An investigation of carbon cycle dynamics since the Last Glacial Maximum: Complex interactions between the terrestrial biosphere, weathering, ocean alkalinity, and CO₂ radiative warming in an Earth system model of intermediate complexity

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Abstract

Proxy reconstructions and modeling studies of the glacial-interglacial changes in the global carbon cycle have led to a stimulating debate in the paleoclimate literature about the mechanisms leading to a 90-100 ppmv increase in atmospheric CO₂. In this paper, we used the University of Victoria Earth System Climate Model v. 2.9 to simulate the carbon cycle response to ice sheet retreat and Milankovitch (insolation) forcing from the Last Glacial Maximum (LGM) to the present. In addition, we conducted sensitivity studies to address the contributions of CO₂ radiative forcing, atmospheric carbon content, and weathering rates to climate and carbon cycle changes since 21 kyr BP. The simulations show that ice sheet and orbital changes by themselves do not lead to a notable increase in atmospheric CO₂ over the course of deglaciation. However, with the application of CO₂ radiative forcing and different weathering rates, the simulated atmospheric CO₂ variations ranged over ~35 ppmv. Virtually all of the simulated net global vegetation carbon uptake since the LGM is attributable to CO₂ fertilization rather than greater land availability or warmer temperatures. Furthermore, the ‘greening’ from CO₂ fertilization significantly enhances total deglacial warming (by 0.14°C) and contributes to warmer intermediate and deep ocean temperatures during the interglacial period. We also found that CO₂ radiative forcing was the dominant factor allowing for greater outgassing at the ocean surface and an earlier ventilation of deep-ocean DIC. The downwelling of high-alkalinity surface waters stimulated by a stronger, earlier overturning circulation led to greater deep sedimentation (alkalinity removal), which, in turn, permitted CO₂ to continue to increase through much of the simulation period.
1 Introduction

Significant changes in the carbon cycle following glacial terminations are well-documented in ice core records of atmospheric CO$_2$ (Petit et al. 1999; Monnin et al. 2001; Marcott et al. 2014); however, identifying the mechanisms that dominate these changes has proven to be a challenge. Different hypotheses suggest that oceanic, terrestrial, and geological (volcanic) sources of carbon may have contributed to the observed 90-100 ppmv increase in atmospheric CO$_2$ over the course of deglaciation (Simmons et al. 2016, in preparation).

With respect to oceanic sources of carbon, large changes in ocean stratification (Adkins et al. 2002; Duplessy et al. 2002; Hodell et al. 2003), vertical mixing (Watson and Naveira Garabato 2006) and water mass distributions (Marson et al. 2014) between the LGM and the present suggest that major circulation changes may have occurred during deglaciation, leading to a large ventilation and outgassing of glacial deep-ocean respired CO$_2$ to the atmosphere (e.g., Galbraith et al. 2007; Jaccard et al. 2009). The greater vertical mixing and upwelling of carbon-rich glacial deep water may have been enhanced dynamically by poleward-shifted (Toggweiler et al. 2006) or stronger (D’Orgeville et al. 2010) Southern Hemisphere westerlies during deglaciation, as well as reduced Southern Ocean seasonal ice cover (Stephens and Keeling 2000; Roche et al. 2012). Furthermore, increasing sea surface temperatures from the LGM to the pre-industrial era would have decreased the solubility of CO$_2$ in sea water, contributing to a net ~10-20 ppmv increase in atmospheric CO$_2$ after accounting for the compensatory effect of ocean freshening (Kohfeld and Ridgwell 2009; Brovkin et al. 2012; Menviel et al. 2012). This gradual surface ocean warming may have also caused a decrease in the remineralization depth of falling
organic matter, leading to more respiration nearer to the ocean surface and less trapping of
respired CO$_2$ in the deep ocean (Menviel et al. 2012). In addition, less dust/iron transport to and
nutrient availability in the surface ocean following a glacial termination (Lambert et al. 2008)
may have led to a weaker biological pump during the deglacial period (Parekh et al. 2008;
Galbraith and Jaccard 2015). Finally, the flooding of continental shelves would have led to
greater shallow-water sedimentation (Brovkin et al. 2012) and coral reef formation (Ridgwell et
al. 2003), allowing for long-term ocean alkalinity decreases, which would play its own role in
reducing ocean CO$_2$ uptake during the interglacial period.

Other studies have highlighted terrestrial and volcanic contributions to the observed
deglacial increase in atmospheric CO$_2$ concentrations. For example, recent estimates of changes
in terrestrial carbon stocks (Peterson et al. 2014; Ciais et al. 2012) suggest that the biosphere
sequestered a net ~330 Pg C between the LGM and pre-industrial period. At the same time,
peatland uptake (400-600 Pg C) since the beginning of the Holocene (~11000 BP) (Yu 2011) and
vegetation uptake (~550-694 Pg C in Prentice et al. 2011) were probably much greater. Thus, the
relatively modest net uptake of ~330 Pg C since the LGM suggests that a large terrestrial release
to the atmosphere occurred during the deglacial period, and likely from a passive soil carbon
reservoir, followed by a larger resequestration by the terrestrial biosphere during the interglacial
(Ciais et al. 2012). Much of this carbon released during the deglacial period was likely sourced
from glacial peatlands and permafrost (which extending to the mid-latitudes during the Last
Glacial Maximum) and vegetation growing along now-flooded continental shelves (see
Montenegro et al. 2006; Prentice and Harrison 2009; Zech 2012; Köhler et al. 2014; Simmons et
al. 2015; Brovkin et al. 2016). In addition, proxy records suggests that enhanced volcanic
activity and CO$_2$ emissions (associated with mantle decompression from ice sheet retreat) may
have provided a significant source of carbon to the atmosphere during the early Holocene
(Huybers and Langmuir 2011), potentially helping maintain elevated atmospheric CO$_2$
concentrations in the wake of accelerated terrestrial uptake from peatland and boreal forest
expansion (Roth and Joos 2012; Broecker et al. 2015).

In this paper we analyze the results of a transient run of the UVic ESCM v. 2.9 for the
entire period from the LGM to the present, with the purpose of evaluating the carbon cycle
response to dominant, well-documented physical and thermodynamic changes in the Earth
System over the period: ice sheet extent/height, insolation, and atmospheric CO$_2$-related
radiative forcing and carbon content. In particular, we discuss simulations that (1) allow the
model’s climate and carbon cycles to respond to the changes in the model-calculated CO$_2$, (2)
include (solely) the CO$_2$ radiative forcing from ice cores and (3) include a prescribed
atmospheric carbon content (which injects enough carbon into the model atmosphere to satisfy
the demands of other carbon reservoirs and produce the observed increase in atmospheric CO$_2$
content). The distinction between the three types of modeling experiments helps distinguish
processes related to CO$_2$ warming and CO$_2$ carbon content that contribute to climate changes that
enhance initial increases in atmospheric CO$_2$.

The results presented here build on past work using the UVic model, which explored
relatively short-term glacial (e.g., Schmittner et al. 2007; Schmittner and Galbraith, 2008) and
deglacial changes (Huiskamp and Meissner 2012; D’Orgeville et al. 2010). Because we have
performed a transient simulation for the entire deglacial period using a model with sediments
(Eby et al. 2009), we have been able to investigate the alkalinity feedbacks on the deglacial CO$_2$
rise. In particular, we carried out sensitivity experiments to explore deglacial and interglacial
changes with two different (constant) weathering rates. These simulations were compared with
other modeling experiments where alkalinity was held constant (weathering set equal to
sedimentation), thus isolating the collective contribution of alkalinity to CO$_2$ changes.

Below we provide a brief description of the model (Section 2.1) and the details of our
equilibrium simulation for the LGM (Section 2.2). The experimental design for transient
simulations of the period is given in Sections 2.3-2.5. Then, in Section 3, we discuss the results
for a suite of simulations, including a comparative analysis of the relative contribution of
weathering rate, CO$_2$ radiative forcing, and CO$_2$ carbon content to the global carbon cycle. The
major conclusions are given in Section 4.

2 Model and Methodology

2.1 Model Description

The University of Victoria Earth System Climate Model (UVic ESCM, or simply UVic
model) v. 2.9 provides a simplified but detailed representation of the Earth’s climate system and
carbon cycle and is classified as an Earth system Model of Intermediate Complexity (EMIC)
(Claussen et al. 2002). All components of the model operate on a grid of 1.8º latitude by 3.6º
longitude. The core of the model is a three-dimensional primitive equation ocean general
circulation model (the Modular Ocean Model v. 2.0, see Pacanowski (1995)) with 19 vertical
levels, coupled to a dynamic-thermodynamic sea ice module (Weaver et al. 2001). Eddy
transport is parameterized according to Gent and McWilliams (1990), and diapycnal ocean
(vertical) mixing is accomplished through a time-independent horizontally-constant diffusivity
profile (approximately 0.3 $10^{-4}$ m$^2$ s$^{-1}$ at the pycnocline). In its present configuration, the
model’s ocean regime is defined by unchanging present-day bathymetry and sea level, and thus
some important features of LGM and deglacial topography, such as continental shelves above
sea level, are not featured in the simulations discussed here. However, a closed Bering Strait is a feature of these simulations.

