Life-cycle modification in open oceans accounts for genome variability in a cosmopolitan phytoplankton

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Emiliania huxleyi is the most abundant calcifying plankton in modern oceans with substantial intraspecific genome variability and a biphasic life cycle involving sexual alternation between calcified 2N and flagellated 1N cells. We show that high genome content variability in Emiliania relates to erosion of 1N-specific genes and loss of the ability to form flagellated cells. Analysis of 185 E. huxleyi strains isolated from world oceans suggests that loss of flagella occurred independently in lineages inhabiting oligotrophic open oceans over short evolutionary timescales. This environmentally linked physiogenomic change suggests life cycling is not advantageous in very large/diluted populations experiencing low biotic pressure and low ecological variability. Gene loss did not appear to reflect pressure for genome streamlining in oligotrophic oceans as previously observed in picoplankton. Life-cycle modifications might be common in plankton and cause major functional variability to be hidden from traditional taxonomic or molecular markers.

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Introduction

Marine phytoplankton is responsible for roughly half of Earth’s primary productivity and thus is a key driver in the global carbon cycle (Field et al., 1998). How phytoplankton adapt to environmental variability is a critical factor determining the feedbacks of oceanic ecosystems on biogeochemical cycling and climate. Among phytoplankton, coccolithophores are the main calcifiers. One taxon, Emiliania huxleyi, has colonized most ocean surface waters since first arising only 291 kya (Raffi et al., 2006) to become the most abundant modern coccolithophore. It forms dense blooms (103–105 cells per ml) of calcified cells in fjordic, coastal and open ocean temperate to subpolar waters of both hemispheres as part of annual productivity cycles (Paasche, 2001). Calcified E. huxleyi cells are also important components of phytoplankton communities in tropical and subtropical open oceans (see, for example, Hagino and Okada, 2004; Beaufort et al., 2008; Siokou-Frangou et al., 2010), despite rarely or never forming blooms in these more stable and oligotrophic zones where cells are 100–1000-fold more dilute. This cosmopolitan species represents a highly relevant model for assessing phytoplankton adaptation to contrasting environments.

Laboratory studies have indicated that high physiological variability exists within E. huxleyi, with, for instance, contrasting responses of different strains to experiments simulating ocean acidification (Riebesell et al., 2000; Iglesias-Rodriguez et al., 2008; Langer et al., 2009). A high level of genomic variability was also suggested through genomic sequencing of several strains (Read et al., 2013;
Kegel et al., 2013). *E. huxleyi* has a biphasic life cycle consisting of non-flagellated diploid (2N) cells that produce calcite plates (coccoliths) and haploid (1N) cells that are flagellated but not calcified (Klaveness, 1972; Green et al., 1996; von Dassow et al., 2009). Both cell types are capable of asexual reproduction by mitosis, and are assumed to be connected by meiosis and syngamy (that is, sexual reproduction), as in other coccolithophores (Billard and Inouye, 2004). Sex might provide genetic advantages for adaptation to new environments (Kaltz and Bell, 2002; Becks and Agrawal, 2010, 2012), whereas biphasic life cycling might facilitate adaptation to heterogeneous environments through niche partitioning (Hughes and Otto, 1999; Coelho et al., 2007) or provide escape from specific biotic pressures such as parasites or viruses (Correa and McLachlan, 1991; Frada et al., 2008). Consistent with ecological niche partitioning, 1N and 2N *E. huxleyi* show important physiological and transcriptomic differences (Houdan et al., 2005; von Dassow et al., 2009; Rokitta et al., 2011, 2012). 1N *E. huxleyi* appear resistant to specific lytic viruses (*Emiliania huxleyi* viruses (EhVs)) that attack 2N cells (Frada et al., 2008) and serve as major biological agents controlling *E. huxleyi* bloom dynamics in relatively productive temperate to subpolar areas (Brussaard et al., 1996; Wilson et al., 2002; Cooilen, 2011). We hypothesized that the life cycle of *E. huxleyi* has differentially adapted to the contrasting ecological pressures of habitats in which the species blooms versus those in which it forms more stable and dilute populations.

