Constraining the evolution of Neogene ocean carbonate chemistry using the boron isotope pH proxy

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Over the course of the Neogene, the Earth underwent profound climatic shifts from the sustained warmth of the middle Miocene to the development of Plio-Pleistocene glacial–interglacial cycles. Major perturbations in the global carbon cycle have occurred alongside these shifts, however the lack of long-term carbonate system reconstructions currently limits our understanding of the link between changes in CO2, carbon cycling, and climate over this time interval. Here we reconstruct continuous surface ocean pH, CO2, and surface ocean aragonite saturation state using boron isotopes from the planktonic foraminifer Triloculina trilobus and we perform a sensitivity analysis of the key variables in our calculations (e.g. δ11Bsw, [Ca]sw, CCD). We show that the choice of δ11Bsw influences both seawater pH and CO2 while [Ca]sw reconstructed dissolved inorganic carboxy exerts a significant influence only on CO2. Over the last 22 Myr, the lowest pH levels occurred in the Middle Miocene Climate Optimum (MMCO; 17–14 Myr ago) reaching ~7.6 ± 0.1 units in all our scenarios. The extended warmth of the MMCO corresponds to mean CO2 and aragonite saturation state levels of 470–630 ppm and 2.7–3.5, respectively. Despite a general correspondence between our CO2 record and climate, all CO2 scenarios show a peak at ~9 Ma not matched by corresponding changes in climate reconstructions. This may suggest decoupling (i.e. significant CO2 change without a discernible climate response) for a limited interval in the Late Miocene (11.6–8.5 Ma), although further refinement of our understanding of the temporal evolution of the boron isotopic composition of seawater is necessary to fully evaluate the nature of the relationship between CO2 and climate. Nonetheless, from our long-term view it is clear that low-latitude open ocean marine ecosystems are unlikely to have experienced sustained surface pH and saturation levels below 7.7 and 1.7, respectively, during the past 14 million years (66% CI).

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1. Introduction

Boron isotopes (δ11B) in marine carbonates have been exploited as a proxy for the marine carbonate system (i.e. pH, pCO2) on multiple timescales within the Cenozoic (e.g. Sanyal et al., 1995; Palmer et al., 1998; Pearson and Palmer, 2000; Foster et al., 2012; Pennan et al., 2013; Anagnostou et al., 2016; Gutjahr et al., 2017; Chalk et al., 2017). Plio-Pleistocene foraminiferal δ11B derived reconstructions exhibit glacial–interglacial variability in pH and pCO2, in good agreement with independent records (e.g. ice cores) (Seki et al., 2010; Bartoli et al., 2011; Hönisch et al., 2009; Henehan et al., 2013; Martínez-Boz et al., 2015a; Chalk et al., 2017). However, application of this proxy beyond the Plio-Pleistocene is not straightforward as calculation of pH and pCO2 requires records of seawater chemistry (e.g. δ11Bsw, [Ca]sw) and a second carbonate system parameter (e.g. total alkalinity [ALK], or dissolved inorganic carbon [DIC]) which currently have considerable uncertainties (Horita et al., 2002; Brennan et al., 2013; Pälike et al., 2012; Raitzsch and Hönisch, 2013; Greenop et al., 2017). Current reconstructions of the major ion composition of seawater, which impacts boron isotope reconstructions through influencing key dissociation constants and estimates of carbonate (CO32−) concentration, rely on low-resolution fluid inclusion measurements (Horita et al., 2002; Brennan et al., 2013) and improved estimates await development of higher resolution records from a range of proxies. For the early Cenozoic, the likely upper and lower bounds of δ11Bsw have been estimated using the δ11B–pH relationship, whilst estimates of the second carbonate system parameter

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have been determined using Earth System modelling (Anagnostou et al., 2016; Gutjahr et al., 2017). According to Anagnostou et al. (2016), the early Eocene CO2 concentration was 1400 ± 470 ppm decreasing to 550 ± 190 ppm in the early Oligocene. The uncertainty window on these estimates reflects uncertainty in calcite saturation and δ11Bw estimates, δ11B measurement error, and possible atmosphere-ocean disequilibrium. Despite these relatively large uncertainties, it is clear that early Cenozoic pCO2 was much higher than in the Plio-Pleistocene and CO2 change was an important driver of climate change in the early Cenozoic. However, application of the δ11B-pCO2 proxy during the more recent Neogene (2.6 to 23 Myrs ago) transition to a bipolar icehouse world is complicated by the fact that most reconstructions indicate a relatively dramatic shift in δ11Bw in the late Miocene to Pliocene. While there is first-order agreement between different records for the late Miocene δ11Bw rise, the exact timing, magnitude and rate of change of this shift remains uncertain (Pearson and Palmer, 2000; Lemarchand et al., 2002; Raitsch and Hönisch, 2013; Greenop et al., 2017), with implications for the structure of δ11B-based reconstructions of pCO2 during the period. Furthermore, the history of carbonate chemistry (i.e. DIC and alkalinity) in this interval is poorly understood and currently is limited to model estimates that use pCO2 estimates from GEOCARB carbon cycle models and/or paleo-pCO2 proxy estimates paired with estimates of the calcite compensation depth (CCD) (Tyrrell and Zeebe, 2004; Ridgwell, 2005).

During the Neogene period, the Earth underwent profound changes from the warmth of the Middle Miocene Climatic Optimum (~14–17 Myrs ago; MMCO) to the expansion of the Antarctic ice sheet at 13.9 Ma (Lear et al., 2010). Dynamic changes in the carbon cycle have been proposed to be associated with these climatic shifts, involving changes in silicate weathering, volcanic emissions and the organic carbon subcycle (Derry and France-Lanord, 1996; France-Lanord and Derry, 1997; Raymo and Rudiman, 1992; Raymo, 1994; Foster et al., 2012). The evolution of the carbonate system during this time interval therefore represents a critical gap in our understanding of climate-carbon cycle interactions. Previous CO2 reconstructions derived from a disparate set of proxies (e.g. plant stomata, marine alkenones, paleosols, marine carbonate δ11B) have produced a wide range of estimates, yet with a general consensus that MMCO CO2 levels were higher than preindustrial, and that CO2 subsequently declined across the mid-Miocene climate transition (Kürschner et al., 2008; Foster et al., 2012; Zhang et al., 2013). Since CO2 and pH are highly correlated variables in the ocean carbonate system, the discrepancies between existing CO2 reconstructions (Pagani et al., 1999a, 1999b; Foster et al., 2012; Zhang et al., 2013) imply a lack in understanding of the ocean’s history of pH change and the evolution of the carbon cycle and also, to some extent, an uncertainty in proxy systematics. Further, the degree of coupling between climate and CO2 in the late Miocene is poorly constrained, limiting our ability to determine the importance of CO2 variability in setting the evolution of Miocene climate (Pagani et al., 1999a, 1999b; LaRiviere et al., 2012). Thus, the lack of available records that constrain the ocean’s carbonate chemistry is a critical impediment to establishing the relationship between the long-term Earth System evolution of carbon cycle and climate (Hönisch et al., 2012). Additionally, without detailed knowledge of long-term changes in seawater carbonate chemistry we are blind to its role in driving evolutionary adaptation and resilience of marine organisms to environmental change (Hönisch et al., 2012; Hain et al., 2015).

Here we use a new dataset of 40 planktic boron isotope measurements in conjunction with existing benthic data and new constraints on the evolution of seawater δ11B (Lemarchand et al., 2002; Raitsch and Hönisch, 2013; Greenop et al., 2017) to reconstruct surface pH and ocean carbon content for the past 22 million years. Combining our surface pH record with independent reconstructions of seawater calcium and magnesium concentrations, changes in the calcite compensation depth, deep-sea pH, and temperature, we calculate both atmospheric CO2 and surface aragonite saturation state (Ωaragon). We also discuss the relative importance of the variable sources of uncertainty in these parameters on our reconstructions. With extensive uncertainty propagation using a Monte Carlo approach and sensitivity analysis we reconstruct long-term pH, aragonite saturation state and atmospheric CO2 since the MMR (0–17 Myrs ago). Our new record of ocean carbonate chemistry allows us to examine the role of CO2 in the climate system over the course of the Neogene and contextualize future carbon emission pathways against this geological perspective.