Concerning ocean chemical properties, inorganic carbon chemistry and air-sea exchanges of CO$_2$ (controlled by temperature, salinity, DIC, alkalinity, wind speed, and sea ice cover) are described in Ewen et al. (2004). Contributions of the biological pump to the ocean carbon cycle are calculated by a Nutrient-Phytoplankton-Zooplankton-Detritus (NPZD) module, which includes both nitrates and phosphates as well as nutrient recycling through microbial respiration and differential treatment of dissolved and particulate organic carbon (POC) (Schartau and Oschlies 2003; Schmittner et al. 2005; Schmittner et al. 2008). An important update to the 2.9 version of the UVic ESCM is the inclusion of an oxic-only sediment module from Archer (1996), which allows the model to evaluate secondary changes in ocean chemistry and atmospheric CO$_2$ due to sediment respiration and calcium carbonate compensation, although without the effect of coral reefs (Eby et al. 2009). In this model, the dissolution of CaCO$_3$ in falling detritus occurs with an e-folding depth of 3500 m (Schmittner et al. 2008). Once reaching the ocean floor, total CaCO$_3$ dissolution depends on the respired CO$_2$ content of overlying waters and the CaCO$_3$:POC rain ratio at the sediment level (which is calculated as the amount of unrespired organic carbon to calcite content of falling detritus that reaches the ocean floor). Changes in the rain ratio are constrained in the model to lie between 0.55 and 1.82 due to the buffering capacity of sediments (Ridgwell 2003) and the neglect of CaCO$_3$ ballasting effects in the current version of the model. In our simulations, however, the rain ratio does not actually vary significantly (1.05-1.27). It should be noted that all sedimentation occurs at or below a depth of 1240 m, and thus variations in shallow water sedimentation (with increasing sea level) are not parameterized in these simulations.
The model atmosphere is simplified to a two-dimensional Energy-Moisture Balance Model (EMBM) (Fanning and Weaver 1996; Weaver et al. 2001). Surface wind fields must be prescribed in order to drive horizontal moisture advection and the wind stress over the ocean. All simulations in this paper use a dampened version of the “wind feedback” in Weaver et al. (2001), which provides a thermal-wind adjustment to 20th-century reanalysis winds (Kalnay et al. 1996) based on the model-generated spatial temperature gradient at each time step. The atmospheric Δ^{14}C was kept constant at 0‰ for all simulations, and thus the oceanic Δ^{14}C profiles generated by the model are mostly a function of ocean ventilation changes. The fractionation ratio δ^{13}C is not traced in the current version of the model for either the atmosphere or the ocean.

Vegetation changes on land are driven by the Top-down Representation of Interactive Foliage and Flora Including Dynamics (TRIFFID) Module, which represents five plant functional types (PFTs): C_3-photosynthesis grasses (manifested as mid-latitude prairie/steppe and very high latitude tundra in the model), C_4-photosynthesis grasses (mostly tropical and subtropical savannah in the model), broadleaf trees (tropical and subtropical forest), needleleaf trees (predominately boreal and high-elevation forest), and shrubs (tropical/Mediterranean bushy vegetation and tundra-boreal forest transition regions) (Cox 2001; Meissner et al. 2003; Matthews et al. 2004). The distribution of PFTs is determined by temperature and soil moisture criteria, and in regions where several types of vegetation can grow, Lotka-Volterra equations (predator-prey interspecies competition based ultimately on a tree-shrub-grass dominance hierarchy and net primary production) determine the predominant species for each grid cell. The soil model (MOSES) stores and respires terrestrial carbon litter in a single layer with 1 m depth (Cox et al. 1999). There is no representation of peatlands in the current version of the model.
The UVic model also incorporates prescribed continental ice sheets and non-conducting marine ice shelves (represented as a “lid” over the ocean). Their thickness and areal coverage at the LGM and over the course of deglaciation are determined by the ICE-4G dataset (Peltier 2002). The geographical coverage and retreat of these ice sheets is prescribed every 1000 years according to the above data, with no ice melting (except of accumulated snow on or near the ice sheets) as a result of insolation or temperature changes generated by the model. As a result, ice sheet and ice shelf retreat does not produce a freshwater flux into the ocean or onto nearby land surfaces.

2.2 LGM Equilibrium Simulation

Slowly-varying atmospheric CO$_2$ and land ice cover make the LGM perhaps the most appropriate time period to generate a model equilibrium climate with constant forcing parameters. We selected a model spin-up start date of 21000 B.C. (~22950 BP), approximately 2000 years before the standard date for the LGM in order to ensure that glacial conditions were adequately captured by the model in transient simulations. Initiated from pre-industrial conditions (the default restart file for the UVic ESCM v. 2.9), this equilibrium simulation was run for 9700 model years with fixed CO$_2$ (191.1 ppm), orbital forcing (the Milankovitch forcing for 22950 BP), and land ice sheets (which are unchanged in the database from LGM-extent ice sheets at 21000 BP) and with the model’s wind feedback adjustment to NCEP reanalysis winds. The carbon content of the oceanic, terrestrial, and sediment reservoirs were equilibrated to maintain the constant atmospheric CO$_2$. As these studies focus only on CO$_2$, no adjustments to the CO$_2$ radiative forcing were made to account for lower concentrations of other greenhouse gases (namely CH$_4$ and N$_2$O). In order to maintain the assumption of an equilibrium carbon cycle just prior to the LGM, the weathering flux was set equal to the sedimentation rate in the spin-up.
simulation (i.e., constant ocean alkalinity and no nascent carbonate compensation at the beginning of transient simulations).

2.3 Transient Forcing

This equilibrium simulation for 22950 BP was then used as the initial conditions for a series of transient simulations exploring deglaciation with time-evolving orbital (Milankovitch) forcing and retreating ice sheets (ICE-4G) from 23000 BP to the present (ice data was interpolated between data points every 1000 years). A complete list of these experiments is provided in Table 1. Incorporating changes in land ice and insolation, Free Carbon (FC) transient simulations allowed the model’s carbon reservoirs to evolve freely without any other prescribed forcing beyond evolving ice sheets and insolation changes. The only CO$_2$ radiative forcing in these FC experiments was provided by model-generated atmospheric CO$_2$ concentration and not the observed record from ice cores. Prescribed Carbon (PC) simulations, however, forced the model’s carbon reservoirs to equilibrate to the observed ice core record of atmospheric carbon content, following the Vostok record (Petit et al. 1999) from 23000 BP to 7950 BP, Taylor Dome (Indermühle et al. 1999) from 7950 BP to 944 BP and from Law Dome (Etheridge et al. 1996) after 944 BP (as in Simmons et al. 2013). In contrast to the FC experiments, carbon was injected into (or removed from) the atmosphere in the PC simulations to balance the net sources and sinks from other carbon reservoirs (oceans, biosphere, sediments) in order to maintain the interpolated ice core CO$_2$ concentration for that date. A third series of simulations, testing model sensitivity to deglacial CO$_2$ radiative forcing and denoted “CO2rad” in the results discussion, allowed the model carbon cycle to evolve freely but applied the radiative forcing (warming effect) of CO$_2$ according to the ice core record. The CO$_2$ radiative forcing increased from 0 Watts m$^{-2}$ at 20954
BP and was calculated from the same ice core record (Vostok and Taylor/Law Domes) as in the PC simulations using the traditional formula (ΔF = 5.35 × ln(C/C₀), where C represents the linearly-interpolated CO₂ between data points from the ice core record and C₀ represents the CO₂ value from Petit et al. (1999) at ~20954 BP).

Table 1: A summary of the principal transient simulations discussed in this paper. The acronyms FC CA (free carbon, constant alkalinity), FC HW (free carbon, higher weathering rate), FC LW (free carbon, lower weathering rate), CO₂rad CA (CO₂ radiative forcing, constant alkalinity), CO₂rad HW (CO₂ radiative forcing, higher weathering rate), CO₂rad LW (CO₂ radiative forcing, lower weathering rate), PC CA (prescribed (atmospheric) carbon, constant alkalinity), and PC HW (prescribed (atmospheric) carbon, higher weathering rate) are described in Sections 2.3 and 2.4.

