The Genetic Diversity of *Mesodinium* and Associated Cryptophytes

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Ciliates from the genus *Mesodinium* are globally distributed in marine and freshwater ecosystems and may possess either heterotrophic or mixotrophic nutritional modes. Members of the *Mesodinium major/rubrum* species complex photosynthesize by sequestering and maintaining organelles from cryptophyte prey, and under certain conditions form periodic or recurrent blooms (= red tides). Here, we present an analysis of the genetic diversity of *Mesodinium* and cryptophyte populations from 10 environmental samples (eight globally dispersed habitats including five *Mesodinium* blooms), using group-specific primers for *Mesodinium* partial 18S, ITS, and partial 28S rRNA genes as well as cryptophyte large subunit RuBisCO genes (*rbcL*). In addition, 22 new cryptophyte and four new *M. rubrum* cultures were used to extract DNA and sequence *rbcL* and 18S-ITS-28S genes, respectively, in order to provide a stronger phylogenetic context for our environmental sequences. Bloom samples were analyzed from coastal Brazil, Chile, two Northeastern locations in the United States, and the Pribilof Islands within the Bering Sea. Additionally, samples were also analyzed from the Baltic and Barents Seas and coastal California under non-bloom conditions. Most blooms were dominated by a single *Mesodinium* genotype, with coastal Brazil and Chile blooms composed of *M. major* and the Eastern USA blooms dominated by *M. rubrum* variant B. Sequences from all four blooms were dominated by *Teleaulax amphioxeia*-like cryptophytes. Non-bloom communities revealed more diverse assemblages of *Mesodinium* spp., including heterotrophic species and the mixotrophic *Mesodinium chamaeleon*. Similarly, cryptophyte diversity was also higher in non-bloom samples. Our results confirm that *Mesodinium* blooms may be caused by *M. major*, as well as multiple variants of *M. rubrum*, and further implicate *T. amphioxeia* as the key cryptophyte species linked to these phenomena in temperate and subtropical regions.

**Keywords:** *Mesodinium*, *Teleaulax*, cryptophytes, ciliates, acquired phototrophy, mixotrophy, red tides, ciliate genetic diversity
INTRODUCTION

Species belonging to the genus *Mesodinium* are among the most widely distributed and abundant marine ciliates in coastal and estuarine ecosystems (Leppanen and Bruun, 1986; Sanders, 1995; Bell and Laybourn-Parry, 1999). Red water blooms of *M. rubrum*-like ciliates (= Myrionecta rubra) have been recorded since Darwin's journey on the Beagle (Darwin, 1906), and are recurrent features in many coastal ecosystems (Lindholm, 1978; Crawford et al., 1997; Herfort et al., 2011a). While *M. rubrum* has historically been reported as a single species from numerous global locations (Taylor et al., 1971), cryptic diversity has been suspected within the *M. rubrum* morphospecies for some time, due primarily to variability in cell size (Leegaard, 1920; Lindholm, 1985; Rychert, 2004). Previously, only one environmental study of the genetic diversity of the Mesodiniidae has been published, and it focused on the *Mesodinium rubrum/major* complex within the Columbia River Estuary (Herfort et al., 2011b). Phylogenetic analysis of rRNA genes spanning the internally transcribed spacer region (ITS) have demonstrated that *M. rubrum* is actually a species complex, composed of at least six major clades (Herfort et al., 2011b; Garcia-Cuetos et al., 2012). One of these clades was described as a new species, *M. major*, based on molecular and ultrastructural characteristics, and is larger and has more plastids than *M. rubrum* (Garcia-Cuetos et al., 2012). Additionally, a new mixotrophic species, *M. chameleon*, was described from primarily benthic habitats, with unique cilia/kinetid structures as well as plastid type and organization of cryptophycean organelles (Moestrup et al., 2012).

Cryptophytes are known to be important components of phytoplankton communities in coastal ocean ecosystems, especially in low light, estuarine, or high latitude environments (Buma et al., 1992; Mallin, 1994; Adolf et al., 2006). While little is known regarding temporal or spatial trends of cryptophyte diversity, *Teleaulax*, *Plagioselmis*, and *Hemiselmis* are commonly encountered in marine environments (Hill et al., 1992; Cerino and Zingone, 2007; Metfies et al., 2010). Despite the well-established connection between *Teleaulax*-like cryptophytes and growth of *M. rubrum* (Gustafson et al., 2000; Johnson and Stoecker, 2005; Smith and Hansen, 2007), relatively few studies have documented their relationship in nature. Single cell PCR of *M. rubrum* from both coastal Japan and the Columbia River Estuary, have revealed predominanlty *Teleaulax amphioxeia* plastids (Nishitani et al., 2010; Herfort et al., 2011b).

All photosynthetic *Mesodinium* spp. are thought to harbor only cryptophyte organelles, which they acquire through feeding on free-living prey (Gustafson et al., 2000). When acquiring organelles from cryptophyte prey, *M. rubrum*-like ciliates also retain mitochondria, cytoplasm, and the nucleus (Johnson, 2011). The nucleus, referred to as a kleptokaryon, remains transcriptionally active and appears to facilitate functional control and division of stolen organelles (Johnson et al., 2007; Lasek-Nesselquist et al., 2015). However, relatively few strains of the *M. major/rubrum* complex have been successfully cultured and studied in detail. *Mesodinium* spp. are well-documented to form blooms in numerous coastal regions, particularly in estuaries (Crawford et al., 1997; Herfort et al., 2011a) and coastal upwelling zones (Ryther, 1967; Packard et al., 1978). Blooms of *Mesodinium* are highly productive (Smith and Barber, 1979) and form dynamic aggregations within the water column (Crawford and Purdie, 1992). *M. rubrum*-like ciliates can form thin layers in stratified surface waters (Sjöqvist and Lindholm, 2011), concentrate around down-welling frontal regions (Packard et al., 1978), and are capable of self-retention within estuarine systems by vertically migrating to avoid tidal flushing (Crawford and Purdie, 1992). These behaviors are due to *M. rubrum*'s astounding motility, which allow the ciliate to move at speeds of ~400 body lengths s⁻¹ through jumping (Fenchel and Hansen, 2006). One of the best-studied recurrent *M. rubrum* bloom locations is within the Columbia River estuary in the Pacific Northwest of the United States. Blooms in this system have been described to occur annually during late summer, and appear to be caused by only one of the five known genotypes of *M. rubrum* found in the estuary (Herfort et al., 2011b).

Here we present an analysis of the genetic diversity of *Mesodinium* and cryptophyte algal communities from both bloom and non-bloom conditions, in order to shed light on which species and variants of each group are associated with the “red tide” phenomenon. We designed new *Mesodinium* primers are capable of amplifying all known species from the Mesodiniidae family, rather than the *M. rubrum/major* complex only (Herfort et al., 2011b). We also designed one new cryptophyte rbcL primer in order to better anneal with major marine groups of these flagellates, since previous studies (Hoef-Emden, 2005) were focused on genera dominant in freshwater as well. This study establishes a framework for assessing the biogeography of Mesodiniidae genetic diversity and provides new insights into the cryptic diversity of these ciliates and cryptophyte algae.