2. Materials and methods

The boron isotope proxy relies on the principle that the ratio of the dominant boron species in seawater, borate ion and boric acid, varies as a function of pH (Dickson, 1990), which also controls the isotopic composition of the borate ion that is incorporated into a marine carbonate such as a planktic foraminiferal calcite (Vengosh et al., 1991; Hemming and Hanson, 1992; Foster and Rae, 2016). The foraminifer-derived pH record may then be used to estimate CO2 levels and Ωaragon using assumptions about other carbonate system parameters. Early attempts at reconstructing pH and atmospheric CO2 from fossil foraminifera over this interval (Spivack et al., 1993; Palmer et al., 1998; Pearson and Palmer, 2000) are qualitatively useful but have been superseded by a revised understanding of boron isotope systematics and analytical developments (see discussion in Seki et al., 2010; Foster et al., 2012; Anagnostou et al., 2016).

2.1. Sample locations and site details

In this study we analysed the CaCO3 shells of a single planktic foraminifera species, *Trilobatus trilobus*. This species was formerly assigned to the genus *Globigerinoids* but is no longer considered part of that genus (Suesszaferri et al., 2015). The species is sometimes referred to as *T. sacculifer*, especially those individuals that bear a final sac-shaped final chamber, but the feature is ecophenotypic (André et al., 2013) and the name *trilobus* has priority. In this study we avoided individuals with sac-shaped chambers. Usually for planktic foraminifera, *T. trilobus* displays no significant cryptic genetic diversity (André et al., 2013). It lives in the upper part of the water column in association with dinoflagellate symbionts, although it probably sinks towards the end of its life cycle at gametogenesis, somewhat complicating the shell chemistry (e.g. Birch et al., 2013). The species is widely used in boron isotope studies because it is one of the few to survive almost the entire Neogene and Quaternary periods (Suesszaferri et al., 2015) and a species-specific calibration for δ11B-pH exists (discussed further below; Sanyal et al., 2001; Seki et al., 2010; Foster et al., 2012; Greenop et al., 2014).

We analysed the δ11B of 40 samples of this species from geographically diverse locations including Ocean Drilling Program (ODP) Site 926 (n = 26; western tropical North Atlantic), ODP Site 761 (n = 3; tropical Indian Ocean), ODP Site 872 (n = 7; tropical Pacific Ocean), and ODP Site 1000 (n = 4; Caribbean Sea). This dataset supplements previously published *T. trilobus* analyses from ODP Site 668 (Hönisch et al., 2009), ODP Site 761 (Foster et al., 2012; Greenop et al., 2014), ODP Site 926 (Foster et al., 2012), ODP Site 999 (Bartoli et al., 2011; Seki et al., 2010; Caribbean Sea), ODP 1264 (Stap et al., 2016) and outcrops of the Blue Clay formation in Malta (Badger et al., 2013) and previously published *Globigeri-
noides ruber analyses from ODP 999 (Martinez-Boti et al., 2015a; Chalk et al., 2017) (Fig. 1). Sea surface temperatures (SSTs) were reconstructed from these sites. ODP Site 761 (16°3′S, 115°3′E, 2179 m water depth) is situated in the Indian Ocean on the Wombat Plateau. ODP Site 926 (3°N, 42°W, 3598 m water depth) in the western equatorial Atlantic Ocean on Ceara Rise. ODP Site 872 (10°N, 162°E, 1287 m water depth) is in the tropical north Pacific gyre on the sedimentary caps of flat-topped seamounts. ODP Site 1000 (16°N, 79°W, 927 m water depth) is situated in the Caribbean Sea on the Nicarguan Rise on a drowned carbonate platform.

Our sites are far removed from coastal influences and sources of upwelling, and likely remained close to equilibrium with atmospheric CO2 and therefore variations in atmospheric CO2 are likely the dominant control on δ13C variability at all study sites (Pearson and Palmer, 2000; Takahashi et al., 2009, 2017; Hönisch et al., 2009; Bartoli et al., 2011; Seki et al., 2010; Foster et al., 2012; Badger et al., 2013; Greenop et al., 2014; Martínez-Boti et al., 2015a; Chalk et al., 2017) (Fig. 1). Although foraminifer shells from these sites (except Malta) exhibit the recrystallized textures common in deep sea pelagic sites (e.g. Sexton et al., 2006), existing evidence suggests that the boron isotopic signal remains intact within the foraminifer test during recrystallisation, hence the process does not strongly affect δ11B (Foster et al., 2012; Badger et al., 2013; Edgar et al., 2015).

2.2. Age model

Our age model for ODP Site 761 uses biostratigraphic datums (Holbourn et al., 2004) on the CK95 timescale (Cande and Kent, 1995; Supplementary Table S1). The age model for ODP Site 926 is a polynomial fit through nanofossil and planktic foraminifer biostratigraphic datums based on the shipboard report (Site 926 Shipboard Scientific Party, 1995) and post-cruise refinement (Chaisson and Pearson, 1997; Pearson and Chaisson, 1997) adapted to the CK95 timescale. Ages for ODP Site 872 was calculated by linear interpolation between reliable biostratigraphic datums (Pearson, 1995) from the early to mid-Miocene. Note that for ODP Site 872, age estimates for sample depths 105.71 and 106.79 mbsf have large uncertainty (±1 Ma) as evidence for reworking is present in this section, however this is not significant at the resolution of this study for the early Miocene. Nanofossil and planktic foraminifer biostratigraphic datums tied to the CK95 timescale were used for the age model at ODP Site 1000A (Site 1000 Shipboard Scientific Party, 1997a, 1997b). The age models from published boron isotope records from ODP Sites 761 (Foster et al., 2012; Greenop et al., 2014), 926 (Foster et al., 2012), and the Blue Clay Formation (Badger et al., 2013) were put on the CK95 timescale in a similar manner using shipboard reports, published biostratigraphic datums (Holbourn et al., 2004; Site 926 Shipboard Scientific Party, 1995; Site 999 Shipboard Scientific Party, 1997a, 1997b; Hilgen et al., 2009). Previously published records from the Plio-Pleistocene (<5 Myr) (Hönisch et al., 2009; Seki et al., 2010; Bartoli et al., 2011; Martínez-Boit et al., 2015a; Stap et al., 2016; Chalk et al., 2017) are on their original orbitally tied age models.

2.3. Analytical methodology

Between 100 and 150 shells of T. trilobus were picked from the 300–355 μm size fraction and cleaned following the oxidative cleaning method (Barker et al., 2003). Samples were dissolved in 0.5M HNO3 and stored for boron isotope analysis in Teflon vials. Following Foster (2008) and as detailed in Foster et al. (2013), boron separation from the calcium matrix was performed using the boron specific resin Amberlite IRA 743. Boron isotope analysis was performed on a Thermo-Fisher Scientific Neptune multi collector inductively coupled plasma mass spectrometer (MC-ICPMS) at the University of Southampton with a sample-standard bracketing routine (Foster et al., 2013; Henehan et al., 2013).

External reproducibility is calculated following Rae et al. (2011) as follows:

\[ 2\sigma = 2.25 \exp^{-23.01[11B]} + 0.28 \exp^{-0.64[11B]} \]  
\[ 2\sigma = 33450 \exp^{-168.21[11B]} + 0.311 \exp^{-1.477[11B]} \]  

where \([11B]\) is the intensity of \(^{11}B\) signal in volts and equation (1) and equation (2) used with 10\(^{-11}\) and 10\(^{-12}\) rulers, respectively (δ\(^{11}\)B\(_\text{planktic}\) ± 0.22–1.00%). In house standards (JCP-1; Hathorne et al., 2013) were processed with each column batch (n = 4) and these and boric acid standards repeatedly analysed during each analytical session were consistent with the long-term reproducibility (Supplementary Table S2).

