Cadmium and phosphorus cycling in the water column of the South China Sea: The roles of biotic and abiotic particles

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1. Introduction

Cadmium, considered a toxic metal for terrestrial organisms, exhibits nutrient type distribution in the ocean (Boyle et al., 1976). Similar to major nutrients like nitrate and phosphate, internal cycling of cadmium in the ocean is controlled by biological activity, depleted by algal uptake in the euphotic zone and regenerated by bacteria mostly in surface waters (Boyle et al., 1976; Martin et al., 1976; Bruland, 1980; Knauer and Martin, 1981). In addition to its vertical distribution, the horizontally segregation of Cd concentrations with increasing water age clearly shows that its horizontal cycling in the ocean also behaves like major nutrients (Bruland and Lohan, 2004). Cadmium appears to be the most eligible trace metal to join the Redfield ratio (Bruland et al., 1991; Kuss and Kremling, 1999; Ho et al., 2003; Ho, 2006). Relatively, phosphorus, a well-known essential major nutrient mainly used for reproduction and energy transfer in all organisms, has been traditionally regarded as a long term limiting nutrient (Redfield, 1958; Tyrrell, 1999). Its importance on limiting phytoplankton growth in the euphotic zone and its transformation processes in marine water columns have thus been relatively ignored...
(Benitez-Nelson, 2000). More recent studies have gradually revealed the importance and complexity of phosphorus in controlling algal growth and community structure in the ocean (e.g., Wu et al., 2000; Sañudo-Wilhelmy et al., 2001; Dyhrman et al., 2006).

With the highly correlated relationship between Cd and P in seawater, Cd/Ca ratios in the hard parts of benthic foraminifera have been used as proxies to reconstruct P concentrations in past oceanic environments (Boyle, 1988). Cd/P ratios in deep water, theoretically, are determined by biological fractionation of plankton in oceanic surface waters over long time scales. However, culture studies have revealed complicated interactions between phytoplankton and bio-active trace metals that may influence the correlation and cycling of Cd and P in marine water columns, especially in oceanic surface waters. For example, Cd/P ratios in phytoplankton may be affected by physiological state of microalgae, phytoplankton phylum, the relative concentrations of Cd and P in seawater, or even the relative concentrations of other trace metals. Laboratory studies have revealed that Cd may be used as an alternative nutrient to replace Zn in carbonic anhydrase in marine diatoms when Zn is not available (Lane and Morel, 2000; Lane et al., 2005). Under low Zn and low pH conditions, both laboratory and field studies have found that diatoms may take up more Cd (Lane and Morel, 2000; Price and Morel, 1990; Cullen and Sherrell, 2005). Culture and field studies also show that different algal species can have very different Cd/P quotas (Ho et al., 2003, 2004; Abe, 2005; Finkel et al., 2007) and resistance to Cd toxicity (Payne and Price, 1999). Coccolithophore and dinoflagellates with cysts tend to have relatively high Cd quotas. Phytoplankton’s community structure becomes an important factor in influencing Cd cycling and the Cd/P ratio both in the surface oceans and deep waters. Algal Cd quota would also be elevated in regions with relatively high Cd concentrations, such as upwelling regions (Lösch et al., 1998; Sunda and Huntsman, 2000; Cullen and Sherrell, 2005). The interactive relationship among Cd, Zn, Mn, and Fe can be important in deciding algal uptake of Cd (Sunda and Huntsman, 1996, 1998, 2000; Cullen and Sherrell, 2005; Cullen, 2006). These results imply that the Cd/P ratios in deep waters might change in a long term when conditions affecting the Cd/P ratios of biogenic particles in oceanic surface water change.

