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Rapid reorganization in ocean biogeochemistry off Peru towards the end of the Little Ice Age


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Abstract. Climate and ocean ecosystem variability has been well recognized during the twentieth century but it is unclear if modern ocean biogeochemistry is susceptible to the large, abrupt shifts that characterized the Late Quaternary. Time series from marine sediments off Peru show an abrupt centennial-scale biogeochemical regime shift in the early nineteenth century, of much greater magnitude and duration than present day multi-decadal variability. A rapid expansion of the subsurface nutrient-rich, oxygen-depleted waters resulted in the present-day higher biological productivity, including pelagic fish. The shift was likely driven by a northward migration of the Intertropical Convergence Zone and the South Pacific Subtropical High to their present day locations, coupled with a strengthening of Walker circulation, towards the end of the Little Ice Age. These findings reveal the potential for large reorganizations in tropical Pacific climate with immediate effects on ocean biogeochemical cycling and ecosystem structure.

1 Introduction

Paleoclimate evidence for large, rapid shifts in climate and biogeochemistry associated with Dansgaard-Oeschger events during the last glacial period indicate that the ocean-atmosphere climate system was susceptible to rapid changes in circulation and biogeochemical cycling (Schmittner et al., 2007). The Holocene appears more stable, however, large changes in tropical rainfall patterns at the Holocene thermal maximum (Haug et al., 2001), during the Medieval Climatic Anomaly (MCA) and the Little Ice Age (LIA) have been documented (Graham et al., 2007; Rein et al, 2004; Haug et al., 2001; Newton et al., 2006).
The LIA was associated with lower solar radiation and enhanced volcanic activity, resulting in large expansions of mountain glaciers in Europe and North America during the fifteenth to nineteenth centuries (Cronin et al., 2002; Broecker et al., 2000), and by drier conditions in the Northern Tropics (Haug et al., 2001). During the LIA, stronger northern trades may have resulted from enhanced latitudinal temperature gradients under the cooler Northern Hemisphere conditions, displacing southward the mean ITCZ position and its associated precipitation belt (Koutavas and Lynch-Stieglitz, 2004). Alternatively, large-scale cooling by solar and/or volcanic activity could have reduced the zonal temperature gradient in the Equatorial Pacific, again driving the marine ITCZ southward, particularly in the Eastern Pacific (Peterson and Haug, 2006; Mann et al., 2005).

Seasonal and interannual changes of the ITCZ and the trade winds are the main sources of biogeochemical variability in the Eastern Tropical South Pacific (ETSP), modulating biological productivity and oxygen concentrations (Barber and Chávez, 1983; Pennington et al., 2006). Here we examine the relationships between biogeochemical cycles and centennial-scale climate modes during the past ~700 years in the ETSP, and focus particularly on the transition between the LIA and the modern warmer climate. Given evidence for ecosystem changes associated with current warming trends (Field et al., 2006), the potential for crossing attractor thresholds or “tipping points” and undergoing large regime shifts (Mantua, 2004) is of major interest, particularly in regions that are heavily dependent on living marine resources.

We develop time series of multiple proxies from cores collected in the laminated sediments of the upper Peruvian margin to reconstruct the paleoceanographic and paleoecological conditions. High resolution sedimentary paleo-archives are preserved in certain topographic areas of the Peruvian margin within the oxygen minimum zone (OMZ) (Fig. 1; Reinhardt et al., 2003). The sediment cores were retrieved from mud lenses off Callao (12° S, 184 m, Krissek et al., 1980) and off Pisco (14° S, 300 m, Gutiérrez et al., 2006).

2 Materials and Methods

The Callao (~12° S) and Pisco (~14° S) areas were selected as sites for box coring based on previous sedimentological and geochemical information from exploratory surveys and the literature. Two Soutar-box cores were collected: one from the shelf off Callao (B0405-13, 12°00′S, 72°42′S, 184 m) and the other one from the upper slope off Pisco (B0405-06, 14°07′S, 76°30′S, 299 m) from the R/V José...
Olaya Balandra (IMARPE) in May 2004 (Fig. 1). Subsampling for proxy determinations was performed following the stratigraphy, i.e. according to the geometry of couplets of laminae or bands (Gutiérrez et al., 2006; Morales et al., 2007).

The quantitative mineralogical composition, including lithic and biogenic-derived compounds, was obtained by Fourier Transformed Infrared Spectrometry (FTIR) (Bertaux et al., 1998). More detailed explanations on FTIR methodology are given in Sifeddine et al. (2008). Total organic carbon was determined from total carbon measurements with a Thermo Electron CNS elemental analyzer, corrected for carbonate content. $\delta^{15}N$ was determined by mass spectrometry after acidification at the Department of Geosciences, University of Arizona (USA). Analyses of Molybdenum (Mo) and Cadmium (Cd) were carried out by ICP-MS and ICP-AES (Ultramass Varian), respectively, after hot plate acid digestion (combination of acid HF, HNO$_3$ and HClO$_4$), which was used to eliminate organic matter and to remove silicates (Cho et al., 1999).