<table>
<thead>
<tr>
<th>Simulation Name</th>
<th>Milankovitch forcing</th>
<th>ICE-4G prescribed land ice</th>
<th>Ice core CO₂ radiative forcing starting at 19004 B.C.</th>
<th>Prescribed atmospheric CO₂ from ice cores</th>
<th>Weathering Rate</th>
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</thead>
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<tr>
<td>FC CA</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>sedimentation rate</td>
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<tr>
<td>FC HW</td>
<td>X</td>
<td>X</td>
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<td>144.0 Tg C/yr or 12.0 Tmol/yr</td>
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<tr>
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<td>120.7 Tg C/yr or 10.1 Tmol/yr</td>
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<td>CO₂rad CA</td>
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<td>CO₂rad HW</td>
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<td>144.0 Tg C/yr or 12.0 Tmol/yr</td>
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2.4 Weathering Rates

Changes in weathering rates can have important implications for ocean alkalinity. However, modeling studies investigating the influence of weathering changes are limited (Brovkin et al. 2012). A comprehensive weathering module is not available in the 2.9 version of the model, but the effects of weathering can be tested indirectly by using different constant weathering rates or by setting the weathering rate equal to the ocean sedimentation rate (the latter of which is determined by the model’s coupled ocean biogeochemistry). In simulations where weathering is set equal to sedimentation, denoted henceforth as CA for “constant alkalinity,”
there can be no carbonate compensation because weathering inputs directly balance precipitation to the sediments. Simulations with constant alkalinity allow for purely physical mechanisms (and their climate responses) to be isolated from those related to chemical changes in the ocean (alkalinity changes and carbonate compensation). The UVic ESCM, however, does not remove carbon from the terrestrial carbon reservoir (soil erosion) in order to maintain this weathering flux but rather generates new bicarbonate to inject into the ocean. Thus, as the model is currently configured, total carbon conservation is not possible for the constant weathering simulations.

Outside of the CA experiments, one set of simulations uses a “higher” weathering rate (HW) of 12.0 Tmol (Alk)yr\(^{-1}\) (or 4566 kg C s\(^{-1}\)), which is the equilibrium weathering rate obtained at the end of the spin-up simulation (Section 2.2). However, in the first 2000 years of both the FC and PC transient simulations (leading up to the actual LGM at 21000 BP), the sedimentation rate dropped abruptly before stabilizing to a lower rate at 21000 BP of ~10.1 Tmol yr\(^{-1}\). Thus, for a second series of “low weathering (LW)” experiments, we used 10.1 Tmol yr\(^{-1}\) as a constant weathering rate for the simulation from 21000 BP onward. The range of weathering rates used here may be relatively modest compared to fluctuations in weathering since the LGM.

**3 Results and Discussion**

**3.1 Atmospheric CO\(_2\)**

The range of processes considered here resulted in CO\(_2\) differences of up to 35 ppmv, though none of the simulations reproduced the full magnitude of the observed CO\(_2\) increase. The largest increase in CO\(_2\) (~25 ppm) from the LGM to the late Holocene is comparable to that obtained by an AOGCM study with a free carbon cycle (Chikamoto et al. 2012). Fig. 1 illustrates the results for transient experiments with the freely-evolving model carbon cycle (FC). Atmospheric CO\(_2\) (Fig. 1a) does not increase in any of the simulations without prescribed
radiative forcing, indicating that the large changes in land ice (the ice-albedo feedback) and
northern hemisphere insolation do not by themselves trigger mechanisms in the model to
increase CO$_2$. In fact, for the simulation with constant alkalinity (FC CA) and the higher
weathering rate (HW), atmospheric CO$_2$ declines 5-10 ppm from 21000 BP to present.

The results also suggest that the terrestrial carbon reservoir plays an intimate role in the
evolution of atmospheric CO$_2$ over the course of these simulations. The decrease in atmospheric
CO$_2$ over the period in the FC CA simulation is compensated by a 200 Pg C net increase in the
terrestrial carbon since the LGM (Fig 1b, blue line), a figure less than the ~330 Pg C increase
estimated by Ciais et al. (2012) and (Peterson et al. 2014). In the same simulation, a
compensating transfer of ocean carbon (~195 Pg C) to the atmosphere (Fig. 1b, blue line) is
partially hindered by an increase in export production (not shown). The lower constant
weathering rate (LW), however, produced a slightly greater atmospheric CO$_2$ (195 ppm) by the
mid-Holocene (Fig. 1a, green line). The simulation with the higher constant weathering rate
(HW) yielded, in turn, the lowest atmospheric CO$_2$ value just above 180 ppm at the end of the
Holocene (Fig. 1a, purple line). The greater weathering is an alkalinity source to the oceans and
causes atmospheric CO$_2$ to decrease unless sedimentation (the alkalinity sink) outpaces
weathering, whereas lower weathering leads to a decrease in ocean alkalinity if sedimentation
outpaces the combined effect of weathering and sediment dissolution (alkalinity sources).

The dip in CO$_2$ in most experiments after 6000 BP (Fig. 1a) occurred in response to a
large reduction in Antarctic marine ice shelves. Simmons et al. (2013) showed that, with the
disappearance of Antarctic marine ice shelves over the course of an interglacial, open ocean
increasingly expanded into the (prescribed) katabatic wind environment around Antarctica,
leading to more air-sea exchange, sea ice formation, and brine rejection. This resulted in an
enhancement of bottom water formation near the coast of Antarctica, which in turn led to a greater prominence of deep, high-DIC Antarctic bottom waters globally. These denser abyssal waters also led to poorer abyssal ventilation, resulting in an accumulation of DIC at depth and a corresponding drawdown in atmospheric CO$_2$ of about 5 ppm. In a sensitivity study (not shown) of the FC LW simulation with a slightly more extensive ice shelf distribution (as in Simmons et al. 2013), the atmospheric CO$_2$ concentration rose to 199 ppmv between 6000 BP and present rather than the declining further as shown here.

For the simulations including CO$_2$ radiative forcing, a total post-glacial CO$_2$ rise of approximately 15 ppm occurred in the CO2rad CA simulation (Fig. 1a, red line). Compared to the same constant alkalinity simulation without radiative forcing (Fig. 1a, blue line), the net effect of this prescribed warming is 20 ppm more CO$_2$ in the atmosphere. This 20 ppm difference represents the combined effect of both outgassing (from warmer ocean surface temperatures) and deep-ocean ventilation of deep DIC in the model’s atmospheric CO$_2$ balance (Fig. 1b, red line). Furthermore, by allowing alkalinity to respond to ocean ventilation changes, the influence of CO$_2$ radiative forcing on the HW and LW profiles led to an even greater increase in CO$_2$ (to 215-230 ppm) due to a notable decrease in both DIC and alkalinity associated with a better-ventilated ocean (Fig. 1a, orange and brown lines).

### 3.2 Terrestrial Carbon

While atmospheric CO$_2$ did not increase markedly in the simulations described above, a more detailed analysis of the changes in vegetation and terrestrial carbon yields some surprising finds. Terrestrial carbon (Fig. 1c) increased just over 200 Pg C in all FC simulations, although Fig. 1d demonstrates that vegetation carbon (above-ground biomass) actually decreased in many
runs from the LGM to the early Holocene (11000 BP). The net terrestrial increase after 15000 BP was largely driven by a global increase in soil carbon as ice sheets retreated (Fig. 2c,d). The simulations with CO$_2$ radiative forcing (Fig. 1d, red and orange lines) produced a strong decline in vegetation carbon from the LGM through the early Holocene, in spite of vegetation expansion over formerly ice-covered areas in all of these simulations.

The changes in the spatial distribution of vegetation carbon stocks (Fig. 2a,b) provides additional insight into terrestrial carbon changes. From Figs. 2a,b, we note that terrestrial vegetation gains in mid and high-latitude regions were unable to outweigh larger losses in the tropics (until, at least, the end of the simulation in this FC LW case). The disappearance of ice sheets and related warming slowly destabilized some tropical and subtropical ecosystems. This is particularly true where subtropical boreal forests were replaced by grasslands (such as the present-day southeastern United States, Fig. 2a), resulting in new PFT distributions that accumulated biomass more slowly due to the low atmospheric CO$_2$ in these simulations. Even by the end of the HW simulation (Fig. 2b,d), the model only yielded substantial vegetation and soil carbon gains in areas covered by ice sheets during the LGM; other parts of the world demonstrated equivalent or slightly reduced carbon storage. It should be noted that the UVic’s wind feedback is unable to generate a significantly-enhanced African monsoon vegetation (Fig. 2a); thus, the model likely overemphasized the vegetation loss in the Sahel region in the early Holocene. However, the net effect on vegetation carbon should be similar, as most of the African vegetation gains in the mid Holocene were likely lost by the late Holocene (Indermülhe et al. 1999). Other modeling results also suggest that African forests were replaced more readily by low-density forests and shrublands (Fig. 2a) under low glacial CO$_2$ conditions (Jolly and Haxeltine 1997; Street-Perrott et al. 1997). The sensitivity of subtropical and tropical vegetation
to less CO₂ fertilization thus appears to have had a significant impact on vegetation and terrestrial carbon losses in these simulations, implying that the greater CO₂ fertilization over the course of deglaciation was instrumental in maintaining high biomass density in the tropics.

