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Application of sediment core modelling to interpreting the glacial-interglacial record of Southern Ocean silica cycling

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Abstract. Sediments from the Southern Ocean reveal a meridional divide in biogeochemical cycling response to the glacial-interglacial cycles of the late Neogene. South of the present-day position of the Antarctic Polar Front in the Atlantic sector of the Southern Ocean, biogenic opal is generally much more abundant in sediments during interglacials compared to glacials. To the north, an anti-phased relationship is observed, with maximum opal abundance instead occurring during glacials. This antagonistic response of sedimentary properties provides an important model validation target for testing hypotheses of glacial-interglacial change against, particularly for understanding the causes of the concurrent variability in atmospheric CO$_2$. Here, I illustrate a time-dependent modelling approach to helping understand climates of the past by means of the mechanistic simulation of marine sediment core records. I find that a close match between model-predicted and observed down-core changes in sedimentary opal content can be achieved when changes in seasonal sea-ice extent are imposed, whereas the predicted sedimentary response to iron fertilization on its own is not consistent with sedimentary observations. The results of this sediment record model-data comparison supports previous inferences that the changing cryosphere is the primary driver of the striking features exhibited by the paleoceanographic record of this region.

1 Introduction

The Southern Ocean can be though of as the Achilles Heel of the modern global carbon cycle, with changes in: biological productivity (Martin, 1990; Watson et al., 2000), vertical mixing and stratification (Francois et al., 1997; Toggweiler, 1999; Toggweiler et al., 2006), and sea-ice cover (Stephens and Keeling, 2000) all suspected of playing a key role in modifying atmospheric CO$_2$ on glacial-interglacial time scales. Changes in the surface ocean environment of this region will be manifested in interlinked properties of sedimentary material deposited to the ocean floor, thus recording the mechanism(s) at work in controlling the observed CO$_2$ variability. However, the paleoceanographic evidence from the sediments of this region has not proved possible to interpret unambiguously (Anderson et al., 2002; Elderfield and Rickaby, 2000; Sigman and Boyle, 2000). The difficulty faced in quantitative reconstruction of the surface environment directly from sedimentary properties is a consequence of the different nonlinear processes that interact to determine a particular sediment state (i.e., the measured proxy value).

A promising modelling methodology is available to address this model-data divide – working forwards from an explicit description of the biogeochemical processes involved towards the observations. By recording the properties of biogenic material preserved in deep-sea sediments within an ocean-sediment carbon cycle model, sediment core records can be generated (Heinze, 2001; Ridgwell, 2007$^1$). This relatively data-friendly model output can then be contrasted rather directly with the observed sediment record. Here I illustrate the potential of this approach by evaluating the consequences of two possible drivers of glacial-interglacial environmental change in the Southern Ocean against measured changes in sedimentary opal content.

2 Modelling methodology

In this study, I use the ocean-sediment carbon cycle model of Ridgwell (2001). This is based on a zonally-averaged representation of ocean circulation, shown schematically in Fig. 1. To better resolve the effect on sediment composition

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$^1$Ridgwell, A.: Interpreting transient CCD changes by marine sediment core modeling, Paleoceanography, submitted, 2007.
of changes in sea-ice cover in the Southern Ocean, the two 7.5° zones south of 55° S in the parent model are sub-divided into a total of six sub-zones of width 2.5° as described in Ridgwell (2001). Each of these sub-zones is assumed to be characterized by the same vertical velocity as its parent zone with meridional velocities obtained by linear interpolation between 55° S and 62.5° S and extrapolation south of 62.5° S.

Overall, ocean-atmosphere biogeochemistry is similar to that used in previous work on the atmospheric CO₂ implications of glacial-interglacial (Watson et al., 2000) and anthropogenic (Ridgwell et al., 2002) changes in aeolian iron supply to the ocean. Tracers advected in the ocean component include total dissolved inorganic carbon (DIC), dissolved oxygen (O₂), alkalinity, temperature, and salinity. Of these, CO₂ and O₂ are exchanged with a “well-mixed” atmosphere across the air-sea interface. The stable isotopes of carbon (¹³C and ¹²C) are treated separately, with all major fractionation processes between them taken into account. Three nutrients potentially limiting to biological activity in the ocean are considered; phosphate (PO₄), silicic acid (H₄SiO₄), and total dissolved iron (Fe). Nutrients, together with DIC and ALK, are taken out of solution in the sunlit surface ocean layer (euphotic zone) through biological action, and exported as particulate organic matter, CaCO₃, and opal to deeper layers. As this particulate material settles through the water column, it is subject to remineralization processes, resulting in the release of dissolved constituent species to the ocean, or it may be buried in accumulating sediments. To accomplish this, the ocean is everywhere underlain by a series of discrete sediment modules handling ocean-sediment interactions, in which the preservation of biogenic CaCO₃ and opal reaching the sediment surface is explicitly predicted. Loss of material through burial in the sediments is balanced over the long-term by prescribed inputs to the ocean representing continental weathering and geothermal processes (in the case of DIC, ALK, and H₄SiO₄) as well as aeolian input at the surface (for Fe). For full description and evaluation of atmosphere-ocean-sediment carbon cycling in the model, readers are referred to Ridgwell (2001) and Ridgwell et al. (2002).

