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1	Plio-Pleistocene climate sensitivity	from a new high-resolution CO ₂	record
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2 M.A. Martinez-Boti^{1,a}, G.L. Foster^{1,a*}, T. B. Chalk¹, E.J. Rohling^{2,1}, P.F. Sexton³, D.J. Lunt^{4,5}, R.D.

3 Pancost^{5.6}, M.P.S. Badger^{5,6}, D.N. Schmidt^{5,7}

¹Ocean and Earth Science, University of Southampton, National Oceanography Centre Southampton,
 Southampton, SO14 3ZH, UK

- ²Research School of Earth Sciences, The Australian National University, Canberra 2601, Australia
- ³Centre for Earth, Planetary, Space & Astronomical Research, The Open University, Milton Keynes, MK7 6AA,
 8 UK
- ⁴School of Geographical Sciences, University of Bristol, University Road, Bristol, BS8 1SS, UK
- 10 ⁵The Cabot Institute, University of Bristol, UK
- ^bOrganic Geochemistry Unit, School of Chemistry, University of Bristol, BS8 1SS, UK
- ¹² ⁷School of Earth Sciences, University of Bristol, Wills Memorial Building, Bristol, BS8 1RJ, UK
- 13 *corresponding author
- ^a These authors contributed equally to this work.

15 Theory and climate modelling suggest that the sensitivity of Earth's climate to changes in radiative forcing could depend on background climate. However, palaeoclimate data have thus far been 16 17 insufficient to provide a conclusive test of this prediction. Here we present new atmospheric CO_2 18 reconstructions based on multi-site boron-isotope records through the late Pliocene (3.3 to 2.3 19 Myr ago). We find that Earth's climate sensitivity to CO₂-based radiative forcing (Earth System 20 Sensitivity) was half as strong during the warm Pliocene as during the cold late Pleistocene (0.8 to 21 0 Myr ago). We attribute this difference to the radiative impacts of continental ice-volume 22 changes (ice-albedo feedback) during the late Pleistocene, because equilibrium climate sensitivity 23 is identical for the two intervals when we account for such impacts using sea-level reconstructions.

We conclude that, on a global scale, no unexpected climate feedbacks operated during the warm Pliocene, and that predictions of equilibrium climate sensitivity (excluding long-term ice-albedo feedbacks) for our Pliocene-like future (with CO₂ levels up to maximum Pliocene levels of 450 ppm) are well described by the currently accepted range of 1.5 to 4.5 K per CO₂ doubling.

Since the start of the industrial revolution, the concentration of atmospheric CO₂ (and other 28 29 greenhouse gases; GHGs) has increased dramatically (from ~280 to ~400 ppm)¹. It has been known 30 for over 100 years that changes in GHG concentration will cause the surface temperature of the Earth to vary². A wide range of observations reveals that the sensitivity of Earth's surface 31 32 temperature to radiative forcing amounts to ~ 3 K warming per doubling of atmospheric CO₂ 33 concentration (with a 66% confidence range of 1.5 to 4.5 K; e.g. ref. 1,3), due to direct radiative 34 forcing by CO₂ plus the action of a number of fast-acting positive feedback mechanisms, mainly 35 related to atmospheric water vapour content and sea-ice and cloud albedo. Uncertainty in the 36 magnitude of these feedbacks confounds our ability to determine the exact equilibrium climate 37 sensitivity (ECS; the equilibrium global temperature change for a doubling of CO_2 on timescales of 38 about a century, when all 'fast' feedbacks have had time to operate; see ref. 3 for more detail). Although the likely range of values for ECS is 1.5 to 4.5 K per CO₂ doubling, there is a small but finite 39 possibility that climate sensitivity may exceed 5 K (e.g. ref. 1). Understanding the likely value of ECS 40 clearly has important implications for the magnitude, eventual impact and potential mitigation of 41 42 future climate change.

Any long-range forecast of global temperature (i.e. beyond the next 100 years) must also consider the possibility that ECS could depend on the background state of the climate^{4,5}. That is, in a warmer world, some feedbacks that determine ECS could become more efficient and/or new feedbacks could become active to give additional warmth for a given change in radiative forcing (such as those relating to methane cycling⁶, atmospheric water vapour concentrations^{5,7,8}, in addition to changes in the relative opacity of CO₂ to long wave radiation^{5,9}). One approach to identify whether ECS depends on climate background state is to reconstruct ECS during periods in the geological past when Earthwas warmer than today.

51 The Pliocene (2.6 to 5.3 Myr ago) is one such time, with the warmest intervals between 3.0 and 3.3 Myr ago ~3 K globally warmer than pre-industrial times^{10,11}, while mean sea level stood 12-32 m 52 above the present level^{12,13}. Although most of this warmth is commonly ascribed to increased 53 atmospheric CO₂ levels¹⁴, it has been suggested that simple comparisons of the observed 54 temperature change in the geological record with the climate forcing from CO₂ alone are unable to 55 constrain ECS¹⁰. Instead, a parameter termed Earth System Sensitivity (ESS) is defined – the change 56 57 in global temperature for a doubling of CO₂ once both fast and slow feedbacks have acted and the 58 whole Earth system has reached equilibrium (in contrast, ECS excludes the slow feedbacks; for a 59 discussion of fast versus slow feedbacks, see ref. 3). The most important slow feedbacks are those related to ice-albedo and vegetation-albedo changes. Because of these slow feedbacks, Pliocene ESS 60 is thought to have been ~50 % higher than ECS^{10,15}, with some existing geological data suggesting a 61 Pliocene ESS range of 7-10 K per CO₂ doubling¹⁶, which greatly exceeds a modern ESS estimate of ~4 62 K per CO₂ doubling¹⁰. If ECS was similarly enhanced, then that would imply that either extra positive 63 fast feedbacks operated, or that existing positive fast feedbacks were more efficient, thus increasing 64 the temperature response for a given level of CO₂ forcing. 65

Understanding past climate sensitivity critically depends on the accuracy of the CO₂ data used. 66 Despite a tendency toward increased agreement between different CO₂ proxies¹⁷, individual pCO₂ 67 estimates for the Pliocene still range from ~190 to ~440 µatm (Fig 1a,b) and there is little coherence 68 69 in the trends described by the various techniques (Fig 1a,b). This hinders any effort to accurately 70 constrain Pliocene ECS or ESS. To better determine Pliocene CO₂ levels, we generated a new record, based on the boron isotopic composition ($\delta^{11}B$) of the surface mixed-layer dwelling planktic 71 foraminiferal species Globigerinoides ruber from ODP Site 999 (Caribbean Sea, 12°44.64' N, 72 73 78°44.36' W, 2838 m water depth; Extended Data Figure 1) at more than 3× higher temporal

resolution (1 sample every ~13 kyr; Fig. 1c) than previous δ^{11} B records (1 sample every 50 kyr; Fig. 74 1b). The δ^{11} B of *G. ruber* is a well-constrained function of pH¹⁸ and seawater pH is well correlated 75 with $[CO_2]_{aq}$, as both are a function of the ratio of alkalinity to total dissolved carbon in seawater. In 76 the absence of significant changes in surface hydrography, [CO2]aq is largely a function of 77 atmospheric CO₂ levels and δ^{11} B-derived CO₂ has been demonstrated to be an accurate recorder of 78 atmospheric CO₂ (Extended Data Figure 2a)¹⁸⁻²⁰. Today, the surface water at Site 999 is close to 79 equilibrium with the atmosphere with respect to CO_2 (expressed here as $\Delta pCO_2 = pCO_2^{sw} - pCO_2^{atm} =$ 80 +21 µatm; Extended Data Figure 1)^{18,21} and has remained so for at least the last 130 kyr (Extended 81 Data Figure 2)¹⁸. ODP Site 999 also benefits from a detailed astronomically calibrated age model²² 82 and high abundance of well-preserved planktic foraminifera throughout the past 4 million years^{23,24}. 83 During our study interval it is also unlikely to have been influenced by long-term oceanographic 84 changes such as the emergence of the Panama Isthmus ~3.5 Myr ago (see detailed discussion in ref. 85 86 23). To increase confidence that atmospheric CO_2 changes are driving our pH (and hence our p CO_2^{sw}) 87 record for ODP Site 999 and that the air:sea CO₂ disequilibrium remained similar to modern values, we also present lower-resolution δ^{11} B data from *G. ruber* from ODP Site 662 (equatorial Atlantic, Fig. 88 89 1c; 1°23.41'S, 11°44.35°W, 3821 m water depth; Extended Data Figure 1), where current mean annual ΔpCO_2 is +29 µatm with a seasonal maximum of +41 µatm²¹. Analytical methodology and 90 91 information detailing precisely how pCO2^{sw} is calculated, with full propagation of uncertainties, can be found in the Methods section (with full δ^{11} B and pCO₂ in Supplementary Information Tables 1&2). 92