Subsamples for foraminifera and fish scales were heated with hydrogen peroxide and sodium pyrophosphate to remove the organic matter and then were sieved through a 125 $\mu$m mesh and 355 $\mu$m mesh. The retained material in the 355 $\mu$m mesh was sorted for fish remains (bones and scales) while foraminifers were counted and identified from the smaller size fraction with stereo-microscopy. Diatom valves and other siliceous remains were determined from splits that were acid cleaned with HCl 10% and H$_2$O$_2$ 30% (Batterbee, 1989). Quantitative slides were made using 100 $\mu$L of acid-clean residue of about 0.1 g of wet-sediment (equivalent of 0.02–0.03 g of dry-sediment) diluted on a 20-mL vial. A 22×22 mm coverslip was permanently fixed over a 1″×3″ glass slide with Zrax® (R.I.=1.74) to enhance diatom taxonomic features. Identiﬁcations were done with contrast-phase microscopy techniques.

Mass accumulation rates (MAR) for the past $\sim$135 years were determined from downcore profiles of $^{241}$Am and excess $^{210}$Pb. Radiocarbon ages from bulk organic sediments were used to solve the geochronology for the whole record (see supplementary information: http://www.biogeosciences.net/6/835/2009/bg-6-835-2009-supplement.pdf). It has been suggested that $^{14}$C dating of bulk organic carbon provides reliable results for the Peruvian margin sediments, by comparison of dated bulk organic matter and alkenone $^{14}$C dates (Higginson and Altabet, 2004). In this study, geochemical and petrographic inspection of the samples ensured that the dated organic matter in the age model was dominated by homogenous and granular amorphous organic matter, which is associated with high productivity linked to upwelling (Boussafir et al., 1995; Pichevin et al., 2004; Valdés et al., 2004). This was done to minimize possible bias of the age models in the older part of the cores due to the presence of reworked material in the samples.

The conventional radiocarbon ages were calibrated taking into account global and local reservoir effects that were estimated (supplementary information: http://www.biogeosciences.net/6/835/2009/bg-6-835-2009-supplement.pdf). A linear regression between these calibrated $^{14}$C ages and accumulated mass was then used to estimate MAR for the lower section of the cores. The final age models combined the $^{210}$Pb-based chronology with the above mentioned age-mass regression fit, and also took into account stratigraphic features (Fig. 2).
3 Results and discussion

3.1 Lithology and geochronology

The Pisco and Callao cores show an almost identical shift in core lithologies at about 35 cm (Fig. 3a, j), as observed in dry bulk density and X-ray grey level, despite differences in location (300 km of distance) and depth within the OMZ. There is a striking similarity in the core lithologies and downcore patterns of lithic, geochemical and biogenic proxies between the two sites (Fig. 3), which indicates that regional to large-scale processes largely determine the temporal variations. Terrigenous sedimentation dominates below 35 cm, whereas diatom/siliceous and organic matter are major constituents above that depth.

The MAR varied by a factor of two, with the highest average values after ca. 1950 AD in both cores (Callao: 0.034±0.001 g cm⁻² y⁻¹; Pisco: 0.036±0.001 g cm⁻² y⁻¹) and lowest values before ca. 1820 AD (Callao: 0.017±0.001 g cm⁻² y⁻¹; Pisco: 0.022±0.001 g cm⁻² y⁻¹) (Table 1). The cores spanned the last 700±30 years approximately. Dating and sampling resolution resolve variability at sub-decadal to inter-decadal time-scales (Fig. 2; see also supplementary material: http://www.biogeosciences.net/6/835/2009/bg-6-835-2009-supplement.pdf). The shift in core lithologies is dated near 1820 AD (Fig. 3b, k).

The period since the late nineteenth century to the present show sedimentation rates of 1.8–2.1 mm y⁻¹ for Callao, and 1.9–2.3 mm y⁻¹ for Pisco, very similar to those of Koidzumi and Goldberg (1982) and to Rein et al. (2004), who studied cores near our study sites using ²¹⁰Pb or ¹³⁷Cs (Table 1). For the period before the late nineteenth century our sedimentation rates (0.6–0.7 mm y⁻¹) are similar to previous studies based on ¹⁴C measurements in the sedimentary organic fraction (Rein et al., 2004; Agnihotri et al., 2006; Skilbeck and Fink, 2006), even though there are differences in local reservoir ages used.

3.2 Multidecadal to centennial variability in multiple proxies

The multiple geochemical and biogenic proxies developed in the study are used to reconstruct past changes of water column oxygenation, sediment redox conditions and productivity, according to their source and downcore behaviour (Fig. 3b, k). In order to constrain the interpretation of our records, we compare our results with some records developed by Sifeddine et al. (2008) as Total Organic Carbon (TOC), Molybdenum (Mo) and Lithogenic input. The variation pattern of fluxes of these proxies is also evident with concentrations, indicating that flux variability represents true changes in the input and not only changes in sedimentation rate (Sifeddine et al., 2008). The same applies for the multiple proxies introduced here. Nevertheless, we preferred to use fluxes instead of concentrations in order to control dilution effects in the sedimentary matrix.