The configuration of deep-sea sediments is similar in nature to that of earlier box modelling work (Munhoven and Francois, 1996; Walker and Opdyke, 1995). Sediments are considered at 15 discrete water depths, spanning the depth of the ocean interior from 6000 m up to base of the euphotic zone (100 m), for a total of 315 separate sedimentary modules (i.e., 15 for each of the 21 ocean grid points). Each sediment module comprises a single, 5 cm thick homogeneous surface layer, underlain by a series of 1 cm thick sub-layers, illustrated in Fig. 2. The surface layer represents the upper zone of the sediment where solid composition is near homogeneous with depth and where the primary diagenetic processes take place. Excess solid material is exported out of this surface layer (i.e., buried) and stored in a stack of sedimentary sub-layers lying immediately below. The presence of the stack enables the progressive erosion of material during times of net dissolution at the sediment surface to be accounted for, a process which is critical to the millennial-scale buffering of atmospheric CO₂.
Fig. 2. Schematic of global biogeochemical cycling in the model and structure of the sediment modules. The initial state of the marine silica cycling is indicated, with fluxes (of Si as either silicic acid or biogenic opal) in units of Tmol Si yr\(^{-1}\). The positioning of the 15 sediment modules are indicated by dotted boxes with their structure shown to the right of the figure alongside the prescribed biodiffusion mixing profile. Each comprises a 5 cm thick surface layer in which diagenesis is assumed to take place, underlain by a stack of (1 cm thick) storage or sediment compositional “memory” layers. The preservation of calcium carbonate (CaCO\(_3\)) in the sediments is also calculated (see, Ridgwell, 2001; Ridgwell and Hargreaves, 2007) but not shown in this schematic. (Ridgwell and Hargreaves, 2007). It also provides a means of simulating how changes in the solid and isotopic composition of the sediments in response to global environmental change (glacial-interglacial cycles in this example) are recorded in the marine record.

In addition to the advective transfer of solids between the surface layer and the stack arising from net sedimentary accumulation/erosion, a transfer of material is prescribed between sub-layers representing the vertical mixing of solids by the action of benthic animals – bioturbation. The effect of bioturbation is modeled here as a quasi-diffusive process (Pope et al., 1996), with the mixing rate between pairs of layers in the sediment stack determined by a biodiffusion coefficient which decreases with an e-folding depth of 1 cm and has a maximum mixing rate at the top of the stack of 16 cm\(^2\) kyr\(^{-1}\) adapted from Peng et al. (1979) (and see Ridgwell, 2001). Mixing also occurs between the surface layer and upper-most sediment stack layer. The prescribed biodiffusion profile is shown in Fig. 2.

One advantage that modelled sediment records have over the data is that the precise age of material deposited to the sediment surface is known. A numerical tracer of the time of deposition is used to tag material reaching the sediment surface, allowing a mean age for each sediment sub-layer to be tracked (Ridgwell, 2001). In this way, an internal age-scale is generated, alleviating the need for the tuning of a \(\delta^{18}O\) stratigraphy to an orbital template (e.g., Bassinot et al., 1994).

