# Carbon cycle responses to changes in weathering and the long-term fate of stable carbon isotopes

Aurich Jeltsch-Thömmes<sup>1</sup> and Fortunat  $Joos^1$ 

<sup>1</sup>University of Bern

December 7, 2022

#### Abstract

The causes of the variations in  $CO_2$  of the past million years remain poorly understood. Imbalances between the input of elements from rock weathering and their removal from the atmosphere-ocean-biosphere system to the lithosphere likely contributed to reconstructed changes. We employ the Bern3D Earth system model of intermediate complexity to investigate carbon-climate responses to step-changes in the weathering input of phosphorus, alkalinity, carbon, and carbon isotope ratio ( $\delta^{13}C$ ) in simulations extending up to 600,000 years.

 $CO_2$  and climate approach a new equilibrium within a few ten thousand years, whereas the equilibration lasts several hundred thousand years for  $\delta^{13}C$ . These timescales represent a challenge for the initialization of sediment-enabled models and unintended drifts may be larger than forced signals in simulations of the last glacial-interglacial cycle. Changes in dissolved  $CO_2$  change isotopic fractionation during marine photosynthesis and  $\delta^{13}C$  of organic matter. This mechanism and changes in the organic matter export cause distinct spatio-temporal perturbations in  $\delta^{13}C$  of dissolved inorganic carbon.

A cost-efficient emulator is built with the Bern3D responses and applied in contrasting literature-based weathering histories for the past 800,000 years. Differences between scenarios for carbonate rock weathering reach around a third of the glacial-interglacial CO<sub>2</sub> amplitude, 0.05 exceed reconstructed variations in marine carbonate ion. Plausible input from the decomposition of organic matter on shelves causes variations of up to 10 ppm in CO<sub>2</sub>, 4 mmol m<sup>-3</sup> in CO<sup>2-</sup><sub>3</sub>, and  $0.09\delta^{13}$ C. Our results demonstrate that weathering-burial imbalances are important for past climate variations.

## Carbon cycle responses to changes in weathering and the long-term fate of stable carbon isotopes

1

2

3

## A. Jeltsch-Thömmes and F. Joos

Climate and Environmental Physics, Physics Institute and Oeschger Centre for Climate Change Research,
 University of Bern, Bern, Switzerland

6	Key Points:
7	• Earth system's response to weathering changes is probed in 600,000 year-long sim-
8	ulations with the Bern3D model and a cost-efficient emulator
9	• Equilibration timescales are order 10,000 years for $CO_2$ and 100,000 years for $\delta^{13}C$ ,
10	requiring careful model initialization
11	• Literature-based weathering scenarios for the past 800,000 years reveal significant
12	responses in $CO_2$ , carbonate ion, and $\delta^{13}C$

Corresponding author: A. Jeltsch-Thömmes, aurich.jeltsch-thoemmes@unibe.ch

#### 13 Abstract

The causes of the variations in  $CO_2$  of the past million years remain poorly understood. 14 Imbalances between the input of elements from rock weathering and their removal from 15 the atmosphere-ocean-biosphere system to the lithosphere likely contributed to recon-16 structed changes. We employ the Bern3D Earth system model of intermediate complex-17 ity to investigate carbon-climate responses to step-changes in the weathering input of 18 phosphorus, alkalinity, carbon, and carbon isotope ratio ( $\delta^{13}$ C) in simulations extend-19 ing up to 600,000 years. CO<sub>2</sub> and climate approach a new equilibrium within a few ten 20 thousand years, whereas the equilibration lasts several hundred thousand years for  $\delta^{13}$ C. 21 These timescales represent a challenge for the initialization of sediment-enabled mod-22 els and unintended drifts may be larger than forced signals in simulations of the last glacial-23 interglacial cycle. Changes in dissolved CO<sub>2</sub> change isotopic fractionation during ma-24 rine photosynthesis and  $\delta^{13}$ C of organic matter. This mechanism and changes in the or-25 ganic matter export cause distinct spatio-temporal perturbations in  $\delta^{13}$ C of dissolved 26 inorganic carbon. A cost-efficient emulator is built with the Bern3D responses and ap-27 plied in contrasting literature-based weathering histories for the past 800,000 years. Dif-28 ferences between scenarios for carbonate rock weathering reach around a third of the glacial-29 interglacial CO<sub>2</sub> amplitude, 0.05% for  $\delta^{13}$ C, and exceed reconstructed variations in ma-30 rine carbonate ion. Plausible input from the decomposition of organic matter on shelves 31 causes variations of up to 10 ppm in CO<sub>2</sub>, 4 mmol m<sup>-3</sup> in CO<sub>3</sub><sup>2-</sup>, and 0.09‰ in  $\delta^{13}$ C. 32 Our results demonstrate that weathering-burial imbalances are important for past cli-33 mate variations. 34

#### <sup>35</sup> Plain Language Summary

Data from ice cores and marine sediments document large changes in atmospheric 36 carbon dioxide  $(CO_2)$  and climate during the past million years. Carbon isotopes and 37 other proxies can help to understand underlying processes. In this study, we investigate 38 Earth's response to plausible changes in the input of carbon and other elements from the 39 weathering of rocks or the decomposition of previously accumulated organic matter with 40 the help of a computer model. Results show significant variations in  $CO_2$ , carbon iso-41 topes, marine chemistry, marine biological productivity, and burial fluxes of biogenic par-42 ticles to the lithosphere. The adjustment time to changes in input flux is several ten thou-43 sand years for  $CO_2$  and climate, and several hundred thousand years for carbon isotopes. 44 As it is computationally challenging to simulate such long time periods with complex 45 models, we used our results to build an emulator. Such emulators, representing the re-46 sponses of spatially-resolved and process-based models, are useful for studies address-47 ing Earth's history over many millions of years. Simulating a million years with the em-48 ulator takes seconds, whereas it takes about three months with the complex model. In 49 conclusion, our work highlights the role of weathering fluxes and their possible contri-50 bution to past climate-carbon cycle swings. 51

## 52 1 Introduction

<sup>53</sup> Chemical weathering of rocks and deposits eventually provides a continuous flow <sup>54</sup> of carbon and other elements to the ocean (e.g. Walker et al., 1981; Berner, 1990; Suchet <sup>55</sup> & Probst, 1995; Hartmann et al., 2014; Compton et al., 2000; Lacroix et al., 2021), thereby <sup>56</sup> influencing climate, atmospheric CO<sub>2</sub>, carbon isotopes, biogeochemical cycles, and the <sup>57</sup> oxidative capacity at the Earth's surface on millennial and longer time scales (Kump & <sup>58</sup> Alley, 1994; Hayes & Waldbauer, 2006). However, Earth's system responses to changes <sup>59</sup> in weathering remain uncertain and their representation in Earth system simulations ex-<sup>60</sup> tending over many thousands or even millions of years challenging.

Proxy records show climate and biogeochemical cycles to vary on orbital time scales
 (Lisiecki & Raymo, 2005; Lisiecki, 2014). Over the past 800 thousand years (kyr), ice

core records document past changes in atmospheric CO<sub>2</sub> (e.g., Neftel et al., 1982; Siegen-63 thaler et al., 2005; Lüthi et al., 2008; Marcott et al., 2014; Bereiter et al., 2015) and in 64 its stable carbon isotope ratio,  $\delta^{13}C_{atm}$  (<sup>13</sup>C/<sup>12</sup>C in permil units), (e.g., Lourantou et 65 al., 2010; Schmitt et al., 2012; Schneider et al., 2013; Eggleston et al., 2016). Variations 66 in the greenhouse gas  $CO_2$  strongly contributed to glacial-interglacial climate swings and 67  $\delta^{13}$ C records constrain related biogeochemical processes. Marine records of  $\delta^{13}$ C (e.g., 68 Raymo et al., 1997; Hoogakker et al., 2006; Oliver et al., 2010; Schmittner et al., 2013; 69 Lisiecki, 2014; C. D. Peterson et al., 2014; C. D. Peterson & Lisiecki, 2018), carbonate 70 preservation (e.g., L. C. Peterson & Prell, 1985), and other proxies reach further back 71 in time and cover the Pleistocene epoch, although with partly sparse spatial coverage 72 and lower temporal resolution. 73