For both cores several of the proxies show slightly higher levels before ca. 1400 AD, followed by low values until 1820 AD, when most of the geochemical and biogenic proxies exhibit a rapid change towards higher values, within one to two decades given chronological errors. A sudden decrease of lithic fluxes occurred also around 1820 AD, suggesting that the western flank of the Peruvian Andes shifted from a wetter to a drier condition at this time (Sifeddine et al., 2008).

3.2.1 Water column biogeochemistry

δ¹⁵N of organic matter exhibited remarkably similar levels between sites at the different centennial periods (Fig. 3c, i). Values from 1400 to 1820 AD (4.8±0.6‰), corresponding to the LIA, were about 1‰ lighter than before 1400 AD. The LIA period bears multidecadal variability; timing of periods with lighter δ¹⁵N values at both sites were about 1480–1530 AD, the mid seventeenth century and 1780–1810 AD. Around 1820 AD a rapid change towards heavier values occurred at both sites within one or two decades. δ¹⁵N reached an average of 6.7±0.6‰ for the third period, about 2‰ heavier than the preceding centuries.

The δ¹⁵N from our records agrees with the δ¹⁵N reported by Agnihotri et al. (2008) in a 2300-year late Holocene sedimentary record at 11°S off Peru. However, different chronological approaches preclude a direct comparison between our 700-year record and the 11°S record reported by Agnihotri et al. (2008). These authors employed an overall reservoir age correction of 420±170 years, implying a local reservoir effect (∆R) from negligible levels to ca. 170 years, while in this study, the ∆R was estimated to be 188±79 years for Pisco and 279±53 years for Callao, consistent with previous estimates reported in the literature (Taylor and Berger, 1967; Jones et al., 2007, see also supplementary information: http://www.biogeosciences.net/6/835/2009/bg-6-835-2009-supplement.pdf).

Downcore variations of δ¹⁵N can reflect changes in organic matter source, in diagenesis or in the δ¹⁵N of the nitrate taken up by phytoplankton (Altabet et al., 1999; Martínez et al., 2006, DePol-Holz et al., 2007). Analysis of the organic fraction (palynofacies) indicates that over 95% of the organic matter is marine in the Callao core, and over 90% is marine in most of the Pisco core (Suppl. Fig. SF2, see http://www.biogeosciences.net/6/835/2009/bg-6-835-2009-supplement.pdf). Terrestrial plant organic matter is typically high (>20) in C:N ratios and light in δ¹³C (Meyers, 1997). Our downcore C:N records were closer to the marine Redfield ratio and did not show any significant change around 1820 AD. Downcore C:N was 10.2±0.7 in Callao, whereas in Pisco it was higher before 1600 AD (10.9±0.5) than after this date (8.9±0.8). The C:N ratio confirms that there were no significant changes in the
Table 1. Mean sedimentation rates (SR) in the central Peruvian upper margin. Results of this study (including MAR) and of previous studies are compared.

<table>
<thead>
<tr>
<th>core</th>
<th>latitude (S)</th>
<th>water depth (m)</th>
<th>MAR (g cm⁻² y⁻¹)</th>
<th>mean SR (mm y⁻¹)</th>
<th>time window (year AD)</th>
<th>dating method</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>1228</td>
<td>11°04'</td>
<td>252</td>
<td>0.4</td>
<td>1430 - 1950 AD</td>
<td>(^{14}C) (n = 1)</td>
<td>Skibb &amp; Fink (2006)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.7</td>
<td>&lt;0 - 1430 AD</td>
<td>(^{14}C) (n = 6)</td>
<td>Skibb &amp; Fink (2006)</td>
<td></td>
</tr>
<tr>
<td>1229</td>
<td>10°59'</td>
<td>151</td>
<td>0.3</td>
<td>1300 - 1950 AD</td>
<td>(^{14}C) (n = 1)</td>
<td>Skibb &amp; Fink (2006)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1.0</td>
<td>&lt;0 - 1300 AD</td>
<td>(^{14}C) (n = 6)</td>
<td>Skibb &amp; Fink (2006)</td>
<td></td>
</tr>
<tr>
<td>W7706-40</td>
<td>11°18'</td>
<td>186</td>
<td>0.7</td>
<td>&lt;0 - 1650</td>
<td>(^{14}C) (n = 6)</td>
<td>Agricolti et al. (2008)</td>
<td></td>
</tr>
<tr>
<td>B0413</td>
<td>12°00'</td>
<td>184</td>
<td>0.034 ± 0.001</td>
<td>&gt;1952 AD</td>
<td>(^{234}U)</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.031 ± 0.003</td>
<td>1870 - 1952 AD</td>
<td>(^{234}U)</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.021 ± 0.005</td>
<td>1818 - 1870 AD</td>
<td>(^{137}Cs) Pb</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.017 ± 0.001</td>
<td>ca. 1300 - 1818 AD</td>
<td>(^{14}C) (n = 6), stratigraphy</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td>2312</td>
<td>12°02'</td>
<td>194</td>
<td>3.3</td>
<td>1946 - 1974</td>
<td>(^{206}Pb) - CFCS</td>
<td>Koide &amp; Goldberg (1982)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1.5</td>
<td>1870 - 1946</td>
<td>(^{206}Pb) - CFCS</td>
<td>Koide &amp; Goldberg (1982)</td>
<td></td>
</tr>
<tr>
<td>106KL</td>
<td>12°03'</td>
<td>184</td>
<td>1.3</td>
<td>1900 - 1960 AD</td>
<td>(^{137}Cs) Pb</td>
<td>Rein et al. (2004)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.9</td>
<td>ca. 1400 - 1900</td>
<td>(^{14}C) (n = 2)</td>
<td>Rein et al. (2004)</td>
<td></td>
</tr>
<tr>
<td>B0406</td>
<td>14°07'</td>
<td>296</td>
<td>0.036 ± 0.001</td>
<td>&gt;1952 AD</td>
<td>(^{234}U)</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.033 ± 0.009</td>
<td>1860 - 1952 AD</td>
<td>(^{239}Pu) Pb</td>
<td>This study</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>0.025 ± 0.006</td>
<td>1960 - 1946 AD</td>
<td>(^{137}Cs) Pb</td>
<td>This study</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>0.022 ± 0.001</td>
<td>ca. 1300 - 1818 AD</td>
<td>(^{14}C) (n = 6)</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td>1909</td>
<td>14°39'</td>
<td>183</td>
<td>3.2</td>
<td>1946 - 1974</td>
<td>(^{206}Pb) - CFCS</td>
<td>Koide &amp; Goldberg (1982)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1.5</td>
<td>1870 - 1946</td>
<td>(^{206}Pb) - CFCS</td>
<td>Koide &amp; Goldberg (1982)</td>
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</tbody>
</table>