Although particles play a vital role in controlling trace metal distribution and cycling in the ocean (Turekian, 1977; Li, 1991), the role of particles on Cd and P cycling in marine water columns have been relatively ignored. Most previous studies on Cd and P have emphasized their correlation in dissolved form. In terms of vertical cycling, large biogenic particles, mostly composed of sizable plankton such as large diatoms, zooplankton and their hard parts, detritus, and fecal pellets, serve as the dominant components of sinking particles and thus are principal agents in transporting both major and trace elements from oceanic surface waters to deep waters (Fowler and Knauer, 1986; Allredge and Jackson, 1995). Since the cycling of Cd and P is driven and transformed by biogenic particles generated in the eutrophic zones (Martin and Broenkow, 1975; Bruland et al., 1978; Martin and Knauer, 1973), elemental concentrations in biogenic particles in oceanic surface waters not only reflect the interaction between plankton and seawater, but also provide fundamental information to study and model their vertical cycling processes in marine water columns (Martin and Knauer, 1973; Collier and Edmond, 1984; Bruland et al., 1991; Ho et al., 2003, 2007). In this study, we focus on determining the concentrations of Cd, P, and Al in particles, including phytoplankton and zooplankton in the surface water, and the sinking particles collected by sediment traps at three depths (120, 600, and 3500 m) in the northern South China Sea. Aluminum is used to evaluate the lithogenic source of trace metals in suspended particulate material and plankton samples (Ho et al., 2007). We especially examine the elemental composition of plankton larger than 10 μm due to their importance in vertical transport of elements. We also determined the vertical fluxes of the two elements through sinking particles to elucidate their temporal and vertical cycling features and their relationship with biogenic and lithogenic particles. The published data of the dissolved concentrations of Cd and P (Wen et al., 2006) will be integrated with the data from this study to elucidate the roles of biotic and abiotic particles on the cycling of Cd and P and their residence time in different zones of the water column.

2. Methods

2.1. Sampling site

The sampling site, the South-East-Asian-Time-Series (SEATS), was located in the northern South China Sea basin (bottom depth 3783 m) (Fig. 1). The northern South China Sea annually experiences the

![Fig. 1. Location of the sampling site in the South China Sea.](image-url)
northeastern monsoon in winter and southwestern monsoon in summer. Particles from biomass burning and anthropogenic materials from eastern and southern China and Southern Asia are brought to the region periodically (Lin et al., 2007). Our previous study revealed that the composition of many other bioactive trace metals in the size-fractionated plankton collected in the surface waters of the SEATS site were from extracellular inorganic particles adsorbed on cells (Ho et al., 2007). Several lines of evidences suggest that the extracellular metals were originally derived from anthropogenic aerosols containing highly soluble trace metals (Ho et al., 2007). Occasionally, the winter monsoon may also induce dust storms originating from the deserts of central China and input lithogenic particles into the surface water. This semi-enclosed deep water basin also lies close to regions with high fluvial input from the shelf, mainly from the Pearl River and Mekong River. Lateral transport of lithogenic particles from the continental shelf and break could be significant.

