# Targeted metagenomics and ecology of globally important uncultured eukaryotic phytoplankton

Marie L. Cuvelier<sup>a,b</sup>, Andrew E. Allen<sup>c,1</sup>, Adam Monier<sup>a,1</sup>, John P. McCrow<sup>c</sup>, Monique Messié<sup>a</sup>, Susannah G. Tringe<sup>d</sup>, Tanja Woyke<sup>d</sup>, Rory M. Welsh<sup>a</sup>, Thomas Ishoey<sup>c</sup>, Jae-Hyeok Lee<sup>e</sup>, Brian J. Binder<sup>f</sup>, Chris L. DuPont<sup>c</sup>, Mikel Latasa<sup>g</sup>, Cédric Guigand<sup>b</sup>, Kurt R. Buck<sup>a</sup>, Jason Hilton<sup>b</sup>, Mathangi Thiagarajan<sup>c</sup>, Elisabet Caler<sup>c</sup>, Betsy Read<sup>h</sup>, Roger S. Lasken<sup>c</sup>, Francisco P. Chavez<sup>a</sup>, and Alexandra Z. Worden<sup>a,b,2</sup>

<sup>a</sup>Monterey Bay Aquarium Research Institute, Moss Landing, CA 95039; <sup>b</sup>Rosenstiel School of Marine and Atmospheric Science, Miami, FL 33149; <sup>c</sup>J. Craig Venter Institute, San Diego, CA 92121; <sup>d</sup>US Department of Energy Joint Genome Institute, Walnut Creek, CA 94598; <sup>e</sup>Department of Biology, Washington University, St. Louis, MO 63130; <sup>f</sup>Department of Marine Sciences, University of Georgia, Athens, GA 36072; <sup>g</sup>Institut de Ciències del Mar (CSIC), E-08003 Barcelona, Spain; and <sup>h</sup>Department of Biological Sciences, California State University, San Marcos, CA 92096

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Among eukaryotes, four major phytoplankton lineages are responsible for marine photosynthesis; prymnesiophytes, alveolates, stramenopiles, and prasinophytes. Contributions by individual taxa, however, are not well known, and genomes have been analyzed from only the latter two lineages. Tiny "picoplanktonic" members of the prymnesiophyte lineage have long been inferred to be ecologically important but remain poorly characterized. Here, we examine pico-prymnesiophyte evolutionary history and ecology using cultivation-independent methods. 18S rRNA gene analysis showed picoprymnesiophytes belonged to broadly distributed uncultivated taxa. Therefore, we used targeted metagenomics to analyze uncultured pico-prymnesiophytes sorted by flow cytometry from subtropical North Atlantic waters. The data reveal a composite nuclear-encoded gene repertoire with strong green-lineage affiliations, which contrasts with the evolutionary history indicated by the plastid genome. Measured pico-prymnesiophyte growth rates were rapid in this region, resulting in primary production contributions similar to the cyanobacterium Prochlorococcus. On average, picoprymnesiophytes formed 25% of global picophytoplankton biomass, with differing contributions in five biogeographical provinces spanning tropical to subpolar systems. Elements likely contributing to success include high gene density and genes potentially involved in defense and nutrient uptake. Our findings have implications reaching beyond pico-prymnesiophytes, to the prasinophytes and stramenopiles. For example, prevalence of putative Ni-containing superoxide dismutases (SODs), instead of Fe-containing SODs, seems to be a common adaptation among eukaryotic phytoplankton for reducing Fe quotas in low-Fe modern oceans. Moreover, highly mosaic gene repertoires, although compositionally distinct for each major eukaryotic lineage, now seem to be an underlying facet of successful marine phytoplankton.

comparative genomics | primary production | prymnesiophytes | marine photosynthesis | haptophytes

**G** lobal primary production is partitioned equally among terrestrial and marine ecosystems, each accounting for  $\approx$ 50 gigatons of carbon per year (1). The phytoplankton responsible for marine primary production include the cyanobacteria, *Prochlorococcus* and *Synechococcus*, and a multitude of eukaryotic phytoplankton, such as diatoms, dinoflagellates, prasinophytes, and prymnesiophytes (2–4). Most oceanic phytoplankton are "picoplanktonic" (<2–3 µm diameter) and have high surface area to volume ratios, an advantage in open-ocean low-nutrient conditions (5–8). Despite the importance of eukaryotic phytoplankton to carbon cycling only six genomes have been sequenced and analyzed comparatively, all being from diatoms and prasinophytes. These revealed greater differentiation than anticipated on the basis of 18S rRNA gene analyses (9–11). The observed genomic divergence is associated with major differences in physiology and niche adaptation (10).

