The evolution of deep ocean chemistry and respired carbon in the Eastern Equatorial Pacific over the last deglaciation

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Key Points:
- Combined [CO³⁻], δ¹³C and U/Ca evidence indicates increased respired carbon storage in the last glacial maximum Eastern Equatorial Pacific
- Along with reduced deep-water ventilation, this evidence points to enhanced soft-tissue pump efficiency driven by ocean circulation changes
- Modest [CO³⁻] change suggests calcite dissolution as a further contribution to atmospheric CO₂ drawdown via increased ocean alkalinity
Abstract

It has been shown that the deep Eastern Equatorial Pacific (EEP) region was poorly ventilated during the Last Glacial Maximum (LGM) relative to Holocene values. This finding suggests a more efficient biological pump, which indirectly supports the idea of increased carbon storage in the deep ocean contributing to lower atmospheric CO₂ during the last glacial. However, proxies related to respired carbon are needed in order to directly test this proposition. Here we present *Cibicides wuellerstorfi* B/Ca ratios from ODP Site 1240 measured by laser-ablation inductively-coupled-plasma mass spectrometry (LA-IPCMS) as a proxy for deep water carbonate saturation state (Δ[CO₃²⁻], and therefore [CO₃²⁻]), along with δ¹³C measurements. In addition, the U/Ca ratio in foraminiferal coatings has been analysed as an indicator of oxygenation changes. Our results show lower [CO₃²⁻], δ¹³C and [O₂] values during the LGM, which would be consistent with higher respired carbon levels in the deep EEP driven, at least in part, by reduced deep-water ventilation. However, the difference between LGM and Holocene [CO₃²⁻] observed at our site is relatively small, in accordance with other records from across the Pacific, suggesting that a ‘counteracting’ mechanism, such as seafloor carbonate dissolution, also played a role. If so, this mechanism would have increased average ocean alkalinity, allowing even more atmospheric CO₂ to be ‘sequestered’ by the ocean. Therefore, the deep Pacific Ocean very likely stored a significant amount of atmospheric CO₂ during the LGM, specifically due to a more efficient biological carbon pump, but also to an increase in average ocean alkalinity.

1. Introduction

Several marine carbon cycle mechanisms have been invoked to explain, at least, a major part of the atmospheric CO₂ drawdown observed over the last glacial period [e.g. Kohfeld and Ridgwell, 2009; Sigman et al., 2010]. These mechanisms can be categorised broadly as deriving primarily from: 1) changes in whole ocean carbonate chemistry through global alkalinity (ALK) increase, for example due to ‘carbonate compensation’ or carbonate dissolution [e.g. Archer and Maier-Reimer, 1994]; 2) changes in the conditions or efficiency of air-sea gas exchange, i.e. changes in the ‘solubility pump’ [e.g. Volk and Hoffert, 1985]; or 3) changes in the accumulation of respired organic carbon (in the form of dissolved inorganic carbon, DIC) in the ocean interior, i.e. changes in the global average efficiency of the ‘soft-tissue’ component of the ‘biological carbon pump’ [e.g. Volk and Hoffert, 1985], either in absolute terms or relative to the global average efficiency of its ‘carbonate pump’ counterpart.

The efficiency of the soft tissue carbon pump was initially defined in terms of the gradient of major nutrient concentrations between the deep ocean and the shallow ocean, which arises due to the interplay of biological export from the surface ocean and the large-scale overturning circulation [Volk and Hoffert, 1985]. Due to the association of carbon with biologically fixed nutrients, the efficiency of the soft tissue pump relates directly to the DIC concentration gradient between the surface- and deep ocean (note that the two may become uncoupled due to air-sea
gas exchange effects, which affect carbon but not major nutrients). A more efficient glacial soft-tissue pump would therefore result in greater carbon storage in the deep sea and a lower surface ocean- and atmospheric pCO₂. This would occur as long as the soft tissue pump remained dominant over its carbonate pump counterpart (as it is in the modern ocean), as an efficient carbonate pump draws down surface ocean alkalinity and thus increases atmospheric CO₂. Changes in the relative strength of the soft tissue and carbonate pumps would be driven by e.g. biological community structure driven DIC:ALK 'rain ratio' changes or water column remineralisation profile changes [Archer and Maier-Reimer, 1994; Kwon et al., 2009]. Notably, a global average increase in soft-tissue pump efficiency can occur as a result of: 1) an increase in the biological flux of nutrients and carbon from the surface ocean to the deep interior, representing increased soft-tissue pump 'strength' [e.g. Martinez-Garcia et al., 2014]; and/or 2) a reduction in the return flux of remineralised nutrients and respired carbon from the deep interior to the surface ocean, for instance due to a weaker large-scale overturning circulation, representing reduced soft-tissue pump 'leakiness' [e.g. Toggweiler, 1999; Stephens and Keeling, 2000].

While a categorization of the various mechanisms for atmospheric CO₂ drawdown, as proposed above, might be useful for organising our thinking about past marine carbon cycle changes, it is important to emphasize that these mechanisms need not have operated in isolation from each other. For example, if an increase in soft-tissue pump global efficiency was achieved via a change in the ocean circulation, this would very likely co-occur with a change in the conditions of air-sea gas exchange and therefore the solubility pump. Furthermore, regardless of whether enhanced soft-tissue pump global efficiency is achieved via an increase in biological export rates or a reduction in the overturning circulation rate, it would lead to more respired carbon storage in the deep ocean, which in turn would lower deep ocean carbonate ion concentration ([CO₃²⁻]), and [O₂]. If [CO₃²⁻] was reduced enough to cause seafloor carbonate dissolution (a mechanism also referred to as ‘respiratory calcite dissolution’ [Archer, 1991, 1996; Archer and Maier-Reimer, 1994]), it would eventually increase the whole ocean alkalinity budget and lead to further atmospheric CO₂ drawdown.