2.2. Sampling of size-fractionated plankton

The plankton samples were collected aboard two respective cruises during early spring (March 21 to 30, 2002) and fall (October 19 to 20, 2006). Plankton samples were collected using 20 L Teflon-coated Go Flo bottles (General Oceanics) mounted on a Teflon-coated rosette equipped with a modified epoxy coated Seabird CTD and Kevlar wire. Sampling of the size-fractionated plankton was especially focused at chlorophyll maximum depths. Six replicate casts were carried out at the chlorophyll maximum within 24 h. For the oligotrophic waters like the South China Sea, where the highest total SPM is less than 0.5 mg/L in the surface water, filtering large volumes of seawater is required to obtain a sufficiently large amount of plankton biomass for trace metal analysis. We used a novel trace metal clean filtration device to gently filter 120–200 L of seawater and collect diverse sized plankton through two gravity filtration steps (Ho et al., 2007). Step one involved concentrating the plankton in the sampling bottles by guiding the seawater into the filtration apparatus. The gravity filtration plankton sampler is made of polycarbonate material and equipped with 150-, 60-, and 10-μm aperture changeable Nitex nets and three 100-μL polyethylene sampling bottles in sequence, and acid-washed before each use. The design is shown in Ho et al. (2007). Before drawing seawater, the water spigots of six Go Flo bottles on the elevated rosette were first connected to the acid-washed C-flex tubes. The other ends of the C-flex tubes were then connected to the inflow ends of the size-fractionated filtration apparatus. The seawater in the bottles flowed through the six directional PTFE inlets and the water level was raised over the top bottle in the device, indicating that the intact size-fractionated biogenic particles had been gently separated through the nets; the particles then sank into the 100-μL polyethylene bottles fixed at the bottom of the nets. The gentleness of the filtration process was witnessed by the live plankton observed in the sampling bottles after collection. Before processing the second filtration, the nets were sprayed with trace metal-clean filtered seawater to rinse out any plankton stuck on the inner surface of the nets. The concentrated plankton samples collected in the 100-μL polyethylene bottles were then filtered again through an acid-washed filtration apparatus with pre-weighted acid-washed 47-mm 10-μm pore size polycarbonate filters. The samples in the two larger fractions were then passed through the filters in less than a minute under gravity or very low vacuum. The biogenic particles on the filters were quickly rinsed with Milli-Q water and collect two larger fractions of plankton. Any other swimmers, which passed through the filters, were then passed through 1 mm nylon mesh to remove swimmers and large zooplankton. After being freeze-dried and weighed, the filters with the plankton particles were digested in 1 mL 8 N super pure nitric acid (Seastar) in 20 mL Teflon vials at 120 °C on a hot plate for 24 h (Ho et al., 2003). After diluting the digested solution to 10 mL with Milli-Q water and removing the filter, the solution was centrifuged to remove any particulate residues. In terms of sinking particles, the supernatant in each cup was first removed by plastic syringe and stored in a refrigerator for further processing. Particle samples were divided into six portions using a perimatic peristaltic pump dispenser (Jencons Ltd.). The subsamples were then passed through 1 mm nylon mesh to remove swimmers and large zooplankton. Any other swimmers, which passed through the sieve, were manually picked. A small portion of well-mixed particle samples were sampled and filtered with 0.2-μm acid-washed polycarbonate filter under vacuum. The samples on the filters were quickly rinsed with Milli-Q water a couple of times to remove sea salts. After being freeze-dried and weighed, the filters were soaked in a mixture of 0.5-mL super pure nitric acid and 0.1-mL super pure hydrofluoric acid (Seastar) in 10-mL microcentrifuge Teflon vials. The filters with particles and the blank filters were digested in a microwave oven (MARS, CEM) at 180 °C for 15 min with a 10-minute ramp from room temperature to 180 °C. After complete digestion, the samples were evaporated to be almost dry on hot plates at 80 °C under a trace metal-clean hood. The dried samples were added with super pure nitric acid and diluted sequentially with Milli-Q water to obtain samples in 3% nitric acid solution, which were ready for ICPMS analysis.

Due to consideration given for determining Hg species by other researchers in the sediment trap study, saline solution without Hg was used as the trap cup solution. It should be noted that even with preservative added, significant amounts of Cd and P in particles can become soluble from sinking particles during long term storage in sediment traps (Knauer and Martin, 1981). It is thus essential to determine and include the soluble portion of P and Cd in the supernatant to obtain accurate total particle fluxes (Knauer and Martin, 2003).
We observed significant amounts of P and Cd existing in the supernatant solution, particularly in the samples from shallow depths. Supernatant samples were first diluted 100-fold and treated with 1200 W UV radiation for 12 h to decompose soluble organic matter in the diluted supernatant. Phosphorus concentrations in the diluted supernatant solution were analyzed by both a spectrophotometer and ICPMS. Using the corresponding concentrations obtained from the two methods as the data for x and y axes, the slope of the correlation line was 1.1, showing good accuracy for P analysis in the supernatant. The concentrations of Al, P, and Cd in the various particle samples and the concentrations of Cd in the diluted supernatant solution were determined by using a sector field high resolution ICPMS (Element 2 or Element XR, Thermo Scientific), fitted with a desolvation system (ESI). The analysis was conducted with sensitivity higher than $10^6$ counts per second for 10 ppb In under medium resolution. The sensitivity and stability of the instrument were adjusted to optimal conditions before analysis. External and internal standards were both applied for concentration quantification. The details of the precision, accuracy, and detection limit of the method were described in Ho et al. (2003, 2007).