Weathering fluxes enter the ocean through riverine transport in the form of dis-74 solved inorganic and organic compounds and particulate organic matter (Meybeck, 1982; 75 Regnier et al., 2013; Tréguer et al., 2021; Lacroix et al., 2021). For example, phospho-76 rus is primarily delivered to the ocean as particulate organic phosphate (POP), medi-77 ated by land biosphere processes, and attached to surfaces of iron-manganese and hy-78 droxide particles, in addition to dissolved inorganic and dissolved organic phosphorus 79 (Compton et al., 2000; Lacroix et al., 2021). Estimates suggest that there are weather-80 ing hotspots and that 70% of today's global weathering fluxes derive from only 10% of 81 the land area (Hartmann et al., 2014). Carbon is also released to the climate system via 82 outgassing of magmas at seafloor hydrothermal vents, hot-spot and island arc volcanoes, 83 and by other forms of volcanism (Hayes & Waldbauer, 2006). Chemical weathering rates 84 are thought to be governed by lithology, hydrology and runoff, rates of physical erosion, 85 soil properties and shielding of rocks by soils, and temperature. (e.g., Hartmann et al., 86 2014; Colbourn et al., 2013). All these parameters changed over glacial-interglacial cy-87 cles and Earth's history. 88

The input of carbon and nutrients from the lithosphere is roughly balanced by the burial of biogenic particles and other material in the lithosphere, yet imbalances remain. For example, the input by weathering and the input-burial balance was strongly perturbed over past glacial-interglacial cycles (L. C. Peterson & Prell, 1985; Broecker & Peng, 1987; Kump & Alley, 1994; Cartapanis et al., 2016, 2018). Input-burial imbalances directly affect the inventories and concentrations of carbon, nutrients, and alkalinity in the ocean. In turn, the production and burial of biogenic particles are changing.

The burial of biogenic particles is mediated by the marine carbon and biogeochem-96 ical cycles (e.g., Emerson & Bender, 1981; Sarmiento & Gruber, 2006; Tschumi et al., 97 2011). Marine ecosystems remove dissolved inorganic carbon (DIC), nutrients, and al-98 kalinity from surface waters to eventually generate dissolved organic matter and parti-99 cles of calcium carbonate  $(CaCO_3)$ , opal, and organic matter. The particles sink through 100 the water column towards the ocean floor. This biogenic material is mainly remineral-101 ized to DIC and inorganic nutrients within the water column and surface (reactive) sed-102 iments, but a fraction is buried in consolidated sediments. Rates of particle sinking, rem-103 ineralization, and burial depend on particle size and composition, and environmental pa-104 rameters. These parameters include, for example, temperature, oxygen, viscosity, or the 105 saturation state of water for  $CaCO_3$  in the water column and reactive sediments. Large 106 amounts of carbon and nutrients get buried in organic forms in the coastal zone and on 107 continental shelves (Regnier et al., 2013; Wallmann et al., 2016). This material may be 108 released again from exposed shelves during low glacial sea level and contribute, together 109 with changes in weathering and burial, to imbalances between fluxes from and to the litho-110 sphere (Wallmann, 2014; Wallmann et al., 2016). Changes in surface ocean DIC and al-111 kalinity from input-burial imbalances and altered particle cycling change atmospheric 112  $CO_2$ , forcing climate to change. Input-burial imbalances also strongly affect  $\delta^{13}C$  of car-113 bon in the atmosphere, ocean, ocean sediments, and the land biosphere (Broecker, 1970; 114

Schrag et al., 2013; Roth et al., 2014; Mills et al., 2017; Jeltsch-Thömmes & Joos, 2020;
Komar & Zeebe, 2021).

Modeling the responses to input-burial imbalances is computationally challenging. 117 The response time scales to input-burial imbalances range from several millennia to hun-118 dreds of thousands of years for  $CO_2$  (e.g. Archer et al., 1998; Colbourn et al., 2015) and 119  $\delta^{13}$ C (Roth et al., 2014; Jeltsch-Thömmes & Joos, 2020). Spatial gradients within the 120 ocean are important and influence, for example, the isotopic composition of the burial 121 flux of organic and  $CaCO_3$  particles into the lithosphere (Jeltsch-Thömmes & Joos, 2020). 122 123 Earlier model studies on glacial-interglacial change considered the effect of sedimentary carbonate burial and dissolution on perturbations in atmospheric CO<sub>2</sub>, but typically ne-124 glected organic matter burial (e.g., Broecker & Peng, 1987, 1989; Emerson & Archer, 1992; 125 Archer & Maier-Reimer, 1994; Sigman et al., 1998; Archer et al., 2000; Sigman & Boyle, 126 2000; Sigman et al., 2010; Brovkin et al., 2012; Ganopolski & Brovkin, 2017; Willeit et 127 al., 2019). Further studies show that feedbacks between organic matter burial, marine 128 nutrient concentrations, and biological productivity are important (Tschumi et al., 2011; 129 Menviel et al., 2012; Cartapanis et al., 2018; Jeltsch-Thömmes et al., 2019; Komar & Zeebe, 130 2021) and can strongly amplify perturbations in atmospheric CO<sub>2</sub> on glacial-interglacial 131 time scales (Roth et al., 2014). Input-burial imbalances arising from changes in the burial 132 fluxes of opal and carbonate or the weathering rates of silicate and carbonate rocks (e.g., 133 Munhoven & François, 1996; Munhoven, 2002; Clark et al., 2006; Willeit et al., 2019; Börker 134 et al., 2020; Köhler & Munhoven, 2020), or changes in CO<sub>2</sub> outgassing associated with 135 volcanism (Huybers & Langmuir, 2009; Roth & Joos, 2012) have been invoked as pos-136 sible contributing processes to explain glacial-interglacial variations and the Neogene cool-137 ing (23 to 2.6 million years ago) (e.g., Caves et al., 2016; Rugenstein et al., 2019). On 138 even longer timescales, covering Earth's history, the coupled carbon-silica cycle is thought 139 to stabilize climate on Earth via negative feedbacks in weathering of carbonate and sil-140 icate rocks (e.g., Walker et al., 1981; Berner, 1990; Ridgwell & Zeebe, 2005; Stolper et 141 al., 2016; Kasting, 2019; Isson et al., 2020). Net organic matter accumulation in sedi-142 ments and the accumulation of the oxidizing capacity at Earth's surface is reconstructed 143 from  $\delta^{13}$ C recorded in carbonate deposits (Broecker, 1970; Schrag et al., 2013; Mills et 144 al., 2017). Models applied to quantitatively interpret the  $\delta^{13}$ C carbonate records typ-145 ically treat the fast exchanging reservoirs (atmosphere, ocean, ocean sediments, and land 146 biosphere) as a single boxes to facilitate long simulations (Berner, 2006; Bergman et al., 147 2004). 148

Cost-efficient substitute models, which capture the spatial and temporal responses 149 of more complex models, could be constructed and applied to study input-burial imbal-150 ances instead of box models. The substitute (or emulator) can be used to explore responses 151 over long time scales or to run many sensitivity studies, which are computationally in-152 accessible with more comprehensive and therefore more expensive models. The spatio-153 temporal response of a complex model, its Green's function, can be captured in an ide-154 alized model simulation where the forcing is changed in a step- or pulse-like manner (e.g., 155 Maier-Reimer & Hasselmann, 1987; Joos & Bruno, 1996; Thompson & Randerson, 1999; 156 Hooss et al., 2001; Joos et al., 2013; Metzler et al., 2018; Strassmann & Joos, 2018; Bas-157 tiaansen et al., 2021). Idealized response simulations allow for a better understanding 158 of underlying processes as they reveal the characteristic timescales and spatial patterns 159 of the system's adjustment to an external perturbation, e.g., a change in weathering. While 160 the processes and timescales affecting  $CO_2$  have been investigated in several step-change 161 experiments (e.g., Archer & Maier-Reimer, 1994; Sigman et al., 1998; Tschumi et al., 2011), 162 to our knowledge no studies so far have systematically investigated the effect of changes 163 in weathering input fluxes on both, carbon and carbon isotope budgets. 164

Here, we use the Bern3D Earth system model of intermediate complexity to perform idealized, up to 600 kyr long simulations. The weathering input fluxes of alkalinity, nutrients, carbon, and  $\delta^{13}$ C are changed in a step-wise manner. The aim is to generate Green's functions and to understand the spatial and temporal responses in carbon inventories and  $\delta^{13}$ C in the atmosphere and ocean, and changes in marine biogeochemical cycling as well as the evolution of input-burial imbalances. We quantify the contribution to the carbon and carbon isotopic perturbations from the organic carbon and CaCO<sub>3</sub> cycles and highlight the role of vertical gradients in  $\delta^{13}$ C in the ocean. The response is used to build a cost-efficient substitute model to simulate the response in CO<sub>2</sub>, CO<sub>3</sub><sup>2-</sup>,

and  $\delta^{13}C_{\text{DIC}}$  to different weathering scenarios over the past 800 kyr.

