Deep Atlantic carbonate ion and CaCO₃ Compensation during the Ice Ages

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Higher alkalinity is compensation for reduced CaCO₃ burial in the deep ocean in response to increased carbon sequestration in the deep ocean. This process accounts for about half of the reduction in glacial atmospheric CO₂. To date our understanding of this process comes from benthic carbon isotope and CaCO₃ burial records. Here we present a 1.5 My orbitally resolved deep ocean calcite saturation record (ΔCO₂) derived from benthic foraminiferal B/Ca ratios in the North Atlantic. Glacial ΔCO₂ declines across the mid-Pleistocene transition (MPT) suggesting increased sequestration of carbon in the deep Atlantic. The magnitude, timing, and structure of deep Atlantic Ocean ΔCO₂ parallels changes in %CaCO₃ and contrasts the small amplitude, anti-phased swings in IndoPacific ΔCO₂ and %CaCO₃ during the mid-to-late Pleistocene questioning the classic view of CaCO₃ compensatory mechanism. We propose that the increasing corrosivity of the deep Atlantic causes the locus of CaCO₃ burial to shift into the equatorial Pacific where the flux of CaCO₃ to the seafloor was sufficiently high to overcome low saturation and establish a new burial “hot spot”. Based on this mechanism, we propose that the persistently low ΔCO₂ levels at Marine Isotope Stages (MIS) 12, set the stage for the high pCO₂ levels at MIS 11 and subsequent interglacials via large swings in ocean alkalinity caused by shifts in CaCO₃ burial. Similarly, the development of classic (“anti-correlated”) CaCO₃ patterns was driven by enhanced ocean stratification and an increase in deep ocean corrosivity in response to MPT cooling.

Keywords: B/Ca, benthic foraminifera, calcite saturation, North Atlantic, carbon dioxide, Pleistocene, CaCO₃ compensation

Key Points:
- North Atlantic deep ocean calcite saturation is reconstructed using B/Ca ratios in benthic foraminifera across the mid-to-late Pleistocene
- Deep ocean ΔCO₂ levels declined across the mid-Pleistocene and G/IG ΔCO₂ amplitude increased as ΔCO₂ became persistently under-saturated 400 ky ago
- Shifts in the locus of CaCO₃ burial maintains the global CaCO₃ balance on G/IG timescales

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1. **Introduction**

Mechanisms proposed to explain Pleistocene changes in atmospheric pCO$_2$ call for a transfer of carbon dioxide from the atmosphere to the glacial deep ocean by way of increased efficiency of the biological pump (Broecker & Peng, 1982; Broecker & Peng, 1987; Boyle et al. 1988). In contrast, upwelling of respired CO$_2$ and its transfer back to the atmosphere would explain the pCO$_2$ increase during the interglacials. However, as previously pointed out biological mechanisms cannot account for the full G/IG amplitude and duration of atmospheric pCO$_2$ during the late Pleistocene (Broecker and Peng, 1982; Boyle, 1988; Broecker and Peng, 1989; Archer and Maier-Reimer 1994; Toggweiler, 1999; Sigman & Boyle, 2000). This raises the likelihood that calcium carbonate (CaCO$_3$) compensation in response to changes in the amount of carbon sequestered in the deep ocean might have had substantial contribution to the G/IG changes in pCO$_2$ (Broecker and Peng, 1982; Boyle, 1988; Broecker and Peng, 1987; Toggweiler, 1999).

In this scenario, the sequestration of excess glacial metabolic CO$_2$ reduces carbon ion (CO$_3^{2-}$) in the deep ocean. As the deep ocean CO$_3^{2-}$ is controlled by the balance between the supply of ingredients for CaCO$_3$ (calcium (Ca$^{2+}$) and bicarbonate (HCO$_3^-$)) into the ocean and removal via burial of CaCO$_3$ in the sediments, this imbalance promotes CaCO$_3$ dissolution until a new steady state is reached via adjustments in the lysocline, the depth in the ocean where CaCO$_3$ preservation (above) gives way to dissolution (below). The shoaling of the lysocline raises ocean alkalinity and acts to further lower atmosphere CO$_2$. Current CaCO$_3$ compensation theory predicts, the ocean's CO$_3^{2-}$ response should be fairly fast, within the range of 5-10 ky and atmospheric pCO$_2$ would change on the timescale set by the CO$_3^{2-}$ response time (Broecker & Peng, 1987; Keir, 1988; Archer et al. 1997; Kohler & Fischer, 2006). If this mechanism did operate on G/IG timescales across the Pleistocene, we would expect to find evidence for relevant oscillations in deep ocean carbon storage, CO$_3^{2-}$ content, lysocline in tandem with CaCO$_3$ preservation and dissolution patterns.

A second mechanism proposed to explain G/IG changes in deep ocean CO$_3^{2-}$ and CaCO$_3$ burial patterns invokes sea level driven shifts in carbonate deposition a.k.a coral reef hypothesis (Berger, 1982; Berger and Keir, 1984; Opdyke and Walker, 1992). In this scenario, during glaciation, sea level drop reduces the shelf area and CaCO$_3$ deposition on continental shelves. This would act to increase whole ocean alkalinity and deep ocean CO$_3^{2-}$ and plays a critical role in reducing atmospheric CO$_2$. This idea implies that on G/IG
timescales whole ocean alkalinity and pCO$_2$ is dictated by sea level driven shifts in the partitioning of CaCO$_3$ deposition between continental shelf and deep ocean, or more specifically enhanced glacial preservation is tied to alkalinity supplied from the shelf areas during this period of sea level fall. If this hypothesis did operate on G/IG timescales then there should be a correlation between timing and amplitude of long-term changes in deep ocean CO$_3^{2-}$ and sea level in the past.

Most of our knowledge for changes in Pleistocene deep ocean G/IG carbon inventory and sedimentary carbonate processes derives from benthic carbon isotope records ($\delta^{13}$C$_b$) and CaCO$_3$ burial and preservation fluxes. $\delta^{13}$C$_b$ records have been used to track the transfer of carbon between the organic and inorganic pools in the ocean, and records spanning the Pleistocene have also been interpreted to reflect large G/IG shifts in water mass distributions with implications for carbon storage in the deep ocean (Raymo et al. 1997; Clark et al. 2006; Skinner et al. 2009). However, interpretations of $\delta^{13}$C$_b$ records in terms of carbon storage and circulation are confounded as these proxies are impacted by a range of biogeochemical and physical processes, such as changes in preformed $\delta^{13}$C$_b$ signals of the source water (Lynch-Stieglitz et al. 1995; Raymo et al. 1997; Yu et al. 2008 and references herein). Pleistocene %CaCO$_3$ records from the deep Atlantic show enhanced dissolution during glacial periods relative to interglacials, whereas Pacific %CaCO$_3$ records indicate enhanced preservation during glacials relative to interglacials (Arhenius, 1952; Farrell & Prell, 1989; Howard & Prell, 1994; deMenocal et al. 1997; Hodell et al. 2001; Sexton & Barker, 2012 and references here in). This classic ‘seesaw’ CaCO$_3$ preservation pattern has persisted across the last 800 ky and has been suggested to vary in response to vertical repositioning of the Atlantic lysocline (Berger & Winterer, 1974; Boyle, 1983; Farrell & Prell, 1989; Howard & Prell, 1994; deMenocal et al. 1997). However, interpretations of %CaCO$_3$ records in terms of variations of deep ocean CO$_3^{2-}$ are confounded by the fact that in addition to dissolution they also reflect carbonate production and dilution by terrigenous material and thus cannot directly be related to changes in atmospheric pCO$_2$.