**Materials and methods**

Clonal axenic 2N and 1N *E. huxleyi* strains (RCC1216 and RCC1217, respectively) obtained from the Roscoff Culture Collection (RCC: www/roscoff-culture-collection.org) and originating from the same genetic background (that is, a clonal 2N strain isolated from temperate coastal waters near New Zealand that formed 1N cells in culture) were grown under identical conditions for comparative analyses of gene expression. Previous analysis of normalized Sanger-sequenced complementary DNA (cDNA) libraries (von Dassow et al., 2009) was complemented here by non-normalized 454-sequenced cDNA libraries (Supplementary Table S1) and microarray expression analysis. Growth conditions of *E. huxleyi* strains for harvesting of RNA for transcriptome sequencing by 454 have been previously described (von Dassow et al., 2009). 454 sequences have been submitted to the European Nucleotide Archive (ENA) database (http://www.ebi.ac.uk/ena; study accession number ERP008543). For microarrays, cells were harvested at midday and midnight from cultures in early exponential growth (50 000–100 000 cells per ml) on a 14:10 light/dark cycle at 100 μmol photons m⁻² s⁻¹ at 17°C. A total of 28 306 clusters of Sanger expressed sequence tags (ESTs; 39 091 single EST reads from RCC1216 and RCC1217 (von Dassow et al., 2009) and 72 513 from CCMP1516 downloaded from http://genomeportal.jgi.doe.gov) were represented by 84 881 60-mer probes (2–3 probes/cluster) on 105K microarrays (Amadid: 022065) for two-color (Cy3 and Cy5) competitive array hybridizations (Agilent, Santa Clara, CA, USA). The same arrays were also used for competitive genome hybridization to compare genome content of RCC1216 and CCMP1516, using protocols previously defined for *E. huxleyi* with an earlier microarray system (Kegel et al., 2013). Illumina whole-genome data sets from strains CCMP1516, 92A, EH2 and 92F have been previously described (Read et al., 2013). The genomes of two newly isolated strains (CHC428 and CHC307;ENA study accession number PRJEB7726) were sequenced by Illumina-technology (Illumina, San Diego, CA, USA) at the Leibnitz Institute for Age Research (Jena, Germany) using the same methods.

The CCMP1516 JGI (Joint Genome Institute) whole-genome assembly (Read et al., 2013) and the Illumina genomic resequencing contig data sets were queried with full-length axonemal and cytoplasmic dynein heavy chain (aDHC and cDHC) genes of *Chlamydomonas reinhardtii* by TBLASTN (Altschul et al., 1997). Each contig with DHC homology was analyzed by BLASTX (Altschul et al., 1997) against Swiss-Prot (Boutet et al., 2007). Top reciprocal alignment (to *C. reinhardtii* homolog) assigned DHC paralog class, determined completeness of the DHC homolog and mapped conserved DHC functional modules, including the 6 AAA ATPase modules and stalk region, onto the *E. huxleyi* sequences. Illumina contigs of strain 92A encoded complete homologs of DHCs in all but one case, in which the complete homology was encoded on two contigs (see Supplementary Information). TBLASTN queries were conducted using the 92A contigs as queries against the CCMP1516 genome assembly and the Illumina data sets. The existence of major deletions affecting DHC homologous regions in the CCMP1516 genome was tested by mapping of the independently generated Illumina genome contigs of CCMP1516 against the JGI whole-genome assembly and, in five cases, by targeted PCR and resequencing. PCR and reverse transcriptase-PCR followed by end sequencing and mapping of 454 cDNA reads tested whether corresponding DHC homologous regions in RCC1216 and RCC1217 were intact and only expressed in the 1N phase.

To extend the genome content survey for presence/absence of two DHC genes (DHC1β and cDHC) crucial for flagellar formation found in RCC1216, RCC1217, 92A and 92F, but not in CCMP1516 and EH2, targeted PCR was used on genomic DNA from other *E. huxleyi* strains (a final total of 185 distinct parental genotypes, listed in Supplementary Table S17, including 86 new strains isolated in October–November 2011 and July
2013 from unenriched Southeast Pacific samples (locations specified in Supplementary Information) using a novel flow cytometer technique to distinguish calcified cells (von Dassow et al., 2012) (Supplementary Table S18). All strains were analyzed extensively by light microscopy for flagellated cell formation (Supplementary Information). In the PCR tests, two to four independent primer pairs were used. A gene was considered potentially absent only if none of these primer pairs amplified a product, while primers for a control gene (elongation factor 1α) successfully amplified product of expected size from the same DNA extract run in simultaneous PCRs.

Discriminant analyses based on 15 ocean variables derived from satellite chlorophyll, particulate inorganic carbon, sea surface temperature, spatial variability (decorrelation scales) and bathymetry were used to unveil the biogeographic/ecological preferences of the 99 RCC E. huxleyi strains checked for the presence/absence of flagellar genes (see Supplementary Figures S12 and S13, Supplementary Note and Supplementary Data S4 for detailed results). MODIS/Aqua satellite 2002–present level 3 data were downloaded from the Goddard Space Flight Center Ocean Biology Processing Group's OceanColor website (http://oceancolor.gsfc.nasa.gov) and ocean water depth from the ETOPO5 digital elevation database available at http://www.ngdc.noaa.gov/mgg/global/relief/ETOPO5.

