Diatoms and their influence on the biologically mediated uptake of atmospheric CO₂ in the Arabian Sea upwelling system

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Abstract. Sediment trap experiments have been carried out in order to study processes controlling shifts from diatom to non-diatom dominated systems in the western Arabian Sea. One of our major problems was to link sediment trap records to surface ocean processes. Satellite-derived observations on upper ocean parameters were helpful to reduce this problem in the past and gain a new quality by combining it with results obtained during the Joint Global Ocean Flux Study (JGOFS) in the Arabian Sea. The new results imply that intense grazing can decline or impede the development of large diatom blooms in a silicon-enriched near shore upwelling system off Oman. In the open western Arabian Sea diatom blooms recover within the offshore advecting upwelled water and lead to peak organic fluxes into the deep sea but only during the later phase of the upwelling season. During onset of the upwelling season grazing favoured by eolian iron inputs causing the formation of thinner diatom shells seems to prevent the development of a large diatom bloom within the silicon-enriched offshore advecting upwelled water. An increased relevance of diatoms and diatom-grazing copepods in the planktonic community as well as oligotrophic conditions seem to raise the ratio between organic carbon production and precipitation of calcium carbonate (rain ratio) in the surface water. The decomposition of organic matter in the water column reduces the rain ratio within in the sinking matter especially in the oligotrophic region dominated by cyanobacteria and reduces also the variation of the carbon to nutrient uptake ratios seen in the surface water.

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Possible effects of changing Redfield and rain ratios on the atmospheric CO$_2$ concentration were quantified using GCMs. It was estimated that an increase of the C/P ratio by 30% (122:1 to 158.6:1) could lower the atmospheric CO$_2$ concentration by $\sim$72 ppm (Heinze et al., 1991). A doubling of the global mean rain ratio from 4 to 8 could reduce the atmospheric CO$_2$ concentration by 28.5 ppm (Heinze et al., 1991), and raising the rain ratio from 5 to 16.6 could, however, reduce the atmospheric CO$_2$ concentration by 70 ppm (Archer et al., 2000). These changes explaining a major proportion of the glacial/interglacial variation of the atmospheric CO$_2$ concentration were achieved by assuming that diatoms outcompete the carbonate-producing coccolithophorids and drive the export production. Due to such a competition the rain ratio is directly linked to biogenic opal production in some models used to study the feedback impact of the carbonate production in the ocean on increasing CO$_2$ concentrations in the atmosphere (Heinze, 2004).

Since diatom growth is believed to be limited by the availability of silicon at lower latitudes (Dugdale and Wilkerson, 1998; Rixen et al., 2000), changes of the global silicon cycle and re-organisation of the marine silicon cycle were proposed as being possible mechanisms to fertilise lower latitudes with silicon during glacial times (Froelich et al., 1992; Harrison, 2000; Conley, 2002; Ridgwell et al., 2002). The re-organisation of the marine silicon cycle is suggested to be triggered by an enhanced eolian iron input lowering the Si/N uptake ratio of diatoms in the Southern Ocean and, subsequently, enhancing the silicon export from higher to lower latitudes (Matsumoto and Sarmiento, 2002).

In addition to coccolithophorids competing with diatoms, there are also carbonate-producing heterotrophs like foraminifera and pteropods which feed on diatoms. In the Arabian Sea sediment trap experiments showed that coccolithophorids contribute only $<15\%$ to the carbonate export into the deep sea and that the peak flux of foraminifera into the deep sea coincides with that of diatoms during the highly productive upwelling season (Haake et al., 1993b, 1993a; Zeltner, 2000). This suggests strongly that effects of foraminifera and pteropods on the rain ratio should be taken into consideration, as these organisms are important or even the main carbonate producer in the ocean (Table 1). The total fluxes of calcium carbonate and organic carbon measured by deep moored sediment traps in the Arabian Sea (Lee et al., 1998; Honjo et al., 1999; Rixen et al., 2002) and ratios between calcium carbonate dissolution and organic carbon remineralization in the water column (Hupe and Karstensen, 2000) have been used to calculate rain ratios (Rixen et al., 2005). In this study these calculated rain ratios ranging between 2 and 3.5 will be compared with rain ratios derived from total dissolved inorganic carbon concentrations (DIC), total alkalinity (TA), and CO$_2$ partial pressure differences ($\Delta p$/CO$_2$) between the atmosphere and surface water (Goyet et al., 1998b, 1999; Millero et al., 1998). Furthermore plankton counts (Garrison et al., 2000; Schiebel et al., 2004) will be evaluated in conjunction with nutrient (Morrison et al., 1998) and iron concentrations (Measures and Vink, 1999) in order to study factors influencing shifts in the planktonic community structure and associated changes of the composition of sinking particles.