Pigment-based estimates indicate that prymnesiophytes, also known as haptophytes, are broadly distributed and abundant.

Oceanic prymnesiophytes are thought to be small owing to high levels of prymnesiophyte-indicative pigments in regions where most Chl *a* (representing all phytoplankton combined) is in the <2-µm size fraction (6, 12). Six picoplanktonic prymnesiophytes exist in culture (6, 7) but prymnesiophyte 18S rDNA sequences from <2–3-µm size-fractioned environmental samples typically belong to uncultured taxa (6, 13–15). As a whole, this lineage reportedly diverged from other major eukaryotic lineages early on, 1.2 billion years ago (16), and their overall placement among eukaryotes is uncertain (4, 16). They are extremely distant from phytoplankton with published genomes. Thus, although inferences exist regarding their importance and evolutionary history, uncertainties surround even the most basic features of oceanic pico-prymnesiophytes, such as cell size, biomass, growth rates, and genomic composition.

One approach for gaining insights to uncultivated taxa is metagenomics. However, unicellular eukaryotes have larger genomes and lower gene density than marine bacteria and archaea and are less abundant, making efficient sequence recovery difficult by seawater filtration. Parsing of eukaryotic data from diverse communities is particularly problematic owing to the paucity of relevant reference genomes. Selection of DNA from an uncultivated target microbe(s) (e.g., by fosmid sequencing or using cells sorted by flow cytometry) obviates bioinformatic parsing issues and has revealed unique gene complements in uncultured prokaryotes (17, 18).

To address uncertainties regarding pico-prymnesiophyte ecology, we integrated a suite of cultivation-independent methods. Targeted metagenomics was developed to investigate diversity and genomic features of uncultivated pico-prymnesiophytes. Growth rates were measured and used to assess primary production in the same region. Building on this contextual dataset, biomass contributions were evaluated across provinces spanning tropical to subpolar systems, providing a comprehensive view of global importance and latitudinal variations.

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Data deposition: The sequences reported in this paper have been deposited in the Gen-Bank database (accession nos. HM581528–HM581638 and HM565909–HM565914). Other scaffolds with predicted genes from this Whole Genome Shotgun/454 project have been deposited at DNA Data Bank of Japan/European Molecular Biology Laboratory/GenBank under the accession no. AEAR00000000. The version described in this paper is the first version, AEAR0100000.

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<sup>&</sup>lt;sup>1</sup>A.E.A. and A.M. contributed equally to this work.

<sup>&</sup>lt;sup>2</sup>To whom correspondence should be addressed. E-mail: azworden@mbari.org.

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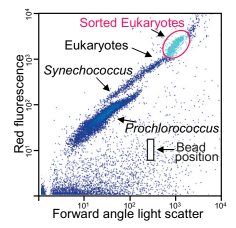
## **Results and Discussion**

**Eukaryote-Targeted Metagenomics Approach and Diversity Context.** Photosynthetic picoeukaryote populations were sorted by flow cytometry (hereafter "sorted" or "the sort") based on scatter and autofluorescence characteristics from two subtropical North Atlantic samples collected in the Florida Straits (Fig. 1, Fig. 2 *Inset*, and *SI Materials and Methods, Sections 1 and 2*). Whole-genome amplification (19) was performed on sorted target populations (*SI Materials and Methods, Section 3*), providing unprecedented access to pico-prymnesiophyte DNA.

The sorted pico-prymnesiophytes were distinct from cultured taxa but closely related to environmental sequences from native populations. 18S rDNA clone libraries built from the sort DNA were analyzed within the context of  $<2-3 \mu m$  size-fractionated clone libraries from the Florida Straits, the Sargasso Sea, the Pacific Ocean (Fig. 2, SI Materials and Methods, Section 4, Fig. S1A, and Table S1), and published data. Similar to other studies (6, 13, 15), the vast majority of sequences were from uncultured prymnesiophytes. The majority of prymnesiophyte sequences from the Florida Straits Station 04 sort belonged to environmental group 8 (111 clones, 99-100% identity; Fig. S1B), also present in the easterly Station 08 sort. Lower levels of group 3 were detected in the Station 04 sort (7 clones, 99-100% identity), along with sequences at the tip of the tree (e.g., group 15; SI Materials and Methods, Section 5). Group 8 was also seen in the Sargasso Sea (14) and on multiple dates in the Florida Straits (Table S2). Group 8 had only 93% 18S rDNA identity to any cultured organism (to Chrysochromulina), and phylogenetic placement was unresolved (no bootstrap support).