MATERIALS AND METHODS

Collection of Cell Material

Environmental samples analyzed for *Mesodinium* and cryptophyte diversity were opportunistically gathered from various sources. In most cases (BR, NC, Bar-M4, GF-L3a, GF-XVI, TV), samples were collected from surface water preserved using 5–10% Lugol’s fixative. However, samples were also collected onto a 0.2 µm Sterivex™ filter (EMD Millipore, Billerica, MA USA) (CL) or centrifuged to form a cell pellet (LIS), and kept frozen until analysis. Previous research has shown that Lugol’s preserved material is sufficient for DNA extraction and quantitative (q) PCR assays, and while sensitivity of the assays decreased overtime, positive amplification was still possible for several months of sample storage at room temperature (Bowers et al., 2000). The sample from SGI, however, was unpreserved and shipping of the sample was slow due to the geographical isolation of the collection site. Upon the arrival of the SGI sample it was immediately centrifuged and the pellet was frozen until DNA was extracted.
Cultures Used for Phylogenetic Analysis
Cryptophyte and *Mesodinium* spp. cultures were grown in the lab for DNA extraction and sequencing of rbcL or rRNA gene fragments for phylogenetic comparison with our environmental data. Cryptophyte cultures including *Teleaulax acuta* (SCCAP K-1486), *T. amphioxeia* (CCMP 1170, GCEPO1), *Gemmignera cryptophila* (CCMP 2564), *Chroomonas* sp. (CCMP 270), *Hemiselmis pacific* (CCMP 706), *H. andersonii* (CCMP 439), *H. rufescens* (CCMP 440), *Hemiselmis* sp. (SUR21-C3), *Hemiselmis* sp. (NR11), and *Hanusia phi* (CCMP 325) were grown in the lab in F/2-Si at 18°C on a 14:10 L:D cycle, as were *M. rubrum* cultures AND-A0711 and CBJR05. Additionally, the *M. rubrum* cultures were maintained with GCEPO1, by feeding the ciliate weekly with a 1–5% volume addition of log-phase cryptophyte culture. A *Falcomonas* sp. culture (CCMP 2293) was also grown on F/2 without added Si (F/2-Si), but at 4°C and constant light. All of the above cryptophyte and *Mesodinium* cultures were harvested for DNA extraction by centrifugation of 10 ml of dense culture within a 15 ml Falcon tube at 4000 RPM for 15 min, the supernatant decanted, and the cell pellet frozen until processed. Cultures of *Teleaulax gracilis* (Cr6EHU), *T. minuta* (Cr8EHU), *T. cf. merimbula* (Cr59EHU), *aff. Teleaulax* (Gr22EHU), *Plagioselmis nanoplanctica* (Cr50EHU), *P. cf. prolonga* (Cr10EHU, Cr127EHU, Cr143EHU, Cr194EHU) and *Urgorri complanatus* (Cr1EHU) were grown in the same light and temperature conditions described above, on a modified F/2-Si (enriched with soil extract and selenium at 0.006 mM Na2SeO3 final concentration) at a salinity of 30 (*P. nanoplanctica* strain grown in freshwater), and were preserved in ethanol prior to DNA extraction and analysis of the rbcL gene. A live sample of *M. rubrum* culture MR-INO2007 were also acquired for analysis. For all cultures, approximately 10–20 ml of dense stationary-phase culture was collected by centrifugation (4000 RPM, 10 min) and frozen until extraction.

DNA Extraction and Gene Fragment Amplification and Sequencing
Nucleic acids of environmental samples were extracted from frozen cells collected by centrifugation (4000 RPM, 10 min) or on Sterivex™ filters (EMD Millipore) using either the PowerWater Sterivex DNA Isolation Kit (MO BIO Laboratories, Inc.) or a hot detergent lysis method as described by Gast et al. (2004), modified to exclude zirconia-silica bead disruption. All cultures as well as the North Carolina Bloom sample were extracted using the Qiagen DNeasy Plant Mini Kit. PCR was conducted using GoTaq (Promega) or GoTaq G2 Hot Start mix in 50 ml reactions, with a final concentration of 2.5 mM MgCl2, 200 µM dNTPs, 2.5 U GoTaq Flexi polymerase, and 0.1 µM primers for normal and 0.2 µM for hot start. Primers for *Mesodinium* spp. (Table 1) were designed to amplify the majority of the SSU and LSU rRNA genes, and the entire ITS region, resulting in a ~1880 bp amplicon. Gene fragments of the large subunit of the ribulose-1,5-bisphosphate carboxylase oxygenase gene (rbcL) were PCR amplified from DNA extracts using primers targeting cryptophyte plastids. A new primer, crypt_rbcLR2 (5'-CAGTGAATACCACCTGAAGCTA rbcL: 1185), designed using an alignment of cryptophyte rbcL sequences (See Supplementary Data Sheet 1 for accession numbers) was used in combination with L2F (Hoef-Emden et al., 2005). PCR conditions were: 95°C for 5 min followed by 40 cycles of 95°C for 60 s, 55°C for 60 s, and 72°C for 90 s followed by 72°C for 7 min. The genus-specific primers Meso245F and Meso28S_R were also used to amplify a combined fragment of the *Mesodinium* spp. 18S-ITS-28S genes. PCR conditions were: 95°C for 5 min followed by 40 cycles of 95°C for 60 s, 57°C for 60 s, and 72°C for 90 s followed by 72°C for 7 min. PCR products were visualized by agarose gel electrophoresis and later excised and purified from the gels using the Zymoclean Gel DNA Recovery Kit (Zymo Research). Clone libraries were constructed from gel purified fragments using the pGEM-T Easy Vector in the pGEM-T Easy Vector System II cloning kit (Promega Corporation) according to the manufacturer’s protocol. Selected clones were submitted for Sanger sequencing with a single primer to either Beckman Coulter Genomics (Single Pass Sequencing) or the W. M. Keck Ecological and Evolutionary Genetics Facility at the Marine Biological Laboratory (Woods Hole). Full-length Sanger sequencing of select *Mesodinium* clonal clones were run at Genewiz (Boston) (see Supplementary Data Sheet 1 for accession numbers). Sequences were edited and assembled into contigs with Sanger sequencing with a single primer to either Beckman Coulter Genomics (Single Pass Sequencing) or the W. M. Keck Ecological and Evolutionary Genetics Facility at the Marine Biological Laboratory (Woods Hole). Full-length Sanger sequencing of select *Mesodinium* clonal clones were run at Genewiz (Boston) (see Supplementary Data Sheet 1 for accession numbers). Sequences were edited and assembled into contigs using Sequencer (Gene Codes Corporation). We sequenced and analyzed 687 cryptophyte rbcL clones (accession numbers in progress) from all stations. For *Mesodinium* spp., a total of 903 clones were sequenced (accession numbers in progress) from all stations. Using a sequence similarity criterion of 98 and 99% for cryptophytes and *Mesodinium*, respectively, we constructed independent contigs and generated consensus sequences for use in our global alignment for each sample. This process helped to reduce the number of sequences used in our phylogenetic analyses while maintaining meaningful diversity data.