It has been proposed that increased export production in the EEP over the last glacial period played a role storing more respired carbon in the deep ocean than during the Holocene [e.g. Pichevin et al., 2009; Robinson et al., 2009; Doss and Marchitto, 2013]. However, other studies have argued against this [e.g. Winckler et al., 2016; Costa et al., 2016]. Glacial [O₂] decrease in the deep EEP supports higher respired DIC accumulation [Bradtmiller et al., 2010], although without elucidating the mechanism behind its variation. In this respect, it is now apparent that the EEP was less ventilated over the last part of the last glacial period, becoming better ventilated at the onset of the Heinrich Stadial 1 [de la Fuente et al., 2015; Umling and Thunell, 2017]. This would indirectly support the idea of a more efficient biological pump during the last glacial due to a longer residence time for carbon in the deep ocean, thus reducing the leakiness of the biological pump and enhancing the deep ocean’s respired carbon inventory.

In this study we set out to test the proposition that the respired carbon content of deep waters was indeed greater in the EEP during the LGM, as suggested by radiocarbon ventilation ages from this area [e.g. de la Fuente et al., 2015].
approach is to combine several respired carbon related proxies in the same sediment core ODP Site 1240 (hereafter referred to as ODP1240). Thus measurements of δ¹³C and B/Ca have been performed on the calcite shells of the benthic foraminifer species *Cibicides wuellerstorfi* as a means of reconstructing respired nutrient and [CO₃²⁻] changes, respectively. B/Ca analyses have been proposed as a method for reconstructing past deep-water [CO₃²⁻] changes on the basis of an empirical correlation with deep water carbonate saturation state (Δ[CO₃²⁻]) [Yu and Elderfield, 2007], and on the assumption of invariant local saturation [CO₃²⁻] ([CO₃²⁻]ₛₑₐ). In addition, sedimentary redox conditions (i.e. estimates of changing [O₂]) have been reconstructed from analyses of U/Ca in the coatings (U/Ca₃ [Boiteau et al., 2012; Gottschalk et al., 2016]) of the planktonic species *Neogloboquadrina dutertrei*, and have been compared to existing authigenic uranium (aU; [Cochran et al., 1986; Bradtmiller et al., 2010; Jaccard et al., 2016]) measurements from the same area [Kienast et al., 2007; Bradtmiller et al., 2010].

Neither U/Ca₃ nor aU have so far been shown to quantify changes in O₂, and despite the fact that each of these approaches has its own limitations, we propose that their combination may offer robust qualitative information on glacial-interglacial bottom-water/pore-water [O₂] changes.

2. Materials and methods

2.1. EEP hydrographic settings

Sediment core ODP1240 was recovered from the southern edge of the Panama Basin in the EEP (0°01.31' N, 86°27.76' W) at 2,921 m water depth (Figure 1). The chronostratigraphy for this sediment core was previously described by de la Fuente et al. [2015], and is based on sediment depth-age modeling using calibrated radiocarbon dates on *Neogloboquadrina dutertrei* that have been corrected for a variable reservoir age offsets, constrained through stratigraphic alignment of the U₋³⁷ record from ODP1240 with Greenland δ¹⁸O records. The hydrography of the EEP, in common with the whole tropical Pacific, is highly influenced by Southern Ocean water masses at depth [Strub et al., 1998; Kessler, 2006], with the depth of ODP1240 location, in particular, currently bathed by a mixture of Upper Circumpolar Deep Water (UCDW) and Pacific Deep Water (PDW; Figure 1). PDW refers to the water mass formed at depth in the North Pacific through very slow vertical mixing of bottom and intermediate waters of southern origin and extremely long residence times, and UCDW represents the upper layer of the deep water mass flowing within the Antarctic Circumpolar Current (ACC), a mixture of deep and re-circulated waters from the Atlantic, Indian and Southern Oceans [Mantyla and Reid, 1983; You, 2000]. Although these deep water masses have different origins, both occupy approximately the same density and depth range in the Pacific [Talley et al., 2011] as a consequence of the extremely weak deep circulation in the Pacific Ocean. An additional feature of the modern Panama Basin hydrography is a degree of ‘biogeochemical aging’ in its deepest part. Although this points to the possibility of highly localized influences on respired carbon in the Panama Basin, such influences do not affect the basin as a whole and are relatively small, with hardly any influence the at location of ODP1240 today.
2.2. B/Ca analysis by LA-ICPMS and $[\text{CO}_3^{2-}]$ estimates

The epibenthic foraminifer *C. wuellerstorfi* was picked from 15 samples from the top three meters of sediment core ODP1240 covering the last ~30 kyr. Each sample was composed of ~6 specimens (>212 μm) that were rinsed in milliQ water and secured on a double-sided carbon tape on a glass slide for LA-ICPMS analysis. Between 2-3 depth-profiling ablations were made per shell using the RESOlution M-50 prototype 193nm ArF laser ablation system (featuring a two-volume Laurin LA cell) coupled to an Agilent 7500ce quadrupole ICPMS at Royal Holloway University of London [Müller et al., 2009]. Ablation was performed using a laser spot size of 96 μm, repetition rate of 2 Hz and fluence of ~4 J/cm². Data quantification follows [Longerich et al., 1996] with the transformation of raw count rates to concentration achieved using $^{43}\text{Ca}$ as an internal standard and NIST610/612 as a calibration standard. The accuracy and precision of B/Ca measurements using these conditions is <5 and <10% respectively (see Evans et al. [2015]). Several steps to exclude any possible contaminant phases from the data were applied [Boyle, 1983; Bolton et al., 2011; Marr et al., 2011], and spots with B/Ca values higher than the individual foraminifer average plus 2SD were removed and not included in further palaeoclimatic interpretations. A more detailed discussion of the methodology can be found in the Supporting Information [Eggins et al., 1998; Jochum et al., 2006; Jochum et al., 2011; Evans and Müller, 2013; Müller and Fietzke, 2016].