#### <sup>175</sup> 2 Model Description and Experimental Set up

## 2.1 Model Description

The Bern3D v2.0s intermediate complexity model couples a single layer energy-moisture 177 balance atmosphere with a thermodynamic sea-ice component (Ritz et al., 2011), a 3D 178 geostrophic-frictional balance ocean (Edwards et al., 1998; Müller et al., 2006) with an 179 isopycnal diffusion scheme and Gent-McWilliams parameterization for eddy-induced trans-180 port (Griffies, 1998), and a 10-layer ocean sediment module (Heinze et al., 1999; Tschumi 181 et al., 2011; Jeltsch-Thömmes et al., 2019). The horizontal resolution across Bern3D model 182 components is  $41 \times 40$  grid cells and 32 logarithmically spaced depth layers in the ocean 183 (Roth et al., 2014). Wind stress is prescribed from the NCEP/NCAR monthly wind stress 184 climatology (Kalnay et al., 1996), and gas exchange at the ocean surface and calcula-185 tion of carbonate chemistry follow OCMIP-2 protocols (Najjar & Orr, 1999; Wanninkhof, 186 2014; Orr & Epitalon, 2015), with an adjusted gas transfer dependency on wind speed 187 (Müller et al., 2008). In the ocean, marine productivity is restricted to the euphotic zone 188 (75 m) and calculated as a function of light availability, temperature, and nutrient con-189 centrations (P, Fe, Si; Parekh et al., 2008; Tschumi et al., 2011). 190

The sediment module covers the top 10 cm and dynamically calculates the trans-191 port, redissolution/remineralization, and bioturbation of solid material, the pore water 192 chemistry, and diffusion (see Tschumi et al., 2011). Burial (loss) of phosphorus, silica, 193 carbon, and alkalinity from the sediment to the lithosphere is balanced by a variable in-194 put flux to the coastal surface ocean during spin-up. These weathering input fluxes are 195 set equal to burial fluxes at the end of the spin-up for transient simulations. Iron is added 196 to the model ocean by prescribed fluxes from aeolian deposition and continental mar-197 gins and removed by particle scavenging (Parekh et al., 2008). Iron is not included in 198 the sediment module. 199

The model is coupled to a 4-box representation of the land-biosphere carbon reservoirs (Siegenthaler & Oeschger, 1987). Here, the only purpose of this 4-box carbon reservoir model is to represent the dilution of atmospheric isotopic perturbations by the land biosphere, while other processes and changes in land carbon stocks are not considered.

204

176

## Implementation of ${}^{13}C$ in the Model

 $^{13}$ C is implemented as a tracer in all Bern3D model components such that  $^{13}$ C fluxes 205 and inventories can be explicitly simulated across the atmosphere-ocean-land biosphere-206 reactive ocean sediments (AOBS) system. Fractionation of <sup>13</sup>C is considered for atmosphere-207 ocean gas transfer, carbonate chemistry, the formation of CaCO<sub>3</sub>, POC and DOC, and 208 during photosynthesis on land. Fractionation during photosynthesis on land is kept fixed; 209 the simple 4-box model, applied here for computational reasons, does not represent changes 210 in C3 and C4 plants nor changes in their discrimination. This is an uncertainty. No frac-211 tionation is considered for the remineralization of organic carbon in the ocean and on 212 land. Formulations on how fractionation is calculated and corresponding references are 213 summarized in table 1. 214

**Table 1.** Equations describing the fractionation in the model. T denotes sea surface temperature in Kelvin, and  $[CO_{2,aq}]$  aqueous  $CO_2$  in  $\mu$ mol kg<sup>-1</sup>.

References: <sup>a</sup> Siegenthaler and Muennich (1981), <sup>b</sup> Mook (1986), <sup>c</sup> Freeman and Hayes (1992), <sup>d</sup> Siegenthaler and Oeschger (1987)

domain	formulation
air-sea <sup><i>a</i>,<i>b</i></sup>	$\alpha_{a \to s} = \left(1 - 0.0005 - 0.0002\right) \cdot \left(1 - \frac{0.373}{T} + 0.00019\right)$
$\begin{array}{c} \text{carbonate} \\ \text{chemistry}^b \end{array}$	$\alpha_{\rm CO_{2,aq}\leftrightarrow HCO_{3}^{-}} = \left(1 - \frac{9.866}{T} + 0.02412\right)$
	$\alpha_{\rm HCO_3^- \leftrightarrow CO_3^{2-}} = \left(1 - \frac{0.867}{T} + 0.00252\right)$
	$\alpha_{\mathrm{CO}_{2,aq}\leftrightarrow\mathrm{DIC}} = \frac{\mathrm{DIC}}{[\mathrm{CO}_{2,aq}] + \frac{[\mathrm{HCO}_{3}^{-}]}{\alpha_{\mathrm{CO}_{2,aq}\leftrightarrow\mathrm{HCO}_{3}^{-}}} + \frac{[\mathrm{CO}_{3}^{2^{-}}] \cdot \alpha_{\mathrm{HCO}_{3}^{-}\leftrightarrow\mathrm{CO}_{3}^{2^{-}}}}{\alpha_{\mathrm{CO}_{2,aq}\leftrightarrow\mathrm{HCO}_{3}^{-}}}$
sea-air <sup><math>b</math></sup>	$\alpha_{s \to a} = (1 - 0.0005 - 0.0002) \cdot \alpha_{\text{CO}_{2,aq} \leftrightarrow \text{DIC}}$
$CaCO_3$ formation <sup>b</sup>	$\alpha_{\rm HCO_3^- \to CaCO_3} = \left(1 - \frac{4.232}{T} + 0.0151\right)$
$ \begin{array}{l} \text{marine} \\ \text{photosynthesis}^c \end{array} \\$	$\alpha_{[\rm CO_2] \to C_{\rm org}} = (1.00119 + 0.01203 \cdot \log_{10} ([\rm CO_{2,aq}]))$
$\begin{array}{c} \text{terrestrial} \\ \text{photosynthesis}^d \end{array}$	constant fractionation of -18.1 $\%$

#### Implementation of Weathering Input Fluxes in the Model

215

The loss of carbon (C), silica (Si), alkalinity (ALK), and phosphorus (P) from the 216 open ocean through sedimentary burial is counteracted by the input of weathered ma-217 terial to the open ocean. In the model, this is implemented in an idealized way: the loss 218 fluxes of tracers through burial of material in the lithosphere are, in steady state, bal-219 anced by input fluxes of equal size. The elemental ratios (<sup>12</sup>C:<sup>13</sup>C:Si:ALK:P) of the in-220 put flux to the open ocean are therefore dictated by the elemental ratios of the global 221 burial flux at the end of the spin up. Throughout the manuscript, these fluxes to the ocean 222 will be referred to as input fluxes. 223

C, Si, P, and ALK input are uniformly added to the corresponding dissolved in-224 organic tracer concentrations of the near-coastal surface grid cells in Bern3D. Thus, we 225 do not represent reprocessing of elements, e.g., by the land biosphere or in estuaries, nor 226 different forms of input such as dissolved and particulate organic matter, or P attached 227 to surfaces of iron-manganese and hydroxide particles (Compton et al., 2000; Lacroix et 228 al., 2021). Atmospheric deposition of C, Si, ALK, and P may be viewed as included in 229 the input fluxes. This simplification appears justified as the different forms of input even-230 tually enter the inorganic pools on time scales shorter than the multi-millennial timescales 231

addressed in this study. We attribute the elemental input fluxes conceptually to differ ent processes such as silicate and carbonate weathering.

Input from 1 mol CaSiO<sub>3</sub> adds one mole of Si, two moles of alkalinity, and no carbon to the model ocean. This is a short-circuit of the atmosphere following the simplified reaction for weathering of tectosilicates (e.g., Colbourn et al., 2013):

$$2\mathrm{CO}_{2(aq)} + \mathrm{H}_{2}\mathrm{O} + \mathrm{CaSiO}_{3(s)} \longrightarrow \mathrm{Ca}_{(aq)}^{2+} + 2\mathrm{HCO}_{3(aq)}^{-} + \mathrm{SiO}_{2(aq)}$$
(1)

Two moles of carbon are removed from the atmosphere-ocean system in the form of  $CO_2$ and two moles of alkalinity and carbon in the form of  $HCO_3^-$  are added to the ocean. On the timescales considered here, the atmosphere and surface ocean can be considered well mixed and we do not represent the outgassing of  $CO_2$  and the related interhemispheric ocean carbon transport arising from the riverine carbon inputs (Sarmiento et al., 1992; Lacroix et al., 2020).

