Evolutionary inheritance of elemental stoichiometry in phytoplankton

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The elemental composition of phytoplankton is a fusion of the evolutionary history of the host and plastid, resulting in differences in genetic constraints and selection pressures associated with environmental conditions. The evolutionary inheritance hypothesis predicts similarities in elemental composition within related taxonomic lineages of phytoplankton. To test this hypothesis, we measured the elemental composition (C, N, P, S, K, Mg, Ca, Sr, Fe, Mn, Zn, Cu, Co, Cd and Mo) of 14 phytoplankton species and combined these with published data from 15 more species from both marine and freshwater environments grown under nutrient-replete conditions. The largest differences in the elemental profiles of the species distinguish between the prokaryotic Cyanophyta and primary endosymbiotic events that resulted in the green and red plastid lineages. Smaller differences in trace element stoichiometry within the red and green plastid lineages are consistent with changes in trace elemental stoichiometry owing to the processes associated with secondary endosymbioses and inheritance by descent with modification.

Keywords: phytoplankton; evolution; stoichiometry; endosymbiosis; trace metals; Redfield ratio

1. INTRODUCTION

Oxygenic photosynthesis by phytoplankton has significantly contributed to the oxidation state of the ocean and atmosphere [1–3]. Based on inferred atmospheric and oceanic oxygen concentrations, biomolecular and isotopic evidence, phytoplankton may have been present 2.7–2.4 Ga ago ([4–6]; but see [7]). These would probably have been the Cyanophyta, the only extant prokaryotic group of oxygenic phototrophs [6]. Almost a billion years later, organic walled fossils attributable to eukaryotic phytoplankton appeared in the fossil record [6,8,9]. Three lineages of photosynthetic organisms became established as a result of primary endosymbiosis: the green plastid lineage (chlorophyll (chl) b-containing green algae and land plants, Viridiplantae), the red plastid lineage (chl c-containing, Rhodophyta) and the Glaucocystophyta (also known as Glaucophyta) [10–12].

Later, secondary endosymbiotic events occurred in both the green and red plastid lineages. In the green lineage, this gave rise to the Chlorarachniophyceae and Euglenophyceae. In the red plastid lineage, secondary endosymbiosis gave rise to the Cryptophyceae, the Heterokontophyta (including the Bacillariophyceae and Chrysophyceae), the Haptophyceae, the chl c-containing Dinophyceae and several other groups (figure 1). This diverse assemblage of eukaryotic secondary chl c-containing red phytoplankton is collectively referred to as the Chromalveolates [12–14]. In the Dinophyceae, secondary endosymbiotic events also gave rise to chl a and b (e.g. Gymnodinium chlorophorum) and chl a and c with peridinin plastid group (figure 1). As a result of tertiary endosymbiosis, chl a and c with fucoxanthin-containing dinoflagellates arose making them the most diverse and complex of all the phytoplankton groups (e.g. [15,16]). Prior to the Mesozoic, the fossil record indicates that the Cyanophyta and members of the eukaryotic green plastid lineage probably dominated phytoplankton assemblages; by contrast, the red plastid lineage dominates the contemporary ocean [8,11,17]. Coincident with the major evolutionary changes in the phytoplankton are shifts in the oxidation state of the atmosphere and ocean and the availability of macro- and micro-nutrients [18–21].

Taxonomic differences in macro- and trace element to P (or to C or N) ratios in phytoplankton associated with fundamental biochemical differences or unique phenotypic strategies in response to their environment have been revealed from decades of careful field and laboratory work [22–26]. Unfortunately, well-established variability in elemental stoichiometry in response to growth rate and environmental conditions has made it difficult to quantify taxonomic differences in multi-element composition from multiple studies. A statistical analysis of a multi-element study conducted under defined nutrient-replete environmental conditions on 15 eukaryotic marine phytoplankton species revealed significant variability between species and distinctive elemental profiles associated with the green and red plastid lineages [27]. Phytoplankton in the green plastid lineage (products of a primary endosymbiotic event: Chlorophyceae and Prasinophyceae) generally had higher C, N, Fe, Zn and...
Cu to P ratios than those in the red plastid lineage (products of a secondary engulfment event: Haptophyceae, Bacillariophyceae and Dinophyceae) when grown under identical conditions. Based on these results, the authors hypothesized that the distinctive elemental signatures across phytoplankton hierarchies under nutrient-replete conditions reflected evolutionary histories in response to the different selective forces present at the time of their evolution and radiation, combined with fundamental biochemical and genetic constraints [11,27].

