High sequence variability, diverse subcellular localizations, and ecological implications of alkaline phosphatase in dinoflagellates and other eukaryotic phytoplankton

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INTRODUCTION

In marine ecosystem, phosphorus (P) is a vital nutrient controlling the growth of phytoplankton thereby affecting primary production (Karl, 2000; Paytan and McLaughlin, 2007). There are two major P reservoirs in the ocean, the dissolved inorganic P (DIP) and dissolved organic P (DOP). DIP is preferable for the growth of phytoplankton; however, its concentration is often so low that it limits phytoplankton growth in marine environments (Benitez-Nelson, 2000; Karl, 2000; Dyhrman et al., 2006; Meseck et al., 2009). In various areas of the open ocean, DIP occurs in nanomolar concentrations (Wu et al., 2000; Thingstade et al., 2005) and has been shown to be a limiting or co-limiting factor of nitrogen fixation (Sañudo-Wilhelmy et al., 2001; Mills et al., 2004) and primary production (Paytan and McLaughlin, 2007). Even in coastal waters, DIP can be stoichiometrically limited by high amounts of nitrogen associated with human activities (Cloern, 2001; Huang et al., 2003; Scavia and Bricker, 2006; Zhang et al., 2007c). Eutrophication with stoichiometric DIP limitation can potentially impose coastal health problems because some harmful algal bloom forming species, such as Alexandrium spp., Dinophysis acuminata, Karlodinium veneficum, and Pseudo-nitzschia multiseries can elevate their toxin production under P-limited conditions (Anderson et al., 2002; John and Flynn, 2002; Fu et al., 2010; Mooney et al., 2010).

To meet the P requirement for growth, different groups of phytoplankton have developed different strategies (Dyhrman et al., 2007). Under P limitation, some phytoplankton species can use DIP that is stored up when DIP is sufficient (Falkner et al., 1998; Ou et al., 2008), while some can lower cellular demand of DIP by using non-phosphorus lipids in the membrane in response to phosphorus scarcity (Van Mooy et al., 2006, 2009), and others maximize DIP uptake efficiency through high affinity P transport systems (Scanlan and Wilson, 1999; Moore et al., 2005; Orchard et al., 2009). Yet, the most important mechanism to cope with DIP deficiency is to hydrolyze various forms of DOP found in the ocean through a battery of enzymes to yield phosphate (Dyhrman et al., 2009).
Alkaline phosphatase (AP) genes have been isolated and characterized for active sites to hydrolyze the various phosphorus esters (Luo et al., 2005; Nicholson et al., 2006; Huang et al., 2007; Duhamel et al., 2010, 2011). AP genes also have been isolated and characterized from various types of molecules (e.g., nucleotides, proteins, and alkaloids) and is most effective in an alkaline environment such as seawater (pH typically around 8). Marine bacterial AP has been widely studied and three different forms, Clade I, II, and III, have been recognized (e.g., Luo et al., 2009; Sebastian and Huerta, 2010), and a putative AP gene has been identified (Quisel et al., 1996; Moseley et al., 2006). However, it is believed that hydrolysis of phosphorus esters by alkaline phosphatase (AP) is the most common mechanism of DOP utilization (Labry et al., 2003; Nicholson et al., 2006; Huang et al., 2007; Duhamel et al., 2010, 2011).

Alkaline phosphatase (EC 3.1.3.1) removes phosphate groups from various types of molecules (e.g., nucleotides, proteins, and alkaloids) and is most effective in an alkaline environment such as seawater (pH typically around 8). Marine bacterial AP has been widely studied and three different forms, phoA, phoX, and phoD, have been recognized (e.g., Luo et al., 2009; Sebastian and Ammerman, 2009; White, 2009). These three forms are localized in different regions of the cell and require different metal ions for active sites to hydrolyze the various phosphorus esters (Luo et al., 2009). AP genes also have been isolated and characterized from some marine cyanobacteria (Ray et al., 1991; Orchard et al., 2009). In contrast, relatively little is known about AP genes in marine eukaryotic phytoplankton. An AP gene (putative phoX) was isolated from the green alga Volvox carteri which was a Ca$^{2+}$-dependent extracellular phosphatase responding to inorganic phosphate starvation (Hallmann, 1999). From the genome sequence of the freshwater chlorophyte Chlamydomonas reinhardtii, a putative phoX gene was identified which is likely to encode the previously characterized Ca$^{2+}$-dependent extracellular AP (Quisel et al., 1996; Moseley et al., 2006). An AP gene (ehap1) was isolated and characterized in the haptophyte Emiliania huxleyi (Xu et al., 2006). Putative AP has also been identified from the pelagophyte Aureococcus lagunensis (Sun et al., 2012). Gene models also predict the presence of AP genes in the genomes of diatoms Thalassiosira pseudonana and Phaeodactylum tricornutum (Armbrust et al., 2004; Bowlwar et al., 2008), especially a number of putative APs in T. pseudonana (Dyrhman et al., 2012). However, structural features and expression patterns of these putative AP genes remain to be characterized. There has been little evidence of DOP-utilizing proteins in dinoflagellates other than an AP-like protein found in Proorocentrum minimum (Dyrhman and Palenik, 1997) until recent detection of putative AP genes in two dinoflagellate species, Amphidinium carterae (Lin et al., 2011) and Karenia brevis (Morey et al., 2011; Lin et al., 2012).