243

Weathering of carbonate rocks  $(CaCO_3)$  is given by (e.g., Goodwin & Ridgwell, 2010):

$$\operatorname{CaCO}_{3(s)} + \operatorname{CO}_{2(aq)} + \operatorname{H}_2 O \longrightarrow \operatorname{Ca}_{(aq)}^{2+} + 2\operatorname{HCO}_{3(aq)}^{-}.$$
 (2)

In the model, two moles of alkalinity for each mole of carbon are added. This is again a short-circuit of the atmosphere.

Input fluxes (F) of P, Si, ALK, DIC, and  ${}^{13}C(DIC)$  are attributed to the weath-246 ering flux from carbonate rocks,  $F_{CaCO_3}$ , silicate rocks,  $F_{CaSiO_3}$ , an assumed conceptual 247 input flux releasing the elements of former organic material,  $F_{\rm org}$  (from hereon referred 248 to as organic input flux), and a remainder flux,  $F_{\text{remainder}}$ , to close the carbon and car-249 bon isotopic budget of the model in equilibrium.  $F_{\text{remainder}}$  is thought of as CO<sub>2</sub> emis-250 sions from volcanism and tectonic outgassing. The input  $F_{\text{org}}$  should not be confused 251 with a flux of organic material as all input fluxes are added to the inorganic tracer pools 252 in Bern3D. At steady state, the global elemental fluxes of  $F_{\rm org}$  are equal to those of the 253 global burial flux of particulate organic matter. The fluxes are related to each other in 254 molar units: 255

$$F_{\rm P} = \mathcal{R}_{\rm P:C} \cdot F_{\rm org} \tag{3}$$

$$F_{\rm Si} = F_{\rm CaSiO_3} \tag{4}$$

$$F_{\text{ALK}} = 2 \cdot F_{\text{CaSiO}_3} + 2 \cdot F_{\text{CaCO}_3} + \mathcal{R}_{\text{ALK:P}} \cdot \mathcal{R}_{\text{P:C}} \cdot F_{\text{org}}$$
(5)

$$F_{\rm DIC} = F_{\rm CaCO_3} + F_{\rm org} + F_{\rm remainder} \tag{6}$$

$$F_{\text{DIC}-13} = F_{\text{CaCO}_3} \cdot {}^{13} R_{\text{CaCO}_3} + F_{\text{org}} \cdot {}^{13} R_{\text{org}} + F_{\text{remainder}} \cdot {}^{13} R_{\text{remainder}}$$
(7)

The burial fluxes of P, Si, ALK, DIC, and <sup>13</sup>C(DIC) are diagnosed at the end of 259 the model spin-up and prescribed as the input fluxes for these components ( $F_{\rm P}$ ,  $F_{\rm Si}$ ,  $F_{\rm ALK}$ , 260  $F_{\text{DIC}}$ ,  $F_{\text{DIC}-13}$ ) at the start of transient simulations.  ${}^{13}R_{\text{CaCO}_3}$  and  ${}^{13}R_{\text{org}}$  are diagnosed 261 from the signatures of the CaCO<sub>3</sub> and particulate organic carbon (POC) burial.  ${}^{13}R_{\text{remainder}}$ 262 is the signature needed to close the <sup>13</sup>C budget of the model in equilibrium after the spin-263 up. We assume no fractionation for CaSiO<sub>3</sub> weathering. All input is assumed to be free 264 of <sup>14</sup>C. Eq. 3 to 7 are solved for the steady-state fluxes ( $F_{\text{org}}$ ,  $F_{\text{CaSiO}_3}$ ,  $F_{\text{CaCO}_3}$ ,  $F_{\text{remainder}}$ ), 265 and the signature  ${}^{13}R_{\text{remainder}}$ . Table 2 shows steady state fluxes and inventories of CaCO<sub>3</sub>, 266 POC, and opal, as well as  $\delta^{13}$ C signatures.  $\delta^{13}$ C of the export flux is not diagnosed in 267 the current model setup. 268

<b>Table 2.</b> Export, deposition, and burial fluxes as well as sediment inventories of $CaCO_3$ , opal,
and particulate organic carbon (POC) and $\delta^{13}$ C signatures, determined as the mean of the first
100 years after a pre-industrial spin-up in the control run, and corresponding literature estimates.
References: <sup>a</sup> Battaglia et al. (2016), <sup>b</sup> Tréguer et al. (2021), <sup>c</sup> Sarmiento and Gruber (2006), <sup>d</sup>
Milliman and Droxler (1996), $^{e}$ Feely et al. (2004)

variable	units	Bern3D	observational estimates	$\begin{array}{c} \delta^{13}\mathrm{C} \\ \mathrm{in}\% \end{array}$
Export				
$\overline{\text{CaCO}_3}$	$GtC yr^{-1}$	0.98	$0.72 - 1.05^{a}$	
opal	$Tmol Si yr^{-1}$	110.02	$112^{b}$	
POC	$GtC yr^{-1}$	11.93	$6.5 - 13.1^c$	
Deposition				
$\overline{\mathrm{CaCO}_3}$	$GtC yr^{-1}$	0.49	$0.5^d$	2.91
opal	$Tmol Si yr^{-1}$	77.6	$84^{b}$	
POC	$GtC yr^{-1}$	0.65	$1.7-3.3^{c}$	-20.39
Burial				
$\overline{\mathrm{CaCO}_3}$	$GtC yr^{-1}$	0.22	$0.1 - 0.14^{e}$	2.88
$opal/F_{Si}$	$Tmol Si yr^{-1}$	6.72	$9.2^{b}$	
$POC/F_{org}$	$GtC yr^{-1}$	0.24	$0.12 \text{-} 0.26^{c}$	-20.42
F <sub>ALK</sub>	Gt eq. $yr^{-1}$	0.41		
$F_{ m DIC}$	$GtC yr^{-1}$	0.46		-9.09
Reactive ocean sediments				
$CaCO_3$	$\operatorname{GtC}$	939		3.05
opal	Tmol Si	$20,\!658$		
POC	$\operatorname{GtC}$	516		-20.13
Input				
$\overline{F_{\text{CaCO}_3}}$	$GtC yr^{-1}$	0.14		2.88
$F_{ m CaSiO_3}$	$Tmol Si yr^{-1}$	6.72		
$F_{ m org}$	$GtC yr^{-1}$	0.24		-20.42
$F_{ m remainder}$	$GtC yr^{-1}$	0.08		2.88

#### 269

#### 2.2 Experimental Set-up and Analysis

The model is spun up over 60 thousand years (kyr) under 1765 CE boundary con-270 ditions. An atmospheric  $CO_2$  concentration of 277.8 ppm with an isotopic signature of 271  $\delta^{13}C_{atm} = -6.305 \%$  and  $\Delta^{14}C_{atm} = 0 \%$  is prescribed. From the 1765 CE steady state, 272 at nominal year -100, 100.1 kyr long experiments are started. After 100 years, i.e., in nom-273 in al year 0 of the experiments, step-changes in  $F_{\text{CaCO}_3}$  and  $F_{\text{org}}$  from -80% to +80% in 274 steps of 20% are applied. The corresponding amounts of carbon,  $^{13}C$ , alkalinity, and P 275 (Eqs. 3-7) are then added to the coastal surface ocean. A subset of the above experi-276 ments, namely the changes of +40%, -40%, and the control run, are extended by an ad-277 ditional 500 kyr simulation period in order for the model to establish new equilibrium 278 after the step-change in carbon isotopes. 279

For organic input, an additional 100 kyr sensitivity experiment is performed to investigate the effect of differences in the  $\mathcal{R}_{P:C}$  ratio. To this mean, 30% of the P entering marine sediments is immediately dissolved back into the ocean. This P leaching changes the sedimentary and therefore also the input flux  $\mathcal{R}_{P:C}$  to about 1 : 152. Our setup for changing  $F_{\text{org}}$  is different from a step-change in the ocean P inventory as applied by Tschumi et al. (2011) and others. In the experiments presented in this study, the change in  $F_{\text{org}}$ implies a sustained change in the input fluxes of P, C, <sup>13</sup>C, and ALK.

Results presented in this study are relative to a control run to account for any drift over the 600 thousand year (kyr) simulation period. Generally, drift is small, and amounts, for example in the case of  $CO_2$ , to well below 0.01 ppm kyr<sup>-1</sup> in the control simulation. The output frequency for marine 3-D tracer fields after the perturbation is every 10 years during the first 1 kyr, every 200 years until 10 kyr, every 1 kyr until 100 kyr, and every 5 kyr thereafter.