*C. wuellerstorfi* B/Ca ratios have been experimentally shown to linearly relate to $\Delta[\text{CO}_3^{2-}]$ based on global core-top measurements in the modern ocean, as described by equation 1 below [Yu and Elderfield, 2007], thereby allowing downcore $[\text{CO}_3^{2-}]$ to be estimated from equation 2, albeit with some caution.
$\Delta[\text{CO}_3^{2-}] = (B/\text{Ca} - 177.1)/1.14 \text{ (in } \mu\text{mol/mol})$  \hspace{1cm} (1)\\

$\Delta[\text{CO}_3^{2-}] = [\text{CO}_3^{2-}] - [\text{CO}_3^{2-}]_{\text{sat}}$  \hspace{1cm} (2)

Seawater $[\text{CO}_3^{2-}]_{\text{sat}}$ in the modern ocean is estimated from water column $[\text{CO}_3^{2-}]_W/\Omega_{\text{calcite}}$ data, where $\Omega_{\text{calcite}}$ is the calcite saturation state in seawater ($\Omega_{\text{calcite}}$ = $[\text{Ca}^{2+}]_W/\text{K}_{\text{sp}}$, and $\text{K}_{\text{sp}}$ is the solubility product of calcite $([\text{Ca}^{2+}]_W[\text{CO}_3^{2-}]_{\text{sat}}$ at a given temperature, salinity and pressure). However, seawater $[\text{CO}_3^{2-}]_{\text{sat}}$ in the deep ocean is typically assumed to have remained constant over the late Pleistocene since the influence of pressure, bottom water temperature and salinity in the past are thought to have exerted little effect on $[\text{CO}_3^{2-}]_{\text{sat}}$ over this time period [Allen et al., 2015]. Thus, the general equation to estimate downcore $[\text{CO}_3^{2-}]$ through B/Ca ratios measured in the calcite shell of $C. \text{wuellerstorfi}$ would be:

$$[\text{CO}_3^{2-}] = ((B/\text{Ca} - 177.1)/1.14) + ([\text{CO}_3^{2-}]_{\text{(mod)}}/\Omega_{\text{calcite (mod)}})$$ \hspace{1cm} (3)

In this study, modern $[\text{CO}_3^{2-}]$ and $\Omega_{\text{calcite}}$ have been inferred from instrumental measurements of ALK and DIC available from the GLODAP dataset (WOCE P19C transect; Tsuchiya and Talley, [1998]). For these estimates, the CO2SYS software package [Lewis et al., 1998; Pierrot et al., 2006] and the equilibrium constants from Mehrbach et al. [1973] and Dickson and Millero [1987] were used. The obtained modern values at a depth of 2,888 m were 67.7 µmol/kg for $[\text{CO}_3^{2-}]$ and 0.9 for $\Omega_{\text{calcite}}$, leading to a modern $[\text{CO}_3^{2-}]_{\text{sat}}$ of 75.2 µmol/kg (station: 18373; LAT: 0.004° N; LONG: 85.84° W).

Since the initial calibration of Yu and Elderfield [2007] was published, several studies have provided additional core-top measurements, but only one provided B/Ca results analysed by LA-ICPMS [Raitzsch et al., 2011]. In order to add further support to this technique, B/Ca ratios in $C. \text{wuellerstorfi}$ specimens have been analysed here in nine core-top samples across the Atlantic Ocean by LA-ICPMS, following the same scheme described above for downcore analyses. Estimates of modern $\Delta[\text{CO}_3^{2-}]$ were obtained using either in situ data from the corresponding cruises when available, or data from nearby GLODAP sites [Key et al., 2004], in both cases by deriving values for modern $[\text{CO}_3^{2-}]$ and $\Omega_{\text{calcite}}$ (Table S1 in the Supporting Information).

### 2.3. $\delta^{13}$C measurements

Down-core $\delta^{13}$C measurements were performed on 46 samples from ODP1240, covering the last 25 kyr and consisting exclusively of $C. \text{wuellerstorfi}$ individuals from the >212 µm fraction size. The analyses were performed using a Thermo Kiel device attached to a Thermo MAT253 Mass Spectrometer in dual inlet mode at the University of Cambridge. The sample size was ~100 µg, with the exception of a few samples of 20 µg, which is close to the detection limit of this machine. A few additional $\delta^{13}$C measurements in $C. \text{wuellerstorfi}$ from ODP1240, consisting of four to five specimens each (also from the >212 µm size fraction) were obtained at the Scientific and Technological Center of the University of Barcelona (CCiT-UB) by using a FinniganMAT252 mass spectrometer fitted with a Kiel Carbonate Device I. All $\delta^{13}$C values presented here are reported as per mil relative to the international
standard Vienna Pee Dee Belemnite (VPDB), with a typical external reproducibility estimated to be equal or better than ±0.06‰ and ±0.03‰ for the Universities of Cambridge and Barcelona, respectively.

2.4. U/Ca analysis in planktonic foraminiferal coatings

U/Ca ratios measured in the foraminiferal shell coatings have been suggested to reflect changes in the sedimentary redox conditions and therefore potentially linked with changes in the [O$_2$] in bottom waters [Boiteau et al., 2012]. This oxygen related proxy is based on the redox chemistry of U in seawater, shifting from the soluble form U(VI) in oxygenated waters to the insoluble U(IV) species as the medium depletes in oxygen [Langmuir, 1978; Cochran et al., 1986; Morford and Emerson, 1999].