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94 A new record of Pliocene pCO₂ change

Where our data for both sites overlap in time, reconstructed pCO₂^{atm} values between 2.3 and 3.3 Myr ago agree within uncertainty (Fig 1d; Extended Data Figure 3), and are consistent with most independent records (see Fig 1a,b; Extended Data Figure 2b,c), confirming that the variations we observe are predominantly driven by changes in atmospheric CO₂ concentrations. However, the 99 enhanced resolution of our $\delta^{11}B$ -pCO₂^{atm} record (Fig. 1d) also reveals a hitherto 100 undocumented^{16,23,25,26} level of structure in the CO₂ variability during the 1 million year period 101 investigated, including a transition centred on 2.8 Ma, spanning ~200 kyr, where average pCO₂^{atm} 102 undergoes a decrease of ~65 µatm (Fig 1d).

103 Detailed atmospheric CO₂ measurements from ice cores show orbital-scale (~100 kyr) oscillations in pCO_2^{atm} with a peak-to-trough variation of ~80-100 µatm through the late Pleistocene (90 % of the 104 pCO₂ values lie between +36 and -41 μ atm of the long-term mean; Extended Data Figures 2, 4)²⁷⁻²⁹. 105 106 Once the long-term trend is removed from our Plio-Pleistocene data (thick blue line in Fig 1d), and 107 we have taken into account our larger analytical uncertainty (see Methods), we observe orbital-scale variations in our δ^{11} B-pCO₂^{atm} record of only slightly smaller amplitude than the ice-core pCO₂^{atm} 108 record (0-0.8 Myrs) and for the last 2 Myrs in other δ^{11} B-based records^{19,20,30} (Extended Data Figure 4 109 and Methods), which is in clear contrast with the benthic δ^{18} O which shows increasing variability 110 111 over the last 3 Myrs (Fig 1e and Extended Data Figure 4).

112 Given the different amplitudes of climate variability, the observed similarity between Pliocene and late Pleistocene pCO₂^{atm} variability seems counter-intuitive given the notion that CO₂ is a key factor 113 in amplifying glacial-interglacial climate change^{27-29,31,32}. This is illustrated by a well-defined non-114 linear relationship in a cross plot between deep-sea benthic δ^{18} O and $\ln(CO_2/C_0)$ (where C₀ = pre-115 industrial $CO_2 = 278 \mu atm$), which accounts for the logarithmic nature of the climate forcing by CO_2 116 (Fig. 2b). Note also the clear overlap between Pleistocene (0-2.2 Myrs) ice-core CO₂ measurements 117 and δ^{11} B-based CO₂ reconstructions in this plot (Fig. 2b; Extended Data Figure 2). A similar 118 relationship is also evident in raw δ^{11} B-space (Fig. 2a). Below an inflection at about 275±15 µatm 119 pCO_2^{atm} (equating to $ln(CO_2/C_0) \approx 0$) benthic $\delta^{18}O$ shows a steeper relationship with CO_2 -based 120 121 forcing than it does above this value (Fig. 2). This likely reflects some combination of: (i) growth of larger northern hemisphere ice sheets at pCO₂^{atm} below 275±15 µatm³³ increasing radiative ice-122 123 albedo feedback and amplifying climate forcing by CO_2 change; (ii) an increase in oxygen isotope

124 fractionation in precipitation with increasing size of the ice sheets, which leads to a proportionally 125 greater ¹⁸O enrichment in seawater³⁴; and (iii) potentially stronger deep-sea cooling at low pCO_2^{atm} 126 due to the high-latitude-focussed influences of the ice-albedo feedback process. These findings 127 highlight the profound impacts of northern hemisphere ice-sheet growth on climate variability in the 128 Pleistocene ^{31,32}, relative to the Pliocene (Fig. 2b).

129 Our new data show that the ~275±15 µatm threshold was first crossed at ~2.8 Ma during Marine Isotope Stage (MIS) G10 (Fig. 1d, horizontal dashed line), and - more persistently - during 130 131 subsequent Marine Isotope Stages G6 (2.72 Myr ago), G2 (2.65 Myr ago), and 100 (2.52 Myr ago), when values as low as 233^{+63}_{-53} µatm (95% confidence) were reached and when intervening 132 interglacial values also seem to have been suppressed (Fig. 1c,d). These isotope stages are significant 133 in that they are associated with an increase in the amplitude of glacial-interglacial sea-level 134 oscillations (Extended Data Figure 5b)^{12,13,35} and coincide with the timing of the first substantial 135 136 continental glaciations of Europe, North America and the Canadian Cordillera, as reconstructed by Ice-rafted debris and observations of relic continental glacial deposits³⁶⁻³⁸. Hence, our new high-137 resolution pCO2^{atm} record robustly confirms previous hypotheses^{16,23,25,39} (based on low-resolution 138 CO₂ data) that the first substantial stages of glaciation on the northern hemisphere, as well as a 139 recently recognised deep-sea cooling during the late Pliocene/early Pleistocene¹³, coincided with a 140 significant decline in mean atmospheric pCO2^{atm} at 2.7-2.9 Ma of ~40-90 µatm (mean_{3.0-3.2Ma} -141 142 mean_{2.4-2.7Ma} = $66 \pm 26 \mu atm; p < 0.001$ (two-tailed), n=40).

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144 Efficiency of climate feedbacks

The high fidelity of the boron isotope pH/pCO_2^{atm} proxy (Extended Data Figure 2), coupled with the high resolution of our new pCO_2^{atm} record, offers an opportunity to examine the sensitivity of Earth's climate system to forcing by CO_2 during a period when Earth's climate was, on average, warmer than today⁴⁰. For this exercise, global temperature estimates are also needed. We consider two approaches for this. The first is an estimate of global mean annual surface air temperature change (Δ MAT) over the last 3.5 million years, from a scaling of the northern hemisphere climate required to drive an ice-sheet model to produce deep ocean temperature and ice-volume changes consistent with benthic δ^{18} O data (Fig 3a,b)³⁵. This approach produces a continuous record of global temperature that agrees well with independent constraints for discrete time intervals (see ref. 35).

154 We supplement Δ MAT with a record from a second approach, which is independent from benthic δ^{18} O values. For this, we generated a sea surface temperature stack (SSTst) from 0 to 3.5 Myr ago 155 (Fig. 3c,d), comprising 10 high-resolution (average ~4 kyr) SST records based on $U_{37}^{k'}$ alkenone 156 unsaturation ratios, from latitudes between 41 °S and 57 °N. The selected sites (see Extended Data 157 158 Figure 1b) all offer near-continuous temporal coverage of the last 3.5 Myr (see Methods). Our SSTst record agrees well with independent, higher density compilations of global SST change^{32,41} (Fig 3c 159 160 blue line), indicating that SSTst offers a reliable approximation of global SST change (see Methods for 161 more details). Moreover, our SSTst allows us to directly compare the major SST changes, within the 162 same archives, between the Plio-Pleistocene and late Pleistocene.

163 When comparing temperature records from the two approaches considered, it must be emphasized 164 that Δ MAT reflects global mean annual surface air temperature change, while SSTst approximates 165 global mean sea surface temperature change. Hence, their amplitudes of variability will be different, 166 mainly because SSTst does not include temperature changes over land. Approximately, Δ SST = Δ MAT 167 * 0.66 (ref. 32,42), but direct conversion is not needed here, as we merely aim to contrast Pliocene 168 climate behaviour with that for the Pleistocene, within the same data types.