A robust assessment of deep ocean CO$_3^{2-}$ across Pleistocene G/IG cycles awaits the development of basin-wide records of ocean carbonate chemistry to fully resolve the nature of the ocean buffering capacity and its role in driving G/IG pCO$_2$ cycles. As the CO$_3^{2-}$ of the deep ocean is regulated by changes in mean ocean alkalinity, records of deep ocean CO$_3^{2-}$ and %CaCO$_3$ provide insights into the compensation mechanism, coral reef hypothesis, and the
ability of the ocean to buffer changes in atmospheric CO₂. Also, the CO₃²⁻ of the deep ocean’s resired carbon pool should track atmospheric pCO₂ which makes it a good proxy to investigate mechanisms affecting the carbon cycle and atmospheric pCO₂ (Broecker & Peng, 1987; Yu et al. 2010). Here we use boron to calcium ratios in the benthic foraminifer *Cibicidoides wuellerstorfi* as a means to investigate Pleistocene G/IG changes in deep North Atlantic Ocean CO₃²⁻.

Boron to calcium (B/Ca) ratios in epifaunal benthic foraminifera have been used to reconstruct seawater calcite saturation state ($\Delta[CO_3^{2-}] = [CO_3^{2-}_{in\,sinu}] - [CO_3^{2-}_{sat}]$) based on available empirical calibrations (Yu & Elderfield, 2007; Brown et al. 2011; Rae et al. 2011; Raitzsch et al. 2011; Yu et al. 2013). Overall, the available evidence from B/Ca-based $\Delta$CO₃²⁻ records show a divergent response related to depth and ocean basin across the last glacial cycle (Yu et al. 2010; Yu et al. 2013; Doss & Marchitto, 2013; Yu et al. 2014; Allen et al. 2015; Yu et al. 2016). Currently available, long Pleistocene records are mostly of low resolution documenting long-term trends in $\Delta$CO₃²⁻ but not resolving the G/IG cyclicity (Rickaby et al. 2010; Elmore et al. 2015; Kender et al. 2016) except to two recent studies. In the North Atlantic, Lear et al (2016) put forth a suite (n=80) of trace metal records (Cd/Ca, B/Ca, U/Ca) across the mid-Pleistocene (0.4 to 1.1 Ma) in Deep Sea Drilling Program (DSDP) site 607 and show a decrease in glacial B/Ca values across the MPT. As G/IG changes in δ¹³Cᵢ occur alongside shifts in nutrient content and deep ocean CO₃²⁻, the authors attributed the cause to a change from dominantly northern water mass to dominantly southern source waters across the MPT and an increase in resired carbon pool of the glacial Atlantic.

Beyond this they suggest that the increased corrosivity of bottom waters during glacialcs might have increased the ocean alkalinity and lowered atmospheric CO₂ (Lear et al. 2016). Additionally a study by Kerr et al. (2017), using orbitally resolved Indo-Pacific B/Ca records show that deep Pacific CO₃²⁻ concentration slightly increased during the glacial intervals while falling during the interglacials across the last 500 ky with similar patterns to the sediment CaCO₃ content. They argue that a global increase in CO₃²⁻ during glacial intervals is caused by the reduction of carbonate deposition on the shelves during low sea level stands (a.k.a ‘coral reef hypothesis’) across the last 450 ky (Berger, 1982; Opdyke & Walker, 1992). Here, we explore alternative mechanism, whereby the post-MPT Pacific and Atlantic CO₃²⁻ records are driven by basin-to-basin CaCO₃ compensation.
In this paper we extend the DSDP site 607 North Atlantic record of Lear et al. (2016; n=80) and increase the number of samples by 280 analyses, alongside including a nearby piston core record (Chain 84-24-23 PC) to provide a complete orbitally resolved benthic foraminifera B/Ca reconstruction for the last 1.5 My through the mid to late Pleistocene. Combined with existing IndoPacific B/Ca records (Kerr et al., 2017), carbon isotope, and %CaCO$_3$ records (Ruddiman et al. 1989; Raymo et al. 1997) we investigate mechanisms controlling the deep ocean carbon budget of the North Atlantic and examine the physical and biogeochemical processes that can explain the observed G/IG changes in ΔCO$_3^{2-}$. The focus on the deep Atlantic stems from geochemical and modeling evidence suggesting the deep North Atlantic has played a significant role in carbon sequestration on G/IG timescales during the late Pleistocene (Toggweiler, 1999; Hain et al. 2010; Yu et al. 2016; Howe et al. 2016). We discuss the importance of the CaCO$_3$ cycle for the G/IG pCO$_2$ change in light of this 1.5 My Atlantic ΔCO$_3^{2-}$ stack and propose that the enhancement of atmospheric CO$_2$ by CaCO$_3$ compensation is due to a change in the locus of CaCO$_3$ burial between the Atlantic and IndoPacific in response to the transfer of alkalinity between these ocean basins. Our new record also provides new insights into the mechanisms associated with the mid-Pleistocene transition (MPT) and mid-Brunhes Event (MBE).

2. Core material and methods

2.1 Hydrographic conditions & Age Model

We present benthic foraminiferal B/Ca records for Deep Sea Drilling Program (DSDP) site 607 (41°00’N; 32°57’W, 3427 m) on the western flank of the Mid-Atlantic Ridge supplemented with measurements from a nearby piston core (Chain 82-24-23 Piston Core (referred to hereafter as CHN82-24-23PC; 43°N, 31°W; water depth 3406 m; Figure 1). Both sites are situated within the path of North Atlantic Deep Water (NADW; T=2.6 °C, S=34.1) close to its northern sources. In the modern ocean, bottom water at these sites reflects the influence mostly of NADW (low nutrient, high δ$^{13}$C$_b$ and ΔCO$_3^{2-}$) and to a lesser extent Antarctic Bottom Water (AABW; enriched nutrient, low δ$^{13}$C$_b$ and ΔCO$_3^{2-}$) (Yu et al. 2008) (Figure 2). Modern ΔCO$_3^{2-}$ (33.2 µmol/kg) are estimated from nearby GEOSECS data. The age model for both sites are based on LRO4 stack Lisiecki & Raymo (2005) and previously used in Sosdian & Rosenthal (2009).
2.2 Analytical Method

Well-preserved benthic foraminifera of the species *Cibicidoides wuellerstorfi* were picked from >150 μm fraction at sampling intervals of 15 cm providing an average temporal resolution of ~3 ky. Picked samples restricted to the morphology of *C. wuellerstorfi sensu stricto* for B/Ca analyses throughout the core (Rae et al. 2011). Pleistocene records of B/Ca should mainly reflect deep ocean ∆CO$_3^{2-}$ variability due to the relatively long oceanic residence time of B (~10 My) and Ca (~1 My) and their conservative nature (LeMarchand et al., 2002). *C. wuellerstorfi* B/Ca were converted to deep sea ∆CO$_3^{2-}$ using a sensitivity of 1.14 µmol/mol per µmol/kg specific to this species obtained from the global core-top calibration from Yu et al. (2007) where B/Ca (µmol/mol)=1.14(±0.04) x ∆CO$_3^{2-}$ (µmol/kg) +177.(±1.0).