While low similarity among the AP gene sequences from different organisms was noted, gene sequence divergence and phylogenetic relationships, particularly in relationship to the characterized bacterial APs, have not been examined. Further, in light of the diverse subcellular localizations of the different types of bacterial APs, whether the diverging AP gene sequences in eukaryotic phytoplankton might bear similar cytological and hence ecological significance remains to be explored. To better understand these questions, we isolated a putative AP full-length cDNA from a strain of Alexandrium, the latest-diverging dinoflagellate genus documented so far (John et al., 2003), and compared it with putative APs from the phylogenetically relatively basal and intermediate dinoflagellate species A. carterae and K. brevis (Zhang et al., 2007a), respectively, as well as AP-like sequences identified in other eukaryotic phytoplankton. Sequence variability, phylogenetic relationship, and putatively different subcellular localization of the AP sequences were analyzed.

**MATERIALS AND METHODS**

**ALGAL CULTURE AND ENZYME LABELED FLUORESCENCE LABELING**

Alexandrium strain ACHK-NT was kindly provided by the Collection Center of Marine Bacteria and Algae (CCMBA), Xiamen University, China. This strain was originally isolated from Hong Kong and identified as “A. catenella” morphologically. Based on the small subunit ribosomal RNA gene (SSU rDNA) isolated from this strain (unpublished) and phylogenetic analysis guided by information on the intragenomic SSU rDNA variation of “Alexandrium tamarenses complex” (Miranda et al., 2012), this strain belongs to Clade IIC, which is exclusively composed of Asian “A. catenella”. The culture was grown in K medium (Keller and Guillard, 1985; Keller et al., 1987) with 0.22 μm-filtered autoclaved seawater (28 PSU) at 20°C under a L:D = 14 h:10 h photocycle with a photon flux of 100 ± 10 μE m$^{-2}$ s$^{-1}$. One million cells were collected during exponential stage by centrifugation, resuspended in Trizol reagent (Invitrogen, Carlsbad, CA, USA) and stored at −80°C for subsequent RNA extraction.

The cells of three different dinoflagellate species (“A. catenella”, A. carterae, and K. brevis) under P limitation in the late stationary growth stage (DIP below detection limit of 0.2 μM in the medium) were collected by centrifugation and the cell pellets were then incubated in 1 mL 70% ethanol for 30 min. The samples were centrifuged and the supernatant was aspirated off. The pellets were incubated in 100 μL 0.22 μm-filtered sterile seawater with ELF®-97 phosphatase substrate (Invitrogen, Carlsbad, CA, USA) in a final concentration of 0.25 mM for 30 min in the dark. The samples were washed three times using sterile seawater and stored in 100 μL sterile seawater in the dark (González-Gil et al., 1998; Ou et al., 2010). The stable green fluorescent precipitates produced in situ by the AP activity and red autofluorescence of the live cells produced by chloroplast were both observed under Nikon 90i epifluorescence microscope (green fluorescence excited by UV, 340–380 nm; Nikon Corporation, Tokyo, Japan).

**RNA ISOLATION AND cDNA SYNTHESIS**

Total RNA was extracted as previously reported (Lin et al., 2011) and stored at −80°C. The 1st strand cDNA was synthesized from ~400 ng of the RNA using M-MLV reverse transcriptase (Promega, Madison, WI, USA) with a modified oligo-dT primer then purified using Zymo DNA Clean and Concentrator (Zymo Research, Orange, CA, USA).

**cDNA CLONING OF THE ALKALINE PHOSPHATASE GENE FRAGMENT AND RAPID AMPLIFICATION OF cDNA ENDS**

Based on the nucleotide (nt) and amino acid (aa) sequence alignments (see Figure 1) of A. carterae and K. brevis AP genes (Lin et al., 2011, 2012), degenerate primers were designed to obtain AP gene fragments from “A. catenella” under the following PCR

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FIGURE 1 | Comparison of AP conserved regions in dinoflagellates (A) and other eukaryotic phytoplankton (B). (A) Boxed are regions used to design the degenerate primer for amplifying AlecaAP fragment. (B) Example conserved domains (boxed) identified from alignment of amino acid sequences from *A. carterae* (ACAAP), *K. brevis* (KBRAP), *A. catenella* (AlecaAP), diatoms *Phaeodactylum tricornutum* (Phatr) and *Thalassiosira pseudonana* (Thaps), pelagophyte *Aureococcus anophagefferens* (Auran), and coccolithophorid *Emiliania huxleyi* (Emihu). Identical residues are black-shaded while chemically similar ones gray-shaded.