Calibration studies have shown that the foraminifer species is predominantly incorporated in the calcium carbonate shell and thus the boron isotopic composition of foraminifer can be used to estimate ocean pH (Vengosh et al., 1991; Hemming and Hanson, 1992; Sanyal et al., 2001). To calculate pH from the δ\(^{11}\)B of T. trilobus we account for the species-specific offset between δ\(^{11}\)B\(_\text{borate}\) in seawater and δ\(^{11}\)B of the foraminifer calcite due to vital effects (equation (3)) (Foster and Rae, 2016; Henehan et al., 2016). We use the calibration slope from Sanyal et al. (2001) calculated following Martínez-Boit et al. (2015b) with the intercept based on core-top δ\(^{11}\)B T. suecicus 300–355 μm data from Seki et al. (2010) with estimates of surface water borate from Henehan et al. (2016):

\[ \delta^{11}\text{B}_{\text{borate}} = (\delta^{11}\text{B}_{\text{trilobus}} - 2.69)/0.833 \]  

This species-specific calibration curve is used to estimate the δ\(^{11}\)B of borate ion. The measurement uncertainty of δ\(^{11}\)B\(_\text{trilobus}\) estimated by the external reproducibility is fully propagated with an appropriately scaled normal distribution.
Here we compile our *T. trilobus* δ\(^{11}\)B data with previously published δ\(^{11}\)B records from the Miocene (Foster et al., 2012; Badger et al., 2013; Greenop et al., 2014) and Plio-Pleistocene (Fig. 2B; Hönsch et al., 2009; Bartoli et al., 2011; Seki et al., 2010; Martínez-Boití et al., 2015a; Stap et al., 2016; Chalk et al., 2017). To estimate δ\(^{11}\)B\(_{\text{borate}}\) and surface ocean pH, we applied size-fraction and species-specific δ\(^{11}\)B\(_{\text{borate}}\)-δ\(^{11}\)B calibrations (Supplementary Table S3). As previously published δ\(^{11}\)B analyses in this compilation were performed using a range of techniques, we apply inter-instrument offsets to the published negative thermal ionization mass spectrometry (N-TIMS) data to correct to the MC-ICPMS and in doing so account for the well-documented instrument dependent offset in these NTIMS data (Foster et al., 2013; Farmer et al., 2016) due to differences in chemical composition of sample (i.e. B/Ca) and other reasons that are currently unclear (Farmer et al., 2016). Some δ\(^{11}\)B data from Hönsch et al. (2009) were performed on two different TIMS, the NBS design instrument at Stony Brook and Thermo Scientific Triton at LDEO. To account for this and correct the data to the MC-ICPMS, we use an offset of \(-1.1\%\), as determined from an interlaboratory comparison of five modern and fossil samples measured on both the NBS design and Triton NTIMS (Hönisch et al., 2009). To align the δ\(^{11}\)B datasets collected on the Triton (Hönisch et al., 2009; Bartoli et al., 2011; Stap et al., 2016) with the Neptune MC-ICPMS used here, we adopt the \(-1.12\%\) offset presented in Farmer et al. (2016) based on average of 5 fossil samples measured on both the Triton NTIMS and Neptune (Supplementary Fig. S1). Using this approach, the degree of the δ\(^{11}\)B\(_{\text{borate}}\) overlap between datasets from both instruments is still relatively poor in the region 2.5–3.0 Ma. To overcome this, we estimate pH and then tune the NTIMS offsets for overlapping pH datasets until there is a good agreement between NTIMS and MC-ICPMS. The best fit is achieved when we apply an additional \(-1.0\%\) offset to the Triton planktic δ\(^{11}\)B datasets. This additional offset suggests that further work is needed to provide a mechanistic understanding of the offset between MC-ICPMS and NTIMS techniques. We do not include the dataset of Stap et al. (2016) in the compilation as even with this treatment of the NTIMS offset the δ\(^{11}\)B\(_{\text{borate}}\) values exhibit substantial scatter and do not align with any of the other records (Supplementary Fig. S1). We applied no offsets to the benthic δ\(^{11}\)B data from Greenop et al. (2017) and Raitzsch and Hönsch (2013), measured on the MC-ICPMS and NTIMS, respectively, as the data agree well in their raw state (see Greenop et al., 2017).

Tandem trace element analyses were performed on a small aliquot (<10%) of each sample measured here on the Thermo-Fisher Scientific Element 2 single collector inductively coupled plasma mass spectrometer at University of Southampton following the method outlined in Henahan et al. (2013). Over the duration of this study, analytical reproducibility for Mg/Ca was ±2.7% (2σ). *T. trilobus* Mg/Ca values were converted to sea surface temperatures (SST) using the calibration of Anand et al. (2003) with a correction for seawater Mg/Ca (Horita et al., 2002; Brennan et al., 2013) following Hasiuk and Lohmann (2010) using a power constant of 0.41 (Delaney et al., 1985; Evans and Mueller, 2012).

![Fig. 2](image_url)

**Fig. 2.** (A) Benthic foraminiferal oxygen isotope (δ\(^{18}\)O) record reflecting changes in global climate (Zachos et al., 2008); Labels indicate the timing of the Middle Miocene Climate Optimum (MMCO), Middle Miocene Climate Transition (MMCT), and Northern Hemisphere Glaciation (NHG); (B) Boron isotopic composition (δ\(^{11}\)B) of planktonic foraminifer *T. trilobus* (300–335 μm) record from this study (blue) and published data from *T. trilobus* and *G. ruber* (Hönisch et al., 2005; Seki et al., 2010; Bartoli et al., 2011; Foster et al., 2012; Badger et al., 2013; Greenop et al., 2014; Martínez-Boití et al., 2015a; Chalk et al., 2017) (grey). Note the inverted axis with low δ\(^{11}\)B corresponding to low pH and pCO\(_2\). Error bars denote analytical external reproducibility (2 s.d.); (C) Sea surface temperatures (SSTs) derived from foraminiferal Mg/Ca from this study (blue) and published SST data derived from Mg/Ca (Hönisch et al., 2009; Bartoli et al., 2011; Foster et al., 2012; Badger et al., 2013; Greenop et al., 2014; Martínez-Boití et al., 2015a; Stap et al., 2016; Chalk et al., 2017) and alkenone U\(^{37}\)F values (Seki et al., 2010) (grey). Sites are coloured as in (B); (D) δ\(^{11}\)B\(_{\text{borate}}\) reconstructions from Lemarchand et al. (2012) with ±1% confidence interval, Raitzsch and Hönsch (2013) using the fractionation factor of Klochko et al. (2006) and 2 sigma uncertainty, and Greenop et al. (2017) with 95% confidence intervals. These reconstructions are referred hereafter as \(5\delta^{11}B_{\text{borate}}\text{-LO2}^{\circ}\text{-LO2}\), \(5\delta^{11}B_{\text{borate}}\text{-RH13}\text{ and 5}\delta^{11}B_{\text{borate}}\text{-G17}\). (E) Surface ocean pH reconstructions using the three \(5\delta^{11}B_{\text{borate}}\) reconstructions in (D). Note there is an additional \(5\delta^{11}B_{\text{borate}}\text{-RH13}\) reconstruction presented in Raitzsch and Hönsch (2013) not shown here calculated using an apparent fractionation factor. Here we use the Klochko et al. (2006) derived \(5\delta^{11}B_{\text{borate}}\) value to be consistent with the \(5\delta^{11}B\) calculations in this study. 66% confidence intervals are plotted. Dashed lines represent \(5\delta^{11}B_{\text{borate}}\) estimates from >17 Ma to highlight uncertainties associated with the \(5\delta^{11}B_{\text{borate}}\) reconstruction. Pre-industrial and 2017 levels are highlighted by blue and black asterisks respectively on the pH axis. Pleistocene and Eocene pH estimates are shown as vertical bars on the right side of panel E (Hönisch et al., 2009; Foster, 2008; Anagnostou et al., 2015). (For interpretation of the colours in the figure(s), the reader is referred to the web version of this article.)
mented Fig. S2). SST estimates were calculated in a similar manner for previously published Mg/Ca records (Hönisch et al., 2009; Seki et al., 2010; Bartoli et al., 2011; Foster et al., 2012; Badger et al., 2013; Greenop et al., 2014; Martínez-Botí et al., 2015a; Chalk et al., 2017). The following equations were used to correct for Mg/Ca seawater and estimate temperature (in units of °C) from foraminiferal Mg/Ca ([Mg/Ca]sw):

\[
T(°C) = \ln \left( \frac{[Mg/Ca]_{foram}}{0.38[Mg/Ca]_{sw,c}} \right) / 0.99
\]

\[
[Mg/Ca]_{sw,c} = \left( \frac{[Mg/Ca]_{sw,a}}{[Mg/Ca]_{sw,m}} \right)^{0.41}
\]

where equation (4) is the generic planktic foraminifera calibration of Anand et al. (2003) with a correction applied to the seawater Mg/Ca ratio to account for past Mg/Ca changes. This correction, given in equation (5), requires the seawater Mg/Ca for the age of the sample ([Mg/Ca]_{sw,a}) and modern seawater Mg/Ca for which the calibration was established ([Mg/Ca]_{sw,m}) to yield the corrected ratio for the paleotemperature equation [Mg/Ca]_{sw,c}. Trace element data were also used to check for foraminiferal cleaning efficiency (Barker et al., 2003). In all cases Al/Ca was <100 μmol/mol and in the majority of cases Mn/Ca was <1000 μmol/mol indicating cleaning was efficient. Some Mn/Ca values were higher than our 1000 μmol/mol threshold, for example at ODP site 926, but in all cases they do not covary with Mg/Ca or δ¹³B (Supplementary Fig. S3). The ODP site 761 Mg/Ca presented in this study is of too low resolution to judge the influence of contamination through correlation of Mg/Ca and Mn/Ca so instead we look to benthic foraminiferal Mg/Ca data from the same site which shows a lack of correlation between Mn/Ca and Mg/Ca (see Lear et al., 2015 for further discussion); ODP site 872 shows no positive correlation between Mn/Ca and Mg/Ca.