Samples were gently crushed between two glass plates and metal oxides visible within chambers were removed from the sample to minimize contamination. The fragments were transferred into acid-cleaned, dry 0.5 ml micro-centrifuge tubes. Crushed foraminiferal samples contained 3-15 individuals and ideally weighed from 0.200 mg to 0.400 mg, but due to low species abundance in certain intervals samples occasionally weighed < 0.200 mg. Samples were cleaned using the reductive or ‘long’ cleaning protocol (Boyle and Keigwin, 1985/6; Rosenthal et al., 1997). The effect of different cleaning methods on B/Ca ratios is minimal (Yu et al., 2007). Samples were cleaned and dissolved using reagents prepared with boron-free water as prepared by passing double distilled water over an anion exchange resin (Aldrich, amberlite IRA-743) to remove boron. Following cleaning, samples were dissolved in trace metal clean 0.065 N Nitric Acid (SEASTAR) and 100 µl of this solution was diluted with 300 µl of trace metal clean 0.5 N Nitric Acid to obtain a Ca concentration of about 3±1 mmol mol$^{-1}$. Diluted samples were analyzed for a suite of trace metal ratios using the Rutgers Sector Field Inductively Coupled Plasma-Mass Spectrometer (ICP-MS), on a Finnigan MAT Element-1 and Element-2 XR located at the Department of Marine and Coastal Sciences using methods adapted from Rosenthal et al., (1999). The method was slightly modified to accommodate for measurement of B/Ca ratios, specifically, ammonia gas was introduced in the Teflon-spray chamber during sample analysis to remove the memory effect associated with analysis of boron (Babila et al., 2014). Long-term precision, based on analysis of consistency standards across a range of B/Ca ratios is ±4%. Analysis runs including CHN82-24-23PC B/Ca records were offset from the long-term mean B/Ca, as determined by the long-term average of the consistency standards, and to account for this we applied an analytical
offset to CHN82-24-23PC B/Ca values of 5% prior to including them in Figure 3. Al, and Ti were used as contaminant indicators and no long-term correspondence in trend exists between B/Ca and each elemental ratio. However, across MIS 25, high B/Ca values correspond with elevated Al and Ti and were removed from the final record.

2.3 Cross-Spectral Analysis & Trends in glacial and interglacial maxima

Here we use the ARAND package (Howell, 2001) to evaluate the phase and coherency between elemental and isotopic records from DSDP 607 and compare variables relative to benthic δ18O in 100-ky world (i.e. from 140 to 750 ka, 100 ky periodicity). We interpolated all records to even intervals of 3-ky resolution prior to cross-spectral analysis and used the inverse of benthic δ18O in the analysis. Proxy phase relationships are compared and presented in the representative time window during the “100 ky world” of the late Pleistocene.

To explore the trends in glacial and interglacial extremes in B/Ca ratios, we define glacial and interglacial maxima across the last 1.5 My from the benthic oxygen isotope record. ΔCO3²° maxima and minima were picked within a ka window around the interglacial or glacial peak, to allow for uncertainty associated with identifying the interglacial or glacial maxima for each site, the tuning approach, and to account for changes in the length and structure of interglacial periods (Tzedakis et al., 2009). G-I amplitude is taken as the absolute value of the difference between successive average glacial/interglacial values (e.g. G1-IG1, G2-IG1, G2-IG2).

3. Results

3.1 Down core B/Ca record

During the last 1.5 My, C. wuellerstorfi B/Ca at site 607 exhibits G/IG variability with a decrease in average B/Ca minima of 21 µmol/mol across the Pleistocene mostly driven by changes in the glacial values primarily decreasing at MIS 23 from 180 µmol/mol to an average 161 across MIS 20-18. Glacial B/Ca then increase to 179 µmol/mol at MIS 14. At MIS 12, glacial B/Ca values decrease by 25 µmol/mol and was at this level also during MIS 10 and 6 (Figure 3C, 4). In contrast, interglacial B/Ca ratios show a small long-term decline throughout the record resulting in the G/IG amplitude varying across the Pleistocene; during the pre-MPT interval (MIS 49 to 25, 1500 to 950 ka), the average G/IG amplitude in B/Ca is ~24 µmol/mol (IG: 227±4; G: 203±5; ±1 standard errors of the mean, SEE). The G/IG
amplitude increases to 48 (IG: 224±3; G: 170±1; ±1 SEE) µmol/mol across MIS 23 to 13 and throughout the late Pleistocene interval. Interglacial B/Ca values decrease at MIS 23 from average steady values of 227 to 206 µmol/mol at MIS 19 and increase at MIS 17 to average IG value of 227 µmol/mol until MIS 11 where it declines to average IG value of 200 µmol/mol where it remains steady for the rest of the Pleistocene (Figure 4). Notably, the B/Ca record shows a remarkable correlation with the benthic foraminiferal δ13C record both on G/IG and longer time scales (Ruddiman et al. 1989; Figure 3B,C).

Our piston core record encompasses the 165 to 75 and 55 to 10 ka interval. The CHN82-24-23PC B/Ca variability generally agrees with available records across these overlapping time intervals. For comparison, we look toward the published *C. wuellerstorfi* B/Ca records from Yu et al. (2008, 2016) from the deep North Atlantic across the LGM to Holocene (BOFS 8K; water depth 4045 m) and from 60 to 90 ka (MD95-2039; MD01-2446; Figure 3C). CHN82-24-23PC B/Ca tracks the rise in MIS 5 and decline into MIS 4 as shown in records from MD01-2446 and MD95-2039. The BOFS 8K record shows a similar amplitude change in B/Ca (ΔG/IG=43; IG: 213; G:170) µmol/mol to the G/IG change exhibited across the last 400 ky from site 607 (Figure S3). LGM CHN82-24-23PC B/Ca values do not reach the low values of BOFS 8K as the record is of lower resolution and likely not capturing the minimum in B/Ca.

The record of deep Atlantic B/Ca at site 607 from this study agrees well with the low resolution MPT record presented by Lear et al. (2016; Figure 3C). Therefore, we compile their data alongside our record and use this to examine the spectral properties of the record alongside examining deep ocean ΔCO3^2− changes. The B/Ca record exhibits orbital frequencies similar to the δ18O record with a strong 100 ky peak present in the 140 to 700 ka time interval, following the MPT (Figure S2). Prior to the MPT, from 1000-1500 ka, the record shows precession-like periodicities (18, 21 ky) but does not resolve a 41 ky cyclicity, possibly due to small gaps in the record. Also, B/Ca changes lag δ 18Ob by 3 ky similar to the δ 13Cb and %NCW record in the 100 ky world (Figure S3; Table S1) whereas %CaCO3 lags δ 18O by 8 ky.

