EVOLUTIONARY TRAJECTORIES AND BIOGEOCHEMICAL IMPACTS OF MARINE EUKARYOTIC PHYTOPLANKTON

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Key Words coccolithophores, diatoms, dinoflagellates, phylogenetic trees, carbon cycle

Abstract The evolutionary succession of marine photoautotrophs began with the origin of photosynthesis in the Archean Eon, perhaps as early as 3.8 billion years ago. Since that time, Earth’s atmosphere, continents, and oceans have undergone substantial cyclic and secular physical, chemical, and biological changes that selected for different phytoplankton taxa. Early in the history of eukaryotic algae, between 1.6 and 1.2 billion years ago, an evolutionary schism gave rise to “green” (chlorophyll b–containing) and “red” (chlorophyll c–containing) plastid groups. Members of the “green” plastid line were important constituents of Neoproterozoic and Paleozoic oceans, and, ultimately, one green clade colonized land. By the mid-Mesozoic, the green line had become ecologically less important in the oceans. In its place, three groups of chlorophyll c–containing eukaryotes, the dinoflagellates, coccolithophorids, and diatoms, began evolutionary trajectories that have culminated in ecological dominance in the contemporary oceans. Breakup of the supercontinent Pangea, continental shelf flooding, and changes in ocean redox chemistry may all have contributed to this evolutionary transition. At the same time, the evolution of these modern eukaryotic taxa has influenced both the structure of marine food webs and global biogeochemical cycles.

INTRODUCTION

Phytoplankton comprise a diverse, polyphyletic group of single-celled and colonial aquatic photosynthetic organisms that drift with the currents (Falkowski & Raven 1997). Fewer than 25,000 morphologically defined forms are distributed
among at least eight major divisions or phyla. In contrast, nearly all photosynthetic organisms on land belong to a single clade (Embryophyta) that contains approximately 275,000 species (Table 1). Marine phytoplankton constitute less than 1% of Earth’s photosynthetic biomass, yet they are responsible for more than 45% of our planet’s annual net primary production (Field et al. 1998). Their evolutionary trajectories have shaped trophic dynamics and strongly influenced global biogeochemical cycles. In this paper, we examine the macroevolutionary histories of the major eukaryotic phytoplankton taxa that dominate the contemporary oceans, and consider their relationships to biogeochemical cycles.

ORIGINS OF PHOTOSYNTHESIS AND EUKARYOTIC PHYTOPLANKTON: MOLECULAR EVIDENCE

The origins of photoautotrophy are uncertain. Sedimentary microstructures as old as 3,500 Mega-annum (Ma) have been interpreted as cyanobacteria (Schopf 1993, Schopf 2002), but these reports have been questioned (Brasier et al. 2002). Carbon isotope measurements of reduced carbon in successions as old as 3,800 Ma seem to indicate widespread autotrophy in the oceans (Rosing 1999); however, the processes that led to the isotopic fractionation are not well understood. Perhaps more convincingly, fossil lipid biomarkers indicate that oxygenic photoautotrophs were present in the oceans by ca. 2,800 Ma (Brocks et al. 2003) and by ca. 2,300 Ma they had oxidized Earth’s atmosphere (Bekker et al. 2004). Oxygenic photosynthesis subsequently spread to eukaryotes via cyanobacterial endosymbiotic associations that evolved to keep the process localized in membrane-bound organelles called plastids (Yoon et al. 2004). A series of primary, secondary, and possibly even tertiary endosymbiotic associations spread photoautotrophy to five of the eight major extant eukaryotic clades, giving rise to multiple new eukaryotic phyla (Figure 1) (Baldauf 2003, Delwiche 1999, Palmer 2003, Yoon et al. 2004).

The earliest photosynthetic eukaryotes arose from one and possibly two successive endosymbiotic events in which the derived organelles were originally photosynthetic bacteria. One clade of early eukaryotes engulfed an ancestral cyanobacterium that was transformed into a plastid. More controversial is the proposal that an earlier, ancestral eukaryote incorporated the antecedent of an extant purple nonsulfur bacterium, which conferred an anoxygenic photosynthetic pathway on the host cell (Taylor 1987). The engulfed bacterium ultimately became the mitochondrion and lost its ability to photosynthesize, perhaps because of environmental oxidation. To accommodate the production of molecular oxygen within this new host-cell complex, the nascent mitochondrion would have operated its electron transport pathway in reverse (Osyczka et al. 2004), thereby giving rise to an oxygen-dependent respiratory electron transport chain with extremely high energy-conversion efficiencies. In this view, the evolution of the only two organelles that originated via endosymbiotic events were “motivated” by photosynthetic carbon-acquisition pathways.
This original symbiotic engulfment of a cyanobacterium into a host cell was a "primary" endosymbiotic process. The engulfed cyanobacterium contained a full complement of genes that allowed the organism to replicate itself. The symbiotic association was accompanied by the transfer of coding genes from the cyanobacterial genome into the eukaryotic nucleus. Assuming the genomic composition of extant cyanobacteria is representative of the ancestral symbiont genome, then 90% to 99% of the genes in the nascent plastid were subsequently lost or transferred to the host cell nucleus (Grzebyk et al. 2003, McFadden 1999, Palmer 2003). As a result, although they retain a core set of genes and limited autonomous capabilities for some housekeeping molecular processes, plastids cannot live longer than hours and cannot replicate outside of the host cell.

Three extant phytoplankton phyla (Chlorophyta, Rhodophyta, and Glauco-phyta) result from primary endosymbiosis. Various datasets of the eukaryotic host cell [18S ribosomal RNA gene (Figure 1), protein genes (Baldauf et al. 2000), or mitochondria (Gray et al. 1998)] indicate that these clades were derived from a common heterotrophic ancestor. Plastid gene phylogenies, such as those inferred from 16S ribosomal genes (Figure 2), also show the initial division of the plastid cluster according to the three types of primary plastids, typically surrounded by two membranes: cyanelles, green plastids (chloroplasts), and red plastids (rhodoplasts). Thus, despite their physiological differences, the three phyla may have originated from a single endosymbiotic event. Primary green plastids appear to have differentiated from the other two early in their evolutionary history. The accessory pigment, chlorophyll \( b \) (chl \( b \)), is the hallmark of the “green” plastid lineage, which became the forerunner of all chlorophyte algae and (subsequently) all higher plants that colonized terrestrial ecosystems in the Paleozoic. Cyanelles and rhodoplasts have many structural, biochemical, and genetic features in common; they contain chlorophyll \( a \) (chl \( a \)) only and have phycobilin pigments located in phycobilisomes, a simple carotenoid composition with zeaxanthin and \( \alpha \)-carotene or \( \beta \)-carotene similar to cyanobacteria, and plastid genomes that retain numerous genes lost in chloroplasts.

Secondary endosymbiosis is a process that involves two eukaryotic cells and appears to be unique to plastid evolution. A heterotrophic cell first engulfed an alga with primary plastids. This complex mix of organelles was subsequently "enslaved," and the core photosynthetic machinery was salvaged and reduced to a new plastid. Secondary endosymbiotic plastids are present in a number of algal phyla. Two of them are characterized by secondary green plastids (Euglenophyta and Chlorarachniophyta, Figure 1). Plastids in four other algal lineages—cryptophytes, haptophytes (including coccolithophores), dinoflagellates (division Dinophyta), and heterokonts (Figures 1 and 2)—originated from a symbiotic association that appears to have differentiated from rhodophytes. Heterokonts include multiple algal classes, such as diatoms (Bacillariophyceae), brown algae (Phaeophyceae), and classes with species that form harmful algal blooms (Raphidophyceae and Pelagophyceae).
TABLE 1  The higher systematic groups of oxygenic photoautotrophs, with estimates of the approximate number of total known species, their distributions between marine and freshwater habitats, and the type of plastid and major accessory pigments they contain