conditions: 94˚C for 1 min followed by 5 cycles of 95˚C for 15 s, 52˚C for 30 s, and 72˚C for 1 min, and 30 cycles of 95˚C for 15 s, 56˚C for 30 s, and 72˚C for 1 min. Two PCRs were carried out with the 1st PCR performed using forward primer DinoAPNF4a (5′-TTTGGWATGGGWGCMGCRGARCA-3′) paired with reverse primer DinoAPNR6 (5′-CTTRTCRTCRCCTTCRTYMGCKGT-3′). The PCR product was diluted 100 or 1000 fold and used as template for the 2nd (nested) PCR, in which DinoAPNR6 was paired with DinoAPNF4b (5′-GGTATGGGWGCMGCRGARCA-3′) as the primer set. The PCR product with expected size was purified by Zymo DNA Clean and Concentrator (Zymo Research, Orange, CA, USA) and directly sequenced (Invitrogen, Guangzhou, China).

Based on this partial sequence, specific primers were designed for both 5′- and 3′-rapid amplification of cDNA ends (RACE) to obtain the full-length cDNA by nested PCR. In the 5′-RACE, reverse primers AtaapR2 (5′-TGATGGACAAGACGAGCATGG-3′, outside) and AtaapR3 (5′-AGTTTACCTTGGCCGTGCTTAC-3′, inside) coupled with forward primer DinoSL (Zhang et al., 2007b) were used in nested PCRs. In the 3′-RACE, forward primers AtaaF3 (5′-GAGCAGTCCAAGGTATGTGAAC-3′, outside) and AtaaF4 (5′-CTGGACATTGCCAGTGGAC-3′, inside) were paired separately with modified oligo-dT primer in nested PCRs. The PCR products obtained from RACE were either directly sequenced, or cloned into a T-vector and several resultant plasmid clones were sequenced as reported (Lin et al., 2012). The sequences obtained were analyzed using Blastx against GenBank non-redundant protein sequences to confirm that the sequences are of the AP gene.

SEARCH OF GENBANK, JGI GENOME, AND CAMERA METAGENOMIC DATABASES FOR ALKALINE PHOSPHATASE GENES

To recruit as many eukaryotic putative AP genes as possible for phylogenetic analysis, AP sequences of *A. carterae* and *K. brevis* were used as queries using basic local alignment search tool (program tblastn) against GenBank database. We also used the keyword search with “alkaline phosphatase” against GenBank database to recruit any reported sequences which were not hit by tblastn
A set of programs were used to spatially characterize the deduced protein sequence. In addition, to find AP sequences from the natural plankton assemblages, AP aa sequences of the two dinoflagellates were used as query to tblastn against the Bermuda Atlantic Time-Series Study (BATS) and Hawaii Ocean Time-Series (HOT) 454-sequencing reads as well as Sargasso Sea open reading frame datasets in the Community Cyber infrastructure for Advanced Marine Microbial Ecology Research and Analysis (CAMERA) databank. E-value cutoff was set at 1e-3, and 100 hits were set as the target for retrieval. A previous study showed that BATS water is DIP-poor whereas HOT water contains DIP concentrations which stoichiometrically fit the Redfield ratio (Wu et al., 2000).

PHYLOGENETIC ANALYSIS

To examine the relationship between “A. cate
tenella” AP and counterparts from other organisms, deduced aa sequence of the “A. caten
tella” full-length cDNA was aligned with AP sequences from the representative organisms using ClustalX (Thompson et al., 1997). The environmental sequences retrieved from CAMERA were too short and therefore were not included in the analysis. Sequences of different forms of APs (phoA, phoD, phoX, phoV) from cyanobacteria, a group of organisms whose APs have been relatively well studied, were collected and compared with putative dinoflagellate APs (Dino-APs) separately. The phylogenetic analyses were done only to sequences belonging to the same type (phoX) because no reliable alignment could be achieved for a mixed-type dataset. Alignment was performed for each of these datasets using ClustalW1 with default setting (Gapopen 15, Gapext 6.66, Gapdist 8, Maxdiv 40). The alignments were then inspected and corrected manually. Neighbor-joining (NJ) analysis (Saitou and Nei, 1987) was run with 1000 bootstraps, and Bayesian analysis (Huelsenbeck and Ronquist, 2001) was run with model jumping setting for 725,000 generations (two independent runs were performed and the final standard deviation of split frequencies was 0.01), with trees sampled every 100 cycles and the first 25,000 cycles discarded (burn-in).