### Table 1
Primer Design and Use for Amplifying the SSU, ITS, and LSU rRNA Genes of *Mesodinium* spp. and rbcL Gene Fragments of Cryptophytes in This Study

| Primer          | Sequence | Gene: position (bp) |
|-----------------|----------|----------------------|
| **Mesodinium spp. primers** |          |                      |
| Meso245F        | CGACTGAGACGTCCGG | 18S: 246            |
| Meso580R        | CGTCCTGACTGTAAGTC | 18S: 585            |
| Meso1200F       | ATCCCGTAAAGCAAGGAC | 18S: 1217          |
| Meso1440F       | AACTGGAATGTCGTAAGGC | 18S: 1446         |
| Meso580R        | GACTGACAGATACGGAGGC | 18S: 603            |
| Meso586R        | AGCTCTGCTTCTTGTGA | 18S: 1017           |
| Meso1480R       | CTAACAACCTCAGCTGGAG | 18S: 1545          |
| Meso28S_R       | AGACTTGGATGACATTTTACC | 28S: 298          |
| **Cryptophyte primers** |          |                      |
| rbcL2F-800      | AGGAGGAAWAYATGTCTCAAT CCG rbcL: 1 |                      |
| Crypt_rbcLR2    | CAGTGAATACCCACTGGAAGCTA | rbcL: 1185       |
Phylogenetic Analysis and Species Assignment

Consensus sequences from assembled contigs were used for separate alignments of *Mesodinium* spp. and cryptophytes, in combination with available sequences in Genbank and sequences generated from cultures in our laboratory (see above). Alignments were constructed using the Clustal X algorithm (Larkin et al., 2007), and refined by eye using MacClade 4.08a (Maddison and Maddison, 2000). All maximum likelihood (ML) phylogenetic trees were executed with PhyML 3.0 (Guindon et al., 2010) using 100 bootstrap replicates and the general time reversible (GTR) substitution model, while estimating the gamma distribution parameter, proportion of invariable sites, and the transition/transversion ratio. Phylogenetic trees were constructed using TreeDyn 198.3 (Chevenet et al., 2006). Assignment of *Mesodinium* and cryptophyte species was determined by their nearest neighbor match of a known species within ML phylogenetic trees.

RESULTS

Sample Characteristics

Bloom Samples

The St. George Island (SGI; Alaska) bloom in the Bering Sea was observed within the main harbor, creating dense patches of red water (Table 2). The presence of *M. rubrum*-like ciliates was confirmed by microscopy at the time of sampling. A massive *Mesodinium* bloom was sampled from Ilha Bela, São Paulo, Brazil (BR) in the South Atlantic, and microscopic images from samples of the bloom confirmed the presence of large cells (not shown). This bloom stretched for over 100 km of the Brazilian coastline, and was visible from space (Carlowicz et al., 2014). A bloom sample was analyzed from the North Atlantic, in Oyster Bay, Long Island Sound (LIS), and was also documented using satellite imagery (Dierssen et al., 2015). This bloom revealed *M. rubrum* to be present at ~1000 cells ml⁻¹. Another North Atlantic red water bloom was sampled from the Outer Banks of North Carolina (NC), revealing *M. rubrum* at 1750 cells ml⁻¹. Finally, streaks of red water were sampled in the South Pacific, ~25 km off the Chilean coast (CL) during an oceanographic cruise. However, samples from this bloom were only quickly collected from the surface for nucleic acid extraction, with no additional documentation.

Non-bloom Samples

Samples from the Baltic Sea included one coastal Finland (Tvärminne) and two offshore Gulf of Finland samples (Table 2). A non-bloom subsurface sample from the Barents Sea (Bar-M4) was also analyzed and was documented to have 10 *Mesodinium* ml⁻¹. A sample taken during a research cruise in the North Pacific within the California Current (CC) was also analyzed, and had low levels of *M. rubrum* present (Table 2).

Mesodiniidae Phylogeny

Using primers designed to amplify a partial region of the 18S, the entire ITS, and a partial region of the 28S ribosomal RNA genes of all known Mesodiniidae taxa, we recovered sequence data from both heterotrophic and plastidic species from eight locations and analyzed their phylogenetic relationships (Figure 1). We also analyzed new sequence data from four cultures of *M. rubrum* and one culture of *Mesodinium pulex* (Table 3). As shown previously (Garcia-Cueto et al., 2012) the Mesodiniidae formed four well-supported and distinct clades, represented by *M. pulex*, *M. pupula*, *M. chameleon*, and the *M. major/rubrum* complex, respectively, and all comparisons between these major clades revealed the greatest p-distance (>7%) within the dataset (Table 4). Relatively few environmental sequences were recovered from the *M. pulex*, *M. pupula*, and *M. chameleon* clades, and intracladal diversity within these groups was lower than within the *M. rubrum/major* complex (Figure 1A). Compared to the *M. major/rubrum* variants, the mixotrophic ciliate *Mesodinium chamaeleon* shared 89.6–92% similarity, while *M. pulex* had 83.2–87.7%, and *M. pupula* only 81.2–86% (Table 4). *M. chamaeleon* had low similarity to the heterotrophic *M. pulex* (86%) and *M. pupula* (82.9%), as did the latter two species to one another (83.8%).

### TABLE 2 | Location and description of samples for analysis of *Mesodinium* and cryptophyte community diversity.

| Region          | Description                        | Abbrev. | Date     | Lat   | Long | Sal PSU | Temp °C | Meso² cells ml⁻¹ | Sample notes<sup>a</sup> |
|-----------------|------------------------------------|---------|----------|-------|------|---------|---------|------------------|--------------------------|
| South Atlantic  | Ilha Bela, BZ                       | BR      | 2/15/14  | −23.8 | −45.22 | -       | 24      | 37.5             | bloom                    |
| North Atlantic  | Long Island Sound                   | LIS     | 9/24/12  | 40.91 | −73.60 | 27.9    | 21.9    | 1003             | bloom; Oyster Bay       |
| North Atlantic  | Wavenes, NC                         | NC      | 10/6/08  | 35.56 | −75.44 | 28.63   | 22.5    | 1750             | bloom                    |
| North Atlantic  | Barents Sea; station M4             | Bar-M4  | 6/27/11  | 74.53 | 30.11 | 35.06   | 4.9     | 10              | 60m                      |
| Baltic Sea      | Gulf of Finland; station LL3a       | GF-LL3a | 7/8/12   | 60.07 | 26.34 | 5.26²   | 17.4²   | -                | depth integrated         |
| Baltic Sea      | Gulf of Finland; station XVI        | GF-XVI  | 7/8/12   | 60.25 | 27.25 | 4.52²   | 17.5²   | -                | depth integrated         |
| Baltic Sea      | Tvärminne Zoological Station       | TV      | 7/31/13  | 59.63 | 23.25 | 5.5     | 18.5    | -                | Hanko, Fl; conc.⁴        |
| South Pacific   | Chile Coast                         | CL      | 11/23/10 | −19.97| −70.72 | 34.8    | 19.5    | -                | bloom                    |
| North Pacific   | Bering Sea; St. George Is.          | SGI     | 9/15/12  | 56.57 | −169.68| -       | 6.8     | -                | bloom                    |
| North Pacific   | California Current                  | CC      | 7/5/13   | 36.33 | −123.14| 33.8    | 12.8    | 0.5              | 2m                       |