3. Results and discussions

3.1. Cd, P, and Al in SPM and plankton

Aluminum and phosphorus, with their contrasting known abundance in both lithogenic and biogenic particles correspondingly, are used to evaluate the relative biotic and abiotic sources for trace metals in SPM and plankton samples (Ho et al., 2007). By assuming constant trace metal composition in both biogenic and abiotic particles, their quantitative relationship can be obtained in the following mass balance formula:

$$[M] = a[P] + b[Al]$$

- $[M]$ total concentration of metal in particulate matter
- $[P]$ phosphorus concentration in biogenic particles
- $[Al]$ aluminum concentration in abiogenic particles
- $a$ P-normalized metal quotas in biogenic particles
- $b$ Al-normalized metal quotas in abiogenic (assuming lithogenic) particles

Although biogenic hard parts (e.g., CaCO$_3$ and SiO$_2$) contribute a significant portion in biogenic particles in terms of mass, the concentrations of trace metals and P in the hard parts are so low that the contribution of these elements in the hard parts to the overall biogenic particles are ignorable. The M/P ratios in biogenic organic particles may reasonably represent the overall ratios in biogenic particles. The relative influence of biogenic and lithogenic sources on Cd in the SPM is first evaluated by examining the vertical distribution patterns for particulate Al and P, coupled with the concentrations of SPM and Chlorophyll-a in the SPM of the surface waters (Fig. 2). The vertical concentration of particulate Al increased with depth in the top 200 m. The total SPM, chlorophyll-a, and particulate P exhibited a similar distribution pattern, all with a maximum at depth of 60 m, showing that most of the particles at the chlorophyll maximum depth were composed of phytoplankton (Ho et al., 2007). The concentrations of Cd in the SPM also exhibited a similar distribution pattern, all with a maximum at depth of 60 m, showing that most of the particles at the chlorophyll maximum depth were composed of phytoplankton, particularly in the samples from shallow depths.

The relationship among the ratios of Cd/P and Al/P is shown in Fig. 3, which includes the data obtained from the cruises of March 2002 and October 2006 (Table 1). The three points in each figure represent the averaged metal quotas in each size fraction, from the

$$\frac{[M]}{[P]} = a + b\frac{[Al]}{[P]}$$

![Fig. 2. The vertical profiles of P, Cd, Al, and Chlorophyll-a in SPM, and the total SPM collected in the surface water of the SEATS site.](image)

![Fig. 3. The correlation of Cd/P or Zn/P to Al/P (mmol/mol) in the three size-fractionated plankton, 10–60 μm, 60–150 μm, and larger than 150 μm, which were collected at the chlorophyll maximum depth of the SEATS site in March 2002 and October 2006. Each point with 1 standard deviation as error bars (n = 3 to 5) stands for the average M/P ratios for each size fraction. The smaller the plank sizes were the higher the Al/P ratios were. The dotted lines in the figure stand for the averaged Zn and Cd quotas in marine plankton assemblages obtained from reliable field and laboratory studies (Ho, 2006).](image)
smallest to the largest with the increasing Al/P ratios. Our previous study found that the metal composition in the phytoplankton collected in the same sampling site was mainly dominated by extracellularly adsorbed abiotic forms and the metals were highly likely derived from anthropogenic aerosols containing abundant dissolvable trace metals (Ho et al., 2007). Using Zn as an example, the Zn quota in plankton increased with increasing abiotic Al/P ratios, showing that Zn was strongly associated with abiotic Al (Ho et al., 2007). The trace metals distributed in the same way as Zn included: Ti, V, Cr, Mn, Fe, Co, Ni, and Cu. In contrast, the correlation of Cd with Al was different from other metals. Despite Al/P ratios increasing from 14, 76, to 248 mmol/mol in the three fractions of plankton in 2004, respectively, the Cd quotas were fairly consistent, with the average of 0.26 ± 0.04 (1SD), 0.26 ± 0.03, to 0.34 ± 0.02 (mmol/mol P) for the plankton from the largest to the smallest. In 2006, when Al/P increased from 8, 62, to 287, the Cd quota only varied from 0.12 ± 0.003, 0.16 ± 0.05, to 0.19 ± 0.03, respectively. Compared to the one order of magnitude increase for Zn/P quota, the Cd/P quota was fairly consistent, insignificant or slightly higher in the smallest fraction of the plankton. The relatively consistent Cd/P quota in the diverse plankton with increasing Al indicates that most of the Cd in the suspended particles was located in plankton intracellularly but not adsorbed extracellularly, and the Cd bio-accumulation through the microbial food web was ignorable. Although there was no significant variability for the Zn/P and Al/P ratios in individual plankton fractions between the two cruises, the Cd/P ratios were lower in the samples collected at the three depths in October 2006.

Table 1

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a n.a.: data not available.