RESULTS

CONSERVATION OF ALKALINE PHOSPHATASES IN DINOFLAGELLATES

After assembling the initial partial AP sequence with the 5′ and 3′ RACE sequences, a full-length cDNA sequence was obtained from “A. catenella,” with an open reading frame of 2,181 bp (GenBank Accession No. JQ806383). Its deduced aa sequence (named as AlcA) was used to search against the GenBank protein database and the top two hits were A. carterae (with Expect value [E-value] to be 0.0) and K. brevis (E-value 2e-155), verifying that this is an AP in bacteria and a homolog from E. huxleyi (EHAP1, GenBank accession number AB151308) that has been shown to be extracellular (Xu et al., 2006). Included in the test was also AP from humans, whose cell surface distribution has been shown (Cutler et al., 1974). For dinoflagellates, a total of 98 protein sequences were tested, including representative proteins associated with cytoskeleton (actin), light harvesting in the chloroplast (peridinin-chlorophyll a-binding protein or PCP) and cell membrane (rhodopsin), carbon fixation in the chloroplast (ribulose-1, 5-bisphosphate carboxylase/oxygenase or Rubisco), and the cell cycle related protein proliferating cell nuclear antigen (PCNA), a well-characterized primarily nuclear protein. No model gave 100% correct prediction and no one failed 100% either, and in most cases predictions shared by two or more of these four models were correct (Table S1 in Supplementary Material). Therefore, we decided to use all four models so that the reliability of the predictions could be assessed, with the likelihood of the predicted AP localization ranked according to the number of the models that supported the prediction.

1 http://clustalw.ddbj.nig.ac.jp/
2 http://www.ch.embnet.org/software/TMPRED_form.html

COMPUTATIONAL ANALYSIS OF MAJOR STRUCTURAL FEATURES AND PREDICTION OF ALKALINE PHOSPHATASE SUBCELLULAR LOCALIZATION

A set of programs were used to spatially characterize the deduced protein. The signal peptide was determined by TargetP V1.1 (Emanuelsson et al., 2000) and SignalP V4.0 (Petersen et al., 2011). GPI-SOM (Fankhauser and Mäser, 2005) and PredGPI (Pierlenoi et al., 2008) were used to locate the glycosylphosphatidylinositol (GPI) anchor in the deduced protein sequence. TMpred2 was used to predict transmembrane domains.

Four existing bioinformatics models were adopted to predict subcellular localizations of AP in eukaryotic phytoplankton. Interpretable Subcellular Localization Prediction (YLoc; Briesemeister et al., 2010) predicts localization of a protein among nine possible subcompartments of the cell: extracellular space, plasma membrane, cytoplasm, mitochondrion, chloroplast, nucleus, Golgi apparatus, endoplasmic reticulum, and peroxisome. The YLoc- algorithm in the package was used with plant selected as the target organism. The second program was Euk-mPLoc2.0 (Chou and Shen, 2010), which was designed specifically for predicting subcellular localization of eukaryotic proteins in 22 different compartments (Chou and Shen, 2010), including the nine in YLoc. The other two programs were WoLF PSORT (Horton et al., 2007) and CELLO (Yu et al., 2006). These programs have been used in various studies, such as functional annotation in C. reinhardtii and the subcellular localization prediction of the secretome for fungi (Soanes et al., 2007; Ghamsari et al., 2011; Rai et al., 2012). Because it has not been verified that these models suit APs or other proteins in eukaryotic phytoplankton. We first ran a set of proteins whose locations of functions are known to test the accuracy of the predictions. For AP, we tested a set of APs from the chlorophyte algae V. carteri and C. reinhardtii that are known to be related to phoX, an extracellular AP in bacteria and a homolog from E. huxleyi (EHAP1, GenBank accession number AB151308) that has been shown to be extracellular (Xu et al., 2006). Included in the test was also AP from humans, whose cell surface distribution has been shown (Cutler et al., 1974). For dinoflagellates, a total of 98 protein sequences were tested, including representative proteins associated with cytoskeleton (actin), light harvesting in the chloroplast (peridinin-chlorophyll a-binding protein or PCP) and cell membrane (rhodopsin), carbon fixation in the chloroplast (ribulose-1, 5-bisphosphate carboxylase/oxygenase or Rubisco), and the cell cycle related protein proliferating cell nuclear antigen (PCNA), a well-characterized primarily nuclear protein. No model gave 100% correct prediction and no one failed 100% either, and in most cases predictions shared by two or more of these four models were correct (Table S1 in Supplementary Material). Therefore, we decided to use all four models so that the reliability of the predictions could be assessed, with the likelihood of the predicted AP localization ranked according to the number of the models that supported the prediction.
Table 1 | Comparison of major features of AlecaAP, ACAAP, and KBRAP:

| Gene (protein) | cDNA (bp) | Amino acid | pH | Mw (kDa) | G+C% | Signal peptide* | GPI anchor |
|---------------|-----------|------------|----|----------|------|----------------|------------|
| acaap (ACAAP) | 2115      | 704        | 4.31 | 75       | 54   | +, 0.784        | +          |
| kbrap (KBRAP) | 2061      | 686        | 4.66 | 72.9     | 49   | +, 0.903        | +          |
| aleca-ap (AlecaAP) | 2181 | 726 | 4.48 | 76.6 | 61 | +, 0.770 | + |

*“+” represent “signal peptide = yes” predicted by SignalP V 4.0 (left); “numbers” represent the score predicted by TargetP 1.1.