<sup>a</sup>Measurements from 1 m, while all others at same depth of sample; <sup>⁴</sup>Due to the patchy distribution of *M. rubrum* and its ability to form thin layers, these concentrations are at best approximation; <sup>²</sup>Sample depths are surface unless indicated otherwise. Depth integrated samples are from 0.5, 3, 5, 7.5, and 10 m; <sup>⁴</sup>Sample concentrated with a 10 µm net.
FIGURE 1 | Maximum likelihood phylogenies of the Mesodiniidae using Mesodinium spp. sequences from 10 global locations: the Baltic Sea (stations GF-LL3a, GF-XVI, TV are grouped together in this figure), Barents Sea (Bar-M4), California Current (CC), St George Island (SGI) harbor, coastal Chile (CL), coastal Brazil (BR), Long Island Sound (LIS), and coastal North Carolina (NC). Phylogeny of the Mesodiniide based on a sequence, composed of a partial SSU rRNA, internally transcribed spacer region (ITS), and a partial region of the LSU rRNA gene. (A) Phylogeny of the Mesodiniidae, focusing on community sequences related to M. chamaeleon and heterotrophic Mesodinium spp. (B) Phylogeny of the M. major/rubrum complex. Letters on branches of tree (A–H) refer to variants within the complex, and cartoons of the ciliates (oral view) depict the relative cell size of various isolates (Table 3). Red text is used for genotypes dominating bloom samples and numbers in parentheses refer to the number of clones representing each genotype.
In contrast, the predominantly phototrophic *M. major/rubrum* species complex formed eight subclades (Figure 1B) of which six have been described previously (Herfort et al., 2011b; Garcia-Cuetos et al., 2012). The *M. major/rubrum* complex was well-resolved, with four of the clades having 100% bootstrap support within a maximum likelihood phylogeny and the most autapomorphic characters (10) within the analyzed variation was found within variants A, D, F, G, and H (<0.5%), while the greatest variation (>1%) was found within clades C and B (Table 4). Polymorphisms between the different *M. major/rubrum* variants are shown in Table 5 (modified from Herfort et al., 2011b), revealing the sequence variation that shape clade phylogeny within this group. For instance, several shared polymorphisms are found between clades A, F, and G at bases 167, 206, 479, 556, and 569 while clades C and E share four such polymorphisms (bp: 69, 75, 556, and 574). Clades B, D, and H share polymorphisms with each other at bases 191, 212, and 556, and at additional loci with clades C and E at 167, 206, and 479. The most divergent variants within the *M. major/rubrum* group were A and C, sharing 95.7 and 95.9% similarity, respectively, with the other six variants, relative to the average among all groups of 96.6%. Of all *M. major/rubrum* variants, group A had the most autapomorphic characters (10) within the analyzed

### TABLE 3 | Origin and cellular dimensions of *Mesodinium rubrum* variant isolates.

| Variant | Strain | Origin/year | Length (µm) | Width (µm) | L/W | Volume (µm³) | Cryptophyte prey | Type reference |
|---------|--------|-------------|-------------|------------|-----|-------------|----------------|----------------|
| A       | NCMA 2563 | McMurdo Sound, AN 1996 | Mean 23.1 (2.6) | 22.4 (2.7) | 1.1 (0.1) | 5707 (1826) | GC | 1 |
| B       | MR-MAL01 | Gomso Bay, KR 2001 | Mean 21.0 | 13 | 1.6 | 1857 | TA | 2 |
| C       | MR-IN200702 | Inokushi Bay, JP 2007 | Mean 25.8 (3.8) | 21.8 (3.8) | 1.2 (0.1) | 6719 (3008) | TA | 3 |
| D       | AND-A0711 | Huelva, ES 2007 | Mean 16.2 (1.7) | 13.0 (1.0) | 1.3 (0.1) | 1374 (308) | TA | 4 |
| E       | Mr-DK2007 | Frederikssund, DK 2007 | Mean 31 | 21 | 1.5 | 7154 | TA | 5 |
| F       | CBJ05 | James River, US 2011 | Mean 21.7 (3.0) | 16.0 (1.7) | 1.4 (0.2) | 2937 (914) | TA | Here |

All measurements made here unless otherwise noted. Mean values presented with standard deviation in parentheses. Values report % sequence similarity and color displays results as a heat map (green: similar; red: dissimilar).

### TABLE 4 | Genetic p-distance matrix of a partial ‘18S–28S’ rDNA region for variants within the *Mesodinium major/rubrum* complex and other *Mesodinium* spp.

|          | A     | B     | C     | D     | E     | F     | G     | H     | MC    | MPUL  | MPUP  |
|----------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| **M. rubrum/major complex** |       |       |       |       |       |       |       |       |       |       |       |
| A        | 99.8  |       |       |       |       |       |       |       |       |       |       |
| B        | 95.4  | 98.6  |       |       |       |       |       |       |       |       |       |
| C        | 94.2  | 95.6  | 99.0  |       |       |       |       |       |       |       |       |
| D        | 95.6  | 97.8  | 95.7  | 100   |       |       |       |       |       |       |       |
| E        | 94.9  | 97.3  | 97.1  | 98.5  | 99.2  |       |       |       |       |       |       |
| F        | 97.5  | 96.4  | 96.1  | 96.6  | 96.1  | 99.8  |       |       |       |       |       |
| G        | 95.8  | 95.5  | 95.9  | 95.9  | 95.6  | 97.8  | 99.8  |       |       |       |       |
| H        | 96.6  | 98.5  | 96.4  | 98.5  | 98.0  | 97.6  | 96.4  | 100   |       |       |       |
| MC       | 91.0  | 90.8  | 89.6  | 91.8  | 91.0  | 91.9  | 91.3  | 92.3  | 99.7  |       |       |
| MPUL     | 86.5  | 86.2  | 86.1  | 86.5  | 86.4  | 86.7  | 83.2  | 87.7  | 88.6  | 100   |       |
| MPUP     | 82.8  | 83.2  | 81.2  | 83.9  | 82.8  | 83.2  | 86.0  | 83.9  | 82.9  | 83.8  | 98.1  |

Values report % sequence similarity and color displays results as a heat map (green: similar; red: dissimilar).
|   | A | F | G | B | D | H | C | E |
|---|---|---|---|---|---|---|---|---|
| 69 | T | T | T | T | T | T | A | A |
| 75 | G | G | G | G | G | G | A | A |
| 80 | C | T | T | T | T | T | T | T |
| 92 | A | A | A | A | A | A | T | A |
| 97 | C | C | C | C | C | C | T | C |
| 101 | A | A | A | A | A | A | G | R |
| 102 | G | G | G | G | G | G | A | G (A) |

**TABLE 5** Details of genetic variation for the *Mesodinium major/rubrum* complex within the partial ‘18S–28S’ rDNA region for all available sequence data.