The model is run over approximately four glacial-interglacial cycles (400 kyr) following an initial 150 kyr spin-up. Ideally, ocean biogeochemical processes would be informed by a suite of surface environmental boundary conditions (such as dust flux and sea-ice extent) generated internally within an integrated Earth system model. Instead,
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Fig. 3. Influence of glacial-interglacial variability in aeolian Fe supply on the sedimentary opal record of the Southern Ocean. Warm (interglacial) and cold (peak glacial) marine isotope stages (Bassinot et al., 1994) are highlighted in orange and blue, respectively. Left-hand panel: dust flux forcing (Watson et al., 2000) applied to the region 47.5° S to 70° S in the model. The middle 3 panels show observed (thick green line, top axis scale) and model-predicted (red line, bottom axis scale) down-core variability in wt% opal content for sediment cores lying either side of the APF. Observed data is based on cores: RC13-259 (53.9°, 04.9° W), RC13-254 (48.6° S, 05.6° W), and V22-108 (43.2° S, 03.2° W), all taken from the Atlantic Sector of the Southern Ocean (Charles et al., 1991; Mortlock et al., 1991). Core RC13-259 exhibits an anomalous pattern through Stages 3 and 4 compared to other cores south of the APF in this area (Mortlock et al., 1991), a result of a hiatus occurring around this level (Anderson, R. F., personal communication). Data from the nearby core RC11-77 (L. H. Burckle, personal communication) is therefore spliced into the interval ~12–90 kyr BP, marked in dark green. Each synthetic core is recovered from a grid point having approximately the same frontal coordinate (Chase et al., 2003), that is, a similar relative latitude to the position of the APF in the model (taken to be 55° S) as the real cores (where the observed APF in the Atlantic sector of the Southern Ocean is ~50° S (Mortlock et al., 1991)). The simulated sediment core wt% opal signal is plotted using an internal age stratigraphy (Ridgwell 2001). Far right-hand panel shows observed (black dotted line with blue circles marking the data points) (Petit et al., 1999) and model simulated (red line) atmospheric CO₂.

because this particular model lacks an interactive climate component, a time history of surface environmental boundary conditions is generated by the transformation of paired paleoclimatic reconstructions (made at discrete time slices) by a suitable continuous proxy signal – e.g., see Ridgwell (2001), Watson et al. (2000), or Köhler and Fischer (2006).
3 Testing hypotheses for G-I change using synthetic sediment cores

Records of biogenic opal content in deep-sea sediments from the Atlantic sectors of the Southern Ocean reveal a pronounced meridional asymmetry (Charles et al., 1991; Mortlock et al., 1991; Kumar et al., 1995; Anderson et al., 1998), with glacial-interglacial changes in composition of sites lying approximately either side of the position of the present-day Antarctic Polar Front (APF) being opposite in sign (e.g., see Fig. 3). To the south, interglacial-age sediments are characterized by higher opal contents than that which occurs during glacial periods, while to the north, it is glacial sediments that are the more opal-rich. These changes are paralleled by indicators of Si utilization (De La Rocha et al., 1999), suggesting that the antagonistic sedimentary response in wt% opal primarily reflects a change in the surface ocean environment and biogenic opal export. I now force the model with time varying glacial-interglacial changes in surface ocean boundary conditions and analyse the resulting signals contained in the simulated sediment core records to help interpret these observations.

3.1 Aeolian iron supply

Enhanced aeolian iron supply to the glacial Southern Ocean has been hypothesized to drive a stronger biological pump and lower the concentration of CO$_2$ in the atmosphere (Martin, 1990). Although considerable uncertainties remain regarding the operation of the iron cycle in this region (Ridgwell and Watson, 2002), results from ocean carbon cycle models go some way to supporting the iron hypothesis and suggest that changes in dust supply may have been responsible for anywhere between 5 and 45 ppm of the observed glacial-interglacial variability in atmospheric CO$_2$ (Archer et al., 2000; Bopp et al., 2003; Watson et al., 2000). Changes in iron availability at the ocean surface may be potentially intertwined with glacial-interglacial changes in opal accumulation in the underlying sediments as follows. The physiological effects of enhanced Fe availability on diatom cellular composition and more efficient utilization of silicic acid (Watson et al., 2000) has been hypothesized to lead to “leakage” (increased transport) of H$_4$SiO$_4$ to lower latitudes during glacials (Brzezinski et al., 2002; Matsumoto et al., 2002). This might potentially help explain the reduction in opal accumulation observed to the south of the APF (Sigman and Boyle, 2000), but increased to the North. To test whether aeolian iron supply plays a fundamental controlling role in the export of opal from the surface ocean of this region, the model is forced with a varying dust flux signal to the Southern Ocean (47.5°–70.0° S) following Watson et al. (2000). The forcing function together with the resulting predicted changes in the last 250 thousand years of sediment accumulation in the Southern Ocean are shown in Fig. 3.

South of the APF, a change in aeolian iron supply has little effect on sedimentary opal content, while to the north, model sediments are strongly anti-phased with the data. Although organic carbon export in the model is enhanced by ∼100% south of the APF as a result of increased Fe availability during the last glacial, the model-parameterized effect of Fe availability on Si utilization efficiency by diatoms (manifested in the Si:C export ratio (Ridgwell, 2001)) restricts the increase in opal export to just ∼34%. The increased opal rain to the sediments is partly offset by a 14% draw-down of the glacial silicic acid (H$_4$SiO$_4$) inventory compared to the subsequent (present) interglacial, which enhances opal dissolution rates. There is also an increase in the dust dilution of opal during the glacial which further decouples the change in wt% opal recorded in the sediments from the opal rain rate. The net result is that there is relatively little imprint of higher productivity on the sedimentary opal record at this location.

