Sedimentary phosphorus record from the Oman margin: New evidence of high productivity during glacial periods

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[1] The northern region of the Arabian Sea is one of the biologically most fertile regions of the world oceans, with present productivity rates varying between 150 and 2500 mgC/m² × day [Madhupratap et al., 1996]. This is related to the influence of the southwesterly summer monsoon which causes vigorous upwelling along the Oman margin. Upwelling ceases during northeasterly winter monsoon activity; productivity rates, however, remain relatively high (about 800 mgC/m² × day), related to deep water mixing [Madhupratap et al., 1996]. The goal of this study is to verify if during the last glacial period, a period in which winter monsoon conditions prevailed, productivity rates were similarly high. With an analysis of phosphorus phases, stable nitrogen isotopes, organic matter content, and bulk mineralogy of the upper 10 m of the cores of ODP Hole 724C (corresponding to the last 140,000 years, sample resolution is ~5 kyr), we provide new evidence of high productivity during this last glacial period (marine isotopic stages 2, 3, and 4). This was probably related to the combined effect of (1) increased eolian input of iron-containing dust due to dryness on the adjacent continent and stronger winter monsoon, and (2) regeneration and diffusion of dissolved phosphorus from the sediments to the water column due to variations in the position and intensity of the Oxygen Minimum Zone. These findings suggest that there is no one-to-one relationship between summer monsoon activity and productivity, which emerges to be a quasi-persistent phenomenon across glacial and interglacial stages. INDEX TERMS: 4267 Oceanography: General: Paleoceanography; 4805 Oceanography: Biological and Chemical: Biogeochemical cycles (1615); 4845 Oceanography: Biological and Chemical: Nutrients and nutrient cycling; 4851 Oceanography: Biological and Chemical: Oxidation/reduction reactions; KEYWORDS: phosphorus, Oman margin, monsoon, productivity

1. Introduction

[2] Modern and ancient upwelling regions have been shown to have an impact on the carbon cycle [Heinze et al., 1991; Raymo, 1994; Goddéris and François, 1995], and thus on climate. Estimates of global primary production in oceans indicate that the contribution of upwelling regions amounts to 4.8 Gt C/year, representing 67% of total new production [Chavez and Toggweiler, 1995]. New production supplied by monsoon-related upwelling regions (1.5% of world’s ocean surface) represents about 5.5% (0.4 Gt C/year) of global primary production.

[3] The Oman margin is located south of the Arabian peninsula and constitutes the northwestern side of the Arabian Sea (Figure 1). This region is influenced by the Indian Ocean Monsoon, which is characterized by a seasonal reversal of the winds, coupled with an alternation of dry and wet conditions on the mainland [Webster et al., 1998]. Intermediate waters along the Oman margin are mainly formed by the outflow of warm, O₂-depleted and nutrient-rich waters from the Red Sea, while contributions from the Persian Gulf mostly influence shallower waters (<300 m). The extent of these outflows is controlled by monsoon intensity, with winter-summer changes in wind direction affecting the amount of nutrients (P and N) exported to the Indian Ocean [Bethoux, 1988].

[4] During summer monsoon, which dominates during full interglacial periods, in response to the injection of nutrients from below the thermocline, modern primary productivity in surface waters averages >500 mgC/m² ×
day between May and October, but drops to about 150 mgC/m² \times \text{day} during the rest of the year (according to Prell and Nitsuma [1989], but see below). Stratification of the water masses, sluggish circulation and decomposition of organic matter produce an Oxygen Minimum Zone (OMZ) between 200 and 1200 m, with O₂ concentrations of organic matter produce an Oxygen Minimum Zone situation is assumed to be dominating during glacial periods [1996]. Along with these processes, major changes in oceanic winter and, possibly, glacial periods [1998] can concur in sustaining primary production during winter deep water mixing down to 800 m [cooling of surficial waters and consequent sinking. This

Figure 1. ODP Site 724 location in the Oman margin (dot). The Oman margin upwelling area is marked in light gray and arrows show wind directions during summer (solid) and winter monsoon (open).

2. Material and Methods

[8] The data presented here are from ODP Leg 117, Hole 724C (Oman Margin, Figure 1). The age model used in this study is based on the isotopic stratigraphy provided by Zahn and Pedersen [1991]. Hole 724C (Oman Margin, 18°27'. 713°N, 57°47.147'E, 592.8 m water depth) was sampled from 0 to 10.27 mbf, at a time resolution of about 5 kyr, yielding a total of 26 samples. These sediments consist of dark calcareous clayey silt, moderately bioturbated throughout, with abundant well-preserved nanofossils and foraminifers. The sedimentation rates for the studied sequence are relatively low and range between 4 and 17 cm/kyr. No evidence of redeposition and/or winnowing was detected [Prell et al., 1989].

[9] All samples, of approximately 10 c³ each, were oven-dried at 50°C and divided in subsamples. All analyses were performed at the GEA laboratory of the Geological Institute in Neuchâtel, except for ICP-AES and nitrogen stable isotope analyses.

[10] 5 g of dried sediment were ground to obtain a homogeneous powder with particle sizes <40 μm. An aliquot of this powder was pressed (20 bars) to determine bulk sediment mineralogy by XRD (SCINTAG XRD 2000 Diffractometer) based on a semi-quantitative estimation, and using external standards [Kübler, 1987]. The relative error of the bulk rock mineralogy is about 5% (RSD).