3.2. Cd and P fluxes

Seasonal variability of primary production has been reported at the sampling site (Chen, 2005; Tseng et al., 2005), with highest production during winter time and lowest during the summer season. Seasonal patterns of organic matter fluxes were recorded in the traps at 120 m, with a sharp increase for both Cd and P from December 2004 to February 2005 when primary production was highest (Fig 4). The depth of the first trap was close to the bottom depth of the euphotic zone (Chen, 2005), where the organic carbon fluxes estimated from sinking particles may reasonably represent new production in the euphotic zone (Pace et al., 1987). New production in the euphotic zone was estimated by multiplying the export P fluxes obtained from the 120 m traps with the Redfield ratio, which ranged from 0.05 to 0.85 gC m⁻² d⁻¹ among the lowest and highest seasons, compatible to the new production estimated by nitrate uptake at the same sampling site, which ranged from 0.03 to 0.26 gC m⁻² d⁻¹ during the lowest to highest production periods (Chen, 2005). It should be noted that new production estimated by nitrate uptake was based on 1 to 2 cruise studies in each season, and these may not have been able to catch the maximum production period as shown in Fig 4. In general, these two new production estimates were consistent with each other while considering the possible uncertainties in both analyses.

At the two deeper depths, the fluxes in P and Cd decreased exponentially. The average flux of P in the 600 m trap was only about one tenth of flux at 120 m, ranging from 5 to 50 μmol/m²/d. Relatively, Cd flux decreased even more sharply, ranging from 1 to 10 nmol/m²/d overall. The Cd and P fluxes in the deepest trap decreased to 0.4–2.4 (nmol/m²/d) and 4–12 (μmol/m²/d), respectively. Except for elevated fluxes during December 2004 to January 2005 in the middle trap, there was no strong seasonal variability for fluxes observed in the deeper water. An abnormally high Al flux was observed during the end
of December 2004 when the South Asia Tsunami occurred. Although the abiotic P portion estimated from Al fluxes only accounted for 10% of the total sinking P pool (Fig. 5), the fluxes of P, but not Cd, were significantly elevated in the two deeper traps (Fig. 4), indicating a preferential removal of P during the tsunami event. Based on the sinking fluxes and the standing stock in seawater (Wen et al., 2006), the residence time of biogenic Cd and P in the zonation of 0–120 m, 0–600 m, and the whole water column are 0.10 and 0.20, 250 and 100, 9100 and 5000 years respectively according to box modeling. The residence time shows that the sinking rate of Cd in the euphotic zone was higher than P but the removal rates of P in the thermocline and the deep water were higher than Cd.

The organic carbon estimated from the biogenic P fluxes of the deepest trap may represent the net organic carbon export of the water column vertically. The portion of abiotic P in the deeper water needs to be excluded while estimating organic carbon fluxes (Fig. 5). The percentage of abiotic P and Cd ranged from less than 2% in the traps at 120 m to about 20–30% in the bottom trap (Fig. 5). After subtracting the abiotic portion, the annual biogenic P flux in the deepest trap was 1.9 mmol/m²/yr, obtained by multiplying 365 by the averaged daily biogenic P flux 5.3 mmol/m²/d, which is equivalent to about 0.20 mmol/m²/yr or 6 g/m²/yr organic carbon export to the sediment.

3.3. Cd/P ratios in sinking particles and seawater

Dramatic seasonal variabilities of Cd/P ratios were also observed in sinking particles collected by the traps at 120 m (Fig. 6), ranging from 0.1 to 1.2 (mmol/mol). In particular, there was a sharp increase from early December of 2004 to late February of 2005. Despite the low Cd/P ratio being seemingly related to the relatively high Al fluxes around December, the relative contribution of lithogenic Al fluxes on Cd/P ratios were ignorable in the traps at 120 m (Fig. 6). The lithogenic P and Cd calculated by using Al fluxes shows that lithogenic Cd and P

![Image](https://example.com/image.png)

**Fig. 5.** (a) The percentage of the abiotic Cd and P in the sinking particles, estimated by Al concentrations in the sinking particles using the Cd/Al and P/Al standard ratios in crustal composition, which are 0.0006 and 11 (mmol/mol to Al), respectively (Taylor, 1964). The P/Al ratios in the sediments of the South China Sea were reported by Calvert et al. (1995). The averaged P/Al ratio in the sediments collected from two stations (GGC1 and GGC2 in Table 1), close to our sampling site, was 11.2±2.7, extremely close to the standard ratio. (b) The comparison of the Cd/P ratios in the sinking particles with Al fluxes.