Moving forward we combine all B/Ca records from this study (DSDP 607; CHN82-24-23PC) and previous studies (DSDP 607; BOFS 8K; MD95-2039; MD01-2446) from the deep North
Atlantic to produce a composite Atlantic B/Ca record. (Figure 4D). Applying the calibration of Yu et al. (2007) yields a long-term change in $\Delta CO_3^{2-}$ of 20 µmol/kg with a G/IG amplitude shift of 23 µmol/kg prior to the MPT (1000-1500 ka), and 38 µmol/kg post MPT (0-750 ka; Figure 5D; calibration uncertainty ±9 µmol/kg). Interglacial $\Delta CO_3^{2-}$ values decrease by 10 µmol/kg from the pre- to post- MPT with average interglacial levels of the last 4 cycles similar or slightly higher than modern $\Delta CO_3^{2-}$ of 33 µmol/kg at site 607. Glacial $\Delta CO_3^{2-}$ levels decreased by 27 µmol/kg from pre- to post- MPT conditions. Our $\Delta CO_3^{2-}$ record shows distinct variations on G/IG timescales indicating modifications in deep North Atlantic Ocean carbon storage across the Pleistocene. The range of deepwater $\Delta CO_3^{2-}$ variability (>30 µmol/kg), recorded at Site 607 exhibits changes beyond the modern $\Delta CO_3^{2-}$ range of the NADW and AABW end-members suggesting that in addition to shifts in water mass distribution, other processes (discussed section 4.1) are driving the observed G/IG cycles in $\Delta CO_3^{2-}$.

Changes in CaCO$_3$ content in deep North Atlantic sediments have been variably related to changes in biological production at the surface, dissolution/preservation on the seafloor and dilution by terrigenous inputs (Ruddiman et al. 1989). LGM CaCO$_3$ content in various sites in the deep Atlantic has been variably linked to lower CaCO$_3$ production and higher dissolution alongside a higher terrigenous input (Crowley, 1983; Francois et al. 1990; Francois & Bacon, 1991; Barker et al., 2004; Sexton & Barker, 2012). However, at site 607 the $\Delta CO_3^{2-}$ records change in parallel with %CaCO$_3$ suggesting that $\Delta CO_3^{2-}$ ion variations are largely associated with the dissolution and preservation cycles in the deep North Atlantic (Figure 5). Deep Atlantic $\Delta CO_3^{2-}$ values observed across the Pleistocene are consistent with Atlantic carbonate preservation records showing high preservation during interglacials and enhanced dissolution during glacials (Crowley, 1983; Ruddiman et al. 1989; Sexton & Barker 2012). Below (see section 4.2), we further explore the relationship between CaCO$_3$ preservation dissolution cycles and deep ocean $\Delta CO_3^{2-}$ content.

4. Discussion

4.1 Atlantic Deep ocean $\Delta CO_3^{2-}$ evolution

Benthic $\delta^{13}C$ in the deep North Atlantic varies in response to changes in the amount of organic carbon/nutrient, the influence of NADW, air-sea gas exchange and shifts in mean ocean $\delta^{13}C$ (Raymo et al., 1997; Raymo, 2004). In present-day, site 607 is bathed by NADW, a well-ventilated water mass with high $\delta^{13}C_b$ (>1.0‰) and in saturated conditions
(ΔCO$_3^{2-}$=33 µmol/kg). In comparison, southern sourced water mass, Antarctic Bottom Water (AABW), is relatively undersaturated (ΔCO$_3^{2-}$=0 µmol/kg) and has lower δ$^{13}$C$_b$ (∼ 0.4‰) due to higher nutrient content. The ΔCO$_3^{2-}$ and δ$^{13}$C$_b$ records closely covary across the last 1 My and display similar shifts during the 1000 to 800 ka interval, thereby suggesting that waxing and waning of northern and southern component waters played a key role in deep Atlantic ΔCO$_3^{2-}$ as previously proposed for the last glacial maximum and recently for the Pleistocene (Marchitto et al. 2005; Yu et al. 2008; Lear et al. 2016; Figure 5). Pena & Goldstein (2014), using ε$_{Nd}$ proxy, show a decrease in the relative proportion of NADW versus southern sourced water from 1000 to 900 ka, supporting the notion of a greater contribution of Southern Component Water (SCW) at Site 607. However, the benthic δ$^{13}$C and ΔCO$_3^{2-}$ composition of source waters is also impacted by air-sea exchange driven by temperature, and CO$_2$ outgassing and invasion in the Southern Ocean and North Atlantic (Lynch-Stieglitz et al. 1995). Considering the impact of air-sea gas exchange across the MPT, Lear et al (2016) estimated that the increase in δ$^{13}$C$_b$ partially reflects reduction of outgassing due to an increase in sea ice extent in the Southern Ocean leading to a decline in SCW δ$^{13}$C$_b$ and ΔCO$_3^{2-}$ alongside the increase in SCW at expense of Northern Component Water (NCW) at site 607.

The decrease in δ$^{13}$C$_b$ from 1000 to 800 ka is associated with a decrease in ΔCO$_3^{2-}$, which suggests the incursions of SCW were associated with lower ΔCO$_3^{2-}$ and increased nutrient content, supported by higher Cd/Ca during glacials from a reconstruction at Site 607 (Lear et al. 2016). To examine this further and determine whether this reflects changes in corrosivity of southern sourced deep waters or changes in the volume of SSW, we compare our ΔCO$_3^{2-}$ record to a %NCW record (Raymo et al. 1997; Lang et al. 2016; Figure 5A, D). The % NCW record documents the circulation component at site 607 scaled in terms of relative changes in preformed δ$^{13}$C$_b$ composition of NADW (ODP Site 982) and δ$^{13}$C$_b$ composition of deep Pacific (ODP Site 849).

From 1000 to 800 ka, the glacial decrease evident in the ΔCO$_3^{2-}$ record, with a pronounced decline mirrored in both the decrease of δ$^{13}$C$_b$ and %NCW record, suggest an increase in SCW penetration to this site, as supported by ε$_{Nd}$ records from Pena & Goldstein (2014). Pleistocene vertical δ$^{13}$C$_b$ gradients between intermediate and deep waters in the South Atlantic show that since 1100 ka a sharp chemocline developed existed between mid-depth
and deep waters during glacial periods. This sharp chemocline separated well-ventilated intermediate waters from poorly ventilated deep waters (Hodell et al. 2003) consistent with the decrease in $\Delta CO_3^{2-}$ recorded at Site 607. Using a stacked magnetic susceptibility record Schmieder et al. (2000) show that during the MPT interim state (from 900 to 600 ka), CaCO$_3$ accumulation rates in the subtropical South Atlantic are notably reduced suggesting greater influence of Antarctic bottom waters. Using planktonic foraminifera fragmentation index, Groger et al. (2003) show the enhancement of G/IG preservation and dissolution cycles at that time. These lines of evidence indicate that changes in our B/Ca-$\Delta CO_3^{2-}$ record across the MPT are related to an increase of SCW at the expense of NCW.