Fig. 3. Multiple proxies developed in the cores collected off Callao (upper panels: a–i) and off Pisco (lower panels: j–r). (a, j) Dry bulk density (white circles; g cm⁻³) and X-ray radiography gray level units (gray lines) are shown as a function of depth. All other proxies are shown as a function of time: (b, k) Flux of the sum of lithic components (feldspar, quartz and clays; Sifeddine et al., 2008); (c, l) yellow circles: \(\delta^{15}N\) values of sedimentary organic matter (%); line: C:N molar ratios; (d, m) flux of elemental Cadmium (\(\mu g\) cm⁻² y⁻¹); (e, n) bars: total foraminiferal abundance (Nr ind. g⁻¹); blue circles: percent of *Bolivia seminuda* in the benthic foraminiferal assemblage; (f, o) flux of elemental Molybdenum (\(\mu g\) cm⁻² y⁻¹) reported by Sifeddine et al. (2008); (g, p) flux of total organic carbon (mg C cm⁻² y⁻¹) (Sifeddine et al., 2008); (h, q) bars: diatom accumulation rate (\(10^{6}\) values cm⁻² y⁻¹), white circles: flux of biogenic silica (mg cm⁻² y⁻¹); (i, r) bars: fish scale deposition rates (Nr x 1000 cm⁻² y⁻¹), gray circles: deposition rates of fish bones and vertebrae (Nr x 1000 cm⁻² y⁻¹). A shift in DDB and grey level at 35 cm depth and its corresponding date (1820 AD) is indicated in each panel by dotted lines and connected with an arrow.
source of organic matter, and it also rules out the possibility that diageneisis was the controlling factor for δ15N, since downcore variations of δ15N and C:N ratios were independent. The stable C:N ratios indicate that there was not a preferential removal of more labile N-rich organic compounds during prolonged periods with low δ15N; increasing C:N ratios would be associated with diageneisis of organic matter (Martínez et al., 2006).

In the oxygen-deficient waters of the ETSP nitrate is used as an electron donor for respiration during degradation of organic matter. Hence settling organic matter δ15N will reflect 15N-enriched nitrate taken up by phytoplankton from upwelled oxygen-depleted water masses (Altabet et al., 1999; De Pol et al., 2007). Therefore our δ15N records ultimately reflect changes in the intensity of nitrate reduction in the source waters for primary producers, i.e. changes in water column oxygenation. The same conclusions have been reached for other sedimentary δ15N records in the ETSP (De Pol et al., 2007; Agnihotri et al., 2008) and this interpretation is consistent with our other proxies of oxygenation and productivity.

Cd exhibits nutrient-like characteristics in the water column; dissolved Cd is normally depleted in surface waters by phytoplankton uptake. Equatorial Subsurface Waters (ESSW), which are the main source of upwelling in the ETSP, are enriched in dissolved Cd (Valdés et al., 2006; Takesue et al., 2004). Cadmium is delivered to sediments in association with organic matter and becomes fixed in reducing sediments (Tribovillard et al., 2006; Böning et al., 2004). Cd in sediments can then be considered a proxy of paleoupwelling and paleoproductivity (Dean et al., 2006). In our records, Cd flux decreases from the fourteenth to the fifteenth century, and then remains at very low levels (Fig. 3d, m). Like δ15N, Cd exhibits a step-like increase in baseline values near 1820 AD, and later on, multidecadal oscillations without any particular trend. The similarity in the behaviour of Cd and δ15N can be interpreted as a rapid change in source water properties towards a new biogeochemical regime with lower oxygen and higher nutrient concentrations. Sifeddine et al. (2008) determined that primary productivity, as inferred by Total Organic Carbon (TOC), was lower and constant before the shift (Fig. 3g, p). It increased rapidly with the shift, showing a non-linear transition until 1870 AD. Thereafter, it exhibited a significant positive trend until the present.