The addition of CO₂ radiative forcing to the constant alkalinity experiment led to less vegetation carbon storage than some of the other FC simulations (Fig. 1d), indicating that the global temperature rise associated with both retreating ice sheets and increasing CO₂ did not lead to more vegetation carbon storage. The PC simulation (Fig. 3) provides an important comparison, as it injected carbon into the model’s carbon reservoirs to force the increase in CO₂, but with radiative forcing equivalent to the CO2rad experiment. The physical presence of more carbon in the atmosphere caused substantial net increases in vegetation carbon stocks (by ~230 Pg C, see Fig. 3b), without the decrease in vegetation carbon produced between the LGM and the early Holocene in the FC simulations (Fig. 1d). The total increase in terrestrial carbon of ~600 Pg C in the PC simulations (Fig. 3a) from the LGM to the late Holocene is also comparable in magnitude to that seen in other modeling studies (for example, 550-694 Pg C in Prentice et al. (2011)). The effect of CO₂ fertilization turned net terrestrial carbon losses in the tropics in the FC experiments (Fig 2c,d) into carbon gains (Fig 3e,f) and also enhanced carbon storage in subpolar regions.

A secondary effect of the denser and more expansive vegetation resulting from the CO₂ fertilization effect led to greater absorption of incoming solar radiation, as seen in comparing the divergence in surface air temperature between the CO2rad HW and the PC HW simulations in Supplementary Fig. 1g (both of which have equivalent CO₂ radiative forcing). The biogeophysical impact of greater CO₂ fertilization in the PC HW simulation increased SAT globally by ~0.14°C (Supplementary Fig. 1g) and contributed to centennial-scale ocean.
ventilations that lead to a higher depth-integrated ocean potential temperature (0.12°C) (Supplementary Fig. 1h). The denser tropical vegetation, particularly in South America, Africa, and Southeast Asia (as shown in Fig. 3d), appears to drive much of the fertilization-related increase in global temperature (up to 0.3°C locally, Fig. 4a), as these regions experience a substantial loss in vegetation density without CO₂ fertilization. Fig. 4b shows that the temperature difference between the PC CA and CO₂rad CA has significant variability in the high latitudes (related to different centennial-scale fluctuations in sea ice and the meridional overturning circulation (MOC)) and an overall zonal-average tropical warming of 0.07-0.2°C. Furthermore, because the tropics are important oceanic upwelling/outgassing regions, the additional warming of the tropics due to the fertilization effect may have a disproportionate influence on the outgassing of CO₂. The model’s wind feedback also suggests that this warm anomaly driven by CO₂ fertilization increases zonal wind speeds in the subtropics (not shown), which adds to the wind stress and dynamical upwelling off the west coasts of South America and Africa (not shown). A deglacial strengthening of the Hadley Cell after the LGM is also suggested by the proxy record (Thompson et al. 1998) and could contribute to poleward shifts in the mid-latitude westerlies (Lamy et al. 2001; Toggweiler et al. 2006).

In summary, the FC simulations (including CO₂ radiative forcing) suggest that, without the full CO₂ fertilization associated with the observed ice core CO₂ trend, modeled vegetation carbon stocks would have been lower following deglaciation than they were during the LGM (contrary to proxy evidence). Total terrestrial carbon did increase (200-250 Pg C) in all free carbon simulations starting at the Pleistocene-Holocene transition, but this feature was largely driven by greater soil carbon storage at high latitudes. The only simulations that support a substantial increase in the photosynthetically-active carbon pool on the scale suggested in Ciais
et al. (2012) are the prescribed carbon dioxide (PC) simulations (Fig. 3a). Thus, the results presented here strongly suggest that the carbon sequestration in response to the deglacial rise in CO$_2$ is the most important factor in increasing vegetation biomass between the LGM and present, with vegetation carbon gains at the end of deglaciation and into the Holocene driven by greater CO$_2$ fertilization instead of ice sheet or temperature changes; without a major increase in atmospheric carbon content, vegetation actually decreased during deglaciation and the early Holocene in the UVic model.

3.3 Physical and Dynamical Ocean Changes

Over the course of the simulations presented here, the oceans gradually become a net source of atmospheric CO$_2$. The most significant oceanic release of carbon to the atmosphere resulted from the greater warming in the CO2rad simulations. For example, the CO2rad CA scenario yielded 20 ppm more atmospheric CO$_2$ than the FC CA simulation by the mid-Holocene. A comparison of these two simulations reveals that there was ~50 Pg C less terrestrial carbon during deglaciation in the simulation with CO$_2$ radiative forcing (Fig. 1c, red line vs. blue line), and only a small fraction of this was absorbed by the oceans (Fig. 1b). The decreasing ability for the oceans to take up atmospheric CO$_2$ (greater outgassing than uptake) was modulated by warming ocean temperatures (Fig. 5b) by over 2°C in the CO2rad CA simulation versus only 0.5°C in the FC CA simulation. This warming was most pronounced at depth, as surface waters vary slightly less over the course of these simulations (1.5-1.6°C for the CO2rad CA and 0.4-0.5°C for the FC CA simulation) (Supplementary Figure 2). Solubility considerations (~9 ppm per 1°C warming) mandate that the net effect should be ~9-11 ppm of the ~20 ppm difference in atmospheric CO$_2$ between the FC and CO2rad simulations. These ocean properties do not incorporate the ice melting/ocean freshening effect favouring greater
**CO₂ uptake during deglaciation, as global salinity only decreased by 0.002 ppt as a result of precipitation changes in these experiments.**

Complementing the CO₂ solubility effect, changes in ocean ventilation also contribute to the higher CO₂ levels in the CO2rad simulations. Fig. 5c shows the global maximum overturning streamfunction, which is a general indicator of the strength of North Atlantic Overturning circulation and NADW formation. The PC (Supplementary Fig. 1e) and CO2rad simulations equivalently produced a two-stage increase in the meridional overturning circulation (MOC) strength from LGM values (~15 Sv) to modern values (~21 Sv) by 15500 BP. While these simulations lack of freshwater fluxes from retreating ice sheets, the strengthening NADW reflects similar timing as the North Atlantic MOC rebound at the end of Heinrich Event 1 (HE1) around 14500 BP as demonstrated by Pa/Th proxy data (McManus et al. 2004). These modeling results suggest that higher CO₂ radiative forcing may have helped accelerate the North Atlantic MOC rebound after HE1. The unforced FC experiments also produced a somewhat stronger (17-18 Sv) meridional overturning circulation around the same time period, but most of these simulations lagged significantly behind the CO2rad MOC recovery, and none of them achieved present-day overturning values. However, all the CO2rad simulations followed the MOC profile from the PC simulation (Supplementary Fig. 1e), indicating that the strength of the MOC rebound to the model’s typical modern values was mostly a product of the temperature increase due to the radiative forcing effect of CO₂.

The divergence in the Δ¹⁴C trend lines between the FC and CO2rad simulations (Fig. 5b), which lagged the divergence in air temperature between the FC and CO2rad simulations by approximately 1000 years, provides further evidence of a link between CO₂ and ventilation changes. The initial increase in CO₂ (Fig. 1a) stimulated greater ventilation of deep water from
the glacial period, and the ventilation (and warming) of CO₂-rich deep waters subsequently led to
less oceanic absorption of atmospheric CO₂ and eventually a net release (achieved by 9000 BP
according to Fig. 1b) from the ocean to the atmosphere from this effect. Fig. 6 illustrates the
deep-ocean DIC change for the CO₂rad HW simulation at 14500 BP, the approximate timing of
post-HE1 recovery and ventilation of the North Pacific (Galbraith et al. 2007), and at 9000 BP,
by which time the NADW transport essentially reached its peak and plateaued (McManus et al.
2004). The simulations correspondingly demonstrate a gradual decrease in the DIC content of the
Atlantic Ocean relative to the LGM state, with the NADW becoming more prominent and
promoting the decrease of DIC in the Indian and South Pacific with time (compare Fig. 6a to Fig.
6b). Related to this ventilation change, proxy evidence of low Δ¹⁴C excursions during the
Mystery Interval (~17500 BP to ~14500 BP) indicate an important ventilation (or series of
flushes) of the glacial deep ocean during the late Pleistocene. What is new here is that CO₂
warming appears to enhance this ventilation.

The model results thus reveal a potential positive feedback relevant to the deglacial rise
in CO₂, in which an initial increase in temperature due to rising CO₂ stimulated a greater
ventilation of carbon-rich deep waters, which helped produce an even larger increase in CO₂.
One potential fast-response mechanism in this feedback loop is sea ice, which is strongly
sensitive to warming temperatures and influences overturning. As air-sea gas exchanges are not
possible through sea ice in the UVic model, much greater annual sea ice extent in both the
northern and southern hemispheres (Fig. 5e-f) could limit convection in both regions (Fig 7).
Even without freshwater fluxes, sea ice extent in the North Atlantic was more expansive in the
FC simulations after the LGM compared to the CO₂rad simulations (Fig 5e), which limited the
strength of North Atlantic overturning and deep-water convection in upwelling regions. The FC
CA simulation in particular demonstrated less outgassing in tropical upwelling regions at 9000 BP (Fig. 7e), whereas the CO2rad CA simulation showed greater upwelling in the tropics (Fig. 7b,e). Similarly, more sea ice in the Southern Ocean in the FC simulations limited the air-sea exchange of upwelling water masses near Antarctica (Fig. 7e-f).