The elemental composition of each individual species is a function of its unique evolutionary history and we expect a large variation in elemental composition between species. The evolutionary inheritance hypothesis predicts that there may be some broad similarities in elemental composition for some elements at higher taxonomic levels based on fundamental genetic differences between the groups, most probably associated with engulfment events. To test the evolutionary inheritance hypothesis, we extended the elemental analysis presented in Quigg et al. [27] with the present study to take in 14 more species including Cyanophyta, Glaucocystophyta and a range of species from the red and green lineages representing both primary and secondary endosymbiotic events (electronic supplementary material, table S1, and figure 1). We quantify the taxonomic difference in elemental stoichiometry on the combined dataset (present study [26,27]). If the elemental composition of different algal groups reflects evolutionary history, then we would predict that the Cyanophyta would have an elemental signature more similar to the taxonomic groups established by a primary endosymbiotic event and most different from the groups established by secondary endosymbiotic events. In general, we would expect relatively less difference in elemental composition within a lineage than between lineages, i.e. members of the green plastid lineage would have an elemental composition more similar to one another than members of the red plastid lineage, but within the lineages, subsequent endosymbiotic events could lead to further divergence in elemental profiles.

2. MATERIAL AND METHODS

(a) Culturing and growth conditions

Culturing, sampling and elemental analysis were all performed essentially as described in Quigg et al. [27] and Ho et al. [26]. Ten marine phytoplankton
supplementary material, table S1) were grown in defined nutrient-replete medium under standard conditions (19 ± 1°C, 12 L : 12 D cycle, 250 µ mol quanta m⁻² s⁻¹; modified Aquil). *Trichodesmium* sp. was grown at 26°C, 85 µ mol quanta m⁻² s⁻¹ (12 L : 12 D cycle) in YBCII medium [28]. Freshwater cultures were grown in Fraquil [29]. Nutrients (N, P and Si), EDTA, trace metals and vitamin stock solutions were added to Fraquil in concentrations defined for the modified Aquil recipe [26].

(b) Growth rates and cell size

Replicate exponentially growing cultures (n ≥ 3) for each species were maintained in semi-continuous batch cultures. Growth rates for most marine species were followed by measuring daily changes in cell density using a calibrated Coulter Multisizer particle counter. Growth rates of the freshwater species were followed by measuring daily changes in *in vivo* chlorophyll fluorescence. *Synechococcus* sp. and *Cyanophora* sp. were counted daily using a Neubauer haemocytometer. Specific growth rates (µ, d⁻¹) were calculated from the initial (F₀) and final (F) relative fluorescence values according to 
\[
u = \frac{\ln F - \ln F₀}{t - t₀},
\]
where *t* was time measured in days or using \(\mu = \frac{\ln C - \ln C₀}{t - t₀}\), where *C* was the cell concentration (cells ml⁻¹). *Trichodesmium* growth rates were measured according to Berman-Frank et al. [30].

(c) Sample preparation and analysis

Detailed information on sample preparation and analysis was provided in Quigg et al. [27] and Ho et al. [26]. Cellular C and N concentrations were measured with a CHN elemental analyser while all other elements were measured using sector field high-resolution inductively coupled plasma mass spectrometry (HR-ICP-MS). Phyttoplankton were filtered gently (less than 130 kPa) onto acid-cleaned polycarbonate filters and rinsed with chelated 0.45 M NaCl [26] and Milli-Q H₂O for marine and freshwater species, respectively. Optima grade HNO₃ was used in all HR-ICP-MS sample preparation and analysis. In order to further improve HR-ICP-MS calibration accuracy and precision, both external and internal standard curves were included with each sample run in this study according to Cullen et al. [31]. Concentration differences were less than 3 per cent for most elements between the two standard curves.