Thus the large shift around 1820 AD most likely reflects a large-scale depletion in oxygen below the mixed layer and surface nutrient enrichment. The shift must have been associated with an expansion of the nutrient-rich, oxygen-poor ESSW that primarily feeds coastal upwelling in the ETSP (Strub et al., 1998). In their 2300-year record, Agnihotri et al. (2008) determined four short centennial periods of “high production and intense denitrification”, with δ15N values similar to those from our records after 1820 AD. The short ‘high production’ periods punctuated longer, persistent periods with “moderate productivity and denitrification”, with δ15N values in the same range as we determined for the LIA. The shift in δ15N of our records is approximately 2‰. Though lower than the δ15N change during the last deglaciation in the Eastern Tropical and Extratropical South Pacific records (5‰; Higginson and Altabet, 2004; De Pol-Holz et al., 2006), the shift towards the end of the LIA and others during the late Holocene are of the same order of magnitude and sign as the δ15N millenial shift at 30° S off Chile during the Younger Dryas (De Pol-Holz et al., 2006; Fig. 2). Therefore the modern biogeochemical regime in the ESTP established since ca. 1820 AD is an excursion from the background Holocene ocean climate, probably as significant as millennial-scale events, leading to an expansion of the OMZ throughout the ETSP.

3.2.2 Reduction-oxidation (redox) conditions in surface sediments

While higher oxygenation in the water column characterized the LIA period prior to 1820 AD, sufficiently dysoxic conditions at the sediment-water interface still permitted the preservation of fine laminations in the deeper Pisco site (Fig. 1), as flux varied between terrigenous and organic sources (Gutiérrez et al., 2006). Fine laminations were not preserved at the Callao site during the LIA, probably due to its shallower water depth that would result in more exposure to bottom oxygenation. However, changes in foraminiferal preservation, as inferred by presence/absence of foraminifera, and changes in the benthic foraminiferal assemblage, support the view of less intensely reducing conditions in the surface sediments at both sites during the LIA and increasingly sulphidic conditions after the LIA.

Dissolution of calcitic foraminiferal tests can occur at lower alkalinity, a condition that is met at the surface when sulphate reduction occurs deeper in the sediments (Froelich et al., 1988). Foraminiferal tests were preserved before 1400 AD, disappeared in the sediment accumulated during the LIA, and re-appeared after 1820 AD. The lack of foraminifers in the LIA layers is attributed to post-depositional dissolution of the tests due to reduced alkalinity. This interpretation is supported by the reduced sulphate reduction during the LIA inferred from Mo content (Sifeddine et al., 2008), by the finding of tests in the Pisco core just below a slump dated at the LIA (Morales et al., 2006; see also Fig. 3e, n), and by the presence of test linings in core slab thin-sections around the same period. In addition, biological mixing and slower sedimentation rates likely enhanced test dissolution in the Callao core.

Among the benthic foraminiferal species present in the Peruvian margin, Bolivina seminuda is increasingly dominant as conditions become more dysoxic (Resig, 1981). Therefore we use the percent contribution of B. seminuda to the benthic foraminiferal assemblage as a proxy of bottom water oxygen and redox conditions at the sediment/water interface (Fig. 3e, n). B. seminuda was present in the records
before the LIA, comprising between 25 and 75% of the total abundance. After the 1820 AD shift, it rapidly increased with time at both sites attaining about 90% dominance at ca. 1950 AD, and then exhibited decadal oscillations until the present. The behaviour of B. seminuda after the shift resembles the variability of Mo records (Sifeddine et al., 2008). Mo displays a quasi-conservative behaviour in the water column, and it is scavenged when free sulphide and elemental sulphur species are present in the water column, in pore waters, or even in sediment anoxic microniches (Tribovillard et al., 2006, 2008). On the Peruvian margin the highest Mo enrichment occurs in organic-rich sediments within the OMZ and on the shelf (Böning et al., 2004; McManus et al., 2006). The increases of dominance of B. seminuda and Mo concentrations at ca. 1820 AD (Fig. 3f, o) are therefore consistent with a rapid upward expansion of the anoxic layer within the surface sediments.

3.2.3 Preservation of proxies of siliceous and fish productivity

Various proxies of primary and tertiary productivity further support an abrupt shift toward higher productivity and reduced oxygenation. Diatoms and amorphous silica exhibited similar patterns throughout the records (Fig. 3h, q), most notably lowest values during the LIA, and an abrupt increase occurred near 1820 AD. Persistently higher primary productivity after 1820 AD may not be apparent in the individual records of diatoms (or amorphous silica) as there is a negative correlation between diatom valves and TOC \(\frac{r=−0.79^{**}}{n=37}\); and \(r=−0.69^{**}, n=42\), for Callao and Pisco, respectively). Higher alkalinity due to enhanced sulphate reduction near the sediment interface leads to an increase of dissolution of diatom valves (Froelich et al., 1988). Thus, the increasing sulphidic conditions associated with higher TOC fluxes after the shift, as inferred by Mo and other redox-sensitive records, may have increased the dissolution of biogenic silica, biasing the record of siliceous productivity after 1820 AD.