Nucleotide positions are based on a similar table in Herfort et al. (2011b). Boxed regions represent autapomorphic character states and are summed in the final row. Letters in parentheses represent minor exceptions of character states.
region, followed by groups G (5) and C (4) (Table 5), and these variants were the only to have 100% boot strap support (Figure 1B).

**Mesodinium Diversity**

The most widely encountered variant was D (*M. major*), found at 6 of the 10 sampling sites, while variants A and F were found at 5 (Figure 2). Sequences from variants C and H, as well as the heterotrophic *M. pulex* were each only recovered from one station, while no clade E sequences were found in this study. Analysis of samples from red-water events of *Mesodinium* revealed that the bloom phenomenon within this species complex is not associated with one particular variant. Sequences from subclades B and D dominated (>85%) clones recovered from the four blooms analyzed in this study, and in two cases sequences consisted only of *M. major* (Figure 2). Variant B dominated two blooms from the eastern United States, and was not found at non-bloom sites (Figure 2). Clones from variants F and H dominated the bloom sample from SGI, however, the sample was not preserved when shipped and thus the relative proportion of *M. major/rubrum* variants is uncertain. Samples from the Baltic Sea were from coastal and open water stations, and were not associated with bloom events. Collectively, the Baltic Sea had the greatest diversity of Mesodiniidae sequences, with clones recovered from heterotrophic *M. pupula*, the mixotrophic *M. chamaeleon*, as well as from multiple *M. rubrum/major* clades. The CC sample was also diverse, with three variants of *M. major/rubrum* as well as *M. chamaeleon* (Figure 2). In contrast, a non-bloom sample from an off shore station in the Barents Sea was composed entirely of *M. major*.

**Cryptophyte rbcL Phylogeny**

Our cryptophyte *rbcL* gene data set included new sequences from 22 cryptophyte cultures, comprised largely of TPG and *Hemiselmis* species. Analysis of our environmental clones resulted in 39 distinct contigs, at 99% sequence similarity, from the nine sample sites (one Baltic site was not assessed for cryptophyte *rbcL* diversity). Of these, 21 consensus sequences were determined to be from the *Teleaulax/Plagioselmis/Geminigera* (TPG) clade (Figure 3). Our maximum likelihood tree had strong support (>90 boot strap) for the cryptophyte clades comprised of TPG, *Proteomonas, Rhodomonas*-like cryptophytes, *Guillardia/Hanusia, Urgorri*, and *Falcomonas*, and *Cryptomonas*, while the clade comprised of *Hemiselmis* and *Chroomonas* had weaker support (78%) (Figure 3). Culture sequences from *T. amphioxeia* and *P. cf. prolonga* were polyphyletic, forming a species complex that appears to be in need of taxonomic revision. The position of *Proteomonas* as sister of TPG was strongly supported. No environmental sequences were recovered from the *Cryptomonas* or *Proteomonas* clades. Of all the recovered environmental sequences, those clustering with *Falcomonas* sp. appeared to represent the most novel lineages of uncharacterized cryptophyte species (Figure 3).

**Cryptophyte Plastid rbcL Diversity**

Community plastid *rbcL* sequences were analyzed for each sample in order to test the hypotheses that *M. rubrum* blooms are associated with TGP-like cryptophyte sequences and that the number of cryptophyte phylotypes are greater in non-bloom samples. Since these samples are unfractionated community
FIGURE 3 | A maximum likelihood phylogeny of cryptophytes using partial plastid LSU RuBisCO gene (rbcL) sequences from 10 global locations: the Baltic Sea (GF-LL3a, GF-XVI, TV), Barents Sea (Bar-M4), California Current (CC), St George Island (SGI) harbor, coastal Chile (CL), coastal Brazil (BR), Long Island Sound (LIS), and coastal North Carolina (NC). Red text refers to genotypes dominating bloom samples and numbers in parentheses refer to the number of clones representing a genotype. Major cryptophyte clades are boxed by color: TPG: Teleaulax/Plagioselmis/Geminigera group; Falc: Falcomonas-like species; CH: Chroomonas/Hemiselmis group; Rho: Rhodomonas-like cryptophytes; GH: Guillardia/Hanusia group; Urg: Urgoniella-like cryptophytes.
samples, it is not possible to determine the proportion of cryptophyte sequences that are from free-living cells or associated with a particular protist species (e.g., _M. rubrum_). However, while these samples may not specifically represent the diversity of the free-living cryptophyte community _per se_, they are ultimately derived from it. Overall 74% of _rbcL_ sequences from all sample sites originated from _T. amphioxeia/P. cf. prolonga_-like sequences, while a striking 82% were from the _TPG_ clade. Bloom samples were dominated by _TPG_ cryptophytes (92%), with samples from BR, CL, LIS, and NC sites mostly (>90%) comprised of _T. amphioxeia/P. cf. prolonga_-like sequences. In contrast to other bloom samples, the SGI bloom was dominated by a phylotype branching near _G. cryophila_. From non-bloom stations (343 sequences), 56% were _T. amphioxeia/P. cf. prolonga_-like sequences, while 72% belonged to the _TPG_ clade. The Bar-M4 sample was primarily composed of _T. amphioxeia/P. cf. prolonga_-like sequences, _T. acuta_, and _Rhodomonas_ sp., while _T. amphioxeia/P. cf. prolonga_-like sequences, while 72% belonged to the _TPG_ clade. The Bar-M4 sample was primarily composed of _T. amphioxeia/P. cf. prolonga_-like sequences, _T. acuta_, and _Rhodomonas_ sp., while _T. amphioxeia/P. cf. prolonga_-like sequences dominated the Baltic Sea and CC samples. One of the Baltic Sea sites, station XVI in the Gulf of Finland, was also rich in a _Hemiselmis_ sp. and _Hanusia phi_. Sequences from _Falcomonas_ spp. were found to comprise a minor component of cryptophyte genetic diversity from eight of the nine sampling sites. _U. complanatus_, which was originally described from Atlantic estuaries in southwestern Europe (Laza-Martínez, 2012), was detected in the Baltic and Barents Seas.

**DISCUSSION**

This study is the first to assess the environmental genetic diversity of both non-photosynthetic and plastidic marine _Mesodinium_ species, across a broad geographical range of sampling sites. Our findings expand upon the known diversity of the _M. major/rubrum_ species complex, and further solidify the relationship of _Teleaulax amphioxeia_ cryptophytes with their bloom events. Further, we show novel diversity of cryptophyte algae, particularly among _Falcomonas_-like species, and provide evidence for the widespread dominance of the _Teleaulax/Plagioselmis/Geminigera_ (TPG) cryptophyte group.