### Table 1. Contribution of carbonate-producing organism to the pelagic marine carbonate production.

<table>
<thead>
<tr>
<th></th>
<th>%</th>
<th>$10^{15}$ g C</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total carbonate production</td>
<td>100</td>
<td>0.60–0.86</td>
<td>Milliman and Droxl (1996)</td>
</tr>
<tr>
<td>Foraminifera</td>
<td>23–56</td>
<td>0.14–0.48</td>
<td>Schiebel (2002)</td>
</tr>
<tr>
<td>Pteropods</td>
<td>$\sim$10</td>
<td>0.06–0.09</td>
<td>Schiebel (2002)*</td>
</tr>
<tr>
<td>Coccolithophorids</td>
<td>4–38</td>
<td>0.02–0.33</td>
<td>Schiebel (2002)*</td>
</tr>
</tbody>
</table>

* and references therein.

2 Study area

The Arabian Sea is strongly influenced by the Asian monsoon. This climatic feature is driven by the sea-level pressure difference between the Asian landmass and the Indian Ocean (Ramage, 1971, 1987). During the boreal winter the sea level pressure over Asia exceeds that over the Indian Ocean due to a stronger cooling of the landmass. Following the pressure gradient and deflected by the Coriolis force, the wind blows from the NE (NE monsoon) over the Arabian Sea. This situation reverses when the summer heating of the Asian landmass leads to the formation of one of the strongest atmospheric lows on Earth. This low attracts the SE trade winds, and after crossing the equator the former SE winds blow as SW winds over the Arabian Sea due to associated changes of the Coriolis force. The SW winds (SW monsoon) form a tropospheric jet (Findslater Jet) extending almost parallel to the Arabian coast (Fig. 1, Findlater, 1977; Rixen et al., 1996). The monsoon winds and the deserts surrounding the western and northern parts of the Arabian Sea lead to dust inputs into the Arabian Sea, which are among the highest in the world ocean (Tegen and Fung, 1994, 1995).
Biological productivity within the Arabian Sea is determined by the interplay between the euphotic zone and mixed layer depth (MLD), whose deepening is caused by winter cooling and wind mixing (Rixen et al., 2002). The interplay between the euphotic zone and the MLD regulating the availability of light and nutrients is well known from the temperate ocean. In the Arabian Sea this leads to early and late NE monsoon blooms. During the SW monsoon the Findlater Jet creates one of the most productive upwelling areas in the ocean (Antoine et al., 1996) and a hot spot for CO$_2$ emission along the Arabian coast (Körtzinger et al., 1997; Goyet et al., 1998b, 1998a; Sabine et al., 2000). Diatom blooms are common during the later phases of the NE and SW monsoon (Haake et al., 1993b; Rixen et al., 2000).