2.4. Solving for pH and the carbonate system

To convert our reconstruction of surface δ¹¹B_{Borate} into pH, CO₂ and carbonate saturation state requires three steps: (1) calculate the acid/base chemistry equilibrium constants based on independently known temperature, salinity and seawater major ion composition; (2) calculate surface water pH using the appropriate equilibrium constants, reconstructed δ¹¹B_{Borate} and independently estimated seawater δ¹³B; and (3) calculate surface water carbonate chemistry based on reconstructed pH and an independently known carbonate system parameter. The uncertainties associated with the various input parameters need to be propagated through the full set of calculations. The input parameters are presented in Fig. 3 and below we describe these calculations in more detail.

All required acid/base chemistry equilibrium constants, including the boric acid dissociation constant K_{b,b}, are calculated using the MyAMI model (Hain et al., 2015) based on reconstructed SST, sea surface salinity (SSS), water depth, and seawater major ion composition. We note that the calcium-to-bicarbonate interaction coefficient β^0(Ca:HCO₃) used in MyAMI was updated from Harvie et al. (1984: \(β^0(Ca:HCO₃) = 0.4\)) and Pytkowicz and Hawley (1974: \(β^0(Ca:HCO₃) = 2.977\)), see their Table 1) to He and Morse (1993: \(β_{HCO₃}^{(i)} = 0.2\) and \(β_{HCO₃}^{(i)} = 0.3\), see their Table 5) to bring MyAMI into agreement with experimental data of Pytkowicz and Hawley (1974). SST estimates are derived from Mg/Ca ratios from T. trilobus for ODP Sites 761, 926, 872, 1000A, and Malta from this study and previously published records (Foster et al., 2012; Badger et al., 2013; Greenop et al., 2014). SST estimates for ODP Site 999 (Seki et al., 2010) are based on a published record derived from the U_{147} proxy. In our CO₂ calculations we assign an uncertainty in SST of ±1 °C (1σ), which is propagated into
these calculations. Salinity has minimal influence on pH calculations ($\pm 1$ psu = $\pm 0.006$ pH units) and thus we assume modern salinity at each site, estimated from nearby GLODAP stations (Key et al., 2004), with an uncertainty of $\pm 1$ psu (1$\sigma$). The MyAMI calculated conditional equilibrium constants used for carbon chemistry calculations account for secular seawater [$Ca^{2+}$] and [$Mg^{2+}$] changes. As further described in section 3.2 below, we carry out all calculations for two distinct [$Ca^{2+}$] and [$Mg^{2+}$] scenarios: (a) constant modern seawater major ion composition, and (b) seawater [$Ca^{2+}$] and [$Mg^{2+}$] changing with time according to the reconstructions of Horita et al. (2002) and Brennan et al. (2013). In both seawater major ion scenarios we attach a nominal uncertainty of 2.25 mmol/kg (1$\sigma$) to both [$Ca^{2+}$] and [$Mg^{2+}$].

To calculate surface pH from our boron isotope data we follow previous work (e.g., Vengosh et al., 1991; Foster and Rae, 2016) using eqns. (6) and (7):

$$\delta^{11}B = \left[ \frac{^{11}B^{10}B_{\text{sample}}}{^{11}B^{10}B_{\text{NIST505}}} - 1 \right] \times 1000$$

$$pH = pK_B^{*} - \log \left( \frac{\delta^{11}B_{\text{sample}} - \delta^{11}B_{\text{borate}}}{\delta^{11}B_{\text{SW}} - \left( \sigma_B \times \delta^{11}B_{\text{borate}} \right) - 1000 - \left( \sigma_B - 1 \right)} \right)$$

where $\delta^{11}B$ is the ratio of $^{11}$B and $^{10}$B expressed in delta notation (in permil, ‰), $^{11}B^{10}B_{\text{NIST505}}$ is the isotopic ratio of the NIST SRM 951 boric acid standard ($^{11}B^{10}B = 4.04367$) (Catanazzo et al., 1970). The boric acid dissociation constant $pK_B^{*}$ is calculated as described above. The boron isotope fractionation factor $\sigma_B$ is taken to be 1.0272 (Klochko et al., 2006). The isotopic composition of borate, $\delta^{11}B_{\text{borate}}$, is based on our $T.\ \text{trilobus}$ data and calculation equation (3) and previously published planktonic $\delta^{11}B$ data and specified calibration equations (Supplementary Table S3). As further described in section 3.2 below, we carry out all calculations for three distinct scenarios for secular change in seawater boron isotopic composition ($\delta^{11}B_{\text{SW}}$) using the reconstructions by (a) Lemarchand et al. (2002) (“L02”), (b) Raitzsch and Hönsch (2013) (“RH13”) and (c) Greenop et al. (2017) (“G17”). Each record is used with its respective reported uncertainty assuming normal distribution.

To calculate CO$_2$ and carbonate saturation state ($\Omega_{\text{calcite}}$) from our pH reconstruction requires independent knowledge of ambient dissolved inorganic carbon concentration DIC, or any other carbonate system parameter:

$$\text{CO}_2 = \frac{\text{DIC}}{K_0^{*}} \times \frac{[H^+]^2}{[H^+]^2 + [H^+]K_1^* + K_1^*K_2^*}$$

$$\Omega_{\text{calcite}} = \frac{[Ca^{2+}]^{2+}}{K_{\text{CaCO}_3}^{*}} \times \frac{\text{DIC}}{[H^+]^2 + [H^+K_1^* + K_1^*K_2^*]}$$

The different $K^*$ terms represent the equilibrium constants governing carbonate chemistry, which we can calculate using the MyAMI model as described above. The [$Ca^{2+}$] concentration is internally consistent with the MyAMI calculations for both seawater major ion composition scenarios described above (i.e., constant modern or based on Horita et al., 2002 and Brennan et al., 2013).

The dissolved inorganic carbon concentration of seawater, DIC, is not generally known through Earth history, and given residence time constraints it may have experienced long-term changes over our study interval (Ridgwell, 2005). As estimating past DIC change in detail is beyond the scope of this study here we devise a data-driven approach to account, to the first order, for long-term drift in DIC in lieu of simply presuming constant DIC across the study interval. To this end we published benthic foraminiferal $\delta^{11}B$ data (Raitzsch and Hönsch, 2013; Greenop et al., 2017) to generate a deep ocean pH record using equation (7), using the same three scenarios for $\delta^{11}B_{\text{SW}}$ (i.e. L02, RH13, G17) that we also use to generate our surface ocean pH records (Lemarchand et al., 2002; Raitzsch and Hönsch, 2013; Greenop et al., 2017). We then combine deep ocean pH with estimates of deep ocean carbonate ion concentration ([CO$_3^{2-}$]) to solve for deep ocean DIC. Deep ocean [CO$_3^{2-}$] is calculated based on past changes in the CCD (using Sime et al., 2007 and Pälike et al., 2012 as two scenarios) in conjunction with the [$Ca^{2+}$] defined by the two scenarios for seawater major ion changes (i.e., constant modern seawater composition versus trends based on data by Horita et al., 2002 and Brennan et al., 2013). We define deep ocean carbonate ion concentration as:

$$[\text{CO}_3^{2-}]_{\text{deep}} = K_{\text{CaCO}_3}^{*} \cdot \Omega_{\text{calcite}} \cdot [Ca^{2+}]$$

where $K_{\text{CaCO}_3}^{*}$ is calculated using the MyAMI model for the hydrostatic pressure at the reconstructed depth of the CCD and the calcite saturation index $\Omega_{\text{calcite}}$ assumes a nominal value of 1. Hence, the implications of past changes in the CCD for carbonate chemistry are represented via the strong pressure dependence of $K_{\text{CaCO}_3}^{*}$, which increases by ~16% per kilometre water depth. We then combine deep ocean carbonate ion concentration and the benthic foraminiferal-based pH record to yield estimated DIC:

$$\text{DIC} = \frac{[H^+]^2 + [H^+K_1^* + K_1^*K_2^*]}{K_1^*K_2^*} \cdot [\text{CO}_3^{2-}]_{\text{deep}}$$

Our data-driven approach makes the assumption that there is no difference in the carbonate ion concentration between the CCD and the core sites of the deep ocean pH reconstruction. This treatment is a significant simplification and certain to introduce error in the DIC estimate. However, we note that observed modern ocean deep ocean gradients in carbonate chemistry are relatively small when compared for example to the depth dependence of $K_{\text{CaCO}_3}^{*}$, that we include to represent past CCD changes. Furthermore, by assuming a nominal $\Omega_{\text{calcite}}$ of 1 at the pressure of the CCD we effectively introduce systematic bias towards greater estimated DIC because bottom waters at the CCD exhibit moderate undersaturation ($\Omega < 1$) so as to facilitate the near complete dissolution of calcite on the seafloor. With these caveats in mind we note that under all scenarios the DIC estimated from the youngest data is lower than deep ocean DIC in the modern ocean (Supplementary Fig. 4), in particular in the scenarios based on the $\delta^{11}B_{\text{SW}}$ record of Raitzsch and Hönsch (2013). Therefore, we caution that the approach taken here to estimate DIC appears to be biased towards lower absolute values by about 10–20% depending on scenario. However, as is evident from the comparison of the different sensitivity scenarios, the change in estimated DIC over the course of the Neogene is dominantly driven by the counteracting effects of secular [$Ca^{2+}$] decline and the tendency of deep ocean pH to increase, which act to raise and lower estimated DIC, respectively.

