomiya et al., 2011).

Scanning electron microscopy established that this strain belonged to the species *Triparma laevis* and transmission electron microscope observations showed the typical ultrastructure of photosynthetic Stramenopiles, with two endoplasmic reticulate membranes surrounding the chloroplast, a girdle and two to three thylakoid lamellae as well as a mitochondrion with tubular

cristae. Phylogenetic analyses based on 18S ribosomal rRNA sequences from the new strain demonstrated that *T. laevis* was closely related to Bolidophyceae (Ichinomiya et al., 2011), rather than part of Chrysophyceae as hypothesized initially (Booth and Marchant, 1987). Phylogenetic analyses using plastidial and mitochondrial encoded genes from *T. laevis* also confirmed its sisterhood with Bolidophyceae and diatoms (Tajima et al., 2016).

Recent phylogenetic analyses using nuclear, plastidial, and mitochondrial genes from several novel strains, including a flagellate form very closely related to the silicified strains, led to a taxonomic revision (Table 1) in which the order Parmales was included within the class Bolidophyceae and *Bolidomonas* species were transferred to the genus *Triparma* (Ichinomiya et al., 2016).

GENETIC DIVERSITY

Clade Diversity

The analysis of full-length nuclear 18S rRNA gene sequences from public databases revealed the existence of two environmental clades (Env. clade I and II) in addition to the group corresponding to the genus *Triparma* (Ichinomiya et al., 2016). These clades are only formed by environmental sequences and no sequences from cultures or isolates are available. Within the *Triparma* group, sub-clades formed by sequences from strains and the environment corresponded to the species *Triparma eleuthera*, *Triparma pacifica*, and *Triparma mediterranea*. The “*T. laevis*” sub-clade, including the species *T. laevis* f. *inornata*, *T. laevis* f. *longispina*, *Triparma strigata*, *Triparma* aff. *verrucosa* and the flagellated strain *Triparma* sp. RCC1657. Other molecular markers (plastid 16S rRNA and *rbcL*, nuclear ITS rRNA and mitochondrial *nad1*) revealed the presence of two distinct sub-clades within the “*T. laevis*” sub-clade, hereafter called for convenience *Triparma* clade I with the two forms of *T. laevis* (f. *inornata*, and *longispina*) and *Triparma* clade II with *T. strigata*, *T. aff. verrucosa* and the flagellated strain RCC1657.

In order to review the current state of the diversity of Bolidophyceae, we analyzed existing GenBank sequences as well as metabarcodes obtained from a range of recent studies (Table 2) focusing on the V4 region of the 18S rRNA gene (see Supplementary Material for Methodology). The phylogenetic analysis of the newly obtained V4 sequences (Figure 2) recovered the two major environmental clades previously described (Env. clade I and II, Ichinomiya et al., 2016), but also revealed the existence of a third environmental clade (called Env. clade III) within which two sub-clades IIIA and IIIB can be clearly separated. Each environmental clade contained sequences from clone libraries (GenBank) as well as identical or nearly identical metabarcode sequences from different surveys suggesting that these environmental clades are not artefactual. These environmental clades may, for some of them, correspond to species of Parmales (e.g., from genera *Tetraparma*

TABLE 1 | Current taxonomy of Bolidophyceae.

class, order, family, genus, species, subspecies, forma (= synonym, basyonym)	Reference
Class Bolidophyceae Guillou et Chrétiennot-Dinet emend. Ichinomiya et Lopes dos Santos	Guillou et al., 1999a ; Ichinomiya et al., 2016
Order Parmales Booth et Marchant emend. Konno et Jordan emend. Ichinomiya et Lopes dos Santos	Booth and Marchant, 1987; Konno and Jordan, 2007; Ichinomiya et al., 2016
Family Pentalaminaceae Marchant emend. Konno et Jordan	Booth and Marchant, 1987; Konno and Jordan, 2007
Genus <i>Pentalamina</i> Marchant	Booth and Marchant, 1987
<i>Pentalamina corona</i> Marchant	Booth and Marchant, 1987
Family Triparmaceae Booth et Marchant emend. Konno et Jordan emend. Ichinomiya et Lopes dos Santos (= "Octolaminaceae" Booth et Marchant)	Booth and Marchant, 1987; Booth and Marchant, 1988; Konno and Jordan, 2007; Ichinomiya et al., 2016
Genus <i>Tetraparma</i> Booth emend. Konno et Jordan	Booth and Marchant, 1987; Konno and Jordan, 2007
<i>Tetraparma catinifera</i> Konno et al.	Konno et al., 2007
<i>Tetraparma gracilis</i> Konno et al.	Konno et al., 2007
<i>Tetraparma insecta</i> Bravo-Sierra et Hernández-Becerril emend. Fujita et Jordan	Bravo-Sierra and Hernández-Becerril, 2003; Fujita and Jordan, 2017
<i>Tetraparma pelagica</i> Booth et Marchant	Booth and Marchant, 1987
<i>Tetraparma silverae</i> Fujita et Jordan	Fujita and Jordan, 2017
<i>Tetraparma trullifera</i> Fujita et Jordan	Fujita and Jordan, 2017
Genus <i>Triparma</i> Booth et Marchant emend. Konno et Jordan emend. Ichinomiya et Lopes dos Santos (= <i>Bolidomonas</i> Guillou et Chrétiennot-Dinet)	Booth and Marchant, 1987; Guillou et al., 1999a ; Konno and Jordan, 2007; Ichinomiya et al., 2016
<i>Triparma columacea</i> Booth	Booth and Marchant, 1987
<i>Triparma columacea</i> f. <i>convexa</i> Konno et al.	Konno et al., 2007
<i>Triparma columacea</i> f. <i>fimbriata</i> Konno et al.	Konno et al., 2007
<i>Triparma columacea</i> f. <i>longiseta</i> Fujita et Jordan	Fujita and Jordan, 2017
<i>Triparma columacea</i> subsp. <i>alata</i> Marchant	
<i>Triparma eleuthera</i> Ichinomiya et Lopes dos Santos (= " <i>Bolidomonas pacifica</i> var. <i>eleuthera</i> ")	Ichinomiya et al., 2016
<i>Triparma laevis</i> Booth	Booth and Marchant, 1987
<i>Triparma laevis</i> f. <i>fusiformis</i> Fujita et Jordan	Fujita and Jordan, 2017
<i>Triparma laevis</i> f. <i>inornata</i> Konno et al.	Konno et al., 2007
<i>Triparma laevis</i> f. <i>longispina</i> Konno et al.	Konno et al., 2007
<i>Triparma laevis</i> f. <i>mexicana</i> (Kosman) Bravo-Sierra et Hernández-Becerril (= <i>Triparma laevis</i> subsp. <i>mexicana</i> Kosman)	Kosman et al., 1993; Bravo-Sierra and Hernández-Becerril, 2003
<i>Triparma laevis</i> subsp. <i>pinnatilobata</i> Marchant	Booth and Marchant, 1987
<i>Triparma laevis</i> subsp. <i>ramispina</i> Marchant	Booth and Marchant, 1987
<i>Triparma mediterranea</i> (Guillou et Chrétiennot-Dinet) Ichinomiya et Lopes dos Santos (= <i>Bolidomonas mediterranea</i> Guillou et Chrétiennot-Dinet)	Guillou et al., 1999a ; Ichinomiya et al., 2016
<i>Triparma pacifica</i> (Guillou et Chrétiennot-Dinet) Ichinomiya et Lopes dos Santos (= <i>Bolidomonas pacifica</i> Guillou et Chrétiennot-Dinet)	Guillou et al., 1999a ; Ichinomiya et al., 2016
<i>Triparma retinervis</i> Booth	Booth and Marchant, 1987
<i>Triparma retinervis</i> f. <i>tortispina</i> Fujita et Jordan	Fujita and Jordan, 2017
<i>Triparma retinervis</i> subsp. <i>crenata</i> Booth	Booth and Marchant, 1987
<i>Triparma strigata</i> Booth	Booth and Marchant, 1987
<i>Triparma verrucosa</i> Booth	Booth and Marchant, 1987

Adapted from Ichinomiya and Kuwata (2017) with permission.

or *Pentalamina*) that have not yet been isolated in cultures.

Diversity Within the Genus *Triparma*

We explored the level of inter- and intra-clade diversity within the genus *Triparma* by analyzing the folding pattern of the ITS2 (see **Supplementary Material** for Methodology) from 14 strains previously described in Ichinomiya et al. (2016). The general ITS2 secondary structure of Bolidophyceae proposed contains the four-helices domains known in many eukaryotic taxa in addition

to helix B9 (**Figure 3**). We located in Helices II and III the universal hallmarks proposed by Mai and Coleman (1997) and Müller et al. (2007): the pyrimidine-pyrimidine (Y-Y) mismatch in helix II and YRRY (pyrimidine – purine – pyrimidine) motif on the 5' side of Helix III, respectively, at alignment positions 95 and 148 and between nucleotides 205 and 208 (**Figures 3, 4**). In all strains analyzed, the Y-Y mismatch was represented by the pair U x C, with the exception of *T. pacifica* strains (U x U), and the YRRY motif of helix III by the sequence UGGU (**Figure 3**).