To determine the sensitivity of global SST and Δ MAT to CO₂ forcing in the Pliocene and Pleistocene, we use time series of forcing calculated from our new and existing CO₂ records (Fig 3e to h), and regress these against both Δ MAT and SSTst (Fig 3a to d; Supplementary Information Tables 1-3). The regression slopes then describe the average temperature change (Δ T in K) per W per m² of forcing 173 (ΔF) for each time interval. These gradients therefore approximate the commonly used sensitivity parameter (S = $\Delta T / \Delta F K W^{-1} m^{2}$) for describing global temperature change for a given forcing³. In this 174 scheme, a doubling of atmospheric CO_2 is equivalent to a forcing of 3.7 W m⁻², so that for the 66% 175 confidence interval of modern climate sensitivity quoted by ref. 1, the present-day equilibrium value 176 of S (S^{*a*}, after ref. 3) is 1.5/3.7 to 4.5/3.7 = 0.4 to 1.2 K $W^{-1} m^2$. However, using palaeoclimate data it 177 is not possible to determine the direct equivalent of S^{a} ; instead, such studies constrain a 'past' 178 parameter (S^{ρ}), which includes the combined action of both fast and slow feedbacks³. Note that 179 Earth System Sensitivity, ESS (in K) = $S^{p} \times 3.7$. Explicit accounting for slow feedback processes in 180 determinations of S^{p} can make it approximate S^{a} (ref. 3). Following ref. 3, an S^{p} estimate after 181 accounting for carbon-cycle feedback is indicated by S_{CO2}, and one accounting for both carbon-cycle 182 183 and land-ice-albedo feedbacks is $S_{CO2,LI}$, where the latter gives a useful approximation of S^{a} . We follow this approach, using $S^p = \Delta MAT/\Delta F$ and $S^{p,SST} = \Delta SST/\Delta F$, both in K W⁻¹ m². Note that our 184 determinations of the sensitivity parameter are based on our entire reconstructed time series, 185 186 rather than on a simple comparison between a limited Pliocene average and the modern average, as was done in previous studies^{3,16}. Since we calculate a S^{p} (and $S^{p,SST}$) for the Pliocene and compare this 187 to the late Pleistocene S^p (and S^{p,SST}), we also avoid complications due to independent changes in 188 boundary conditions (such as topographic changes)³⁹ because we assess sensitivity within each 189 relatively short time window (2.3 to 3.3 Ma vs. 0 to 0.8 Ma). In addition, our approach emphasizes 190 relative changes in CO₂ levels and temperature over the intervals considered, rather than absolute 191 192 values. This improves accuracy because relative changes are much better constrained than absolute temperature and pCO_2^{atm} values from proxy data (see Methods for further discussion). 193

194 Preliminary regression of Δ MAT against Pliocene pCO₂^{atm} identified one data point (at 2362 kyr; 195 white circle in Fig. 1d & 2) with a particularly large residual and significant leverage on the least 196 squares regression (a high Cook's distance). With interglacial-like pCO₂^{atm} values but glacial-like δ^{18} O 197 values (Fig 2), this point may reflect a chronological error, or a short period of unusually high air:sea 198 disequilibrium with respect to CO₂ at ODP Site 999. To avoid the influence of this one point on subsequent linear regressions, we have removed it from our $\delta^{11}B$ -pCO₂^{atm} record. The remaining 199 pCO₂^{atm} data (73 points) were interpolated to a constant resolution (1 kyr), smoothed with a 20 kyr 200 201 moving average to reduce short-term noise and resampled back to the original data spacing (~1 202 sample every 13 kyr). A Monte Carlo approach was followed to determine uncertainties for this smoothed record given the uncertainty in the δ^{11} B-derived pCO₂^{atm}. Radiative forcing changes due to 203 pCO₂^{atm} changes are calculated using $\Delta F_{CO2} = 5.35* ln(CO_2/C_0)$ W m⁻²; where C₀ = 278 µatm (Fig 3)⁴³. 204 We ignore mean annual forcing by orbital variations because it is small (<0.5 W m^{-2} with a periodicity 205 of 100 to 400 kyr) $^{^{31,32}}$ and averages out over the length of our records. Linear regressions of ΔMAT 206 and SSTst versus ΔF_{CO2} were performed using an approach that yields a probabilistic estimate of 207 208 slope, and hence sensitivity to CO_2 forcing ($S_{CO2} = \Delta T / \Delta F_{CO2,LI} = \Delta T / \Delta F_{CO2,LI}$), which fully accounts for uncertainties in both X and Y variables (see Methods; Fig 4). Fig 5a-d displays 209 210 probability distribution functions (pdfs) of the determinations of slope for each time interval. This 211 analysis reveals that, irrespective of the global temperature record used (Δ MAT or SSTst), the average global sensitivity of Earth's climate to forcing by CO₂ only (S_{CO2}) is approximately 2x higher 212 for the Pleistocene than it is for the Pliocene (Fig 4&5). This validates previous inferences of a strong 213 additional feedback factor during the Pleistocene (at pCO_2^{atm} levels below ~280 µatm), which likely 214 arises from the growth and retreat of large northern hemisphere ice sheets and their role in 215 changing global albedo^{31,32}. 216

Given that, to a first order, the Earth system responds to radiative forcing in a consistent fashion, largely independent of the nature of that forcing⁸, we can determine the climate forcing arising from continental ice albedo changes via a relatively simple parameterisation of sea-level change ($\Delta F_{LI} =$ sea-level change (m) × 0.0308 W m⁻²; following ref. 31,32). Several reconstructions of sea-level change partially or completely span the last 3.5 Ma (e.g., ref. 13, 35, 44, 45, and 46 recalculated by 12), and we explore the implications of each of these independent records. Cross-plots of combined 223 CO₂ and ice-albedo forcing ($\Delta F_{CO2} + \Delta F_{LI} = \Delta F_{CO2,LI}$) versus ΔMAT and ΔSST st are shown in Fig 4 for the Pliocene and Pleistocene. Fig 5e-h displays the influence of choices of temperature and sea-level 224 225 record on our determinations of $S_{CO2,LI}$ (= $\Delta T/\Delta F_{CO2,LI}$). In contrast to S_{CO2} , $S_{CO2,LI}$ is similar for both the 226 Pliocene and Pleistocene, regardless of temperature record or other parameter choices (Fig 5). This 227 robustly indicates that the apparent difference between Pliocene and Pleistocene climate sensitivity 228 arises almost entirely from ice-albedo feedback influences. It also implies that all of the other 229 feedbacks that amplify climate forcing by CO₂ (e.g. sea-ice and cloud albedo, water vapour, vegetation, aerosols, other GHGs) must have operated with rather similar efficiencies during both 230 the Pliocene and Pleistocene. Thus, we find no evidence that additional (unexpected) positive 231 feedbacks had become active to amplify Earth system sensitivity to CO2 forcing during the warm 232 233 Pliocene. Alternatively, if additional positive feedbacks did become active (e.g. increase in steady-234 state methane concentration or changes in cloud properties), then their effect must have been negated by the loss of other amplifying feedbacks (e.g. Arctic sea-ice) or the addition of more 235 236 negative feedbacks. This finding is at odds with previous studies (e.g. ref. 16,47) most likely because 237 of differences in our approach to determine Pliocene climate sensitivity (i.e. we determine a within-238 Pliocene sensitivity) and shortcomings in the proxy systems used by the earlier investigations, both 239 in terms of CO₂ and temperatures (e.g. see ref. 48). For instance, Fig 1d (and Extended Data Figure 2) indicate that both orbital-scale variability in pCO_2^{atm} and the major decline at 2.7-2.9 Ma are 240 absent from the previously used¹⁶ alkenone-based pCO₂^{atm} records and as a result regressions of 241 242 temperature and alkenone-derived forcing are poorly defined (Extended Data Figure 2d-f).