Fig. 7 also shows that downwelling regions in the North Atlantic (Fig. 7a-b) shifted northeastward over the course of deglaciation in concert with reduced sea ice extent (Fig. 7c-d), whereas upwelling regions (carbon sources to the atmosphere) intensified in the Southern Ocean along retreating sea ice margins, particularly in the CO2rad experiments. Proxy evidence also suggests that this region was likely critical during the upwelling of glacial deep water (Anderson et al. 2009). During the late Holocene, abrupt increase in SH sea ice after 6000 BP in most simulations (Fig. 7f) is a response to a rapid decrease in Antarctic marine ice shelves in the database (described further in Simmons et al. (2013)). This transition was followed by a reduction in atmospheric CO2 associated with a weaker NADW, slowly-ventilating AABW and reduced upwelling (more sea ice) in the Southern Ocean.

In summary, these model results indicate that warming temperatures due to increasing CO2 may have contributed to a decrease in annual-mean sea ice in both hemispheres, which coincided with a faster overturning of model deep waters through the Southern Ocean and stimulated a further rise in CO2. These findings suggest that the Southern Ocean sea ice mechanism described in Stephens and Keeling (2000) may have had some influence in enhancing an ongoing release of carbon from the deep ocean to the atmosphere, especially considering that winter sea ice extent at the LGM was likely much greater than at present (Gersonde et al. 2005; Roche et al. 2012). After accounting for reduced gas solubility with warming SSTs, the sensitivity of atmospheric CO2 to Southern Ocean sea ice area seems similar.
in magnitude to the 6.2 ppm obtained in the AOGCM study of Chikamoto et al. (2012). Fletcher et al. (2007) showed that most OGCMs in OCMIP-2 underestimate both the present-day outgassing of CO$_2$ in the high-latitude Southern Ocean by as much as 0.4 Pg C yr$^{-1}$ and the CO$_2$ uptake in the mid-latitude Southern Ocean by -0.2 Pg C yr$^{-1}$. Therefore, the true effect of sea ice changes in this region may have been larger than that modeled here. In particular, Brovkin et al. (2012) cited a 20 ppm sensitivity to SH sea ice after including increased diffusivity with their zonally-averaged ocean model.

**3.4 Alkalinity response to ocean ventilation**

Through a more vigorous ventilation of deep water, the CO$_2$ radiative forcing promoted a shift in the total DIC content (Fig. 6, discussed above) and alkalinity in the deep ocean (Fig. 8). These results are consistent with Galbraith et al. (2007), who documented a thorough ventilation of glacial deep waters in the North Pacific around 14500 BP, which coincided with a 10 ppm increase in atmospheric CO$_2$ in the ice core record. However, the replacement of glacial deep waters containing high levels of respired DIC with surface waters that have relatively low DIC and higher [CO$_3^{2-}$] would have contributed further to a long term increase in atmospheric CO$_2$ by supporting more CaCO$_3$ sedimentation, thereby decreasing net ocean alkalinity (Galbraith et al. 2007).

While alkalinity-induced changes in the $p$CO$_2$ of the oceans are not possible in the CA transient setup (by definition), the differences between the FC HW and CO2rad HW experiments help quantify this effect. The only distinction between these two simulations was the radiative forcing of CO$_2$ imposed on the CO2rad HW experiment. The FC HW run produced a slight increase in alkalinity between the LGM and the Holocene; carbonate compensation in favour of
sediment burial was unable to outpace the weathering (alkalinity) inputs into the ocean until the late Holocene, contributing to the lower atmospheric CO$_2$ (~180-182 ppm) toward the end of the simulation. The CO2rad HW experiment showed the opposite trend, with a significant net decrease in alkalinity (40 μmol kg$^{-1}$, see Fig. 8a) and a pronounced increase in CO$_2$ to 215 ppm during the Holocene (Fig. 1a), outpacing the CO2rad CA simulation by 10 ppm by the mid-Holocene. This is aided by both a decrease in DIC content of the deep oceans due to greater ventilation (Fig. 6a-b) accompanied by a larger increase in precipitated calcite (Fig. 8c).

The alkalinity of the CO2rad HW and FC HW simulations began to diverge between 17000 BP and 16000 BP (Fig. 8a, purple line vs. orange line), coincident with the MOC recovery in the CO2rad simulations and reduced DIC storage in the deep Atlantic Ocean. This AMOC-induced change in alkalinity is portrayed by the average column depth-integrated alkalinity differences in Fig. 9a-b, which shows Atlantic waters losing alkalinity with time (consistent with the evolution discussed in Yu et al. 2014). Proxies tend to relate changes in alkalinity more specifically to [CO$_3^{2-}$] (for example, Catubig et al. (1998); Yu et al. (2014); Yu et al. (2010); Rickaby et al. (2010), and references therein), and subtracting alkalinity (Alk) from DIC in the model’s spatial output allows a rough approximation of this quantity (Zeebe and Wolf-Gladrow 2001). By 9000 BP, the CO2rad HW simulation provides Δ[Alk-DIC] changes since the LGM below 2990 m of +26-42 μmol kg$^{-1}$ in the North Atlantic (above current estimates), +5-10 μmol kg$^{-1}$ in most of the Pacific (comparable to Yu et al. (2010)), little net change in the Southern Ocean, and -10 μmol kg$^{-1}$ in the Weddell Sea (below current estimates). A decrease in [Alk-DIC] in intermediate waters was also produced in the Atlantic (not shown), as in Yu et al. (2010).
However, Fig. 9 shows that $[\text{CO}_3^{2-}]$ during deglaciation (Fig. 9c) and the early interglacial (Fig. 9d) was also greater in the CO2rad HW simulation compared to the neo-glacial FC HW simulation in the Atlantic, with much smaller changes in the Pacific. The contrast between the CO2rad HW and FC HW in the North Atlantic (+20 to +25 μmol kg$^{-1}$) are similar to the +20 μmol kg$^{-1}$ LGM-to-Holocene change given in Yu et al. (2010), and the simulated change (-24 μmol kg$^{-1}$) in the deep Weddell Sea is comparable to that given in Rickaby et al. (2010) (-25 μmol kg$^{-1}$). Although the proxy record of glacial-interglacial changes in $[\text{CO}_3^{2-}]$ at sufficient resolution is highly limited and prevents firm conclusions, the spatial distribution in Fig. 10c-d agrees qualitatively with some evidence available to date. In other words, the CO$_2$-ventilation feedback causes changes in the alkalinity and DIC in the model ocean that match the sign and magnitude of the proxy-derived changes in $[\text{CO}_3^{2-}]$ in the deep Atlantic and Weddell Sea (Yu et al. 2010; Rickaby et al. 2010).

In the simulations considered here, however, deep ocean DIC changes were more pronounced in the Atlantic and Southern Indian than they are in the Pacific, which remains relatively poorly-ventilated until the mid-Holocene. This is in contrast to proxy evidence in Galbraith et al. (2007), which documented an important deglacial ventilation in the North Pacific starting around 14500 BP. As a consequence, the earlier real-world ventilation of Pacific DIC during the Bølling-Allerød would have led to an even greater long-term decrease in alkalinity than modelled here through enhanced sedimentation in the Pacific basin following the replacement of deep waters (Galbraith et al. 2007). The model did, in fact, slowly increase the circulation in CO2rad HW until a renewal of deep waters in the North Pacific was completed abruptly between 8000 BP and 6000 BP in a manner similar to that illustrated in Simmons et al. (2013). This Holocene renewal of deep Pacific waters also stimulated a further reduction in
ocean alkalinity (Fig. 8a). The distribution of Antarctic marine ice shelves (the interpolated ICE-14G database) influenced the timing of the ventilation of North Pacific deep waters (see Simmons et al. 2013); an earlier ice shelf transition might force a more accurate North Pacific flush of DIC, which would in turn permit an earlier alkalinity-induced increase in atmospheric CO$_2$ during the Holocene. The ice configuration around Antarctica merits further study in this regard in order to determine whether there was a more abrupt and earlier ventilation of the North Pacific as proxy evidence suggests (Galbraith et al. 2007).

3.5 Sensitivity to Weathering and Carbonate Compensation

Another important factor affecting ocean alkalinity is the chemical weathering rate. Despite the model’s inability to produce a deglacial increase in atmospheric CO$_2$ without CO$_2$ radiative forcing, it demonstrated significant sensitivity to the choice of weathering rate. The difference in weathering rate explains the approximately 10-15 ppm difference between the FC LW and FC HW simulations. Fig. 8a shows that the lower weathering rate is associated with a lowering of mean alkalinity (by -33 μmol kg$^{-1}$) and the higher weathering rate results in a net increase in alkalinity (by +10 μmol kg$^{-1}$). These findings may seem rather intuitive, but they demonstrate the relative importance of carbonate compensation, weathering, sedimentation flux, ocean ventilation, and deglacial atmospheric CO$_2$ changes.