Changes in bottom water oxygen concentration may affect the preservation of fish scales (Salvatteci, 2008). Fish scales have more fissures and altered colorations in the centuries immediately preceding the shift, indicating that greater degradation may be partly responsible for the reduced number of fish scales. Additionally, fish scales show more signs of degradation in the shallower Callao site, located closer to the oxycline, making direct comparisons of fish scale abundance and flux between the two cores difficult. Therefore, following DeVries (1979), we consider fish bones and vertebrae as more reliable indicators of variability in local pelagic fish abundance as they are more resistant to degradation or dissolution than fish scales.

3.2.4 Variability after the 1820 regime shift

There are two different patterns in the biogeochemical proxies during and after the 1820 shift. The first one, typified by \(\delta^{15}N\) and foraminifera preservation, shows a step-like change at ca. 1820 AD to persistent higher oxygen depletion and redox conditions. The second pattern, typified by TOC, fish remain fluxes, percent abundance of B. seminuda and Mo fluxes shows a rapid increase after 1820 AD that persisted for a few decades, followed by a return towards pre-shift conditions from ca. 1845 to 1865 AD, and then a long-term trend towards higher levels. Cd fluxes exhibit an intermediate behaviour, probably because their records are influenced by sediment redox conditions. However, fluxes of diatom and biogenic silica are actually higher during 1845–1865 AD due to high abundances of Skeletonema costatum, a species which blooms during upwelling relaxation (Alvarez et al., 2005) and which forms aggregates at warmer temperatures (Thornton and Thake, 1998). Thus this bidecadal period likely corresponds to conditions with fewer upwelling events, more frequent intrusions of oceanic waters and more stratification in the coastal fringe, allowing episodic S. costatum blooms, but with oxygen and nutrient concentrations in the upwelling source waters at post-shift levels.

3.3 Climatic driver for the biogeochemical shift

Seasonal to interannual variability of water column structure, upwelling intensity and productivity in the ETSP are coupled to the strength of the Walker circulation, the expansion/contraction of the South Pacific Subtropical High (SPSH) and the position of the Intertropical Convergence Zone (ITCZ) (Barber and Chávez, 1983; Strub et al., 1998). Generally, changes in Walker circulation, SPSH and ITCZ are correlated during El Niño (EN) events, whereby the coast of Peru is characterized by reduced productivity and an increase in subsurface oxygenation due to a weakening of the equatorial thermocline and Walker circulation (Morales et al., 1999). While coral records from the mid-Equatorial Pacific suggest enhanced ENSO activity during part of the LIA (Cobb et al., 2003), more comprehensive SST reconstructions, multiproxy records and historical information indicate that the expression of EN events, particularly along the Peru coast was less frequent and/or less intense during the LIA (d’Arrigo et al., 2005; Gergis and Fowler, 2006; Ortlieb, 2000) and therefore a higher frequency of EN events is an unlikely explanation of the LIA conditions reported here.

We hypothesize that the centennial-scale shift in the Peruvian upwelling system around 1820 AD was driven by a northward displacement of the ITCZ and the northern rim of the SPSH from their southward condition during the LIA to their modern positions, coupled with an enhancement of the Walker circulation. Comparison of the \(\delta^{15}N\) records off Peru with Indo-Pacific records of paleotemperature and paleosalinity shows that a relatively abrupt change occurred in
Fig. 4. Comparison of (a) $\delta^{15}$N time-series in Pisco (this study) with paleo records developed in other regions: (b) $\delta^{15}$N time-series in Mejillones Bay, which reflects alongshore winds intensity and influence of ESSW in coastal upwelling off Northern Chile (Vargas et al., 2007), (c) Indo-Pacific paleo-temperatures and paleo-salinities based on Mg:Ca and $\delta^{18}$O/$\delta^{16}$O ratios of foraminiferal tests (Newton et al., 2006), (d) percent sedimentary Titanium concentration from the Cariaco Basin taken as an indicator of regional runoff and precipitation (Haug et al., 2001). The gray bar envelopes the shift period, within dating uncertainties.

the Indo-Pacific region at the same time than the biogeochemical shift off Peru (Fig. 4). The large-scale expansion of ESSW and associated increase in productivity imply a basin-scale adjustment of the equatorial thermocline/pycnocline tilt and stronger equatorial upwelling (Pennington et al., 2006; Fiedler and Talley, 2006). Taken together, the records also imply enhanced eastern trades, expansion of the cold tongue, and a stronger zonal SST gradient across the Pacific.

Thus, the lower productivity and less oxygen-depleted regime during the LIA was probably maintained by an ITCZ located southward of its modern position and reduced influence of the SPSH along the Peruvian margin. This combination of factors also diminished equatorial upwelling and hence both the zonal and meridional SST gradients in the Eastern Tropical Pacific (Haug et al., 2001; Koutavas and Lynch-Stieglitz, 2004). A northward migration of the ITCZ, as inferred from the Cariaco basin (Fig. 4), might have caused a “tipping point” that established the Walker circulation that is evident in modern times.