The observed level of group 8 18S rDNA divergence has important implications for gene content. The diatoms *Thalassiosira pseudonana* and *Phaeodactylum tricornutum*, with 90% 18S rDNA identity, share only 30–40% of their genes and occupy fundamentally different niches (10). Of the four picoeukaryote genomes (all prasinophytes), two *Micromonas* isolates, with 97% 18S rDNA identity, have 69.5% DNA identity over aligned genome regions, sharing at most 90% of their protein-encoding genes (11). Although an unpublished genome of the coccolit thophore *Emiliania huxleyi* is available, this prymnesiophyte is larger than the pico-prymnesiophytes and expected to occupy a distinct niche (8, 16). Furthermore, the soft-bodied prymnesiophytes in the sort were distant from coccolithophores (Fig. 2), a group known for their calcium carbonate plates (16).

Given the differences between the group 8–enriched sort population and cultured taxa, we used targeted metagenomics to discover genomic features of environmentally relevant picoprymnesiophytes. Station 04 sort DNA was sequenced by 454-FLX



**Fig. 1.** Forward angle light scatter and chlorophyll-derived red fluorescence characteristics of picophytoplankton groups (at Florida Straits Station 04; Fig. 2 *Inset*, blue arrow), including the metagenome population (magenta circle) and showing the position of 0.75- $\mu$ m bead standards (black box, not added to actual sort to avoid contamination).

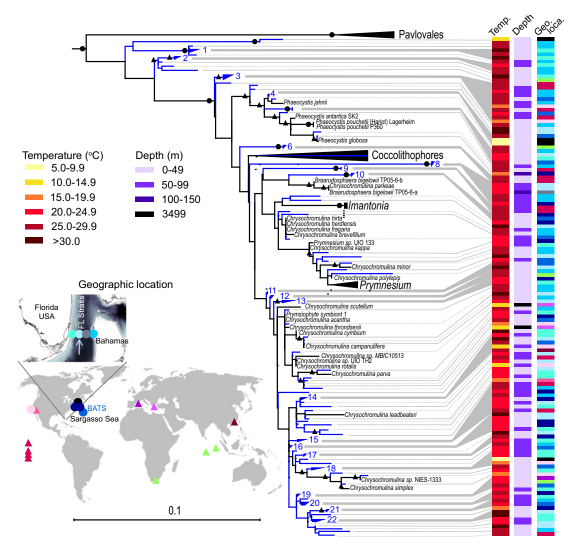
and Sanger technologies (*SI Materials and Methods, Section 5*). Genes were modeled on assembled scaffolds and then screened phylogenetically using the *E. huxleyi* nuclear genome. For selection, half the identifiable genes on a scaffold had to clade directly with *E. huxleyi*, to the exclusion of gene sequences from all other taxa (*SI Materials and Methods, Section 6*). This ensured that only prymnesiophyte-derived scaffolds were further analyzed, because the stramenopile *Pelagomonas* fell partially in the same flow-cytometric population (*SI Materials and Methods, Section 5*). Seventy-one percent of genes on selected scaffolds were sistered by *E. huxleyi* genes, demonstrating screening rigor; only this scaffold subset (2 MB of genes on pico-prymnesiophyte scaffolds seemed to be missing from *E. huxleyi*, supporting their distance from coccolithophores.

**Gene Density and Genome Size Predictions.** Prymnesiophytes are currently placed in the controversial eukaryotic Supergroup Chromalveolata along with stramenopiles (e.g., diatoms) and alveolates (e.g., dinoflagellates) but separate from the Supergroup Archaeplastida, which includes prasinophytes and all green-lineage organisms (e.g., land plants). However, their genome characteristics are unknown and phylogenetic placement uncertain. We found high gene density in the pico-prymnesiophyte metagenome (Fig. 3*A* and *SI Materials and Methods, Section 7*), akin to pico-prasinophytes, which purportedly underwent genome streamlining to optimize life in oligotrophic niches (9, 11). GC content was 60%.