3 Data base, methods and results

Nutrients, DIC, TA, temperature, and salinity profiles measured at the sampling sites S1–S15 during the U.S. JGOFS cruises ttn49 (18 July 1995–13 August 1995) and ttn50 (14 August 1995–13 September 1995) were obtained from the U.S. JGOFS database (Figs. 1, 2). The mixed layer depth was defined as the depth at which a pronounced temperature decrease and nutrient increase occurred within profiles. Subsequently all data were averaged for the mixed layer depth (Table 2). The resulting mean mixed layer temperatures and salinities increase generally from the coastal upwelling zone towards the open ocean (Fig. 2). During both cruises slightly reduced temperatures and salinities occurred between stations S5 and S7, approximately 500 km offshore. As shown by satellite-derived SSTs charts (Fig. 2), this anomaly was associated with the filament that extended almost parallel to the transect, perpendicular to the coast towards the open Arabian Sea. Filaments are cold water structures caused by an accelerated advection of upwelled water. While moving offshore the upwelled water get mixed with the warmer and saltier surface water that was formed during the preceding oligotrophic intermonsoon season (Fischer et al., 2002; Weller et al., 2002). Mixing of two water masses is indicated by a linear correlation between temperature (T) and salinity (S) if latent and sensible heat fluxes between ocean and atmospheres are negligible. Temperature und salinity (T/S) data derived from the sampling sites along the Oman transect are not correlated (Fig. 3) and T/S data obtained from
the sites S2–S14 deviate from the line (mixing line) that connects T/S data obtained from the sampling site S1 and S15. In order to calculate energy fluxes required to explain these deviations from the mixing line (ΔSST) the time during which the surface water was in contact with the atmosphere must be known. The ages of the surface water can be calculated at each sampling site by dividing the distance to the coast by the mean advection velocity and considering that the upwelled water was already a few days old prior to it left the coast. Since the lower temperatures and the reduced salinity indicate an accelerated advection of upwelled water within the filament an age correction has been applied. The distance from the coast towards the most offshore station (S13) and the station closest to the coast (S1) is ∼1100 km and ∼26 km, respectively. At these two stations the salinity was 36.6 and 35.7 (Table 2). Based on these two points defined by the distance to the coast and the salinity a linear regression was developed which allowed us to derive corrected distances from the salinity measured at each sampling site (distance [km]=1135.1×Salinity−40451.8). The time since the water mass was in contact with the atmosphere was obtained by dividing the corrected distance by the mean advection velocity ranging between 0.2 and 0.8 m s⁻¹ (Rixen et al., 2000). Additionally it was assumed that coastally upwelled water was already 4 days old prior to it left the coast. The energy fluxes required to explain ΔSST have been calculated as follows: \( \rho \times \text{MLD} \times \text{Cp} \times \Delta \text{SST} \) / (age of the water), whereas ‘‘\( \rho \)’’ is the sea water density in ‘‘kg m⁻³’’, MLD is the mixed layer depth in ‘‘m’’, ‘‘\( \text{Cp} \)’’ is the specific heat of the water (∼3980 J kg⁻¹ K⁻¹), \( \Delta \text{SST} \) and the age of the water are given in ‘‘°C’’ and seconds, respectively. The calculated data show that a heat flux of 90 to 584 W m⁻² could have caused the ΔSST at sampling site S7 (Table 3). Energy fluxes of 150 to 200 W m⁻² determined at the U.S. JGOFS surface mooring site which is close to S7 (Weller et al., 1998) fall within this range implying that heat fluxes could have caused ΔSST at station S7 and most probably also at the other water sampling sites for which no data on heat fluxes are available.