<sup>293</sup> Changes in the carbon isotopic fluxes and inventories are expressed in [GtC %] by <sup>294</sup> multiplying inventories or fluxes of carbon [GtC] with their corresponding  $\delta^{13}$ C signa-<sup>295</sup> tures [%]. We attribute the change in the isotopic burial minus input flux to changes <sup>296</sup> ( $\Delta$ ) in carbon burial (B) and input (I) fluxes and in the isotopic signature of the burial <sup>297</sup> flux ( $\delta_B$ ) relative to the equilibrium state after the spin-up (subscript 0) for both POC <sup>298</sup> and CaCO<sub>3</sub>:

$${}^{13}B - {}^{13}I = (B_0 + \Delta B) \cdot (\delta_0 + \Delta \delta_B) - (I_0 + \Delta I) \cdot \delta_0$$
  
=  $\Delta (B - I) \cdot \delta_0 + B_0 \cdot \Delta \delta_B + \Delta B \cdot \Delta \delta_B$  (8)

Note that the steady state carbon and isotopic burial-input fluxes vanish with  $B_0 - I_0 = 0$  and  $\delta_{B,0} = \delta_{I,0} = \delta_0$ .

## 2.3 Green's Function Substitute Model

301

Green's functions for a change in input flux (weathering) are directly obtained from 302 the step-change simulations with Bern3D. Generally, the Green's function can be deter-303 mined by prescribing a step-like change in forcing (e.g., the magnitude of weathering flux) 304 in a model previously spun up to equilibrium. Then the model is run towards a new equi-305 librium and the simulated change  $\Delta y(\vec{x},\tau)$  of any variable of interest (e.g.,  $\delta^{13}C_{\text{DIC}}$ ) is 306 monitored. The response  $\Delta y$  is normalized by the magnitude of the step change to yield 307 the Green's function,  $r^y(\vec{x},\tau)$ .  $\tau$  is the time passed since the step-like change and  $\vec{x}$  in-308 dicates location. 309

Any forcing history, F(t), may be approximated by a series of small step-like changes,  $\Delta F(t_i)$ . Then, the response,  $\Delta y(\vec{x}, t)$ , in variable y at time t and location  $\vec{x}$  to this forcing history is for a linear system:

$$\Delta y(\vec{x},t) = \sum_{i} \Delta F(t_i) \cdot r^y(\vec{x},t-t_i).$$
<sup>(9)</sup>

 $t-t_i$  represents the time  $\tau_i$  that has passed since the step change  $\Delta F(t_i)$  at time  $t_i$ . The sum is over all step changes used to approximate the forcing history from its beginning up to time t. The system is assumed in equilibrium at the start of the forcing time series.

Instead of running a complex 3-dimensional dynamic model, its response is approximated by Eq. 9, a great efficiency gain. For further ease of application, a Green's function can be approximated using Principal Component-Empirical Orthogonal Functions (PC-EOF). The temporal component of the PC-EOF can be approximated by a sum of exponential terms and the substitute model (Eq. 9) can be represented by a series of box models and spatial patterns for further computational efficiency. (see Joos et al., 2001; Strassmann & Joos, 2018; Jeltsch-Thömmes & Joos, 2020).

The dynamics of a linear system are fully characterized by its Green's function. Although the carbon cycle and the Earth system are not linear, the linear approximation of Eq. 9 is often useful. Here, we determined Greens's function over a wide range of step changes (-80% to +80% of the unperturbed flux) to evaluate non-linearities. In the discussion in section 4.2, we will apply Eq. 9 for different weathering scenarios, using directly the (normalized) model output from the step simulations described in section 2.2.

#### 330 3 Results

#### 331 332

#### 3.1 Earth System Response to a 40% Increase in the CaCO<sub>3</sub> Weathering Input Flux

We first address Earth system responses to changes in CaCO<sub>3</sub> weathering. The prescribed 40% step-like increase in the CaCO<sub>3</sub> weathering flux translates into an increase in the inputs of carbon and alkalinity in a 1:2 molar ratio (Fig. 1a-b). The  $\delta^{13}$ C signature of the total carbon input is also increased by 1.3% (Fig. 1c) because the relative share of the isotopically heavy CaCO<sub>3</sub> weathering input becomes larger. The individual  $\delta^{13}$ C signatures of the CaCO<sub>3</sub> and POC input remain unchanged in our setup.

In response to the step-increase in carbon and alkalinity (ALK) input, a new equilibrium of the climate-carbon system is approached with an e-folding timescale ( $\tau$ ) of about 10 thousand years (kyr, Fig. 1d-l; 2a). In other words, concentrations and fluxes typically change most rapidly immediately after the step and then slowly approach a new equilibrium following roughly an exponential curve. As a remarkable exception,  $\delta^{13}C$  isotopic signatures continue to change over a few 100 thousand years (Fig. 1m). Possible fits of the responses in  $\Delta CO_2$  and  $\Delta \delta^{13}C_{DIC}$  are provided as equations in Fig. 1.

Atmospheric  $CO_2$  decreases, by almost 32 ppm (Fig. 1d), as the surface ocean con-346 centration of ALK increases about twice as much as that of dissolved inorganic carbon 347 (DIC) (Fig. 1f,g). Surface air temperature (SAT) decreases by about 0.5 °C (Fig. 1e) 348 in response to lower atmospheric  $CO_2$ . DIC and ALK increase by about 37 and 68 mmol 349  $m^{-3}$  on ocean average (Fig. 1f,g). Due to this "ocean alkalinization", ocean pH, and car-350 bonate ion concentration increase in the surface and deep ocean, and the calcium car-351 bonate saturation horizon deepens on average by about 700 m in the North Pacific (Fig. 352 1h-j). In turn, marine biogenic  $CaCO_3$  is better preserved in ocean sediments, and the 353 net transfer of  $CaCO_3$  from the ocean to the sediments, and eventually the lithosphere, 354 increases by about 25% (Fig. 11). The equilibrium increase in CaCO<sub>3</sub> burial of 25% cor-355 responds to the implied increase in the  $Ca^{2+}$  input of 25% in our scenario which results 356 from constant  $F_{CaSiO_3}$  and a 40% increase in  $F_{CaCO_3}$ . The modeled saturation horizon 357 in the equatorial Atlantic was already relatively close to the ocean bottom before the step-358 change in input, leaving little room for further deepening (Fig. 1i). The timescale of this 359 carbonate compensation process (~8-10 kyr, e.g. Archer et al., 1997, 1998; Jeltsch-Thömmes 360 & Joos, 2020) is reflected in the timescale of the  $CO_2$  perturbation of ~10 kyr. 361

Changes in global export fluxes of particulate organic matter (POM),  $CaCO_3$ , and 362 opal are comparably small (<3%, Fig. 1k) and result in only small declines in the sed-363 imentary POM and opal inventories (Fig. 11). Export production generally decreases as 364 a result of lower temperatures and increased sea-ice area, with the largest changes in the 365 Southern Ocean. The higher input of ALK to the surface ocean (Fig 1a), together with 366 the slight reduction in  $CaCO_3$  export (Fig 1k), causes a larger increase of ALK in the 367 surface than the deep ocean (Fig 1g), while the surface-to-deep gradient in DIC remains 368 almost unaffected by the increase in  $CaCO_3$  weathering (Fig 1f). 369

We establish budgets for the carbon (Fig. 2a) and alkalinity perturbations by com-370 paring accumulated burial-input imbalances for  $CaCO_3$  and POM with changes in the 371 major reservoirs, atmosphere, ocean, and sediments. Carbon and ALK fluxes are sto-372 ichiometrically linked. C:ALK molar ratios of 1:2 and 117:-17 are applied for  $CaCO_3$  and 373 POM and the molar weight of carbon is  $12.01 \text{ g mol}^{-1}$ . Carbon and alkalinity storage 374 in the ocean increases by about 600 GtC and 91 Petamol eq, respectively. The major-375 ity of the perturbation in the combined atmosphere-ocean system ( $\sim$ 521 GtC; 91 Pmol 376 eq) and the reactive ocean sediments ( $\sim 334$  GtC; 60 Petamol eq) is driven by imbalances 377

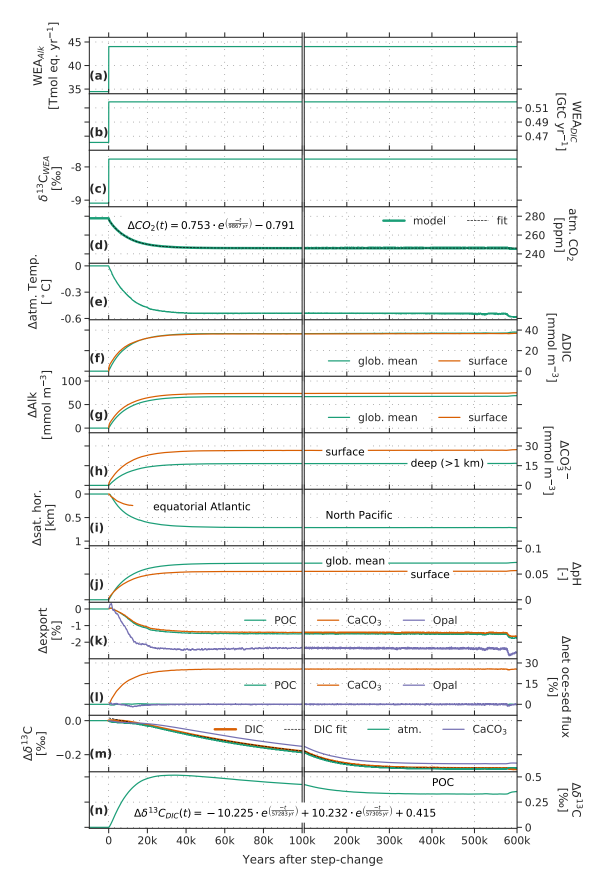


Figure 1. Timeseries evolution of the forcing (a-c) and a set of Earth system properties (d-n) for a 40% step increase in the  $CaCO_3$  input flux. For better visibility, data in panels e, k, l, and m (only  $CaCO_3$ ) is shown as 100 yr running means. Surface refers to upper 100 m.