TABLE 2 | List of metabarcoding studies using the V4 region of the 18S rRNA genes that have been used for the phylogenetic (**Figure 2**) and the biogeography analyses (**Figures 6, 7**).

Data set	Region	Samples #	Bioproject	Sequencer	Clustering	Reference
OSD – LGC – 2014	Ocean	157	PRJEB8682	Illumina	0.97	Kopf et al., 2015
MALINA – Monier 2014	Arctic Ocean	24	PRJNA202104	454	0.98	Monier et al., 2013, 2014
ACME – Comeau – 2011	Arctic Ocean	11	SRA029114	454	0.98	Comeau et al., 2011
Nansen Basin – Metfies – 2016	Arctic Ocean	17	PRJEB11449	454	0.97	Metfies et al., 2016
Southern Ocean – Wolf – 2014	Southern Ocean	6	PRJNA176875	454	0.97	Wolf et al., 2014
Fildes Bay – Luo – 2016	Southern Ocean	10	PRJNA254097	Illumina	0.97	Luo et al., 2015
Fram Strait – Kilius – 2013	Arctic Ocean	5			454	Kilius et al., 2013

See *Supplementary Material* for details.

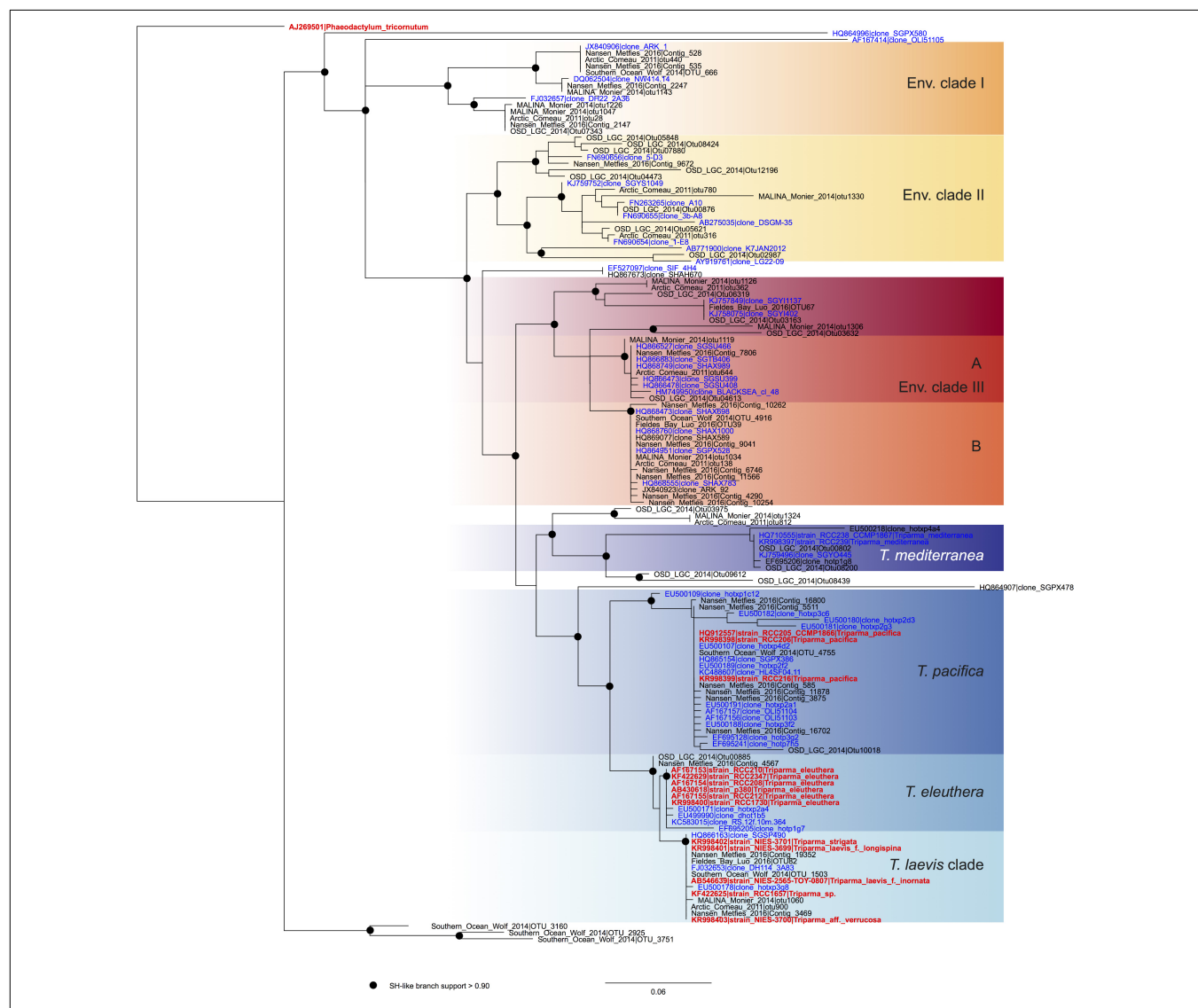


FIGURE 2 | ML phylogenetic tree based on the V4 region of the 18S rRNA gene based both on GenBank sequences available from the PR² database (Guillou et al., 2013) and on metabarcodes OTUs obtained from the studies listed in **Table 2** (see Methodology in **Supplementary Material**). The tree was constructed with phyML and dots correspond to nodes with SH-like branch support > 0.90. GenBank sequences from environmental samples are colored in blue and those from cultures, in red.

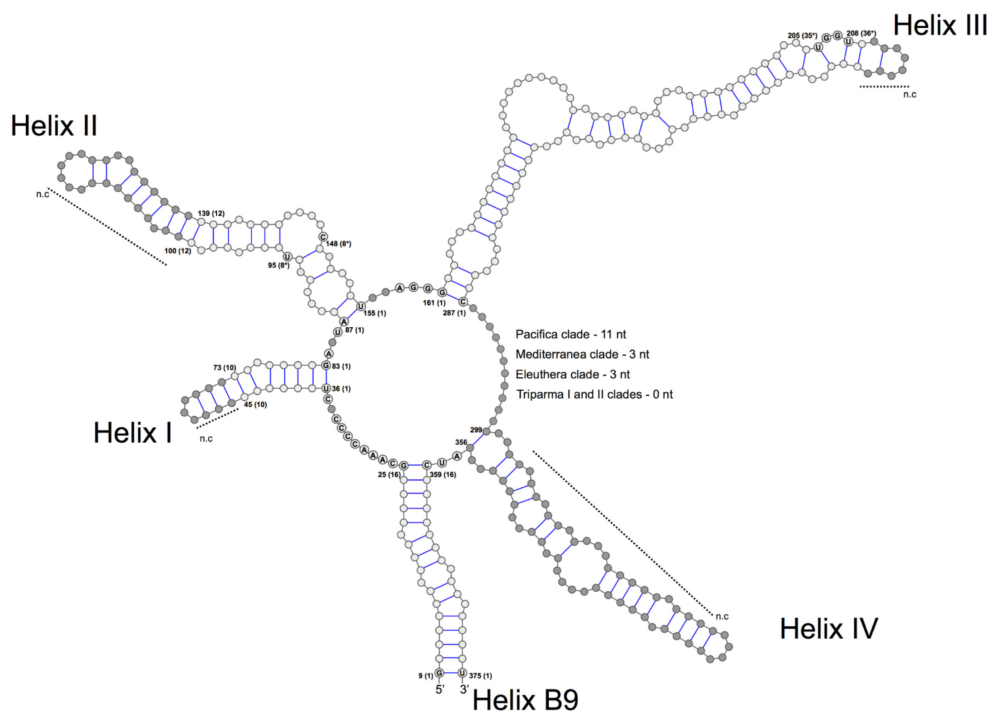


FIGURE 3 | General structure model of the ITS2 molecule of the *Triparma* clades. The four major helices are labeled as Helix I – Helix IV and the interaction region of 5.8S and 28S rRNA as B9. Numbers refer to the alignment positions and those between brackets to the compared positions in each helix. The nucleotides that are 100% conserved in the helices spacers, hallmarks positions in helices II and III and first two base pairs of the helices are shown. The gray dots represent segments that display length and sequence variation (see Methodology in **Supplementary Material**).