Following the MPT, from MIS 20 to 14, both records show similar shifts in glacial values with glacial $\Delta CO_3^{2-}$ increasing until MIS 12 paralleling the increase in glacial $\delta^{13}C_b$ values. Interglacial $\Delta CO_3^{2-}$ slightly increase from MIS 21 to 17 and then plateau in tandem with the interglacial $\delta^{13}C_b$ trend (Figures 4, 5). Both features suggest a tight coupling on G/IG timescales. In contrast, the %NCW record shows only a small (~10%) increase in glacial maxima across this time interval. Thus, the inferred small decrease in the volumetric contribution of SCW likely contributed negligibly to the increase $\Delta CO_3^{2-}$ during this time interval. Instead, the close correspondence between the glacial increase in $\Delta CO_3^{2-}$ (by about 20 $\mu$mol/kg) and the increase in glacial $\delta^{13}C_b$ suggests that both were driven by whole ocean changes in carbon. From MIS 15 onwards, Lear et al. (2016) propose that the interglacial increase in $\delta^{13}C_b$ is related to reduced interglacial CO$_2$ evasion into North Atlantic source waters, impacting both the $\delta^{13}C_b$ and $\Delta CO_3^{2-}$ records. Following the MPT, $\Delta CO_3^{2-}$ remains at relatively similar levels suggesting air-sea gas exchange minimally impacted the interglacial $\Delta CO_3^{2-}$ North Atlantic source waters (Figure 4).

Stacked planktic and benthic $\delta^{13}C$ foraminifera records, show a decrease by 0.35‰ around 900 ka, gradual return to pre-1 Ma values at 500 ka and a final rapid decrease at 420 ka (Wang et al. 2004; Hoogakker et al. 2006; Lisiecki, 2010). The global long-term increase of 0.6 per mil lasting until 500 ka and the subsequent decrease at MIS 12 likely reflect changes in the carbon burial to the deep ocean modified by changes in the rain ratio, bottom water ventilation, and surface ocean production (Wang et al. 2004; Hoogakker et al. 2006; Lisiecki, 2010). The similarity between the $\Delta CO_3^{2-}$ and $\delta^{13}C_b$ records both in the long-term trends and G/IG variability throughout this time window suggests it is recording changes in oceanic carbon-carbonate system and changes in deep ocean circulation.
4.2 Deep ocean CO$_3^{2-}$ and CaCO$_3$ preservation/dissolution cycles

A recent study, using orbitally resolved IndoPacific B/Ca records show that over the past 500 ky, deep Pacific $\Delta$CO$_3^{2-}$ G/IG variability parallels changes in sediment %CaCO$_3$ content, apparently in contrast with the expectation of anticorrelation between these properties derived from the classic compensation theory (Kerr et al., 2017). To explain the apparent contradiction between the expectation from the compensation theory and their observed records, Kerr et al., (2017) argue that a global increase in CO$_3^{2-}$ during glacial intervals is caused by the delivery of alkalinity from the shelves due to the reduction of carbonate deposition on the shelves during low sea level stands (a.k.a ‘coral reef hypothesis’) across the last 450 ky (Berger, 1982; Opdyke & Walker, 1992). Here, we propose an alternative mechanism, whereby the post-MPT Pacific and Atlantic CO$_3^{2-}$ records are driven by basin-to-basin alkalinity transfer, rather than the classic vertical CaCO$_3$ compensation.

Over the past 400 ky, the Atlantic $\Delta$CO$_3^{2-}$ record shows G/IG swings of 30-40 µmol/kg in the 100-ky band which decline into the glacial and recover across the terminations (Fig. 5D; 6D). In contrast, the IndoPacific records show smaller (<10 µmol/kg) changes in $\Delta$CO$_3^{2-}$ as reported by Kerr et al. (2017) (Figure 6C), which rise into the glacial and shoot down across the terminations. This behavior is also evident in the %CaCO$_3$ records from the Atlantic, IndoPacific, and Southern Ocean, in support of our deep ocean $\Delta$CO$_3^{2-}$ records (Figure 5C;6A,B). The nature of the deep ocean $\Delta$CO$_3^{2-}$ records contrasts the expected changes as estimated from box models and predicted by CaCO$_3$ compensation theory (Boyle, 1988; Broecker & Peng, 1989; Toggweiler, 1999; Toggweiler, 2008).

Using a seven-box model Toggweiler (2008), estimates that in response to deep ocean ventilation the response to expected change in whole ocean $\Delta$CO$_3^{2-}$ below 1000 m, is on the order of ~9 µmol/kg, significantly lower than $\Delta$CO$_3^{2-}$ swings observed at DSDP site 607 across the last 400 ky (Figure 7). Also, the structure of the observed $\Delta$CO$_3^{2-}$ changes lacks the expected spike up on the terminations and down on glacial initiations followed by rapid return to the steady state baseline as expected from the classic compensation theory. Additionally, the large swings in Atlantic $\Delta$CO$_3^{2-}$ are not compensated quickly as expected and the deep Atlantic CO$_3^{2-}$ remains above or below its long-term mean for periods substantially greater than the assumed response time of ~5-10 kyrs (Figure 5D;6D; Broecker & Peng, 1987; Keir, 1988; Archer et al. 1997; Kohler & Fischer, 2006). It is noteworthy that the $\Delta$CO$_3^{2-}$ reconstructions show that both the deep Atlantic and Pacific were undersaturated.
with respect to calcite during glacial intervals suggesting that dissolution was not the only control on CaCO$_3$ preservation in the deep glacial ocean. The difference in the structure of G/IG change, magnitude, and relative timing of G/IG cycles between the Atlantic and Pacific deep $\Delta$CO$_3^{2-}$ records questions the classic view of CaCO$_3$ compensatory mechanism and calls for a different approach (Toggweiler, 1999; 2008).

We propose that over the past ~1 My, CaCO$_3$ compensation occurred in response to changes in the places where CaCO$_3$ is buried rather than vertical shifts in the lysocline. This explains the Atlantic-Pacific difference and the contrast to the box model predictions. We hypothesize that deep ocean $\Delta$CO$_3^{2-}$ and $\%$CaCO$_3$ were driven by changes in corrosivity of the deep North Atlantic and interbasin migration of calcite burial or ‘hot spots’ across G/IG cycles (Dunne et al. 2012). A "hot spot" is an area of the ocean where the enhanced production of CaCO$_3$ at the surface overlies a favorable burial environment on the sea floor. The net result is a local burial flux of CaCO$_3$ that is higher than average. The concept is important because the cumulative burial in a few large hot spots could, in principle, balance the delivery of Ca$^{2+}$ and HCO$_3^-$ ions to the ocean from rivers. At present, the main hotspot or locus of CaCO$_3$ burial is in the deep Atlantic and much of the global burial of CaCO$_3$ occurs there due to the formation of the youngest, least corrosive deep waters in this region (see Fig. 1c in Dunne et al. 2012). In contrast, the Pacific Ocean has a less propensity for CaCO$_3$ burial due to the oldest, undersaturated deep waters residing there. Also, in the present day the production and flux of CaCO$_3$ to the sea floor is not uniform across the ocean and burial of CaCO$_3$ is more of a kinetic process than a purely equilibrium process (Li et al. 1969; Broecker & Peng, 1982; Dunne et al. 2012). In this sense, favorable burial environments can also include areas where the water above the sea floor is mildly undersaturated with respect to calcite and enhanced fluxes of CaCO$_3$ to the sea floor overwhelm the dissolution in the sediments (due to the slow kinetics of the dissolution reactions). This non-uniformity, we claim, creates the potential for "hot spots" of CaCO$_3$ burial and preservation, which result when enhanced fluxes to the seafloor coincide with a relatively shallow seafloor and/or the presence of NADW. In this instance, all the CaCO$_3$ production in areas outside the hot spots tends to dissolve.