A closer look at the FC LW simulation reveals that the sedimentation rate increased with time and exceeded the weathering rate for the entire simulation after 19000 BP (Fig. 8c, green line vs. light grey line), associated with an increase in calcifiers. Because sedimentation (an alkalinity sink) surpassed weathering (alkalinity source) for virtually the entire simulation, the greater removal of carbonate and bicarbonate in the surface ocean led to a downwelling of waters.
that have low [CO$_3^{2-}$], supporting a shallower lysocline and greater (oxic) dissolution of sediments and calcite rain. This is reflected in Fig. 8c, which shows that the FC LW sediment flux dipped below the FC CA sediment flux after 15000 BP.

The limited impact of carbonate compensation for the lower weathering rate is illustrated in Fig. 8d by changes in the CCD (defined here as the level where the rate of sedimentation equals the rate of dissolution). While the CCD was shallowest in the FC LW experiment compared to other runs, it still deepened during ventilations and did not rise above the original depth until 5500 BP. The rapid shoaling in the late Holocene appears to be aided by the increasing prevalence of more corrosive, high DIC southern-sourced deep waters after the Antarctic marine ice shelf disappearance (6000 BP-5000 BP) (Simmons et al. 2013). However, before this transition, a comparison of Fig. 8d with Fig. 5c reveals that the CCD deepening in the simulation was stimulated by greater ventilation (i.e., a stronger alkalinity pump to the deep ocean via the NADW). Although the partial recovery of the NADW driven by land ice and orbital forcing is much more limited without the addition of CO$_2$ radiative forcing, the somewhat more intense downwelling of low-DIC, high-alkalinity North Atlantic surface waters appears to be sufficient to support a deepening of the CCD. Without these changes in the MOC, a CCD shoaling would be supported throughout the simulation based on carbonate compensation.

Therefore, a combination of low weathering (low alkalinity input), more calcite rain, and greater ventilation (which flushes deep DIC and lowers the CCD) helped drive down ocean alkalinity and supported a long-term increase in atmospheric CO$_2$.

The FC HW simulation shows a different pattern of carbonate compensation; from the beginning of the simulation to 11000 BP, the net sediment flux (Fig. 8c, purple line) remained below the weathering rate (Fig. 8c, dark purple line), and the greatest deepening of the CCD
occurred during this time (Fig. 8d). In this case, the more intense ventilation after 16000 BP (Fig. 5c) and carbonate compensation (Fig. 8c) together favored a deepening of the CCD up to the beginning of the Holocene. This allowed for more sediment preservation and caused the sediment flux to drift above that of the FC CA simulation (Fig. 8b,c, blue line vs. purple line). Then after 11000 BP (approximately the beginning of the Holocene), the calcite flux surpassed the weathering rate, resulting in increasing dissolution with time by the mid-Holocene (in Fig. 8c, the purple line re-approaches blue line), although the long-term shoaling of the CCD associated with this effect (after 7000 BP) was nearly negligible (Fig. 8d). The deepening of the CCD accompanied by a larger sediment flux contributed to the decrease in alkalinity during the Holocene in this simulation, although this change is not substantial enough to counter the increase in ocean alkalinity during the late Pleistocene associated with a greater alkalinity source (weathering) than sink (sedimentation).

By contrast, when CO$_2$ radiative forcing was added to HW experiment, the sediment flux surpassed the weathering flux much earlier (16500 BP) and exceeded it by a much greater quantity (57 Tg C yr$^{-1}$) than in the simulation without radiative forcing (17 Tg C yr$^{-1}$). Correspondingly, the greater dissolution supported after 16500 BP drove the sediment flux significantly below the constant alkalinity experiment (CO2rad CA) after 11000 BP (Fig. 8c, light orange line vs. red line). Carbonate compensation in the CO2rad HW simulation should, in turn, support a more pronounced shoaling of the CCD than in the FC LW simulation (in which the sedimentation flux only surpassed weathering by a maximum of 29 Tg C yr$^{-1}$). However, the CCD in the CO2rad HW experiment remained below the original LGM level until 5000 BP and did not shoal above the FC LW CCD. The implication is that the greater ventilation stimulated by the CO$_2$-ventilation feedback, favouring the replacement of low-alkalinity, high-DIC deep
waters with higher-alkalinity, lower-DIC surface waters, dominated the evolution of ocean alkalinity for most of the simulation.

These results are generally consistent with the proxy data presented in Yu et al. (2010), which suggest that abyssal oceans demonstrated a decrease in deep-ocean DIC content and increase in \([\text{CO}_2^{2-}]\) into the early Holocene in the deep Equatorial Pacific and North Atlantic in response to a ventilation of the deep ocean. Moreover, they support the conjecture in Rickaby et al. (2010) that, for a lower weathering rate for the LGM, an increase in sedimentation flux (due to a low CCD) can by itself work to decrease alkalinity and initiate an increase in \(\text{CO}_2\). Our modeling results also suggest, however, that the same net effect (greater abyssal alkalinity and a deeper CCD) can be obtained from more ventilation of the deep ocean for both weathering rates, including in cases where carbonate compensation opposes the ventilation-induced alkalinity change. Further more, proxy records (Foster and Vance 2007; Vance et al. 2009; Crocket et al. 2012; Crocket et al. 2013) suggest that increased weathering flux into the North Atlantic coincides with periods of deeper overturning in the same region, which could accelerate deep sedimentation in NADW-dominated deep ocean basins more than modelled in the current study.

In summary, weathering dominated ocean alkalinity changes in our simulations, except in simulations with stronger, early deglacial deep-ocean ventilations. Greater ocean ventilation, in turn, had a more immediate impact on alkalinity than carbonate compensation, except for very long time scales (~15000-20000 years). Greater ocean alkalinity in the HW simulation drove \(\text{CO}_2\) (180-182 ppm) slightly below the CA simulation (183 ppm), whereas the relatively higher end-result \(\text{CO}_2\) (195 ppm) obtained in the LW simulation was driven by the reduction in ocean alkalinity due to lower alkalinity inputs. Only the larger ventilation change in the CO2rad HW (with intensified calcite sedimentation) led to an earlier downwelling of high-alkalinity surface
waters, which allowed greater deep sedimentation and a decrease in alkalinity. Given the enhanced ventilation of the CO$_2$ radiative forcing simulation, weathering would have to increase markedly for the oceans to be able to gain alkalinity as in the HW simulation.

### 3.6 The Alkalinity Response to Holocene Terrestrial Uptake

Broecker et al. (1999) proposed that terrestrial uptake during the early Holocene (boreal forests and expanded monsoonal vegetation) contributed to a deglacial-interglacial alkalinity change by sequestering CO$_2$ from the atmosphere and surface ocean, thus increasing the sea surface concentration of carbonate relative to dissolved CO$_2$. The downwelling of these low $p$CO$_2$, high [CO$_3^{2-}$] waters in the mid-Holocene would have deepened the CCD, allowing more sedimentation of falling CaCO$_3$ shells and ultimately reducing the [CO$_3^{2-}$] of the ocean. Broecker et al. (1999) documented the removal of alkalinity through the decrease in [CO$_3^{2-}$] from the size distribution of CaCO$_3$ shells in western tropical Atlantic and western tropical Pacific cores, which implied a late Holocene 11±2 μmol kg$^{-1}$ decrease in [CO$_3^{2-}$] and over 20 ppm atmospheric CO$_2$ increase due to greater sedimentation during the early and mid-Holocene. In subsequent modelling studies, Joos et al. (2004) attributed 4-11 ppm of the Holocene CO$_2$ increase to an alkalinity decline in response to a sediment preservation event; Kleinen et al. (2010) argued that there was no significant difference due to this effect from sensitivity simulations including and excluding terrestrial changes, whereas Menviel and Joos (2012) attributed about +5.3 ppm in response to terrestrial uptake during the Holocene.

Of the results presented in this study, the PC HW simulation (Fig. 10a-b) provides the best illustration of this effect, as it imposes a decrease in atmospheric CO$_2$ (Supplementary Fig. 1a) between the early and mid-Holocene, as documented in the ice core record in Petit et al. (1999). This caused 125 Pg C to be extracted from the oceans (Supplementary Fig. 1b).
However, the resulting total ocean alkalinity decrease of 4 μmol kg$^{-1}$ (Fig. 10a, pink line) could not account for the pace of the CO$_2$ increase after the mid-Holocene, requiring continued injections of carbon into the atmosphere in order to reproduce the late Holocene rise in CO$_2$. Fig. 10c-d show the change in the alkalinity and [Alk-DIC] patterns for the PC HW simulation between 7000 B.C and 5600 BP, just prior to the Antarctic ice shelf transition (occurring in the model at 5500 BP) and before renewed acidification from external CO$_2$ sources (Supplementary Fig. 1b). It shows the greatest alkalinity losses are in the Pacific Ocean (Fig. 12a), where deep Δ[Alk-DIC] is relatively high (Fig.12b), whereas a low [Alk-DIC] anomaly drove down deep ocean values in the Atlantic. The carbonate losses in the Atlantic had already begun by this date (-5 to -10 μmol kg$^{-1}$), whereas in the Pacific they had not yet been achieved (+5 μmol kg$^{-1}$). By 3000 BP (not shown), the low carbonate anomaly began to move to the Equatorial Pacific (-5 μmol kg$^{-1}$), whereas the Atlantic anomaly (-25 μmol kg$^{-1}$) was overestimated (due in part to the more prominent AABW after the aforementioned ice shelf transition).