Genome size was calculated using multiple methods accommodating the composite nature of the sorted pico-prymnesiophyte population (*SI Materials and Methods, Section 7*). The first involved a model linking the size distribution of euKaryotic Orthologous Groups of proteins (KOG), allocation of functions across those families and total gene content for 12 protistan reference genomes. This rendered an average genome content of 12,711 ( $\pm$ 1,145) genes (Fig. S2.*A* and *B*), such that the 1,624 pico-prymnesiophyte nuclearencoded genes identified constituted 13% of those in a representative genome. Results were similar (11,600 total genes; *SI Materials and Methods, Section 7*) using a method based on counts of 132 near single-copy core genes in other eukaryotes. Diatom genome content is comparable,  $\approx$ 10,000–14,000 genes (10), whereas the pico-prasinophytes contain 8,000 to just over 10,000 genes (9, 11).

Mosaic Gene Repertoires. Evolutionary and functional aspects of chloroplast and nuclear genomes were analyzed using gene content. 16S rRNA genes revealed pico-prymnesiophyte chloroplast genome scaffolds; the largest (scaffold C19847), containing 45 protein-encoding genes among others (Tables S3 and S4), seemed to be from group 8 (SI Materials and Methods, Sections 5 and 6, and Figs. S1B and S3A). Phylogenetic analysis of 22 concatenated plastid genome-encoded genes conserved across all lineages placed this pico-prymnesiophyte within the Chromalveolata, between cryptophytes and stramenopiles, and directly sistered by E. huxleyi (SI Materials and Methods, Section 7, and Fig. S3B), the only published prymnesiophyte plastid genome (20). This placement corresponded to that for E. huxleyi in other plastid phylogenies (21), but with greater support. Nucleotide level comparison of C19847 with E. huxleyi showed large-scale rearrangements akin to the divergent diatoms (Fig. S3C). Gene-level synteny was relatively conserved with E. huxleyi (Fig. S3D).

In contrast, global analysis of the 1,624 predicted picoprymnesiophyte nuclear-encoded genes revealed a distinctive evolutionary signature relative to plastid genome-encoded elements. Phylogenomic analyses of nuclear-encoded genes indicated that these phytoplankton are evolutionarily situated between the greenlineage and stramenopiles, with greater apparent affinity to the former. The gene repertoire shared a higher degree of overlap with prasinophytes (Fig. 3B). Even among pico-prymnesiophyte genes with homologs in both prasinophytes and stramenopiles, similarity was generally higher to prasinophytes. Of 352 phylogenetic trees constructed herein that contained at least one green-lineage organism and one stramenopile, the genetic distance between the



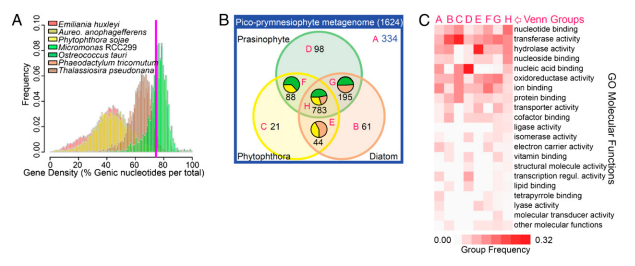
**Fig. 2.** Maximum-likelihood reconstruction of (blue lines) environmental and (black) cultured prymnesiophyte 18S rDNA sequences from two sorts and 25 size-fractionated samples from discrete depths, dates, and locations (circles; Table S1) and previous publications (triangles) (*SI Materials and Methods, Section 4*). The Station 04 sort (blue arrow) was in the Gulf Stream core. A single representative was used for redundant sequences within each library. Supported clades composed of only environmental sequences were collapsed after tree building when of  $\geq$ 99% identity (blue groups 1–22; Table S2). Node support is shown for (black circles) 100% and (black triangles)  $\geq$ 75% support. Uncultured prymnesiophytes have also been seen recently in the South Pacific (40).

pico-prymnesiophyte and the former was shorter twice as often (Fig. S4A). Across all trees, the mean relative genetic distance was 6.5% shorter between pico-prymnesiophytes and Archaeplastida, compared with stramenopiles, calling into question the current, largely plastid genome-based, phylogenetic placement of this lineage.

More than 75% of pico-prymnesiophyte genes assigned to KOGs were in the largest KOG families (top 20%), whereas for other marine phytoplankton this percentage was lower (SI Materials and Methods, Section 7). This high level of KOG redundancy reflected the mosaic nature of pico-prymnesiophyte gene repertoires. The functional KOG repertoire straddled prasinophytes and stramenopiles, such that several expansions in the metagenome seemed to be absent from one or both of these other lineages. Presence of more than one pico-prymnesiophyte taxon in the sort does not explain this redundancy, because it would require disproportionate sampling of the same KOGs, to the exclusion of others, from each pico-prymnesiophyte genome. Families with multiple representatives included nudix hydrolases and arylsulfatases, not found in prasinophytes, but present in metazoa and bacteria (arylsulfatases are also in stramenopiles) (Table S5). Polyketide synthases were also found and present in prasinophytes and bacteria but missing from diatoms.