Marine metagenomic sequence data were downloaded from CAMERA (158 metagenomes; http://camera.calit2.net/), National Center for Biotechnology Information/Sequence Read Archive (NCBI/SiAR) and GenBank (2 metagenomes; http://www.ncbi.nlm.nih.gov/), classified by region and searched by BLASTN using 8 genomic sequences of EhV viruses (NCBI: EhV84 (JF974290.1), EhV86 (NC_007346.1), EhV88 (JF974310.1), EhV201 (JF974311.1), EhV202 (HQ634145.1), EhV203 (JF974291.1), EhV207 (JF974317.1) and EhV208 (JF974318.1), all sequences low-complexity filtered (JF974291.1), EhV207 (JF974317.1) and EhV208 (JF974318.1), all sequences low-complexity filtered by DUST). BLASTN alignments ≥ 150 nt with ≥ 75% read coverage and ≥ 95% identity were considered as positive detections of EhV in metagenome data. Metagenome database hits were used as BLASTN queries against the NCBI nt/nt database to verify closest homology to EhV sequences.

Detailed methods are included in Supplementary Information.

**Results**

**Differential gene expression between life cycle stages**

Both 454 and microarray analyses revealed large expression differences between 1N and 2N cells. A complete discussion of the 1N vs 2N differences revealed is beyond the scope of this study, but we briefly highlight some of the substantial functional differences between the life-cycle stages to show that these new results are consistent with and strengthen previous studies (Supplementary Figures S1 and S2, Supplementary Table S2–S5 and Supplementary Note).

One of the striking differences between cell types is the presence of flagella in 1N cells (Klaveness and Pauxse, 1971; Klaveness, 1972; Green et al., 1996). As expected, the 82 genes previously identified as homologs of proteins with highly flagellar-specific functions (von Dassow et al., 2009) showed evidence of 1N-specific expression in microarray and 454 data. The differential expression was statistically highly significant for 56 genes in the microarray data and 46 genes in the 454 data.

The most obvious 2N-specific character is the intracellular precipitation of calcite coccoliths in large membrane-bound vesicles that are subsequently secreted (Paasche, 2001). Genes previously associated with these processes (von Dassow et al., 2009; Mackinder et al., 2011), including a putative HCO₃⁻ transporter in the SLC4 family, a CA₂⁺-Na⁺ exchanger (NCKX1) and a t-SNARE homolog that might be specifically involved in the exocytosis of coccoliths, all showed expression of highly specific to calcified 2N cells. The list of highly 2N-specific genes identified here included six additional SLC4 homologs, two more NCKX homologs and two more syntaxin/SNARE homologs (Supplementary Information) as well as other genes related to exo- and endocytosis, suggesting that 2N-specific versions of proteins involved in endo- and exocytosis might be involved in membrane trafficking specific to coccolith secretion.

**Erosion of haploid genome content in some diploid strains**

A significantly larger proportion of 1N-specific ESTs from RCC1217 were found to be missing in the CCMP1516 genomic data compared with ESTs that were specific to 2N (RCC1216) or nonspecific (Figure 1, Supplementary Figure S3 and Supplementary Table S6). Comparative genome hybridization, whole-genome Illumina resequencing and targeted PCR of ploidy-specific transcripts confirmed that genes displaying clear 1N-specific expression were more than twice as likely than 2N-specific or nonspecific genes to be absent or underrepresented in the genomic DNA of CCMP1516 (Figure 1, Table 1, Supplementary Tables S7–S11 and Supplementary Data S1). Overall, 1N-specific genes accounted for 60% of the genes that were present in RCC1216/1217 but undetected in CCMP1516, whereas highly 2N-specific genes accounted for 20% (Table 1 and Supplementary Table S10). Whole-genome Illumina resequencing of three other E. huxleyi strains revealed that 1N-specific genes were more likely to be lost in one 2N strain (EH2, formation of flagellated cells in culture not observed) but retained in two strains (92A and 92F, known to
form flagellated cells in culture) (Figure 1 and Supplementary Table S12).