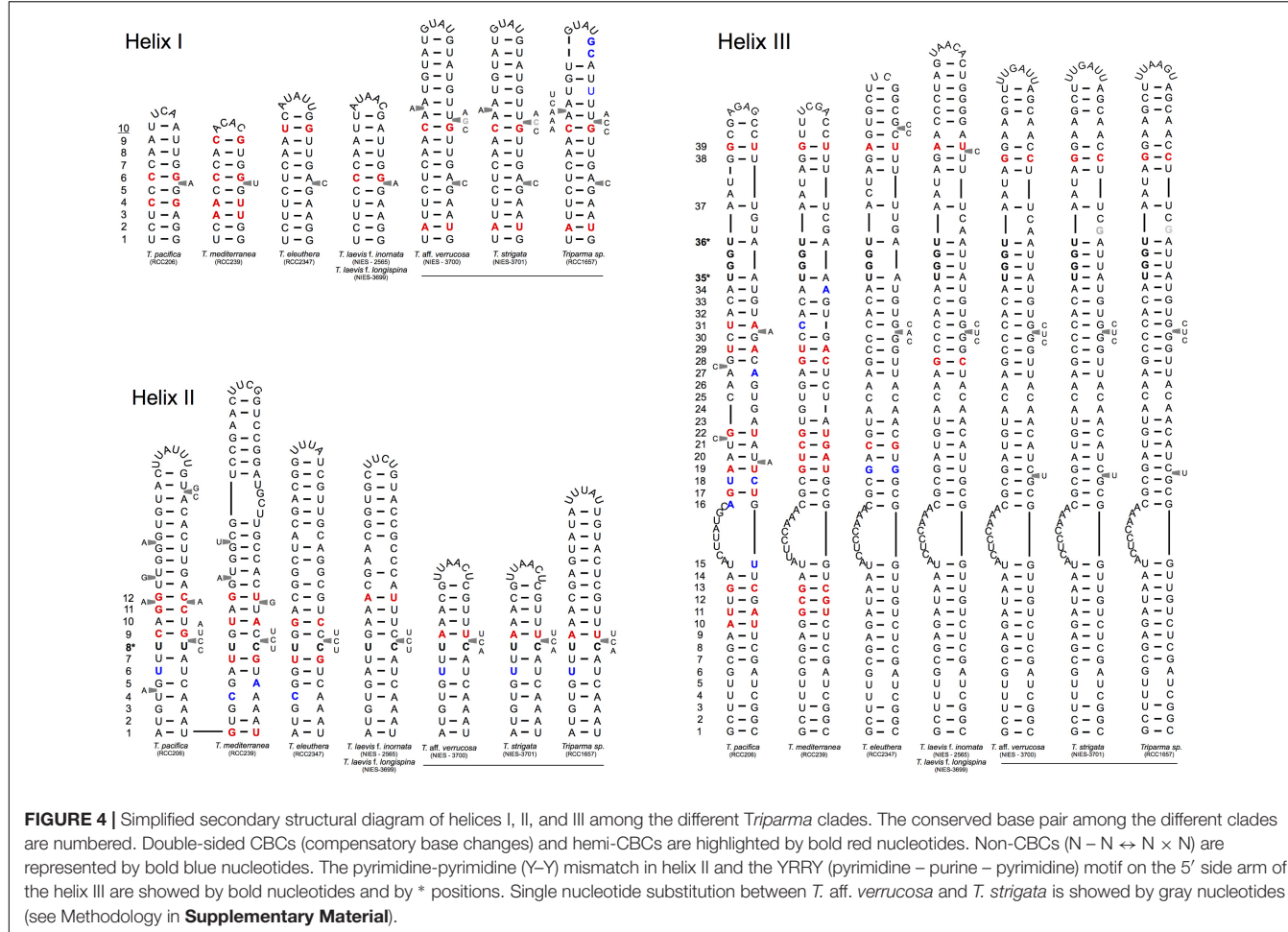
The spacers between helices B9 and I, I and II, II and III, and IV and B9 were conserved in length and sequence among the clades (Figure 3), as well as the first two base pairs of helices I, II, and III. In contrast, the spacer between helices III and IV showed greater variation between Bolidophyceae clades but it was conserved at the intra-clade level (Figure 3). Helices B9 (a region of the 5.8S and 28S rRNA interaction) and III showed good intra and inter-clade conservation (Figure 3 and Supplementary Figure 1). The ITS2 sequence from *T. aff. verrucosa* is incomplete and the 3' side arm of helix B9 could not be determined (Supplementary Figure 1).

The identification of CBCs in Bolidophyceae ITS2 secondary structure was based on the phenetic approach which relies on a base pair sequence comparison of all CBCs between two sequences without direct reference to their evolutionary origin (Müller et al., 2007; Coleman, 2009). The phylogenetic approach method which considers the status of a given base pair in the ancestor of two sister taxon could not be applied for Bolidophyceae given the conflicting branching pattern among phylogenies (for more details see Ichinomiya et al., 2016).

Putative CBCs, hCBCs, and non-CBCs type changes were identified in the conserved regions of the helices B9, I, II, and III within each clade and between clades (Figure 4 and Supplementary Figure 1). Helix IV (Figure 4) was not included in the inter-clade analysis given its known variable nature (Coleman, 2007). Several CBCs and hCBCs were identified at inter-clade level suggesting that each clade within *Triparma*

genus (*T. pacifica*, *T. mediterranea*, *T. eleuthera*, *Triparma* I and II, *sensu* Ichinomiya et al., 2016) is composed by at least one species (Figure 4).

At the intra-clade level, no CBC, nor hCBCs were identified between the two forms of *T. laevis*, f. *inornata* and f. *longispina* (*Triparma* clade I), that differ by the plate morphology, suggesting that these two forms may belong to the same species, although the absence of CBCs is not an absolute indicator that two organisms belong to the same species (Müller et al., 2007; Caisová et al., 2011, 2013). However, at least one CBC is a good indicator (93.1% of confidence for plants and fungi) that in most of the cases, two organisms represent distinct species (Müller et al., 2007). For *Triparma* clade II (*T. aff. verrucosa*, *T. strigata*, and *Triparma* sp. RCC1657), no CBC or hCBCs were identified between the morpho species *T. aff. verrucosa* and *T. strigata*. The ITS operon sequences from these two strains, including the two internal transcribed spacers and 5.8S rRNA, are nearly identical (99.1%) differing only by six nucleotides. Although three of these substitutions are within the ITS2, none correspond to a nucleotide pair in the ITS2 secondary structure (Figure 4). However, between *Triparma* sp. RCC1657 on the one side and *T. aff. verrucosa* plus *T. strigata* on the other side, 1 hCBC (helix B9, position 6, Supplementary Figure 1) and 1 CBC (helix IV, box, Supplementary Figure 1) were identified, suggesting that *Triparma* clade II is composed by at least two species, one corresponding to *Triparma* sp. RCC1657 and the other by *T. aff. verrucosa* and *T. strigata*.



ECOLOGY

Oceanic Distribution

We explored the distribution of Bolidophyceae in the ocean using data obtained by SEM observation, environmental sequencing and metabarcoding. By compiling available records of observed silicified morphological species of Bolidophyceae in natural waters (**Supplementary Table 1**), we mapped the geographical and specific distribution pattern of each morphological species (**Figure 5**). *Tetraparma pelagica*, and the *Triparma* species, *T. laevis*, *T. columnacea*, *T. retinervis*, and *T. strigata* are widely distributed from polar to subtropical regions. In contrast, *Pentalamina corona*, *Tetraparma gracilis*, *Tetraparma catinifera*, and *Triparma verrucosa* are restricted to polar or subpolar regions. *T. gracilis* was observed in both, Arctic and Antarctic regions while *P. corona* seems endemic to the Antarctic and, *T. verrucosa* and *T. catinifera* to the subarctic region. *Tetraparma insecta* and the recently described species *Tetraparma silverae* and *Tetraparma trullifera* seem to be restricted so-far to subtropical regions.

Using both available environmental GenBank sequences and 18S rRNA V9 metabarcodes acquired during the Tara Oceans expedition, Ichinomiya et al. (2016) established the

oceanic distribution of the major *Triparma* species and of environmental clades. *T. mediterranea* metabarcodes dominated in the Mediterranean Sea while *T. pacifica* and *T. eleuthera* were co-dominant in the tropical and sub-tropical oceans. The *T. laevis* clade was clearly associated with cold Antarctic waters but was also found near the Costa Rica dome. Bolidophyceae sequences were most abundant in the picoplanktonic fraction (0.8–5 μm) of the Tara Oceans samples and represented at most 4% of the photosynthetic reads and less than 1% on average (Ichinomiya et al., 2016).

In order to obtain a more complete image of the Bolidophyceae distribution, we used the large data set of 18S rRNA V4 metabarcodes described above. This data set includes a range of studies (**Table 2**) including OSD (Ocean Sampling Day) that sampled an extensive set of coastal stations (Kopf et al., 2015) and several from Arctic and Antarctic waters that were not covered by the Tara expedition.