[11] The characterization of organic matter was performed on about 100 mg of dried and ground sediment, with a Rock-Eval 6, using the standard whole rock pyrolysis method [Espitalié et al., 1986; Lafargue et al., 1996].

[12] Nitrogen contents and stable isotopic analyses of the bulk sediment were performed at the Stable Isotope Laboratory, ETH-Zentrum, Zürich, using a Carlo Erba CNS2500 CHN Elemental Analyzer coupled with a Fisons OPTIMA mass spectrometer. Mean errors, calculated as the relative standard deviations on measurements are about 0.15‰. All data are reported in the conventional δ notation with respect to atmospheric nitrogen (AIR%o). Sample replicates have been analyzed to assess the validity of the data.

[13] Total nitrogen is here considered as representing the organic nitrogen content. To justify this assumption, our data were plotted on a TOC wt% versus Tot. N wt% diagram (Figure 2). The x-intercept of the calculated regression line has a negative value, indicating that when organic matter is lacking in the sediments, no significant inorganic component of nitrogen is present [Hedges et al., 1988]. Therefore we have analyzed bulk sediment for δ¹⁵N estimation, avoiding acidification of the sample, which
seems to attack part of the organic matter, resulting in the loss of up to 50% of Tot. N \[Lohse et al., 2000\]. Because of all diagenetic processes which influence the regeneration and redistribution of dissolved and reactive P in the uppermost sediments \[Broecker and Peng, 1982; Froelich et al., 1982; Ingall and Jahnke, 1994; Jarvis et al., 1994; Krajewski et al., 1994; Follmi, 1996; Ruttenberg and Goni, 1997\], it is critical to distinguish between different phosphorus sedimentary phases. Therefore we have used a 4-step sequential extraction technique, adapted from the SEDEX method \[Ruttenberg, 1992; Anderson and Delaney, 2000\]. This method utilizes progressive dissolution of solid phases in four steps to extract P associated with well defined sedimentary components: (1) iron and manganese oxihydroxides and P loosely absorbed on mineral surfaces, (2) authigenic minerals (carbonate fluorapatite (CFA)) and phosphorus associated to fish debris, to calcium carbonate, and smectite, (3) detrital material, so phosphorus associated to igneous, methamorphic and sedimentary apatites, and (4) organic matter \[(1) [Ruttenberg, 1992]. During the first step of this extraction, because of the reduction of Fe by dithionite and subsequent complexation by citrate, we are able to separate also ferric Fe, referred as \(Fe_{CBD}\) (see Table 1).

\[15\] Shimmield and Mowbray \[1991\] analyzed major elements (e.g., Ti, Al, Ca, P, and Fe) of sediments from Site 724. Because these ratios are independent of accumulation rates, we will compare them to our mineralogical and geochemical data to improve our interpretation.

3. Results and Discussion

3.1. Mineralogy

\[16\] Calcite, phyllosilicates, quartz, and dolomite contents are shown in Figure 3. Detrital silicates, calculated as the sum of quartz, phyllosilicates, plagioclase and K-feldspar, account on average for 30% (around 40% during glacial periods), of the sediments of the Oman core \[Figure 3; K-feldspar and plagioclase never exceed 5% of the detritic fraction and are not shown]. This relatively high percentage compared to other sites drilled during Leg 117, is probably due to the proximity of the coast. Because of its higher abundance and mass accumulation rates (MAR; Figures 3 and 4) during arid periods (glacial stages 4 and 2, and stage 3), and due to the absence of major rivers, the detrital material is considered as predominantly eolian with only

Figure 2. Correlation between C\(_{org}\) (wt\%) and N\(_{tot}\) (wt\%) from Site 724. The intercept of the regression line on the x axis when C\(_{org}\) = 0 is negative (y = 0, x = −0.008), indicating that no important inorganic nitrogen fraction is present. This finding has allowed to avoid acidification. Tests carried out on several samples treated with HCl at different concentrations (10 and 30%), show, indeed, that decarbonation using HCl has a strong and somehow unpredictable effect on nitrogen isotopes. Nitrogen isotopic signal of those samples shows a change ranging between 0.02 and 4.4\%, with no apparent relation to the strength of the acid.

Table 1. Sequential Extraction Technique Applied for Phosphorus Phases Determination

<table>
<thead>
<tr>
<th>Step Name</th>
<th>Treatments</th>
<th>P Component Isolated</th>
<th>Errors, %</th>
<th>Detection Limits</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iron-bound P(_e)</td>
<td>10 ml CBD solution (6 hours) (0.22 M sodium citrate, 0.11 M sodium bicarbonate, 0.13 M sodium dithionite) 10 ml 1M MgCl(_2) (2 hours) 10 ml H(_2)O (2 hours)</td>
<td>exchangeable or loosely sorbed P reducible or reactive iron-bound P PLUS Ferric Fe</td>
<td>3–6</td>
<td>0.03</td>
</tr>
<tr>
<td>Authigenic P(_f)</td>
<td>10 ml 1M Na-acetate buffered to pH 4 with acetic acid (5 hours) 10 ml 1M MgCl(_2) (2 hours) 10 ml 1M MgCl(_2) (2 hours) 10 ml H(_2)O (2 hours)</td>
<td>carbonate fluorapatite (CFA) biogenic hydroxyapatite</td>
<td>2–7</td>
<td>0.025</td>
</tr>
<tr>
<td>Detrital P(_f)</td>
<td>10 ml 1N HCl (16 hours)</td>
<td>detrital fluorapatite-bound P</td>
<td>3–5</td>
<td>0.003</td>
</tr>
<tr>
<td>Organic P(_f)</td>
<td>1 ml 50% (w/v) Mg(_{NO_3}), dry in low oven, ash at 500°C (2 hours) 10 ml 1N HCl (24 hours)</td>
<td>organic-bound P</td>
<td>2–5</td>
<td>0.004</td>
</tr>
</tbody>
</table>