Further analyses of the nature of the genes showing reduced competitive genome hybridization signals and no significant matches in the CCMP1516 genome confirmed that this strain has lost the ability to form functional motile 1N cells. Of the previously mentioned 82 genes from RCC1216/1217 coding for proteins involved in eukaryotic cilia or flagella (von Dassow et al., 2009) and displaying expected 1N-restricted expression patterns (Supplementary Tables S4 and S5), 19 (23%) were missing from the CCMP1516 genome (Supplementary Table S14).

A similar pattern of loss of flagella-related genes was observed in the EH2 strain, whereas all 82 flagellar genes were detected in the 92A and 92F strains (Supplementary Tables S13–S15).

Detailed examination of DHC homologs provided further evidence that CCMP1516 and EH2 have lost the ability to produce the flagellated 1N cell stage. A typical eukaryotic flagellum contains at least 10 paralogous aDHCs and 1 cDHC that are large proteins (4000–4600 amino acids) with a highly conserved modular structure: the 3000 amino-acid DHC motor domain consists of 6 AAA ATPase modules with a stalk (S) between domains A5 and A6, whereas the N-terminal 1000–1500 amino acids participate in protein interactions specific to each
regions of >97% nucleotide identity to this 92A contig were identified exclusively on scaffolds 68 and 529 in the CCMP1516 assembly. Scaffold 529 generally shows high homology and synteny to a section of the larger scaffold 68 (Figure 2a). Scaffold 68 completely lacks sequence sections for modules A5 and A6 of the motor domain and the N-terminal and C-terminal regions, and contains only a small section for the catalytic ATPase (A1). Scaffold 529 includes a shorter predicted gene where almost the entire motor domain and most of the N-terminal tail domain have been excised. Illumina contigs from CCMP1516 (Figure 2a) and targeted PCR (Figures 2b–d) confirmed these structures. Both long-range PCR with end sequencing and normal PCR (with independent primer sets) confirmed the section missing from CCMP1516 scaffold 529 was present in RCC1216 (2N) genomic DNA (Figure 2e and Supplementary Information). These sections were only expressed in the flagellated 1N strain RCC1217 (Figures 2b–f). No CCMP1516 Illumina contigs matched the entire region of 92A, paired-conig_3082, or specifically to the high DHC-homologous regions missing from scaffolds 68 and 529.

Thus, three independent methods agreed that the CCMP1516 genome has no complete outer arm aDHCα gene, but the corresponding gene is complete in strains RCC1216/1217, 92A and 92F.

Extending this strategy confirmed that the CCMP1516 genome does not contain a single complete DHC ortholog; all loci detected appear to be pseudogenes that have suffered independent deletions of large portions of the corresponding DHC genes that are complete in strains 92A and 92F. For scaffold pairs 48/399 and 22/722, both CCMP1516 Illumina genome resequencing contigs and targeted PCR confirmed the alternate structures of the DHC loci indicated by the JGI whole-genome assembly (Figure 3). Furthermore, CCMP1516 Illumina genome resequencing contigs were also consistent with the JGI whole-genome assembly at scaffold pairs 67/682 (inner arm DHC1b), 31/53 (inner arm DHC1b) and 43/329 (cDHC) (Supplementary Information). Querying the EH2 Illumina genome resequencing contigs found only small sections of DHC genes; in many cases the contigs appeared to exhibit excisions of major sections of DHC homology (Supplementary Information). The CCMP1516 and EH2 genomes thus appear to have lost the capacity to form flagellated cells but retain pseudogene ‘fossils’, suggesting loss of function in the recent evolutionary past.

**Loss of flagella among other E. huxleyi strains**

To unveil the evolutionary and ecological patterns of the apparent irreversible loss of the flagellated 1N life cycle in *E. huxleyi*, we applied a targeted PCR-based survey to 99 RCC strains (Supplementary Table S17 and Supplementary Data S2). DHC1b and a cDHC amplified from all 20 strains observed to produce flagellated 1N cells, despite wide

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**Table 1** Ploidy-dependent genomic difference between *Emiliania huxleyi* strains RCC1216/1217 and CCMP1516

|                | 1N specific | 2N specific | No ploidy specificity |
|----------------|-------------|-------------|-----------------------|
| Consensus absent | 471 (10.3%) | 153 (3.4%) | 154 (0.8%)            |
| Present, lower CNV | 804 (17.5%) | 538 (11.8%) | 1102 (6.1%)          |
| Others          | 3314 (72.2%) | 3661 (84.8%) | 16728 (93.0%)        |