While our approach to estimating past DIC has unaccounted bias, the first-order changes in the DIC estimates over time can be attributed to trends in the data-driven input parameters and is reasonable first approach. Therefore, we carry out linear regression on the output of a 10,000 member Monte-Carlo simulation of the deep ocean input data, extracting for each scenario the estimated DIC rate of change and the intercept uncertainty. Based on these regression results, we estimate surface ocean DIC change relative to its known modern concentration for each respective scenario of $\delta^{11}B_{\text{SW}}$, CCD and seawater major ion composition. That is, we set
surface DIC to its modern reference value of 2000 μmol/kg plus the regression change over time plus a nominal uncertainty based on the regression intercept value (Supplementary Fig. S5) (i.e., ± 1σ of 190 to 270 μmol/kg depending on scenario; see Supplementary Table S4 for regression results). We note that in the early Miocene interval (>17 Myr) of the Greenop et al. (2017) Δ11B record large uncertainties exist in the Δ11Brecon value for the oldest samples which cause an edge effect in the smoothing algorithm used in that study (Greenop et al., 2017). Thus, for the regressions using the Greenop et al. (2017) Δ11Brecon scenario we only use benthic Δ11B data <16 Ma to regress DIC and extrapolated further back in time. This is necessary because this possible bias would have been propagated into the regression of DIC change, thus affecting our reconstruction for the entire record.

We note that applying the DIC change regressed from the benthic data to the surface we implicitly assume a constant DIC gradient (ΔDIC). An alternative would be to regress deep alkalinity and assume a constant alkalinity gradient (ΔALK) instead, but we find that these two alternative approaches yield nearly identical results (see Supplementary Fig. S6). We argue that the surface pH that we reconstruct is the dominant driver of reconstructed pCO2 and surface carbonate saturation because it is a strong constraint on the ΔALK-to-ΔDIC difference. Whether ΔDIC or ΔALK are assumed to be constant therefore makes only a minimal difference for our purpose, and the deviation between these two assumptions falls well within the range of our propagated uncertainties.

DIC estimates vary between Δ11Brecon scenarios across the Neogene: both L02 and G17 show a long-term DIC rise (+20 to +37 μmol/kg per million years, depending on scenario, Supplementary Table S5) in contrast to RH13 scenario, which results in essentially constant DIC (+3 to +11 μmol/kg per million years, depending on CCD scenario, Supplementary Table S5) throughout the study period. That is, the scenarios based on the three available Δ11Brecon reconstructions collectively support a modest rate of long-term DIC increase over the study period, which integrated over the duration of the Neogene could amount to a significant expansion of the active carbon pool of the climate system. Along these lines, our assertion of significant Neogene DIC pool expansion agrees with the previously published model results from Tyrrell and Zeebe (2004), Ridgwell (2005), and Caves et al. (2016) (Supplementary Fig. S7).

The approach taken here is similar to Foster et al. (2012) which used a simple model anchored by paleo-pH depth profile and CCD reconstructions to estimate surface alkalinity and calculate pH and pCO2. Using this approach, Foster et al. (2012) estimate surface alkalinity at 1293 ± 200 umol/kg at a single data point (12.72 Ma) and extrapolate across the mid-Miocene. The resultant average mid-Miocene DIC ~1100 ± 180 umol/kg are much lower than estimates derived in this study (Supplementary Fig. S7). Building on this, Greenop et al. (2014) adopted the approach of Foster et al. (2012) but increased the uncertainty to ±300 umol/kg to account for likely rapid CCD changes not resolved in available CCD reconstructions. Based on our new data-driven analysis and existing model based estimates (Ridgwell, 2005; Tyrrell and Zeebe, 2004) we argue that the approach by Foster et al. (2012) yielded DIC that was too low. Consequently, due to the higher DIC estimates applied here, the Mid-Miocene pCO2 estimates calculated from planktic δ13C data of Foster et al. (2012) and Greenop et al. (2014), as well as from our new data from ODP site 761 and 926, are significantly higher than previously published. While highlighting again the caveats in our estimates of surface DIC we argue that this approach is an improvement in deriving long-term DIC change for the boron isotope-based reconstruction of pCO2 and carbonate saturation across the Neogene.

3. Results and discussion

3.1. Δ11B record of T. trilobus from the Neogene

Our δ11B record displays a long-term trend in which low δ11B is associated with high Mg/Ca in the same planktic foraminifera, and with low benthic oxygen isotope ratios (δ18O). All else being equal, this suggests that low pH (and thus high CO2) is associated with high sea surface temperatures and some combination of relatively warm bottom waters and/or low global ice volume (Fig. 2). Reasonable agreement between δ11B from geographically diverse sites suggests the record mainly reflects global carbon cycle variations rather than complications relating to diageneis and variations in the extent of surface water (dis)equilibrium with respect to atmospheric CO2 (Fig. 1). The broad correlations evident in Fig. 2 alone suggest a tight coupling between the carbon cycle (that predominantly drives δ11B change) and climate (that predominantly drives δ18O change) through the Neogene. Our new δ11B data presented in this study shows similar variability to previously published δ11B records from the Miocene (Foster et al., 2012; Badger et al., 2013; Greenop et al., 2014) and Plio-Pleistocene (Hönisch et al., 2009; Seki et al., 2010; Bartoli et al., 2011; Martínez-Botí et al., 2015a; Chalk et al., 2017) and fills in data gaps in the early Miocene and late Miocene (Fig. 2; Supplementary Fig. S1). From here on we compile δ11B estimates from this study alongside previously published δ11B records (with appropriate analytical offsets as required, see above) and calculate carbonate system variables to examine Neogene changes in the carbon cycle.

3.2. Scenarios and sensitivity testing

The calculations carried out in order to reconstruct seawater carbonate chemistry and atmospheric CO2 require independent knowledge of the histories of seawater boron isotope composition (δ11Bsw) and major ion composition (especially [Ca2+] and [Mg2+]) as well as the CCD. For each of these parameters we make at least two different choices and carry out the full set of calculations, including Monte-Carlo uncertainty propagation, for all twelve combinations. In this section we describe all scenarios and explain the agreements/disagreements between them. The output data for all scenarios is provided as a tabulated dataset in the supplemental material (Supplementary Table S6). In the section below, we show that the uncertainties in our reconstructions are dominated by the δ11Bsw reconstructions, not the CCD, and hence only consider these scenarios in our main discussion (see section 3.3).

For our consideration of changes in δ11Bsw, we use the δ11Bsw reconstructions of Lemarchand et al. (2002) (“L02”), Raitzsch and Hönisch (2013) (“RH13”), and Greenop et al. (2017) (“G17”) (Fig. 2). Specifically, RH13 uses benthic δ11B and assumptions about deep ocean pH, G17 uses pH gradients in the ocean derived from planktic and benthic δ11C, whereas L02 is based on geochemical modelling of boron fluxes. Overall, all three records show a general increase through time from the middle Miocene to present, however, the δ11Bsw records diverge in the early Miocene and late Miocene (Greenop et al., 2017; Fig. 2D). The divergence in the early Miocene (>17 Myr) and late Miocene amongst the three δ11Bsw scenarios yields divergent pH and pCO2 reconstructions for these time intervals but has only minor impact on our reconstruction of surface δ13Corg (Fig. 4). The G17 reconstruction has likely too low δ11Bsw before 17 Myr due an edge effect (Greenop et al., 2017) and thereby yields anomalously high pH and low pCO2 for this interval. Conversely, the RH13 reconstruction agrees well with the G17 record from the middle Miocene to present but is likely too high in the early Miocene as these authors assume a linear increase in surface pH through time (Fig. 2D). Further, because Raitzsch and Hönisch (2013) presuppose knowledge of pH in
their approach, caution should be used when interpreting reconstructions of pH and pCO2 that are based on their $\delta^{11}$Bw record (Greenop et al., 2017). Both measurement-based records, RH13 and G17, fall within the uncertainty envelope of L02 modelled reconstruction. However, L02 does not show the same multi-million year variability as RH13 and G17 across the Late Miocene, suggesting that the ocean’s input and output of boron that the Lemarchand et al. (2002) model is based on requires further refinement. Overall, we are encouraged by the good agreement between the three independent $\delta^{11}$Bw records from the Middle Miocene to present, which builds confidence in our reconstructions, but we caution that further refinement is needed in $\delta^{11}$Bw of the early and late Miocene. Carbonate system data for all $\delta^{11}$Bw scenarios are provided in the output table (Supplementary Table S6) and discussed further below (see Section 3.3). As the $\delta^{11}$Bw – G17 > 17 Myr reconstruction is biased by the edge effect we use a dashed line to represent this portion of the pH, CO2 and $\Omega_{\text{arag}}$ reconstructions.