To the north, a slight reduction in advected Si supply coupled with greater Si utilization efficiency during glacial times combine to produce a substantial decrease in opal accumulation, contrary to the data. While different assumptions regarding the dependence of Si:C export ratios on Fe availability could conceivably improve the simulation, it is unlikely that a simultaneous match could be achieved to the data both north and south of the APF.

Of the total ∼90 ppm amplitude of glacial-interglacial CO$_2$ variability, 27 ppm is explained by this mechanism. The model predicted phasing between decreases in dust forcing and initial increases in CO$_2$ is consistent with observations as previously found (Watson et al., 2000) despite arguments that an apparent lag in response of up to ∼5 kyr exists (Broecker and Henderson, 1998). This can be understood in terms of a highly non-linear response of the carbon cycle to dust (Ridgwell and Watson, 2002; Gaspari et al., 2006). Thus, although the effects of dust are consistent with the timing of the observed intra-glacial variability in atmospheric CO$_2$, the sedimentary (opal) record is not consistent with the iron hypothesis as a sole driver of biogeochemical changes in the Southern Ocean between glacials and interglacials.

3.2 Sea-ice extent

Reconstructions of the cryosphere at the time of the Last Glacial Maximum (LGM) indicate that seasonal sea-ice cover was much more extensive than today in the Southern Ocean. In addition to being suggested as exerting a strong control on atmospheric CO$_2$ (Stephens and Keeling, 2000), this surface environmental change is also central to explanations for the observed paleoceanographic features of this region (Anderson et al., 2002; Charles et al., 1991; Crosta and Shemesh, 2002; Sigman and Boyle, 2000). However, the highly non-linear nature of sedimentary opal preservation (Archer et al., 2000; Ridgwell et al., 2002) and the unobvious coupling relationship between opal export and sea-ice cover (Chase et al., 2003) point to the need for analysis
within a numerical model to quantify whether changes in sea-ice extent could give rise to the observed features of the opal record.

As an illustration of how the cryosphere can modulate the sedimentary record, seasonal sea-ice limits are now modified in the model with present-day wintertime and summertime fractional sea-ice coverage of each model grid point taken from CLIMAP (CLIMAP, 1976) and interpolated at intermediate months according to monthly insolation. The maximum and minimum seasonal limits are then varied over the course of approximately four glacial-interglacial cycles, taking information regarding the timing and rate of change from the Vostok temperature record (Petit et al., 1999) and taking the absolute amplitude of the envelope from the difference between present-day and LGM CLIMAP reconstructions (Fig. 4). This assumes that a first-order correspondence exists between sea-ice cover and Antarctic air temperature, an assumption supported by coupled climate model results (Gildor and Ghil, 2002).

In contrast to the large decrease in atmospheric CO$_2$ (>45 ppm) reported in a box model of the ocean carbon cycle (Stephens and Keeling, 2000), a relatively muted response (~7 ppm) is found here to increased glacial sea-ice cover in the Southern Ocean (Fig. 4). This disparity may

![Fig. 4. The influence of glacial-interglacial variability in the seasonal limits of sea-ice extent on sedimentary opal content. Left-hand panel: change in maximum/wintertime (continuous line) and minimum/summertime (dotted line) fractional sea-ice cover applied to the model Southern Ocean (shown averaged over all model grid points lying south of the APF). Other panels as per Fig. 3. Note the shift in the wt%opal scale for the simulated core at 51.25° S relative to observations (RC13-254). The effect on wt%opal variability and atmospheric CO$_2$ of applying an alternative forcing assumption of CLIMAP winter-time sea-ice extent but with fixed modern summer-time extent is shown as a red dotted line.](image-url)
be a consequence of the lack of any explicit representation of equatorial up-welling zones (regions of intense CO₂ outgassing in the modern ocean (Takahashi et al., 1997)) in more highly idealized box models which can result in the Southern Ocean becoming the dominant source of CO₂ to the (modern) atmosphere (Stephens and Keeling, 2000). While the sea-ice forcing has little apparent impact on the global carbon cycle it does drive a marked change in sediment composition as shown in Fig. 4. Some of the primary features of the opal record are now well reproduced, particularly the transitions into and out of cold glacial stages (e.g., 2.2, 4.2, 6) and immediately following glacial inception in RC13-259. Crucially, the characteristic antagonistic variability either side of the APF is captured. This is a simple consequence of increased seasonal sea-ice cover south of the APF restricting biogenic export during glacial, with previously utilized nutrients advected northwards and fuelling greater (mainly diatom) export to the north (Anderson et al., 2002).