**Contingency analysis of absence**

|                | X², d.f., P-value | Rel. risk, 1N vs 2N | Rel. risk, 1N vs no ploidy-spec. | Rel. risk, 2N vs no ploidy-spec. |
|----------------|-------------------|---------------------|-------------------------------|-------------------------------|
|                |                   |                     |                               |                               |
| X², d.f., P-value | 1170, 2, <0.0001 | 3.05 (2.56–3.65, <0.0001) | 2.0 (10.0–14.3, <0.0001) | 4.39 (3.15–4.90, <0.0001) |

**Contingency analysis of present, lower CNV**

|                | X², d.f., P-value | Rel. risk, 1N vs 2N | Rel. risk, 1N vs no ploidy-spec. | Rel. risk, 2N vs no ploidy-spec. |
|----------------|-------------------|---------------------|-------------------------------|-------------------------------|
|                |                   |                     |                               |                               |
| X², d.f., P-value | 763, 2, <0.0001  | 1.60 (1.44–1.77, <0.0001) | 3.16 (2.90–3.44, <0.0001) | 1.98 (1.80–2.18, <0.0001) |

Abbreviations: CNV, copy number variation; Rel., relative; spec., specific.

Ploidy-specific expression is determined by 50% difference in expression level between 1N and 2N cells (P<0.05: analysis of variance (ANOVA) for microarray results and Audic–Claverie statistics for 454 read counts). Apparent absence is indicated when the best BLASTN hit scores <100 against both JGI genome assembly and CCMP1516 Illumina paired-end read contigs and competitive genome hybridization (CGH) results gave <2-fold lower signal from CCMP1516 genomic DNA (gDNA) compared with RCC1216 gDNA. The global X² test calculates the significance of the difference among all expression categories. Relative risk ratios are given (with 95% confidence interval and P-value for x² or 2 x 2 tests) for 1N-specific versus 2N-specific and 1N-specific versus not ploidy-specific genes. Overall, ploidy-specific genes (considering both 1N specific and 2N specific) were more likely than not-ploidy-specific genes to be absent from the CCMP1516 genome by all three measures (χ² = 775, P < 0.0001) and more likely than not-ploidy genes to exhibit lower copy number (χ² = 628, P < 0.0001). The proportions of genome content differences attributable to 1N-specific genes (60%) and 2N-specific genes (20%) were calculated from the number of 1N-specific genes (471 genes) or 2N-specific genes (153 genes) found to be absent from CCMP1516 divided by the total genes identified as absent from CCMP1516 by the consensus of JGI, Illumina and CGH measures (775 genes).

DHC paralog (Asai and Koonce, 2001). Absence of a single DHC paralog leads to flagellar defects (Kamiya, 2002). In all, 12 distinct aDHC homologs and 1 cDHC homolog were expressed in RCC1217 (1N) cells (von Dassow et al., 2009), only 9 of which mapped to the CCMP1516 genome assembly. In addition, 19 loci encoding partial DHC homologs were identified in the CCMP1516 genome assembly (Supplementary Table S16), yet none was long enough to encode a complete DHC protein, thus appearing to be pseudogenes (Figure 2 and Supplementary Figures S4–S10). In all, 14 loci harboring DHC pseudogenes occurred on 8 pairs of homologous scaffolds, 3 occurred on a triplet of homologous scaffolds and 2 loci occurred on regions of large scaffolds that were not highly homologous to other scaffolds (Supplementary Table S16).

A complete homolog of *C. reinhardtii* outer arm DHCα (DYHA_CHLRE) was encoded on a single large contig in the 92A Illumina data set. Extensive
differences in geographic origin (Supplementary Data S3). In contrast, these genes failed to amplify from 37 2N *E. huxleyi* strains that have never been observed to produce flagellated 1N cells, even when two to four distinct primer pairs were used for each gene. Phylogenetic analysis based on the mitochondrial *cox1* and *cox3* genes for 83 strains indicated that loss of 1N-specific genes has occurred recently.
and independently in several lineages of the *E. huxleyi* haplotype group α (‘warm-water’ clade) (Beaufort et al., 2011; Hagino et al., 2011; Bendif et al., 2014) (Figure 4, Supplementary Figure S11).