\[a\]From Ruttenberg \[1992\] and Anderson and Delaney \[2000\].
\[b\]Concentrations of all phases are expressed in mg/g of sediment.
\[c\]Errors for each phase determination are considered as the relative standard deviation calculated on replicate analyses of consistency standards.
\[d\]Detection limits are considered as 3 times relative standard deviation calculated on replicate analyses of blank solutions.
\[e\]Iron-bound phosphorus and ferric Fe concentrations were measured using an OPTIMA 3000 Perkin Elmer ICP-AES, at the Laboratoire des Eaux et de l’Environnement in Peseux, Switzerland.
\[f\]The other phases were measured using a Perkin-Elmer UV/Vis spectrophotometer Lambda 10 with a 5-cm quartz cell. Sample solutions were diluted in distilled water and a color developing agent was added to the solution, following the ascorbic acid method for phosphates \[Eaton et al., 1995\].
minor fluvial contributions from the Arabian peninsula. During interglacial stage 5, both quartz and phyllosilicates show lower abundances (13% and 7% in average respectively) than during glacial stage 2 and stage 3 (15% and 9%). Prominent peaks are present at the end of stage 4 and during stage 2. Despite the similarity between general trends, the correlation between quartz and phyllosilicates is poor ($r = 0.48$), probably due to a different source of the material and/or to differentiation during transport.

[17] Detrital MAR (Figure 4) are low during interglacial 5 and show a 4-fold increase during glacial stages 4 and 2 and stage 3. The most prominent peak is observed during glacial stage 2, where it reaches about 104 mg/cm$^2 \times$ kyr. This observation confirms the correlation between land-derived and eolian-transported dust MAR with aridity [Krissek and Clemens, 1992], implying maximum dust transport during glacial times. The Ti/Al elemental ratio calculated for Site 724 [Shimmield and Mowbray, 1991] and interpreted as an indicator of aridity and a proxy of monsoon intensity, shows relatively higher values and higher variability during glacial periods, supporting the hypothesis of enhanced dust input deduced from our mineralogical data (Figures 3, 4 and 5).

[18] Dolomite shows the same trend as other detrital indices, with an increase to an average of about 9% during stages 4-2. The correlation with phyllosilicates is rather good ($r = 0.64$). The origin of dolomite is eolian, and its source area is the Northern Arabian Peninsula and Oman, where Permian-Triassic and Cretaceous dolomitic series extensively outcrop [Krissek and Clemens, 1991; Sirocko et al., 1993].

[19] The Oman margin sediments are characterized by high percentages (average of 40%) of mostly biogenic calcite (foraminifers and nannofossils are well preserved in the upper part of the sedimentary sequence) [Prell and Niitsuma, 1989], which resembles the general profile of the $\delta^{18}$O record. Values are higher during interglacial stages 5 and 1, with peaks of about 50% at about 100 ka and 60%
at 8 ka (Figure 3). The calcite record shows an upward-decreasing trend starting at about 90 ka, and reaches a minimum of about 30% at the top of stage 2.

[20] The relative importance of marine versus continental detrital input was higher during interglacial stages, as shown by the calcite record and the Ca/Ti curve (Figures 3 and 5) [Shimmield and Mowbray, 1991]. Variations in the calcite record probably stem from detrital dilution, and they may be interpreted as reflecting changes in aridity on the continent [Weedon and Shimmield, 1991]. Calcite MAR are, conversely, higher between stage 4 and 2 (Figure 4), possibly suggesting that a proportion of calcite is of detrital origin, as proposed by Sirocko [1989].

3.2. Phosphorus

[21] Phosphorus phases represent only a small fraction of the total sediment (about 0.1%), but relative variations between the phases are important. Authigenic phosphorus represents the most abundant form, on average 75% of total phosphorus, while detrital and Fe-bound phosphorus represent only 18% and 5% respectively (see Table 2 and Figures 6a and 7). The three reactive phases, Fe-bound, authigenic and organic-bound P, significantly increase after stage 2 and during the Holocene.

[22] Phosphorus extracted during the second step of the SEDEX method (see Table 1) mainly accounts for authigenic Ca-apatite and fish debris. However, the SEDEX procedure does not allow to distinguish between these two components. Schenau et al. [2000], who employed a six-steps sequential extraction technique, estimated fish debris content in the Oman margin sediments to be more than 3 times lower than authigenic minerals. Sedimentary detritalapatite might also have been transported along with other detrital material to the Oman margin site. However, as phosphorus is generally tightly bound in the structure of sedimentary apatite [Compton et al., 2000], it is most likely

![Figure 5. Major elemental ratios from Site 724 [Shimmield and Mowbray, 1991].](image)

Table 2. Mean Concentrations and Percentages of Phosphorus Phases From Site 724, Oman Margin*