There is considerable evidence for a southward displacement of the ITCZ during the LIA, attributed to the cooling of the Northern Hemisphere (Koutavas and Lynch-Stieglitz, 2004). Terrigenous input to the Peruvian continental margin was most probably driven by precipitation and runoff-driven erosion of fine-grained material, followed by river discharge and dispersion by ocean currents (Scheiddeger and Krissek, 1982; Rein et al., 2007). Therefore, the higher lithic fluxes to the Central/Southern Peruvian margin sediments during the LIA are in agreement with continental records that indicate enhanced humidity and monsoonal precipitation in the Western Peruvian Andes (Jomelli et al., 2007; Unkel et al., 2007), when dry climatic conditions characterized the Cariaco Basin (Fig. 4; Haug et al., 2001). Lower salinity in Makassar Strait, a condition that prevailed during the LIA, occurs nowadays during austral summer, associated with higher precipitation and monsoonal winds but not during El Niño events (Gordon et al., 2003; Newton et al., 2006). Figure 5 illustrates how the inferred precipitation patterns during the LIA are consistent with a persistent austral summer-like southward migration of the ITCZ and the associated precipitation belt, rather than a strong zonal (El Niño-like) shift of precipitation eastward.

The southward position of the ITCZ was likely associated with a latitudinal contraction of the SPSH during the LIA, as lower paleotemperatures ($U^K_{37}$) off Concepcion ($36^\circ$ S) suggest an increased latitudinal SST gradient across the Eastern Subtropical South Pacific (Vargas et al., 2007). An enhanced latitudinal SST gradient and Hadley circulation also characterized the LIA in the subtropical South Pacific until around 1870 (Hendy et al., 2002).

The $\delta^{15}$N records off Mejillones at $23^\circ$ S, Northern Chile, exhibit a delayed and gradual increase of $\delta^{15}$N from 1820 to 1870 AD (Vargas et al., 2007) compared to the step-like change in the Peruvian margin records. It was postulated that $\delta^{15}$N variability off Central Chile during the Holocene is modulated by the formation rate of the Eastern South Pacific Intermediate Waters (ESPIW), which in turn is governed by wind forcing and latitudinal SST gradient at mid-latitudes (DePol-Holz et al., 2006, 2007). Currently this water mass ventilates only a thin layer over the thermocline up to Northern Chile (Schneider et al., 2003), but during the Late Glacial Maximum and most of the Holocene it could have been much more developed ventilating the subsurface layer (DePol-Holz et al., 2006). Higher ventilation during the LIA can be expected here if the northward displacement of subtropical fronts is associated to lower SST gradient at mid-latitude. Therefore the slower $\delta^{15}$N rise during the period 1820–1870 AD in Mejillones might be due to persistent ESPIW-driven ventilation, counteracting the increased oxygen depletion originating at low-latitudes. Certainly this hypothesis needs to be examined with more subtropical and mid-latitude late Holocene $\delta^{15}$N and paleotemperature records from the South Eastern Pacific.

The variability in the Peru records between 1820 and 1870 AD can be attributed to changes in the intensity of the SPSH. A continued strengthening of the SPSH and/or coastal winds along the Peruvian margin in the twentieth century may account for the trend towards greater TOC and redox conditions (Sifeddine et al., 2008; see also Fig. 3). This trend contrasts with the variability of water column oxygen and nutrient properties that only show multidecadal-scale fluctuations after 1820, as inferred from $\delta^{15}$N and Cd. This
decoupling of productivity (TOC) and water column nitrate reduction ($\delta^{15}$N) differs from the findings of Agnihotri et al. (2008), who suggest that productivity changes and water column nitrate reduction in the late Holocene were tightly coupled at centennial time-scales.

3.4 Ecosystem responses to the biogeochemical shift

Diatom analysis suggests that both cool-upwelling and warmer-offshore species were favoured after the 1820 AD shift. While the abrupt increase in diatoms was dominated by the cool-water upwelling diatom assemblage, other groups increased as well (Fig. 6). In particular, oceanic diatoms off Pisco actually make up a greater portion of the total...
assemblage during several periods of the high productivity regime.

The response of pelagic fish species is similar, with the increase in productivity after 1820 AD being favourable for both cool-upwelling species as well as species that live in or beyond the upwelling front waters (Fig. 6). The records of fish remains from different locations in Fig. 3 reveal how different species have expanded and contracted both in abundance and population range. It should be noted that diatoms and the conglomerate of pelagic fish shown in Fig. 3 are generally selected in highly productive upwelling ecosystems and that cooler and warmer in this context refers to relative changes within the generally productive condition.

The present day nucleus of the Peruvian anchovy or anchoveta, *Engraulis ringens*, is from 6° to 12° S and the Callao site lies within this range (Pauly and Tsukayama, 1987). Since anchoveta is the dominant species in terms of fish scales, we can then assume that vertebrae and other bones in the records are dominated by anchoveta as well. The higher fluxes of fish remains in Callao than in Pisco during the twentieth century is consistent with the modern spatial distribution for this species (Figs. 3i, r and 6).