Acid residues (6.7 and 5.4%, respectively). A signal peptide was detected with a probability of 0.770 by TargetP 1.1 and confirmed by SignalP V4.0. The GPI anchor located at C-terminus (residues 721–726), which ensures AP to attach to cell membrane, was predicted by GPI-SOM and further confirmed by PredGPI (Table 1).

High divergence of alkaline phosphatase amino acid sequences revealed by phylogenetic analyses

Alkaline phosphatase aa sequences are highly variable among the eukaryotic phytoplankton. No remarkable similarity was observed between the Dino-APs and other well-characterized eukaryotic APs such as EHAP1 from E. huxleyi (Landry et al., 2006; Xu et al., 2006, 2010) or phoX from V. carteri and C. reinhardtii (Hallmann, 1999; Moseley et al., 2006). Nevertheless, tblastn of Dino-APs against E. huxleyi CCM1516 genome data (IGI) produced several hits with E-value lower than 1e-31. As predicted using TMpred, both the putative Dino-APs and other phytoplankton APs examined all contained 3–7 transmembrane domains. Interestingly, Dino-APs showed 15–30% of aa similarity with the APs from cyanobacteria, including a characterized atypical phoA gene (C-terminal half of the gene) from Synechococcus sp. PCC7942 (Ray et al., 1991); the phoA gene with GenBank accession number AAA27331. Further sequence comparison showed that V. carteri, C. reinhardtii also possessed phoA-like AP genes in addition to the phoX type mentioned above. The putative phoA-like AP sequences from P. tricornutum, T. pseudonana, V. carteri, C. reinhardtii, Aureococcus anophagefferens, cyanobacteria and bacteria share 10–16 conserved domains with Dino-APs (Figure 1B). The phylogenetic tree of phoA-like AP from different organisms showed that all of the eukaryotic phytoplankton APs clustered together, in sister relation to APs from marine bacteria (Figure 2).

Alkaline phosphatase gene sequences from environmental samples

No putative AP sequences were found in the Sargasso Sea ORF, likely due to its relatively small database size. Five significant hits were obtained from HOT and 15 from BATS sharing several conserved regions with Dino-APs. When these retrieved sequences were blasted against GenBank database using blastp, several of them matched bacterial AP as their top hit, giving two HOTs and five BATS sequences as eukaryotic AP as they best matched APs of dinoflagellates or diatoms (Table 2). Given that the metagenomic data were obtained from microbial plankton, the AP genes detected were probably restricted to pico- or nano-plankton. This might explain why so few sequences of AP genes were found and partially why none were found from the Sargasso Sea ORF dataset.

Computationally predicted different subcellular localizations of alkaline phosphatases in dinoflagellates and other eukaryotic phytoplankton

The test runs using protein sequences from various organisms showed that most of the proteins were predicted correctly when the prediction was supported by two or more of the four models used. APs in humans were predicted to be extracellular by YLoc+, WoLF PSORT, and CELLO and cell membrane by Euk-mPLoc, consistent to observed extracellular space distribution of this protein (Cutler et al., 1974). Like phoX in bacteria, the phoX-like APs in chlorophyte C. reinhardtii and V. carteri have been shown to be extracellular (Hallmann, 1999; Moseley et al., 2006); both of them were predicted to be extracellular by two of the models (Table S1 in Supplementary Material). Among dinoflagellate proteins, actin was uniformly predicted to be cytoskeleton by Euk-mPLoc, and to be chloroplastic by YLoc+ and CELLO, the two models that do not have the cytoskeleton category. PCP was correctly and uniformly predicted to be chloroplastic by all four models. Rhodopsin, a known transmembrane protein, was predicted as a plasma membrane protein in all the 19 cases by Euk-mPLoc and CELLO and 11 of these are supported by three or more of the models. All 16 Rubisco sequences, from various species of dinoflagellates, were predicted as chloroplast proteins by three or more models. All 16 PCNA genes examined were predicted to be nuclear proteins, and 5 of these were supported by three or more models. All these results confirmed that the joint-multiple prediction models could be applied to dinoflagellates and other phytoplankton proteins by the majority rule (≥2 models). Predicting results based on this rule are shown in Table 3. Putative AP in “A. catenella” would be localized in the extracellular space (2 models) and chloroplast (2 models), with equal probability. Putative AP in A. carterae would most likely be in the extracellular space (plus endoplasmic reticulum, which can be considered as transporting and secreting out of the cell; 3 models) but cytoplasm (1 model) and chloroplast (1 model) are possible as well. In K. brevis, putative AP was predicted to be most likely extracellular (also plus Endoplasmic Reticulum; 4 models) or cell surface (cell wall plus membrane; 2 models) with a small probability to be found in other intracellular compartments (1 model each).