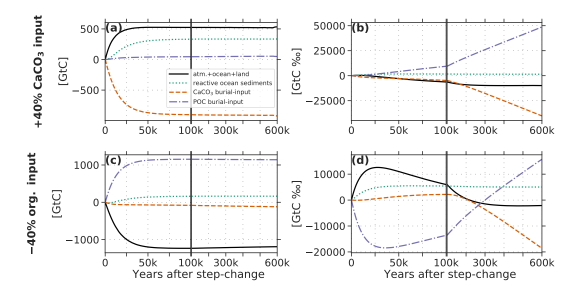


Figure 2. Carbon (a,c) and carbon isotopic (b,d) budget for step-changes of +40% and -40% in the CaCO<sub>3</sub> (a,b) and organic (c,d) input flux.

in the CaCO<sub>3</sub> burial-input flux ( $\sim$ -912 GtC; 152 Petamol eq). Contributions from imbalances associated with the POM cycle are small ( $\sim$ 53 GtC; -0.6 Petamol eq) (Fig. 2a). Accordingly, the majority of change in the reactive ocean sediments is explained by an increased CaCO<sub>3</sub> content ( $\sim$ 361 GtC; 60 Petamol eq) with little contribution from decreased POM storage ( $\sim$ -27 GtC; 0.3 Petamol eq). The values refer to means over the period 500-550 kyr after the step change.

## 384 Carbon Isotopes

After an initial small increase, resulting from the higher  $\delta^{13}$ C signature of the weathering input after the step-change (Fig. 1c),  $\delta^{13}$ C decreases gradually after the step in the atmosphere, ocean, land biosphere (AOB), and sedimentary CaCO<sub>3</sub> reservoirs.  $\delta^{13}$ C in these reservoirs stabilizes at about -0.28 ‰ lower values after several hundred kyr (Fig. 1n). In contrast,  $\delta^{13}$ C of sedimentary POC increases relatively fast to peak at around 30 kyr after the step. Afterwards, the anomaly decreases slightly to stabilize at about 0.35 ‰ after several hundred kyr (Fig. 1m).

The  $\delta^{13}$ C perturbation budget is shown in (Fig. 2b). The accumulated isotopic per-392 turbation in the AOB system stabilizes at  $\sim$ -9800 GtC  $\infty$  after several hundred kyr (Fig. 393 2b, black line). The change is mainly linked to the decrease in the mean isotopic signa-394 ture of the constituents of the AOB system (Fig. 1m); The contribution from the change 395 in the carbon inventory of the AOB system is small (not shown). The perturbation in 396 reactive sediments accumulates to  $\sim +1500$  GtC % (Fig. 2b, green dotted). This pos-397 itive perturbation is mainly due to the decrease in the sedimentary POC inventory, with 398 further contributions from the change in POC signature and from CaCO<sub>3</sub>. 399

The equilibration of  $\delta^{13}$ C after the change in CaCO<sub>3</sub> input is controlled by burialinput imbalances in both the CaCO<sub>3</sub> and POC cycles (Fig. 2b). This is different than for carbon for which burial-input imbalances of CaCO<sub>3</sub> dominate the response (Fig. 2a). The  $\delta^{13}$ C perturbation in the AOBS system is stabilized by the canceling effects of POC versus CaCO<sub>3</sub> burial-input flux. While the POC burial-input flux removes <sup>13</sup>C from the system, the CaCO<sub>3</sub> burial-input flux adds <sup>13</sup>C (Fig. 2b, magenta versus orange lines).

For further understanding, we attribute the net isotopic fluxes of POC and CaCO<sub>3</sub> burial-input according to Eq. 8 (Fig. 3a-c). The three terms considered are linked to changes

in the net burial-input carbon flux  $(\Delta(B-I)\cdot\delta_0)$ , in the  $\delta^{13}$ C signature of the burial 408 flux  $(\Delta \delta_B \cdot B_0)$ , and the perturbations in the burial carbon flux and signature  $(\Delta B \cdot B_0)$ 409  $\Delta \delta_B$ ) (Fig. 3). The majority of the carbon isotopic perturbation in response to a 40% 410 increase in CaCO<sub>3</sub> input is caused through changes in the  $\delta^{13}$ C signatures of the POC 411 and  $CaCO_3$  burial fluxes (Fig. 3b). Lower atmospheric  $CO_2$  (Fig. 1d) and therefore lower 412 surface water  $[CO_{2,aq}]$  reduces fractionation during marine photosynthesis (see Table 1). 413 As a result,  $\delta^{13}C$  of the POC export flux increases, and the mean  $\delta^{13}C$  of sedimentary 414 POC peaks about 30 kyr after the step change in CaCO<sub>3</sub> input (Fig. 1n). Consequently, 415  $\delta^{13}$ C of the POC burial flux increases, and  $^{13}$ C is removed by POC burial-input. This 416 removal causes  $\delta^{13}$ C of DIC to decrease (Fig. 1m; 2b). The negative perturbation in  $\delta^{13}$ C<sub>DIC</sub> 417 is incorporated into newly formed CaCO<sub>3</sub> and POC, explaining the gradual decrease in 418  $\delta^{13}$ C of the CaCO<sub>3</sub> sediment inventory (Fig. 1m) and burial flux, and the post-peak de-419 crease in the  $\delta^{13}$ C signature of the POC sediment inventory (Fig. 1n). 420

<sup>421</sup> Changes in the carbon isotopic budget due to the carbon burial-input imbalance <sup>422</sup>  $(\Delta(B-I) \ \delta_0)$  are comparably small (Fig. 3a). They are larger for CaCO<sub>3</sub> than for POC <sup>423</sup> as a result of the larger cumulative change in the burial-input flux for CaCO<sub>3</sub> than for <sup>424</sup> POC (Fig. 2a). For the same reason, the contribution of  $\Delta B \cdot \Delta \delta$  is substantially larger <sup>425</sup> for CaCO<sub>3</sub> (~ 8000 GtC‰) than POC (~ 0 GtC‰) (Fig. 3c).

The perturbation in  $\delta^{13}C_{DIC}$  varies spatially within the ocean. The surface-to-thermocline 426 gradient in  $\delta^{13}C_{DIC}$  is reduced after the step in CaCO<sub>3</sub> input. In Fig. 5a horizontally-427 averaged  $\Delta \delta^{13} C_{\text{DIC}}$  is plotted versus time and depth.  $\Delta \delta^{13} C_{\text{DIC}}$  is on average slightly 428 less negative in the thermocline than in surface and deep ocean waters. The differences 429 of  $\Delta \delta^{13} C_{\text{DIC}}$  in surface versus  $\Delta \delta^{13} C_{\text{DIC}}$  of thermocline and deep waters are small, <0.05 430  $\infty$  on average (Fig. 5c). They are a consequence of reduced fractionation during ma-431 rine photosynthesis resulting from lower  $[CO_{2,aq}]$ . Thus, the POC flux transports a less 432 negative isotopic signal from the surface to the thermocline, where most of the POC is 433 remineralized. Changes in circulation could contribute to a different  $\delta^{13}C_{DIC}$  gradient 434 but variations in modeled ideal age of water masses are very small (<10 years, not shown). 435 The response to a 40% decrease in CaCO<sub>3</sub> input yields a very similar response but with 436 inverse sign (Fig. 5b,c) and is not further discussed here. 437

In summary, the carbon-climate system equilibrates typically within a few tens of kyr after the step change in CaCO<sub>3</sub> input, whereas it takes several hundred kyr to establish a new equilibrium for  $\delta^{13}$ C signatures. Changes in proxy-related variables are a decrease in CO<sub>2</sub> and SAT, an increase in carbonate ion concentration with deepening of the lysocline and higher carbonate preservation, near absent or small responses in POM and opal bulk fluxes, and a long-term decrease in  $\delta^{13}$ C of atmospheric CO<sub>2</sub> and DIC with a slightly reduced surface-to-thermocline gradient in  $\delta^{13}$ C<sub>DIC</sub>.