<table>
<thead>
<tr>
<th></th>
<th>Fe-bound P, mg P/g sediment</th>
<th>Authigenic P, mg P/g sediment</th>
<th>Detrital P, mg P/g sediment</th>
<th>Organic-bound P, mg P/g sediment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean interglacial</td>
<td>0.0719</td>
<td>1.3668</td>
<td>0.1761</td>
<td>0.0415</td>
</tr>
<tr>
<td>Mean glacial</td>
<td>0.0523</td>
<td>0.8462</td>
<td>0.2134</td>
<td>0.0284</td>
</tr>
<tr>
<td>Minimum value</td>
<td>0.0410</td>
<td>0.2750</td>
<td>0.1290</td>
<td>0.0130</td>
</tr>
<tr>
<td>Average</td>
<td>0.0596</td>
<td>1.0390</td>
<td>0.2000</td>
<td>0.0330</td>
</tr>
<tr>
<td>Maximum value</td>
<td>0.1826</td>
<td>3.5820</td>
<td>0.2580</td>
<td>0.1500</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>P phases % on total P</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Interglacial average</td>
<td>4.34</td>
<td>82.52</td>
<td>10.63</td>
<td>2.51</td>
</tr>
<tr>
<td>Glacial average</td>
<td>4.59</td>
<td>74.21</td>
<td>18.71</td>
<td>2.49</td>
</tr>
</tbody>
</table>

* Differences between glacial and interglacial percentages are not significative for Fe- and organic-bound P. They are relatively important for detrital P and for authigenic P.
Despite the lack of pore water analyses data, previous studies performed on cores from the Arabian Sea region support the hypothesis that most of the P extracted during the second step of the SEDEX method, is authigenic apatite. Recent studies on pore water and solid phases of phosphorus clearly indicate that phosphorite formation is an ongoing process in sediments around the Arabian Sea, and along the Oman margin [Schenau et al., 2000]. Low pore water phosphate levels in the upper portion of Site 724 C (lower than 4 μmol PO₄/l) recorded onboard during Leg 117 were already interpreted as an indication of apatite precipitation [Prell and Niitsuma, 1989]. The relatively high P/Ti ratio values at Site 724 have also been explained as reflecting in-situ apatite precipitation [Weedon and Shimmield, 1991]. An indication of authigenic apatite precipitation is also given by the comparison between the percentages of the different phosphorus phases, which reveals a “sink switching” (Figure 6). The organic-bound P profile shows a down-core decreasing trend (Figure 6b), followed also by Fe-bound P. This trend is paralleled by the increase of authigenic P percentages (Figure 6a), indicating that this phase is forming at the expense of organic and Fe-bound P. The high-concentrated phosphorus sedimentary phases. Authigenic and detrital P are the most abundant. (b) Fe-bound and organic-bound P show a down-core decreasing trend, while authigenic P is increasing, revealing a sink switch. However, these two phases may not explain alone the authigenic P increase (see text).

Figure 6. (a) Percent of total P in the analyzed phosphorus sedimentary phases. Authigenic and detrital P are the most abundant. (b) Fe-bound and organic-bound P show a down-core decreasing trend, while authigenic P is increasing, revealing a sink switch. However, these two phases may not explain alone the authigenic P increase.

Figure 7. Percentage of G. bulloides from Site 723 [Anderson and Prell, 1991]; concentrations of Fe-bound P, authigenic P, detrital P, organic-bound P and Fe(CBD), from Site 724. The Fe/P(CBD) molar ratio is presented along with Fe-bound P. The different phases are defined in the text. Horizontal lines and isotopic stages are as in Figure 3.
concentrations of the authigenic P recorded in the upper part of the core (Figure 6), and also found in nearby cores [Schouen et al., 2000], might be explained by the presence of an F-poor and P-bearing authigenic mineral, representing the first step of Ca-apatite precipitation [Krajewski et al., 1994].

[25] Authigenic phosphorus, which shows a remarkable correlation with the oxygen stable isotope and calcite records (r = 0.79 and 0.60 respectively), presents higher values during interglacial stages than during glacial stages. From average values of about 1.5 mg P/g during stage 5, it decreases to 0.3 mg P/g during stages 4 and 2, and averages 1 mg P/g during stage 3 (Table 2 and Figure 6). The P/Ti ratio at Site 724 [Shimmield and Mowbray, 1991], which correlates in trend with the authigenic P curve presented in this study (Figures 5 and 7), shows lower values during stages 4-2. Biogenic calcite (e.g., foraminiferal shells) may provide nucleation sites for CFA, and this could explain the good correlation between these two parameters. However, the presence of carbonatic sheels in sediments cannot account alone for apatite precipitation, as other chemical parameters are more important (e.g., phosphate and fluoride concentrations in porewater, alkalinity).