As for the history of CCD change, which factors into our estimation of Neogene DIC change, comparing equivalent scenarios with contrasting CCD reconstruction (either Pâlike et al., 2012 or Sime et al., 2007) we find only small differences in estimated DIC change (Supplementary Fig. S8), with the scenarios based on the Sime et al. (2007) record consistently resulting in slightly greater rates of long-term DIC increase because of the tendency of long-term CCD deepening in that record (see Supplementary Table S4; Supplementary Fig. S5). This difference between CCD scenarios has only very modest implications for ocean carbon chemistry, consistent with earlier findings (Tyrrell and Zebe, 2004; Ridgwell, 2005). Consequently, regardless of which CCD reconstruction is used, we arrive at essentially identical reconstructions of CO2 and $\Omega_{\text{arag}}$ (Supplementary Fig. S8). This is because the long-term trend in CCD over our study interval (i.e., <500 m) implies only a ~8% change in the deep ocean calcium-to-carbonate product, DIC, CO2 and surface saturation state, whereas long-term fractional changes in $[\text{Ca}^{2+}]$, benthic $[\text{H}^+]$ and surface $[\text{H}^+]$ are all much greater. Due to the minor impact the CCD reconstruction has on our results we focus our discussion below on only the scenarios using the CCD reconstruction of Pâlike et al. (2012), referred hereafter simply as ‘Pâlike’, because it incorporates data from an array of Pacific DSDP and ODP sediment cores, has better age control, and is of higher resolution.

For our consideration of seawater major ion composition, we carry out our calculations using $[\text{Ca}^{2+}]$ and $[\text{Mg}^{2+}]$ reconstructions of Horita et al. (2002) and Brennan et al. (2013) as well as the hypothetical scenario that both concentrations remained unchanged at their modern seawater value throughout the entire study interval (Fig. 3). The reconstructed $[\text{Ca}_{\text{bw}}]$ derived from fluid inclusions are consistent with calcium isotope model-data comparisons suggesting that $[\text{Ca}_{\text{bw}}]$ likely decreased through the Neogene (Horita et al., 2002; Heusser et al., 2005; Brennan et al., 2013). Consistent with Hain et al. (2015) we find that the leading order impact of seawater major ion composition operates through $[\text{Ca}^{2+}]$ being a direct factor in the CaCO3 solubility product, not through ion-ion interactions and ion-complex formation affecting the equilibrium constants. As described above, using reconstructed deep ocean pH and CCD (the latter as a constraint on the deep ocean calcium-to-carbonate ion product) the reconstructed ~40% secular decline in seawater $[\text{Ca}^{2+}]$ over the course of the Neogene translates to a reciprocal secular increase in DIC (Equation (10), Supplementary Fig. S4). Conversely, the general tendency of reconstructed deep ocean pH increase imparts a secular decline in estimated DIC because at higher pH less DIC is needed to maintain a given carbonate ion concentration. That is, in the scenarios that include seawater major ion change the opposing effects of $[\text{Ca}^{2+}]$ and benthic pH on estimated DIC partially cancel out, leaving a modest residual long-term DIC rise (Supplementary Fig. S9; Supplementary Table S5; 20 μmol/kg and 27 μmol/kg per million years in the G16 and L02 $\delta^{11}$Bw scenarios, respectively) or effectively no net DIC change in the case of the RH13 $\delta^{11}$Bw scenario (3 μmol/kg per million years). Such secular DIC rise during the Neogene is consistent with previous work.
(Supplementary Fig. S7; Tyrrell and Zeebe, 2004; Ridgwell, 2005; Caves et al., 2016). As a point of comparison, the hypothetical scenario of time-invariant seawater major ion composition eliminates the calcium-driven tendency of secular increase in DIC, yielding instead long-term estimated DIC decline driven by reconstructed long-term benthic pH rise. With regard to our reconstruction of surface carbon chemistry, which is the product of both our extensive surface pH reconstruction and the long-term trends in DIC that we infer from deep ocean data, the opposing trends in DIC change between the “fluid inclusions” and “constant” scenarios impart opposing tendencies upon reconstructed atmospheric CO2, with the more realistic “fluid inclusions” scenario exhibiting a reduced secular CO2 decline (Supplementary Fig. S9). In contrast, our reconstruction of surface carbonate saturation state is virtually insensitive to these scenarios of changing seawater major ion composition (including calcium concentration) (Supplementary Fig. S9). This is largely because our estimate of DIC change is anchored by deep ocean carbonate saturation state via the CCD, which already accounts for seawater major ion composition changes and the strong effect of [Ca2+] on the carbonate saturation product. In the discussion below we focus exclusively on the “fluid inclusions” scenario because the “constant” scenario is intended only as a sensitivity test and to illustrate our approach to the calculations.

3.3. Neogene pH, CO2, and ωarag reconstructions

The calculation of pH and pCO2 requires records of seawater chemistry (e.g. δ18Osw, [Ca]sw) and a second carbonate system parameter (here derived via the CCD) which currently have considerable uncertainties. Based on our sensitivity analysis (Section 3.2), we will discuss three scenarios which use the L02, and RH13, G17 δ18Osw reconstructions along with ‘Pälike’ CCD, and fluid inclusion [Ca2+] and [Mg2+]. Late Miocene pH (∼8.1) are in agreement with earlier model-based estimates (Ridgwell, 2005). The δ18B-derived CO2 record from each scenario are in relatively good agreement with published records from a range of proxies (e.g., alklenones, stomata, paleosols) (Zhang et al., 2014; Foster et al., 2017; Supplementary Fig. S10). However, our record shows more structure, with peaks in CO2 during the MMCO, late Miocene, and late Pliocene (Fig. 4; Supplementary Fig. S10). A particular strength of the δ18B proxy and our approach is that we also calculate specific secular changes in seawater carbonate chemistry.

The most acidic surface conditions (pH∼7.6–7.7) occurred in the MMCO where inter-sample variability likely reflects orbital timescale (∼100-kyr) cycles associated with large (∼200 ppm) CO2 variations reaching >800–1100 ppm (i.e. max CO2 value: 891 ± 165 ppm:L02; 1103 ± 87 ppm:RH13; 774 ± 232 ppm:G17) and an average of ∼470–630 ppm (i.e. 510 ppm:L02; 630 ppm:RH13, 470 ppm:G17) (Fig. 4; Greenop et al., 2014). Our MMCO surface-water pH estimates are similar to the low values reported for the late Eocene (7.6–7.8; Anagnostou et al., 2016), a time of global warmth, high CO2 and low ice volume (Lear et al., 2000). Although the rest of the record is at lower resolution and contains sample aliasing of orbital-scale variability, some long-term trends are clearly discernible. Following the MMCO, CO2 decreased and surface pH increased to ∼8.1 at 12.5 Ma, similar to pre-industrial (Figs. 2, 4). Surface pH was ∼8.0–8.1 between 9 and 3 Myr and subsequently increased towards the late Pleistocene glacial/interglacial range of ∼8.3 to ∼8.1 (Foster, 2008; Hönisch et al., 2009; Henehana et al., 2013; Chalk et al., 2017).