Remarkably, changes in CaCO<sub>3</sub> input cause changes in  $\delta^{13}$ C of DIC and atmospheric 445  $CO_2$  of several tenths of a permil. Such  $\delta^{13}C$  changes are comparable to reconstructed 446 changes over glacial-interglacial cycles. Also remarkable, the change in  $\delta^{13}$ C of CaCO<sub>3</sub> 447 and DIC is with -0.2% opposite to the change in the signature of the input flux of +1.3%. 448 This is different to expectations of equal change in  $\delta^{13}$ C signatures when treating the 449 climate system as one box (e.g., Eq. 1 in (Schrag et al., 2013)) and neglecting changes 450 in fractionation. The contribution of the POM cycle, in particular a smaller isotopic frac-451 tionation during marine photosynthesis under lower  $CO_2$ , is key to understanding the 452 isotopic changes. These  $\delta^{13}$ C changes become only evident when considering the reor-453 ganization of the carbon cycle within the climate system instead of treating the climate 454 system as a single reservoir and when considering POC input and burial fluxes in ad-455 456 dition to CaCO<sub>3</sub> input-burial imbalances.

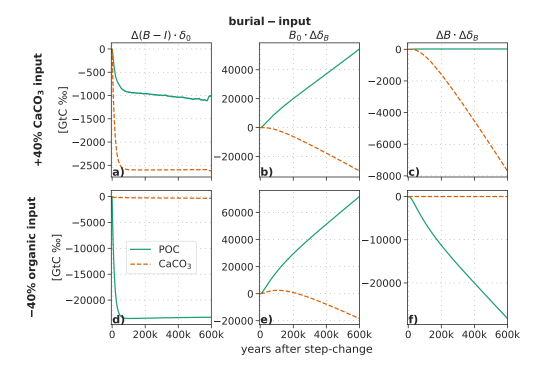


Figure 3. Cumulative contribution of the burial-input flux to the carbon isotopic perturbation for a (a,b,c) 40% increase in the CaCO<sub>3</sub> and a (d,e,f) 40% decrease in the organic input flux. The contribution is split into changes in the (a,d) carbon flux ( $\delta_0 \cdot \Delta F$ ), the (b,e)  $\delta^{13}C$  signature  $(\Delta\delta\cdot F_0)$  of the flux, and their (c,f) combination  $(\Delta\delta\cdot\Delta F)$ .

#### 3.2 Earth System Response to a 40% Decrease in the Organic Input Flux

A 40% decrease in the organic input flux reduces P and carbon inputs, increases 458 the alkalinity input, and leads to an enrichment in the  $\delta^{13}$ C signature of the total car-459 bon input (Fig. 4a-d). Similar as for the change in CaCO<sub>3</sub> input, most carbon cycle fluxes 460 and tracers adjust after the step-change towards a new equilibrium on time scales of or-461 der  $\sim 7.5$  kyr (Fig. 4e-m). Exceptions are the CaCO<sub>3</sub> ocean-to-sediment flux which is 462 only temporarily perturbed (Fig. 4m, orange lines), and  $\delta^{13}$ C signatures which show changes 463 over several hundred kyr (Fig. 4n,o). The e-folding timescale for the fit of the atmospheric 464  $CO_2$  perturbation is different from the  $CaCO_3$  step-change experiment, as the under-465 lying processes differ. While in the case of the  $CaCO_3$  step-change experiment,  $CaCO_3$ 466 compensation is the controlling process, in the case of the organic step-change, an in-467 terplay between changes in POC export production and burial rates, mediated by changes 468 in the phosphorous inventory are at play, as described in detail below. 469

457

In response to the step-reduction in carbon input, DIC, atmospheric CO<sub>2</sub>, and temperature decrease (Fig. 4e-g). In the steady state solution of the model, the organic input flux contributes about half of the carbon input (Table 2). Modeled changes in the atmospheric and marine carbon inventory are thus a direct result of reduced carbon input after the step-change.

The marine P inventory decreases as a result of the reduced P input, limiting biological productivity. Production and export of organic material, as well as of CaCO<sub>3</sub>, decreases by about 23% (Fig. 41). In turn, the POC flux to sediments and eventually POC burial decreases by 40%, thereby exactly offsetting the reduction in organic matter input at the new equilibrium. Opal export is only reduced by a couple of percents. At the

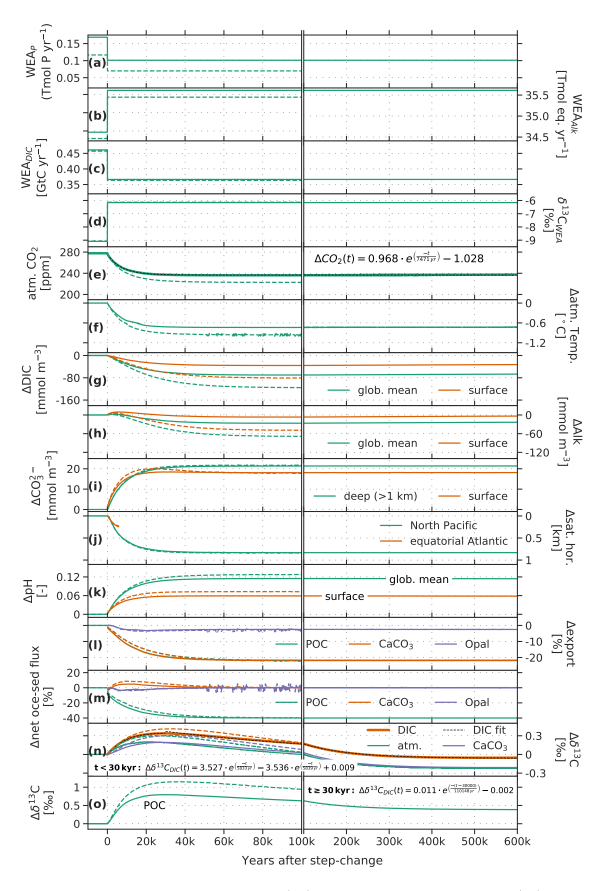


Figure 4. Timeseries evolution of the forcing (a-d) and a set of Earth system properties (e-o) for a 40% decrease in the organic input flux. Dashed colored lines show results from the sensitivity experiment with a sedimentary C:P ratio of ~152:1. For better visibility, data in panels f, l, m, and n (only CaCO<sub>3</sub>) is shown as 100 yr running means. Surface refers to upper 100 m.

end of the simulation, opal export is reduced in high latitudes as a result of increased sea-ice cover, and increased in low- and mid-latitudes (not shown).

In response to reduced DIC, the carbonate ion concentration increases as  $[CO_3^{2-}]$ 482  $\sim$  [ALK]-[DIC]) (Fig. 4g-i). ALK input and ALK in the ocean changes only little dur-483 ing the first 10 kyr (Fig. 4b,h). The shift to a higher  $[CO_3^{2-}]/[DIC]$  ratio goes hand in hand with an increase in pH (Fig. 4k). The increase in  $[CO_3^{2-}]$  causes a deepening of the 484 485 saturation horizon by about 800 m in the North Pacific (in the equatorial Atlantic the 486 saturation was already relatively close to the ocean bottom before the step-change in in-487 put, leaving little room for further deepening Fig. 4j). In turn, more CaCO<sub>3</sub> is transferred from the ocean to the sediments in the first 50 kyr after the step change (Fig. 4m), de-489 spite the  $\sim 23\%$  reduction in CaCO<sub>3</sub> export. This CaCO<sub>3</sub> transfer lowers the ocean's ALK 490 inventory (Fig. 4h), and the temporary perturbation in the  $CaCO_3$  flux to sediments van-491 ishes again (Fig. 4m). 492

These marine carbon cycle changes affect the surface-to-deep ocean gradients in DIC and ALK (Fig. 4g,h; red versus green lines). The decrease in DIC and ALK is larger in the deep ocean than in the surface, and the surface-to-deep gradient in DIC is reduced on global average. These changes in gradients are linked to the reduction in POC and CaCO<sub>3</sub> export, that tend to remove less carbon and slightly less alkalinity from the surface ocean.