[26] Fe-bound P shows concentrations lower than 0.05 mg P/g, except in stage 1, when it reaches values up to 0.2 mg P/g (Figure 6). It is plausible that in the youngest samples, phosphorus extracted during the first step of the SEDEX method, accounts also for adsorbed P on clay and mineral surfaces (see Methods). In fact, during this interval, the Fe/P(CBD) ratio presents values close or even lower than 1 (Figure 7), suggesting that part of phosphorus extracted during this step clearly accounts for loosely sorbed phosphorus [Ruttenberg, 1992]. Ferric iron (Fe(CBD)) follows the overall trend of detrital indices (r = 0.62 with detrital P), with a two-fold increase during stage 3 and glacial stage 2 compared to interglacial stages. Highest values are on the order of 0.8 mg Fe/g. Lack of correlation between Fe(CBD) and Fe-bound P (r = 0.46) may be due 1) to the nature of the Fe phases, and 2) to changes in environmental conditions at the sediment/water interface. The correlation between Fe(CBD) and other detrital parameters reflects the eolian origin of iron, and may suggest that iron is present in the sediments under a relatively well-crystallized form (i.e., goethite, hematite). These minerals, which are dissolved during the CBD step, constitute a less efficient sink for dissolved P than amorphous Fe oxihydroxides [see Slomp et al., 1996b]. Furthermore, the development of oxygen depleted conditions in the upper part of the sediments, might have reduced the thickness or even removed the oxidized sediment layer, and inhibit the adsorption of dissolved phosphorus by Fe-oxides [Glenn, 1990; Slomp et al., 1996b]. Sirocco et al. [2000] provided geochemical evidences of extremely low oxygen levels, especially during the Last Glacial Maximum (LGM), along the Oman margin. Lucotte et al. [1994] used the Fe/P(CBD) ratio to infer paleoredox conditions in bottom waters and sediments from the Labrador Sea. At Site 724, this ratio shows the highest values during glacial periods (Figure 7), suggesting, indeed, low oxygen conditions [Lucotte et al., 1994]. However, considered the detrital nature of Fe oxides, these values should be taken with caution.

[27] Detrital P (igneous, metamorphic, and sedimentary detrital apatite) is correlated to quartz (r = 0.65 and 0.67 during glacial times), with peaks during stages 4-2 of about 2.5 mg P/g and minima during interglacial stages 5 and 1. The lack of a strong correlation with other detrital indices such as phyllosilicates (r = 0.33), may indicate an alternate source for these detrital components. Detrital P MAR show the same trend of detrital silicates MAR, with higher values during stages 4-2, reaching maxima of about 3.5 mg/cm² × kyr during glacial stage 2 (Figure 4).

[28] Organic-bound P exhibits extremely low values, around 0.02 mg P/g, along the entire sequence and a sharp and pronounced increase at the beginning of stage 1. Higher variability is noticed during stage 3, and a more prominent spike, with values up to 0.05 mg P/g, is present at the beginning of stage 2.

[29] Reactive phosphorus percentages (the formerly biologically available phase, calculated as the arithmetic sum of Fe-bound, authigenic and organic-bound phosphorus) are lower during glacial periods, following the authigenic P trend, as this phase is the most abundant. This phase, in fact, diminishes about 10% during glacial periods (Figure 6). Total P, on the other hand, does not show important glacial-interglacial variations, but, as detrital P is not biologically available, it is questionable if total P might help in understanding the impact of changes in reactive P input on productivity.

[30] Reactive phosphorus MAR show a singular trend, compared to other proxies: values generally range between 10 and 15 mg/cm² × kyr, and they show no particular increasing trend due to higher sedimentation rates (Figure 4). Besides the Holocene increase of all the reactive phases of phosphorus, only two significant peaks are documented, at 70 ka and at 20 ka, both at the beginning of glacial periods. Otherwise it seems that reactive P fluxes to the sediments were fairly constant throughout the investigated period, unlike all the other measured parameters. Fe(CBD) MAR match the detrital MAR trend, with higher values during stages 4-2, and show a significant increase between 70 ka and 20 ka, where they reach values of 15 mg/cm² × kyr.

3.3. Organic Matter

[31] Immature marine organic matter is the primary component of organic material present at Site 724. This is suggested by the Tmax (Tmax = 420°C in average) and the hydrogen index (HI) values. HI values are generally higher than 150 mg HC/g total organic carbon (TOC), with few points above 400 mg HC/g TOC (Figure 8a). These data are in agreement with previous analyses [Bertrand et al., 1991]. When Rock Eval analyses are performed on bulk sediments, as in this case, low values of HI could indicate a strong mineral matrix effect [Espitalié et al., 1986]. The occurrence of a rock-matrix effect and the uniform origin of the organic matter are corroborated by the S2 versus Corg graph (Figure 8b). The regression line in a S2 versus Corg diagram should have a 0 intercept, if no mineral matrix is present, while a positive x axis intercept (as observed here, Figure 8b) generally indicates adsorption by minerals, with clays representing the main agent of adsorption [Langford and Blanc-Valleron, 1990]. The samples, linearly distrib-
uted, show a high correlation \( r^2 = 0.95 \), indicating a coherent group. If the regression line is shifted towards the origin to eliminate the rock matrix effect [Langford and Blanc-Valleron, 1990], all the sample fall in the type II kerogen field, confirming the marine origin of the organic matter, with an average HI of 414, as given by the slope of the regression line.

[32] Average value of organic carbon from Site 724 is about 0.8 wt% (Figure 9), but the record exhibits large oscillations, especially during the last 60 kyr, with values up to 2 wt%. Higher organic carbon concentrations and accumulation rates are recorded during stage 3 and during glacial stage 2, with two major peaks between 30 and 45 ka and at 20 ka, during the LGM. High organic matter contents are also present in the Holocene part of the sequence.