In order to quantify the U concentration in the foraminiferal coatings, clays were removed and an acid leach step to remove any adsorbed ions was performed in 20 N. dutertrei samples, each one composed of ~30 specimens collected from the 212-300 µm fraction. No oxidative or reductive step was performed in order to preserve the thin coating layer [Boiteau et al., 2012]. Cleaned samples were dissolved in 0.1M HNO$_3$, diluted to 10 ppm Ca$^{2+}$ and analysed by ICP-MS at the University of Cambridge using a Thermo Element XR, with a typical analytical reproducibility of ~7%.

3. Results

3.1. C. wuellerstorfi B/Ca core-top measurements by LA-ICPMS

In Figure 2, C. wuellerstorfi B/Ca analyses from the Atlantic Ocean core-tops are compared to their corresponding Δ[CO$_3^{2-}$] values, as well as to other core-top data available from the literature [Yu and Elderfield, 2007; Brown et al., 2011; Rae et al., 2011; Raitzsch et al., 2011; Yu et al., 2013].
Figure 2. Bottom water $\Delta [CO_3^{2-}]$ vs B/Ca ratios in *C. wuellerstorfi* from core-top samples analysed by solution-ICPMS [Yu and Elderfield, 2007; Brown et al., 2011; Rae et al., 2011; Yu et al., 2013] and by LA-ICPMS [Raitzsch et al. [2011] and core-tops from the Atlantic Ocean analysed in this study (red stars)]. Black solid line represents the linear fit from Yu and Elderfield [2007] by solution-ICPMS, while grey solid line represents the linear fit from Raitzsch et al. [2011] by LA-ICPMS. Black dashed line represent the linear fit from all the records together including the core-tops analysed in this study.

The core-top measurements presented here fit well within the two existing linear regressions from Yu and Elderfield [2007] (A=1.14; B=177.1) and from Raitzsch et al. [2011] (A=1.37; B=170.9), and combining all of the data results in an indistinguishable new calibration equation (~230 core top samples including this study; A=1.07; B=177.7), thus providing support for B/Ca as a proxy for $\Delta [CO_3^{2-}]$ and for LA-ICPMS as a suitable technique for this type of analysis. The original calibration equation from Yu and Elderfield [2007] is adopted here for downcore $\Delta [CO_3^{2-}]$ estimates in ODP1240 for better comparison with previously published data.

3.2. B/Ca-$[CO_3^{2-}]$, $\delta^{13}C$ and U/Ca, downcore ODP1240

Downcore B/Ca ratios from *C. wuellerstorfi* in ODP1240 core are ~10 µmol/mol lower over the end of the last glacial period (~18-23 kyr), with an average of ~172±5 µmol/mol (±1SE), than across the mid-late Holocene (~0-8 kyr), which
exhibits values of ~182±4 µmol/mol (±1SE) (Figure 3a). When converted into [CO$_3^{2-}$] the difference between glacial (~71 µmol/kg) and interglacial (~80 µmol/kg) is ~9 µmol/kg, which is not as large as might be expected based on observed radiocarbon ventilation changes, possibly as a result of other processes that we discuss in the next sections. C. wuellerstorfi δ$^{13}$C measurements from ODP1240 are also characterised by lower values during the LGM compared to the Holocene of ~0.4 ‰, showing good consistency between measurements performed in the two laboratories (Figure 3b). Although both [CO$_3^{2-}$] and δ$^{13}$C seem to be in good agreement and exhibit a similar pattern across the deglaciation, a ‘bump’ just before the termination onset is observed only in the [CO$_3^{2-}$] record. This may reflect processes that have an impact exclusively in [CO$_3^{2-}$] without modifying δ$^{13}$C (i.e. changes in seafloor carbonate dissolution), or analysis artefacts in those specific samples. Although we cannot discard any of these possibilities unequivocally, the observed change in [CO$_3^{2-}$] before the deglaciation seems to be too rapid to be explained by variations in calcite dissolution. U/Ca$_C$ measurements in the planktonic species N. dutertrei show higher values at the LGM compared to the Holocene suggesting, very likely, a more reducing environment at the water-sediment boundary at that time, and therefore lower [O$_2$], in the deep EEP (Figure 3c).

Figure 3. Deep water respired carbon related records across the first 450 cm of the sediment core ODP1240 (last ~30 kyr). a) B/Ca ratios in C. wuellerstorfi measured by LA-ICPMS and [CO$_3^{2-}$] derived estimates (grey solid line/dots). Uncertainties represent ±1 SE in [CO$_3^{2-}$] (grey envelope). [CO$_3^{2-}$] was
estimated by using the calibration equation from *Yu and Elderfield* [2007] and assuming a constant [CO$_2^-$]$_{aw}$ value of 75.2 μmol/kg (estimated from modern [CO$_2^-$] and δC$_{DIC}$ values from the GLODAP dataset WOCE P19 transect, see main text). b) Benthic δ$^{13}$C from *C. wuellerstorfi* (black line/solid dots and crossed dots analysed at the Universities of Cambridge and Barcelona, respectively). c) U/Ca ratios in the coatings of *N. dutertrei* (grey dashed line/asterisks). All lines are B-spline smoothed. Vertical dashed lines delimit relevant climatic periods, and light grey vertical bands highlight the coldest periods of the last 30 kyr. HS1 and HS2: Heinrich Stadial 1 and 2 respectively; BA: Bølling-Allerød; YD: Younger Dryas.