Another approach to evaluating terrestrial-alkalinity interactions as proposed by Broecker et al. (1999) is to free the carbon cycle in the PC simulations after 8000 BP (i.e, following the extraction of DIC from the ocean). We did this for both the constant alkalinity experiment (PC CA, Fig. 11, dark blue line) and the higher weathering rate (PC HW, Fig. 11a, dark purple line) to isolate the effect of the alkalinity change on atmospheric CO$_2$. Fig. 11a shows that there is virtually no difference in the end-result atmospheric CO$_2$ concentrations for these two simulations (251.3 ppm). Furthermore, the alkalinity change following 6000 BP. in the freed PC HW simulation (Fig. 11b, dark purple line) is small (5 μmol kg$^{-1}$), becoming roughly constant after 6000 BP. Therefore, it appears that the greater downwelling of more corrosive Southern Ocean-generated waters in these simulations after 6000 BP exactly counterbalanced the slowly
decreasing alkalinity in response to DIC removal between 10000 BP and 8000 BP, leading to unchanged alkalinity in the freed PC HW simulation. Redoing these two simulations (freed PC CA and freed PC HW) with more extensive ice shelves (Fig. 11b, light blue and light purple line respectively) as in Simmons et al. (2013) led to less AABW formation and a more prominent NADW. As Fig. 11b reveals, the more prominent NADW caused alkalinity to continue decreasing during the late Holocene in the PC HW simulation, allowing atmospheric CO$_2$ to be 10 ppm higher than in the PC HW simulation without ice shelves and 7 ppm higher than the constant alkalinity experiment with the same more-extensive ice shelves. The significantly greater atmospheric CO$_2$ concentrations in the PC HW simulation with Antarctic marine ice shelves suggests that a dominant NADW is required in order for ocean alkalinity to continue decreasing in response to the early Holocene terrestrial uptake. The mechanism proposed by Broecker et al. (1999) thus appears to be strongly dependent on the ocean circulation state and the proportion of AABW to NADW at depth in the model.

Results from the FC simulations also suggest a limited impact of the terrestrial uptake on ocean alkalinity without the inclusion of more extensive Antarctic ice shelves. Ciais et al. (2012) scaled back the terrestrial uptake to ~300 Pg C between the LGM to the pre-industrial Holocene, which is only slightly greater than the ~200-250 Pg C terrestrial uptake between the early and late Holocene in our FC and CO2rad simulations. The FC HW simulation, which experienced a terrestrial uptake of ~130 Pg C between 14500 BP and 8000 BP (Fig. 1c), induced a slight decrease in alkalinity into the later Holocene (Fig. 8a, purple line), although a significant fraction of this signal seems to be associated with more ventilation (Fig. 5c) and calcite rain (Fig. 8c) during this time period. While terrestrial uptake in our simulations has a limited impact on interglacial alkalinity, the CO2rad results do provide for a substantial alkalinity-induced increase
in atmospheric CO$_2$ during the Holocene in the CO2rad HW simulation. Furthermore, this increase occurred in spite of decreased vegetation carbon storage between the LGM and the early Holocene. In particular, the terrestrial carbon storage in the CO2rad HW scenario (with higher CO$_2$) only began to (slowly) surpass the FC HW simulation at ~9500 BP and was only 20 Pg C larger than the FC HW simulation at 8000 BP, at which point (comparing the CO2rad CA and CO2rad HW trends in Fig. 1a) the alkalinity-induced increase in atmospheric CO$_2$ in the CO2rad HW simulation was already well underway. A general decrease in [CO$_3^{2-}$] over the course of the late Holocene was also obtained in this experiment (Fig. 12a,b), again perhaps somewhat overestimated due to the influence of the Antarctic ice shelf transition on AABW but also on the scale of the changes given in Broecker et al. (1999). Thus, in our results, the effects of the greater ocean ventilation of respired deep-ocean DIC (the CO$_2$-ventilation feedback) appears to be the more prominent factor in promoting a decrease in ocean alkalinity.

4 Conclusions

In this paper, we presented a suite of simulations using the UVic ESCM v. 2.9 to test the model’s transient response to land ice changes, orbital parameter changes, CO$_2$ radiative forcing, and alkalinity inputs (from different constant weathering rates) from the LGM to the present. In the simulations without high latitude terrestrial carbon storage, we focused on important oceanic processes. These simulations failed to reproduce the full deglacial rise in atmospheric CO$_2$. Depending on the processes included, we simulated both a decline in atmospheric CO2 of ~10 ppm to an increase of up to 25 ppm. Modeling experiments with only Milankovitch and ice sheet forcing did not produce a full recovery in the Atlantic meridional overturning circulation, leading to higher oceanic carbon storage and the lowest values of atmospheric CO$_2$ (unchanged or lower than the LGM value, depending on the weathering rate). Those modeling experiments that
included CO2 radiative forcing produced a warmer ocean, a more thorough ventilation of the deep ocean. This suggests that the warming effect provided by higher atmospheric CO2 acts as a positive feedback on the deglacial rise in CO2.

This CO2-ventilation feedback occurred in the model in response to stronger overturning circulation in the North Atlantic and greater outgassing in the Southern Ocean, both enhanced by reduced sea ice extent. This ventilation of deep-ocean DIC allowed atmospheric CO2 to increase by ~10 ppm. The alkalinity decrease in the ocean in response to greater ventilation contributed an additional 10 ppm in our simulations, mostly over the course of the late deglacial and interglacial periods due to the downwelling of high-alkalinity, low-DIC waters in the North Atlantic. These circulation features are largely driven by a reorganization of the Atlantic and Indian oceans, where the NADW has more influence in the model. In terms of alkalinity changes, a comparison with simulations without CO2 radiative forcing reveals that the CO2-ventilation feedback in the model may by itself potentially explain deep ocean variations in [CO3^2-] on the order of those given in Yu et al. (2010) and Rickaby et al. (2010) in the deep Atlantic and Weddell Sea.

The ocean alkalinity and atmospheric CO2 during deglaciation are also strongly influenced by weathering. When only ice sheets and Milankovitch forcing are prescribed, the total ocean alkalinity was largely driven by the weathering rate, with higher (lower) weathering rates leading to greater (less) ocean alkalinity and lower (higher) atmospheric CO2. Only the thorough, early ventilation of deep-ocean DIC that occurred in the CO2 radiative forcing experiments were able to produce a net lowering of total ocean alkalinity under higher weathering rates. This alkalinity decrease occurred in response to a lower CCD in the Atlantic, Indian, and South Pacific basins in response to stronger NADW formation and overturning. The
results suggest that an early ventilation event is important for stimulating a decrease in alkalinity during the deglacial and interglacial periods to maintain higher atmospheric CO$_2$. However, the constant weathering rates used for this experiment assume an equilibrium between weathering and sedimentation for two different points during the late glacial period (23000 BP and 21000 BP). Rickaby et al. (2010), on the other hand, suggested that the weathering rate may have been substantially lower than the sedimentation rate during the LGM. This would have resulted in an increased atmospheric CO$_2$ during the early deglacial period. While the present set of simulations do not model this differentiation in weathering and sedimentation rates during the LGM and early deglacial, they support the hypothesis that lower weathering rates and greater deep ocean ventilation worked together to allow a net decrease in alkalinity.

In summary, our results suggest that there are potential oceanic contributors to the atmospheric CO$_2$ rise during deglaciation:

(1) Lower weathering rates during the early deglacial maintained a slowly increasing atmospheric CO$_2$ (providing that the sedimentation rate exceeded the weathering rate).

(2) Ventilations of glacial deep water, the reinvigoration of the NADW, and stronger overturning supported an outgassing of respired CO$_2$ from the deep ocean.

(3) The ventilation of high-CO$_2$ deep waters and replacement with surface waters with higher carbonate concentrations allowed alkalinity to continue to decrease (through greater sedimentation) in the late deglacial and early interglacial period, maintaining the initial increases in CO$_2$ even for higher weathering rates.

The three steps together did not account for the full deglacial rise in CO$_2$ in the simulations presented here. This is likely due to the fact that our experimental setup does not account for
several important processes. In particular, during our equilibrium simulation for the LGM, we removed 366 Pg C of terrestrial carbon from the pre-industrial spin-up that would have otherwise been transferred under the ice sheet. Reconfiguring the model to reapportion this carbon to the oceans and terrestrial biosphere (under fixed CO$_2$) during the equilibrium simulation would have increased the DIC content of the oceans and enhanced the ventilation-alkalinity response during deglaciation. Furthermore, the downwelling of CO$_2$ would have increased sediment dissolution and provided an even lower equilibrium weathering rate than the HW and LW rates used here. Huiskamp and Meissner (2012) found that the upwelling of DIC in their simulations for the Mystery Interval could only explain ~58% of the CO$_2$ rise through ocean processes, perhaps also because of the lack of transfer of terrestrial carbon to the deep ocean in these experiments. A weakened biological pump during NADW shutdowns (with less calcite rain and sedimentation as a result) would also lead to an increase in alkalinity, and thus greater oceanic absorption of atmospheric CO$_2$. Our results thus indicate that a combination of a lower weathering rate and higher initial DIC content of the oceans could allow for a larger early deglacial increase in atmospheric CO$_2$ than in the present simulations.