<table>
<thead>
<tr>
<th>Taxonomic group</th>
<th>Known species</th>
<th>Plastids</th>
</tr>
</thead>
<tbody>
<tr>
<td>Category</td>
<td>Total</td>
<td>Marine</td>
</tr>
<tr>
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<td></td>
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<tr>
<td>Kingdom</td>
<td></td>
<td></td>
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<tr>
<td>Eubacteria</td>
<td></td>
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<tr>
<td>Subdivision</td>
<td>Cyanobacteria (stricto sensu)</td>
<td>1,500</td>
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<tr>
<td>(=Cyanophytes, blue-green algae)</td>
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<td></td>
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<tr>
<td>Subdivision</td>
<td>Chloroxybacteria (= Prochlorophyta)</td>
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</tr>
<tr>
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<tr>
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<tr>
<td>Division</td>
<td>Dinophyta (Dinoflagellates)</td>
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<tr>
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<td>Dinophyceae</td>
<td></td>
</tr>
<tr>
<td>Kingdom</td>
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<tr>
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<td>Prasinophyceae</td>
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<td>Ulvophyceae</td>
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<td></td>
<td>Charophyceae</td>
<td>12,500</td>
</tr>
<tr>
<td>Division</td>
<td>Date</td>
<td>Number of species</td>
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<tr>
<td>--------------------------</td>
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</tr>
<tr>
<td>Bryophyta (mosses, liverworts)</td>
<td>22,000—1,000</td>
<td>Lycopsida (ferns)</td>
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<tr>
<td>Magnoliophyta (flowering plants)</td>
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<td>Monocotyledoneae</td>
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<td>Subkingdom Euchromista</td>
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<td>Division Cryptophyta</td>
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</tr>
<tr>
<td>Class Cryptophyceae</td>
<td>200—100—100</td>
<td>Division Haptophyta</td>
</tr>
<tr>
<td>Class Heterokontophyta</td>
<td></td>
<td>S (red)</td>
</tr>
</tbody>
</table>

Kingdom Fungi

Division Ascomycotina (lichens) | 13,000 | 15 | 20

\*The difference between the number of marine and freshwater species, and that of known species, is accounted for by terrestrial organisms. Dashes indicate that no species are known (by us) for their particular group in this environment. Abbreviations are as follows. Plastid type: \(P=\) primary plastid (cyanelle, red, or green); \(S=\) secondary plastid (from the green or red plastid lineage). Accessory pigments: \(B=\) bilipigments; \(b=\) chlorophyll \(b; c_1/c_2/c_3=\) chlorophyll \(c\) type 1, 2 or 3; \(db=\) divinyl chlorophyll \(b.\) Carotenoids: \(\alpha=\) \(\alpha\)-carotene; \(\beta=\) \(\beta\)-carotene; \(A=\) antheraxanthin; \(Aljalloxanthin; BF=19'\)-butanoyloxyfucoxanthin; \(Dd=\) diadinoxanthin; \(Di=\) dinoxanthin; \(Dt=\) diatoxanthin; \(Fu=\) fucoxanthin; \(HF=19'\)-hexanoyloxyfucoxanthin; \(L=\) lutein; \(N=\) neoxanthin; \(P=\) prasinoxanthin; \(Pe=\) peridinin; \(V=\) violaxanthin; \(Va=\) vaucheraxanthin; \(Z=\) zeaxanthin. One-digit abbreviations indicate pigments already present in cyanobacteria or primary plastids; two-digit abbreviations indicate pigments appearing in secondary plastids.

There is ongoing debate over the number of endosymbiotic events in the evolutionary history of eukaryotic phytoplankton. Early molecular phylogenetic analyses suggested that the two secondary green plastid clades and four secondary red phyla were derived from independent endosymbiosis events. The nuclear and mitochondrial genomes of these two secondary green phyla are not closely related (Figure 1) (Baldauf 2003, Gray et al. 1998). Similarly, nuclear and plastid gene phylogenetic analyses indicate that the cryptophytes, haptophytes, dinoflagellates, and heterokonts arose from independent host cells. Plastid gene phylogenies indicate multiple origins of secondary plastids within rhodophytes (see Figure 2) (Bhattacharya & Medlin 1998, Müller et al. 2001, Oliveira & Bhattacharya 2000, Yoon et al. 2002a), and eukaryotic gene sequence analysis indicates that secondary host cells were unrelated or, at best, that heterokonts and alveolates share an ancient common ancestor (Figure 1) (Baldauf 2003, Baldauf et al. 2000, Bhattacharya & Medlin 1998). However, it has been proposed that secondary endosymbiosis occurred only twice: once in the green algal lineage and once in the red algal lineage (Cavalier-Smith 2003). A single endosymbiotic origin of plastids in the “green” euglenids and chlorarachniophytes seems unlikely, but some phylogenetic studies support the single-event hypothesis for phyla that contain secondary red plastids (Fast et al. 2001, Harper & Keeling 2003, Yoon et al. 2002b). This would indicate that all extant basal heterokonts and alveolates must have then lost their plastids to return to the heterotrophic nutritional mode.

Endosymbiotic events transformed both the eukaryotic host and symbiotic cells. The most noticeable impact concerned the composition of accessory photosynthetic pigments (Table 1). Most extant marine cyanobacteria contain only chl $a$, open-chain tetrapyrroles (phycobilins), and two carotenoids (zeaxanthin and $\beta$-carotene). However, the cyanobacterial division of Prochlorophyta also contains chl $b$ or a divinyl derivative as a major accessory pigment. Within the Eucarya, chl $b$ is present only in green plastids (hereafter, the “green” line), whereas phycobilins are present only in red plastids. For that reason, the ancestor of plastids was suggested to have been a cyanobacterium that contained both chl $b$ and phycobilins (Tomitani et al. 1999). However, that hypothesis appears to be unlikely because extant prochlorophytes are genetically very distant from plastids (Hess et al. 2001). After primary endosymbiosis, rhodophytes retained the cyanobacterial pigments (but with $\alpha$-carotene instead of $\beta$-carotene), and glaucophytes acquired an additional minor carotenoid. In contrast, primary green plastids greatly diversified in carotenoid pigments; they added four major pigments that absorb light in the wavelength range 400 to 520 nm (versus the less energetic wavelength range of 480 to 580 nm absorbed by rhodophytes). Whereas one may ask if the “primitive” pigment composition of rhodophytes limited the evolution of this phylum, interestingly, accessory pigments diversified in the red plastid lineage after secondary endosymbiosis. The new set of pigments, although specific to secondary red plastids, conferred similar properties to new plastids that mimic those provided by accessory pigments of primary green plastids. Chlorophyllide $c$ and its derivatives, which are not present in any extant photosynthetic prokaryotes or in rhodophytes,
absorb light in the 450 to 480 nm range, which nearly overlaps the absorption range by chl \( b \). Secondary red plastid carotenoids, some of which are unique to a phytoplankton phylum, absorb light from 400 nm up to 580 nm. We refer to the secondary red plastid phyla that contain chl \( c \) as the “red plastid lineage.”

**EARLY PHYTOPLANKTON EVOLUTION:**

**FOSSIL EVIDENCE**

Fossils of probable eukaryotic origin first appear in abundance in rocks dated as approximately 1,700 to 1,900 Ma (Han & Runnegar 1992, Javaux et al. 2001, Knoll 1994). This age should be regarded as a minimum date for eukaryotic diversification because the paleontological record of older strata is sparse; molecular biomarkers suggest that both cyanobacteria and at least stem group eukaryotes existed by 2,700 Ma (Brocks et al. 1999, Summons et al. 1999), and the former, at least, may have evolved earlier (Knoll 2003). The systematic affinities of the earliest eukaryotic fossils are poorly constrained, but by 1,200 Ma, multicellular red algae lived along tidal flats, attached to hardgrounds (Butterfield 2000).

The presence of rhodophytes in mid-Proterozoic rocks places a minimum constraint on the timing of the red-green plastid schism. Reds radiated early as multicellular, benthic constituents of nearshore communities (Xiao et al. 2004). In contrast, early branching greens are planktonic, the paraphyletic “prasinophytes” that still contribute to photosynthesis in open ocean environments (Melkonian & Surek 1995). Xanthophyte fossils in uppermost Mesoproterozoic and Neoproterozoic rocks (Butterfield 2002, German 1990) indicate that secondary endosymbiosis followed relatively closely on the heels of primary events.

The functional as well as phylogenetic schism of reds and greens suggests that early members of the two lineages became adapted for different environmental regimes; the greens proliferated in oligotrophic open oceans underlain by anoxic waters, whereas the reds differentiated in coastal waters where both nutrients and fully oxygenated waters were more abundant (Anbar & Knoll 2002). Cyanobacteria appear to have dominated primary production in early to middle Proterozoic oceans, but by the end of the era (543 Ma), eukaryotic algae had become important constituents of marine ecosystems (Knoll 1989, Knoll 1992, Lipps 1993, Tappan 1980).

Many fossils of Proterozoic (and younger) protists cannot be assigned to extant taxa with any degree of certainty. Closed, organic-walled structures that resemble (at least broadly) dinocysts or prasinophyte phycomata are called “acritarchs,” a grouping that makes them easier to deal with, but not to understand. Acritarchs first appeared as a minor component of the fossil record approximately 1700 to 1900 Ma (Summons et al. 1992, Zhang 1986); a moderate increase in diversity occurred approximately 800 to 900 Ma (Knoll 1994). This diversification coincides with the early stages of Rodinia rifting; however, no firm causal link has been established between the two events (Figure 3).
In the Cambrian and Ordovician, eukaryotic phytoplankton underwent a second diversification in concert with the early Paleozoic radiations of marine invertebrates. The affinities of Early Paleozoic phytoplankton remain a subject for debate (Molyneux et al. 1996). Molecular biomarkers indicative of dinoflagellates occur in association with some Cambrian microfossils (Moldowan & Talyzina 1998), but morphological and ultrastructural analyses also show that prasinophyte green algae played a larger role in the continental shelf phytoplankton than they have in more recent times (Colbath & Grenfell 1995, Talyzina & Moczydlowska 2000). Preserved phytoplankton diversity peaked in the mid-Paleozoic and declined rapidly in the Late Devonian to Early Mississippian.