Latent heat fluxes do not affect the relationship between salinity and nutrient concentrations as changes in the amount of water in the mixed layer increase both the salinity and the nutrient concentration. Since in addition to that salinity and nutrient concentration are unaffected by sensible heat flux, salinity instead of temperature was used to define the end-members and to calculate the mixing ratios “a” and “b” within the two-end-member mixing analysis at sampling sites S1–S13 (Salinity\( _{\text{S1}−\text{S13}} \approx a_{\text{S1}−\text{S13}} \times \text{Salinity upwelled water} + b_{\text{S1}−\text{S13}} \times \text{Salinity oligotrophic water} \)), whereas \( a_{\text{S1}−\text{S13}} + b_{\text{S1}−\text{S13}} = 1 \). Station S1 was defined as the upwelling and station S13 as the oligotrophic end-member because there are hardly any discernible variations in the salinity between stations S13.
A mixing ratio “b” of zero indicates pure upwelled water whereas a mixing ratio “b” of one implies that no upwelled water is present. The mixing ratios “a” and “b” determined for each station and the mean nutrient concentrations could be expected if mixing were the only factor controlling the nutrient concentration. Deviations of measured phosphate, inorganic nitrogen, and silicon concentrations from the mixing line can be attributed to biological consumption. In order to determine error ranges of the derived biological consumption caused by analytical methods, a relative percentage error of 0.5% and 0.012% of nutrient and salinity data (U.S. JGOFS data base documentation) was considered. Within these error ranges random errors were produced by applying the standard fortran 77 random number generator (F77-RNG). Subsequently, the mixing analysis was performed 500 times by using the produced errors. The mean ratios of inorganic nitrogen and phosphate consumptions (N/P ratios) vary between ~7 and 24 (Fig. 5b) and fall within the range of N/P ratios determined in marine particulate matter and phytoplankton (~3–34, Geider and La Roche, 2002; Klausmeier et al., 2004). The resulting standard deviations of the N/P ratios range between 0.1 and 0.4.

The DIC and TA data were treated in the same way as nutrient data, but prior to calculating the biological consumption the DIC concentrations were corrected for CO2 emission. The CO2 emission into the atmosphere have
been derived from the \(\Delta pCO_2\) data published for each water sampling site in the U.S. JGOFS database (Table 2). The required wind-dependent gas transfer velocity “k” was calculated using eight different formulations (Liss and Merlivat, 1986; Wanninkhof, 1992; Wanninkhof and McGillis, 1999; Nightingale et al., 2000; Feely et al., 2001). Satellite-derived wind speeds (see Fig. 1) were taken from Rixen et al. (1996). The standard deviation and the resulting mean C/P and rain ratios and their standard deviations are given in Fig. 5a. The mean C/P uptake ratios vary between 80 and 150 and are well within the range of published C/P ratios observed in marine particulate matter and phytoplankton. The same holds true for the rain ratios which vary between \(\sim 2.3\) and \(\sim 8.1\) except at station S11.

### Table 3. Sampling site, the mean mixed layer temperature (determined), the temperature as suggested by the mixing line indicate in Fig. 4 (expected), the difference between the mixed layer temperature and the expected temperature (\(\Delta TST\)), and energy fluxes required to explain the temperature deviation. The energy flux has been calculated by assuming a mean advection velocity of 0.8 and 0.2 m s\(^{-1}\). 

<table>
<thead>
<tr>
<th>Sampling site</th>
<th>Temperature [(\circ)C]</th>
<th>Energy fluxes [W m(^{-2})]</th>
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<tr>
<td></td>
<td>determined</td>
<td>expected deviation</td>
</tr>
<tr>
<td><strong>tn 49</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S12</td>
<td>27.19</td>
<td>26.86</td>
</tr>
<tr>
<td>S11</td>
<td>27.28</td>
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<td>S10</td>
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<tr>
<td>S9</td>
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<td>S8</td>
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<td><strong>S7</strong></td>
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</tr>
<tr>
<td>S6</td>
<td>24.55</td>
<td>22.22</td>
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<td>S5</td>
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<td>S4</td>
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</tr>
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<td>S3</td>
<td>25.98</td>
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<tr>
<td>S2</td>
<td>22.97</td>
<td>21.37</td>
</tr>
<tr>
<td><strong>tn 50</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S11</td>
<td>26.93</td>
<td>25.37</td>
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<td>S9</td>
<td>26.53</td>
<td>24.09</td>
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<tr>
<td>S8</td>
<td>26.10</td>
<td>23.18</td>
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<tr>
<td><strong>S7</strong></td>
<td>26.52</td>
<td>23.89</td>
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<td>S6</td>
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<tr>
<td>S5</td>
<td>25.64</td>
<td>22.38</td>
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<tr>
<td>S4</td>
<td>24.88</td>
<td>21.95</td>
</tr>
<tr>
<td>S3</td>
<td>23.82</td>
<td>21.59</td>
</tr>
<tr>
<td>S2</td>
<td>23.92</td>
<td>21.21</td>
</tr>
</tbody>
</table>