The discontinuity in gene content could help explain puzzling ambiguities in prymnesiophyte evolution. The chloroplast genome would drive the relationship toward a red-lineage secondary endosymbiosis (Chromalveolata; Fig. S3B), whereas the nuclear genome retains features of a green host (Fig. 3B and Fig. S44). Of pico-prymnesiophyte genes that appeared more similar to Archaeplastida than to stramenopiles, 55% were closer to streptophytes, particularly early diverging plants, suggesting a strong ancestral green-lineage influence in the prymnesiophyte host organism's gene pool; only 45% were closer to green algae (Fig. S4B). Alternatively, like diatoms (22), prymnesiophytes may have obtained green-lineage genes from an ancient cryptic endosymbiont. Although the paucity of eukaryotic phytoplankton genomes may influence results, nuclear-encoded marker genes from larger cultured prymnesiophytes also show strong green-lineage affiliations (3). We anticipate that E. huxleyi genome analysis will confirm that the mosaic gene repertoire reported here is a lineage-wide characteristic.

**Functional Gene Repertoire.** Pico-prymnesiophyte nuclear-encoded genes also showed differences in functional composition (Fig. *3C*). Although all transcription factor families recovered were



**Fig. 3.** Characteristics of the pico-prymnesiophyte metagenome. (*A*) Gene density histograms as the proportion of nucleotides in genes over all chromosomes within an incrementally sampled sliding window in prasinophytes and photosynthetic (diatoms and *Aureococcus*) and nonphotosynthetic (*Phytophthora*) stramenopiles. Average gene density on pico-prymnesiophyte scaffolds was 74% (magenta line). (*B*) Venn diagram of pico-prymnesiophyte genes in relation to prasinophyte (green, *Micromonas* and *Ostreococcus*), diatom (beige), and *Phytophthora* (yellow) genes by BLASTP (e-value  $\leq 1.0e^{-9}$ ). Numbers indicate gene counts in each Venn group (magenta letters), whereas pie charts show relative best BLAST proportions for overlapping groups. Some pico-prymnesiophyte genes (D) comparison of functional profiles of pico-prymnesiophyte-specific genes with those from each Venn group by high-level Gene Ontology (GO) molecular functions. Relative proportions are shown for each GO function within a Venn group such that each column represents 100%.

also in Archaeplastida and stramenopiles, higher proportions of specific regulators were observed, such as chromatin remodeling genes (Tables S5 and S6), which play roles in meiosis and "life-style" changes. These included divergent SET-domain proteins and SNF2/helicases as chromatin remodeling factors (23). Distinct SUV39 subfamily members (SET) identified were not found in stramenopiles (Fig. S54). This family is responsible for heterochromatin formation (23) and could enable control over invasive entities like transposons and viruses.

Other features corresponded to life in modern oceans, for example, a Ni-containing superoxide dismutase (Ni-SOD). SODs are vital to photosynthetic organisms, scavenging toxic superoxide radicals generated by multiple metabolic pathways including photosynthesis. Isoforms use different metals at the active site, influencing trace metal requirements (24). Presence of sodN, which encodes Ni-SOD, corresponds with absence of Fe-SOD-encoding genes in many marine bacterial genomes, and Ni-SOD seems to be more prevalent in open-ocean strains (25). This results in replacement of some cellular Fe demand with Ni, a valuable adaptation considering 10-fold higher Ni concentrations in oceanic waters (25). We identified putative sodN genes in the pico-prymnesiophyte metagenome, the four pico-prasinophyte genomes (see also ref. 9) and P. tricornutum (Fig. \$5B). T. pseudonana did not seem to encode sodN, but rather a Fe-SOD, correlating with its coastal distribution and higher iron quotas than P. tricornutum (26). Fe-SOD-encoding genes were not found in the sodN-containing phytoplankton genomes. Presence of putative sodN genes in oceanic phytoplankton from three major eukaryotic lineages suggests that replacement of Fe-SOD with Ni-SOD may be a common adaptation to the lower availability of iron in modern oceans than in past times (27).