Fish abundances inferred from bones and vertebrae were dramatically lower prior to 1820 AD off Callao. During warm, less productive periods anchoveta migrate to the Pisco area (Mendo, 1991), which often retains a small upwelling plume even under EN events (Barber and Chávez, 1983). Off Pisco, fluxes of fish bones and vertebrae from 1600 to 1700 AD were as high as during the twentieth century, while in other periods of the LIA they were reduced. Further south, Mejillones Bay (23° S) is within the distributional area of the present day Northern Chile anchoveta stock (Serra, 1983).
Fish remains in Mejillones Bay sediments do not show the centennial shift around 1820 AD, though fish scales from anchoveta increased slightly around this date (Valdés et al., 2008). The combined records suggest that off Central Peru the main pelagic fish populations were reduced during the LIA, and that the anchoveta population had moved southward.

During the 20th century multidecadal periods of low anchovy abundances are often associated with increases in sardine abundances synchronously off California, Japan and Peru (Schwartzlose et al., 1999; Chavez et al., 2003). Although sardine scales are much thicker and more resistant to degradation than anchoveta scales, no scales were found off Callao or Pisco during the low productivity regime, neither were any from larger offshore species such as jack mackerel (*Trachurus murphyi*) or horse mackerel (*Scomber japonicus peruvianus*). At Mejillones Bay, scales of sardine and offshore species were present throughout the record with no signs of degradation, but as with bones and vertebrae, fluxes were not higher prior to 1820. The combined evidence from the three different sites indicates that sardines did not expand during the centennial-scale regime of low productivity. This suggests that the decrease in productivity during LIA was of such magnitude and duration to limit growth of fish populations that in the modern era seem to be adapted to periods of lower productivity.

Within the highly productive regime, a centennial-scale northward expansion of anchoveta is coupled with two multidecadal periods of expansion of sardines (Fig. 6). The former period corresponds to low anchoveta scale fluxes and low fluxes of bones and vertebrae during 1845–1865 AD, which also shows that reduced fluxes of fish debris is not necessarily an artefact of degradation. The latter period corresponds to the multi-decadal “sardine regime” observed in the twentieth century (Chavez et al., 2003). Note that the flux of bones and vertebrae is diminished since the development of the fishery in the 1960s, likely ending up as fish meal rather than passing through the guts of predators and being transported to the sediments.

### 3.5 Implications for climate and ecosystem changes

The reduced Walker circulation during the LIA was probably maintained by a southern displacement of the ITCZ associated with global cooling, in sharp contrast to present-day weakening of the Walker circulation associated with warming across the globe during El Niño periods. Furthermore, the current EN teleconnection that results in precipitation anomalies in Central Chile and Northwestern Peru was absent prior to ca. 1817 AD (Ortlieb, 2000, 2004), likely due to changes in tropical atmospheric circulation at this time. The implication is that we cannot use present-day conditions as predictors of the future and the coupling of important processes (like the ITCZ and Walker circulation) may differ in the future under different climate scenarios.

The abruptness of the regime shift reported here reveals the potential for rapid reorganization in the modern tropical Pacific climate, circulation, ocean biogeochemical cycles and marine productivity. The large and rapid changes in productivity and biogeochemical cycles in the ETSP are of the same magnitude as the gradual centennial to millennial changes during the Holocene. Current global warming favours the expansion of the tropical oxygen minimum areas in the global ocean (Stramma et al., 2008). However, parallel increase of open-ocean stratification, weakening of tropical circulation and trade winds system (Vecchi et al., 2006), and enhancement of coastal upwelling (Bakun, 1990; Bakun and Wecks, 2008) may introduce new instabilities in the present-day biogeochemical regime that started with the end of the LIA.

On the other hand, the ecosystem response after the end of the LIA involved an overall increase of the major groups of diatom and pelagic fish, in contrast to decreased ecosystem productivity in the Eastern Pacific during El Niño and El Viejo (Barber and Chávez, 1983; Chavez et al., 2003). This paradox suggests that the processes at work during recent multidecadal variations may be substantially different from centennial- to millenial-scale changes. Furthermore, without the centennial, climate-driven, increase in bottom-up productivity of the upwelling system the intense anchoveta exploitation during the past decades would not have been possible. Given that looming climate changes may easily exceed variations observed in the twentieth century, major and non-linear regime shifts in the ecosystems that depart from twentieth century paradigms should be expected. Mechanistic understanding is needed to predict future scenarios and to develop adaptive policies.

### 4 Conclusions

Reduced productivity along with greater oxygenation in the water column characterized the ETSP biogeochemical regime during the LIA, from the fifteenth to the early nineteenth century. This regime was likely maintained by a mean southward displacement of the ITCZ relative to its modern range and an associated weakening of the SPSH.

A rapid expansion of nutrient-rich, oxygen depleted waters occurred in the ETSP around 1820 AD, probably driven by the northward shift of the ITCZ, the expansion of the SPSH and a spin-up of tropical circulation that has persisted (and perhaps strengthened) through modern time. The magnitude of the shift in biogeochemical cycling and particularly in the regional marine nitrogen budgets appears similar to other centennial excursions from the background Holocene climate and to millennial-scale change during the Younger Dryas in the Eastern South Pacific. The impact on the ecosystem was an overall increase in primary and fish productivity that has continued during the twentieth century.
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