**Fig. 5.** (a) C/P, (b) N/P and (c) rain (POC/PIC) ratios versus the proportion of oligotrophic water. Error bars are calculated as described in the text. The number in panel (c) indicates the rain ratio at the sampling site S11 which is not plotted as it is out of scale.
where the rain ratio is 22.3 (Fig. 5c). However, due to the low carbon consumption the standard deviation at the sampling sites S10–S12 (proportion of oligotrophic water >0.7) are so high that these data have to be treated with caution within the following discussion.

In order to calculate the new production rates (Dugdale and Goering, 1967; Eppley and Peterson, 1979) the POC-production (see Eq. 3) was integrated over the depth of the mixed layer and subsequently divided by the age of the upwelled water. The comparison with primary and export production rates measured at the same time and at the same stations (Buesseler et al., 1998) show that the resulting new carbon production rates are lower than the primary production rates, as expected (Fig. 7). Since new production exceeds export production rates it is assumed that the biomass is growing in the mixed layer but under different environmental conditions.

New production rates which are consistent with published data on primary and export production rates and reliable rain and Redfield ratios suggest that our approach is suitable for studying biogeochemical processes in the Arabian Sea during the upwelling season. In the following discussion the results obtained by our mixing analysis will, furthermore, be linked to plankton counts and sediment trap data in order to investigate factors influencing changes in the planktonic community structure and associated impacts on the rain ratio.

4 Discussion

4.1 Diatom blooms

The shoaling of the mixed layer and enhanced organic carbon flux measured in the deep sea reveal that the cruise ttn49 and the cruise German JGOFS cruise M32/5 took place at the beginning of the upwelling season in the open western Arabian Sea (Fig. 8). On month later during the cruise ttn50 the high organic carbon fluxes indicate the peak upwelling season. At this time plankton counts (Garrison et al., 2000) show that diatoms dominate the planktonic community from approximately 100 to 400 km offshore (Fig. 9b). Flagellates succeed diatoms in the central Arabian Sea after silicon concentrations in the surface water reach their oligotrophic intermonsoon value of \( \sim 2 \mu \text{mol kg}^{-1} \), (compare Figs. 9b and d). Plankton growth rates generally decrease with decreasing nutrient concentrations after the latter falls below a certain threshold (Lalli and Parsons, 1993). Assuming that such a threshold is close to a silicon concentration of \( \sim 2 \mu \text{mol kg}^{-1} \) diatom growth rates falling below the high grazing rates (Smith et al., 1998) could explain the declining diatom bloom in the open western Arabian Sea where silicon in the surface water is not consumed. This observation confirms experimental data suggesting a silicon-threshold of \( \sim 2 \mu \text{mol kg}^{-1} \) for the dominance of diatoms within the planktonic community (Egge and Aksnes, 1992). Near the coast, at silicon concentration of \( \sim 11 \mu \text{mol kg}^{-1} \) the relatively low contribution of diatoms to the planktonic community (Fig. 9b, d) has been attributed to intense grazing by copepods slowing down the development of diatom blooms in the coastal region off Oman (Smith, 2001). During the early phase of the upwelling season the abundance of diatoms decreased drastically close to the Oman coast as shown by plankton counts.
(Fig. 9a, Schiebel et al., 2004). This decline is accompanied with a transition from a *Chaetoceros* dominated diatom assemblage near the coast to one dominated by *Nitzchia bicoapsitata* further offshore. The decline and the associated transition of the diatom assemblage has been linked to decreasing nutrient concentrations and intense grazing (Smith, 2001; Schiebel et al., 2004).