Here we build upon these observations and propose that incursion of dense corrosive waters from the southern hemisphere into the glacial deep Atlantic decreased Atlantic calcite burial thus raising the ocean’s average CO$_3^{2-}$. We propose this is compensated by a shift in the locus of CaCO$_3$ burial into the equatorial Pacific to compensate for the declining burial in the other
areas. This is exhibited during glacials, when $\Delta \text{CO}_3^{2-}$ and %CaCO$_3$ records show an increase in glacial deep Atlantic corrosivity and undersaturation that inhibits burial across the last 800 ky (e.g. MIS 12) (Fig. 6B, D). Also, at the same time, empirically, it seems that the locus of burial shifts to the tropical Pacific evident from sedimentary carbonate records which shows that the area of CaCO$_3$-rich sediments expands in the Pacific during the LGM and shrinks everywhere else (Farrell & Prell, 1989; Catubig et al. 1998). This suggests that the Pacific became a hot spot for CaCO$_3$ burial despite the undersaturated state during glacials. We propose this is likely caused by the following effects i) increase in CaCO$_3$ export that overcame the apparent corrosivity of the water and ii) the depth at which major CaCO$_3$ dissolution starts occurs is not $\Delta \text{CO}_3^{2-}$=0 but in fact lower. This is consistent with Subhas et al. (2017) suggesting that CaCO$_3$ dissolution starts only at a saturation state of 0.7 not 1. Therefore, it seems that the small increase in deep Pacific $\Delta \text{CO}_3^{2-}$ was sufficient to shift the balance from carbonates dissolution to preservation. In this way, the $\Delta \text{CO}_3^{2-}$ in the deep IndoPacific rises in response to the same mechanism that keeps the $\Delta \text{CO}_3^{2-}$ in the deep Atlantic low and the atmospheric CO$_2$ low.

When the locus of burial shifts to the Pacific, the ocean's average CO$_3^{2-}$ and alkalinity of the ocean must increase to compensate, or rise to a point where the burial in the new hot spot is sufficiently high to overcome the general corrosivity of the deep Pacific. These excursions lead to a larger compensation response that is spread out over longer periods of time and would make the amount of alkalinity and dissolved inorganic carbon in the ocean swing widely over time. This is reflected in the slow rise of $\Delta \text{CO}_3^{2-}$ through stages 4, 3, and 2 in TT013-PC72 (Kerr et al. 2017) and in the late glacial increases in %CaCO$_3$ at site 1089, a stand in for burial in the Indo Pacific (Figure 6A). The gradual increase in alkalinity over this time helps keep the pCO$_2$ of the atmosphere low for tens of thousands of years and the ocean cold, which reinforces the stratification of the deep Atlantic and times when the $\Delta \text{CO}_3^{2-}$ falls over time at site 607 as seen particularly well during MIS 12 and 10. So, instead of the CO$_3^{2-}$ simply ticking up and down within a G/IG transition, as in the classical response, a shift of the locus of CaCO$_3$ burial tends to hold the deep CO$_3^{2-}$ up or down over entire glacial or interglacials.

Conversely, when the locus of burial shifts back to the Atlantic during interglacials, the ocean's average CO$_3^{2-}$ and alkalinity of the ocean falls because it is suddenly easy to bury
CaCO₃ again in the less corrosive Atlantic. This seems to be seen especially well during Terminations II and I at VM 28-122 in Yu et al. (2013) and in the %CaCO₃ decreases at site 1089 (Figure 6A). The net loss of alkalinity from the ocean helps keep the atmospheric pCO₂ high and consequently the ocean warm during interglacials. Also, CO₃²⁻ in the deep Atlantic is “super-saturated” during IG intervals and as such needs to decrease a lot to reach undersaturation and start dissolution as expressed by the large swings in ΔCO₃²⁻ (30-40 µmol/kg).

The response of %CaCO₃ and deep ocean CO₃²⁻ content to variations in the Atlantic meridional overturning circulation (AMOC) strength has been previously explored in several modeling studies (Chikamoto et al. 2008; Yu et al. 2016). Chikamoto et al. (2008) show that during an ‘AMOC shutdown’, initiated by a freshwater perturbation, the average carbonate ion concentration in the North Atlantic is reduced by ~37 µmol/kg and the CaCO₃ burial flux is reduced in the North Atlantic and increased in the Pacific. In addition, Yu et al. (2016) investigate the influence of AMOC changes on deep Atlantic CO₃²⁻ and show that a cessation of NADW leads to a decrease in deep (>3 km) North Atlantic and Equatorial Atlantic CO₃²⁻ by 20–40 µmol/kg. Although these model results provide some justification for our argument at least, the Chikamoto et al (2008) model results have a negligible effect on atmospheric pCO₂ in contrast to our conjecture that G-I compensation changes played a role in atmospheric pCO₂. We suggest here that their model does not fully incorporate regional production differences in CaCO₃, which when in combination with favorable burial environments produces hotspots, does not replicate fully the scenario where the burial of CaCO₃ shifts from one ocean to another and concomitant G-I changes in pCO₂.

In previous studies it has been argued that sea-level variations drive the cycles of CaCO₃ burial and dissolution in the ocean and can explain the low amplitude variations in the Pacific and glacial increase in ΔCO₃²⁻ and covariation in %CaCO₃ and ΔCO₃²⁻ (Yu et al., 2013; Kerr et al. 2017). Kerr et al. (2017) use the broad correlation of the amplitude of glacial increase in CO₃²⁻ with the amplitude of sea-level decline as support. In doing so, the authors suggest that shifts in Atlantic ΔCO₃²⁻ are less sensitive to whole ocean carbonate system changes, as past water mass distribution are strong in this region and impacting deep carbon storage. Furthermore, the ΔCO₃ records presented here resemble the δ¹³Cₗ (proxy of carbon storage and circulation) rather than the δ¹⁸Oₗ record (proxy of ice volume and sea level). This is very
pronounced between MIS 20 and 14 and then 12 and 6 when glacial $\Delta CO_3$ follow closely the $\delta^{13}C_b$ (Figure 5). These lines of evidence suggest that changes in carbon storage in the deep Atlantic are tightly coupled to the very thing that is driving the variations in atmospheric CO$_2$, namely changes in the ventilation of southern-sourced bottom water. Building on this idea, we argue that hypothesized enhanced preservation, due to alkalinity supplied from the shelf areas during sea level fall, could affect the Atlantic but not enough to overcome the dissolution changes caused by shifts in ventilation/carbon storage. Therefore, we argue that CaCO$_3$ dissolution in the Atlantic was the dominant source of alkalinity to the deep Pacific and hence played the key role in controlling atmospheric CO$_2$ across mid-to-late Pleistocene. Indeed, CaCO$_3$ compensation, as viewed from changes in deep water $\Delta CO_3^2$ occurred throughout the past 1.5 My and apparently scaled with the changes in atmospheric pCO$_2$ (Figure 5).