243

244 Constraints on Climate Sensitivity

Using the geological record to directly estimate ECS (and thus S^{*a*}) is problematic because information on the appropriate magnitude of a number of key feedbacks (such as vegetation-albedo) is typically unavailable³. Nonetheless, considerable effort has determined that ECS estimates based on the last glacial maximum fall within the range of ECS estimates from other approaches (1.5 to 4.5 K per CO₂ doubling, or 0.4 to 1.2 K W⁻¹ m²; ref. 1). Our analysis implies that a similar ECS applies to the Pliocene and early Pleistocene (2.3 to 3.3 Ma; Fig 5; Supplementary Information Table 4). In addition, our estimate of Pliocene S_{CO2} using Δ MAT lies within a range of 0.6 to 1.5 K W⁻¹ m² (at 95% confidence), meaning that, once all feedbacks have played out for future CO₂ doubling, ESS (= S_{CO2} x 3.7) will very likely (95% confidence) be <5.2 K and will likely (68% confidence) fall within a range of 3.0 to 4.4 K (Supplementary Information Table 4).

255 In May 2013, atmospheric CO_2 levels crossed the 400 ppm threshold to values last seen during the 256 Pliocene (Fig. 1c). Given current CO₂ emission rates, global temperatures may reach those typical of 257 the warm periods of the Pliocene by 2050¹. Our findings suggest that, if the Earth system behaves in 258 a similar fashion to how it did during the Pliocene as it continues to warm in the coming years, an ECS of 1.5 to 4.5 K per CO₂ doubling¹ likely provides a reliable description of the Earth's temperature 259 260 response to climate forcing, at least for global temperature rise up to 3 K above the pre-industrial 261 level. Studies of even warmer intervals in the deeper geological past (well before 3.3 Myr ago) are needed to determine whether any additional climate feedbacks should be expected as the Earth 262 warms even further into the 22^{nd} Century if CO₂ emissions continue unabated. 263

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383 **Supplementary Information** is linked to the online version of the paper at www.nature.com/nature

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400 online version of the paper. Correspondence and requests for materials should be addressed to GLF401 (gavin.foster@noc.soton.ac.uk).

402

403 Figure Legends

Figure 1. Records of late Pliocene/early Pleistocene pCO₂^{atm}. (a) pCO₂^{atm} based on δ^{13} C of 404 sedimentary alkenones (dark green circles (ODP 999)²⁵; aquamarine squares (ODP 999)²⁶; dark 405 orange (ODP 1208)¹⁶, purple circles (ODP 806)¹⁶; dark red squares (ODP 925)⁴⁹). Error bars are 406 uncertainty in pCO₂^{atm} at the 95% level of confidence. (b) δ^{11} B of planktic foraminifera from ODP 999 407 (blue closed circles for *G. sacculifer* and squares²⁵ for *G. ruber*²⁵; red squares for *G. sacculifer*²³) and 408 stomatal density of fossil leaves (purple filled circle)⁵⁰. Error bars are uncertainty in pCO₂^{atm} at the 409 410 95% level of confidence. (c) New boron isotope data from ODP 999 (blue circles) and ODP 662 (red circles). Error bands for ODP 999 denote 1sd (dark blue) and 2sd (light blue) analytical uncertainty, 411 error bars for ODP 662 show 2sd analytical uncertainty. (d) Atmospheric pCO₂ (µatm) determined 412 413 from data shown in (c) for ODP 999 (blue circles) and ODP 662 (red circles). Error band encompasses 68% (dark blue) and 95% (light blue) of 10,000 Monte Carlo simulations of pCO₂^{atm} using the data in 414 415 (c) and a full propagation of all the key uncertainties (see Methods). For ODP 662 error bars encompass 95% of 10,000 simulations. Dotted lines show the modelled threshold of northern 416 hemisphere glaciation (280 μ atm)³³. (e) Benthic δ^{18} O stack²², prominent marine isotope stages are 417 418 labelled (blue for glacial, red for interglacial stages). Thick lines on several panels are non-parametric smoothers through the data. Blue open circle on (d) highlights the data point that is identified as 419 420 outlier in Fig 2 and not used in subsequent regressions.

421 Figure 2. Relationship between δ^{11} B, climate forcing from CO₂ and δ^{18} O. (a) δ^{11} B vs. δ^{18} O and (b) 422 In(CO₂/C_o) vs. δ^{18} O for data from the last 3 million years. Ln(CO₂/C_o) is defined in the text. Boron data 423 in (a) are from this (blue open and closed circles) and published studies (green circles³⁰; blue 424 triangles²⁰). Ice-core CO₂ data shown as open red circles²⁷⁻²⁹. The vertical dashed line is at a CO₂ of 425 278 µatm. The data point removed from subsequent regression analysis is highlighted as open blue 426 circles. Note that the δ^{11} B-pCO₂ data from ref. 23 are not plotted for clarity. The black line is a non-427 parametric regression through all the data shown. The δ^{11} B data from ref. 30 have been corrected 428 for laboratory and inter-species differences through a comparison between core-top δ^{11} B values.

Figure 3. Pleistocene and late Pliocene time series. (a) and (b) mean annual surface air temperature 429 change (Δ MAT)³⁵, (c) and (d) sea surface temperature change (Δ SST; this study in red and from a 430 stack of a more comprehensive compilation³² in blue). Uncertainty envelopes at 95% confidence for 431 both temperature records are shown in red. (e) ΔF_{CO2} for the Pleistocene from ice-core data²⁷⁻²⁹. 432 (f) ΔF_{CO2} for the late Pliocene calculated using the CO₂ data from this study. (g) $\Delta F_{CO2,LI}$ calculated 433 using data in (e) and published sea-level records (R14¹³, VDW11³⁵ and from ref. 44 for 0-520 kyr and 434 ref. 45 for 520 to 800 kyr, R09+E12). (h) $\Delta F_{co2,U}$ for the late Pliocene calculated using the CO₂ data 435 436 from this study and published sea-level records (ref. 46 recalculated by ref. 12, N09, R14¹³, VDW11³⁵). Error bands in (e) to (h) represent the uncertainty in smoothed CO₂ record and sea-level 437 (68% and 95% confidence in light and dark respectively) propagated using a Monte Carlo approach 438 (n=1000) for each reconstruction. 439

Figure 4. Cross plots of forcing and temperature response. (a) ΔMAT vs. ΔF_{co2} and (b) to (d) ΔMAT 440 vs. $\Delta F_{CO2,LI}$ for the following sea-level records detailed in the caption for Figure 3: (b) R09+E12^{44,45} and 441 N09^{12,46} (c) VDW11³⁵, (d) R14¹³. (e) Δ SST vs. Δ F_{CO2} and (f) to (h) Δ SST vs. Δ F_{CO2,LI} for the same sea-442 443 level records as in panels (b) to (d). In all panels late Pleistocene data (0-800 kyr) are shown as red 444 open circles and late Plio-Pleistocene (2300-3300 kyr) as blue open circles. Regression lines fitted by 445 least-squares regression are also shown in the appropriate colour (shaded bands represent 95% 446 confidence intervals). For (a) to (d) the temperature record is that of ref. 35 and for (e) to (h) it is SSTst from this study. In all cases the slope (m) and standard error uncertainty are determined by 447 448 least squares regression. Also shown are the *p* values for the regressions.

Figure 5. Probability density functions of the slope from regressions of temperature against climate forcing. (a,c,e,g) Δ MAT and (b,d,f,h) Δ SST against Δ F_{CO2} and Δ F_{CO2,LI} for the Pleistocene (a, b, e, f) and Pliocene (c, d, g, h), taking into account the uncertainties on all variables (see text). In (e) to (h) individual pdfs are shown for different choices of sea-level, the combined pdf shown in bold is the sum of these different pdfs and therefore also incorporates uncertainty related to the choice of sea-level record. Also shown and labelled are the median (bold), 68th percentile (dot-dash) and 95th percentiles (dotted).