Among these metabarcodes, the *Triparma* clade was slightly dominating in terms of total reads followed by the three environmental clades III, I, and II, respectively, in this order (**Figure 6A**). Within *Triparma*, *T. pacifica* was most abundant followed by *T. mediterranea*. One environmental subclade (IIIA) was also particularly abundant. The relative contribution of

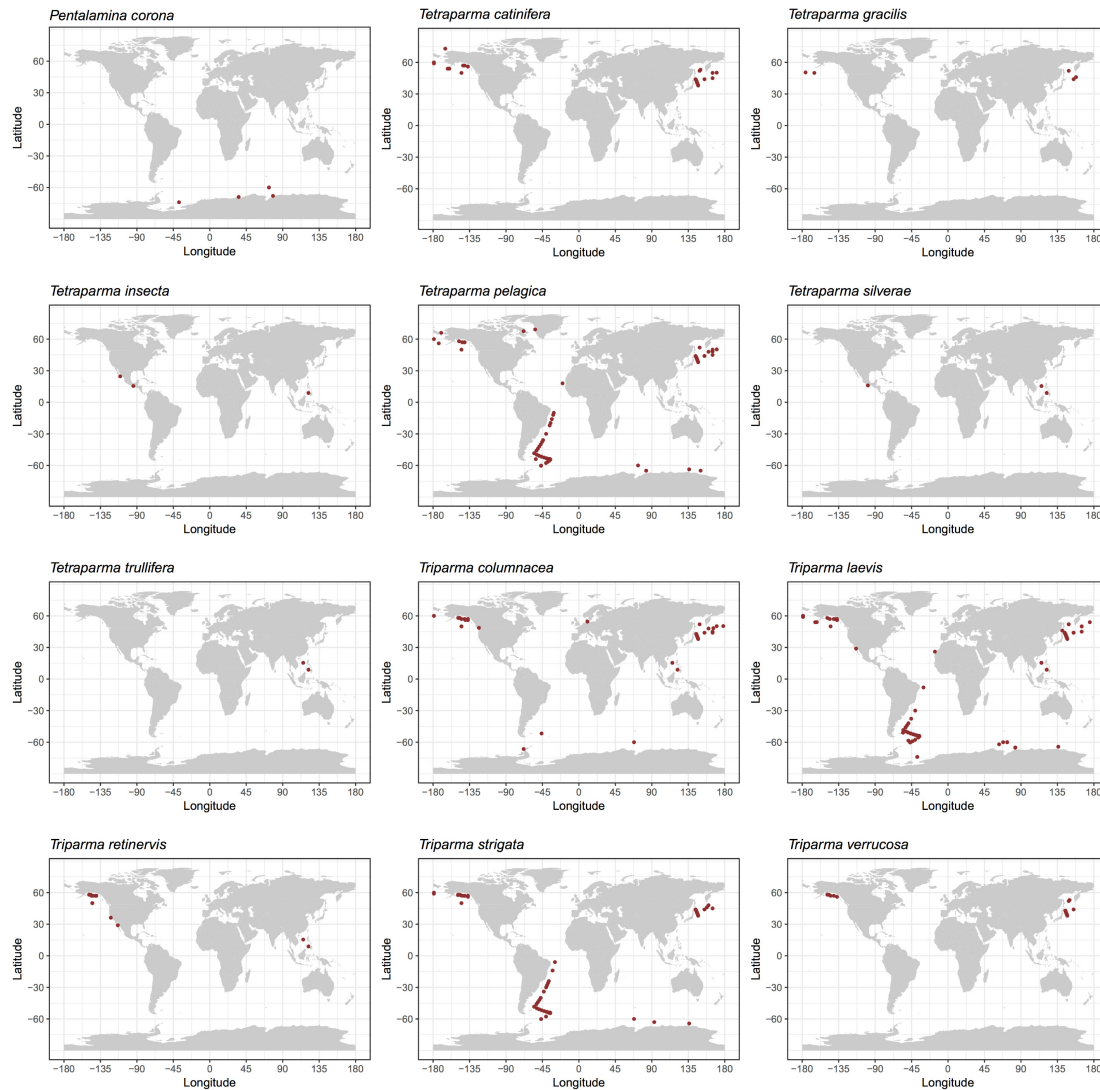


FIGURE 5 | Distribution of silicified Bolidophyceae species based on literature records of SEM observations (see **Supplementary Table 1**).

Bolidophyceae to total metabarcodes at each station varied widely with an average of 0.23% (**Figure 6B**). It was highest, up to 12%, in both Arctic and Antarctic regions as well as around the European coast. In contrast Bolidophyceae were absent at several stations along the East coast of North America and in the Eastern Mediterranean Sea (**Figure 6C**).

The distribution of individual *Tripelta* species and environmental clades confirmed some of the trends observed in the Tara Oceans data (Ichinomiya et al., 2016) but also revealed new features (**Figure 7**). Among the *Tripelta* species, *T. eleuthera* and *T. pacifica* were the most ubiquitous and often co-occurred at the same stations, suggesting that their ecological niches are very close. They did not seem to be present though in really polar waters such as in the Beaufort Sea. In contrast, it was confirmed that *T. mediterranea* was indeed mostly restricted to the Mediterranean Sea while the *T. laevis* clade was only found at high latitudes both in the

Arctic and Antarctic. Some environmental clades had clear biogeographic distributions such as clade IIIA found mostly in temperate latitudes and IIIB only in the Arctic and Antarctic regions. The latter clade seemed particularly prevalent in the high Arctic Ocean. Clade I was also mostly observed at high latitudes, although not restricted to polar waters, in contrast to clade II which was more widespread. These distribution patterns may reflect the genetic diversity within these clades. Clades IIIA and B have very low genetic diversity (**Figure 2**) in contrast clades I or II. The former may therefore correspond to a single species with a narrow niche and the latter to several species or even genera, explaining their wide distribution.

Seasonal Cycle

Ichinomiya and Kuwata (2015) investigated the seasonal influence in the abundance and vertical distribution of the

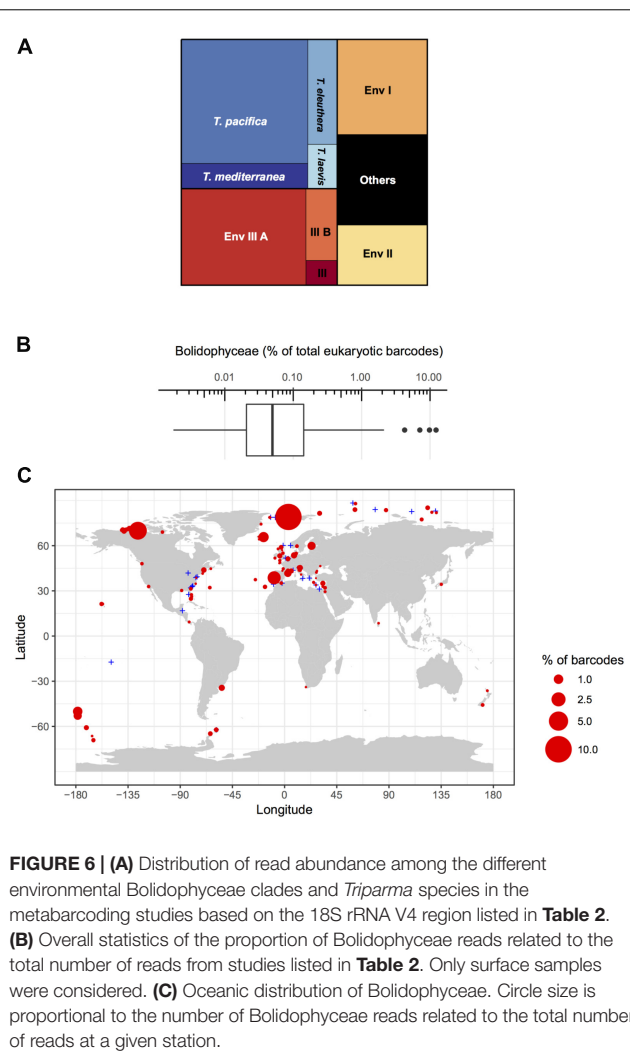


FIGURE 6 | (A) Distribution of read abundance among the different environmental Bolidophyceae clades and *Triparma* species in the metabarcoding studies based on the 18S rRNA V4 region listed in **Table 2**.