These computational predictions of putative Dino-AP localization were verified by enzyme labeled fluorescence (ELF) labeling. In the three dinoflagellate species we have analyzed so far, AP activity as indicated by ELF showed different localization patterns (Figure 3). In “A. catenella,” AP appeared to be located both intracellularly and on the cell surface. In A. carterae, AP activity was more on the cell surface than intracellular but both locations had substantial activities. In K. brevis, AP activity was mostly...

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FIGURE 2 | Phylogenetic tree inferred from AP amino acid sequences of eukaryotic phytoplankton, marine bacteria, and cyanobacteria. Shown is neighbor-joining (NJ) tree; Bayesian analysis (BE) tree has similar tree topology albeit with more uneven branch lengths; support of nodes > 70% in NJ bootstrap values and >0.70 in Bayesian posterior probability is shown. Color backgrounds indicate the type of organisms which the AP sequence is from. For instance, red background indicates sequences from species whose chloroplasts belong to chromalveolates (“Red-type”), green background indicates sequences from green algae, blue background indicates sequences from cyanobacteria.

Among other groups of eukaryotic phytoplankton putative AP was predicted to be in the extracellular space and plasma membrane of *P. tricornutum* and most likely in the cytoplasm of *T. pseudonana* and *A. anophagefferens*. In *E. huxleyi*, the *phaA*-like AP (gi|Emihu1|444279|estExtDG_fgenesh_newKGskg.C_3060016:21-652) was more likely extracellular but could be cytoplasmic as well whereas EHAP1 (ABI51308) was predicted to be extracellular and cell surface (plasma membrane; Table 3). EHAP1 has been experimentally shown to be extracellular (Xu et al., 2006).

**DISCUSSION**

Alkaline phosphatase seems to be common in both prokaryotic and eukaryotic phytoplankton. AP genes have been studied extensively in bacteria and scarcely in eukaryotes, while AP activity has been detected in a wide range of eukaryotic phytoplankton.
Table 2 | BLASTP result of CAMERA datasets.

| Sampling site | Clone ID | Hit organisms* | E-value |
|---------------|----------|---------------|---------|
| HOTS          | HF_READ_03392351/accession=HF_READ_03392351 | Thaps, Ampca, Phatr | −9 to −13 |
| HOTS          | HF_READ_03392351/accession=HF_READ_03392351 | Thaps, Phatr, Ampca | −9 to −14 |
| BATS          | BATS_READ_00574637/accession=BATS_READ_00574637 | Ampca, Thaps | −19 to −26 |
| BATS          | BATS_READ_00574637/accession=BATS_READ_00574637 | Ampca, Thaps | −19 to −26 |
| BATS          | BATS_READ_02761309/accession=BATS_READ_02761309 | Thaps, Ampca | −4 to −8 |
| BATS          | BATS_READ_00987788/accession=BATS_READ_00987788 | Ampca | −6 |
| BATS          | BATS_READ_00987788/accession=BATS_READ_00987788 | Ampca | −6 |

*Ampca: Amphidinium carterae (dinoflagellate); Phatr: Phaeodactylum tricornutum (diatom); Thalassiosira pseudonana (diatom).

(Table S2 in Supplementary Material). The combination of the new molecular data retrieved in this study with literature information on AP activities indicates that AP is an ancient and common feature in marine microbes, attesting to the notion that phosphorus is a common limiting factor of phytoplankton production. The association of AP activity in eukaryotic phytoplankton with DIP limitation in the environment has been widely established. Recently, AP gene expression in dinoflagellates was also found to be regulated by DIP availability (Lin et al., 2011, 2012). To date, AP activity has not been detected in raphidophytes, and the AP gene has not been identified in the genomes of rhodophytes, such as *Synechococcus* spp. confirmed that there are two putative APs, one sharing a conserved domain with the characterized *phoX* from marine γ-proteobacteria, and the other sharing 52% identity to 50% of the translated *phoA* gene from *Synechococcus* PCC7942 (Orchard et al., 2009).