456

457 Methods

458 Sample locations. We present new data from two deep ocean sites: ODP Site 999 (Caribbean Sea, 459 12°44.64'N and 78°44.36'W) and ODP Site 662 (Equatorial Atlantic, 1°23.41'S, 11°44.35'W). Both sites have well-constrained age models for the Pliocene and are part of the Lisiecki and Raymo 460 benthic foraminifera δ^{18} O stack²² (hereafter LR04). Sedimentation rates are comparable between the 461 462 sites (~3 cm/kyr at ODP 999 and ~4 cm/kyr at ODP 662). At ODP Site 999, seventy four samples were analysed at an average temporal resolution of around 1 sample every 13 kyr, targeting several glacial 463 464 and interglacial maxima. ODP Site 662 was analysed at much lower resolution (8 samples in 1000 kyr 465 = 1 sample every 125 kyr on average), and the chosen samples were limited to peak interglacial conditions to avoid potential upwelling influences during glacial periods⁵¹. The extent of the modern 466 air-sea CO₂ disequilibrium at each location is displayed in Extended Data Fig 1a. 467

Analytical methodology. Between 90 and 200 individuals of *Globigerinoides ruber* (~10 μg/shell)
were picked from the 300-355 μm size fraction from ODP Sites 999 and 662. Foraminiferal samples
were crushed between cleaned glass microscope slides and subsequently cleaned according to
established oxidative cleaning methods⁵²⁻⁵⁴. After cleaning, samples were dissolved in ~0.15 M
Teflon-distilled HNO₃, centrifuged and transferred to 5 ml Teflon vials for storage. An aliquot (~20 μl;

~7% of the total sample) was taken for trace element analysis. Boron was separated from the
dissolved samples using Amberlite IRA-743 boron-specific anion exchange resin following established
procedures²⁰. Boron isotope ratios were measured on a Thermo Scientific Neptune multicollector
inductively coupled plasma mass spectrometer (MC-ICPMS) at the University of Southampton
according to methods described elsewhere^{18,20,54}.

478 External reproducibility of δ^{11} B analyses is calculated following the approach of ref. 54, and is 479 described by the relationship:

480
$$2\sigma = 1.87 \cdot \exp^{-20.6 \left[{^{11}B} \right]} + 0.22 \cdot \exp^{-0.43 \left[{^{11}B} \right]}$$
 [1]

481 where [¹¹B] is the intensity of the ¹¹B signal in volts (see ref. 18 for further details).

Trace elements were measured on a Thermo Scientific Element 2 single collector ICPMS at the 482 University of Southampton, following established methods²⁰. Over the period 2012-2013, analytical 483 reproducibility for Mg/Ca was \pm 2.7% (2 σ). Raw Mg/Ca ratios were corrected for changes in the 484 Mg/Ca ratio of seawater (Mg/Ca_{sw}) using the approach of ref. 55 using the power-law modification 485 of ref. 56 and the modelled Mg/Ca_{sw} of ref. 57. Specifically, we use a H value⁵⁶ of 0.41, originally 486 derived for *Globigerinoides sacculifer*⁵⁸, as no species-specific H value is currently available for *G*. 487 *ruber* (for extended discussion, see ref. 48). The following equation^{56,59} was therefore used to derive 488 489 calcification temperatures from our Mg/Ca ratios, which also includes a depth correction to account 490 for the influence of dissolution on shell Mg/Ca ratios.

491
$$T({}^{o}C) = \frac{\ln\left(\left(\frac{Mg}{Ca_{test}}\right)/\left(E \times \left(\left(\frac{Mg}{Ca}\right)_{sw}^{t}\right)^{H}\right)\right)}{0.09} + (0.61 \times Z)$$
[2]

492 Where $\left(\frac{Mg}{Ca}\right)_{sw}^{t}$ is the Mg/Ca ratio of seawater at the time of interest, Z is the core depth in km and E 493 is defined by the following equation⁵⁶:

$$E = \frac{0.38}{\left(\left(\frac{Mg}{Ca}\right)_{sw}^{t}\right)^{H}}$$

Trace element data were also used to check the efficiency of the foraminiferal cleaning
 procedure^{20,54}. All samples had Al/Ca ratios of <100 μmol/mol, and typically <60 μmol/mol.

496 **Determination of pH from** δ^{11} **B of** *G. ruber***.** Boron in seawater exists mainly as two different species, 497 boric acid (B(OH)₃) and borate ion (B(OH)₄⁻), and their relative abundance is pH dependent. There are 498 two isotopes of boron, ¹¹B (~80%) and ¹⁰B (~20%), with a ratio normally expressed in delta notation 499 as:

500
$$\delta^{11} B(\%) = \left[\left(\frac{{}^{11} B / {}^{10} B_{sample}}{{}^{11} B / {}^{10} B_{NIST951}} \right) - 1 \right] x1000$$
 [3]

where ${}^{11}B/{}^{10}B_{NIST951}$ is the isotopic ratio of NIST SRM 951 boric acid standard (${}^{11}B/{}^{10}B$ = 4.04367; ref. 60).

There is a pronounced isotopic fractionation between the two dissolved boron species, with boric acid being enriched in ¹¹B by 27.2‰ (ref. 61). As the concentration of each species is pH dependent, their isotopic composition also has to change with pH in order to maintain a constant seawater δ^{11} B. Calibration studies^{54,62,63} have shown that the borate species is predominantly incorporated into foraminiferal CaCO₃, and therefore ocean pH can be calculated from the δ^{11} B of borate ($\delta^{11}B_{borate}$) as follows:

509
$$pH = pK_{B}^{*} - \log_{\hat{\xi}}^{\hat{\ell}} - \frac{\mathcal{O}^{I1}B_{sw} - \mathcal{O}^{I1}B_{borate}}{\mathcal{O}^{I1}B_{sw} - \left({}^{11-10}K_{B} \cdot \mathcal{O}^{I1}B_{borate}\right) - 1000 \cdot \left({}^{11-10}K_{B} - 1\right)_{\hat{\emptyset}}^{\hat{\tau}}}$$
 [4]

510 where pK_{B}^{*} is the dissociation constant for boric acid at in situ temperature, salinity and pressure⁶⁴, 511 $\delta^{11}B_{sw}$ is the isotopic composition of seawater (39.61%; ref. 65), $\delta^{11}B_{borate}$ is the isotopic 512 composition of borate ion, and $^{11-10}K_B$ is the isotopic fractionation between the two aqueous species 513 of boron in seawater (1.0272±0.0006) (ref. 61).

514 In our calculations, temperature for ODP Site 999 is derived from Mg/Ca ratios measured on aliquots (separated after dissolution) of the same samples used for δ^{11} B analysis and for ODP Site 662 from 515 published records of temperature using the $U_{37}^{k'}$ proxy⁶⁶. Despite the uncertainty in Mg/Ca-derived 516 SST's we have not used published $U_{37}^{k'}$ temperature records for ODP Site 999 because they are of 517 lower temporal resolution and close to saturation (T = 28-29 °C)²⁵. Salinity has little influence on the 518 519 calculations of pH (± 1 psu = ± 0.006 pH units), and therefore is assumed to be constant at 35 psu 520 (similar to the present-day mean annual average at both locations). The uncertainty associated with this assumption is propagated into pCO_2^{atm} calculations. 521

522 Boron has a long residence time in seawater (10-20 Ma; ref. 67), and to account for likely (small)

523 changes in the boron isotopic composition of seawater ($\delta^{11}B_{sw}$) over the last 3 million years, we use a

524 simple linear extrapolation between modern $\delta^{11}B_{sw}$ (39.61‰; ref. 65) and the $\delta^{11}B_{sw}$ determined by

ref. 68 for the middle Miocene (12.72 Ma; $\delta^{11}B_{sw}$ = 37.8‰). This simple estimation yields $\delta^{11}B_{sw}$ =

526 39.2‰ at 3 Ma, which is consistent with available independent constraints, for example based on

527 assumptions of bottom water pH and measured benthic foraminiferal δ^{11} B (ref. 69).

528 Finally, in order to calculate pH from the δ^{11} B of *G. ruber*, it is necessary to account for species-

529 specific differences between $\delta^{11}B_{\text{borate}}$ in ambient seawater and $\delta^{11}B$ in foraminiferal calcite

530 ($\delta^{11}B_{\text{calcite}}$; i.e., "vital effects"). Here we used the species- and size-specific calibration equation of ref.

18 for *G. ruber* 300-355 μm (Equation 5). This equation has been applied in previous studies¹⁸ to

532 produce a δ^{11} B-based atmospheric pCO₂ (pCO₂^{atm}) record for the last 30 kyr that is in very good

agreement with ice-core pCO_2^{atm} records (Extended Data Figure 2).