Recent reconstructions of the past state of the cryosphere have suggested that the minimum (summer-time) limit may have been little changed (e.g., Crosta et al., 1998; Gersonde et al., 2005). I have therefore tested the sensitivity of model predictions to assumptions about LGM sea-ice cover in an additional experiment in which the CLIMAP reconstruction is again used to inform the winter-time extent in the model, but the summer-time extent is now fixed at modern. The resulting recorded variability in wt% opal exhibits the same anti-phased response either side of the APF (Fig. 4). However, the amplitude of the variability is now substantially reduced.

Fig. 5. The effect of both dust and seasonal sea-ice limits combined. Sediment core record and atmospheric CO₂ panels as per Fig. 3.
Simultaneously applying both dust and sea-ice extent forcings leads to a further improvement in down-core opal variability, shown in Fig. 5. Of the total \( \sim 90 \) ppm amplitude of glacial-interglacial \( \delta^{13} \)C variability, \( 35 \) ppm is explained by these two mechanisms together.

### 4 Discussion

The information contained in the model sediments is determined by processes and environments representing large spatial means. In contrast, cores recovered from the deep sea may be influenced by local processes such as up-welling (Bertrand et al., 1996) and sediment redistribution by bottom currents (Bareille et al., 1994). Moreover, in order to maximize the temporal resolution of core data, drilling locations are often chosen where such local processes result in enhanced sedimentation rates. Close model-data correspondence in the absolute values of opal content is thus not expected a priori.

Despite these caveats, the match for core RC13-259 (Fig. 5) is surprisingly good, lending confidence to the plausibility of the model assumptions. It should be noted that the sedimentary response to glacial-interglacial perturbation has not been “tuned” in any way – adjustment of unconstrained parameter values is undertaken only in order to achieve a reasonable present-day ocean-sediment steady state (Ridgwell, 2001). Nor is there anything unique about the particular opal records chosen – alternative choices of pairs of observed and synthetic cores are found to exhibit similar matches. All 6 cores from north of the Antarctic Polar Front (APF) presented by Mortlock et al. (1991), show a relatively coherent glacial-interglacial pattern which is anti-phased with the 5 cores from south of the APF, which also exhibit a coherent pattern between themselves. Thus, it seems unlikely that the model-data comparison presented here is compromised by changes in sediment focusing or lateral transport. Indeed, the differences in \( ^{230} \)Th normalized opal flux between LGM and Holocene presented by Kumar et al. (1995) on the same cores as used here (i.e., RC13-259, RC13-254, V22-108) are consistent with the changes in wt% opal (Mortlock et al., 1991), and exhibit the same anti-phased relationship either side of the APF. However, in locations where lateral transport has been quantified and is important, in principal one could impose a sediment focusing factor in the model on the predicted particulate settling fluxes at the ocean-sediment interface (i.e., prior to calculating opal preservation in the sediments), which would enable the model-data comparison to be still carried out on the same terms as the data.

Combined, the two surface ocean forcings lead to a slightly better simile of the opal data than does changing sea-ice alone. This beneficial interaction is not obvious from the effects of the two mechanisms in isolation, demonstrating the additional value of a mechanistic modelling approach. Isotopic tracers can also be incorporated alongside bulk sediment fractions in the modelled cores (Ridgwell, 2001). For instance, simulations made of down-core variability in planktonic foraminiferal \( \delta^{13} \)C (not shown) suggest that sea-ice can also explain virtually all of the \( \sim 1 \% \) increase associated with the last deglacial transition exhibited by RC13-259 (Mortlock et al., 1991), consistent with interpretations of diatom-bound organic matter \( \delta^{13} \)C (Crosta and Shemesh, 2002). Explicit simulation of \( \delta^{15} \)N would allow the biogeochemical effects of further mechanisms such as ocean stratification (Crosta and Shemesh, 2002; Francois et al., 1997; Sigman and Boyle, 2000) to be disentangled from the sedimentary record.