![Image](https://example.com/image.png)

**Fig. 6.** The vertical and temporal variability of Al fluxes and Cd/P ratios in the sinking particles collected at the three depths of 120, 610, and 3500 m of the SEATS site.
roughly equivalent to 60 mg/m²/d of lithogenic mineral fluxes, and simply assuming P adsorption capacity to be 1 mg P/g clay minerals, the corresponding P adsorption would be around 2 μmol/m²/d, accounting for a significant amount of the total P fluxes observed in the deep water. This estimate appears to be preliminary. However, the results indicate that the role of lithogenic particles on removing soluble P and influencing Cd/P ratios in sinking particles and seawater can be significant, particularly over long time periods. To further test our argument, we have studied the data sets published by Noriki et al. (1999). The sinking particles in the study were collected by sediment traps deployed in the Pacific Ocean at depths ranging from 1500 to 5100 m at three different stations, where the relative percentages of the organic matter and biogenic hard parts in the sinking particles did not change much. Once again, we have observed a strong exponential decay relationship between the Cd/P ratios and the clay percentage (Fig. 7), strongly supporting our argument.

The other factor which may influence the Cd/P ratios in particles and seawater would be the community structure of phytoplankton. Culture studies have clearly shown that the variability of Cd quotas in 5 major marine algae phyla grown in identical medium ranged up to two orders of magnitude, from 0.01 to 1.0 mmol/mol P (Ho et al., 2003, 2004). This two orders of magnitude range was largest among the trace metals measured, including: Fe, Mn, Zn, Cu, Co, Cd, and Mo (Ho et al., 2003). The ratios were generally high for coccolithophores and low for diatom and chlorophyta. Culture study by Sunda and Huntsman (2000) also showed much higher Cd quota in coccolithophores than in diatom. Dinoflagellates with cysts can have high Cd quotas intracellularly (Ho et al., 2003). Once the algal community structure shifts, it is expected that one would observe different Cd/P ratios in plankton and thus resultant sinking organic particles. With relatively low nitrate concentrations in the surface water during the productive winter season, it is known that coccolithophore is the dominant algal species during the winter season in the South China Sea (Chen et al., 2007), which generally happens from January to March. When the northeastern monsoon starts to blow in October and brings the nutrients to the surface water to induce the growth of diatom, the Cd/P ratios in sinking particles, which are supposedly dominated by diatoms, would be relatively low. After nitrate concentrations go down, the succession of coccolithophores would follow and dominate (Chen et al., 2007). The change of Cd/P ratios from December to February 2005 may thus be caused by the succession of microalgal species. Field studies also support the argument. Nolting et al. (1999) reported that Cd removal by phytoplankton during Emiliania huxleyi bloom was more significant than for other bioactive trace metals. Ho (2006) has compiled a few reliable field studies and reported that Cd quotas in the phytoplankton samples collected in various oligotrophic open ocean regions were remarkably consistent, ranging from 0.44 to 0.54 mmol/mol P. These quotas are almost one order of magnitude higher than the Cd quota observed by Martin and Knauer (1973) in the coastal plankton samples known to be dominated by diatom during spring bloom period. Once the biogenic particles formed in oceanic surface waters can reach deep water, the shift of microalgal community structure in oceanic surface water can be a potentially important factor in influencing dissolved Cd/P ratios in deep waters when the structure change lasts long enough, particularly longer than the residence time of Cd and P in deep water.