534 $\delta^{11}B_{borate} = (\delta^{11}B_{calcite} - 8.87 \pm 1.52)/0.60 \pm 0.08 \text{ (uncertainty at } 2\sigma)$ [5]

535 It is important to note that, not only is there generally good preservation of the sites we use^{23,24}, but 536 also the δ^{11} B of *G. ruber* does not appear to be significantly affected by partial dissolution²⁵.

Determination of pCO₂^{atm} from δ^{11} B-derived pH. Another variable of the ocean carbonate system is 537 required besides pH in order to calculate the partial pressure of CO_2 in seawater (p CO_2^{sw})⁷⁰. Here, 538 total alkalinity (TA) is assumed to be constant at values similar to modern at ODP Site 999 (2330 539 μ mol/kg; ref. 20). It is important to note that pCO₂^{sw} estimates are mostly determined by the 540 541 reconstructed pH and that TA has little influence. This is because pH reflects the ratio of TA to DIC (total dissolved inorganic carbon), so when pH is known the ratio of TA:DIC is set, so the effect on 542 pCO₂^{sw} of a large increase/decrease in TA is partially countered by an opposite change in DIC. 543 Indeed, at a given pH, a change in TA by 10% only results in a pCO₂^{sw} change of 10%. For example, 544 modifying TA by $\pm 100 \,\mu$ mol/kg (a range equivalent to modelled variations in TA for the last 2 million 545

546 years; ref. 30) only modifies reconstructed pCO_2^{sw} (when pH is known) by less than ±12 µatm.

pCO₂^{sw} was calculated using the equations of ref. 70, the "seacarb" package of R (ref. 71) and a 547 548 Monte Carlo approach (n= 10,000) to fully propagate the uncertainty in the input parameters (at 95% confidence or full range, where appropriate): δ^{11} B (±analytical uncertainty, calculated using 549 550 Equation [1], and calibration uncertainty in Equation [5]), Mg/Ca-derived temperature (±3 °C), salinity (±3 psu), TA (±175 μ mol/kg), $\delta^{11}B_{sw}$ (±0.4‰). pCO₂^{atm} was then calculated from pCO₂^{sw} using 551 552 Henry's Law and subtracting the modern disequilibria with respect to CO₂ at the two sites (Extended 553 Data Figure 1; Supplementary Information Tables 1&2). Note that for the quoted uncertainty range for temperature, salinity, and $\delta^{11}B_{sw}$ a normal distribution is assumed. However, for TA we have 554 555 assumed a "flat" probability (i.e. an equal probability of TA being any value between 2155 and 2505 µmol/kg). We therefore do not ascribe weight to the assumption that TA remains constant, but 556 rather fully explore the likely range given the available, model based, constraints^{72,73}. It should also 557 be noted that salinity and temperature have little control on our estimated pCO_2^{sw} (+1 psu = +0.2 558 μ atm; +1 °C = +8 μ atm). 559

Comparison with published records of Pliocene pCO2^{atm}**.** Figure 1 and Extended Data Figures 2b&c 560 show a comparison of our new high resolution δ^{11} B-derived pCO₂^{atm} record with published records. 561 As noted in the main text, although the various approaches agree, in detail our new record exhibits 562 more structure. As a consequence, cross plots of the previously published CO₂ data against Δ MAT 563 (or SSTst) are largely incoherent (Extended Data Figure 2d-f). In the case of the stomatal estimates⁵⁰ 564 and the existing δ^{11} B-based records^{23, 25} this is largely a consequence of their low temporal 565 resolution, although analytical issues⁷⁴ and species choice (we use *G. ruber* that spends its entire life 566 cycle in the mixed layer whereas ref. 23 uses G. sacculifer that migrates during its life cycle and 567 whose δ^{11} B, unlike *G.ruber*, is modified by partial dissolution²⁵) may also play a role for the 568 discrepancy with earlier δ^{11} B records (see ref. 25 for further discussion). The lack of variability 569 570 through the Pliocene for the alkenone-based records may be related to changes in the size of the alkenone producers²⁶, fluctuations in nutrient content/water depth of maximum alkenone 571 572 production, and/or variations in the degree of passive vs. active uptake of CO_2 by the alkenone producing coccolithophorids^{49,75}. 573

Continuous records of Pliocene and late Pleistocene global temperature change. Robust records of global temperature change are needed to determine how the Earth's climate has responded to changes in CO_2 . Here we estimate this variable using two independent approaches: (i) we generate a stack of available sea surface temperature records (SSTst); and (ii) following ref. 35 we use a reconstruction of global mean annual surface air temperature change based on a scaling of the northern hemisphere temperature required by a simple coupled ice-sheet-climate model to forward model the benthic δ^{18} O stack of ref. 76 (tuned here to the LRO4 age model; Δ MAT).

For the SST stack (SSTst) we imposed a number of criteria for site selection. These are: (i) the record must be continuous from late Pliocene to late Pleistocene (or nearly so); (ii) the temporal resolution must be relatively high (ideally better than 1 sample per 10 kyr; for ODP Site 1237 we have however accepted a lower resolution to increase spatial coverage) to allow us to fully resolve the dominant 585 orbital-scale variability; (iii) be based on $U_{37}^{k'}$, given that Mg/Ca suffers an unacceptable level of uncertainty on these timescales due to the secular evolution of the Mg/Ca ratio of seawater (e.g., 586 ref. 48); and (iv) the temperatures recorded by the $U_{37}^{k'}$ proxy must be less than 29 °C, above which 587 the proxy becomes saturated and therefore unresponsive⁷⁵. Ten published records meet these 588 589 criteria (ODP Sites 982, 607, 1012, 1082, 1239, 846, 662, 722, 1237 and 1090; ref. 66, 78-85) and the 590 locations of these sites are shown in Extended Data Figure 2b. The average temporal resolution of 591 these records is 1 sample every ~4 kyr (ranging from ~2 to ~13 kyr) and the published age model of 592 each site is either part of the LR04 stack or was tuned to it (see the original publications for details).

593 In order to stack the records, each was first converted to a relative SST record referenced to either 594 the average of the Holocene (0-10 kyr), or mean annual modern SST if the Holocene is missing, and 595 then linearly interpolated to a 5 kyr spacing. These relative records are then averaged to produce a 596 single stacked record of relative SST change (SSTst; Supplementary Information Table 5). The 597 number of sites contributing to SSTst varies but for most of the record is ≥ 8 (Extended Data Figure 598 6a&b). Uncertainty on SSTst is estimated by a Monte Carlo procedure where 1000 realisations are 599 made of each individual SST record with noise added reflecting the magnitude of analytical uncertainty in the $U_{37}^{k'}$ SST reconstruction (± 1 °C at 2 σ ; ref. 75). Since we are using the same proxy 600 601 for each location it is not necessary to consider the calibration uncertainty as this should be the same for each record. Each SST realisation is then averaged to produce 1000 realisations of SSTst. 602 603 The mean of these 1000 realisations is then calculated and the 95% confidence interval is given by 604 the 2.5% and 97.5% percentile (red band on Figure 3). Jacknifing of SSTst (i.e. the sequential removal 605 of one record at a time) indicates that no particular record has undue influence and SSTst remains 606 close to the bounds relating to analytical uncertainty (the grey lines on Extended Data Figure 6c&d). 607 Our aim with SSTst was not to specifically reconstruct global SST change but rather to examine the

608 change in SST at these locations for a given forcing in the Pliocene and Pleistocene. We therefore do

not require SSTst to reflect global SST change. However, in order to assess how well SSTst does
reflect global SST we:

- (i) Examined the mean of historic SST change (1870AD to 2013 AD; from the HadISST
 database; ref. 86) at each location where we have an alkenone palaeo-SST record. This
 comparison is shown in Extended Data Figure 7 (blue circles). Despite exhibiting more
 variability than the mean annual global average (red in Extended Data Figure 7), these
 10 sites clearly capture the global long term trend in global mean SST^{87,88} over the last
 140 years or so (Extended Data Figure 7).
- 617 (ii) Compare SSTst to a multi-proxy and more comprehensive and independent compilation
 618 of ref. 32 that covers the last 100 kyr with >30 sites and the last 278 kyr with >10 sites.
 619 When data for the last 278 kyr are stacked together in a similar way to SSTst, the stack
 620 of ref. 32 (blue on Fig 3c) compares well with SSTst giving us confidence that it closely
 621 reflects global SST change.
- 622 (iii) Compare SSTst to discrete global reconstructions of SST. For the last glacial (20-25 kyr) 623 SSTst gives a Δ SST of -2.2 ± 0.4 K, which is close to the Δ SST of -3.2 K from a recent 624 comprehensive compilation for the LGM⁴² and is within uncertainty of earlier 625 reconstructions (e.g., ref. 85 where Δ SST of -1.9 ± 1.8 K). For the Mid-Pliocene Warm 626 Period (3-3.3 Ma), SSTst gives an average of +2.3 K. A simple mean calculated from the 627 larger multi-proxy PRISM SST compilation of ref. 40 is very similar at +2.6 K. SSTst is 628 slightly warmer than an area weighted mean of the PRISM SST set (+2 K; ref. 40).