**Phylogeny of _Mesodinium_**

_Mesodinium_ spp. have an unusual 18S rRNA gene with numerous deletions and substitutions within conserved regions compared to other ciliates (Johnson et al., 2004), and they appear to lack a separate 5.8S rDNA region, suggesting that it may be fused to the 28S rDNA gene (Herfort et al., 2011b). This characteristic is also found in the microsporidia and bacteria (Hoef-Emden, 2005). While _Mesodinium_ show typical litostome secondary structure in their V4 region, including reduction of helices 23.1, 23.8, and 23.9 and the absence of 23.5 (Strüder-Kypke et al., 2006), phylogenies based on the SSU rRNA gene consistently result in a novel, deep basal branch within the ciliates (Johnson et al., 2004; Chen et al., 2015a). From ultrastructural observations, only the ciliary transition region supports their placement within the litostomes (Garcia-Cuetos et al., 2012). A recent multigene phylogeny of the _Mesodiniumidae_ also suggests a deep basal ciliate branch and strongly implies that they may represent a distinct ciliate class, the _Mesodiniidae_ (Gao et al., 2016). However, a phylogenomic analysis of non-model ciliates using transcriptomic data failed to resolve the monophyly of the _Mesodiniidae_ (Chen et al., 2015b).

Our phylogenetic analysis of taxa within the _Mesodiniidae_ agree with Garcia-Cuetos et al. (2012), with strong support for four major clades distinguishing the two heterotrophic lineages of _M. pulex_ and _M. pupula_, the mixotrophic _M. chamaeleon_, and the predominantly phototrophic _M. major/rubrum_ complex. At least one additional major clade of _Mesodinium_-like ciliates are found in freshwater habitats, and phylogenies based on the SSU rRNA gene group them as a sister group to _M. pulex_ (Bass et al., 2009).

Since we did not sequence samples from freshwater habitats and no cultures of these species were available, comparisons of their full rRNA gene cassette to marine _Mesodiniumidae_ were not made here. Our analysis resolved two novel subclades of _M. rubrum_: (1) variant G was identified from sequence data of a culture from Chesapeake Bay, USA, along with environmental sequences from the North Carolina (NC) and St. George Island (SGI) bloom events, and (2) variant H, which is only represented by environmental sequences from the SGI sample. One sequence from a culture isolated in Spain (Riobó et al., 2013) grouped with a Danish culture from the F subclade (Garcia-Cuetos et al., 2012) as well as clones from the Baltic Sea and SGI. Together, variants F and G grouped strongly with variant A to form one major branch of the _M. rubrum/major_ complex. We found _M. rubrum_ sequences from the Baltic Sea, CC, NC, and SGI that grouped with variant A, which includes a well-studied cultured strain from Antarctica (Gustafson et al., 2000; Johnson et al., 2007) and sequences from coastal Oregon (Herfort et al., 2011b). Cultures isolated from Korea (Yih et al., 2004) and Japan (Nishitani et al., 2008) were found to belong to subclade B, which included sequences from blooms in LIS and NC, as well as the recurrent blooms of the Columbia River Estuary (Herfort et al., 2012). _M. major_ (subclade D) was found in the Baltic Sea, Bar-M4, and in blooms from coastal BR and CL, and has also been found in the Columbia River Estuary (Herfort et al., 2011b) as well as coastal Denmark (Garcia-Cuetos et al., 2012). Two other _M. rubrum_ variants, C and E, remain uncultured and have only been found in the Northeastern Pacific (here and Herfort et al., 2011b). Subclades B, C, D, E, and H formed the second major branch of the _M. rubrum/major_ complex. While C, E, and H have not yet been cultured, subclade B appears to be highly variable in size, both within and between strains (Table 2). A bloom of _M. rubrum_ variant B in LIS was found to contain cells of typical size for _M. rubrum_ cultures (Tables 2 and 3).

**Genetic Diversity of _Mesodinium_**

The studies of Herfort et al. (2011b) and Garcia-Cuetos et al. (2012) revealed that “_M. rubrum_” is a species complex of closely related genetic variants, most of which are morphologically indistinguishable. While cell size clearly distinguishes _M. major_ from _M. rubrum_ variants, cell size within _M. rubrum_ strains appears to be highly variable considerably (Table 2).
Furthermore, sequence variation of rRNA genes and the ITS region in *M. major* relative to *M. rubrum*, is equal to or lower than variation among *M. rubrum* variants (Table 4), making the distinction of species vs. variant somewhat ambiguous based on these sequences alone. While we are beginning to identify trends in genotype distribution of *M. major/rubrum* ciliates and their association with bloom events, we know little about distinctions in the physiological or behavioral diversity among these variants and if they may possess distinct ecological niches.

While our results strongly suggest that heterotrophic species have less intracladal diversity in marine ecosystems, our samples focused on bloom events and are therefore biased toward the mixotrophic species. Stations where heterotrophic species were abundant, such as two Gulf of Finland (Baltic Sea) samples rich in *M. pupula* (>40% of *Mesodinium* clones), revealed essentially identical sequences within and between these populations. Furthermore, *M. pupula* sequences from the Baltic Sea were nearly identical to those from SGI. In addition, *M. pulex* sequences from SGI and *M. chamaeleon* sequence data from SGI and the Baltic Sea were remarkably similar to strains isolated from Danish waters.

**Bloom Samples**

All temperate and subtropical bloom samples were dominated either by variant B or D from the *M. rubrum-major* complex. The only previous bloom where genetic diversity of *Mesodinium* spp. were assessed was in the Columbia River Estuary, and these populations are also associated with subclade B (Herfort et al., 2011b). Annual blooms of *M. rubrum* occur within the Columbia River, developing first near the mouth of the estuary and later within the main channel in coincidence with neap tides and decreased turbulence (Herfort et al., 2011a). While summer *Mesodinium* populations in the Columbia River are dominated by subclade B, they are largely absent in spring samples, which were instead associated with subclades C and D (Herfort et al., 2011b). Our study only provided “snapshots” of community genetic diversity, and it is likely that diversity at our various sites changes seasonally. Danish cultures of variant F have also been isolated from red-water within islands of the Danish straits (Garcia-Cuetos et al., 2012), however, it is unclear if this clade was primarily responsible for the observed bloom.

Blooms of *M. rubrum* along the North Carolina coastline have been periodically observed (Hathaway, 2014), but no published studies are available. The North Carolina bloom was the only preserved bloom sample that had more than one subclade of *M. rubrum* present, with three other subclades comprising ≤5% of the community. Since the bloom sample was collected near Oregon Inlet, which empties the Pamlico Sound into the Atlantic Ocean, some of the strains present may have been washed in from estuarine populations. The sequences of one of these subclades (G) are identical to that of a strain isolated from the James River estuary in nearby Chesapeake Bay, which would support mixing of estuarine populations into this coastal bloom. The Long Island Sound bloom was also dominated by subclade B (Figure 2), and was associated with calm wind speeds (Dierssen et al., 2015). This bloom was detected from the Space Station, using a Hyperspectral Imager for the Coastal Ocean sensor (Dierssen et al., 2015).