(d) Statistical analyses using the combined dataset

To test hypotheses on the evolutionary inheritance of stoichiometry in phytoplankton, we combined the findings from the present study with those in an earlier study (electronic supplementary material, table S1). Patterns in trace element composition were examined using principal component analysis (PCA) on the correlations of log-transformed trace element to phosphorus (P) ratios [32]. We tested for statistical differences in element : P ratios across classes using ANOVA and pairwise t-tests with Holm’s sequential correction [32] using log C, N, Fe, Mn, Zn, Cu, Co and Cd to P ratios. To analyse the hierarchical structure in the elemental profiles of the phytoplankton, we calculated the distance between average log (element : P) ratios and used these distances to construct a dendrogram. The squared distance between observations *i* and *j* was defined as:
\[
d_{ij}^2 = \sum_{k} (x_{ik} - x_{jk})^2, \quad k = \text{Fe, Mn, Zn, Cu, Co, Cd},
\]
where \(x_{ik} = \log(k : P)\) for observation *i* and \(x_{jk}\) were divided by the standard deviation of the observations for all trace elements so that each element ratio contributed equally to the distance. To analyse the distance between groups (Cyanophyta, Chlorella, *Gephyrocapsa oceanica* and *Gephyrocapsa oceanica* accumulate such high concentrations of Ca (e.g. [26,27]) as it is used to construct calcite and aragonite laths. In a study on the growth of the diatom (Bacillariophyceae) *Phaeodactylum*

3. RESULTS

Phytoplankton were grown under nutrient-replete conditions and had specific growth rates ranging from 0.23 to 1.11 d⁻¹ (electronic supplementary material, table S1). Elemental ratios for each of the 14 species examined and each element measured are summarized in electronic supplementary material, table S2. Elemental quotas were normalized to P rather than to C or N for direct comparison with previous studies. Statistically similar patterns were observed whether we normalized to P or C or N (see below) [33,34]. Small differences in the average elemental profiles relative to previous studies reflect differences in the relative number of species and observations and proportion of species representing the primary and secondary endosymbiotic events within the lineages [26,27].

(a) Major nutrients: C, N, P and S

The average C : N : P : S ratio for the 29 species investigated was 132 : 18 : 1 : 0.99, compared with the canonical Redfield ratio of 106 : 16 : 1 : 1.7 [35] and consistent with earlier studies with phytoplankton grown under identical conditions [26,27], and in other laboratories and in the field (see reviews of Duarte [36] and Geider & La Roche [37]). While the ratios of the major nutrients were variable (electronic supplementary material, table S2), patterns between plastid lineages were observed (electronic supplementary material, figure S1). C : P is significantly higher in G1 relative to GLA, R1, R2 and RG2 while the C : P in G2 did not differ from other groups. N : P is significantly higher in G2 relative to R1 and R2, but does not differ from other groups. The green plastid lineage (G1 and G2) also tends to have higher C : N ratios. Our average S : P ratio of 0.75 (electronic supplementary material, table S2) for the 14 species in the current study (and 0.99 for all 29 species investigated) fell between the range of previous reports: 1.7 in Redfield et al. [35], 0.65 in Payne & Price [38], 0.3 in Fraga [39] and 1.3 in Ho et al. [26].

(b) Major cations: K, Mg, Ca and Sr

Despite the ubiquitous use of cations in cellular processes [39], phyttoplankton accumulate them to differing degrees (electronic supplementary material, table S2). High concentrations of Ca : P (94 ± 5 mmol : mol) and Sr : P (107 ± 4 mmol : mol) specifically were observed in the cyanobacterium *Trichodesmium* sp. Typically, only calcifying phytoplankton such as the coccolithophores *Emiliania huxleyi* and *Gephyrocapsa oceanica* accumulate such high concentrations of Ca (e.g. [26,27]) as it is used to construct calcite and aragonite laths. In a study on the growth of the diatom (Bacillariophyceae) *Phaeodactylum*
tricornutum, Hayward [40] found variations in the external concentration of Na, K, Ca and Mg had little effect on the internal concentration of these elements. Thus, differences in the media (modified Aquil, Fraquil and YBCII) cannot be assumed to be primarily responsible for the differences in cation : P ratios among the phytoplankton examined.