Taken together, these comparisons clearly indicate that, although SSTst is made of a limited number of sites, it does appear to closely reflect change in global SST. This conclusion is also supported by the general agreement between the trends (but not absolute values) exhibited by Δ MAT and SSTst through the Pliocene and Pleistocene (Fig. 3), with subtle differences between these two climate records (e.g. at 2.8 Ma) potentially a result of a decoupling between deep and surface water
temperature evolution, small spatial biases in our SST stack, and/or minor age-model inaccuracies.

635 **Regression-based determinations of climate sensitivity** In order to examine the climatic response 636 (expressed as either Δ MAT or Δ SST) to forcing by CO₂ and land-ice albedo changes in both time 637 periods, we used a linear regression approach. Because each variable used (CO₂ and SL, Δ MAT or 638 Δ SST) has an associated uncertainty, however, it is necessary to fully explore the influence of these 639 uncertainties on our estimates of slope determined using least squares linear regression. Due to 640 difficulty of performing least squares linear regression with uncertainty in X- and Y- variables that are 641 not necessarily normally distributed we have used a two stage approach to fully propagate all the 642 uncertainties involved. Firstly, we generated 1000 realisations of each temporal record of each 643 variable (e.g. $\Delta F_{CO2, L}$, $\Delta F_{CO2, Ll}$, ΔMAT or ΔSST) based on a random sampling of each record within its 644 uncertainty envelope. This uncertainty envelope was either a simple normal distribution (e.g. ± 6 645 ppm for ice-core CO_2) or based on other Monte Carlo output (e.g. random sampling the 10,000 simulations of the Pliocene δ^{11} B-pCO₂^{atm} record or the 1000 realisations of SSTst; see above). Then 646 647 the first realisation of the ΔF_{CO2} (or $\Delta F_{CO2,LI}$) record was regressed against the first realisation of the 648 Δ MAT (or Δ SST) with the uncertainty in the slope and intercept of that regression determined using 649 a bootstrapping approach (n=1000; ref. 90). The second realisation of the forcing term and the 650 climate response was then regressed and the 1000 estimates of slope and intercept by 651 bootstrapping were combined with 1000 of the first regression. This continued for all 1000 realisations and a probability density function for the slope and intercept, accounting for X- and Y-652 653 uncertainty, was then constructed from the combined bootstrap estimates for each realisation 654 (n=1000000). The results of this approach are shown in Fig 5.

As noted above, pCO_2^{atm} (and hence ΔF_{CO2}) calculated from boron isotopes is a function of not only the measured $\delta^{11}B$ but also the total alkalinity (TA; or other second carbonate system variable) and, beyond the last 1 million years or so, the boron isotopic composition of seawater ($\delta^{11}B_{sw}$). This is

illustrated in Extended Data Fig 8. Here pCO₂^{atm} is calculated from an artificial δ^{11} B and temperature 658 record (Extended Data Fig 8a), a TA of either 2000 μ mol/kg, 2300 μ mol/kg or 2600 μ mol/kg, a δ^{11} B_{sw} 659 of 38.8, 39.6 (i.e. modern) or 40.4 % (Extended Data Fig 8) and the assumption that pCO₂^{atm} = 660 pCO₂^{sw}. These parameter choices result in a large difference in absolute CO₂ but, although they are 661 extreme and perhaps unlikely for the Pliocene, the slope of a linear regression of global temperature 662 change and ΔF_{co2} is very similar for each set of parameters (Extended Data Fig 8c,d). So much so, 663 even with only a poor knowledge of $\delta^{11}B_{sw}$ (e.g. ± 0.8 ‰) and TA (e.g. ± 300 µmol/kg) the accuracy of 664 665 the relationship between reconstructed ΔF_{CO2} and temperature is not unduly impacted. The residence time of boron in seawater (10-20 Ma) ensures that changes in $\delta^{11}B_{sw}$ across the time 666 interval examined here (1 Myr) are unlikely to be large (<0.1 ‰; ref. 67) and so uncertainty in the 667 absolute value of $\delta^{11}\text{B}_{\text{sw}}$ and any changes across the study interval can be ignored for our 668 determinations of S^{ρ} . In all the previous calculations we assume that TA is randomly distributed 669 670 between 2155 and 2505 μ mol/kg, therefore accounting for all possible trends in TA across the time 671 interval studied within this range. However, to better examine the influence of a large secular shift in TA on our estimates of S^{ρ} we have imposed a 200 μ mol/kg decrease (TAd) or increase (TAi) across 672 673 our Pliocene study interval. The slope for the regressions using one parameter set (VDW11 and sea-674 level from ref. 46 recalculated by ref. 12) but with such a varying TA are shown in Extended Data Fig 675 8e&f. Even this relatively large secular change does not have a major influence on the estimated 676 slope, clearly illustrating that our assumptions regarding TA, both its absolute value and its secular evolution, have little influence on our calculated ΔF_{CO2} and hence our conclusions. 677

Pliocene pCO₂^{atm} variability The apparent cyclicity in our Pliocene CO₂ record can be investigated
using spectral analysis. Extended Data Fig 4c shows the evolutive power spectra for the Pliocene
pCO₂^{atm} and a ~100 kyr cycle is clearly dominant. Our sampling resolution is 1 sample ~13 kyr, which
is not sufficient to resolve cycles of a precessional length (e.g. 19 and 23 kyr) but may be adequate to
resolve obliquity (~41 kyr length) yet these cycles are apparently absent in the generated spectra

683 (Extended Data Fig 4c). To ensure our resolution is not biasing this result we have sampled the LR04 684 benthic δ^{18} O stack at our exact sampling resolution and examined the evolutive power spectra of 685 this sampled record (Extended Data Fig 4d). This analysis reveals the presence of 100 kyr and 41 kyr 686 cycles in the δ^{18} O data, despite our relatively low resolution, supporting the observation that the 687 dominant cycle in Pliocene pCO₂^{atm} is ~100 kyr.

- The magnitude of Pliocene pCO₂^{atm} variability, shown in Extended Data Fig 4a, is similar to that exhibited by published late and mid-Pleistocene δ^{11} B-pCO₂^{atm} records (green and red lines on Extended Data Fig 4a) and by the Late Pleistocene ice core data when noise that is approximately equivalent to our δ^{11} B-pCO₂^{atm} uncertainty is added (± 35 µatm; black dashed line on Extended Data Fig 4a). In contrast, the δ^{18} O variability for these time intervals increases markedly from the Pliocene to late Pleistocene as the magnitude of glacial-interglacial cycles increases (Fig 1e, Extended Data Fig 4b).
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808 Figure Legends for Extended Data Figures

Extended Data Figure 1. Maps of modern mean annual ΔpCO_2 and sea surface temperature

810 **labelled with site locations.** (a) Map of sites used for pCO_2^{atm} reconstruction with the mean annual 811 modern ΔpCO_2 from the reconstruction of ref. 21. (b) Map of the sites (and labelled with their

- depths) used to generate SSTst with mean annual modern SST from the World Ocean Atlas 2013 (ref.
- 813 91). Figures constructed and data visualised in Ocean Data View⁹².