Numerous studies have commented on the size variability in natural populations of *M. rubrum*-like ciliates in diverse geographical locations (Taylor et al., 1971; Lindholm, 1981; Crawford, 1993; Rychert, 2004; Montagnes et al., 2008). Garcia-Cuetos et al. (2012) formally described the largest of plastidic *Mesodinium* spp. as *M. major*, which attains cell dimensions of 50 µm × 40 µm (L × W). Particularly large *M. rubrum*-like cells have been observed in many blooms and coastal ocean communities, particularly those associated with upwelling regions (Packard et al., 1978). Both of our *M. major*-dominated bloom samples were from coastal blooms in Brazil and Chile, and were likely associated with upwelling. Both regions have a rich history of documented *Mesodinium* “red tides,” many of which have been described reaching massive proportions. *Mesodinium* blooms have been previously reported in the coastal region of northern São Paulo, Brazil, where our sample was taken, and have been described as forming thin layers of red water 1–2 m below the surface with a maximum thickness of ~30 cm (Owen et al., 1992). An image of the bloom sampled near Ilha Bela, Brazil, was captured by NASA’s Aqua satellite using the moderate resolution imaging spectrophotometer (MODIS), and the red water formed an offshore patch of staggering proportions, extending 800 km from Rio De Janeiro to south of Florianópolis (Carlowicz et al., 2014). *Mesodinium* blooms in southern Brazil have also been documented to occur near the Itajaí-Açu River (Proença, 2004).

Darwin is thought to have documented the first *Mesodinium* bloom off the Chilean coast while on the H. M. S. Beagle (Darwin, 1906; Hart, 1943), and such occurrences are common in this region during non El Niño periods (Avaria and Muñoz, 1987; Marin et al., 1993). Blooms documented from the Peruvian upwelling zone have reported some of the highest rates of primary production ever measured (Smith and Barber, 1979), and red water patches spanning 100s of square miles (Ryther, 1967). Based on these previous observations and our phylogenetic analysis of the Chilean and Brazilian blooms, it is likely that these recurrent events and perhaps blooms in other upwelling ecosystems are predominantly *M. major*.

**Non-bloom Samples**

*Mesodinium* is abundant in the Baltic Sea (Leppanen and Bruun, 1986) and is known to form occasional blooms in brackish basins around the archipelago region of Åland (Lindholm, 1978). During spring in the northern Baltic Sea, *M. rubrum*-like ciliates dominate ciliate biomass and are thought to account for ~10% of all primary production (Leppanen and Bruun, 1986). In the lower Baltic Sea, both large and small *M. rubrum*-like ciliates dwell in the upper water column during June, with mostly large cells found at greater depths (Rychert, 2004). Within the Åland region, up to three size classes (~20, 40, 60 µm) of *M. rubrum*-like ciliates are observed, with the smallest being dominant during autumnal red water events (Lindholm, 1978). Deep (~70 m) layers of pigmented *Mesodinium* have also been found in the Baltic Sea, below the thermocline (Setälä et al., 2005) and near the anoxic boundary of the Gotland Basin (Weber et al., 2014). Our two samples from the Gulf of Finland (GF) revealed a mixed community of *M. rubrum* subclades A, D, and F. While subclade
D (M. major) is the largest Mesodinium spp., both subclades A and F may have a range of cell sizes, even within a single strain (Table 2). Surprisingly, the largest component of both GF samples was M. pupula, a heterotrophic species that is slightly larger than M. pulex (Garcia-Cuetos et al., 2012). The ecology of M. pupula is poorly known, but has been shown to be one of the most abundant heterotrophic ciliates in coastal Yellow Sea microbial communities (Jiang et al., 2011). While M. pulex has instead been reported as a dominant heterotrophic ciliate species in Baltic communities (Setälä and Kivi, 2003), this species is difficult to discern from M. pupula using traditional light microscopy. The near shore station from Hanko, FI, also had subclades D and F, as well as the mixotrophic M. chamaeleon, which is capable of possessing a variety of cryptophyte plastid types, including those containing phycocyanin (Moestrup et al., 2012). M. chamaeleon differs from M. rubrum-like ciliates, in the structure of its cirrus, its predominantly benthic niche, and in the organization of sequestered cryptophycean organelles (Moestrup et al., 2012). M. chamaeleon-like ciliates, reported as green-blue in appearance, have been observed in the Åland region (Lindholm, 1985), as well as in a coastal Rhode Island estuary (Hargraves, 1991). Our Baltic sequence of M. chamaeleon was essentially identical to an isolate from coastal Denmark, and both were slightly different from sequences found in SGI.

The CC sample was one of the more surprising due to the unexpected presence of M. chamaeleon (13% of clones) in an offshore site. The presence of this “benthic” ciliate in a pelagic environment underscores how little we know about the ecology of this species. The remaining portion of the community was split among variants A, C, and D. Variants C, E, and H are the least distributed and most poorly studied of the M. major/rubrum complex, as none have yet been cultured and all have only been found at only one or two sites along the North Pacific coast of the US.

Cryptophyte Phylogeny

Taxonomic relationships within the cryptophytes, based on traditional morphological and structural traits of their pigments and cell surface, have recently been tested using gene phylogenies (Fitt et al., 2001; Hoef-Emden et al., 2002). These studies have found that while biliprotein type is congruent with molecular phylogenies, the type of inner periplast is not (Hoef-Emden et al., 2002). These phylogenies revealed seven separate lineages within plastid-bearing cryptophytes, all except Cryptomonas, containing representatives from marine environments. An additional lineage, represented by the brackish-water species U. complanatus, was later discovered (Laza-Martinez, 2012). Previously Hoef-Emden et al. (2005) commented on the potential for accelerated evolutionary rates when constructing rbcL phylogenies of cryptophytes, due to shifts in the number of amino acids showing changes in codon usage among early and late diverging taxa. This study was based only on genus Cryptomonas, an exclusively freshwater group containing some highly divergent taxa. Yet, despite finding high evolutionary rates for rbcL, resulting cryptophyte phylogenies were largely congruent with those of 18S rDNA (Hoef-Emden et al., 2005). Our results, which used the most diverse rbcL dataset to date, are also congruent with previously published phylogenies of nuclear 18S rRNA genes (Hoef-Emden et al., 2002). The eight lineages of plastid-bearing cryptophytes were recovered in the rbcL phylogeny. Moreover, the Proteomonas lineage appeared as the sister group of TPG with high support. While major clades are well-supported in previous phylogenies, the relationships among lineages have not been well-resolved. The clustering of Proteomonas and TPG was only partially supported in nucleomorph, but not nuclear, 18S rRNA genes phylogenies in a clade that also included Guillardia/Hanusia (Hoef-Emden et al., 2002).

The traditionally used morphological trait of periplast structure, as a continuous sheet or polygonal, has been shown to vary with life stage in the genera Cryptomonas and Proteomonas and is thus phylogenetically uninformative (Hill and Wetherbee, 1986; Hoef-Emden and Melkonian, 2003). Similar observations of periplast dimorphism have also been made within the Teleaulax group (Garcia-Cuetos, 2011), and this trait has traditionally been used to distinguish this genus from Plagioselmis. Furthermore, the midventral band, a morphological trait used to distinguish Teleaulax (thought to be absent) and Plagioselmis, was recently shown to be polyphyletic within this group (Laza-Martinez et al., 2012). The finding of strains with the T. amphioxeia and P. prolonga morphology sharing nearly identical rbcL sequences can be interpreted from the perspective of a dimorphic species, making both names synonym. The Teleaulax/Plagioselmis dimorphism can also account for incongruences between microscope-based morphological identifications and environmental DNA sequence identities, as in Bazin et al. (2014), where only T. acuta sequences were retrieved from a P. prolonga red tide sample. These observations, combined with the polyphyletic phylogeny of Teleaulax shown here and previously using 18S rRNA (Deane et al., 2002; Rial et al., 2015), suggest that these genera are in need of taxonomic revision.