THE RISE OF THE RED LINEAGE

Mesozoic Expansion of the Red Lineage

In the early Mesozoic, the red eukaryotic phytoplankton began to assume a new ecological importance in the marine realm (e.g., Falkowski et al. 2004a,b). Marine prasinophytes declined during the Jurassic and Cretaceous, although this group of green algae is still represented in the oceans today (albeit as a minor constituent of open-ocean communities). A period of transition to red-line-dominated primary production occurred in the Triassic to Early Jurassic, ultimately resulting in the ecological dominance of coccolithophores, dinoflagellates, and diatoms in the contemporary ocean.

The first unequivocal dinoflagellates appeared in the fossil record as organic-walled cysts preserved in Middle Triassic continental margin sediments (Stover et al. 1996). Studies of molecular biomarkers indicate that dinoflagellates may have existed as far back as the Neoproterozoic (Moldowan & Talyzina 1998, Summons & Walter 1990); however, these biomarkers did not become prominent constituents of marine bitumens until the Triassic (Moldowan et al. 1996, Moldowan & Jacobson 2000), when microfossils more clearly document their expansion and radiation (Fensome et al. 1996, Stover et al. 1996) (Figure 3).

The calcareous nannoplankton (dominated by coccolithophorids) were the second group in the red lineage to radiate in the fossil record. They originated in the Late Triassic (Bown et al. 2004) (Figure 3) at about the same time that molecular biomarkers of coccolithophorids became common (Moldowan & Jacobson 2000). The earliest nannoplankton have been identified in Carnian sediments from the southern Alps (Bown 1998, Janofske 1992) and Nevada (F. Tremolada, personal communication).

The silica-encased diatoms were the last of the three major groups to emerge in the Mesozoic. The siliceous diatom frustules are highly soluble compared with other siliceous fossils (e.g., radiolaria and sponge spicules), which possibly imparts a preservational bias to the fossil record of diatom origin. Reports of diatom frustules in Jurassic sediments (Rothpletz 1896) have proved difficult to replicate by later workers, although molecular biological clock estimates (Medlin et al. 2000)
and molecular biomarkers (Moldowan & Jacobson 2000) indicate that diatoms may have evolved earlier, but remained minor components in the marine realm until the Cretaceous. The first unequivocal fossil records of diatoms document radiations in the Early Cretaceous oceans (Harwood & Nikolaev 1995). Diatom morphologies in the Early Cretaceous were dominated by cylindrical and long-cylindrical forms that show very little variation (Gersonde & Harwood 1990, Harwood & Gersonde 1990); the similarity among these early diatom morphologies argue against a pre-Mesozoic origin (Harwood & Nikolaev 1995). Late Cretaceous diatoms were dominated by discoidal and biddulphioid frustule morphologies (Harwood & Nikolaev 1995). Diatoms were present in nonmarine environments by 70 Ma (Chacon-Baca et al. 2002).

Regardless of the exact timing of the evolutionary origins of coccolithophores, dinoflagellates, and diatoms, fossil and biomarker data document the major expansion of all three groups in the Mesozoic (Bown et al. 2004, Grantham & Wakefield 1988, Harwood & Nikolaev 1995, Moldowan & Jacobson 2000, Stover et al. 1996) (Figure 3). They began their evolutionary trajectories to ecological prominence as the supercontinent Pangea began to break apart in the Late Triassic to Early Jurassic (∼200 Ma), which marked the opening phase of the current Wilson cycle of continental breakup, dispersal, and reassembly (see The Role of the Wilson Cycle, below). Sea level rose as Pangea fragmented and the Atlantic Ocean basin widened, flooding continental shelves and low-lying inland areas (Figure 3). In addition, the fragmentation of the continents and creation of a new ocean basin produced an increase in the total length of coastline where many plankton lived. Nutrients (such as phosphate) that were previously locked up in the large continental interior of Pangea were transported to newly formed shallow seas. At the same time, diversities increased in the three groups of eukaryotic phytoplankton, and in marine invertebrates, paralleling a long-term increase in sea level that began in the Early Jurassic (Haq et al. 1987, Vail et al. 1977) (Figure 3). Greater nutrient availability coupled with expanded habitat area may well have contributed to the radiation of phytoplankton that lived along continental margins. Accordingly, the diversities of eukaryotic phytoplankton of the red lineage parallel sea-level rise through the Mesozoic (Figure 3). In contrast, short-term [1 million year (myr) scale] sea-level changes may have resulted in species turnover events, but had very limited impact on the large-scale evolutionary history of phytoplankton clades.

Life-cycle strategies may also have been an important component that favored the radiation of the red lineage as continental shelves flooded and epicontinental seas became widespread in the Jurassic. Dinoflagellates, coccolithophorids, and diatoms produce resting stages. After a bloom, a small fraction of the phytoplankton population becomes arrested in a specific stage of the cell’s life cycle in which the production of cell armor increases, and the cell sinks to the seafloor until conditions become favorable to bloom (usually the following year). The timing of the bloom depends on environmental conditions such as ocean stratification and day length. The resting stage is associated with gamete formation and gene exchange in the planktonic portion of the life cycle. This life-cycle strategy requires shallow
marine areas and also promotes genetic isolation by reducing the gene flow. Gene transfer, which can occur through sexual recombination, lateral vectors (such as viral infection), or both, is highly attenuated in benthic stages. Over relatively short periods of time, genetic isolation may have increased the tempo of evolution and phenotypic selection, processes that have been observed in contemporary phytoplankton assemblages (Medlin et al. 1997).

Cenozoic Expansion of Diatoms

Bolide impact at the Cretaceous/Tertiary boundary (65 Ma) caused major extinctions (Alvarez et al. 1980) that are recorded in the fossil records of the coccolithophores and, to a lesser extent, the diatoms and dinoflagellates (Figure 3). The environmental factors that favored the expansion of the red lineage through the Mesozoic allowed these phytoplankton to repopulate the marine realm after the Cretaceous/Tertiary boundary (K/T) mass extinction event. Dinoflagellates and calcareous nannoplankton recovered to preextinction diversity levels by the earliest Eocene (~55 Ma), only to decline through the rest of the Cenozoic as long-term sea level began to fall in the mid-Paleogene and the extent of flooded continental areas decreased. Among fossilizable taxa, modern dinoflagellate species diversity has declined to levels comparable to the earliest Middle Jurassic, whereas modern coccolithophorid species diversity has declined to Late Jurassic levels.

In contrast to the other phytoplankton, diatom diversity has increased through the Cenozoic, despite falling sea level. Two pulses of diversification occurred in diatoms: the Eocene/Oligocene boundary interval and the middle to late Miocene. The latter was accompanied by a substantial radiation among pennate species (Strelnikova 1991). Conditions that may have favored the expansion of diatoms in the Cenozoic are discussed in the following section.

SELECTION PRESSURES AND ADAPTATIONS

Resource Acquisition

In contrast to most species of dinoflagellates and coccolithophores, diatoms frequently form extensive blooms along continental margins and in upwelling regions of the contemporary ocean. These organisms are responsible for approximately 40% of the net primary production and more than 50% of the organic carbon that is exported to the ocean interior (Falkowski et al. 2003). Planktonic diatoms have evolved a nutrient storage vacuole that retains high concentrations of nitrate and phosphate (Raven 1997). The storage vacuole allows diatoms to acquire pulses of inorganic nutrients, which can deprive competing taxa of these essential resources, overcome light-dependent nutrient uptake in mixing systems, or both. The storage capacity of the vacuole is sufficient to allow two to three cell divisions without the need for external nutrient resources. Consequently, diatoms thrive best in regions where nutrients are supplied with high pulse frequencies.
Models based on resource acquisition strategies (e.g., Grover 1988; Litchman E, Klausmeier CA, Miller JR, Schofield OM, Falkowski PG, in review; Tilman 1977; Tozzi et al. 2004) suggest that diatoms dominate when brief periods of water-column stability are punctuated by high turbulence, whereas coccolithophores and dinoflagellates dominate when the water column is more stably stratified. After Tozzi and others (2004).