(B) Overall statistics of the proportion of Bolidophyceae reads related to the total number of reads from studies listed in **Table 2**. Only surface samples were considered. **(C)** Oceanic distribution of Bolidophyceae. Circle size is proportional to the number of Bolidophyceae reads related to the total number of reads at a given station.

silicified forms of Bolidophyceae in the western North Pacific using SEM. The area investigated is surrounded by the cold Oyashio current with water temperature below 5–8°C at 100 m (Shimizu et al., 2009). The Bolidophyceae community was mainly composed of *T. laevis* ($64 \pm 22\%$) with only small regional and seasonal differences in contrast to diatoms that display clear seasonality patterns (Takahashi et al., 2008; Suzuki et al., 2011). The vertical distribution of the silicified Bolidophyceae community changed seasonally according to the hydrographic condition. Silicified Bolidophyceae had a wide vertical distribution between 0 and 100 m with high abundance of $10\text{--}10^2$ cells mL^{-1} in March and May at stations where the water column was well mixed or weakly stratified (Figure 8). In contrast, from May to October at stations where the water was stratified Bolidophyceae were absent from the surface layer, but mainly distributed under the pycnocline from 20 to 50 m with lower abundance of $<0.1\text{--}10$ cells mL^{-1} . Komuro et al. (2005) also reported similar seasonal variations in depth distributions of the silicified form at Station KNOT (44°N , 155°E) in the western North

Pacific. They extended from 0 to 100 m in January and May, but were restricted to the subsurface layer from 30 to 100 m in August. Silicified Bolidophyceae have optimal growth temperatures below 10°C, but do not grow above 15°C (see section “Cell Physiology”). These data suggest that silicified Bolidophyceae actively grow during the cold mixing season and maintain their population under the pycnocline during the warm stratified season (Figure 9). Flagellated forms of Bolidophyceae may also be present in the surface layer during stratification since they have been reported in the surface layer during the summer season in the English Channel (Not et al., 2002) and northern South China Sea (Wu et al., 2017) using 18S rRNA-targeted oligonucleotide probes specific of Bolidophyceae detected by *in situ* hybridization and tyramide signal amplification (FISH-TSA). 18S rDNA sequences of Bolidophyceae have also been detected using high-throughput sequencing (Kataoka et al., 2017) at 10 m in summer and autumn in the Oyashio region when silicified forms were absent.

Role in Food Webs

It is not clear how Bolidophyceae contribute to the microbial food web. Materials resembling silicified Bolidophyceae have been reported in fecal pellets of copepods (Booth et al., 1980; Urban et al., 1993) and Antarctic krill (Marchant and Nash, 1986), indicating that they can be grazed by larger predators (Kosman et al., 1993). Bolidophyceae main grazers are expected to be small protozoans, such as choanoflagellates (Taniguchi et al., 1995) although there is no evidence of their direct ingestion by protists.

CELL PHYSIOLOGY

Temperature

Ichinomiya et al. (2013) and Ichinomiya and Kuwata (2015) conducted growth experiments at various temperature ranging from 0 to 15°C, using three silicified strains: *T. laevis f. inornata*, *T. laevis f. longispina*, and *T. strigata*. These silicified Bolidophyceae species were able to grow at 0 to 10°C (*T. laevis f. inornata* and *T. strigata*) and 5 to 10°C (*T. laevis f. longispina*) but not over 15°C (Figure 10). The optimal growth temperatures were 5°C for *T. laevis f. inornata* and 10°C for *T. laevis f. longispina* and *T. strigata*, with growth rates of 0.35 d^{-1} , 0.50 d^{-1} , and 0.69 d^{-1} , respectively. In contrast, strains of naked flagellated forms have higher growth rates and grow at higher temperatures (Figure 10). *T. eleuthera* showed positive growth at 16–24°C with the maximum growth rate of 1.7 d^{-1} at 22°C (Stawiarski et al., 2016) while *T. pacifica* and *Triparma* sp. RCC201 had growth rates of 0.91 d^{-1} and 0.51 d^{-1} at 20°C, respectively (Jacquet et al., 2001; Thomas and Campbell, 2013).

Silica

The growth of diatoms is limited by dissolved silicate (Martin-Jézéquel et al., 2000; Sarthou et al., 2005). Diatoms cell cycle is controlled by silica and silica limitation arrests cells at the $G_1\text{-}S$ boundary (Darley and Volcani, 1969; Okita and Volcani,

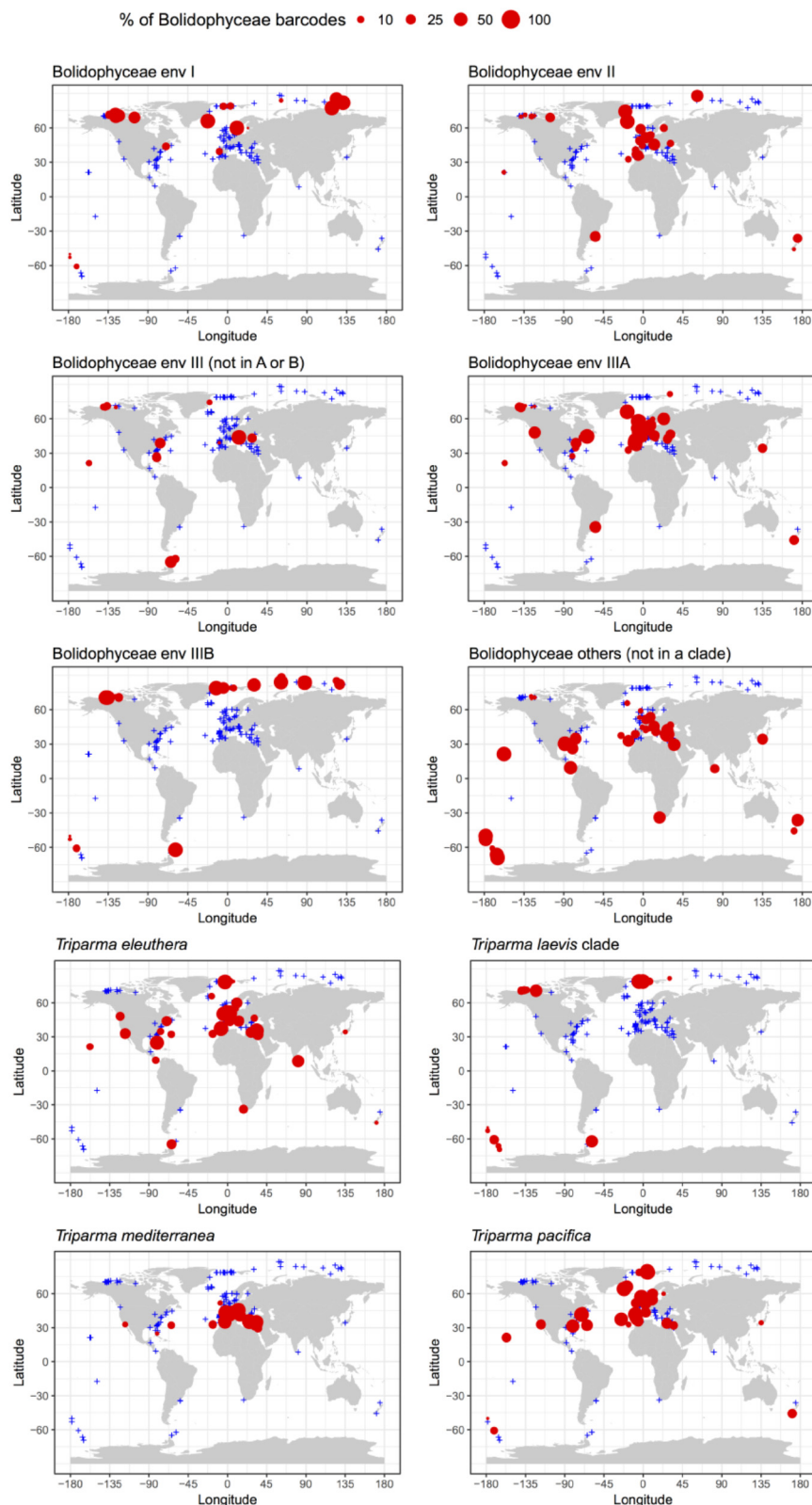


FIGURE 7 | Percentage relative to the total number of Bolidophyceae reads of the different environmental Bolidophyceae clades and *Triparma* species in the metabarcoding studies based on the 18S rRNA V4 region listed in **Table 2**. Only surface samples were considered.