Terrestrial processes also provided important contributions to the simulated deglacial and interglacial carbon cycle. Experiments with only CO$_2$ radiative forcing, ice sheet, and orbital changes produced modest (200-250 Pg C) increases in terrestrial carbon storage from the LGM to the present. Increasing land availability and warmer temperatures were not sufficient mechanisms to stimulate net gains in vegetation carbon, as the lower CO$_2$ levels (and limited fertilization) led to carbon losses in the tropics and subtropics that compensated for greater carbon sequestration at higher latitudes. Simulations with prescribed CO$_2$, in which enough carbon was physically injected into the modeled Earth system to allow CO$_2$ to follow the ice core
trend, produced an increase in terrestrial carbon storage of 600 Pg C, comparable to other recent
modeling estimates (Prentice et al. 2011). Therefore, our simulations suggest that CO₂
fertilization accounts for most of the net vegetation carbon uptake since the LGM. Furthermore,
this fertilization effect (when compared to the low fertilization simulations) provided a
significant increase in temperatures (0.1-0.3°C) in the tropics and may have strengthened
outgassing by warming important upwelling regions and increasing the baroclinic gradient (wind
speeds) and coastal upwelling in the subtropics. The introduction of these temperature anomalies
into a model with a three-dimensional atmosphere would go further in determining the
importance of these fertilization-induced wind anomalies and would be a worthwhile endeavor
for future work.

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Fig. 1: Carbon reservoirs that exchange directly with the atmosphere. These include (a) atmospheric carbon dioxide (ppmv), (b) total ocean carbon (organic and inorganic carbon), (c) total terrestrial carbon (both vegetation and soils), and (d) vegetation (above-ground) carbon. All simulations represented in this figure have a freely-evolving carbon cycle. They include separate transient simulations with constant ocean alkalinity (FC CA, blue line), a higher (12 Tmol yr$^{-1}$) constant weathering rate (FC HW, purple line), and a lower (10.1 Tmol yr$^{-1}$) constant weathering rate (FC LW, green line). These three simulations were repeated with CO$_2$ radiative forcing (denoted by warmer colours): CO2rad CA (red line), CO2rad HW (orange line), and CO2rad LW (tan line).
Fig. 2: The net change in the spatial distribution of above-ground vegetation carbon at (a) 10000 Before Present (BP) and (b) 150 BP, and the net change in total terrestrial carbon (vegetation and soils) at (c) 10000 BP and (d) 150 BP since the Last Glacial Maximum (LGM) climate (defined as 20900 BP) for the FC LW simulation. Vegetation carbon anomalies are plotted on a scale of -800 Tg C to +800 Tg C, whereas terrestrial carbon anomalies are plotted from -1250 Tg C to +1250 Tg C.
Fig. 3: Time series of (a) global terrestrial carbon (vegetation and soils) and (b) above-ground vegetation carbon for a Prescribed Carbon simulation with constant alkalinity (PC CA, black line) and Prescribed Carbon with the higher (12 Tmol yr⁻¹) weathering rate (PC HW, pink line). The FC HW and CO2rad HW results (from Fig. 1) are plotted for comparison. In (c-f), vegetation carbon and total terrestrial carbon changes from the LGM are mapped as in 2(a-d) for the PC CA simulation.
Fig. 4: Surface air temperature differences between the PC CA simulation and the CO2rad CA simulation, both of which have equivalent CO2 radiative forcing. In (a), a time slice is provided for 10000 BP, which corresponds to the terrestrial and vegetation carbon stocks shown in Fig. 3 for the PC CA simulation. In (b), a Hovmöller diagram shows the zonal-average surface air temperature difference between the two simulations with time.
Fig. 5: Time series of (a) global mean surface air temperature (9°C to 15°C), (b) volume-weighted mean ocean potential temperature (2°C to 5°C), (c) global maximum ocean meridional overturning streamfunction value (14 Sv to 25 Sv), (d) volume-weighted mean ocean Δ^{14}C (-200‰ to -150‰) under a constant atmospheric Δ^{14}C of 0‰ for all simulations, (e) mean annual Northern Hemisphere sea ice area (8 - 14×10^6 km^2), and (f) mean annual Southern Hemisphere sea ice area (6 - 18×10^6 km^2). The colours representing each simulation are as in Fig. 1.
Fig. 6: The net change in deep-ocean DIC (below 2990, the deepest 31.2% of the model ocean) in Tg C between the LGM and (a) 14500 BP and (b) 9000 BP in the CO2rad HW simulation, both on a scale from -250 Tg C (blue) to +250 Tg C (red).
Fig. 7: The spatial distribution of the net change in the downward flux of inorganic carbon at the ocean surface between the LGM (-1.5 to +1.5 Tg C yr\(^{-1}\)) and (a) 14500 BP and (b) 9000 BP. Regions with greater outgassing (or less downwelling) than the LGM are shaded in red, whereas areas with greater uptake (or more downwelling) are blue. These are compared to the net change in annual-mean sea ice area since the LGM (-25000 to +25000 km\(^2\)) at (c) 14500 BP and (d) 9000 BP, with area losses relative to the LGM shaded in blue and gains shaded in red. In (e-f), the differences in (e) surface inorganic carbon flux and (f) sea ice area between the FC CA and CO2rad CA simulation at 9000 BP are given. Blue (red) shading represents less (more) outgassing in the FC CA simulation relative to the CO2rad CA simulation in (e) and red indicating greater sea ice area in the FC CA simulation relative to the CO2rad CA simulation at 9000 BP in (f).
Fig. 8: As in Fig. 1, for variables related changes in ocean alkalinity. In (a), changes in total mean ocean alkalinity \((\text{HCO}_3^- + 2\text{[HCO}_3^2^-])\) are given in μmol kg\(^{-1}\) (with a plot range from 2290 to 2390 μmol kg\(^{-1}\)). In (b), the total change in sediment carbon (Pg C) are provided on a different scale (1280 to 1580 Pg C) than in Supplementary Fig. 1d. In (c), weathering fluxes (constant HW and LW rates are represented by a dark grey and light grey line respectively) and sedimentation fluxes (coloured lines) are provided. For the CO2rad CA and FC CA simulations, only the sedimentation rate is plotted, as by definition the weathering rate and sedimentation rates are exactly equal in these simulations. In (d), the carbonate compensation depth (CCD), calculated as the depth where sedimentation flux is equal to the dissolution flux (above which 99-100% of sedimentation occurs in the model) is provided with a scale range from 3950m to 3650 m depth.
Fig. 9: The spatial distribution of alkalinity anomalies (a-b) since the LGM in the CO2rad HW simulation, and Alkalinity-DIC (\(-[\text{CO}_2^\square]\)) differences (c-d) between the CO2rad HW and FC HW simulations. In (a-b), the net change in the total column mean alkalinity (average depth-integrated alkalinity for each grid cell) between the LGM and (a) 14500 BP and (b) 9000 BP is given. In (c-d), the column (depth-integrated mean) Alkalinity-DIC below 2900 m depth in the CO2rad HW simulation is subtracted from that of the FC HW simulation at (c) 14500 BP and (d) 9000 BP.
Fig. 10: The time series for (a) ocean mean alkalinity and (b) weathering and sedimentation fluxes are given, as in Fig. 8a,c. Only the sedimentation flux is plotted for PC CA in (b) because, by definition, weathering rates and sedimentation rates are equivalent in this simulation. For the PC HW simulation, in (c) the mean total column depth-integrated alkalinity change between 9000 BP and 5600 BP is given at each gridcell, whereas in (d) the net change in mean column depth-integrated [Alk-DIC] below 2990 m depth is provided for the same time interval.
Fig. 11: The PC CA and PC HW simulations repeated with a freed carbon cycle after 8000 BP, including different Antarctic marine ice shelf configurations. In the legend, “freed 8000 BP” indicates that no further carbon was injected into the atmosphere (after 8000 BP), whereas “ice shelves” indicates a simulation where marine ice shelves in the interpolated ICE-4G dataset were held fixed after 6000 BP, as in Simmons et al. (2013). Trends in (a) atmospheric CO\textsubscript{2} and (b) ocean alkalinity are portrayed for these simulations. Note that the time scale for both figures is different than in the other time series in this paper.

Fig. 12: The net change between 9000 BP and (a) 5600 BP and (b) 3000 BP in the depth-integrated column (gridcell) average Alk-DIC below 2990 m depth.