Several domains involved in uptake of large substrates, such as proteins and nucleic acids, as well as salvage of nucleosides, were represented more highly in the pico-prymnesiophyte metagenome (Fig. 3C). These could facilitate uptake of otherwise inaccessible nutrients. For example, a member of the amino acid/ polyamine/organocation superfamily is a likely transporter for the nitrogen-rich guanine derivative xanthine, potentially important under nitrogen-limiting conditions. A plant-like putative acid phosphatase (Table S5), which likely cleaves intracellular orthophosphoric-monoesters to phosphate, could also be involved in nutrient scavenging, although this role is speculative. Overall, features of the metagenome suggest adaptations associated with survival in oligotrophic environments.

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**Environmental Importance.** Increased stratification and lower nutrient concentrations predicted under some climate-change scenarios are hypothesized to create conditions favoring picophytoplankton over larger species (11). However, questions remain regarding the ecological importance of pico-prymnesiophytes in today's conditions, against which physiological stressors and future changes can be assessed. To establish their roles, we developed a contextual metadataset for the pico-prymnesiophyte metagenome, including distributions at the subtropical North Atlantic sites (Fig. 2, *Inset*) by FISH. Biomass contributions were compared with the overall picophytoplankton community, specifically, other picoeukaryotes, *Synechococcus* and *Prochlorococcus* (*SI Materials and Methods, Sections 2, 9, and 10*).

Pico-prymnesiophyte contributions in the Sargasso Sea were roughly equivalent at the surface and deep chlorophyll maximum (DCM), representing 23% and 21% of picophytoplankton carbon, respectively. Two pico-prymnesiophyte size classes were evident, cells of  $1.9 \pm 0.4 \times 2.1 \pm 0.3 \mu m$  (n = 89) and  $2.8 \pm 0.6 \times 3.4 \pm 0.5 \mu m$  (n = 127) (Table S7). In the Florida Straits their contributions were typically higher in surface waters, although more nutrients were presumably available at the DCM (Fig. S64). Here, 90%  $\pm$ 9% and 87%  $\pm$  13% of prymnesiophytes were <3  $\mu m$ , averaged over 1 y, at the surface and DCM, respectively. The direct-count– based biomass approach resulted in lower pico-prymnesiophyte DCM contributions than estimated by HPLC (Fig. S6B). However, these methods gave similar surface trends, supporting the HPLCbased inference that picoplanktonic taxa form most prymnesiophyte biomass in open-ocean surface waters (6, 12, 28).

Pico-prymnesiophyte specific growth rates showed that these tiny eukaryotes can grow rapidly, amplifying contributions to primary production. Rates were measured by the dilution method and direct counts (*SI Materials and Methods, Section 11*), which render growth rates close to in situ cell cycle–based rates for taxa amenable to the latter analysis (29, 30). Pico-prymnesiophyte growth rates in the Sargasso Sea were high at 15 m (1.12 d<sup>-1</sup>,  $r^2 = 0.87$ , P < 0.07) and lower at 70 m (0.29 d<sup>-1</sup>,  $r^2 = 0.73$ , P < 0.07). *Prochlorococcus* grew more slowly (0.63 d<sup>-1</sup>,  $r^2 = 0.54$ , P < 0.01) than pico-prymnesiophytes at the surface, but faster (0.60 d<sup>-1</sup>,  $r^2 = 0.61$ , P < 0.001) at depth. Because the pico-prymnesiophyte data constitute the first specific growth rates reported, for experimental validation, we compared *Prochlorococcus* growth rates with previous direct count-based rates from the same region and time of year, which were similar (0.52 d<sup>-1</sup>) (30).

#### Table 1. Average surface biomass of picophytoplankton groups in five biogeographical provinces

Latitude	Ocean area (× 10 <sup>12</sup> m <sup>2</sup> )	Samples (n)	Biomass (µg C L <sup>-1</sup> )			
			Prochlorococcus	Synechococcus	"Non-prym" picoeukaryotes	Pico-prymnesiophytes
60°–45°N	13.22	8	0.1 (0.1)	2.4 (1.1)	3.0 (1.7)	7.0 (4.7)
45°–20°N	47.18	24	3.7 (3.0)	0.5 (0.5)	1.0 (2.7)	2.0 (1.5)
20°N–20°S	122.60	59	8.4 (2.8)	1.0 (1.1)	1.8 (1.7)	1.8 (0.9)
20°–45°S	75.35	19	3.1 (2.0)	0.4 (0.6)	3.7 (1.4)	1.8 (0.9)
45°–65°S	48.69	11	0.3 (0.4)	1.1 (1.9)	3.9 (2.8)	5.5 (4.9)

Area varies over latitudinal zones owing to the influence of land masses; sample number is also shown, with values from sites sampled seasonally averaged and counted here as 1 sample (Table 58). Values in parenthesis reflect SDs.