Among eukaryotic microalgae, only APs from chlorophyte *C. reinhardtii* and *V. carteri* have been categorized with respect to the types recognized in bacteria; *phoX* like sequences occur in both species (Hallmann, 1999; Moseley et al., 2006). The transcriptomic and proteomic study indicated that there are several putative APs in *T. pseudonana* (Dyrhman et al., 2012). Before the analysis of the AP genes from dinoflagellates reported here, the only characterized AP coding gene in marine eukaryotic phytoplankton is *ehap1*, identified in *E. huxleyi* (Xu et al., 2006). The gene product EHAP1, with its signal peptide identified at the N-terminus and without a GPI anchor at the C-terminus, can be released from the cell surface into the medium. But it does not show any similarity to the known types of APs. Because of the very limited knowledge of eukaryotic phytoplankton AP genes, it is difficult to categorize the putative APs from dinoflagellates and other eukaryotic phytoplankton into those recognized types of marine APs. However, our sequence comparison results showed that Dino-APs and other putative AP sequences in other eukaryotic phytoplankton genomes retrieved from the public databases share highest (despite still very low) identifiable sequence similarity with the *phoA* AP from bacteria and cyanobacteria among all the recognized types of AP. These AP amino acid sequences clustered together in the phylogenetic tree (Figure 2). Regardless if these are indeed *phoA* type APs, all our analyses suggest that they are undoubtedly a subset of putative APs.

It is interesting to observe that while cyanobacteria, chlorophytes *C. reinhardtii* and *V. carteri*, and the haptophyte *E. huxleyi* have multiple types of AP, essentially only one form of putative AP sequence has been identified in dinoflagellates. There is little sequence variation in the coding region, most of which involves synonymous nt substitutions (Lin et al., 2011, 2012). In the 2,181-bp long AP cDNA coding region in *A. catenella*, only two sites...
Table 3 | Predicted subcellular localization for alkaline phosphatase in eukaryotic phytoplankton using Yloc, Euk-mPLoc2.0, WoLF PSORT, and CELLO computational models.

|                | Extracellular space | Cell wall | Plasma membrane | Cytoplasm | Nucleus | Mitochondrion | Chloroplast | Golgi apparatus | Endoplasmic reticulum | Peroxisome | Vacuole |
|----------------|---------------------|-----------|-----------------|-----------|---------|--------------|-------------|-----------------|------------------------|-------------|---------|
| **PhoA-like**  |                     |           |                 |           |         |              |             |                 |                         |             |         |
| "Alexandrium catenella" JQ806383 | 2 | 2 | 1 | 1 | 2 | 2 | 1 | 1 | 1 | 1 | 1 | 1 |
| Amphidinium carterae ADT91623 | 3 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Karenia brevis JQ67049 | 3 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Phaeodactylum tricornutum XP_002182197.1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Thalassiosira pseudonana | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| jgi[Thaps3]1179|ggi[Thaps3]1179|fgenesh1_pg_C_chr_1000313:22-645 | 3 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Aureococcus anophagefferens | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| jgi[Auran1]scaffold_13:864419-866137_putative | 3 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Emiliania huxleyi | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| jgi[Emihu1]444279|estExtDG_fgenesh_newKGs_kg.C_3060016:21-652 | 3 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Chlamydomonas reinhardtii Chire_XP_001702318.1 | 2 | 2 | 2 | 2 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Volvox carteri f. nagariensis Volca_XP_002958226.1 | 2 | 2 | 2 | 2 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| **PhoX-like**  |                     |           |                 |           |         |              |             |                 |                         |             |         |
| Chlamydomonas reinhardtii XP_001703098* | 2 | 2 | 2 | 2 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Volvox carteri CAA10030* | 2 | 2 | 2 | 2 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| **Other types** |                     |           |                 |           |         |              |             |                 |                         |             |         |
| Emiliania huxleyi/ ABI51308* | 2 | 2 | 2 | 2 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |

Shade darkness denotes the number of models that support the prediction, from highest (darkest = 3) to lowest (lightest = 1).

*APs that have been previously characterized. The rest of the sequences (except Dino-APs) are tblastn hits using Dino-APs as query.
show changes. This is striking because most of the genes in dinoflagellates that have been studied exist in numerous variations. For instance, *pcna*, which usually occurs in one copy in other eukaryotes, exists in tens of copies in the dinoflagellate *Pfiesteria piscicida*, which contain many synonymous and substantial numbers of non-synonymous nucleotide substitutions (Zhang et al., 2006). Similarly, Rubisco genes in *P. minimum* occur in over 100 copies, which differ from each other by as much as 5%, with both synonymous and non-synonymous changes as well as insertions/deletions (Zhang and Lin, 2003). If the duplication of genes in dinoflagellates has resulted from whole genome duplication, as recently postulated (Hou and Lin, 2009), the negligible level of variation in AP relative to that of other genes may suggest that Dino-APs probably face strong evolutionary purifying selection (Kim et al., 2011). However, if dinoflagellate genes duplicated independently, then the low variability of the gene can simply indicate that the gene copies have only arisen from recent duplication events (Thornton, 2007 and the refs there in).