The results obtained by the plankton counts agree with those derived from deep moored sediment traps and our mixing analysis showing silicon consumption and carbonate biogenic opal ratios which are lower and higher, respectively, during the early than during the later phase of the upwelling season in the open western Arabian Sea (Figs. 9c, 10a). Vice versa, near the coast carbonate biogenic opal ratios reveal a higher contribution of diatoms to the exported matter during the early than during the later phase of the upwelling season (Fig. 9c). This seems to be caused by the export of the declining diatom bloom during the early phase of the upwelling season (Fig. 9a) and the delayed development of the diatom bloom one month later (Fig. 9b).

4.2 The role of iron for the development of diatom blooms

Iron enrichment experiments in the coastal upwelling system off California revealed that iron fertilization favours the growths rates of *Chaetoceros* and other diatoms but leads to the formation of thinner shells as indicated by a low uptake of silicon relative to nitrogen (Hutchins and Bruland, 1998). The iron concentration in the diatom dominated open western Arabian Sea range between 0.5 and 1 nmol l$^{-1}$ during the cruise ttn50 and reach values of >2 nmol l$^{-1}$ during the cruise ttn49 near the coast (Fig. 10b). Off California an increase in the iron concentration from 0.5 to 2.5 halves Si/N uptake ratio (Hutchins and Bruland, 1998). Assuming a similar impact of iron on the Si/N uptake off Oman suggests that *Chaetoceros* blooming during the onset of the upwelling season near the coast off Oman might built thinner
shells than diatoms growing a month later in same region but in water revealing lower iron concentrations. The extremely strong diatom shell is an important protection against predators (Hamm et al., 2003). Since enhanced iron concentrations lead to the formation of thinner shells it is suggested that copepod grazing is favoured in iron-enriched environments. An iron-favoured grazing could explain the decline of the near shore diatom bloom during the onset of the upwelling season (Fig. 9a). On the other hand a less efficient grazing due to lower iron concentrations (Fig. 10b) and thicker shells could slow down but do not prevent the development of a large diatom bloom during the later phase of the upwelling season (Fig. 9b) at which the silicon concentrations are almost equal to those during early phase of the upwelling season (Fig. 10d).

4.3 Rain ratios

Apart from one exception as discussed before the rain ratios derived from mixing analysis range between ~2.3 and 8.1 and are generally higher than those obtained from sediment trap data (2–3.5, Fig. 11). The rain ratios derived from the sediment trap data have been adjusted to a water-depth of 100 m whereas the mixed layer depth is <100 m in the western Arabian Sea (Table 1). POC/PIC ratios tend to decrease within increasing water depth because the decomposition of organic matter is faster than the dissolution of carbonates in the upper water column. Consequently, the difference between rain ratios derived from sediment trap data and the mixing analysis could be caused by differences in the water-depth. Moreover, one should consider that sediment trap data represent a larger area and a longer time interval than data derived form nutrient profiles obtained at a specific site within a relatively short period of time.

However, rain ratios obtained from sediment trap data and the mixing analysis reveal a similar trend in the upwelling region characterized by a proportion of oligotrophic water (pow) <0.6 (Fig. 11). Near the upwelling centres (pow <0.35) the rain ratios increase with an increasing proportion of oligotrophic water until they reach a maximum at a pow of ~0.35. This increase is accompanied with an enhanced silicon consumption (Fig. 10a) indicating a slight recovery of the diatom (Nitzchia bicapitata) bloom after the dramatic decline of Chaetoceros dominated diatom bloom near the coast during the early upwelling season (Fig. 9a). At the transition between the upwelling dominated area and the more oligotrophic sites the rain ratios decrease by ~30 (sediment trap data) and 45% (mixing analysis). This decline is accompanied with decreasing silicon consumption (Fig. 10a) and during the later phase of the upwelling season also with decreasing contribution of diatoms to the plantonic community structure (Fig. 9b).