**Biogeographic distribution of loss of flagella**

Loss of the flagellated phase in *E. huxleyi* was associated with warmer waters and lower amplitude cycles in chlorophyll and particulate inorganic carbon concentrations (Supplementary Figure S12) in the relatively stable low-latitude open oceans, whereas the biphasic life cycle was preserved at high latitudes and coasts (Figure 5). All *E. huxleyi* strains isolated from regions previously observed to exhibit annual EhV-controlled blooms, and/or where EhV sequences were detected in metagenomic databases (Figure 5), retained the genomic capacity to form flagellated 1N cells (Table 2; Fisher’s exact test, *P* < 0.0001). In low latitudes where *E. huxleyi* does not form blooms (see, for example, Moore et al., 2012), loss of the flagellate 1N phase appeared to increase in prevalence away from the coast. This was particularly clear in the Mediterranean Sea: all strains isolated from coastal sites in the Mediterranean retained flagellar genes, whereas 65% of those isolated far from shore lost either or both cDHC and DHC1β genes (Fisher’s exact test, *P* < 0.0001) (Supplementary Figure S14). The cDHC and/or DHC1β homologs were absent from 42% of the *E. huxleyi* strains (33 newly isolated strains from the southeast Pacific and 5 from RCC) isolated from low-nutrient, low-pCO₂ waters >500 km offshore. In contrast, both genes were amplified from 90% (Fisher’s exact test, *P* = 0.0004) of the 53 strains isolated from a coastal
site in the southeast Pacific with high-nutrient/high-pCO₂ upwelling water (Figure 5 and Supplementary Figure S14).

To independently check genome content predictions from this PCR assay, Illumina genome sequencing to \( \geq 50 \) coverage was applied to two oceanic strains (Supplementary Information). A significantly higher proportion of 1N-specific than 2N-specific and nonspecific genes were undetected by Illumina in strain CHC428, a strain from which neither DHC1β nor cDHC successfully amplified (Table 3). In contrast, Illumina sequencing detected 1N-specific and 2N-specific genes at similar levels in strain CHC307, a strain in which
Table 2 Comparison of *Emiliania huxleyi* strain origins with presence/absence of EhV sequences from environmental metagenome data sets

| Region                  | Total reads | EhV reads | Total E. huxleyi strains | No. of strains flagella-less | No. of strains with flagella |
|-------------------------|-------------|-----------|--------------------------|-----------------------------|-----------------------------|
| Norwegian Fjord         | 2 004 956   | 1126      | 1                        | 0                          | 1                           |
| Atlantic North          | 12 850 510  | 220       | 6                        | 0                          | 6                           |
| Atlantic South          | 1 574 807   | 90        | 1                        | 0                          | 1                           |
| North Atlantic European | 6 442 773   | 336       | 12                       | 0                          | 12                          |
| Margin                  |             |           |                          |                             |                             |
| Black Sea               | NA          | —         | 1                        | 0                          | 1                           |
| Pacific Tropical        | 15 788 772  | 0         | 12                       | 10                         | 2                           |
| Tasman Sea              | 1 431 368   | 11        | 5                        | 0                          | 5                           |
| Pacific South           | 2 605 665   | 0         | 16                       | 7                          | 9                           |
| Mediterranean           | 1 211 459   | 0         | 51                       | 27                         | 24                          |
| Humboldt Current System | 13 222 934  | 0         | 53                       | 5                          | 48                          |
| All regions EhV observed | 0           |           |                           |                             | 26                          |
| All regions EhV not observed | 51          |           |                           |                             | 83                          |

Abbreviations: EhV, *Emiliania huxleyi* virus; NA, not available. North Atlantic European Margin includes English Channel, Bay of Biscay, Celtic Sea and North Sea. The proportions of EhV reads/total reads and strains that lost flagellar genes were negatively correlated (Spearman’s coefficient -0.856, \( P = 0.005 \)).

*Non-canonical H4 might represent functions unique to coccolithophores. An F-box homolog (GJ15790), a large family of proteins interacting with the ubiquitin proteosome to play diverse roles in cell-fate decisions in animals and plants, including meiotic development (Lechner et al., 2006), was also not found in all three strains that had lost flagellar genes. A histone H4 gene encoding a non-canonical N-terminus (GJ10238) was previously identified as being present only in *RCC1216* and not in the clonal 1N phase daughter strain *RCC1217* (von Dassow et al., 2009). This non-canonical H4 was also identified in strains 92A, 92F and CHC307, but was not present in *CCMP1516*, EH2 or CHC428. It could be amplified from six further 2N strains that retained both cDHC and DHC1β genes and one 1N strain, but not from three other flagellated (1N) strains or from five calcified (2N) strains that appeared to have lost the key flagellar genes. This non-canonical H4 might be transmitted only in certain 1N mating types.