Competition among diatoms, coccolithophores, and dinoflagellates has been modeled by application of resource acquisition strategies (e.g., Grover 1991; Litchman E, Klausmeier CA, Miller JR, Schofield OM, Falkowski PG, in review; Tilman 1977; Tozzi et al. 2004). The results of these models suggest that diatoms dominate when brief periods of water-column stability are punctuated by high turbulence, such as storm events (Figure 4). In contrast, coccolithophores and dinoflagellates dominate when the water column is more stably stratified. In theory, competitive exclusion could occur if equilibrium conditions were reached. In reality, the coexistence of two or more taxa competing for a single resource is a consequence of the dynamically unstable nature of aquatic ecosystems (Li 2002, Siegel 1998). Margalef (1994) recognized these fundamental differences in physiology and proposed that competition among the three major red lineage taxa could be related to upper-ocean turbulence and the supply of nutrients. The so-called Margalef mandala can be extended over geological time (Prauss 2000) to infer selection processes that led to the early rise of dinoflagellates and coccolithophorids in the relatively stable conditions of the Mesozoic, followed by the rise of diatoms in the Cenozoic.
Increasing latitudinal thermal gradients and decreasing deep-ocean temperatures have contributed to greater vertical thermal stratification through the latter half of the Cenozoic, which increased the importance of wind-driven upwelling and mesoscale eddy turbulence in providing nutrients to the upper ocean. The ecological dominance of diatoms under certain sporadic mixing conditions suggests that their long-term success in the Cenozoic probably can be attributed in part to an increase in event-scale turbulent energy dissipation in the upper ocean (Falkowski et al. 2004a). Sporadic nutrient influx to the euphotic zone may favor diatoms over coccolithophores and dinoflagellates, and a change in concentration and pulsing may favor small diatoms over large diatoms. As a result, a significant decrease in the average size of diatoms occurred in the Cenozoic. Periods of change were concentrated in the middle to late Eocene and early to middle Miocene (ZV Finkel, ME Katz, JD Wright, OM Schofield & PG Falkowski, in review).

Silica Bioavailability

Diatoms precipitate orthosilicic acid in a protein matrix to form extremely strong, structurally intricate biogenic opal shells called frustules. The modern oceans are undersaturated with respect to silica, which is largely the result of the evolution and ecological success of diatoms in removing this element from the dissolved phase, especially in the latter half of the Cenozoic (Conley 2002). Eighty percent of the silica in the oceans is derived from chemical and biological weathering of continental rocks (De La Rocha et al. 2000); the annual net riverine flux is 5.0 Tmol of silica per year (Tréguer et al. 1995). Sustaining the silica flux on geological timescales requires that fresh rock surfaces continuously become exposed to the soil interface, a process that is perpetuated by erosion. The rate of nutrient flux to the oceans on multimillion-year timescales is determined in part by continental elevation. Both orogeny and regression have characterized nearly all of the continents during the Cenozoic, which has increased nutrient fluxes (including silicic acid) (Maldonado et al. 1999) and facilitated diatom expansion (Falkowski et al. 2004a,b). An additional loop in the silica cycle that developed in the Cenozoic also may have accelerated the success of diatoms. The loop involves the coevolution of mammals, grasses, and diatoms. Grasses extract silicic acid from groundwater in soils and store it in opal phytoliths (Conley 2002). Silica can constitute as much as 15% of the dry weight of grasses (Alexandre et al. 1997, Bartoli 1983, Rapp & Mulholland 1992), and phytolith dissolution can release soluble silica two times more efficiently than silicate weathering (Alexandre et al. 1997).

Grasses originated in the Cretaceous, but remained sparse until the Eocene/Oligocene boundary (33.7 Ma) (Kellogg 2000, Retallack 2001), when global climates became more arid as a result of major glaciation in the Antarctic. As grasslands expanded, grazing ungulates evolved and displaced browsers (Janis & Damuth 2000). Hypsodont (high-crown) dentition in ungulates was selected over the brachydont (leaf-eating) dentition in browsing mammals, which coincided with the widespread distribution of silica-rich phytoliths and grit in grassland
forage (Retallack 2001). The rise of grazing ungulates and the radiation of grasses may have acted as a biologically catalyzed silicate weathering process (Falkowski et al. 2004a,b). Phytolith diversity and abundance has increased since the late Eocene (Jacobs et al. 1999, Retallack 2001); almost certainly, grass-mediated silica mobilization from soils increased as the grasses radiated. The subsequent transfer of silica to the oceans (primarily via riverine transport) increased the bioavailability of silica for diatom growth (Falkowski et al. 2004a,b). Accordingly, diatom species diversity, and presumably abundance, increased dramatically at the Eocene/Oligocene boundary (Figure 3). A major expansion of grasslands in the Neogene (Retallack 1997, Retallack 2001) was accompanied by a second pulse of diatom diversification at the species level (Falkowski et al. 2004a,b).

Armor

We can infer the evolutionary trajectories of all three major red lineage clades that appear to have dominated the eukaryotic phytoplankton community since the Mesozoic because they have fossilizable cell walls that also provide armor. This armor likely protected the phytoplankton from grazers long enough for the phytoplankton to form blooms (Banse 1992). Although the composition of the cell walls differs markedly among dinoflagellates, coccolithophorids, and diatoms, grazing by zooplankton may have provided a common evolutionary selection pressure. Unfortunately, virtually no fossil record exists for major modern pelagic zooplankton groups; chitinous crustaceans (e.g., copepods and euphausids) decompose rapidly in the sediments, and virtually all soft-bodied organisms such as salps decompose in the water column before they reach the sea floor. The potential role of cell walls as armor against grazing has long been debated (Smetacek 1999, 2001). The strength and flexibility of diatom frustules makes fracturing them a challenge for invertebrates; in fact, diatoms can pass through a copepod gut intact (Hamm et al. 2003). The chitinous teeth of copepods have a siliceous coating that provides significant compressive strength, and mandible muscles in these crustaceans are highly developed. This feature is a clear example of adaptive evolution between predator and prey in pelagic ecosystems (Adams 2001). However, if diatoms are consumed, zooplankton greatly facilitate the dissolution of the frustule (Bidle & Azam 1999).

Coccolithophores are grazed, but the nutritional benefit derived from ingesting a cell consisting of 30% calcium carbonate is lower than that derived from ingesting a naked cell of the same size. Indeed, zooplankton avoid coccolithophorids if presented with optional, unarmored food sources (Falkowski and Wyman, unpublished data). In addition to armor, thecate dinoflagellates (and some diatoms) have evolved a strategy of vertical migration; they obtain nutrients below the pycnocline at night and rise to the upper portion of the euphotic zone during the day to optimize photosynthesis (Kamykowski 1981, Villareal et al. 1993). This migration strategy is out of phase with that of many zooplankton grazers (Banse 1964).
Ocean Chemistry

Secular changes in seawater chemistry also appear to have influenced phytoplankton evolutionary trajectories. For example, trace elements (including iron, copper, zinc, and manganese) play essential roles in mediating critical biochemical reactions in all phytoplankton. The bioavailability of these elements in seawater is strongly dependent on redox state (Whitfield 2001). Algae with green plastids have substantially higher quotas for iron, zinc, and copper (i.e., Fe:P, Zn:P, and Cu:P ratios) than do red eukaryotes, whereas the latter have higher quotas for cadmium, cobalt, and manganese (Quigg et al. 2003). Furthermore, the redox state of the oceans may discriminate between red and green plastid lineages with respect to fixed nitrogen preferences (Anbar & Knoll 2002; Falkowski & Raven 1997; Litchman E, Klausmeier CA, Miller JR, Schofield OM, Falkowski PG, in review), as well as the availability of phosphate. Members of the green plastid lineage tend to have higher N/P ratios than members of the red plastid lineage (Falkowski et al. 2004b). Hence, the form and availability of macronutrients and trace metals have the potential to be strong selective agents that favor red-or green-plastid-containing taxa.

The Archean biosphere was anoxic, and its initial oxidation in the Paleoproterozoic Era introduced oxygen primarily into the surface mixed layer of the oceans. Complete oxygen depletion appears to have remained common beneath the pycnocline, and sulfidic deep waters were widespread (Canfield 1998) (Figure 3). Under these conditions, early algae likely competed best in the better-oxygenated coastal regions, where rivers delivered essential metals (Anbar & Knoll 2002). As noted above, the evolutionary divergence of red and green plastids took place in the context of this redox heterogeneity and may have contributed to the different evolutionary trajectories of rhodophytes and early chlorophytes. Increasing oxidation of ocean waters may have facilitated the expansion of algae across shelves in latest Neoproterozoic times; undoubtedly, interactions with evolving animals also contributed to Cambrian and Ordovician phytoplankton diversification (Butterfield 1997).