4. Discussion

4.1. A comparison of ocean ventilation and carbonate system changes in the EEP across the last deglaciation

A poorly ventilated deep EEP at the end of the last glacial period compared to the present has been shown by the radiocarbon offsets between benthic foraminifera and atmospheric records (B-Atm) [*de la Fuente et al.*, 2015]. This study also revealed that better ventilation of this area of the Pacific began at the onset of the deglaciation (~17.5 kyr BP), demonstrating a consistency with other ventilation records from the South Pacific [*Sikes et al.*, 2000; *Siani et al.*, 2013; *Skinner et al.*, 2015] and the Atlantic sector of the Southern Ocean [*Skinner et al.*, 2010; *Burke and Robinson*, 2012], all of which broadly co-varied in time with changes in atmospheric Δ$^{14}$C and CO$_2$. Thus, when CO$_2$ increased and Δ$^{14}$C decreased in the atmosphere at the deglaciation onset (Figure 4a,b, respectively), all of these deep water ventilation records start to exhibit enhanced ventilation (Figure 4c, *de la Fuente et al.* [2015] and figures therein). This observation has at least two important implications: 1) that the deglacial increase in atmospheric CO$_2$ might be directly and causally linked to the deglacial evolution of deep ocean ventilation; and 2) that the deep ocean, including the EEP, might have contained a higher concentration of respired CO$_2$ during the glaciation as compared to the late Holocene and thus acted as a source of carbon to the atmosphere during deglaciation.

In the absence of major ocean interior carbon sources, radiocarbon behaves as a semi-conservative tracer that is not significantly affected by biological or chemical reactions within the ocean, and tracks the mean time since a water mass exchanged carbon with the atmosphere. Radiocarbon ventilation estimates therefore allow us to isolate the contribution of ocean ventilation to changes in the global average efficiency of the soft-tissue pump. Increased radiocarbon ventilation ages observed in ODP1240 during the last late glacial period could therefore imply an increase in the amount of remineralised/oxidised organic matter in the ocean interior, i.e. more carbon respired by microorganisms per unit volume due to more time for this respired carbon to accumulate. If this happened, the most noticeable consequences might be, on the one hand, a decrease in seawater δ$^{13}$C due to enhanced lighter carbon isotope (12C) release, and on the other hand, a greater CO$_2$ accumulation in seawater as a product of respiration. Such an increase in seawater [CO$_2$] would have altered the carbonate system equilibrium, leading to a drop in [CO$_3^{2-}$] in order to buffer the change in seawater pH [*Zeebe and Wolf-Gladrow*, 2001; *Sarmiento and Gruber*, 2004; *Williams and Follows*, 2011].
The $[\text{CO}_3^{2-}]$ and $\delta^{13}\text{C}$ results presented here from ODP1240 are in good agreement with these expectations (Figure 4d,e), as well as with a previous study in the area [Doss and Marchitto, 2013], showing lower $\delta^{13}\text{C}$ and $[\text{CO}_3^{2-}]$ during the late glacial, in parallel with reduced radiocarbon ventilation, and an increase in all of these parameters across the deglaciation. Interestingly, when the modern $[\text{CO}_3^{2-}]$ value from this area is taken into account, a drop at the late Holocene is inferred. The existence of such $[\text{CO}_3^{2-}]$ decrease across the late Holocene has been previously suggested on the basis of sediment composition data and modeling studies [Broecker and Peng, 1987; Berelson et al., 1997]. This feature might reflect a restored lower average ocean alkalinity due to enhanced carbonate preservation associated with the release of excess respired carbon from the ocean interior to the atmosphere during deglaciation, which in turn might have helped to maintain the higher CO$_2$ in the atmosphere during the late Holocene. However, another plausible explanation for the late Holocene-modern value ‘mismatch’ observed in our core might be related to the inherent noise contained in the B/Ca-$\Delta[\text{CO}_3^{2-}]$ calibration applied. Indeed, this issue has been observed in previous studies that estimated $[\text{CO}_3^{2-}]$ through B/Ca analysis in benthic foraminifera [Yu et al., 2008, 2013; Allen et al., 2015; Elmore et al., 2015]. In order to avoid this mismatch, some of these studies opted for applying a correction by subtracting the core top value to each measurement downcore, thus forcing the record to fall on the regression line given by the equation: $[\text{CO}_3^{2-}]_{\text{downcore}} = [\text{CO}_3^{2-}]_{\text{pre-industrial}} + \Delta(\text{B/Ca})/1.14$, where $\Delta(\text{B/Ca})$ represents the deviation of each downcore sample relative to the core-top value [Yu et al., 2008, 2010, 2013, 2014b, 2016]. However, in our study, the conventional option of applying the linear correlation from Yu and Elderfield [2007] without correcting for the core-top value has been chosen to avoid applying arbitrary offsets to the dataset. This issue would call for cautious approach when considering absolute $[\text{CO}_3^{2-}]$ values derived from benthic B/Ca analysis, and may suggest that relative changes observed down-core might be more robust than inter-core or absolute value comparisons. It would also call for a careful interpretation when using data for any further carbonate system calculations.
Figure 4. Comparison of deep ocean records from ODP1240 with records of atmospheric CO$_2$ and radiocarbon activity over the last 30 kyr. a) Atmospheric CO$_2$ concentrations from EPICA Dome C (EDC) ice core (for the deglacial period, Monnin et al., [2001]; for the Holocene period, Flückiger et al., [2002]), placed on the age scale of Lemieux-Dudon et al. [2010] (grey line). b) Atmospheric radiocarbon activity ($\Delta^{14}C$) changes (IntCal09 calibration curve; Reimer et al., [2009] (black dotted line). c) Deep-water ventilation reconstruction (B-Atm, from SST alignment in de la Fuente et al. [2015]) (black line/white stars). d) $[\text{CO}_3^{2-}]_{(\text{B} \text{-Ca-based})}$ ($\pm$ SE) (grey solid line/dots). e) $\delta^{13}C$ from C. wuellerstorfi (black line/solid dots and crossed dots analysed at the Universities of Cambridge and Barcelona, respectively). f) Oxygenation proxies: U/Ca$_C$ from N. dutertrei (grey dashed line/asterisks) and $^{238}$U from ME0005-24JC (same location as Site 1240; blue line/stars; Kienast et al., [2007]; Bradtmiller et al., [2010]). All lines are B-spline smoothed. Vertical dashed lines delimit relevant climatic periods, and light grey vertical bands highlight the coldest periods of the last 30 kyr. The grey arrow on the $[\text{CO}_3^{2-}]$ axis represents the modern water value (67.6 µmol/kg) obtained from GLODAP database (WOCE P19C transect; Tsuchiya and Talley, [1998]).