We combined biomass with specific growth and mortality rates to estimate primary production. Despite lower pico-prymnesiophyte abundance, their combined greater cellular biomass and faster growth led to primary production levels comparable to Prochlorococcus at 15 m (1.1 and 1.8 µg C L<sup>-1</sup> d<sup>-1</sup>, respectively; SI Materials and Methods, Section 11). Pico-prymnesiophyte primary production was almost 4-fold higher than that of all other picoeukaryotes (0.27  $\mu$ g C L<sup>-1</sup> d<sup>-1</sup>). Total primary production at the Bermuda Atlantic Time-series Study (BATS; Fig. 2, Inset) in mid-May, before our experiments, and just afterward, in mid-June, was 7.81 and  $0.96\,\mu g\,C\,L^{-1}\,d^{-1},$  respectively, at 20 m (31). Seasonal stratification developed during this period, leading to lower production by mid-June. Over a decade, BATS total primary production occasionally ranged up to 6  $\mu$ g C L<sup>-1</sup> d<sup>-1</sup> at 20 m, but was typically 2–4  $\mu$ g C L<sup>-</sup>  $d^{-1}$ each June; the pico-prymnesiophyte value constitutes 25–50% of that. Likewise, a flow cytometrically defined picoeukaryote population ( $73\% \pm 2\%$  prymnesiophyte cells, over 20 stations), was recently shown to perform  $25\% \pm 9\%$  of primary production in the North East Atlantic, ranging up to 38% (32). Plastid-16S rRNA genes were evaluated at one station in that study and showed uncultured prymnesiophyte taxa, as found herein.

**Global Contributions and Latitudinal Gradients.** Our results, taken with those from the North East Atlantic (32), point to significant pico-prymnesiophyte contributions in the subtropical North Atlantic. However, variations between oceans and latitudinal gradients translate to major biotic differences, and the respective communities will likely respond to change differently. To assess global surface biomass contributions by pico-prymnesiophytes we counted and sized cells in five biogeographical provinces, specifically subpolar (high-latitudes) and subtropical-temperate (midlatitudes) systems, as well as the tropics (low latitudes) (*SI Materials and Methods, Sections 1, 2, 9, and 10*, and Table S8).

Globally, pico-prymnesiophytes averaged  $2.6 \pm 1.8 \ \mu g \ C \ L^{-1}$ when the areal extent of each province was accounted for (Table 1). This amounted to  $\approx 50\%$  of that of *Prochlorococcus* ( $4.7 \pm 2.1 \ \mu g \ C \ L^{-1}$ ), which was less evenly distributed and nearly absent from cold waters. The considerable biomass of pico-prymnesiophytes was again in part due to larger cell size than other picophytoplankton (6), and their contributions were less obvious by abundance alone (Table S8); *Prochlorococcus*, for instance, is orders of magnitude more abundant in low- and mid-latitudes but is also much smaller.

Biomass showed a strong latitudinal signal. In high latitudes, pico-prymnesiophytes dominated, comprising 50–56% of pico-phytoplankton biomass (Fig. 4, Table 1, and Table S8). Relative contributions in mid-latitudes were modified by variations in other groups. Pico-prymnesiophyte biomass per liter was maximal in the northern subpolar province, but the massive extent of the Southern Ocean, relatively unimpeded by land, rendered their greatest contributions to global biomass in the southern subpolar province.

Pico-prymnesiophyte contributions were lowest in the tropics  $(1.8 \ \mu g \ C \ L^{-1})$ , comprising approximately one fifth (21%) of *Prochlorococcus* biomass (Table 1). These in situ observations were at odds with a recent report suggesting that pico-prymnesiophytes are more important than *Prochlorococcus* in low latitudes, based on an algorithm relating satellite surface Chl to prymnesiophyte

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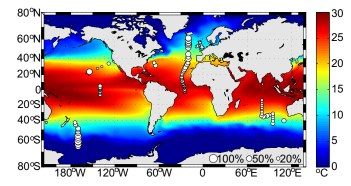
pigments and their contribution to total Chl (15). Validation (of ref. 15) would result in major reevaluation of tropical systems where the streamlined genome, small size, and low nutrient quotas of *Prochlorococcus* seem highly advantageous given extended periods of stratification (see, e.g., ref. 17). The significant discrepancy with our results may stem from issues surrounding the algorithm-based approach used in ref. 15, such as (*i*) the fact that other line ages can contain the prymnesiophyte-indicative marker pigment (28, 33) (*SI Materials and Methods, Section 8*), (*ii*) a variable relationship between a specific pigment content and surface Chl, or (*iii*) not partitioning contributions by organism size and the fact that HPLC samples are not size-fractionated. Furthermore, our tropical surface direct-count-based *Prochlorococcus* biomass data corresponded well with that from HPLC (28, 34), which, for *Prochlorococcus*, is less prone to such caveats.