Black shales are common in Proterozoic marine successions, which indicates oxygen depletion beneath surface water masses in the oxygen-minimum zone (Shen et al. 2002, Shen et al. 2003); black shales also occur episodically through much of the Paleozoic (Arthur & Sageman 1994). Deep-water anoxia may have been particularly pronounced near the end of the Permian (Isozaki 1997). The expanded oxygen-minimum zone persisted through the Early Triassic (Twitchett 1999, Wignall & Twitchett 2002), and may have altered the distributions of many trace elements within the oceans (Whitfield 2001a). Ocean anoxia increases the availability of Fe, Mn, P, and ammonium and decreases the availability of Cd, Cu, Mo, Zn, and nitrate; hence, the green lineage may have been favored over the red lineage at times when subsurface reducing conditions prevailed.

A secular shift in ocean redox conditions in the early Mesozoic changed trace metal availability in the oceans and exerted a selective pressure that favored the red lineage by better meeting their metal requirements (Falkowski et al. 2004a,b).
The last major occurrence of large concentrations of the green algal prasinophytes was in the Early Jurassic, when black shale deposition and ocean anoxia appears to have been widespread (Falkowski et al. 2004a,b). As deep-ocean oxygenation became increasingly permanent through the Mesozoic, Cd, Cu, Mo, Zn, and nitrate availability increased, which allowed the red lineage to expand (Anbar & Knoll 2002, Falkowski et al. 2004a,b). Temporal changes in the availability of redox-sensitive trace metals are consistent with the biological transition to red-lineage dominance of phytoplankton (Falkowski et al. 2004a,b). Thus, long-term changes in ventilation of the world’s oceans ultimately appear to have played a significant role in the rise of the red lineage during the Mesozoic.

The Mesozoic oceans occasionally were punctuated by widespread organic carbon burial events associated with the short-lived (< 1 myr) Oceanic Anoxic Events (OAEs) (e.g., Arthur & Sageman 1994). These events briefly altered ocean redox conditions, but the effects were relatively short-lived. The biological impact of OAEs is debated (e.g., Leckie et al. 2002 versus Bown et al. 2004) despite the substantial effort dedicated to examining the phytoplankton and zooplankton communities across these carbon-burial events (e.g., Erbacher & Thurow 1997, Leckie et al. 2002, Roth 1987). Once the red lineage garnered a more secure ecological advantage in the Late Triassic, the fossil records suggest that the OAEs only had a minor influence on the evolutionary trajectories of eukaryotic phytoplankton (Bown et al. 2004, Falkowski et al. 2004a).

THE ROLE OF THE WILSON CYCLE

The radioactive decay of elements within Earth’s core produces heat that dissipates to the planet’s surface through convection and conduction. Near the surface, the thin oceanic crust conducts heat about three times more efficiently than the thicker continental crust. The differential dissipation of radiogenic heat oceanic basalts versus continental crust leads to the buildup of thermal energy below a fully assembled supercontinent, which causes the continental crust to thin and eventually to fracture and rift apart. A new ocean spreading center and ocean basin form between the continental fragments. Heat-driven convection in the mantle below the lithosphere drives the tectonic plates apart with the fragmented continents attached, and new oceanic crust forms at the intervening spreading center. As it ages, the oceanic crust becomes cooler and denser and eventually subsides as it moves away from the spreading center. When this old crust becomes so dense that it subsides below the adjacent, relatively low-density continental crust, a new subduction zone is created and the ocean basin is consumed as the whole process reverses itself and the continents reassemble. Named after its conceptual discoverer, J. Tuzo Wilson (Wilson 1966), this episodic breakup, dispersal, and subsequent reassembly of supercontinents has become known as the Wilson Cycle, and it occurs over approximately 300-myrr to 500-myrr intervals (e.g., Fischer 1984, Rich et al. 1986, Valentine & Moores 1974, Worsley et al. 1986).
The Wilson Cycle and Evolutionary Trajectories

Several studies have drawn attention to the correlation between evolutionary pulses in the marine realm and the Wilson Cycle (e.g., Damsté et al. 2004, Fischer 1984, Rich et al. 1986, Valentine & Moores 1974, Worsley et al. 1986). Myriad studies since these early publications provide the foundation for speculating on causal rather than casual linkages for the phytoplankton response to the Wilson Cycle.

Global species diversity reflects both the packing of taxa within communities and the distribution of suitable habitat area (MacArthur & Wilson 1967, Rosenweig 1995). A geological proxy for the former is provided by estimates of within-assemblage diversity through time (e.g., Bambach 1977), whereas paleobiogeographic analyses gauge the contribution of the latter. Diversity increases correlate with continental rifting during early iterations of the Wilson Cycle. Acritarchs appear to have radiated as Rodinia rifted in the Late Proterozoic, and diversified again in the Early Paleozoic as Pannotia rifted (Figure 3). Thus, continental separation may have played a role, along with the ecological drivers that govern within-assemblage diversity, in promoting Cambro-Ordovician phytoplankton (and marine invertebrate [Sepkowski 1997; Bambach 1999]) expansion.

Sea-level rise and flooded continental area also are highly correlated with increasing diversity of Mesozoic calcareous nannoplankton (e.g., coccolithophorids) and dinoflagellates (Figure 5), as well as with their declining diversities in the Cenozoic oceans. Although flooded continental area is a small percentage of the total oceanic area suitable for phytoplankton, the shallow seas appear to have contributed proportionally more to niche space because of high nutrient input, high rates of primary production, and habitat heterogeneity. Flooded continental area provides variable, high-nutrient habitat by creating additional upwelling zones and increasing turbulence and nutrient suspension from below the thermocline. In addition, terrestrial nutrient input likely increases because nutrients that were previously sequestered in the large supercontinent interior are more readily transported to the newly opened, nearby oceans. We note that flooded continents increase the availability of fossiliferous sediments and have the potential to impart a taphonomic and sampling component to diversity compilations. This bias may be most pronounced in the older record because most pre-Jurassic ocean crust has been destroyed.

A simple equation captures the empirical relationship between long-term sea level change and diversities in calcareous nannoplankton, dinoflagellates, and diatoms (Figure 5); this equation provides correlation coefficients, but is not a predictor of sea level based on phytoplankton diversities. Although accurate reconstruction of diversities is inevitably biased by differential preservation and the problem of defining species based on morphologic characters, our approach provides an estimate of the relationship between phytoplankton richness (R) and sea level (meters) that solves for coefficients that link the diversity of phytoplankton taxa to change in habitat area caused by sea level change. This relationship is defined as:
Figure 5 Phytoplankton diversity (sources as in Figure 3) as a function of flooded continental area in the Paleozoic (acritarch genera) and sea level in the Mesozoic (calcareous nannoplankton species + dinoflagellates species). The radiation in diatom species diversity alters the relationship between phytoplankton diversity (calcareous nannoplankton species + dinoflagellate species + diatom species) and sea level change in the Cenozoic. Sea level was translated to obtain positive values. Timescales are as in Figure 3.

\[-0.29 \times D_{\text{diatoms}} + 0.42 \times D_{\text{nannos}} + 0.59 \times D_{\text{dinos}} + 25.4 = \text{sea level} \quad (r^2 = 0.82) \quad (1)\]

Our analysis suggests that whereas calcareous nannoplankton and dinoflagellate diversities are highly correlated to sea level, diatom diversity responds to other environmental factors (see Selection Pressures and Adaptations, above) and has rapidly increased over the past approximately 35 myr.

The tectonic processes that drive the Wilson Cycle not only affect available niche space, but also affect ocean chemistry. As seawater cycles through mid-ocean ridges, magnesium is removed and calcium is added. Total ridge length
and seafloor spreading rates can change Mg/Ca ratios in seawater. As a result, high-Mg/Ca ratios tend to occur during times of supercontinent assembly, when ridge length is shortest, characterized by deposition of aragonite and high-Mg calcite (called “aragonite seas”). Low-Mg calcite deposition tends to characterize times of continental breakup (called “calcite seas”) (Hardie 1996, Sandberg 1975). Low-Mg/Ca and high-Ca\(^{++}\) concentration in seawater favors calcification in certain groups of marine organisms (including coccolithophores), which results in a correspondence between these organisms and “calcite sea” intervals (Stanley & Hardie 1998). This relationship is illustrated by the expansion of coccolithophores in the Mesozoic calcite seas (Figure 3), along with other low-Mg calcifiers among marine invertebrates. The massive Cretaceous coccolith chalks (e.g., the White Cliffs of Dover) were deposited on continental shelves when margins were flooded, Mg/Ca ratios were low, and dissolved Ca\(^{++}\) concentrations in seawater were high. This correlation further links the expansion of coccolithophores to the opening phase of the current Wilson Cycle. Chalk deposition continued in the Paleocene after the K/T extinctions, but as the Mg/Ca ratio increased through the Cenozoic, the degree of calcification in coccoliths declined, and coccolith size decreased (Bukry 1971, Houghton 1991).