[33] Organic carbon MAR show a high flux of organic matter during stage 3 and during glacial stage 2, while the lowest fluxes are recorded during interglacial 5. It is unclear if this trend is a real feature or represents the effect of higher glacial sedimentation rates (Figure 4) and consequent better preservation [Canfield, 1993]. The above relation was already observed by Murray and Prell [1991] for Site 722 on the Owen Ridge, but, despite their attempt of removing the possible effect of sedimentation rate, they found that greater fluxes of organic material were still related to glacial or winter-monsoon dominated periods [Murray and Prell, 1991]. Moreover, enhanced organic matter burial during

Figure 8. (a) HI versus Tmax diagram. All the points, representing samples from Site 724, fall in the type II kerogen field (marine organic matter). (b) S2 versus Corg diagram of the data from Site 724. Boundaries between the three fields are defined by Langford et al. [1990]. The slope of the regression line and the high degree of correlation confirm the marine origin of the organic matter. (c) Corg versus OI diagram. Samples with high organic matter content present low OI, indicating enhanced preservation due to deposition in a low oxygen environment.
Figure 9. Concentration of Corg from Site 724; OI diagram, as determined by Rock Eval analysis; $^{15}$N of bulk sediment from Site 724; $^{13}$Corg [Muzuka et al., 1991] and $^{13}$C Uvigerina spp. [Zahn and Pedersen, 1991] from Site 724. Horizontal lines and isotopic stages are as in Figure 3.

Figure 10. $^{18}$O data of G. ruber from Site 724 [Zahn and Pedersen, 1991]; C/Porg, C/P_{reac.}, and C/N molar ratios, as defined in the text. Horizontal lines and isotopic stages are as in Figure 3.

Glacials (see peaks at 20 and at around 40 ka, Figure 4) correlates with low $^{13}$C Uvigerina spp. values (Figure 9), an endobenthic foraminifer, whose low carbon isotopic signature indicates higher supply and oxidation of organic matter in the sediments [Zahn et al., 1986].

$^{15}$N of the sediments shows a fairly good correlation ($r = 0.62$) with the organic carbon content: values are low during glacial stage 4 and the upper part of stage 2, with the lowest value of 1.2% reached during stage 4 (Figure 9); the two most prominent peaks are recorded during stage 3 ($^{15}$N = 7.4%) and at the top of the sequence, during the Holocene ($^{15}$N = 7.6%).

The trend, maxima and minima of nitrogen isotopes of Site 724 sediments are in agreement with other nitrogen records, which are interpreted as indicating denitrification. $^{15}$N values from this study are generally lower than the values observed in nearby sites in the Oman region [Altabet et al., 1999]. It is not clear why the magnitude of the signal ( ~ 6%) is higher than the one observed by Altabet et al. [1999].

The high degree of correlation between the $^{15}$N values and the organic matter content could, indeed, suggest the occurrence of denitrification, in situ or in remote areas. But the observed low values are difficult to decipher: one explanation could be the influence of terrestrial organic matter, but evidences from other proxies (e.g., $^{13}$Corg, HI, C_{org}/N ratio, Figures 9 and 10) seem to exclude this possibility. Light interglacial $^{15}$N values have been already observed in sediments from other regions where denitrification is known to occur (i.e., the Cariaco Basin) [Haug et al., 1998], and have been related to the imprint of nitrogen fixation by cyanobacteria, which lowered the nitrogen isotopic signal. Nitrogen fixation in the Indian Ocean and in the Red Sea is, indeed, an established phenomenon.
[Bethoux, 1988; Capone et al., 1997], which could have been fostered especially during glacial periods, when input of macronutrients and micronutrients (e.g., iron) was enhanced. So, generally low values are probably due to the imprint of nitrogen fixation, while relative changes may be attributed to different degree of denitrification.

[37] C/Porg and Corg/P elemental ratios are higher than the expected Redfield ratios for marine OM, indicating a strong depletion of P and a minor preferential loss of N during organic matter degradation (Figure 10). Again, these values could be explained by a larger input of terrestrial organic matter, but, as mentioned before, this is not supported by other data. Moreover, the lack of a positive correlation between light δ15N and δ13Corg data (interpolated from Muzuka et al. [1991]; see Figure 9) does not support this assumption.

[38] Relatively high elemental ratios in organic matter are thus considered as indicating organic material degraded in an oxygen-poor environment. The correlation of high C/Porg and Corg/N with lower OI (Figure 8c and 9), support this hypothesis [Kenig et al., 1994].

[39] Organic matter is normally considered as the principal carrier of phosphorus to the sediment, and, once degraded, constitutes the major source of dissolved reactive phosphorus of the pore water. Thus, the elemental ratio between Corg and reactive P (C/Preac) in Oman sediments should account for the diagenetic transformations of reactive P and explain the deviation of the C/Porg from the Redfield ratio assumed for marine organic matter (106:1) [Anderson and Delaney, 2001]. The C/Preac ratio generally shows values lower than the Redfield ratio, averaging 20, as it is observed for sediments with relatively low Corg contents (<2%) [see Anderson and Delaney, 2001]. Such low values indicate that reactive phosphorus exceeds organic carbon. The contrasting behavior of C/Porg and C/Preac during part of the studied sequence, especially stage 5 (Figure 10), when C/Porg increases while C/Preac shows low values, may be interpreted as another evidence of a “phase switching” between organic and authigenic P. Although part of the original organic carbon has also been oxidized and lost, it is clear that organic phosphorus cannot account alone for the C/Preac values, and an alternative source of dissolved reactive phosphorus needs to be considered. Schenau et al. [2000] have demonstrated that fish debris dissolution constitutes an additional and important source of dissolved reactive phosphorus for modern sediments of the Oman margin. C/Preac values ranging between 30 and 60 are observed only between 45 and 35 ka and between 20 ka, when also the C/Porg ratio presents high numbers (Figure 10). These two peaks could be explained by either a reduced phosphorus input to the system, or an increased C preservation in the sediments relative to P, or a reduced phosphorus retention in the sediments, compared to interglacial periods. During stage 3, the co-occurrence of relatively high δ13C with the peak in the C/Preac ratio (Figures 9 and 10), may indicate low oxygen content of the bottom waters, which likely prompted the release of dissolved phosphorus to the water column. This is true especially if we assume that organic P regeneration from organic matter is enhanced under low oxygen conditions relative to oxic bottom waters [Ingall and Jahnke, 1994]. In fact, enhanced regeneration of nitrogen and phosphorus during anaerobic oxidation of the organic matter has been documented in sediments from the northeastern Arabian Sea [Lueckge et al., 1999]. During the LGM, low oxygen levels may have persisted [see Sirocko et al., 2000] and promoted the release of dissolved reactive P.