It is puzzling why there are so divergent Cd quotas among different phytoplankton phyla or why some phytoplankton take up Cd strongly in oceanic surface waters. It seems to be unlikely that Cd uptake by phytoplankton is mainly due to Zn replacement. Both field and laboratory studies show that phytoplankton actively take up plenty of Cd even when the supply of Zn is adequate (Nolting et al., 1999; Whitfield, 2001; Ho et al., 2003). Our previous study on the elemental composition in various marine phytoplankton showed that there are two other cellular elements featured with extremely large composition ranges, which are Ca and Sr (Ho et al., 2003). The accidental uptake of elements with similar size charge ratios by phytoplankton may be important since trace metal transporters on algal cellular surfaces are seldom specific. The element with the closest size charge ratio to Cd⁺⁺ ion in the periodic table is Ca²⁺ ion, which is an essential and major element for all phytoplankton especially for coccolithophores and the species with CaCO₃ cysts. It has also been known that Cd can inhibit Ca pump and membrane transport through fish studies (Verboest et al., 1988, 1989a,b); and the influx of Cd was significantly inhibited by a Ca channel blocker in mussels (Wang and Fisher, 1999), suggesting that Cd uptake in these organisms is related to Ca transporters or channels. Field studies in regions where coccolithophore is dominant found high Cd quotas in phytoplankton samples (Kuss and Kremling, 1999; Abe, 2005); culture studies show that calcareous algae are species with high Cd quotas and resistance to high Cd concentrations in seawater (Payne and Price, 1999; Sunda and Huntsman, 2000; Ho et al., 2003). Whether the strong microalgal Cd uptake and the highly varied Cd/P quotas in phytoplankton are caused by accidental uptake through Ca transporters and channels deserves further studies.

The third factor to influence the Cd/P ratios in sinking particles and seawater may be the aeolian supply of the soluble Cd and P in surface water. Dissolved concentrations of Cd and P in seawater were also determined when plankton was collected in 2002. The dissolved Cd/P ratios in the top 1000 m of seawater varied significantly (Fig. 8),

![Fig. 7. The comparison of Cd/P ratios with the clay percentage in the sinking particles collected by the sediment traps deployed in the deep waters of the Pacific Ocean (Noriki et al., 1999). The sampling sites from 1 to 3 are located at 48°N, 13°N, and 0°N, along 175°E, respectively. In addition to clay minerals, opal, calcium carbonate, and organic material were also determined. The major components in the sinking particles were the hard parts, accounting for 80 to 90% of the total mass. The mass percentage of organic matter was generally around 10% with limited variability for most of the samples.](image)

![Fig. 8. The dissolved Cd/P ratios (mmol/mol) in seawater of the sampling site in March 2002 and 2004. The ratios are retrieved from the published data in Wen et al. (2006).](image)
preferential algal uptake of Cd. In deep water, dissolved Cd/P ratios at a depth of chlorophyll-a by coccolithophores during the sampling period of March 2002 (183 nmol/m²/d) for this study site. However, soluble Cd/P ratios of averaged deposition of marine aerosols samples collected from the East China Sea shows that dissolved Cd/P ratios in seawater. The high Cd/P ratio in seawater could thus be caused by the input of aeolian aerosols with highly elevated Cd/P ratios. In the long term, consistent input of elevated soluble Cd and P from aerosols to surface water may gradually influence Cd/P ratios both for biogenic particles and seawater.

The lowest Cd/P ratio in seawater was observed at the depth of chlorophyll-a maximum where size-fractionated plankton was collected. Since microagelal species were possibly dominated by coccolithophores during the sampling period of March 2002 and 2004 (Chen et al., 2007), the low Cd/P ratios in seawater at the depth of the chlorophyll-a maximum may be caused by the preferential algal uptake of Cd. In deep water, dissolved Cd/P ratios in seawater close to 0.3 mmol/mol; the ratios are mainly controlled by preformed deep water based on global seawater horizontal segregation.

4. Conclusion

The major concern in using Cd/P ratios as proxies in paleoceanography is whether the ratios in seawater may remain constant over a geological time scale (Sager and de Baar, 1993). Our study shows that Cd/P ratios in sinking particles varied remarkably with time and depths in the sampling site. In the particles collected at 120 m, the variability of the ratios was associated with changes of primary production and microagelal community structure. At the two deeper depths, increasing lithogenic particles greatly decreased the Cd/P ratios in sinking particles. Several lines of evidences indicate that P adsorption on mineral clays is an important process in influencing the ratios in the sinking particles of the water column. These two factors, including the change of phytoplankton community structure and P adsorption on mineral clays, can be important in influencing Cd/P ratios not only in particles but also in seawater over a long term. Whether P adsorption by lithogenic particles occurs commonly in other marginal seas and open oceans, and how P adsorption by mineral clays quantitatively influences the ratios of Cd/P and Cd/P both in particles and seawater, and P residence time on a global scale all deserve further investigation.

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