**Bottom-Up Control of Marine Invertebrate Fauna**

In his classic essay “Seafood Through Time,” Bambach (1993) identified a series of changes in late Mesozoic and Cenozoic marine faunas—increases in abundance and mechanical strength of top predators, mean size of marine invertebrates, and mean rates of energy consumption. These changes all require a greater nutritional supply to marine life consistent with observed increases in C to P ratios (Martin 2003). Bambach (1993, 1999) suggested that the necessary nutrients were supplied by angiosperm-facilitated increases in erosional runoff from continents, runoff that would have been further augmented by higher erosion rates associated with increasing continental elevations (see Silica Bioavailability, above). As an additional hypothesis, we propose that the Wilson Cycle and the evolutionary shifts in phytoplankton community composition, in conjunction with increases in primary production and the quantity and quality of export production in the Phanerozoic seas, strongly influenced marine invertebrate faunal evolution.

Marine invertebrate faunal diversity (Bambach 1993) is highly correlated with phytoplankton diversity and community composition (Figure 6), but poorly correlated with flooded continental area \((r^2 = 0.14)\) and sea level \((r^2 = 0.05)\). An increase in net primary production increases both the availability of organic matter that can be transferred to higher trophic levels in marine invertebrates and the potential to export food to the seafloor (Falkowski et al. 2003). The fossil record indicates that an evolutionary change in phytoplankton diversity and community composition is a predictor of change in the richness in marine invertebrate genera: (a) diatom genera richness is highly correlated with Cenozoic invertebrate richness \((r^2 = 0.75)\); (b) the combined diversity of calcareous nanoplankton and dinoflagellates is highly correlated with Mesozoic invertebrate diversity
Figure 6  Marine invertebrate diversity (Bambach 1993) and phytoplankton diversity (as in Figure 3) over the Phanerozoic. Timescales are as in Figure 3. Data are available at http://mychronos.chronos.org/~miriamkatz/20040721

There is a strong correlation between marine invertebrate diversity and phytoplankton diversity over the Phanerozoic (r² = 0.89); and (c) a weak correlation exists between the diversity of Paleozoic marine invertebrates and the diversity of the acritarchs (r² = 0.12) (Figure 6).

The radiation of diatoms in the Cenozoic demarcates a large change in food-web structure of the Phanerozoic oceans, which altered the relationship of total phytoplankton diversity to flooded continental area and sea level that had persisted through the Paleozoic and Mesozoic (Figure 5). Total phytoplankton diversity is positively correlated with invertebrate diversity in the Paleozoic and Mesozoic (r² = 0.53), but becomes inversely related to invertebrate diversity in the Cenozoic (r² = 0.61). This pattern indicates that the transitions from the prasinophyte/acritarch-dominated Paleozoic ocean to the nannoplankton/dinoflagellate-dominated Mesozoic ocean to the diatom-dominated Cenozoic ocean were likely associated with changes in total primary production, nutritional quality, export production efficiency, and food availability for heterotrophic consumers.

The linear relationship between phytoplankton and invertebrate diversities can be used to infer changes in food-web structure and efficiency of trophic transfer of primary production into invertebrate species over time. This comparison highlights an increase in the number of invertebrate species relative to phytoplankton species through the Phanerozoic (Table 2): (a) each Paleozoic phytoplankton genus is associated with five invertebrate genera; (b) each Mesozoic dinoflagellate genus and nannoplankton species is associated with 7 and 14 invertebrate genera, respectively; and (c) each Cenozoic diatom genus is associated with 59 invertebrate genera.
### TABLE 2  
Invertebrate diversity supported per phytoplankton species or genera (the slope) over different time intervals

<table>
<thead>
<tr>
<th>Taxa</th>
<th>Time period</th>
<th>Slope</th>
<th>SE(^b)</th>
<th>Intercept</th>
<th>SE(^b)</th>
<th>(r^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diatom genera</td>
<td>Cenozoic</td>
<td>59.0</td>
<td>12.1</td>
<td>2308.0</td>
<td>418.6</td>
<td>0.75</td>
</tr>
<tr>
<td>Nannoplankton spp.</td>
<td>Mesozoic</td>
<td>14.5</td>
<td>1.3</td>
<td>707.5</td>
<td>100.4</td>
<td>0.85</td>
</tr>
<tr>
<td>Dinoflagellate genera</td>
<td>Mesozoic</td>
<td>6.9</td>
<td>0.6</td>
<td>844.1</td>
<td>85.6</td>
<td>0.86</td>
</tr>
<tr>
<td>Nannos + Dinos</td>
<td>Mesozoic</td>
<td>4.9</td>
<td>0.4</td>
<td>766.1</td>
<td>79.9</td>
<td>0.89</td>
</tr>
<tr>
<td>Acritarch genera</td>
<td>Paleozoic</td>
<td>5.2</td>
<td>2.3</td>
<td>454.1</td>
<td>322.9</td>
<td>0.12</td>
</tr>
<tr>
<td>Sum</td>
<td>Phanerozoic</td>
<td>9.0</td>
<td>1.7</td>
<td>187.4</td>
<td>316.0</td>
<td>0.27</td>
</tr>
</tbody>
</table>

\(^a\)Invertebrate diversity \((y)\) is estimated as a linear function of phytoplankton diversity \((x)\) \(\times\) slope + intercept. Diversity of all groups is represented by number of genera, except calcareous nannoplankton diversity, which is represented by number of species. Data from sources outlined in caption for Figure 3.

\(^b\)SE = standard error.

genera. This large change in the richness of invertebrate genera relative to the phytoplankton species and genera may represent an increase in the relative biomass of each of the phytoplankton species caused by an increase in primary production, an increase in the amount of primary production that is exported, or both. In part, the increasing ratio of preserved invertebrate to phytoplankton taxa also reflects the increasing richness of specialized predators through time; this relationship is another consequence of increasing primary production in the oceans, or increasing biological availability of primary production to consumers, or both (Bambach et al. 2002). In the contemporary oceans, diatoms are the major phytoplankton taxa contributing to export production (Dugdale et al. 1998). The increased export efficiency associated with the ecological success of marine diatoms most likely contributed to the increase in marine invertebrate diversity, size, and rate of energy consumption.

**Evolutionary Tempo of Marine Phytoplankton**

We compare evolutionary rates among different taxonomic groups based on background rates of extinction and origination normalized by species richness in the fossil record (Table 3). The results of this analysis show a twofold to fourfold decrease in the average longevity of the calcareous nannoplankton, dinoflagellates, and diatom species from the Mesozoic to the Cenozoic. Using a similar approach, Knoll (1994) described an increase in morphological diversity and a decrease in average species longevity in the acritarchs through the Proterozoic into the early Cambrian, and Barron (2003) documented relatively short longevities (~3 myr) in marine diatoms over the past 18 myr from high-resolution records in the Pacific and Southern Oceans. These results highlight the continuing increase in evolutionary tempo of phytoplankton. Extinction rates in dinoflagellates and calcareous nannoplankton have exceeded origination rates through the Cenozoic, which has resulted in declining diversities.
THE ROLE OF PLANKTON IN BIOGEOCHEMICAL CYCLES

Geological and biological processes modify atmospheric and seawater chemistry through time and influence global climates. Submarine and subaerial tectonics in combination with erosional processes are the primary suppliers of the major elements in geochemical cycles; the oxidation and reduction processes that alter mobile elemental reservoirs are biologically mediated. Hence, geological and biological processes together form feedback loops in various biogeochemical cycles.

As we have shown in the preceding sections, changing environmental conditions selected for different plankton groups through Earth’s history. In turn, phytoplankton have modified various aspects of the environment through time. Perhaps the most notable example of this relationship is the role that plankton played in the oxygenation of Earth’s atmosphere and oceans through the evolution of oxygenic photosynthesis. Increasing oxygen levels led to the oxidation of inorganic substrates, particularly iron and sulfur, which altered seawater chemistry. In turn, primary production became iron-limited as the result of iron oxidation in the oceans. In this section, we explore these types of biogeochemical interactions that have left clues preserved in long-term geological records.
Phytoplankton and the Carbon Cycle

The expansion of the red lineage of phytoplankton that began in the Mesozoic had an impact on the long-term carbon cycle, in part by altering the distribution of carbonate and organic carbon buried on the seafloor. Prior to the Mesozoic, most marine calcifying organisms lived in shallow coastal and shelf regions. Carbonate deposition was concentrated in these areas as a result, and deposition of pelagic carbonates was minimal. Two groups of carbonate-secreting plankton successfully competed for limited carbonate resources and expanded in the Mesozoic oceans—coccolithophores, with their calcite-plated armor, and planktonic foraminifera, with their carbonate tests. As these two groups radiated, the loci of marine carbonate deposition gradually expanded from shallow shelf areas to the deeper ocean (Southam & Hay 1981, Sibley & Vogel 1976) and the carbonate compensation depth (CCD) deepened (e.g., Wilkinson & Algeo 1989). Pelagic carbonate sedimentation has come to dominate as sea level and shelf area has declined since the Late Cretaceous, and the pelagic carbonate reservoir has increased at the expense of the shallow-water carbonate reservoir since the Mesozoic (Wilkinson & Algeo 1989).