Another consequence of the hypothetical increase in respired carbon due to a longer residence time of deep water during the LGM would be a drop in $[O_2]$, as microorganisms consume $O_2$ during respiration of organic matter. Following
reformulations of the traditional Redfield ratios of remineralised organic matter [Redfield et al., 1963], the release of 117 CO₂ molecules consumes 170 molecules of O₂ [Anderson and Sarmiento, 1994]. Our U/Ca₃ results show the expected depletion in O₂ if the respired carbon content of the ocean interior was indeed higher at the LGM compared to the Holocene (Figure 4f). Moreover, our qualitative relative oxygenation changes indicated by U/Ca₃ estimates support previous measurements of authigenic uranium (aU) in core ME0005-24JC [recovered from nominally the same location as Site 1240; Kienast et al., [2007]] showing a similar trend, characterised by higher aU (i.e. lower O₂) during the glacial compared to the Holocene [Kienast et al., 2007; Bradtmiller et al., 2010] (Figure 4f).

Thus, all records presented here, i.e. [CO₃²⁻], δ¹³C and oxygenation proxies, are in good agreement with the deep radiocarbon ventilation in the EEP, and lend support to the hypothesis of a higher efficiency of the biological pump during the LGM due to changes in its leakiness. In the same way, all of these records are in reasonably close agreement through deglaciation, increasing as the deep EEP gets better ventilated at the onset of the last termination. Despite this general agreement, it is interesting to note that both oxygenation proxies, aU and U/Ca₃, do not change synchronously with the rest of the proxies at the beginning of the deglaciation, but later at the termination. A plausible explanation for this apparent mismatch might rely on a decoupling of pore-water and deep water chemistry, for example due to a productivity pulse between ~17-14 kyr [e.g. Calvo et al., 2011] that would have depleted O₂ in the pore-fluids of the sediment (which the oxygenation proxies would record), even after the radiocarbon ventilation and carbonate saturation of the ambient deep-water started to increase. Taken together, all the records presented here point to the deep EEP as a plausible contributor to the decreased atmospheric CO₂ during the LGM, by storing more respired carbon at the expense of the surface ocean and atmospheric carbon inventories, that was posteriorly released during the deglaciation once the ocean interior got better ventilated.

A different interpretation based on similar glacial-interglacial Δ[CO₃²⁻] changes, also from the Panama Basin, has been proposed by Doss and Marchitto, [2013]. This study suggests that the observed glacial Δ[CO₃²⁻] difference found between a sediment core located at the sill depth and sediment cores located deeper in the basin are indicative of an increase in the export production during the last glacial. While we do not discard this possibility, which must eventually be tested against other evidence for increased export production during the last glacial period, we propose that it must have acted in addition to a decrease in ocean ventilation, as indicated by our combined proxy results, including radiocarbon ventilation ages. Below we discuss the available evidence for other possible contributions to the apparent increase in the respired carbon content of the deep EEP, including more direct nutrient-based indicators of export production.

4.2. Export production as a complementary mechanism for a higher biological pump efficiency in the EEP

An increase in the strength of the biological pump (i.e. export production rates) could have operated on the glacial atmospheric CO₂ drawdown independently of,
or in addition to, contributions from changes in ocean ventilation constrained using radiocarbon measurements. The glacial changes observed in deep ocean [CO$_3^{2-}$], δ$^{13}$C and oxygenation records from ODP1240, described in the previous subsection, might indeed be compatible with an increase in surface productivity over the LGM. Thus, a larger flux of organic matter to the deep ocean in this area might have contributed to the observed drop in [CO$_3^{2-}$] and δ$^{13}$C, as a consequence of higher levels of carbon respiration that would have also increased the release of respired CO$_2$ to deep water and the consumption of dissolved O$_2$.

The documented increase in dust-borne iron delivery to the iron-limited low-latitude Pacific Ocean at the last glacial period [McGee et al., 2007] has led to a number of investigations into a potential increase in surface productivity over this period, however, with ambiguous results. On the one hand, lower opal δ$^{30}$Si content in ODP1240 as a proxy for the relative utilization of silica during the glacial has been interpreted as an indication of a decline in the Si:C uptake ratio by diatoms due to an iron-replete glacial ocean, which would be compatible with the assumption of higher productivity over this period [Pichevin et al., 2009]. Similarly, nitrogen isotopes (δ$^{15}$N) from bulk sediments in ODP1240 core, once they are corrected for denitrification effects typical of the Central American margin, showed higher values over the LGM compared to the Holocene suggesting higher local nutrient consumption consistent with higher surface productivity over the LGM [Robinson et al., 2009]. At the same time, higher abundances of organic biomarkers (alkenones and brassicasterol) have been shown in the same core during the LGM, as compared to the present [Calvo et al., 2011]. However, these inferences are contradicted by other studies that do not support the proposed increase in the strength of the biological pump in the EEP as a mechanism for the atmospheric CO$_2$ drawdown during the LGM [Bradtmiller et al., 2006; Kienast et al., 2006]. More recently, for instance, Winckler et al. [2016] reported excess Ba and opal fluxes obtained from sediment cores located along the equatorial Pacific, which do indicate higher productivity in this area (stimulated by an increased Fe supply by upwelling rather than by dust deposition), but with peak export occurring during deglaciation rather than over the LGM. Moreover, Costa et al. [2016] suggested that the glacial dust supply to the equatorial Pacific exerted a negligible effect on surface productivity, arguing that the major nutrient supply with Southern Ocean origin was actually reduced as nutrients were mostly consumed in the Subantarctic Zone.