**Discussion**

Flagella are a conserved, ancestral and complex eukaryotic functional trait that has been lost in only a few major lineages (for example, red algae, seed plants and most fungi; Carvalho-Santos et al., 2011) and it is striking here to observe this functional trait being lost over relatively short evolutionary time-scales in sub-populations of *E. huxleyi*, a relatively young species. This observation challenges the practice of predicting phytoplankton function according to species identification based on either morphology or standard ribosomal DNA barcodes that are identical within the *E. huxleyi* lineage complex and even between *E. huxleyi* and its sister morphospecies *Gephyrocapsa oceanica* (Medlin et al., 1996). Flagellated 1N cells have substantially different functional traits and responses than calcified 2N *E. huxleyi* cells (see, for example, Houdan et al., 2005; Rokitta and Rost, 2012), and yet traditional morphological and molecular classifications group together genotypes that have differential capability to produce these very distinct functional forms. The initial mutation leading to loss of a motile 1N phase (and subsequent gradual accumulation of deletions of 1N-specific genes) may have occurred...
independently in distinct *E. huxleyi* lineages, as suggested by *cox* gene phylogeny and the low overlap between strains of which specific genes have suffered deletions.

Can *E. huxleyi* genotypes that have specifically lost 1N genes engage in meiosis and syngamy? Syngamy has never been observed in *E. huxleyi*, and cues that trigger meiosis or syngamy have not yet been identified in this species. However, non-motile 1N gametes would be severely encounter limited in the dilute planktonic environment of the open ocean. Similarly, syngamy in many other flagellated protists directly involves the flagella and/or flagellar bases (Ferris et al., 2005; Figueroa et al., 2006; Peacock et al., 2014). Both of these considerations imply that the loss of flagellar genes might be associated with loss of sex in *E. huxleyi* sub-populations.

The apparent structure of the CCMP1516 genome might be consistent with long-term absence of meiosis. Most of the DHC loci in the JGI genome assembly of CCMP1516 occurred on pairs of highly homologous and syntenic scaffolds that JGI identified as probably representing alternate structures of homologous chromosome pairs (‘diploid alleles’ in the terminology on the JGI genome portal at http://genome.jgi.doe.gov/). Eight of 11 homologous groups of DHC pseudogene loci in the JGI assembly occurred as pairs of exactly two scaffolds. In each case, the DHC-loci pairs corresponded to only a single highly homologous contig in the 92A and 92F (diploid) Illumina genome databases. The most parsimonious interpretation is that the two sets of homologous chromosomes in the CCMP1516 diploid genome have undergone distinct rearrangements, inversions and hemizygous deletions, although duplications and translocations likely also occurred. Considering the entire JGI genome assembly, the ‘diploid alleles’ scaffold pairs represent 32.9% of all unique structures, and might represent up to 19.7% of each haploid genome present in CCMP1516. Paired scaffold regions also show large gene content differences (that is, presence/absence and gene length differences). Such high divergence between the two haploid genomes in a diploid organism arises under long-term absence of meiosis, as seen in the long-term asexual genomes of bdelloid rotifers (Flot et al., 2013) and *Daphnia* (Xu et al., 2011).

The CCMP1516 genome encodes apparently intact (that is, not pseudogene) homologs of key proteins that mediate meiotic recombination, including *spo11*, *DMC1*, several other *Rad51* homologs, *Rad50* and MRE11 (Supplementary Information). However, *spo11* is also conserved in the genome of bdelloid rotifers, the oldest confirmed asexual eukaryote lineage, that exhibits pronounced ameiotic genome structure (Flot et al., 2013). These genes also appear to be involved in parthenogenetic recombination in the protist *Giardia*, the yeast *Candida* and the metazoan *Daphnia* (Forche et al., 2008; Schurko et al., 2009; Carpenter et al., 2012) that generates hemizygous mutations at rates several orders of magnitude higher than point mutations (Xu et al., 2011). A similar mechanism in *E. huxleyi* might account for some of the structural features of the CCMP1516 genome.

Conclusive evidence of asexuality in *E. huxleyi* genotypes that have lost key 1N genes still awaits. Distinguishing occasionally sexual and permanently asexual *E. huxleyi* populations by population genetics would require a much higher level of sampling than we have been able to achieve to date (only relatively small numbers of strains were successfully isolated from the highly dilute oceanic populations of interest). Alternatively, a reference whole-genome assembly from a strain that produces flagellated cells would be expected to have less structural divergence between homologous chromosomes compared with CCMP1516. Thus, for discussion of possible ecological and evolutionary forces acting to cause 1N-specific gene loss, we keep in mind both the possibility of complete loss of sex and the alternative that the 1N phase has been greatly reduced and modified.