Unlike the oscillations in pH and CO2, our record of low latitude aragonite saturation (ωarag) is broadly constant being in the range 3 to 5 for much of the last 24 million years. In detail however mean low latitude ωarag increased from the MMCO (2.7 ± 0.5:L02; 3.5 ± 0.6:RH13; 3.3 ± 0.6:G17) to relatively stable levels (3.8 ± 0.7:L02; 4.0 ± 0.7:RH13; 3.7 ± 0.7:G17) in the late Miocene with near pre-industrial values (4.1 ± 0.8:L02; 4.3 ± 0.8:RH13; 3.7 ± 0.7:G17) first being reached in the late Pliocene-early Pleistocene (Fig. 4). As with the apparent orbital scale fluctuations in reconstructed MMCO pCO2, the pH fluctuations implied by the Greenop et al. (2014) boron isotope data result in substantial fluctuations in reconstructed saturation state, with the MMCO episodes of low pH corresponding to the lowest saturation levels (1.6–2.1) that we find over the entire Neogene study interval.

It is worth noting that the rather large uncertainty intervals we report for each of a number of scenarios are the result of conservative propagation of uncertainty in the various input parameters. Only a small proportion of that uncertainty is due to the δ18B measurements and the other proxy data required for the boron isotope pH proxy. Instead, structural uncertainty in seawater composition – major ion, δ18Bsw, DIC – are compounded in our approach to reconstruct seawater carbonate chemistry over geologic time-scales much longer than the residence time of each of these constituents. To illustrate that point we consider an alternative scenario for the Plio-Pleistocene (Supplementary Fig. S11), a geologically recent interval where seawater major ion, δ18Bsw, DIC can be anchored at modern values and their structural uncertainty is usefully assessed through the spread in the respective scenarios. For this alternative scenario we propagate the uncertainties due to the spread of δ18Bsw reconstructions and the spread in our scenarios of long-term DIC drift under the ‘fluid inclusion’ major ion scenario, while maintaining the propagation of uncertainties in temperature, salinity and foraminiferal δ18B the same. In that scenario we find uncertainty intervals for pH, CO2 and carbonate saturation states small enough to confidently resolve glacial/interglacial cycles, something not possible with our treatment of the full Neogene dataset (Supplementary Fig. S11; Supplementary Table S7). That is, while our approach to reconstruct seawater carbonate chemistry of the Neogene is able to resolve long-term trends and large magnitude episodes such as the MMCO there remains considerable room for improving the reconstructions by reducing uncertainties related to seawater composition.
3.4. Evaluating Miocene CO$_2$-climate decoupling

Long-term climate trends across the Neogene, as inferred from benthic foraminiferal $\delta^{18}$O$_b$ records, show a transition from an extended period of global warmth of the MMCO to the modern icehouse world marked by the Middle Miocene Climate Transition (MMCT) at 13.9 Ma where Antarctic ice sheets expanded and climate cooled (Fig. 5; Zachos et al., 2001, 2008; Cramer et al., 2009; Lear et al., 2010). Following this global cooling step, the lack of trend in the $\delta^{18}$O$_b$ record suggests that Earth’s climate was relatively stable during the late Miocene (11.6–5.3 Ma) (Zachos et al., 2001, 2008). However, a recent global SST compilation indicates a long-term surface cooling across this interval reaching near modern ocean temperatures between 5 and 7.4 Myrs ago (Herbert et al., 2016). In contrast to the late Miocene, the early Pliocene was a period of prolonged warmth terminated at 2.7 Ma with the initiation of northern hemisphere glaciation and subsequent intensification of glacial interglacial climate variability in the Pleistocene (Lisiecki and Raymo, 2005). Previous CO$_2$ records suggest that the transition from the warm MMCO to the cooler post-MMCT was associated with a decline in CO$_2$ (Kürschner et al., 2008; Foster et al., 2012). The warmth of the early Pliocene has also been previously attributed to higher than pre-industrial atmospheric CO$_2$ levels (Seki et al., 2010; Bartoli et al., 2011; Martínez-Botí et al., 2015a). However, the role of CO$_2$ in evolution of global climate across the Late Miocene is still under debate.

Alkenone-derived CO$_2$ records from Pagani et al. (1999a, 1999b) exhibited low CO$_2$ levels during the early to mid Miocene with no significant shifts in the late Miocene. This led the authors to identify no long-term correlation of climate and CO$_2$ and suggest a late Miocene decoupling between CO$_2$ and climate (Pagani et al., 1999a, 1999b, 2005). However, subsequent work has shown that foraminifera-$\delta^{18}$O based SST estimates used in these late Miocene alkenone-CO$_2$ calculations were too low due to inaccuracies in their estimates of ice volume, calcification depth, and diagenetic overprint (Pagani et al., 2010). Subsequent TEX$_{86}$ derived SST estimates indicate relatively higher temperatures that would suggest higher CO$_2$ levels than previously published, but still supporting a decoupling between climate and CO$_2$ in the late Miocene (Zhang et al., 2014).

Late Miocene climate records suggesting warmer than modern continental and oceanic temperatures and the relatively low alkenone-based CO$_2$ levels are therefore difficult to reconcile with a dominant role of CO$_2$ in setting the evolution of late Miocene climate (Pound et al., 2011; LaRiviere et al., 2012). Specifically, LaRiviere et al. (2012) shows that North Pacific SSTs were significantly higher than today and suggest that in the absence of significantly elevated CO$_2$, a complex mechanism involving a deepened thermocline, expansion of tropical warm pool and increased water vapour may have maintained the late Miocene warmth. Here we use our new Neogene CO$_2$ records to examine the degree of coherency between CO$_2$ and $\delta^{18}$O$_b$ and revisit the idea of climate-CO$_2$ decoupling in the Miocene.

Cross-plots between CO$_2$ climate forcing (i.e., a linear function of the logarithm of CO$_2$, i.e., ln(CO$_2$/278); Myhre et al., 1998) and $\delta^{18}$O$_b$, for all three $\delta^{11}$B$_{sw}$ scenarios, show a correlation across the Mid-Miocene (11.6–16 Ma) and Pliocene (0–5 Ma) (Fig. 6). The good correspondence between CO$_2$ and $\delta^{18}$O$_b$ across these time intervals suggests a dominant role for CO$_2$ in forcing climate. Despite this good coupling between global climate (as recorded by benthic $\delta^{18}$O) and CO$_2$ for the Plio-Pleistocene and Mid Miocene, across the late Miocene (5.3–11.6 Ma) we record CO$_2$ changes that are not strongly matched by corresponding changes in the $\delta^{18}$O$_b$ record (i.e. no variance in $\delta^{18}$O$_b$ that corresponds to the variance in our reconstructed CO$_2$; $R^2 = 0.02–0.09$; Figs. 5; 6). Regardless of the chosen $\delta^{11}$B$_{sw}$ scenario, all our CO$_2$ reconstructions show CO$_2$ peak at ~9 Ma that is not seen in the climate records (Fig. 5). This peak in CO$_2$ at ~9 Ma is also seen in recent cell-sized corrected alkenones and penatate diatoms (diffusive and ACTI-CO models) CO$_2$ records (Bolton et al., 2016; Mejia et al., 2017; Supplementary Fig. S1). However, quantification of the CO$_2$ variations using the diatom method awaits determination of a diatom active carbon uptake model. Nonetheless, the convergence in CO$_2$ trends from 11.6 to 8.5 Ma and the mismatch between CO$_2$ and climate trends support the notion of a late Miocene “climate-CO$_2$ decoupling” (Pagani et al., 1999a, 1999b). This could reflect the significance of oceanographic or palaeogeographic changes and related feedbacks in setting the late Miocene climate, as suggested by others (Pagani et al., 1999a, 1999b; Foster et al., 2010; LaRiviere et al., 2012), which may have masked or counteracted changes in the CO$_2$ forcing at this time.

However, there remain two principle caveats that prevent definite verification of the apparent late Miocene decoupling of CO$_2$
and climate with our current CO₂ record. (1) While we present the highest resolution late Miocene CO₂ reconstruction to date, our record still does not resolve CO₂ change at an orbital timescale and may thus contain aliasing of orbital scale CO₂ variability, and (2) during the late Miocene there is growing evidence for rapid and large magnitude δ¹³Bsw change, evident in the two measurement-based δ¹³Bsw records (RH13 and G17) (Fig. 2). Boron isotope based reconstructions of carbon cycle and CO₂ change during this period critically depends on reconstructions that correctly capture both the pace and timing of this δ¹³Bsw shift. Interestingly, there is a correlation of our planktic δ¹³B to δ¹⁸O_R (R² = 0.22) that is not evident in CO₂ calculated from the boron isotope data (Supplementary Fig. S13). This observation could be used to argue that some intervals of late Miocene carbon cycle change may in fact have changed in tune with climate but this relationship is currently obscured by the uncertainties in the parameters needed to convert δ¹³B to CO₂ (and pH). Therefore, a detailed and robust record of δ¹³Bsw change across the late Miocene is needed to fully elucidate the nature of CO₂-climate coupling at this time.