An increase in organic carbon (C$_{org}$) accumulation rates in the EEP over the LGM has been documented in core ODP1240 [Pichevin et al., 2009], as well as in the nearby core P6 [Pedersen, 1983]. Such higher C$_{org}$ accumulation might have been the result of an increase in surface productivity, but also the result of better preservation of the organic matter due to lower oxygen content in the deep ocean at that time. Thus, in light of the fact that existing proxy reconstructions can neither corroborate nor rule out unequivocally an increase in EEP surface productivity over the LGM, two key questions remain: was the observed increase in the biological pump efficiency in the glacial EEP mainly due to changes in ocean circulation, or changes in the carbon export production in this area; how much did the proposed increase in organic carbon export production affect carbon storage in the deep ocean? Arguably these questions cannot be fully resolved without accurate quantitative estimates of export production and deep ocean apparent
oxygen utilization changes. Nevertheless, it is clear at least that ocean ventilation changes made some contribution, though only qualitative conclusions, such as those drawn above, are possible at present.

4.3. Carbonate seafloor dissolution as a potential counteracting mechanism for $[\text{CO}_3^{2-}]$ EEP changes

A decrease in deep ocean ventilation along with a potential increase in carbon export production in the EEP at the LGM are therefore proposed as the mechanisms responsible for the decrease in deep $[\text{CO}_3^{2-}]$, $\delta^{13}\text{C}$ and oxygenation observed in this area. The action of any of these mechanisms (alone or together) might be expected, in principle, to achieve a relatively large $[\text{CO}_3^{2-}]$ glacial-Holocene difference. For example, spatial correlations of radiocarbon and $[\text{CO}_3^{2-}]$ in the modern deep ocean would predict a change of $\sim 37 \mu\text{mol/kg}$ for the $\sim 960 \mu\text{mol/kg}$ increase in deep water radiocarbon ventilation observed in ODP1240 at the LGM relative to the modern ocean [Key et al., 2004]. However, the results from this study indicate only a small $[\text{CO}_3^{2-}]$ change between these two periods ($\sim 9 \mu\text{mol/kg}$), suggesting that a counteracting mechanism, such as seafloor respiratory calcite dissolution, might have also played a role: when carbonate dissolves in seawater, both $[\text{CO}_3^{2-}]$ and ALK increase [Broecker and Peng, 1989; Zeebe and Wolf-Gladrow, 2001]. If this was the case, a decrease in the CaCO$_3$ content in sediments might be initially expected. However, CaCO$_3$% from ODP1240 indicates little change between 25-15 kyr (Figure 5). Furthermore, the possibility of an increase in the calcium carbonate export production rates as a masking mechanism for the expected CaCO$_3$% drop seems not to be supported by carbonate flux data (expressed as $^{230}\text{Th}$-CaCO$_3$) from this area, which also show little change between last glacial and Holocene (Figure 5).
Figure 5. Carbonate chemistry-related records from core ODP1240 across the past 30 kyr. Upper panel: \([\text{CO}_3^{2-}]_{\text{B/Ca}}\) (grey line/dots; this study). Lower panel: \(^{230}\text{Th}-\text{CaCO}_3\) and \(\text{CaCO}_3\%\) (purple dashed line/open dots and yellow line/open diamonds, respectively; Pichevin et al., [2009]). Vertical dashed lines delimit relevant climatic periods, and light grey vertical bands highlight the coldest periods of the last 30 kyr.

Despite no obvious carbonate dissolution in core ODP1240 being observed during the LGM, it remains possible that \([\text{CO}_3^{2-}]\) was added to deep waters bathing the EEP through carbonate dissolution elsewhere in the Pacific [Anderson et al., 2008]. Indeed, similarly small \([\text{CO}_3^{2-}]\) differences between LGM-Holocene/modern have also been reported in other areas from the Pacific Ocean [Yu et al., 2010; Doss and Marchitto, 2013; Allen et al., 2015; Elmore et al., 2015], and this has been generally attributed to the large buffering capacity of carbonate sediments in this basin [Yu et al., 2010, 2013; Allen et al., 2015]. In Figure 6, a compilation of the averaged \(\Delta[\text{CO}_3^{2-}]\) offsets between the LGM (~18-23 kyr) and the late Holocene (~0-8 kyr) \(\langle \Delta[\text{CO}_3^{2-}]\rangle_{\text{LGM-Hol}}\) from these studies shows the consistency among the Indo-Pacific sites with variations of only ~5 \(\mu\text{mol/kg}\) between these two periods (orange line in Figure 6). By restricting the comparison to LGM-Hol differences, potential biases due to different techniques/calibrations/standards can be avoided, as well as any biases that may arise from assuming constant local \([\text{CO}_3^{2-}]_{\text{sat}}\) across the deglaciation. Thus, negative values represent lower \(\Delta[\text{CO}_3^{2-}]\) at the
LGM compared to the Holocene, while positive ones represent the opposite. The small LGM-late Holocene difference observed in the Indo-Pacific, in contrast with larger offsets detected within the Atlantic and Southern Oceans (blue line in Figure 6) [Yu et al., 2008, 2013, 2014a, 2014b; Gottschalk et al., 2015], still implies lower LGM Δ[CO$_3^{2-}$] for the whole Pacific as well as for the Atlantic and Southern Ocean at depths greater than ~2,500 m water depth. However, the Δ(Δ[CO$_3^{2-}$])$_{LGM-Hol}$ changes present a noticeable difference between basins above ~2,500 m water depth. This might point to the accumulation of respired DIC (i.e. not equilibrated with the atmosphere) in the Pacific, Southern Ocean and Atlantic >2,500m during the last glacial, and the maintenance of 'equilibrium' DIC [Ito and Follows, 2013] in Atlantic water masses shallower than ~2,500 m. The shallow Atlantic would thus have responded more closely to changes in the atmosphere, such that lower atmospheric CO$_2$ would be directly manifested as higher LGM [CO$_3^{2-}$] [Hodell et al., 2001; Yu et al., 2008]. In this situation, seafloor CaCO$_3$ not only might have 'buffered' glacial [CO$_3^{2-}$] changes in the EEP (without affecting δ$^{13}$C significantly), but also might have played a role in the atmospheric CO$_2$ drawdown during the LGM by contributing to increased average ocean alkalinity via [CO$_3^{2-}$] supply to the ocean [Keir, 1995; Sigman and Boyle, 2000]. This would have further reinforced the effects of a higher biological pump efficiency achieved by circulation changes, and perhaps also by surface productivity changes.
Figure 6. Deep water $[\text{CO}_3^{2-}]$ offset between LGM (~18-23 kyr) and Holocene/modern (~0-8 kyr) from several Atlantic/Southern Ocean (in blue) and Indo-Pacific Ocean (in orange) sediment cores. DSDP593Z [Elmore et al., 2015], RR0503-83 [Allen et al., 2015], MW91-9 GGC15, MW91-9 GGC48, WIND 28K and VM28-122 [Yu et al., 2010], TTNO13 PC61, [Yu et al., 2013], BOFS and NEAP cores [Yu and Elderfield, 2007; Yu et al., 2008], RC13-140, RC23-22 and RC23-15 [Doss and Marchitto, 2013], RC16-59 [Broecker et al., 2015], GeoB cores [Raitzsch et al., 2011], MD07-3076 [Gottschalk et al., 2015], TN057-21 [Yu et al., 2014b] and ODP1240 (this study, highlighted in bold). WP: western Pacific; EEP: eastern equatorial Pacific; CP: central Pacific; NEA: North East Atlantic; WEA: western equatorial Pacific; EqA: equatorial Atlantic; SO: Southern Ocean; IO: Indian Ocean.