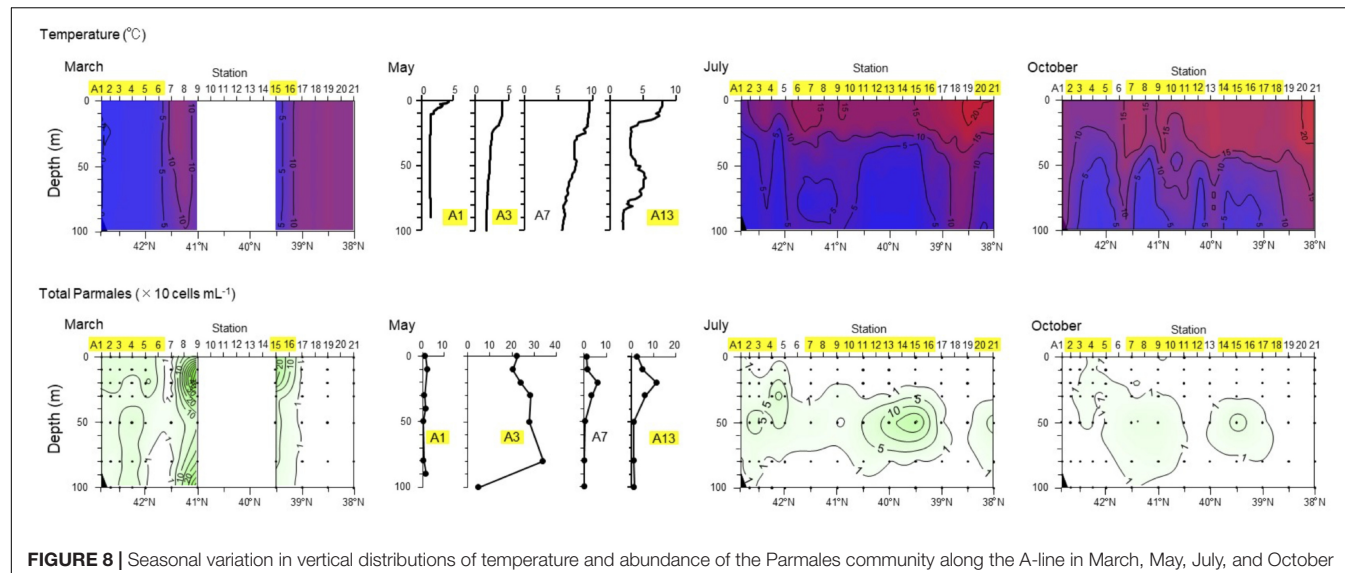


FIGURE 8 | Seasonal variation in vertical distributions of temperature and abundance of the Parmales community along the A-line in March, May, July, and October 2009. Station numbers shaded in yellow indicate the Oyashio region. Redrawn from Ichinomiya and Kuwata (2015) with permission.

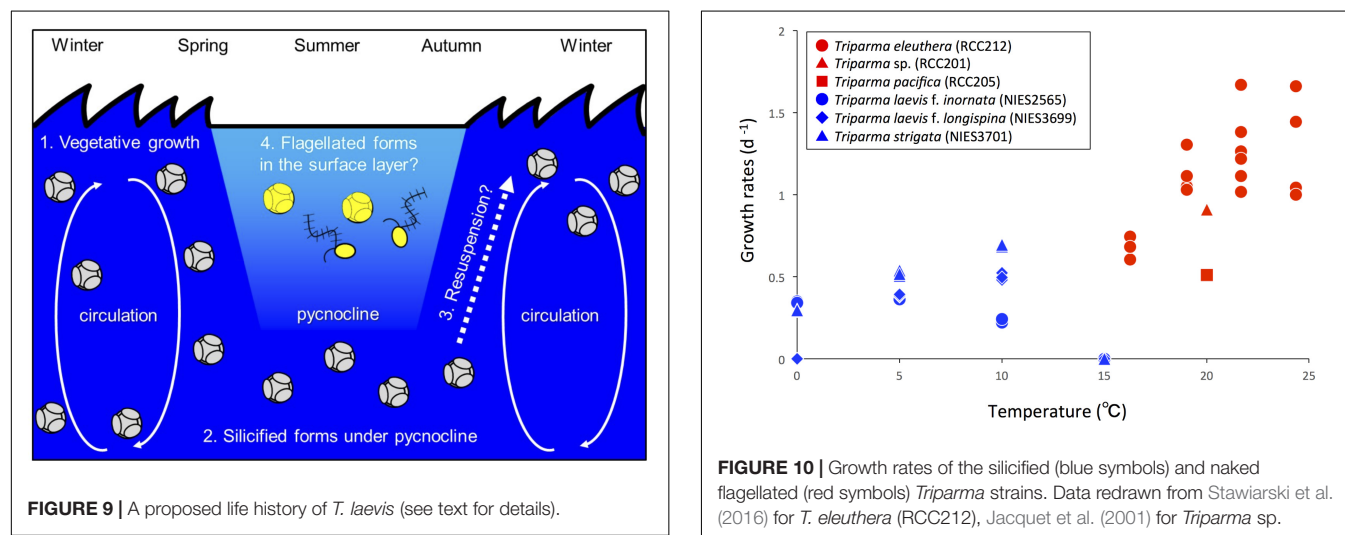


FIGURE 9 | A proposed life history of *T. laevis* (see text for details).

1978; Vaulot et al., 1987) and during the G2–M transition due to silica requirement for DNA replication and cell wall formation, respectively (Vaulot et al., 1987; Brzezinski et al., 1990).

In contrast, Bolidophyceae despite possessing silica plates can grow in the absence of silica (Yamada et al., 2014). *T. laevis* f. *inornata* cells growing under sufficient silicate (100 μ M) are surrounded by eight plates, rounded shield and ventral plates, as well as non-rounded dorsal and girdle plates. However, plate formation becomes incomplete and the fraction of cells lacking dorsal and girdle plates increases at low silicate concentration (10 μ M). Cells finally loose almost all plates at silicate concentrations lower than 1 μ M (Yamada et al., 2014). Other silicified Bolidophyceae strains, *T. laevis* f. *longispina* and *T. strigata*, can also grow under silicate depletion without formation of a silica cell wall (unpublished data). Cell wall

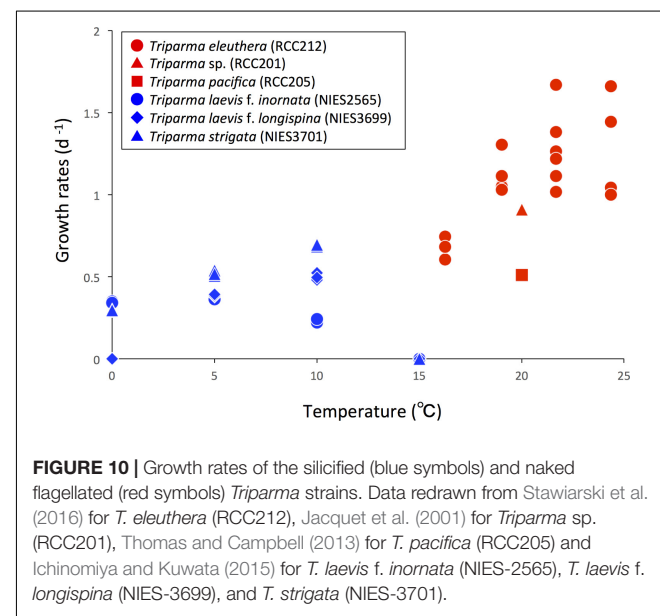


FIGURE 10 | Growth rates of the silicified (blue symbols) and naked flagellated (red symbols) *Triparma* strains. Data redrawn from Stawiarski et al. (2016) for *T. eleuthera* (RCC212), Jacquet et al. (2001) for *Triparma* sp. (RCC201), Thomas and Campbell (2013) for *T. pacifica* (RCC205) and Ichinomiya and Kuwata (2015) for *T. laevis* f. *inornata* (NIES-2565), *T. laevis* f. *longispina* (NIES-3699), and *T. strigata* (NIES-3701).

is restored within a day in about 40% of the naked cells after replenishment of silicate (Yamada et al., 2014). Direct observation of regeneration of the silica cell wall in naked cells after re-supply of silicate using transmission and SEM revealed that shield plates appear first, followed by ventral, dorsal, and girdle plates, in this order. The dorsal and girdle plates are inserted into the space between the previously secreted shield and ventral plates to complete cell wall (Yamada et al., 2016). Similar uncoupling between the formation of silica structures and cell growth has also been observed in other silicified Stramenopiles such as Dictyochales (Henriksen et al., 1993) and Synurales (Leadbeater and Barker, 1995; Sandgren et al., 1996).

TABLE 3 | Mitotic characters of Stramenopiles.