3.5. Geological context of future ocean acidification

Over the last 250 yrs, anthropogenic emissions have increased atmospheric CO₂ levels by >40% to ~400 ppm (Keeling et al., 2001; Betts et al., 2016; https://www.co2.earth) more than 120 ppm above the range of glacial-interglacial cycles of at least the last 800,000 yrs (Petit et al., 1999; Siegenthaler et al., 2005; Luethi et al., 2008; Bereiter et al., 2015). The oceans have absorbed about a third of the CO₂ emitted since the 1750s (Doney et al., 2009; Le Quere et al., 2016), with the CO₂ fertilizing ocean carbonate ion so as to simultaneously decrease pH and calcium carbonate saturation state. Predictions of future pH and saturation state are widely regarded as sufficient to disrupt physiological processes and reduce calcification, development, and fecundity in a range of marine organisms (Gattuso et al., 2015). Furthermore, reduced pH will impact the availability of essential nutrients (i.e. N) and trace metals (i.e. Fe) critical to marine primary producers for biological functions, altering marine biogeochemical cycles (Hutchins et al., 2007; Hoffmann et al., 2012).

In Fig. 7 we compare our geological record of pH and CO₂ to future predictions based on IPCC Representative Concentration Pathway (RCP) scenarios from the stringent (RCP2.6) to the high-carbon business as usual trajectory (BAU; RCP8.5) which extend to 2100 (Bopp et al., 2013). Note we limit our discussion here to MC-ICPMs based pH, CO₂, and Ωˌarag dataset (Seki et al., 2010; Foster et al., 2012; Badger et al., 2013; Greenop et al., 2014; Martínez-Botí et al., 2015a; Chalk et al., 2017; this study), to maintain internal consistency free of poorly defined analytical offsets (see previous discussion). Because ocean acidification will continue beyond 2100, we also make the comparison with updates up to the year 2500 using emission pathways releasing 250 PgC and 2000 PgC (the near equivalent to RCP2.6 and BAU RCP8.5, respectively), 5000 PgC (a possible upper limit including unconventional fossil fuels such as shale gas), and 10,000 PgC (a possible upper limit on all available carbon resources) (Winkelmann et al., 2015; Bruckner et al., 2014). To compare our low latitude specific Ωˌarag to the global average as presented in Winkelmann et al. (2015) model, we adjust model Ωˌarag by a constant offset (i.e. 0.9) based on the pre-industrial difference between low latitude and global model estimates. Current CO₂ reconstructions of 400 ppm (Keeling et al., 2001; Betts et al., 2016; https://www.co2.earth) are similar to the highest values we reconstruct during the interglacials of the Pliocene and in late Miocene. By 2100, pH levels on the RCP8.5 pathway will be <7.8 and CO₂ near 930 ppm, unprecedented for the last 14 million years (66% CI). If continued unabated (following the 2000 PgC emission pathway) conditions will reach a pH as low as 7.7 by 2150 and CO₂ as high 960 ppm, equivalent to some of the lowest MMCO levels (66% CI). Exhaustion of conventional and unconventional (10,000 PgC) fossil fuel reserves could reduce pH to 7.2 and increase CO₂ to ~4200 ppm by 2300 (Bruckner et al., 2014; Winkelmann et al., 2015), unprecedented in at least 22 Myr (at 66% confidence). By 2100 Ωˌarag levels will be <2.75 in the BAU scenarios, lower than any point in the last 8 million years (66% CI) in two scenarios. Further CO₂ emissions predict Ωˌarag to reach globally undersaturated conditions. The only parallels of such extreme and geologically rapid acidification in the geological past is associated with major mass extinction (Clarkson et al., 2015).

On geologic timescales (>10,000 yrs), the balance between the delivery and removal of alkalinity to the oceans via carbonate weathering and burial acts to partially decouple pH and carbonate saturation state (Fig. 4; Hönsch et al., 2012). Previous studies suggest that surface ocean saturation state was relatively constant across the last 65 Myr excluding short-term events such as Eocene–Oligocene Transition and Paleocene–Eocene Thermal Maximum (Tyrrell and Zeebe, 2004; Ridgwell and Zeebe, 2005; Hönsch et al., 2012; Fálik et al., 2012; Zeebe, 2012; Penman et al., 2014). However, these Cenozoic records of Ω rely on carbon cycle models that make significant assumptions about CO₂ across the time interval. Our record of the ocean carbonate system shows broadly constant values for the Neogene in the range 3–5, but with clear minimum Ωˌarag values, lower than pre-industrial levels, during the warmth of the MMCO when pH values are the lowest in our record (Figs. 2, 4). Pacific % carbonate records (Holbourn et al., 2013) indicate that there were large changes in the CCD on glacial–interglacial timescales during the MMCO indicating a dynamic carbon cycle. The fossil record shows significant global and regional extinction of corals and mollusks in the early and middle Miocene, although arguably not an extraordinary occurrence, while calcifying plankton thrived during the MMCO (Budd, 2000; Warren and Bottjer, 2001; Bown et al., 2004; Ezard et al., 2011). Considering the possible consequences of ongoing acidification for marine organisms, however, one must balance these observations with the fact that geological rates of change were much slower leading into the MMCO than current anthropogenic change, allowing for more time for species to adapt (e.g. scleractinian reefs in the middle Miocene; Perrin, 2002; Wilson, 2008; Hönsch et al., 2012).

The existence of past warm climates, combined with a history of glacial–interglacial variability suggests that, if given time, many calcifiers may be able to adapt to slightly more acidic and less saturated waters than occur at present. However there is no clear analogue for the rate and magnitude of change over the coming centuries under BAU emissions scenarios. End-of-century BAU pH predictions are comparable to the long-term pH regime (pH < 7.8) experienced in natural CO₂ seep sites, suggesting future, acclimatized tropical ecosystems may have a lower species diversity with higher number of acidification tolerant species (Hofmann et al., 2011; Fabricius et al., 2011). Reduced pH will alter the biogeochemical cycling and the ocean inventory on nutrients and trace metals, which can significantly impact marine biota function under future CO₂ scenarios (Hutchins et al., 2007; Hoffmann et al., 2012). Moreover, ocean acidification coincides with warming and other environmental stresses, potentially exacerbating these negative effects (Gattuso et al., 2015). The current pH of the surface ocean is already probably lower than at any time in the last 2 Myr and the rate of acidification is increasing (Hönisch et al., 2009; IPCC, 2014). Some organisms are already negatively impacted (Bednarsek et al., 2012). The predicted changes in pH and saturation state constitute a substantial threat to marine life, especially for the higher emissions scenarios in which the chemistry of the open ocean is forced, with unprecedented rapidity, into a state that is clearly well-be-
Beyond the long-term natural baseline of the last millions to tens of millions of years as established in this study (Fig. 7).

4. Conclusions

Here we present a Neogene record of planktic foraminiferal δ13B from which we estimate surface water pH and Ωarag as well as atmospheric CO2. These calculations also incorporate previous reconstructions of seawater δ13B (δ13Bsw), major ion composition and deep sea carbonate burial (i.e., CCD). Our sensitivity analysis demonstrates that different δ13Bsw reconstructions influence our estimates of both pH and CO2, most significantly during the early and late Miocene when δ13Bsw reconstructions diverge. Additionally, the reconstructed secular decline in seawater calcium concentration implies a secular DIC increase that counteracts a portion of reconstructed pH-rise-driven secular CO2 decline but has only a weak effect on reconstructed surface aragonite saturation state. The choice of CCD record used in the reconstruction of DIC has mostly insignificant effects on our reconstructions. Our pH records suggest that the lowest pH of the past 22 Myr occurred in the Middle Miocene Climatic Optimum, when pH levels were similar to those of the early Cenozoic. This interval of global warmth is associated with average CO2 in all our scenarios, reaching >470–630 ppm. Following the MMCO there is a general long-term trend towards lower CO2 as climate cooled towards the Pleistocene. Superimposed on this trend is an apparent CO2 peak at ~9 Ma, reproduced in all scenarios, that is not matched by corresponding changes in SST and δ18O records. However, to definitely resolve the degree climate–carbon cycle coupling of the late Miocene will require more detailed records of the pace and magnitude of δ13Bsw change during that interval. Finally, we show that it is unlikely that marine ecosystems experienced low latitude surface pH and Ωarag below 7.7 and 1.7 respectively during the past 14 million years (66% CI). Under business as usual scenarios MMCO CO2 and pH levels will be reached by 2100 and Ωarag will be lower than any point in the last 14 million years (66% CI). Understanding the full implication of this conclusion requires long-term laboratory and field studies examining physiological and ecosystem responses to the full range of biogeochemical changes associated with ocean acidification as well as additional observations from the fossil record.

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