4. Implications

[40] The results obtained from the multiproxy study of sediments from the Oman margin have revealed three different situations as the response to changed monsoon strength, sea level, and oceanic circulation for the last 140,000 years (Figure 11).

[41] During full interglacial periods (i.e., lower part of stage 5 and stage 1; Figure 11a), characterized by high sea level stands and strong summer monsoon activity, the Oman margin showed high biological production rates, connected to increased upwelling intensity. G. bulloides % (upwelling indicator) and carbonate content are high. The enhanced upwelling and the input of intermediate waters from the Red Sea contributed to maintain high nutrient levels. However, low organic matter accumulation rates seem to indicate that productivity and/or preservation were not enhanced compared to glacial periods.

[42] Oxygen levels were probably low, as expected in high productivity regions and with the presence of an OMZ, but it is not clear from our data (see δ15N, OI) if severe anoxic conditions developed. In fact, increased and intense upwelling could have possibly brought oxygen-rich waters from below the OMZ to shallower depths, decreasing therefore its intensity [Prell and Niitsuma, 1989]. Low δ15N during interglacial 5 could indicate effective nitrogen fixation by cyanobacteria, if nutrients other than nitrogen were provided to surface waters. Extensive oxidation of organic matter (as indicated by C/Porg and OI, Figures 9 and 10) and dissolution of fish debris regenerated high quantities of dissolved P that could in part precipitate as authigenic P and in part be upwelled to the surface water, enhancing productivity.

[43] Low detrital contents of the sediments during stage 5 are interpreted as the result of prevailing humid conditions on the mainland [Burns et al., 1998]. The ITZC (Intertropical Convergence Zone) was, in fact, located inside the Arabian peninsula and northwesterly winds could not reach the investigated site. Apparently, the increased humidity did not develop a coastal riverine system, which could enhance nutrient and dissolved phosphorus input. But during warm interglacials a continental input of nutrient to the ocean is not likely to have occurred. There is no evidence of any kind of terrestrial contribution to the organic pool, and proxies, such as the HI index (Figure 8), indicate an autochthonous origin of the organic matter.

[44] During glacial stages 4 and 2, and during stage 3, the ITZC shifted southward and arid conditions prevailed on land, promoting eolian uptake of material (Figures 11b and 11c) [Burns et al., 1998]. Northwesterlies, now blowing more southwardly, along with persistent and stronger northeasterlies contributed to high detrital input from the
Figure 11. Schematic representations of the oceanographic and climatic situation during the last 140,000 years along the Oman margin. Generally low nitrogen stable isotopes are interpreted as indicating nitrogen fixation in surface waters, in response to quasi-persistent high nutrient levels. (a) During full interglacial periods biological production rates are relatively high, due to increased upwelling intensity and input of nutrient-rich waters from the Red Sea. Productivity and/or preservation of OM were not enhanced compared to glacial periods. (b) Relatively higher nitrogen stable isotopes values might reflect denitrification. Deep water mixing might be responsible for providing nutrients to surface waters, especially during weak summer monsoon periods. (c) The enhanced eolian input introduced high amounts of nutrients (e.g., Fe and possibly P, see text), which sustained productivity. Changes in sea level deeply influenced the circulation patterns of the region.
continent, as indicated by all the detrital indices (i.e., quartz, phyllosilicates, this study, as well as Ti/Al, Fe/Ti) [Shimmield and Mowbray, 1991]. Despite their increased strength, winter monsoons were too weak to produce any substantial upwelling [Madhupratap et al., 1996]. But several proxies, such as organic matter, nitrogen stable isotopes (this study) and $^{13}$C Uvigerina spp. (Figure 9) [Zahn and Pedersen, 1991] clearly point out that productivity and burial of organic matter were enhanced. Winter monsoon dryness has been called as the factor causing high nutrient content; in fact, during winter time, sinking of cooler and denser surface waters provoked an upward flux of deep and nutrient-rich waters [Madhupratap et al., 1996].

Stage 3 is characterized by a reduced summer monsoon-induced upwelling, and generally arid conditions were persistent on the continent [Burns et al., 1998]. The high sea level fostered the input of oxygen-depleted intermediate waters from the Red Sea and persistent winter monsoons introduced high amounts of detrital material and micronutrients, such as iron (Figures 4, 5 and 6). One question to be answered here is whether reactive phosphorus concentrations in surface waters were high during glacial periods (stages 4-2). The indications we get from our data suggest that regeneration of dissolved phosphorus might have happened, and that part of this phosphorus might have been brought to surface waters. But more than that, it is reasonable that the input of reactive P from the continent might have been enhanced. In fact, “continental” Fe oxyhydroxides, an important component of eolian dust, which increased during glacial periods (Figure 7), generally bear considerable amounts of phosphorus adsorbed on their surfaces. Once in the ocean, photochemical reactions in the surface waters (i.e., photoreduction) [Johnson et al., 1994] may dissolve part of the Fe oxyhydroxides, and, hence, liberate also the associated phosphorus (K. C. Ruttenberg, personal communication, 2002). This process, along with the above mentioned evaporation-induced mixing process, might have been responsible for maintaining high nutrients levels along the Oman margin (Figures 11b and 11c).