	Bolidophyceae* ¹	Diatoms	Pelagophyceae	Phaeophyceae	Xanthophyceae	Chrysophyceae/ Synurophyceae	Raphidophyceae	Eustigmatophyceae
Interphase microtubules focus	Centrioles	Microtubule center	No report	Centrioles	Centrioles	Centrioles	Centrioles	No report
Spindle pole	Centrioles	Polar plate	Centrioles	Centrioles	Centrioles	Rhizoplast	Golgi body	barrel/boomerang-shaped nuclear pole body
Extranuclear spindle	+	+	+	-	-	-	-	-
Bundled spindle	+	+	+	-	-	-	-	-
References	Yamada et al., 2017 Pickett-Heaps et al., 1975; Pickett-Heaps, 1991	Pickett-Heaps et al., 1975; Vesk and Jeffrey, 1987	Markey and Wilce, 1975	Massalski et al., 2009	Slankis and Gibbs, 1972; Vesk et al., 1984; Brugerolle and Mignot, 2003		Heywood, 1978	Murakami and Hashimoto, 2009

*¹ Reported in only silicified strain *Tripelta laevis* NIES-2565.

Rounded plates of silicified Bolidophyceae have a structure similar to the valves and scales of auxospores from centric diatoms (Mann and Marchant, 1989; Yamada et al., 2014). In other groups that display silica structures such as diatoms, chrysophytes, synurophytes, and dictyochophytes (Simpson and Volcani, 1981; Knoll and Kotrc, 2015; Finkel, 2016; Marron et al., 2016), silica formation takes place within a specialized membrane-bound compartment termed the Silica Deposition Vesicle (SDV) (Simpson and Volcani, 1981; Preisig, 1994). The origin and location of SDV differ among taxa. Diatoms SDVs for the development of valve, girdle bands of vegetative cells and auxospores scales are formed adjacent to the plasma membrane (Stoermer et al., 1965; Edgar and Pickett-Heaps, 1984; Lee and Li, 1992; Idei et al., 2012), possibly generated from the Golgi body (Lee and Li, 1992). Synurophytes and chrysophytes SDVs are located in the cytoplasmic or chloroplast endoplasmic reticulum. Bolidophyceae SDVs forming the shield and ventral plates are initially produced around the chloroplast and moving toward the plasma membrane like synurophytes and chrysophytes (Yamada et al., 2016). In contrast, SDVs for dorsal and girdle plates are formed adjacent to the plasma membrane like in diatoms. Such differentiation in the development site of SDVs depending on the type of silica plates within a single species has not been previously reported in other organisms (Yamada et al., 2016).

Mitotic Nuclear Division

In eukaryotes, cell division, mitotic process, and related apparatus are often well conserved within high phylogenetic levels (e.g., at the class or phylum levels, Heath, 1980; Schmit and Nick, 2008; De Martino et al., 2009). Among Stramenopiles, the organelles related to the focus of interphase microtubules and spindle poles are the centrioles, like recently observed in Bolidophyceae (Yamada et al., 2017, **Table 3**). However, some classes have unique organelles (**Table 3**), such as the Microtubule Center (MC) and Polar Plate (PP) in diatoms (Pickett-Heaps et al., 1975; Tippit and Pickett-Heaps, 1977; Edgar and Pickett-Heaps, 1984; Pickett-Heaps, 1991; De Martino et al., 2009), the rhizoplast in chrysophytes and synurophytes (Slankis and Gibbs, 1972; Vesk et al., 1984; Brugerolle and Mignot, 2003) or a barrel/boomerang-shaped nuclear pole body in eustigmatophytes (Murakami and Hashimoto, 2009).

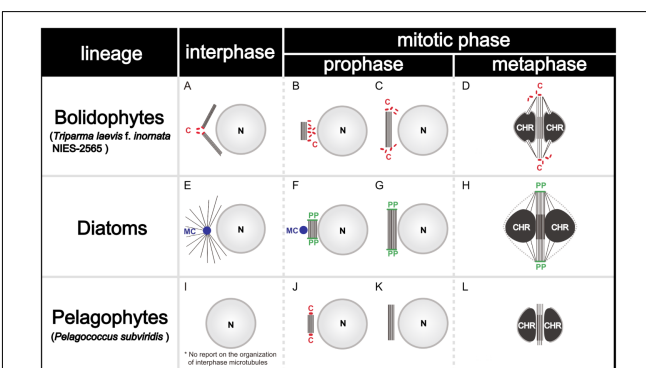


FIGURE 11 | Graphical scheme of the interphase microtubule nucleation and the spindle formation in bolidophytes (A–D), diatoms (E–H), and pelagophytes (I–L). See text for details. Figures (A–D) and (E–H) are adapted from Yamada et al. (2017) and De Martino et al. (2009) with permission.

During the mitosis of *T. laevis* f. *inornata*, the interphase cell has more than four very short centrioles (ca. 80 nm in contrast to 150–500 nm of typical mature centrioles in other Stramenopiles, (**Figure 11A**). In prophase, the spindle bundle forms in the extranuclear region (**Figure 11B**), the centrioles move to the spindle poles (**Figure 11C**) and then it moves to the cytoplasmic tunnel of the nucleus (**Figure 11D**). All along metaphase, the kinetochore microtubules elongate from the spindle poles to the condensed chromatin through the region of partially disintegrated nuclear envelope (**Figure 11D**). Finally, the chromatin is separated to both sides of the cell.

Spindle configuration and formation of *T. laevis* f. *inornata* are very similar to the process found among diatoms and pelagophytes (**Table 3**). They share two conspicuous characters: extranuclear spindle formation (**Figures 11B,C,E,G**) and the bundling of the interpolar microtubules (**Figures 11D,H**). However, the organelle serving as a Microtubule Organizing Center (MTOC) and its behavior differ. *T. laevis* f. *inornata* and pelagophytes have centrioles while diatoms have the specialized MC and PP (**Figures 11E–H**). The centrioles of pelagophytes (reported only in one species, *Pelagococcus subviridis*) appear

TABLE 4 | Comparison of selected characters between Bolidophyceae and diatoms.

Properties	Bolidophyceae		Diatoms
	Silicified species	Flagellated species	
Size (μm)	2–5	1–1.7	2–2000
Level of organization	Unicellular	Unicellular	Unicellular, often form colonies
Silicified cell wall	Yes	No	Yes
Flagellate form	Yes	Yes	Yes in male gametes of centric diatom
Chloroplasts	Lamellae with three thylakoids, girdle lamella		Lamellae with three thylakoids, girdle lamella
Major Pigments	Chl a, Chl c, fucoxanthin, diatoxanthin, diadinoxanthin, b-carotene		Chl a, Chl c, fucoxanthin, diatoxanthin, diadinoxanthin, b-carotene
Mitochondria	Tubular type		Tubular type
Si requirement for growth	No	No	Yes
Position of SDV	Chloroplast ER and plasma membrane	NA	Plasma membrane
Mitotic apparatus	Interphase microtubules focus:Centrioles Spindle pole:Centrioles	NA	Interphase microtubules focus:Microtubule center Spindle pole:Polar plate
Number of species	12	3	30,000–100,000
Oceanic distribution	Ubiquitous, but minor		Ubiquitous, often dominant
Main habitat	Cold eutrophic water (Polar and subpolar region)	Warm oligotrophic water (Tropical or subtropical)	Eutrophic water (Polar, coastal, and upwelling region)

NA, not available.

only during the spindle initiation phase (Vesk and Jeffrey, 1987) while those of *T. laevis f. inornata* occur after central spindle formation in the extranuclear region. Since centriole is the most common organelle serving as MTOC in Stramenopiles (Table 3), the mitotic apparatus of *T. laevis f. inornata* shows more ancestral features than the diatoms.

CONCLUSION – THE EVOLUTIONARY RELATIONSHIPS BETWEEN DIATOMS AND BOLIDOPHYCEAE

Diatoms are highly diverse with 30,000 to 100,000 species (Mann and Vanormelingen, 2013) and constitute one of the top group of primary producers, contributing to about 20% of the photosynthesis on Earth, the equivalent of terrestrial rainforests (Nelson et al., 1995; Falkowski et al., 1998; Mann, 1999). They cover a wide size range from 2 μm to 2 mm and form large blooms in high-nutrient coastal and upwelling systems (Margalef, 1978; Hasle and Syvertsen, 1997). They are the main prey for zooplankton and the carbon that they fix through photosynthesis is efficiently transferred to higher trophic levels highlighted by their role in fish production (Ryther, 1969; Cushing, 1989). From an evolutionary point of view, the appearance of this highly productive group and the resulting increase in oceanic primary production may have driven the evolution of crustaceans, pelagic fish, and whales and shaped modern marine pelagic ecosystems (Parsons, 1979). Although, the origin and early evolution of diatoms remain controversial, the first well-preserved diatom fossils have been dated from \sim 110 Myr ago, in the early Cretaceous (Gersonde and Harwood, 1990; Harwood and Gersonde, 1990), while

molecular-clock-based estimations suggest that the origin of diatoms may have occurred 180 – 250 Myr ago (Medlin, 2011).