Across the MPT around 1.1 Ma, the classic (‘anti-correlated’) CaCO$_3$ patterns (i.e. dissolution in Atlantic and preservation in Pacific during glacials) developed. Sexton & Barker (2012) suggest this was due to the strengthening of deepwater ventilation in the abyssal Pacific during glacials and weakening during IGs, which led to glacial Pacific CaCO$_3$ dissolution to diminish while driving IG Pacific CaCO$_3$ dissolution to intensify. The proposed idea by these authors would, however, let the respired CO$_2$ in the deep Pacific escape to the atmosphere during glacials. Here we propose an alternative scenario whereby the development of extremely poor ventilation and increased stratification in the deep Atlantic provides a better explanation for the Pleistocene CaCO$_3$ observations and atmospheric CO$_2$. Deep ocean cooling starts at Site 607 in the North Atlantic at ~1.2 Ma (Sosdian and Rosenthal 2009, Ford et al., 2016), while based on the record from ODP Site 1123, the deep glacial Southern Ocean was close to freezing throughout this period (Elderfield et al., 2012). While it is possible that some of the oxygen isotopic change of seawater change that Elderfield et al (2012) document at the MPT reflects increase salinity, either way the isotope data reflect an increase in the density of the Southern Ocean deep waters with increased stratification and hence weaker ventilation as seen in the last glacial period in the South Atlantic (Adkins et al. 2001). The cooling of deep water at Site 607 (Sosdian and Rosenthal 2009, Ford et al., 2016) likely reflects increased stratification of the deep Atlantic. A colder ocean led to northward migration of the Antarctic polar fronts and likely resulted in thicker and more expanded southern sea ice (Kemp et al., 2010) that made the deep ocean more saline more stratified and less ventilated over time. We propose the
initial cause of the anti-phased behavior between the Atlantic and Pacific relates to the deep ocean cooling across the MPT that led to changes in ventilation and drove the deep Atlantic into under-saturation consistent with the appearance of anti-phased behavior in %CaCO₃, as pinpointed by Sexton & Barker (2012) at ~1.1 Ma.

Following the cooling at 1.2 Ma, both the δ¹³C₀ and %NCW decline from 1 to 0.8 Ma highlighting changes in ventilation. The coherency between the δ¹³C₀ and Atlantic ∆CO₃²⁻ record across the MPT indicates changes in deep ocean ventilation are driving deep ocean CO₃²⁻ changes (Figure 3; 5). Thus, across the MPT, we propose that the increase in deep ocean corrosivity led to the development of the classic CaCO₃ pattern whereby the locus of CaCO₃ burial shifts to the Pacific during glacial and switches back the Atlantic during interglacials.

4.3 The Mid-Bruhnes Event
Glacial ∆CO₃²⁻ levels drop by ~ 30 µmol/kg after MIS 14, becoming for the first time under-saturated through the last 1.5 My in the deep North Atlantic at MIS 12, when it reaches -20±9 µmol/kg (Figure 5; 6). This period of corrosive deep waters corresponds with the mid-Brunhes dissolution event, beginning at ~600 ka and lasting for 400 ky, peaking around ~400 ka (MIS 11), exhibited by records showing enhanced dissolution in the deep ocean occurring in the North Atlantic (Crowley, 1985; Jansen et al. 1986; Groger et al. 2003; Barker et al. 2006), in the Indian Ocean (Bassinot et al. 1994), and in the Pacific (Farrell & Prell, 1989).

The decrease in the saturation in the deep North Atlantic occurs alongside the development of the largest ice sheets of the Pleistocene, cooling of the deep Southern Ocean and North Atlantic (as recorded in ODP Site 1123 and DSDP Site 607; Elderfield et al., 2012; Sosdian and Rosenthal, 2009) and intensification of the 100 ky cycles in sea level and pCO₂ (Lisiecki & Raymo, 2005; Hönisch et al. 2009; Bereiter et al. 2015). Cooling of the deep Atlantic, which culminated at MIS 12, likely resulted in greater stratification and sluggish ventilation, which enhanced carbon sequestration in the deep Atlantic possible augmented by increased surface ocean productivity (Lawrence et al. 2013).

The % NCW record shows a prolonged period of near 0% levels, the longest duration of reduced NCWs across the last 800 ky (Figure 5). Records of ocean circulation show
exceptionally prolonged (5 ky) collapse of the Atlantic Meridional Overturning Circulation (Riveiros et al. 2013) and increased influence of SSW (Hall & Becker, 2007) at MIS 12 suggesting a longer residence time of seawater in the glacial deep Atlantic. The isolation of these deep waters allowed greater accumulation of respired organic carbon as surface ocean productivity was higher at Site 607/U1313 as indicated by alkenone mass accumulate rates (C$_{37}$ total MAR) and total organic carbon records (Stein et al. 2009; Lawrence et al. 2013). Both records indicate higher surface ocean productivity possibly tied to the more southerly position of the polar front and/or increased dust supply (Stein et al. 2009; Lawrence et al. 2013).

Modeling studies show that a 50% NADW reduction leads to a decrease of ~30 µmol/kg in the deep Atlantic (>3 km) and NADW cessation leads to a greater decrease of 20-40 µmol/kg (Yu et al. 2016). We propose here that additional decrease in ΔCO$_{3}^{2-}$ minima starting at MIS 12, set the stage for the high pCO$_{2}$ levels at the mid-Bruhnes event via an increase in whole ocean alkalinity followed by enhanced CaCO$_{3}$ preservation at MIS 11, 9 and MIS 5. Specifically, we propose that whole ocean alkalinity became higher than "normal" during MIS 12, in response to cooling and reduced ventilation of the deep Atlantic, which caused increase corrosivity. Then, when the deep ocean became better ventilated during termination V, more CaCO$_{3}$ was preserved everywhere for a short period of time. The enhanced removal of alkalinity then pushed the atmospheric pCO$_{2}$ higher. The larger response time than previously assumed contributes to this effect (i.e. the excess alkalinity is not completely removed during the deglaciation but throughout the interglacial maintaining the high pCO$_{2}$ levels). This suggests that IG pCO$_{2}$ levels are partly determined by the preceding G/IG changes and the response time of the compensation. A noteworthy corollary of the proposed mechanism, is that the lower limit of the glacial pCO$_{2}$, which has been ~180ppmV during the past 800 ky, is not strictly set by the ocean alkalinity and consequently the “extra” alkalinity generated at the corrosive glacials (MIS 11, 9, 5) was available for removal during the subsequent interglacials (e.g., MIS 11, 9, 5) leading to higher pCO$_{2}$ levels.