At the same time, the expansion of marine phytoplankton in the Mesozoic resulted in greater export production through time (e.g., Bambach 1993) (see above). The newly-emerged eukaryotic phytoplankton efficiently exported organic matter, and the newly formed Atlantic Ocean margins increased the potential storage area for that organic matter. Substantial amounts of organic carbon were sequestered on the passive continental margins of the Atlantic and on flooded continental interiors (e.g., Arthur et al. 1984, Bralower 1999, Claypool et al. 1977, Jenkyns & Clayton 1997) as Pangea broke apart, export production increased, and organic matter was buried before it could be oxidized. At the same time, sedimentary carbon was both recycled at subduction zones and transferred to orogenic metasediments as the Tethys Sea and Pacific Ocean basins shrank. The circum-Atlantic sediments have not yet been recycled through subduction during the current Wilson Cycle, thereby providing long-term storage of large amounts of isotopically light organic carbon. This biologically mediated increase in organic carbon burial may account for as much as half of the long-term increase in δ^{13}C values recorded in marine carbonates (δ^{13}C_{carb}) (Katz et al. 2004) and organic carbon (δ^{13}C_{org}) (Hayes et al. 1999) from the Jurassic to the Miocene (Figure 7). Increased organic carbon burial contributed to a gradual depletion in CO₂ from the ocean-atmosphere system and a simultaneous increase in the oxidation state of Earth’s surface (Katz et al. 2004).

The long-term depletion of CO₂ acted as a feedback mechanism that was a key factor that selected β-carboxylation and C₄ photosynthetic pathways in marine and terrestrial photoautotrophs. Diatoms have β-carboxylation pathways (Morris 1980, Reinfelder et al. 2000) and dominate carbon export production in the modern ocean (Smetacek 1999). The rapid radiation of diatoms in the latter half of the Cenozoic enriched the ^13C composition of marine organic matter (Figure 7) (Katz et al. 2004).
A decrease in diatom cell size over the Cenozoic may also have affected δ\textsubscript{13}C\textsubscript{org} values (Figure 7) (ZV Finkel, ME Katz, JD Wright, OM Schofield & PG Falkowski, in review). In general, large phytoplankton cells produce high δ\textsubscript{13}C\textsubscript{org} because they tend to have low growth rates and low rates of diffusive flux (Laws et al. 1997, Popp et al. 1998, Rau et al. 1997). Early studies suggested that an increase in diatom size through the Cenozoic might have been responsible for the measured increase in δ\textsubscript{13}C\textsubscript{org} (Hayes et al. 1999). However, the median diatom cell size appears to have decreased through the Cenozoic (ZV Finkel, ME Katz, JD Wright, OM Schofield & PG Falkowski, in review), which contradicts the assertion of Hayes et al. (1999) that a trend toward larger diatoms alone drove the increase in δ\textsubscript{13}C\textsubscript{org}. Therefore, an increase in diatom abundance (rather than size) since the mid-Cenozoic likely contributed to the δ\textsubscript{13}C\textsubscript{org} increase (ZV Finkel, ME Katz, JD Wright, OM Schofield & PG Falkowski, in review; Katz et al. 2004).

Terrestrial ecosystems also contributed to the δ\textsubscript{13}C\textsubscript{org} increase. In the late Miocene (6 to 8 Ma), a global expansion of grasslands was coupled with a shift in dominance from C\textsubscript{3} to C\textsubscript{4} grasses. This shift produced 13\textsubscript{C}-enriched terrestrial biomass (Cerling et al. 1997, Still et al. 2003), some of which was ultimately transferred to and sequestered in the oceans (France-Lanord & Derry 1994, Hodell 1994) at the same time that 13\textsubscript{C}-enriched diatoms continued to expand. These new pathways are responsible for the δ\textsubscript{13}C\textsubscript{org} increase since the mid-Cenozoic, and contributed to the δ\textsubscript{13}C\textsubscript{carb} decrease that began in the mid-Miocene (Figure 7). The abrupt δ\textsubscript{13}C\textsubscript{org} increase occurred without a large change in either the atmospheric oxidation state or an injection of 12\textsubscript{C} from mantle outgassing, and appears to be a unique event in Earth’s history.

Whereas biological fractionation of carbon isotopes by phytoplankton is a major component of the carbon cycle, fractionation of sulfur isotopes occurs primarily through bacterial reduction of sulfate to sulfide. The carbon and sulfur cycles are linked because bacterial sulfate reduction depends on high levels of sedimentary organic matter. Sulfate reduction can result in pyrite burial, which drives δ\textsubscript{34}S of marine sulfate higher. For atmospheric oxygen levels to remain stable, the carbon and sulfur cycles must be counterbalanced so that as one reduced reservoir grows, the other shrinks (e.g., more organic carbon and less pyrite). In the simplest scenario, this hypothetical relationship indicates that intervals of high δ\textsubscript{13}C\textsubscript{carb} should coincide with intervals of low δ\textsubscript{34}S\textsubscript{sulfate}. In reality, this relationship is complicated by other factors, and redox conditions have fluctuated during times when the two cycles were not counterbalanced, especially on short timescales (e.g., Kump 1993, Payton et al. 1998, Strauss 1999). This situation appears to have been the case not only for some brief intervals, but also for the Jurassic to the Neogene, when the long-term trends in both isotope records show increasing values. These trends indicate that there were increases in the sedimentary reservoirs of reduced carbon (= organic matter) and reduced sulfur (= pyrite), which in turn requires corresponding increases in the oxidized species of both carbon and sulfur. This relationship supports the overall increase in the oxidation state of Earth’s surface over this long time period (Katz et al. 2004) (Figure 7).
Primary Productivity and Geochemical Proxy Records

Changes in phytoplankton taxonomic composition can influence the carbon isotope record on shorter timescales as well. Extended intervals of elevated global $\delta^{13}$C$_{\text{carb}}$ values are superimposed on the long-term $\delta^{13}$C increase from the Jurassic to the mid-Miocene (Figure 7). These intervals typically have been attributed to increases in organic carbon burial relative to carbonate burial that resulted from changes in a combination of surface ocean productivity and/or preservation on the seafloor (e.g., Miller & Fairbanks 1985, Scholle & Arthur 1980, Vincent & Berger 1985). Burial of isotopically light organic carbon leaves the remaining mobile carbon reservoir isotopically heavier, which drives $\delta^{13}$C$_{\text{carb}}$ higher. Comparisons among the records of $\delta^{13}$C$_{\text{carb}}$, $^{87}$Sr/$^{86}$Sr, and phosphorous flux may provide further insight into these relationships (Figure 7).

Phosphorous is essential for all cells; this element is required for the synthesis of nucleic acids, metabolism of carbohydrates, and formation of membrane lipids. It is supplied only through continental erosion and riverine delivery to the oceans, where it is either used quickly or authigenically precipitated. Phosphorous can be recycled in the upper ocean as organic matter is oxidized and phosphorous is returned to the surface. Because primary production can be limited by phosphorous availability, we compare proxies for productivity ($\delta^{13}$C$_{\text{carb}}$) (Katz et al. 2004) with phosphorous flux (Follmi 1995) (Figure 7). Episodes of elevated $\delta^{13}$C$_{\text{carb}}$ tend to be accompanied by higher phosphorous fluxes prior to the mid-Miocene, although the inverse is not true (this difference may in part reflect the global $[\delta^{13}$C$_{\text{carb}}$] versus regional [phosphorous flux] nature of the records). The phosphorous flux and $\delta^{13}$C$_{\text{carb}}$ curves decouple most notably during two time intervals: (a) in the Late Cretaceous, when widespread deposition of chalks occurred in shallow waters, where organic carbon must have been oxidized and phosphorous is returned to the surface. Because primary production can be limited by phosphorous availability, we compare proxies for productivity ($\delta^{13}$C$_{\text{carb}}$) (Katz et al. 2004) with phosphorous flux (Follmi 1995) (Figure 7). Episodes of elevated $\delta^{13}$C$_{\text{carb}}$ tend to be accompanied by higher phosphorous fluxes prior to the mid-Miocene, although the inverse is not true (this difference may in part reflect the global $[\delta^{13}$C$_{\text{carb}}$] versus regional [phosphorous flux] nature of the records). The phosphorous flux and $\delta^{13}$C$_{\text{carb}}$ curves decouple most notably during two time intervals: (a) in the Late Cretaceous, when widespread deposition of chalks occurred in shallow waters, where organic carbon must have been oxidized, and (b) in the late Cenozoic, when widespread glaciation increased erosional rates and phosphorous supply to the oceans. These comparisons indicate that phosphorous is not always a limiting nutrient on geological timescales.