**Trace elements: Fe, Mn, Zn, Cu, Co, Cd and Mo**

The major trace elements (and Sr) were present in 1000-fold lower concentrations than the nutrient elements and other major cations (electronic supplementary material, figure S1 and table S2). Mean concentrations for the 14 species were generally consistent with the range of element : P previously reported for phytoplankton growing under similar conditions [26, 27]. All subsequent analysis refers to the combined dataset of 87 observations on 29 species.

The average mmol Fe : Mn : Zn : Cu : Co : Cd to mol P ratio for the whole dataset is 18 : 2.8 : 1.34 : 0.50 : 0.11 : 0.11. Cyanophyta, Glaucocystophyta and the green plastid lineage had the highest Fe : P ratios while the red plastid lineage tended to have lower values (electronic supplementary material, figure S1 and table S2). Relative to other species examined, Cyanophora paradoxa (Glaucocystophyta) and Chlorarachniophycna (Chlorarachniophycna) had the greatest Zn quotas (5.0 + 0.8 and 4.7 + 0.11 mmol Zn : mol P, respectively). Cobalt and cadmium quotas were several orders of magnitude higher in Eutreptiella sp. (Euglenophyceae), which also has significantly lower Zn quotas than the other species examined.

**Statistical analysis of phylogenetic hierarchies**

PCA is a visual aid that can be used to identify taxonomic differences and similarities in log trace elemental : P profiles. The first two principal components of PCA account for 76 per cent of the variance in trace metal composition (46% for the first, 30% for the second). The prokaryotic Cyanophyta, the Glaucocystophyta and the green and red
plastid lineages are clearly separated along the two principal components (figure 2). The green plastid lineage is closer and exhibits more overlap than the red plastid lineage with the Cyanophyta. There is also some evidence of separation between the primary and secondary endosymbiotic events within the red and green plastid lineages. Phycobilisome-containing groups, the Cyanophyta, the Glaucocystophyta and the Cryptophyceae, cluster on the far right end of the first principal component axis.

The analysis of variance for each log element : P shows significant differences in elemental ratios at the phylum and species level. Pairwise t-tests with Holm’s correction document significant differences in most of the log element : P ratios at the lineage or class level (electronic supplementary material, table S3A), except for Co : P. Some of the clearest taxonomic differences in the elemental ratios are: log Fe : P is lowest in the secondary red plastid lineages; log Zn : P is highest in the Glaucocystophyta and the secondary red plastid lineage; and log Cd : P is lowest in the Cyanophyta and highest in the green plastid lineage. The green plastid lineage had, on average, higher Fe, Cu and Cd to P ratios than the red plastid lineage, while Mn : P is higher in the red relative to the green plastid lineage (figure 2). Cyanophyta and Glaucocystophyta often had trace element ratios more similar to the green relative to the red plastid lineage. Within the green plastid lineage Eutetidella sp. (Euglenophyceae, G2) has considerably higher C : P and N : P, Cd : P and Co : P relative to the Chlorarachniophyceae Glaucocystophyta and Cryptophyceae, cluster on the far right end of the first principal component axis.

4. DISCUSSION
Phytoplankton are a polyphyletic group in which primary, secondary and tertiary endosymbiotic engulfment events gave rise to the diversity of species seen today (figure 1). The Cyanophyta are the likely contributor of the ancestral plastid to the green and red plastid lineages and the Glaucocystophyta as a result of primary endosymbiotic engulfment events. The lineages resulting from these primary endosymbiotic events exhibit significant differences in macro- and trace elemental composition (log Fe, Mn, Zn, Cu, Cd to P ratios). Furthermore, there are significant, albeit smaller, differences in the elemental stoichiometry within the green and red plastid lineages (figures 2 and 3). We propose that the larger difference in elemental composition between the major lineages relative to the differences within the lineages supports the hypothesis that there is evolutionary inheritance of micro-element requirements in the phytoplankton.