Pico-prymnesiophytes seem to be highly successful and show signs of optimization to open-ocean conditions. Features within the metagenome remain difficult to relate to niche differentiation given 37% genes of unknown function, similar to many genomes. This lack of functional understanding is perhaps the greatest impediment to connecting genomes to organism physiology and response. The research herein positions us to explore the function of such genes in situ.

#### Conclusions

Soft-bodied prymnesiophytes survived the K/T boundary mass extinctions (16), indicating that taxa akin to those analyzed herein were resilient to perturbation. Surface water warming has now been correlated with increases of picophytoplankton in the Arctic Ocean (35) and will presumably impact lower latitudes. However, the success of small prymnesiophytes and their contributions in future times are linked to evolutionary history and genetic makeup, as well as the rate and extent of perturbations experienced.

Our results, showing genomic features of pico-prymnesiophytes, their rapid growth and significant global contributions, provide



**Fig. 4.** Global surface biomass contributions of prymnesiophytes as percentage of total picophytoplankton carbon, represented by bubble size (scaling at lower right). Sea surface temperature represents 1° increments averaged monthly over 18 y; note differences in the five biogeographical provinces.

several key advancements. Although pico-prymnesiophytes are clearly diverse, aspects of composition, potential streamlining, and functional attributes revealed by targeted metagenomics provide insights into their evolutionary success. Together with data on prasinophytes and stramenopiles (10, 11), our analysis of the picoprymnesiophyte metagenome indicates that mixed-lineage gene repertoires are a transcendent property of successful phytoplankton in modern oceans, not a rare feature. These mosaic gene repertoires seem to be compositionally distinct for each lineage, influenced by disparate sources of the tree of life, or differential gene loss from an ancestor, and presumably factors driving niche differentiation. Furthermore, the complexity of marine microbial communities makes it difficult to determine taxa critical to ecosystem processes and global biogeochemical models; our work highlights the need to prioritize pico-prymnesiophytes. Finally, it opens the door for research on the physiology and response capabilities of uncultivated members of this ancient primary producer lineage.

### **Materials and Methods**

Methods details are in *SI Materials and Methods*. Samples were from the North, Equatorial, and South Atlantic, the North, Equatorial, and West Pacific, and the Indian Ocean(s). 18S rDNA clone libraries (14) were constructed from subtropical Atlantic and North West Pacific size-fractionated samples, or from cells sorted by flow cytometry.

Approximately 300 cells from target populations in the subtropical North Atlantic were sorted by flow cytometry. DNA from two populations was amplified by multiple displacement amplification (19) and used for PCR-based clone libraries and, for one sample, metagenomic sequencing. Metagenomic data were assembled in a two-step high-stringency process and only scaffolds resulting from the second stage considered further. Scaffolds were screened

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by phylogenomic methods using predicted genes and a database containing, among others, 46 eukaryotic genomes including 10 phytoplankton and 2 additional protistan stramenopile genomes. Comparative gene analyses with *E. huxleyi* are prohibited and were not performed. Pico-prymnesiophyte predicted proteins were characterized by profiling and genome-size estimate methods. Chloroplast scaffolds were manually selected and annotated.

Microscopy was used with a prymnesiophyte-specific FISH probe (36) to count and size cells. For some cruises identification was by chloroplast arrangement, flagellar characteristics, and occasional presence of a haptonema; no significant difference (t test, P = 0.43) was detected for Atlantic data from 25° to 35° N (FISH:  $500 \pm 61 \text{ mL}^{-1}$ , n = 26; characteristics-based:  $593 \pm 108 \text{ mL}^{-1}$ , n = 12). Other picophytoplankton groups were enumerated by flow cytometry (37). Biovolume was calculated from cell size for pico-prymnesio-phytes and biomass determined using an established biovolume-based carbon conversion factor, also used for other picophytoplankton groups (38) (Table S7). Dilution experiments were according to (28, 30) and counts by FISH and flow cytometry. HPLC data were analyzed according to ref. 39.

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