Where the three sub Antarctic sectors are resolved in the model (Fig. 1), significant inter-sector differences are predicted with the amplitude of opal variability in the Pacific sector (not shown) only about \( 30-40 \% \) of that occurring at the same latitude in the Atlantic. It is also tempting to speculate on the relationship between the assumptions made regarding glacial summer-time sea-ice extent and the predicted amplitude of opal variability recorded in the sediments. Glacial-interglacial changes in wt% opal of a comparable magnitude to observations requires that glacial summer-time sea-ice extent is not drastically different from the CLIMAP reconstruction in the Atlantic sector of the Southern Ocean. Although the summer limit is known to be over-estimated in CLIMAP (e.g., Geronde et al., 2005), there is evidence for the “sporadic occurrence” of sea-ice reaching almost as far as the modern position of the APF. Making the assumption of little glacial-interglacial change in summer-time sea-ice extent as concluded by Geronde et al. (2005) for the Indian sector, results in little opal variability in the model. This is consistent with the general absence of pronounced differences in opal accumulation across the deglacial transition observed in the Indian sector (Dezileau et al., 2003).

One should recognize that the simplicity of the coarse resolution zonally-averaged model used here severely limits its ability to make deductions about different reconstructions of LGM sea-ice extent. In addition, no account is taken of changes in biogeochemical cycling due to the replacement of open ocean with seasonal ice zone ecosystems (Abelmann et al., 2006). The relationship between summer-time sea-ice extent and opal variability exhibited by model should thus be treated with caution. However, the results presented here hold out the prospect that within a fully sectorally-resolved (3-D) ocean circulation model of marine biogeochemical cycling (e.g., GENIE-1 (Ridgwell et al., 2007; Ridgwell and Hargreaves, 2007)); reconstruction of glacial-interglacial changes in sea-ice extent might be refined by means of simulation of the spatial and temporal patterns of opal accumulation. This would be particularly valuable for the summer-time extent in the Pacific sectors of the Southern Ocean.
which has proved difficult to reconstruct to date (Gersonde et al., 2005). Furthermore, by calibrating the sea-ice/opal accumulation relationship in the model against LGM reconstructions of sea-ice extent and observed sedimentary opal distributions, it might be possible to extend back in time estimates of changes in sea-ice extent through several glacial-interglacial cycles.

5 Conclusions

A full understanding of the mechanisms responsible for glacial-interglacial changes in the global carbon cycle as exemplified by ice-core records of atmosphere CO$_2$ still eludes us. Given the substantial uncertainties in some of the key components of the global carbon cycle and the wide variety of different hypotheses proposed to account for the glacial control of CO$_2$, improved use of data constraints is vital for constraining and informing modelling work. The results presented here demonstrate that comparisons made between synthetic sediment cores and paleoceanographic data can provide important additional insights into the processes involved in glacial-interglacial change. Although in isolation, reduced aeolian iron supply can explain a significant (5 to 45 ppm) fraction of the initial observed deglacial increase in CO$_2$ (Archer et al., 2000; Bopp et al., 2003; Ridgwell, 2001; Watson et al., 2000), the synthetic sediment records highlight the existence of important omissions from this simplistic picture. Changes in the seasonal limits of sea-ice can account for much of the missing sedimentary control. This suggests that the opal record is closely linked to the Antarctic climate signal, mediated by changes in the seasonal limits of sea-ice extent. By generating a model output that is comparable to the nature of the observed data, synthetic sediment cores provide a powerful means of quantifying past climatic changes and compliment more traditional ways of interpreting the paleoceanographic record.

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References

Abelmann A., Gersonde, R., Cortese, G., Kuhn, G., and Smetacek, V.: Extensive phytoplankton blooms in the Atlantic sector of the glacial Southern Ocean, Paleoceanography, 21, 1013, doi:10.1029/2005PA001199, 2006.

Anderson, R. F., Kump, N., Mortlock, R. A., Froelich, P. N., Kubik, P., et al.: Late-Quaternary changes in productivity of the Southern Ocean, J. Mar. Syst., 17, 497–514, 1998.

Anderson, R. F., Chase, Z., Fleisher, M. Q., and Sachs, J.: The Southern Ocean’s biological pump during the Last Glacial Maximum, Deep-Sea Res. Pt. II, 49, 1909–1938, 2002.

Archer, D., Winguth, A., Lea, D., and Mahowald, N.: What caused the glacial/interglacial atmospheric pCO$_2$ cycles?, Rev. Geophys., 38, 159–189, 2000.

Bareille, G., Grouset, F. E., Labracherie, M., Labeyrie, L. D., and Petit, J. R.: Origin of detrital fluxes in the southeast Indian Ocean during the last climatic cycle, Paleoceanography, 9, 799–819, 1994.