Element ratios are known to change in response to growth rate and environmental conditions, including irradiance, temperature, CO₂, pH, the availability of nutrients in the medium and numerous interactions between trace elemental uptake mechanisms. We have not attempted and cannot account for all these myriad effects, but we have minimized them by maintaining the experimental species in exponential growth phase in nutrient-enriched media. Under these conditions, we find significant differences in the elemental profiles at the species level and across the lineages.
consistent with what is known about their unique evolutionary histories, physiological capacities and adaptations to specific environmental conditions [27]. We interpret the coherent statistically significant differences in the average elemental composition of these species within higher taxonomic groupings, such as the green versus red plastid lineages, to represent a residual phylogenetic signal associated with their common evolutionary history. Distances between average trace elemental profiles (figure 3) between the lineages are consistent with phylogenetic trees based on nucleic and protein sequences (e.g. [12–14,47]). Some of the more obvious elemental differences among the lineages are higher Fe, Zn and Cu to P ratios in the green plastid lineage versus the red plastid lineage (figure 2). It has been hypothesized that these differences in trace element composition between the lineages reflect differences in the acquired plastids, termed the plastid imprint hypothesis [27]. A comparison of trace element stoichiometry of species with similar plastids but different hosts and similar hosts but different endosymbionts permits a coarse assessment of the relative importance of the host relative to the plastid in determining trace element composition. There are three lines of evidence that support the plastid imprint hypothesis: (i) the trace element composition of chl ε dinoflagellates with plastids inherited from the red lineage cluster closely with the red plastid lineage while *G. chlorophorum*, a chl b-containing dinoflagellate, clusters with the green plastid lineage based on trace element stoichiometry [27]; (ii) similarity in the trace element stoichiometry in the secondary red plastid lineages despite differences in host lineages (alveolate host associated with the dinoflagellates versus a pre-chromist-type host associated with the Bacillariophyceae (diatoms), Haptophyceae and Cryptophyceae (figure 1)); and (iii) species with phycobilisomes, the Cryptophyta (prokaryotes, no host), the Glaucoystophyta (a product of primary endosymbiosis) and the Cryptophyceae (a product of secondary endosymbiosis), have similar trace element stoichiometry (high Fe : P and Zn : P) (figure 2). There is some evidence that the host may also contribute, in some cases, to trace element stoichiometry. Within the green plastid lineage, *Eutreptiella* sp. (Euglenophyceae, G2) has considerably higher Cd : P relative to the *Chlorarachnion* spp. (Chlorarachniophyceae, G2) (electronic supplementary material, table S2).

What role do the subsequent secondary engulfment events that produce the secondary red and green lineages play in altering the metabolome? The smaller differences in trace element stoichiometry within the red and green plastid lineages (G1–G2 and R1–R2) are consistent with changes in trace elemental stoichiometry, the metabolome [48], owing to the processes associated with secondary endosymbioses and inheritance by descent with modification of trace element stoichiometric profiles over time (figure 3 and electronic supplementary material, table S3B). Although the specific mechanisms controlling the evolution of the metabolome during the process of engulfment and successful endosymbiotic events are unknown, the potential for gene loss, reorganization and radical changes in gene regulation may become increasingly constrained with each subsequent engulfment event. In the primary engulfment event, the host cyanobacteria lost or transferred the majority of its genes to the host nucleus [12–14,47], and new regulation pathways must be established to control metabolite transport between the endosymbiont and the host [12]. In subsequent secondary engulfment events, a eukaryotic symbiont with its own nuclear and plastid genome must be housed within a new eukaryotic host. After the primary endosymbiotic event, subsequent endosymbiotic events may be more likely to maintain established regulatory pathways, and the associated trace metal metabolome, which contribute to successful endosymbiosis, as genes are transferred from the symbiont plastid and nucleus to the host nucleus.

All species require Fe, Mn, Zn, Cu and Co for growth, so differences in trace element stoichiometry probably reflect differences in the relative demand for trace metals in biochemical pathways, transporter capabilities and storage strategies, and not simply the presence or absence of a requirement for a particular metal or metalloenzyme. Cadmium may be an exception. Generally considered a toxin and not required for growth, it has been shown to be used as a metallo-centre in carbonic anhydrase in a diatom [49]. Similarly, nitrogen fixation, present in some species of the Cyanophyta, has a high Fe and Mo requirement owing to the enzyme nitrogenase [30,50,51]. We propose that the combination of gene loss, transfer of plastid genes to the nucleus and the establishment of new regulatory pathways to balance the needs of the plastid in the host [12] under different selection regimes could be responsible for the relatively large differences in the trace metal metabolome among the Glaucoystophyta, green and red plastid lineages.