Recent multigene phylogenetic analyses suggest that bolidophytes, diatoms, pelagophytes, and dictyochophytes form a monophyletic lineage (Riisberg et al., 2009; Yang et al., 2012; Ševělková et al., 2015; Derelle et al., 2016). This lineage, called Diatomista (Derelle et al., 2016) or Khakista (Riisberg et al., 2009), originally only included diatoms and bolidophytes (Cavalier-Smith and Chao, 2006).

Recent success in the isolation of strains of both silicified and naked flagellated Bolidophyceae species allow detailed phylogenetic studies, clarifying the taxonomic position of this group as a sister group of diatoms and revealing the close relationship between silicified and naked strains. Cell wall formation and mitotic division in the silicified species *T. laevis f. inornata* have intermediate features between diatom and more ancient stramenopiles (Figure 11 and Tables 3, 4). Analysis of organellar genomes of this species also suggested that it displays more ancestral characteristics than diatoms (Tajima et al., 2016). Gene contents of the plastid and mitochondrial genomes are similar between *T. laevis f. inornata* and diatoms whereas the gene order of the mitochondrial genome is different. The structure of the mitochondrial genome is also more compact in *T. laevis f. inornata* than in diatoms since the latter species has no introns or repeat regions which are often observed in some diatoms species (Oudot-Le Secq and Green, 2011).

The phylogenetically close relationship between silicified and naked Bolidophyceae strains and recent occasional observation of flagellated cells in cultures of *T. laevis f. inornata* (Ichinomiya et al., 2016) suggest that bolidophytes may have a life cycle that switches between silicified non-flagellated and naked

flagellate stages. This hypothetical life cycle has similarities to centric diatoms for which the diploid vegetative stage produces haploid naked flagellated cells (male gametes or spermatozoa) during sexual reproduction (Drebes, 1977; Vaulot and Chisholm, 1987). The origin, early evolution and key processes for the acquisition of the silica cell wall are not yet been fully understood and Bolidophyceae could play a key role in answering these questions. More comprehensive analyses, including whole genome sequences, observations of the life-cycle, and fossil records, would lead a deeper understanding of the evolutionary relationships between Bolidophyceae and diatoms.

AUTHOR CONTRIBUTIONS

DV analyzed GenBank and metabarcoding sequences and draw distribution maps. MT analyzed the OSD metabarcoding data. ALdS performed ITS secondary structure. MI and AK analyzed the distribution of silicified species. AK, KY, MI, SY, DV, and ALdS wrote the manuscript and contributed to the final editing.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fmars.2018.00370/full#supplementary-material> and <https://doi.org/10.6084/m9.figshare.5549638>.

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Bolidophyceae, a sister picoplanktonic group of diatoms
- a review

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Supplementary Material

Supplementary material is available from <https://doi.org/10.6084/m9.figshare.5549638>

Methodology

Analysis of 18S rRNA V4 region

Long (> 800 bp) Bolidophyceae 18S rRNA sequences deposited in GenBank were downloaded from the latest version of PR² (Guillou et al., 2013) available at https://github.com/vaulot/pr2_database/releases. For 18S V4 metabarcodes, OTU sequence, annotation and read abundance in each sample from the studies listed in **Erreur ! Source du renvoi introuvable.** were either downloaded from repository web sites or obtained directly from the authors. For all OTUs attributed to Bolidophyceae, a BLAST search was performed against PR2 to verify that indeed their closest match was a Bolidophyceae. If it was not the case, the OTU was not considered any further. The pcr.seqs function of Mothur (Schloss et al., 2009) was applied to extract the V4 region of GenBank and OTU sequences using eukaryotic V4 primers (Piredda et al., 2016) with 2 mismatches allowed on each sides. Environmental GenBank and metabbarcode OTUs V4 sequences were dereplicated using the –derep_prefix command of vsearch (Rognes et al., 2016) that selects the longest sequence among identical ones yielding a final set of 141 V4 sequences. Sequences were aligned with MAFFT and a tree was built using PhyML as implemented in Geneious version 10.0.7 (Kearse et al., 2012). Based on this tree, metabarcodes were assigned to one of the environmental clade (see main text) or to one of the genera for which sequences from cultures are available. The alignment is available as Supplementary Material 1.

The relative abundance of Bolidophyceae reads compared to total eukaryotic reads and that of each Bolidophyceae phylogenetic group compared to total Bolidophyceae reads was computed for each

metabarcode sample using R dplyr library. Data were plotted with the following R libraries : treemap, ggplot2 and rworldmap.

ITS2 secondary structure

The use of secondary structure of ITS2 (Internal Transcribed Spacer) in microalgal taxonomy became popular after Coleman et al. (2000) and Muller et al. (2007) suggested a link between the presence of compensatory base changes (CBCs) and species boundaries. Despite the primary sequence and length of the ITS2 vary among different taxa, its secondary structure is conserved as it retains features that are important for its biological function and thought to be universal among eukaryotes (Coleman, 2000, 2009; Joseph et al., 1999; Mai and Coleman, 1997). A double-sided base change of a nucleotide pair retaining the secondary structure is considered a CBC, while a single-side change is called hemi-CBC (hCBC). Other changes, for example N – N ↔ N x N, are considered as non-CBC.

Bolidophyceae ITS2 general folding pattern proposed is based on fourteen ITS2 sequences (accession numbers: MG310179-MG310187) from the strains described at Ichinomiya et al. (2016). The 5.8 and 28S rRNA flanking regions, ITS2 boundaries, were annotated using Hidden Markov Models (HMMs, Keller et al., 2009) and an Eukaryote database as implemented in the ITS2 database annotation tool with the default parameters (<http://its2.bioapps.biozentrum.uni-wuerzburg.de/>, Ankenbrand et al., 2015). RNA secondary structure predictions were performed using the Mfold web interface (Seibel et al., 2008) under the default options with the folding temperature fixed at 37°C, resulting in multiple alternative folding patterns per sequence. The preliminary structure for each sequence was chosen based on the presence of previously defined ITS2 hallmarks defined by Coleman (Coleman, 2000, 2007, 2009; Mai and Coleman, 1997) and similarities among the other structures found within and between the clades. Bolidophyceae ITS2 sequences were aligned with MAFFT using the G-INS-i algorithm (Katoh et al., 2002) and for each sequence within the alignment, the preliminary secondary structure annotated in dot-bracket format was associated, generating a Vienna file which

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was imported to 4SAGE (Seibel et al., 2006, 2008). The resultant alignment and structures were manually edited through extensive comparative analysis of each position (nucleotide) in sequences from the same clade and between clades. The resulting consensus intramolecular folding pattern (secondary structure) for Choropicophyceae was drawn using CorelDRAW X7. The final alignment with the secondary structures in Vienna format is available as Supplementary Material 2.

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Supplementary Files

Supplementary Material 1: Alignment of 18S rRNA V4 sequences from both GenBank and metabarcoding studies, used to build the tree of Figure 2.

Supplementary Material 2: The final ITS2 alignment with the secondary structures in Vienna format.

Supplementary Figure

Helix B9

	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>
16	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>
15	<i>U - A</i>	C - G	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>
14	<i>C - G</i>	<i>C - G</i>	<i>C - G</i>	<i>C - G</i>	<i>C - G</i>	<i>C - G</i>
13	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>
12	<i>G - U</i>	<i>G - U</i>	<i>G - U</i>	<i>G - U</i>	<i>G - U</i>	<i>G - U</i>
11	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>
10	<i>A - U</i>	<i>A - U</i>	<i>A - U</i>	<i>A - U</i>	<i>A - U</i>	<i>A - U</i>
9	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	<i>G - C</i>	G - G
8	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>
7	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>
6	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	U - G
5	<i>A - U</i>	<i>A - U</i>	<i>A - U</i>	<i>A - U</i>	<i>A - U</i>	<i>A - U</i>
4	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>
3	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>	<i>U - A</i>
2	<i>C - G</i>	<i>C - G</i>	<i>C - G</i>	<i>C - G</i>	<i>C - G</i>	<i>C - G</i>
1	<i>G - U</i>	<i>G - U</i>	<i>G - U</i>	<i>G - U</i>	<i>G - U</i>	<i>G - U</i>
	<i>T. pacifica</i> (RCC206)	<i>T. mediterranea</i> (RCC239)	<i>T. eleuthera</i> (RCC2347)	<i>T. laevis f. ornata</i> (NIES- 2565)	<i>T. strigata</i> (NIES-3699)	<i>Triparma sp.</i> (RCC1657)

Helix IV

Supplementary Figure 1 : Simplified secondary structural diagram of helices B9 and IV among the different *Triparma* clades. The conserved base pairs among the different clades are numbered. Double-sided CBCs (compensatory base changes) and hemi-CBCs are highlighted by bold red nucleotides. Non-CBCs ($N - N \leftrightarrow N \times N$) are represented by bold blue nucleotides.

Supplementary Table

Supplementary Table 1: List of published record for morphological species. Available as an Excel file.