Biogeographic comparison of the 185 *E. huxleyi* strains from diverse regions clearly showed that coastal and/or higher latitude, relatively productive and seasonally cycling parts of the oceans are populated with *E. huxleyi* strains that have maintained potential for a biphasic life cycle, whereas strains that have lost the capacity for formation of the 1N flagellated phase of the life cycle tend to originate from lower productivity offshore regions. In environments where EhV viruses are largely responsible for the demise of annual *E. huxleyi* blooms, biotic pressure from EhV might be expected to maintain sexual recombination in the host as part of the ‘Red Queen’ evolutionary arms race between host and pathogen driving positive selection. However, the observation that 1N cells are resistant to EhV (Frada et al., 2008) suggests a simpler mechanism: EhVs that specifically target one life-cycle stage directly select for a biphasic life cycle involving both free-living 2N and 1N phases, providing selective pressure to maintain the full complement of 1N-specific genes.

In subtropical/tropical coastal waters without blooms, biotic and abiotic parameters are highly variable (Uz and Yoder, 2004), conditions expected to favor sexuality (Becks and Agrawal, 2010) and/or niche separation of 1N and 2N phases (Coelho et al., 2007). In the absence of data about the actual ecological niche of 1N *E. huxleyi*, multiple biotic pressures might be invoked to maintain sexuality and/or a biphasic life cycle involving 1N flagellated cells in such environments, including grazing, allelopathy, parasites and perhaps low levels of EhV not detected in metagenomic surveys.

In offshore and lower latitude waters, physiological adaptation to oligotrophy does not seem to be driving the loss of *E. huxleyi* sex and life cycling.
First, genomic changes in the CCMP1516 and EH2 strains reflect accumulation of smaller-scale loss-of-function mutations in genes specific to the 1N stage rather than the massive genome streamlining adapting to oligotrophic life reported in eukaryotic and prokaryotic phytoplankton from the pico-size fraction (that is, \(< 2 \mu m\)) (Worden et al., 2009; Swan et al., 2013). Nuclear DNA contents of CCMP1516 and EH2 were not smaller than those of RCC1216 and 92A (Read et al., 2013) and Supplementary Note). Second, haploids are expected to have an intrinsic physiological advantage over diploids when nutrients are low (Coelho et al., 2007), and ecological distributions of life-cycle stages in other coccolithophores are consistent with this prediction (see, for example, Renaud et al., 2013). Yet, we show here that E. huxleyi apparently blocked in their diploid stages are thriving in oligotrophic open oceanic waters. Extensive studies of growth and survival characteristics under different temperature, nutrient and light conditions would be necessary to determine whether the observed changes are neutral or not with respect to physiological fitness. However, to allow the observed genomic changes to accumulate in low-latitude and offshore populations, deleterious effects must be lower in these environments.

As plankton biomass (chlorophyll) and turnover decreases, biotic interactions will be lower because of reduced encounter rates. Low-latitude open oceans also are more stable, showing lower seasonal variability than high-latitude and coastal regions. Loss of the flagellated 1N phase, and putative loss of sexuality, in E. huxleyi is associated with lower biomass in the open ocean realms and might be consistent with predictions that sex is not advantageous in very large populations experiencing low biotic pressure and low environmental variability (Otto, 2009), or it might be consistent with a lack of selective advantage to a biphasic life cycle in such environments.

In conclusion, major life-cycle alterations affecting functional traits in plankton may occur over relatively short evolutionary timescales in response to changes in biotic pressure. Because of interest in predicting the response of coccolithophores to ocean acidification, many laboratory studies have been conducted on monocultures of 2N E. huxleyi, but effects appear to vary among strains (see, for example, Riebesell et al., 2000; Iglesias-Rodriguez et al., 2008; Langer et al., 2009). The 1N phase may play a role in determining how particular E. huxleyi populations adapt, both as it represents a noncalciﬁed phase reacting diﬀerently to acidification (Rokitta and Rost, 2012), and because it might determine the role of sexual processes in adaptation to a changing ocean.

Conflict of Interest
The authors declare no conflict of interest.

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Author contributions

PvD and CdV initiated and PvD coordinated the research. PvD and IP performed all culture work. UJ, JK and PvD conducted and analyzed competitive genome hybridization (CGH) results. UJ contributed illumina sequencing data. PW and CdS conducted Sanger and 454 sequencing. HO conducted core bioinformatic analyses with targeted help from SA, UJ, PvD, ML and J-MC, E-MB, IP and CdV contributed phylogenetic analyses. PvD, DM-F, M-JGD and YH performed all PCR and targeted resequencing. SD and DMG contributed satellite data analyses and assisted with interpretation. PvD, HO, UJ, CdV and IP wrote the manuscript and all authors revised it.

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