Strontium isotopes may provide another clue to link biogeochemical records. In a simple two-source system (Caldeira 1992), strontium is delivered to the oceans through hydrothermal exchange at midocean ridges (low $^{87}$Sr/$^{86}$Sr values) and continental erosion (high $^{87}$Sr/$^{86}$Sr values); however, the $^{87}$Sr/$^{86}$Sr may be complicated by changes in dominant continental source rock type (e.g., Ravizza 1993) and variable riverine fluxes (e.g., Lear et al. 2003). Over the past 200 myr, four out of five decreases in $^{87}$Sr/$^{86}$Sr values correspond to major episodes of elevated $\delta^{13}$C$_{\text{carb}}$ values (Figure 7). The fifth decrease occurs in the Turonian across a data gap in our $\delta^{13}$C$_{\text{carb}}$ record, but may correlate to a smaller, shorter interval of elevated $\delta^{13}$C$_{\text{carb}}$ documented in published Tethyan records (Jenkyns et al. 1994, Stoll & Schrag 2000). The causal mechanisms behind these correlations are unclear, but may be related to higher pCO$_2$ from increased hydrothermal activity (e.g., Berner 1993) that accelerated the geological and biological components of the global carbon cycle.
SUMMARY

Cyclic tectonic changes superimposed on key secular changes in Earth’s atmosphere, oceans, and even on land have selected for certain phytoplankton clades through time, with an ever-increasing tempo of phytoplankton evolution. In this paper, we have focused on the evolutionary paths that eventually led to the eukaryotic phytoplankton that dominate the contemporary oceans—coccolithophores, diatoms, and dinoflagellates. The earliest primary producers were prokaryotes. For much of the Archaen and Proterozoic Eons, the oceans were dominated by cyanobacteria, with green and perhaps other algae increasing in importance toward the end of the Precambrian. Planktonic algae radiated in the Early Phanerozoic oceans, cyst-forming dinoflagellates and calcareous nannoplankton dominated in the Mesozoic oceans, and, finally, the diatoms rose to prominence in the latter half of the Cenozoic. The primary producers have always been at the base of the food web; hence, the evolution of organisms at higher trophic levels has depended on the evolutionary trajectories of the phytoplankton. The number of invertebrate species relative to phytoplankton species has increased through the Phanerozoic.

This evolutionary succession of marine phytoplankton was a response to a complex system that cannot be explained by a set of ordinary differential equations. Secular shifts in redox seawater chemistry have influenced phytoplankton evolutionary trajectories, both by altering the trace metal availability in the oceans and by changing the balance of fixed nitrogen between the oxidized form (nitrate) and the reduced form (ammonium). In general, the more reducing conditions of the early oceans favored the green lineage, while the higher oxidation states of the later oceans favored the red lineage. Early eukaryotic phytoplankton were best able to compete in the better-oxygenated coastal regions, while green phytoflagellates thrived in open ocean surface waters, where seawater remained Fe-rich and relatively Zn- and Cd-poor (Anbar & Knoll 2002, Whitfield 2001b). Diversity increases in phytoplankton appear to correlate with continental rifting of Rodinia (Late Proterozoic), Pannotia (Early Paleozoic), and Pangea (Jurassic), which ultimately resulted in the three groups of eukaryotic phytoplankton that dominate the modern ocean: coccolithophores, diatoms, and dinoflagellates.

While changing environmental conditions selected for different plankton groups through Earth history, phytoplankton, in turn, influenced biogeochemical components of the environment, often through the biologically mediated oxidation and reduction processes that alter mobile elemental reservoirs. The best example of this process is the role that plankton played in oxygenating Earth’s atmosphere and oceans through the evolution of oxygenic photosynthesis. The subsequent rise of the eukaryotic phytoplankton since the Early Jurassic, coupled with the opening of the Atlantic Ocean basin during the current Wilson Cycle, has increased the efficiency of organic carbon burial and contributed to a gradual depletion of CO₂ from the oceans and atmosphere, with a simultaneous increase in the oxidation state of Earth’s surface. Ultimately, this change favors the red lineage.
ACKNOWLEDGMENTS

We thank Ken Miller, Oscar Schofield, and Scott Wing for their comments on this manuscript, Ben Cramer for his assistance with statistical analyses, and Richard Bambach for discussions and for providing his revised version of Jack Sepkoski’s marine invertebrate database. This study was supported by NSF OCE 00,84032 Biocomplexity: The Evolution and the Radiation of Eukaryotic Phytoplankton Taxa (EREUPT).

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Figure 1  Relationships among photosynthetic eukaryotes inferred from 18S rRNA gene sequences (species names and sequence accession numbers are not shown for the clarity of the figure). Names underlined with one or two strokes indicate taxa that contain primary or secondary endosymbiotic plastids (green or red), respectively. Colored branches indicate taxa that contain photosynthetic plastids. Phylogenetic tree was constructed with the PHYLIP software package, in which genetic distance calculation (F84 substitution model) and the neighbor-joining method were used. Tree displayed is consistent with consensus trees obtained from 1,000 bootstrapped data sets.
Figure 2  Phylogenetic tree of plastids inferred from 16S rRNA gene sequences rooted with anoxygenic photosynthetic bacteria. Tree shows all plastids originating once from cyanobacteria, likely from an ancestral lineage that subsequently became extinct. Data from euglenophytes and chlororachiophytes (secondary green plastids) and from dinoflagellates (two types of secondary red plastids) have been excluded because of high genetic divergences responsible for inconsistent phylogenetic branchings. Phylogenetic tree was constructed with the PHYLIP software package, in which genetic distance calculation (F84 substitution model) and the neighbor-joining method were used. Tree displayed is consistent with consensus tree obtained from 1,000 bootstrapped data sets. Each species name is preceded by the accession number of the DNA sequence used in the phylogenetic analysis. Colors indicate groups of photosynthetic plastids.
**Figure 3** Comparison of eukaryotic phytoplankton diversity curves (after Katz et al. 2004) with sea-level change (Mesozoic-Cenozoic [Haq et al. 1987] and Paleozoic [Vail et al. 1977]), flooded continental areas (Ronov 1994), and marine genus (blue) and family (green) invertebrate diversities (Sepkowski 1997). Phytoplankton species (red) diversities are from published studies (calcaceous nannofossils [Bown et al. 2004], dinoflagellates [Stover et al. 1996], diatoms [Spencer-Cervato 1999], acritarchs [Proterozoic (Knoll 1994)], [Phanerozoic (R.A. MacRae, unpublished data)]). Phytoplankton genus (blue) diversities were compiled for this study from publicly available databases (calcaceous nannofossils and diatoms [Spencer-Cervato 1999], dinoflagellates, and acritarchs [R.A. MacRae, unpublished data]. All records are adjusted to the Berggren et al. (1995) (Cenozoic), Gradstein et al. (1995) (Mesozoic), and GSA (http://rock.geosociety.org/science/timescale/timescl.htm) (Paleozoic) timescales. Taxon-specific biomarkers (Moldowan & Jacobson 2000) and C28/C29 sterane ratios (Grantham & Wakefield 1988) provide a record of increased biomass preservation of eukaryotic phytoplankton in the Mesozoic-Cenozoic. Episodes of supercontinent rifting are shaded. We base our interpretations of acritarch evolution on two compilations: a high-quality Proterozoic and Early Cambrian record of acritarch species (Knoll 1994) and a global, genera-level, Phanerozoic, quality-controlled literature compilation that may include taxonomic synonyms (R.A. MacRae, personal communication). Biological uncertainty exists in the interpretation of different genera and species within the acritarch record. Even though Knoll’s (1994) record underestimates global diversity, and MacRae’s record overestimates it, the long-term patterns of diversification and extinction appear consistent and robust. Data are available at http://mychronos.chronos.org/~miriamkatz/20040721
Figure 7  Geochemical proxy records showing comparisons of phosphorous flux [(Follmi 1995); curve fit (B.S. Cramer, personal communication) uses the SSA-MTM Toolkit from http://www.atmos.ucla.edu/tcd/ssa/(Ghil et al. 2002)], bulk sediment $\delta^{13}$C$_{carb}$ (Katz et al., 2004), strontium isotopes (Howarth & McArthur 1997), sulfur isotopes (Canfield 1998, Payton et al. 1998, Payton et al., 2004, Strauss 1999), and bulk sediment $\delta^{13}$C$_{org}$ (Hayes et al. 1999). Timescales are as in Figure 3. Data are available at http://mychronos.chronos.org/~miriamkatz/20040721