**HIGH ALKALINE PHOSPHATASE SEQUENCE VARIABILITY ACROSS LINEAGES**

The sequence variability of putative AP between lineages is strikingly high. Although it has been shown that 25–30% of the aa residues, especially the active sites, in APs are well conserved among mammals, yeast, and *E. coli* (Murphy et al., 1995), this does not seem to be the case for dinoflagellates and other eukaryotic phytoplankton. Putative Dino-APs did not share obvious similarity with APs from other eukaryotes. There were also no obvious conserved domains observed between putative Dino-APs and *phoV* from *S. elongatus* PCC 7942. When comparing putative Dino-APs with *phoA* from *Trichodesmium* spp. (Orchard et al., 2009) and *Synechococcus* PCC7942 (Ray et al., 1991), more than 10 conserved domains were identified between putative Dino-APs and *phoA*; however, no conserved region was found between putative Dino-APs and *phoX*. The lack of long conserved sequence regions and the overall high variability rendered the alignment of nt sequences of counterparts from different lineages (e.g.,
AP activity in A. lagunensis (Figure 3) to import bulky DOP into the cell. In contrast, APs located inside the cell (i.e., chromalveolate), which as a whole is sister to APs from other bacteria whose chloroplasts belong to the red-type (Lin et al., 2008, 2010). The computational prediction of chloroplast and cell surface localization was suggestive of AP being localized both intracellularly and on the cell surface in most of the species examined (e.g., P. micans, "A. furdyeense," and "A. catenella; Ou et al., 2010). Our ELF labeling patterns were indicative of AP being localized both intracellularly and on the cell surface of "A. catenella," which is consistent with the computational prediction of chloroplast and cell surface localization. Chloroplast in "A. catenella" was highly branched and abundant in the cytoplasm (Hansen and Moestrip, 1998), rendering chloroplast AP ELF labeling as scattered in the cytoplasm (Figure 3). Comparative studies have shown that the diatom S. costatum quickly released the AP into the medium while "A. catenella" and P. donghaiense harbor most of the AP inside the cells and on the cell surface, respectively, so that these algae could use DOP in the ambient water and survive longer under P-limited conditions (Lin et al., 2011). In contrast, K. brevis growth rate declined shortly after DIP was depleted and detectable AP activity increased long after (Lin et al., 2012). Perhaps the more cytoplasmic presence of AP enables A. carterae to access DOP generated inside the cell quickly compared to K. brevis, in which the dominant extracellular/cell surface distribution of AP may force the cells to rely on external DOP generated from cell death and on the import process for the cleaved DIP to enter the cell. Yet, such linkage between AP sequence, subcellular localization, and P nutritional strategy requires much more dedicated studies to verify.

CONCLUSION
Alkaline phosphatase appears to be common in eukaryotic phytoplankton, thus likely an ancient feature, but how many lineages have lost this feature and consequently how they cope with potential phosphate limitation warrants further studies. The high diversity of AP genes in phytoplankton may be related to diversification of subcellular localization and strategies of accessing and utilizing DOP. Currently sequence data are still too limited to infer a definitive evolutionary trend of the gene and functional diversification of the encoded enzyme. Much work is needed to: (1) determine the origin of the AP gene in eukaryotic phytoplankton; (2) elucidate what drives the rapid evolution of this gene in comparison to other genes in the same organism; (3) examine the linkage of the highly variable AP sequences with subcellular localizations and access of the host algae to diverse sources of DOP inside and outside the cells. In addition, on the practical side, the high diversity probably will make the AP gene a promising phylogenetic marker for resolving closely related lineages that cannot be distinguished based on morphology or conserved genes such as those coding for rRNA. Yet convincing evidence ratifying such utility of the AP gene also awaits further studies with broader taxon sampling and more in-depth phylogenetic analyses.

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SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at http://www.frontiersin.org/Aquatic_Microbiology/10.3389/fmich.2012.00235/abstract
Table S1 | Test runs of the four subcellular localization predicting models with AP from dinoflagellates and various other organisms and a variety of other proteins in dinoflagellates.

Table S2 | Eukaryotic phytoplankton in which Alkaline phosphatase activity or coding gene that has been detected.

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Table S1 | Test runs of the four subcellular localization predicting models with AP from dinoflagellates and various other organisms and a variety of other proteins in dinoflagellates.

| Model | Accuracy | Sensitivity | Specificity |
|-------|----------|-------------|-------------|
| 1 | 0.85 | 0.90 | 0.80 |
| 2 | 0.88 | 0.92 | 0.84 |
| 3 | 0.90 | 0.95 | 0.88 |
| 4 | 0.92 | 0.97 | 0.90 |

Table S2 | Eukaryotic phytoplankton in which Alkaline phosphatase activity or coding gene that has been detected.

| Phytoplankton | Alkaline Phosphatase Activity/Coding Gene |
|---------------|----------------------------------------|
| dinoflagellates | yes |
| other eukaryotes | yes |

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