Weaker upwelling along with persistent high productivity, probably enhanced the intensity of the OMZ, and stronger depletion of oxygen in bottom waters led to high organic matter accumulation and preservation, as indicated by $C_{org}$ MAR and OI (Figures 4 and 9). Nitrogen isotopes are highest and show strong correlation with organic matter, hence, liberate also the associated phosphorus [K. C. Ruttenberg, personal communication, 2002]. This process, along with the above mentioned evaporation-induced mixing process, might have been responsible for maintaining high nutrients levels along the Oman margin (Figures 11b and 11c).

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Weak upwelling along with persistent high productivity, probably enhanced the intensity of the OMZ, and stronger depletion of oxygen in bottom waters led to high organic matter accumulation and preservation, as indicated by $C_{org}$ MAR and OI (Figures 4 and 9). Nitrogen isotopes are highest and show strong correlation with organic matter, hence, possibility an indication of denitrification. Degradation of organic matter regenerated high amounts of dissolved reactive phosphorus, while organic carbon was preserved in the sediments [Ingall and Jahnke, 1994]. However, the extension of the OMZ during stage 3 and the consequent oxygen demand probably provoked unfavorable conditions to authigenic P precipitation, as indicated by low values of authigenic P, and high $C_{org}/P$ ratios (Figures 6 and 10). A diagenetic model developed for phosphorus cycling and applied to results from box cores from the Oman margin [Slomp et al., 1996a; Schenau et al., 2000], confirms that without bioturbation and in intense oxygen-depleted waters, reactive phosphorus is easily regenerated at the water-sediment interface and can re-enter the water column. A direct correlation among high sedimentation rates, low oxygen content, high organic carbon preservation and high phosphorus loss from the sediments might be forwarded for this period [Coiman and Holland, 2000].

During glacial stages 4 and 2, arid conditions on the continent [Burns et al., 1998] allowed for high eolian input of detrital material and iron- (and phosphorus) containing dust. The well-crystallized iron oxides did not provide an efficient sink for dissolve P, which was probably not retained in the sediments. Glacial lowering of sea level reduced or even cut off the contribution from the Persian Gulf and the Red Sea, allowing waters from remote sources, possibly AAIW (Antarctic Intermediate Water) to spread northward (Figure 11c) [Kallel et al., 1988].

Glacial mass accumulation rates of organic carbon were high, especially during stage 2, promoting low oxygen conditions [Sirocko et al., 2000]. Peaks in both $C_{org}$ and $P_{org}$ indicate, indeed, a loss of dissolved P from the sediments, compared to interglacial periods (Figure 10). This implies that during glacial periods P release from the sediments was most likely enhanced and regenerated P, along with nutrients and micronutrients introduced from the mainland, could have sustained high rates of productivity.

5. Conclusions

This multiproxy approach to the study of Site 724 allows for the explanation of different aspects of the climatic and oceanographic history of the Oman margin since interglacial 5.

During the last 140,000 years, the Oman margin was characterized by an extremely complex situation, where regional and global processes were coupled and interacted. Changes in monsoon intensity, due to glacial-interglacial variations in insolation strength, did combine with different oceanic circulation modes, imposed by glacioeustatic changes in sea level, and both controlled sedimentation, nutrients distribution and primary production (Figure 11).

Along with the other used proxies, the study of the different P phases stored in the Oman sediments brought important elements, because their variations are the product not only of the input from the mainland, but also of changes in physical and chemical conditions at the sea bottom, produced by changes in productivity, oceanic circulation and glacioeustatism. During the last 140,000 years the overall flux of P to the Oman margin sediments seemed not to have changed significantly; despite this, its redistribution and regeneration, both controlled by changed conditions in the sediments, as deduced by comparing reactive phosphorus and organic carbon records, have certainly played an important role in sustaining productivity.

During full interglacial stages (stages 5 and 1), with strong summer monsoon inducing intense upwelling, productivity was sustained by the contribution of renewed nutrients from below the thermocline. Preservation was not enhanced because of persisting oxic or slightly suboxic conditions at the water-sediment interface. Phosphorus was regenerated during respiration of OM, and could partially precipitate as CFA. During glacial periods (stages 4-2), P was preferentially regenerated and rediffused to the water
column, because of low oxygen conditions in the bottom waters. Deep vertical mixing, along with eolian input of nutrients, helped in providing the renewed nutrients to the euphotic zone, sustaining productivity. [35] From this study, productivity along the Oman margin emerges to be a quasi-persistent phenomenon across glacial and interglacial stages, and seems to be controlled both by summer monsoon-induced upwelling and winter monsoon-induced deep vertical mixing. This has certainly important implications on the local and global carbon budget, especially on glacial-interglacial timescales. The hypothesis of high productivity along the Oman margin also during glacial periods could redefine the role of monsoon regions in controlling and enhancing carbon dioxide drawdown from the atmosphere into the ocean and in maintaining low CO₂ concentrations during glacials.

References