The high abundance of coccolithophorids at approximately 450 km offshore (Fig. 9a) occurred in a water mass...
which is characterized by a pow of ~0.2. The associated rain ratio of ~5 is relatively high (Fig. 11a) and implies in line with sediment trap results that coccolithophorids are only of minor importance for the carbonate export in the Arabian Sea. This in turn implies that changes of the rain ratios as discussed before are mainly caused by variations in the growth and export rates of foraminifera. Enhanced rain ratios associated with even a slightly increased contribution of diatoms in the planktonic community structure could for example results from predators such as copepods competing with foraminifera for diatoms.

High rain ratios seem to characterize the oligotrophic region, in which cyanobacteria dominate the planktonic community (Figs. 11, 9b). These high rain ratios derived from mixing analysis are not mirrored by the sediment trap data which reveal the lowest values in this region (Fig. 11). Since it is generally assumed that cyanobacteria are rapidly remineralized in the upper water column and hardly sink into the deep sea (Karl et al., 1996, 1997) a preferential decomposition of cyanobacterial biomass could explain the difference between the rain ratios derived from the mixing analysis and the those obtained from the sediment trap data. A preferential decomposition of more labile nitrogen containing compounds in the water column (Lee and Cronin, 1982; Wakeham et al., 1997; Lee et al., 2000) is also indicated by C/N ratios which are lower in the organic matter produced in the surface water than those measured in organic matter intercepted by sediment traps in the oligotrophic regions (Fig. 12).

4.4 C/N and C/P ratios

Contrary to the oligotrophic region C/N ratios partly decrease with depth implying a preferential decomposition carbon-enriched material in the upwelling area (Fig. 12). Such carbon-enriched organic matter could be transparent exopolymers (TEP) which are produced by diatoms and foster the formation of fast-sinking marine snow (Passow et al., 1994, 2001). The formation of TEP raises the C/P ratios especially at the end of diatom blooms (Engel et al., 2002) and along the transect at the transition from the upwelling towards the oligotrophic system the CP ratios reveal a drop from values >108 to 78 (Fig. 5a). The huge decrease of the C/P and C/N ratios is not reflected in the C/N ratios of the exported matter in the deep sea which show only a ~7% decrease at the transition from the upwelling towards the oligotrophic system (Fig. 11b). This implies that the preferential decomposition of TEP-like organic matter in the upper water column reduces the impact of the TEP formation on the Redfield ratio in the exported matter during diatom blooms.

5 Conclusions

The evaluation of data collected during the U.S. and German JGOFS expedition in 1995 suggest that an intense grazing decline and impede the development of large diatom blooms in a silicon-enriched near shore upwelling system off Oman. Within the offshore advecting upwelled water mass a large diatom blooms enhancing the organic carbon export into the deep sea develop only during the later phase of the upwelling period. During the onset of the upwelling season no large diatoms blooms occurs in the open western Arabian Sea, despite increased concentrations of dissolved silicon in the surface water. Enhanced iron concentration due to eolian dust inputs into the offshore advecting upwelled water might have favoured grazing at this time because it leads to the formation of thinner diatom shells. The decrease of the rain ratios during transition from upwelling towards the oligotrophic offshore region seems to be caused by foraminifera competing with diatom-grazing copepods for food. The large variability of Redfield ratios and also of the rain ratios in the oligotrophic region is not reflected in the sediment trap record. This suggests that the decomposition of organic matter in the water columns contributes to the relative uniform global mean Redfield ratio and leads to an underestimate of rain ratios derived from sediment traps records in oligotrophic regions dominated by cyanobacteria.

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References


Froelich, P. N., Blanc, V., Mortlock, R. A., and Chillrud, S. N.: River fluxes of dissolved silica to te ocean were higher during glacialis: Ge/Si in diatoms, rivers, and oceans, Paleoceanography, 7(6), 739–767, 1992.


Heinke, C.: Simulating oceanic CaCO$_3$ export production

Biogeosciences, 3, 1–13, 2006 www.biogeosciences.net/bg/3/1/
T. Rixen et al.: Diatom and their role in the Arabian Sea upwelling system

2002.