Interestingly, calculations based on the modern $\delta^{13}$C-$[\text{CO}_3^{2-}]$ relationship in the Panama Basin suggest that the observed glacial-interglacial $\delta^{13}$C difference is
compatible with the observed glacial-interglacial offset in [CO$_3^{2-}$] (when a whole ocean glacial-interglacial δ$^{13}$C change of ~0.3‰ is assumed, due to terrestrial carbon release), which would imply no further [CO$_3^{2-}$] addition from calcite dissolution [Doss and Marchitto, 2013]. Following the calculations of Doss and Marchitto, [2013], a 10 µmol/kg [CO$_3^{2-}$] glacial-interglacial drop would achieve a 30 µmol/kg DIC increase, which in turn would produce a ~0.26‰ change in δ$^{13}$C due to respired carbon addition. If a whole ocean change of ~0.3‰ [Peterson et al., 2014] is assumed to have occurred exclusively due to terrestrial carbon release [Shackleton, 1977], a total change of ~0.56‰ would therefore be expected, which is not very far off the results of Doss and Marchitto, [2013] and our observed glacial-interglacial δ$^{13}$C difference of 0.4±0.06‰. However, it is notable that the observed global average glacial-interglacial δ$^{13}$C change of ~0.3‰ corresponds to a global average radiocarbon ventilation age change of perhaps ~689 14C yr [Skinner et al., 2017], which raises the question of how this average change can be partitioned into a ‘terrestrial carbon component’ and a ‘ventilation/respired carbon component’. Until this issue is resolved, along with the related issue of how sub-aerial volcanism changed across the last glacial period [Broecker et al., 2015], the possibility remains that much of the full glacial-interglacial δ$^{13}$C difference observed in ODP1240 was achieved by respired carbon addition, implying that a bigger drop in [CO$_3^{2-}$] would be expected. This, in turn, would point to an additional process that would have counterbalanced the addition of [CO$_3^{2-}$] from organic carbon respiration, such as CaCO$_3$ dissolution.

5. Conclusions

In this study, an increase in the biological pump efficiency over the LGM through a decrease in its leakiness has been investigated in the EEP as a mechanism for the glacial atmospheric CO$_2$ drawdown. The change in [CO$_3^{2-}$], δ$^{13}$C and oxygenation proxies inferred in this study support this idea by presenting lower values in the EEP at the LGM compared to the late Holocene. However, the decrease in [CO$_3^{2-}$] in this area is relatively small, suggesting that these mechanisms were not the only ones operating, or indeed contributing to the drop in glacial atmospheric CO$_2$. This has also been observed in other Pacific [CO$_3^{2-}$] records and has been suggested to be a consequence of larger carbonate dissolution over the LGM in this basin. Thus, respiratory calcite dissolution not only might explain the relatively small change in deep ocean [CO$_3^{2-}$] between the glacial and late Holocene periods, but also would have likely and directly contributed to the glacial atmospheric CO$_2$ decrease by increasing the ocean average ALK and therefore decreasing the surface ocean pCO$_2$, thus allowing the surface ocean to take up more CO$_2$. In conclusion, the deep EEP, and probably much of the wider deep Pacific, likely sequestered a significant amount of atmospheric CO$_2$ during the LGM, specifically due to a more efficient biological carbon pump caused by lower ventilation rates (and potentially also enhanced export production), but also due to an increase in average ocean alkalinity by respiratory calcite dissolution. However, a comparison with δ$^{13}$C data raises interesting questions regarding the relative contributions to the global marine carbon inventory during the last glacial period, from changes in deep ocean ventilation, export production, carbonate dissolution, terrestrial carbon release, and indeed volcanism. Further investigation in other basins using a
‘biogeochemical fingerprinting’ approach as applied in this study would help to address these questions, e.g. through better constraints on global average ocean chemistry changes at the LGM, which in turn would allow for a more accurate quantification of carbonate dissolution effects and marine CO₂ sequestration over this period.

6. References


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