Bassinot, F. C., Labeyrie, L. D., Vincent, E., Quidelleur, X., Shackleton, N. J., and Lancelot, Y.: The astronomical theory of climate and the age of the Brunhes-Matuyama magnetic reversal, Earth Planet. Sci. Lett., 126, 91–108, 1994.

Bertrand, P., Simmield, G., Martinez, P., Grouset, F., Jorissen, F., et al.: The glacial ocean productivity hypothesis – The important of regional temporal and spatial studies, Mar. Geol., 130, 1–9, 1996.

Bopp, L., Kohfeld, K. E., Le Quere, C., and Aumont, O.: Dust impact on marine biota and atmospheric CO$_2$ during glacial periods, Paleoceanography, 18, 1046, doi:10.1029/2002PA000810, 2003.

Brzezinski, M. A., Pride, C. J., Franck, V., Sigman, D. M., Sarmiento, J. L., et al.: A switch from Si(OH)$_4$ to NO$_3^-$ deple- tion in the glacial Southern Ocean, Geophys. Res. Lett., 29, 1564, doi:10.1029/2001GL014349, 2002.

Broecker, W. S. and Henderson, G. M.: The sequence of events surrounding Termination II and their implications for the cause of glacial-interglacial CO$_2$ changes, Paleoceanography, 13, 352–364, 1998.

Charles, C. D., Froelich, P. N., Zibello, M. A., Mortlock, R. A., and Morley, J. J.: Biogenic opal in Southern Ocean sediments over the last 450,000 years: Implications for surface water chemistry and circulation, Paleoceanography, 6, 697–728, 1991.

Chase, Z., Anderson, R. F., Fleisher, M. Q., and Kubik, P. W.: Accumulation of biogenic and lithogenic material in the Pacific sector of the Southern Ocean during the past 40,000 years, Deep-Sea Res. Pt. II, 50, 799–832, 2003.

CLIMAP project members: The surface of the ice-age Earth, Science, 191, 1131–1137, 1976.

Cristol, X., Pichon, J. J., and Burckle, L. H.: Reappraisal of Antarc- tis seasonal sea-ice at the Last Glacial Maximum, Geophys. Res. Lett., 25, 2703–2706, 1998.

Cristol, X. and Shemesh, A.: Reconciling down core anticorrelation of diatom carbon and nitrogen isotopic ratios from the Southern Ocean, Paleoceanography, 17, 1010, doi:10.1029/2000PA000565, 2002.

De La Rocha, C. L., Brzezinski, M. A., DeNiro, M. J., and Shemesh, A.: Silicon-isotope composition of diatoms as an indicator of past oceanic change, Nature, 395, 680–683, 1999.

Dezileau L., Bareille, G., and Reys, J. L.: The $^{231}$Pa/$^{230}$Th ratio as a proxy for past changes in opal fluxes in the Indian sector of the Southern Ocean, Mar. Chem., 81, 105–117, 2003.

Elderfield, H. and Rickaby, R. E. M.: Oceanic Cd/P ratio and nutri- ent utilization in the glacial Southern Ocean, Nature, 405, 305–310, 2000.

Francois, R., Altabet, M. A., Yu, E.-F., Sigman, D. M., Bacon, M. P., et al.: Contribution of Southern Ocean surface-water strati- fication to low atmospheric CO$_2$ concentrations during the last glacial period, Nature, 389, 929–938, 1997.

Gaspari, V., Barbante, C., Cozzi, G., Cescon, P., Boutron, C. F., et al.: Atmospheric iron fluxes over the last deglacia-
tion: Climatic implications, Geophys. Res. Lett., 33, L03704, doi:10.1029/2005GL024352, 2006.

Gersonde, R., Crosta, X., Abelmann, A., and Armand, L.: Sea-surface temperature and sea ice distribution of the Southern Ocean at the EPLOG Last Glacial Maximum – a circum-Antarctic view based on siliceous microfossil records, Quat. Sci. Rev., 24, 869–896, 2005.

Gildor, H. and Ghil, M.: Phase relations between climate proxy records: Potential effect of seasonal precipitation changes, Geophys. Res. Lett., 29, 1024, doi:10.1029/2001GL013781, 2002.

Heinze, C.: Towards the time dependent modeling of sediment core data on a global basis, Geophys. Res. Lett., 28, 4211–4214, 2001.

Köhler P. and Fischer, H.: Simulating low frequency changes in atmospheric CO$_2$ during the last 740,000 years, Clim. Past, 2, 57–78, 2006, http://www.clim-past.net/2/57/2006/.

Kumar, N., Anderson, R. F., Mortlock, R. A., Froelich, P. N., Kubik, P., Dittrich-Hannen, B., and Suter, M.: Increased biological productivity and export production in the glacial Southern Ocean, Nature, 378, 675–680, 1995.