814 Extended Data Figure 2. Comparisons of boron isotope based-pCO₂^{atm} estimates with other

methodologies and archives. (a) Estimates of pCO_2^{atm} from published $\delta^{11}B$ -records compared to ice-815 816 core CO₂ (red line; ref. 27-29). The dotted line is for $pCO_2 = 278 \mu atm$. In (a) the data of ref. 20 (blue 817 circles) have been recalculated in the same manner as described here for the Pliocene, including using the *G. ruber* δ^{11} B-pH calibration of ref. 18. Error band encompasses 68% (dark blue) and 95% 818 (light blue) of 10,000 Monte Carlo simulations of pCO₂^{atm} (as described in text). Also shown are the 819 *G. sacculifer* based δ^{11} B-pCO₂^{atm} record of ref. 30 (green circles). In this case error bars (± 25 µatm) 820 821 are as determined in that study. Despite similar analytical uncertainty, the smaller error bars for the ref. 30 data result from these authors not propagating the $\delta^{\rm 11} {\rm B}\mbox{-}p{\rm H}$ calibration uncertainty and 822 823 considering a smaller range in temperature, salinity and alkalinity uncertainty than in this study (± 824 0.76 °C, \pm 1 psu, \pm 27 µmol/kg vs. \pm 3 °C, \pm 3 psu, \pm 175 µmol/kg with a flat probability in this study). (b) δ^{11} B-based pCO₂^{atm} record generated here (blue closed circles and 95% and 68% uncertainty 825 band) with pCO₂^{atm} from the δ^{13} C of alkenones from published studies. See caption for Figure 1 for 826 details. (c) δ^{11} B-based pCO₂^{atm} record generated here (blue closed circles and 95% and 68% 827 uncertainty band) with pCO₂^{atm} from previous δ^{11} B-based studies and from plant stomata. See 828 caption for Figure 1 for details. (d - f) Comparison of cross plots of CO₂ forcing and Δ MAT for our 829 high resolution $\delta^{11}B$ -CO₂ record (d), published alkenone-CO₂ (e) and published low resolution $\delta^{11}B$ -830 CO₂(f). In each panel the slopes of regression lines fitted through the data are labelled (± 1 se). In 831 (d) ice-core CO_2 are shown in red open circles and Pliocene $\delta^{11}B$ -CO₂ in open blue circles. In (e) and 832

(f) ice-core CO₂ data are shown in grey for clarity. In (e) alkenone-CO₂ data are from the following sources: ODP 1208 (orange, ref. 16), ODP 806 (purple, ref. 16); ODP 925 (brown, ref. 49); ODP 999 (green circles = ref. 25; green squares = ref. 26). In (c) δ^{11} B-CO₂ are from ODP999 (blue²⁵ and red²³).

836 Extended Data Figure 3. Probability density functions for equivalent aged samples from ODP Site

662 and ODP Site 999. Each panel, labelled with age in ka, shows the probability density function for
a given estimate of pCO₂^{atm} from ODP Site 662 (red) and ODP Site 999 (blue). In most instances
equal age samples are compared, but in some cases either where variability is high and/or
equivalent age samples are absent, we show neighbouring samples from ODP Site 999 (e.g. bottom
left and right). This comparison indicates that although the mean pCO₂^{atm} of ODP 662 tends to be
higher than ODP 999, there is always significant overlap between the estimates from the two sites.

Extended Data Figure 4. Probability density functions of pCO₂^{atm} and benthic δ^{18} O and time series 843 **analysis.** (a) Probability density functions (pdf) of the residuals of $\delta^{11}B$ -pCO₂^{atm} about the long-term 844 trend for the late Pliocene (this study; blue line), the mid-Pleistocene³⁰ (green line) and late-845 Pleistocene^{19,20,21} (red line). Dashed vertical lines show the upper and lower limit (labelled in figure) 846 encompassing 90 % of the data. The residual of the ice-core CO₂ record²⁷⁻²⁹ about the long-term 847 mean for 0 – 0.8 Ma plus a random noise equivalent to \pm 35 µatm (the typical δ^{11} B-CO₂ uncertainty) 848 is shown as a black dashed pdf. (b) Probability density functions of the residual of LR04 benthic δ^{18} O 849 850 from the long-term trend for the late-Pleistocene (red), mid-Pleistocene (green) and late Pliocene 851 (blue). Dashed vertical lines show the upper and lower limit (labelled in figure) encompassing 90% of the data. In contrast to the pCO₂^{atm}, δ^{18} O clearly exhibits an increase in variability over the last 3.3 852 million years. (c) Evolutive power spectral analyses of Pliocene pCO₂^{atm} and resampled δ^{18} O (d). The 853 854 evolutive power spectra was computed using the fast Fourier transform of overlapping segments 855 with a 300,000-year moving window. Before spectral analysis, all series were notch-filtered to 856 remove the long-term trend (bandwith = 0.005), and interpolated to 12 kyr intervals (the real 857 resolution of our record is ~13.5 kyr).

858 Extended Data Figure 5. Summary of sea-level records used to calculate ΔF_{LI} . In (a) and (b) the red curve is from ref. 13 (R14) based on the planktic δ^{18} O from the Mediterranean Sea and the methods 859 developed for the Red Sea by ref. 89. We have removed those intervals identified as possible 860 861 Sapropel events and linearly interpolated across gaps in the original record. The black curve is the 862 sea-level record from an inversion of the benthic oxygen isotope record of ref. 72 (tuned to LR04 here) using an ice sheet model³⁵ (VDW11). The blue curve in (a) is based on the planktic/bulk δ^{18} O 863 from the Red Sea⁴⁴ for the interval 0-520 ka and the paired Mg/Ca and benthic δ^{18} O from the deep 864 South Pacific for the interval 520-800 ka⁴⁵ (R09+E12). The green curve in (b) is based on a scaling of 865 the LR04 δ^{18} O stack to indicators of sea-level from sequence stratigraphy (ref. 46 recalculated by ref. 866 12). In each the uncertainty in the reconstruction at 95% confidence is shown by an appropriately 867 868 coloured error band. Marine isotope stages mentioned in text are labelled.

Extended Data Figure 6. Stacked sea surface temperature record (SSTst). (a) and (b) Number of
records that contribute to SSTst through time. (c) and (d) uncertainty in SSTst due to analytical
uncertainty (at 95% confidence; red band) and showing the influence of jacknifing (i.e. removing one
record at a time; grey lines show maximum and minimum). Note that the jacknifing illustrates that
no one record has undue influence on SSTst.

874 Extended Data Figure 7. Comparison of global SST from HadSST3 dataset with SST HadISST1 from

ODP sites. (a) Historic global mean annual sea surface temperature anomaly from the HadSST3
dataset⁸⁷⁻⁸⁸ (red circles) and mean SST at locations above the ODP sites that make up SSTst from
HadISST1 (blue; local SST). Thick red and blue lines are non-parametric smoothers through both
datasets. (b) Cross plot of global mean annual SST and local SST. The regression line determined
using linear regression has a slope of ~1 and intercept of close to 0, therefore, local SST captures the
global trend well. The shaded blue band in (b) represents the 95% confidence interval of the
regression line.

882	Extended Data Figure 8. The influence of TA and $\delta^{11}B_{sw}$ on determinations of S ^P using linear
883	regression. Artificial δ^{11} B record (a) and temperature record (b). Cross plot and regressions of δ^{11} B-
884	ΔF_{CO2} and global temperature for dramatically varying total alkalinity from 2000 to 2600 $\mu \text{mol/kg}$
885	(TA; c) and $\delta^{11}B_{sw}$ from 38.8 to 40.4 ‰ (d). The slopes of the regressions, which are very similar
886	regardless of parameter choice, are colour coded and listed in the bottom right hand corner of (c)
887	and (d). (e) Probability density function of slope for regressions of Pliocene-aged Δ MAT against
888	ΔF_{CO2} and (f) $\Delta F_{CO2,LI}$, where TA is decreasing by 200 (dashed) and increasing by 200 μ mol/kg
889	(dotted). Note that despite large variations in TA the slope of the regressions do not change
890	significantly.
891	
892	