The strong correlation between phylogenetic differences and trace element stoichiometries is consistent with the engulfment events occurring at different times in oceanic environments with different metal availabilities. The Cyanophyta originated in a reduced ocean with low pO2 in surface and deep waters and high bioavailability of Fe, Mn and Co and lower availability of Zn and Cd [20,52–55], which appears to be reflected in a high Fe : P and lower Cd : P in their particulate matter. Shifts in ocean chemistry from an anoxic Fe-rich ocean to a euxinic/sulphidic and then a more oxic ocean over the last two billion years of Earth’s history have altered the bioavailability of many trace elements [20,52–55]. The euxinic ocean would probably have had lower Fe, Mn, Co and Zn relative to the anoxic ocean, while the modern oxic ocean has relatively higher bioavailability of Zn and Cu but lower Fe, Mn and Co than past reducing oceans. An analysis of proteomes reveals differences in the abundance of Zn-and Fe-binding domains between prokaryotic and eukaryotic proteomes consistent with eukaryotes evolving in a more oxic environment [54].

If the red and green plastid lineages originated contemporaneously in similar environments, we might expect that the trace elemental profiles would be similar; alternatively, the lineages may have originated in contrasting environments. Some evidence suggests eukaryotes may have originated in shallow coastal oxic zones with terrestrial sources of metals, but their distribution and diversification were impeded by the low availabilities of Cu and Mo in the euxinic ocean [6,19,56]. Based on fossil and molecular evidence, it appears that the green plastid lineage (G1) may have been more ecologically
dominant in the Neoproterozoic and Paleozoic (1000–251 Myr) ocean and the red plastid lineage, specifically those lineages produced by a secondary endosymbiotic event, the Bacillariophyceae, Haptophyceae and Dinophyceae (R2), became more ecologically dominant in the Mesozoic and Cenozoic (251 Ma to present) ocean [4, 5, 8]. If the radiation of the lineages was constrained until the elemental needs and physiological strategies of the groups matched the dominant environmental conditions in the ocean surface layers, then the differences in the trace element stoichiometries may better match, or have been secondarily affected by, the environmental conditions when the lineages held ecological dominance and radiated as opposed to when the lineages first originated.

Changes in environmental conditions continue to have differential effects on phytoplankton community composition. Oceanic species of eukaryotic phytoplankton from the same taxonomic group have adapted to grow at much lower concentrations of Fe, Zn, Mn, Cu and Cd than those associated with coastal and estuarine environments [57, 58]. Similar differences may exist in prokaryotic species. A sequence analysis found that a coastal Synechococcus CC9311 has an increased capacity to transport, store or use Fe and Cu compared with an oceanic strain [59]. A potential consequence of phylogenetic differentiation in specific metal requirements is that ongoing environmental change may relegate some taxonomic groups to minor or regional status (e.g. Glaucocystophyta) as the habitats they dominate change from being global to regional.

In summary, there is a great deal of variability in the elemental composition of different phytoplankton species when grown under similar nutrient-replete culture conditions. Although there is variability in elemental composition at the species level, there are similarities in elemental composition in species aggregated into higher taxonomic groupings, which reflect their common evolutionary history. Evolutionary change can occur throughout the history of a lineage, although endosymbiotic events are among the most dramatic and rapid of such changes. The probability of metal profiles of so many different species clustering into the Cyanophyta, Glaucocystophyta and the green and red plastid lineages (and separating on primary and secondary endosymbioses) is low unless much of the trace metal differences were established by the endosymbiotic events (figures 2 and 3). Much more work is required to compile data on the elemental composition of species from diverse taxonomic groups under a variety of environmental conditions to confirm the evolutionary inheritance hypothesis and better understand the evolutionary and physiological controls and ecological consequences of elemental composition in organisms.

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REFERENCES
17 Katz, M. E., Finkel, Z. V., Grzebyk, D., Knoll, A. H. & Falkowski, P. G. 2004 Evolutionary trajectories and


anhydrase from a marine diatom. Nature 435, 42. (doi:10.1038/435042a)