Martin, J. H.: Glacial-interglacial CO$_2$ change; the iron hypothesis, Paleoceanography, 5, 1–13, 1990.

Matsumoto, K., Sarmiento, J. L., and Brzezinski, M. A.: Silicic acid leakage from the Southern Ocean: A possible explanation for glacial atmospheric pCO$_2$, Global Biogeochem. Cy., 16, 1031, doi:10.1029/2001GB001442, 2002.

Mortlock, R. A., Charles, C. D., Froelich, P. N., Zibello, M. A., Saltzman, J., et al.: Evidence for lower productivity in the Antarctic Ocean during the last glaciation, Nature, 351, 220–222, 1991.

Munhoven, G. and Francois, L. M.: Glacial-interglacial variability of atmospheric CO$_2$ due to changing continental silicate rock weathering: A model study, J. Geophys. Res., 101, 21423–21437, 1996.

Petit, J. R., Jouzel, J., Raynaud, D., Barkov, N. I., Barnola, J.-M., et al.: Climate and Atmospheric History of the Past 420,000 years from the Vostok Ice Core, Antarctica, Nature, 399, 429–436, 1999.

Pope, R. H., Demaster, D. J., Smith, C. R., and Seltmann, H.: Rapid bioturbation in equatorial Pacific sediments – Evidence from excess Th$^{234}$ measurements, Deep-Sea Res. Pt. II, 43, 1339–1364, 1996.

Ridgwell, A. J.: Glacial-interglacial perturbations in the global carbon cycle, PhD thesis, Univ. of East Anglia at Norwich, UK (http://www.seao2.org/pubs/ridgwell_2001.pdf), 2001.

Ridgwell, A. J. and Watson, A. J.: Feedback between aeolian dust, climate and atmospheric CO$_2$ in glacial time, Paleoceanography, 17, 1059, doi:10.1029/2001PA000729, 2002.

Ridgwell, A. J., Watson, A. J., and Archer, D. A.: Modelling the response of the oceanic Si inventory to perturbation, and consequences for atmospheric CO$_2$, Global Biogeochem. Cy., 16, 1071, doi:10.1029/2002GB001877, 2002.

Ridgwell, A. and Hargreaves, J.: Regulation of atmospheric CO$_2$ by deep-sea sediments in an Earth System Model, Global Biogeochem. Cy., 21, GB2008, doi:10.1029/2006GB002764, 2007.

Ridgwell, A., Hargreaves, J., Edwards, N., Annan, J., Lenton, T., Marsh, R., Yool, A., and Watson, A.: Marine geochemical data assimilation in an efficient Earth System Model of global biogeochemical cycling, Biogeoosciences, 4, 87–104, 2007, http://www.biogeosciences.net/4/87/2007/.

Sigman, D. M. and Boyle, E. A.: Glacial/interglacial variations in atmospheric carbon dioxide, Nature, 407, 859–869, 2000.

Stephens, B. B. and Keeling, R. F.: The influence of Antarctic sea ice on glacial-interglacial CO$_2$ variations, Nature, 404, 171–174, 2000.

Stocke, T. F. and Wright, D. G.: Rapid changes in ocean circulation and atmospheric radiocarbon, Paleoceanography, 11, 773–795, 1996.

Takahashi, T., Feely, R. A., Weiss, R. F., Wanninkhof, R. H., Chipman, D. W., et al.: Global air-sea flux of CO$_2$: An estimate based on measurements of sea-air pCO$_2$ difference, Proc. Nat. Acad. Sci. U.S.A., 94, 8292–8299, 1997.

Toggweiler, J. R.: Variation of atmospheric CO$_2$ by ventilation of the ocean’s deepest water, Paleoceanography, 14, 571–588, 1999.

Toggweiler, J. R., Russell, J. L., and Carson, S. R.: Midlatitude westerlies, atmospheric CO$_2$ and climate change during the ice ages, Paleoceanography, 21, 2005, doi:10.1029/2005PA001154, 2006.

Walker, J. C. G. and Opdyke, B. C.: Influence of variable rates of netitic carbonate deposition on atmospheric carbon dioxide and pelagic sediments, Paleoceanography, 10, 415–427, 1995.

Watson, A. J., Bakker, D. C. E., Ridgwell, A. J., Boyd, P. W., and Law, C. S.: Effect of iron supply on Southern Ocean CO$_2$ uptake and implications for glacial atmospheric CO$_2$, Nature, 407, 730–733, 2000.