5. Conclusions
To understand the nature of the ocean’s buffering capacity and its role in modulating pCO$_{2}$, reconstructions of the deep ocean carbonate system parameters are necessary.
Pleistocene benthic foraminiferal B/Ca record from DSDP site 607 suggests 30-40 µmol/kg G/IG variability in deep ocean \(\Delta \text{CO}_3^{2-}\), substantially larger than seen in the Indo-Pacific records. The observed timing, magnitude and structure of swings in G/IG deep Atlantic ocean \(\Delta \text{CO}_3^{2-}\) are also larger than expected based on thermodynamic CaCO₃ compensation mechanism. To explain this difference, we suggest that during glacialals, when ventilation weakens, the more corrosive and lower \(\Delta \text{CO}_3^{2-}\) in the deep Atlantic, shifts the locus of CaCO₃ burial to the equatorial Pacific to maintain the long-term CaCO₃ balance via a new carbonate burial ‘hot spot’, where the carbonate flux to the seafloor and the small increase in \(\Delta \text{CO}_3^{2-}\) are sufficient to overcome the under-saturated conditions characteristic of the deep Pacific during interglacial intervals. We argue that, the small increase in IndoPacific \(\Delta \text{CO}_3^{2-}\) is driven by the same mechanism that keeps the deep Atlantic \(\Delta \text{CO}_3^{2-}\) and atmospheric CO₂ low and is not due to the delivery of alkalinity from the shelves during low sea level stands.

Shifts in the global carbon cycle and deep ocean circulation play a role in driving the long-term trend in Atlantic \(\Delta \text{CO}_3^{2-}\) across the mid-to-late Pleistocene. The \(\Delta \text{CO}_3^{2-}\) record from DSDP site 607 shows shifts in G/IG variability associated with the MPT and mid-Brunhes event. The basin-basin fractionation likely initiated across the MPT as the deep Atlantic Ocean began to play a large role in enhancing pCO₂ due to changes in ventilation, likely due to cooling and enhanced stratification, which drove deep Atlantic into under-saturation during glacial intervals. Deep ocean \(\Delta \text{CO}_3^{2-}\) levels reached the lowest levels of the last 1.5 My at Marine Isotope Stage (MIS) 12 and became persistently under-saturated. We propose the low \(\Delta \text{CO}_3^{2-}\) of MIS 12 were possibly initiated by a combination of sluggish circulation and high surface ocean productivity allowing for increased storage of CO₂ in the deep Atlantic. The low \(\Delta \text{CO}_3^{2-}\) of the deep Atlantic set the stage for the high pCO₂ levels at MIS 11 via an increase in whole ocean alkalinity followed by enhanced CaCO₃ preservation at MIS 11 and the subsequent interglacials of MIS 9 and 5.

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Figure 1 Ocean Data Viewer bathymetric map showing modern locations of DSDP site 607 (depth=3.4 km; red symbol) used in this study and from previously published data (grey/black symbols; BOFS8K depth=4.0 km; MD95-2039 depth=3.4 km; MD01-2446 depth=3.6 km; ODP 1089 depth=4.6 km; TT013-PC72 depth=4.3 km) are shown in the map (Schlitzer, 2012).
Figure 2 Deep CO₃²⁻ profiles from (A) the North Atlantic near DSDP site 607 and (B) equatorial IndoPacific, nearby site TT013-PC72, used in Kerr et al. (2017) B/Ca record. In the modern ocean, basin-specific differences in deep ocean CO₃²⁻ occur as a result of the formation of North Atlantic Deep Water (NADW), which fills the deep Atlantic today with preservation-friendly, high-CO₃²⁻ water whereas the deep Pacific is filled with corrosive low-CO₃²⁻ water in response. The saturation state (Δ[CO₃²⁻]) of the deep ocean modulates the depth in the ocean where CaCO₃ preservation (above) gives way to dissolution (below) termed the lysocline whereby its position is deeper in the Atlantic with more CaCO₃ burial per unit area relative to the Pacific (Dunne et al. 2012). Carbonate system data are from nearby GLODAP sites (Key et al. 2004)
Figure 3  DSDP site 607 records of benthic foraminiferal (A) $\delta^{18}$O and (B) $\delta^{13}$C (Ruddiman et al., 1989) and (C) B/Ca ratios from this study (DSDP 607 and CHN82-24-23PC) and previously published deep North Atlantic records from BOFS 8K (Yu et al. 2008), MD95-2039 (Yu et al. 2016), MD01-2446 (Yu et al. 2016) and DSDP site 607 (Lear et al. 2016). All records are on the LRO4 age model and MIS are highlighted based on LRO4 stack alignment (Lisiecki & Raymo, 2005). Error bars show analytical precision for B/Ca (r.s.d. = 4.0%).
Figure 4 Interglacial and glacial B/Ca trends across the mid-to-late Pleistocene determined from the composite DSDP 607 and CHN82-24-23PC records. Data from BOFS 8K are used to estimate G/IG change across the H-LGM. Linear trends in panel A showcase the overall long-term decrease in interglacial and glacial B/Ca values. Note the steeper decrease in glacial relative to interglacial B/Ca values.
Figure 5 DSDP site 607 (A) %NCW at site 607 and the re-occupation site U1313 plotted with 95% confidence intervals (Raymo et al. 1997; Lang et al. 2016) (B) δ¹³C₆ (Ruddiman et al. 1989) (C) %CaCO₃ from DSDP site 607 and VM30-97 (Ruddiman et al. 1989), (D) compiled deep North Atlantic ocean B/Ca and ΔCO₃⁻ from DSDP 607 and CHN82-24-23PC and others (Yu et al. 2008; Yu et al. 2016; Lear et al. 2016) and (E) atmospheric CO₂ reconstructions from ice cores (Petit et al. 1999; Siegenthaler et al. 2005; Luethi et al. 2008; Bereiter et al. 2015), blue ice (Higgins et al. 2015) and boron isotopes (Hönisch et al. 2009; Chalk et al. 2017). Error bars show ΔCO₃⁻ uncertainty window of ±9 µmol/kg from Yu et al. (2007) calibration. Boxes around the 1 Ma data indicate an age uncertainty of ±89ky as presented in Higgins et al. (2015). The Mid-Brunhes Event and Mid-Pleistocene transition are denoted as MBE and MPT.
Figure 6 %CaCO₃ records from (A) ODP 1089 (Hodell et al. 2001) in the deep South Atlantic representing the deep Pacific and (B) DSDP 607 and VM30-97 (Ruddiman et al. 1989) in the deep North Atlantic alongside B/Ca-based ∆CO₂⁻² reconstruction from the (C) equatorial Indo-Pacific (core TT013-PC72; Kerr et al. 2017) and (D) compiled deep North Atlantic ∆CO₂⁻² with (E) atmospheric CO₂ ice core (Petit et al. 1999; Siegenthaler et al. 2005; Luethi et al. 2008; Bereiter et al. 2015) and B-isotope derived reconstructions (Hönisch et al. 2009; Chalk et al. 2017). Dashed line on panel D represents 0 µmol/kg. Error bars show ∆CO₂⁻² uncertainty window of ±9 µmol/kg from Yu et al. (2007) calibration. Note inverted axis in panel B.
Figure 7 Deep CO$_3^{2-}$ cycles across the Pleistocene as estimated by a seven-box model presented in Toggweiler (2008; see Figure 4) showcasing the classic CaCO$_3$ compensation mechanism. Note the time axis is inverted.