Cryptophyte Diversity and its Role in Mesodinium Blooms

About half of our samples were from Mesodinium blooms, and these samples (excluding SGI) were dominated (>95%) by T. amphioxeia, as expected from previous studies (Nishitani et al., 2010, #874; Herfort et al., 2011b, #893). However, clone libraries from non-bloom samples were also largely comprised of TPG sequences, particularly T. amphioxeia (Figure 4). Exceptions were the high latitude BS and SGI samples, which were largely composed of T. acuta and G. cryphila-like sequences, respectively. G. cryphila appears to primarily be a cold-water species, dominating cryptophyte assemblages in Antarctic Dry Valley Lakes (Bielewicz et al., 2011), and was co-isolated with an Antarctic M. rubrum strain (Gustafson et al., 2000). T. gracilis and T. minuta have also been shown to sustain in vitro M. rubrum (clade F) growth (Rial et al., 2015). However, only few sequences of the former and none of the latter were recovered. These results strongly suggest that TPG species may frequently dominate cryptophyte communities in coastal ecosystems, and play critical roles in supporting the productivity of M. major/rubrum.
FIGURE 4 | Community genetic diversity of cryptophyte algal plastids from the Baltic Sea (GF-LL3a, GF-XVI, TV), Barents Sea (Bar-M4), California Current (CC), St George Island (SGI) harbor, coastal Chile (CL), coastal Brazil (BR), Long Island Sound (LIS), and coastal North Carolina (NC).

Community diversity of cryptophytes based on partial sequences of the plastid LSU RuBisCO gene. Categories include the Teleaulax/Plagioselmis/Geminigera (TPG) group, Falcomonas-like species (Falc), the Chroomonas/Hemiselmis group (CH), Rhodomonas-like cryptophytes (Rho), the Guillardia/Hanusia group (GH), and Urgoni-like cryptophytes (Urg). *SGI sample was not properly preserved, so the relative proportion sequences within each category may have been compromised.

Baltic Sea planktonic communities have been shown to be rich in TPG cryptophytes, and are also known to have Hemiselmis and Rhodomonas spp. (Hill et al., 1992; Carstensen and Heiskanen, 2007; Hajdu et al., 2007). While none of our Baltic Sea samples were associated with a Mesodinium bloom, Mesodinium cell counts were not available and thus it is unclear to what degree their plastids may be contaminating the cryptophyte rbcL community signal. In addition to TPG species, we also found Falcomonas, Hemiselmis, and U. complanatus sequences in the Gulf of Finland. U. complanatus was originally described from French, Spanish, and Portuguese estuaries and has a maximum growth rate at a salinity of 10 psu (Lazama-Martinez, 2012); thus its presence within these brackish waters is not surprising. A single clone of U. complanatus was also recovered from our BS sample. While the BS was dominated by T. amphioxeia, M. major was found in this sample at ~10 cells/ml, which would equate to about 400–600 T. amphioxeia plastids ml⁻¹ based on plastid numbers reported by Garcia-Cuetos et al. (2012). Since our community samples were not size fractioned, in this case our clone libraries may have been highly enriched with plastids from this species since it is the preferred prey of the M. major/rubrum complex (Yih et al., 2004; Park et al., 2007; Myung et al., 2011; Hansen et al., 2012).

Peaks in cryptophyte abundance have been found to both proceed (Johnson et al., 2013) and co-occur (Kim et al., 2007; Weber et al., 2014) with high levels of M. rubrum-like ciliates in coastal ecosystems. However, most historical accounts of Mesodinium blooms did not note the abundance or composition of co-occurring cryptophyte communities. While certain aspects of M. major/rubrum blooms have been well-studied within a variety of ecosystems, such as the Southampton estuary (Crawford et al., 1997), upwelling zones from Ecuador to Chile (Ryther, 1967; Avaria, 1976; Jimenez and Intriago, 1987), Baltic Sea Fjords (Lindholm, 1978; Lindholm and Mörk, 1990; Sjöqvist and Lindholm, 2011), and portions of the Gulf of California (Gárate-Lizárraga et al., 2002; Bulit et al., 2004), only within the Columbia River estuary (Herfort et al., 2011a) have studies focused on cryptophyte-Mesodinium dynamics. Within this system, cryptophytes are high in abundance both prior to and during early M. rubrum blooms, but decline as ciliate concentrations increased (Peterson et al., 2013). Observations of M. rubrum cells from these communities revealed what appears to be a novel feeding strategy on cryptophyte algae, with numerous prey cells attached to either the cirri and/or feeding tentacles of the ciliates (Peterson et al., 2013). These observations suggest that some M. rubrum populations may be able to act opportunistically and quickly consume a surplus of cryptophytes in order to sustain their productivity. In the Columbia River, M. rubrum cells from blooms were analyzed for 16S rRNA gene diversity during two consecutive years and found to exclusively possess T. amphioxeia-like plastids (Herfort et al., 2011b). In contrast, surface clone libraries from the estuary were nearly devoid of cryptophyte sequences, suggesting they primarily reside at depth (see Peterson et al., 2013).
CONCLUSION

Our results solidify the link between *Mesodinium* blooms and *T. amphioxea*-like cryptophytes in both temperate and subtropical Atlantic and Pacific ecosystems. While multiple variants of the *M. major/rubrum* complex are linked to estuarine and near-shore blooms, our results suggest that variant B is the most common agent in temperate and subtropical regions, while variants A and F may be involved in higher latitude blooms. The largest variant, *M. major* (D), was shown to be associated with shelf-water and offshore blooms in coastal Pacific regions and is likely responsible for the largest and most productive *Mesodinium* blooms (e.g., Ryther, 1967; Packard et al., 1978; Smith and Barber, 1979) associated with upwelling. Even non-bloom samples in our study were dominated by TPG cryptophyte sequences, implicating this group as playing a critical role in marine microbial foodwebs. Their widespread distribution and abundance, may explain why *Mesodinium* ciliates have adapted to exploit them for fueling their acquired phototrophy niche. We recommend that future studies of *Mesodinium* genetic diversity using targeted PCR should use the primers presented here in order to gain further insights into the diversity of the entire genus.

AUTHOR CONTRIBUTIONS

MJ contributed to sampling, performed data analysis, and wrote the manuscript. DB performed all of the laboratory work for PCR, cloning, and sequencing preparation and contributed to writing the manuscript. DS contributed to data interpretation and writing the manuscript. MJ thanks the funding support of the National Science Foundations Grants NSF-OCE 1031718 and NSF-IOS 1326228. Project IT-699-13 from the Basque Government allows for maintenance of the microalgae culture collection at the University of the Basque Country (UPV/EHU).

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2016.02017/full#supplementary-material

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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