<sup>499</sup> Altogether, the AOB reservoir looses  $\sim 1196$  GtC (Fig. 2c). This change is driven <sup>500</sup> by a cumulative burial-input imbalance in the POC cycle of  $\sim 1142$  GtC with little con-<sup>501</sup> tribution from the CaCO<sub>3</sub> cycle ( $\sim -115$  GtC) (Fig. 2c). The overall increase in carbon <sup>502</sup> storage in reactive ocean sediments (163 GtC) is explained by a gain of CaCO<sub>3</sub> ( $\sim 367$ <sup>503</sup> GtC) which is partly balanced by a loss of POC ( $\sim -203$  GtC). Again, values refer to means <sup>504</sup> over the period 500-550 kyr after the step change.

#### 505 Carbon Isotopes

<sup>506</sup> Less input of isotopically light organic carbon shifts the  $\delta^{13}$ C signature of the atmosphere <sup>507</sup> and ocean to higher values at first. Similarly, the  $\delta^{13}$ C signature of the net ocean-sediment <sup>508</sup> flux of carbon increases. After peaking at around 20 kyr,  $\delta^{13}$ C in the atmosphere and <sup>509</sup> ocean, as well as  $\delta^{13}$ C of sedimentary POC and CaCO<sub>3</sub> (and therefore also the burial <sup>510</sup> flux) decrease slowly to reach new steady-state values after several 100 kyr (Fig. 4n,o).

The driving processes of these changes take place on roughly two timescales (Fig. 511 2d; 3d-f). The timescales are caused by the different response times in the carbon ( $\Delta(I-$ 512 B)) and carbon isotopic budgets. The carbon cycle approaches a new equilibrium after 513 multiple millennia and can be seen as constant after roughly 50 kyr (Fig. 2c). The pos-514 itive cumulative imbalance in the isotopically light POC burial-input (Fig. 2c, dashed 515 purple line) leads to a reduced removal of <sup>13</sup>C from the reactive carbon pools (Fig. 2d, 516 dashed purple line). This causes the initial increase in  $\delta^{13}$ C in these reservoirs (Fig. 4n,o; 517 2d; 3d). This initial, shorter timescale, is reflected in the fit of the  $\delta^{13}C_{\text{DIC}}$  perturba-518 tion for the first 30 kyr after the step-change (equation in panel n, Fig. 4). 519

In contrast, lower atmospheric  $CO_2$  and therefore lower  $[CO_{2,aq}]$  lead to less frac-520 tionation during marine photosynthesis and an increase in  $\delta^{13}$ C of sedimentary POC (Fig. 521 40) and the POC burial flux. Hereby, isotopically heavy carbon  $^{13}$ C is continuously re-522 moved from the combined AOB carbon pool (Fig. 3e; 2d). On timescales of multiple 10 523 kyrs, the change in signature outweighs the effect of the increased cumulative burial-input 524 flux on the carbon isotopic budget and the direction of change in  $\delta^{13}$ C signatures is re-525 verted (Fig. 4n,o; 2d). The longer timescale of this second phase of the  $\delta^{13}C_{DIC}$  pertur-526 bation is reflected in the fit for t > 30 kyr (panel o, Fig. 4). 527

 $\delta^{13}$ C of CaCO<sub>3</sub> follows surface ocean  $\delta^{13}$ C<sub>DIC</sub>. The associated isotopic flux (Figs. 2d; 3e, dashed orange lines) eventually balances the effect of the POC cycle ((Figs. 2d;

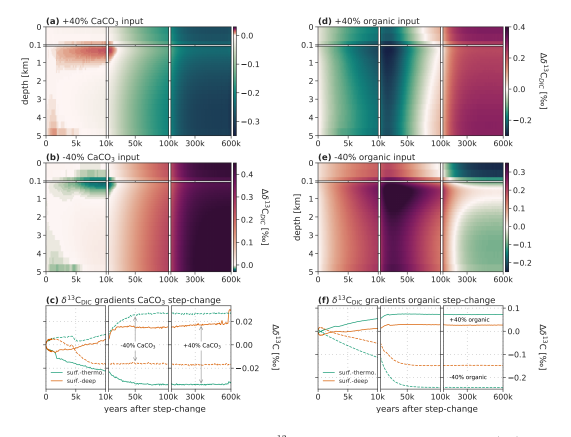


Figure 5. Hovmoeller-type diagram of the  $\delta^{13}C_{DIC}$  perturbation for step-changes of (a,d) +40% in the CaCO<sub>3</sub> and organic input fluxes, (b,e) -40% in the CaCO<sub>3</sub> and organic input fluxes, and (c,f) timeseries of the surface (euphotic zone)-to-thermocline (~600-700 m) and surface-to-deep (2.5-3.5 km) gradient for the respective experiments.

<sup>530</sup> 3d-f), green lines) on the carbon isotopic budget, leading to stable  $\delta^{13}$ C in the AOB sys-<sup>531</sup> tem after 300 kyr (Fig. 4n; 2d).

The perturbation in the POC export causes a strong perturbation in the surface-532 to-deep gradient of  $\delta^{13}C_{DIC}$ . Fig. 5e shows the horizontally-averaged evolution of the 533 perturbation in  $\delta^{13}C_{\text{DIC}}$  after the step-like decrease in input of organic material. Both, 534 the reduction in the export of POC by about 23% (Fig. 4m, green line) and the increase 535 of its  $\delta^{13}$ C signature (Fig. 40) weaken the overall surface-to-thermocline and the surface-536 to-deep gradient in  $\delta^{13}C_{\text{DIC}}$ . Changes in the export of labile dissolved organic matter 537 may also somewhat contribute. Further, changes in input minus burial fluxes also affect 538 spatial isotopic gradients. A clear perturbation in the gradient between the euphotic zone 539 (uppermost two vertical layers in the model) and the underlying waters develops, lead-540 ing to strongly enriched  $\delta^{13}C_{DIC}$  perturbation in thermocline and intermediate depth 541 waters, even reaching down to the ocean floor, as compared to the surface (see dark red 542 blob in Fig. 5e). After about 200 kyr a vertical perturbation pattern with a negative  $\delta^{13}C_{DIC}$ 543 perturbation in the surface, a slightly positive  $\delta^{13}C_{DIC}$  perturbation in the thermocline 544 and intermediate depth waters, and a slightly negative  $\delta^{13}C_{\text{DIC}}$  perturbation in waters 545 below  $\sim 2$  km depth has developed and appears to prevail. The surface-to-thermocline 546 gradient is changed by  $\sim 0.25$  ‰ and the surface-to-deep gradient by  $\sim 0.15$  ‰ on global 547 average. 548

In the case of a 40% increase in organic material input, the spatio-temporal evo-549 lution of  $\Delta \delta^{13} C_{DIC}$  is generally comparable, however the absolute perturbation in global 550 mean  $\delta^{13}C_{DIC}$  is initially smaller and stabilizes at higher levels, and changes in the surface-551 thermocline and surface-deep gradients are smaller. First, the change in the overall  $\delta^{13}$ C 552 signature of weathering carbon input does not change by the same absolute amount for 553 an increase and decrease in organic input. A 40% decrease in organic input changes the 554 overall  $\delta^{13}$ C signature of weathering carbon input by ~2 \%, while a 40\% increase changes 555 it by  $\sim 3$  ‰. The initial perturbation in  $\delta^{13}C_{\text{DIC}}$  caused by the changed input is thus 556 smaller for an increase than a decrease (c.f. Fig. 5d-f). Second, while for a 40% decrease 557 in organic input, the POM export flux is reduced by about  $\sim 23\%$ , in the case of a 40% 558 increase the POM export flux increases by only  $\sim 13\%$ , thus affecting surface-thermocline 559 and surface-deep gradients differently. This non-linearity in the carbon isotopic response 560 has to be kept in mind, when interpreting proxy records in light of these step-change ex-561 periments. 562

## 563 Sensitivity Experiment for Sedimentary P Leaching

Next, we turn to the results from the sensitivity experiment with idealized higher phosphate leaching from sediments (dashed lines in Fig. 4). 30% of the P in organic matter leaving the ocean is immediately "leached back" to the water column. In this model setup, the steady-state input flux of P is reduced in comparison to the standard run (Fig. 4a, solid vs dashed line).

Generally, the response to a reduction in the organic input flux is amplified in the 569 case of the sensitivity experiment. In particular, the reduction in atmospheric  $CO_2$  is 570 30% larger than in the standard model setup, as are the peak perturbations in  $\delta^{13}$ C of 571 DIC and  $CO_2$  around 20 kyr. The 40% reduction in the organic input flux translates into 572 a smaller absolute change in input in the sensitivity compared to the standard run, be-573 cause the steady-state input flux of P is smaller (Fig. 4a). In turn, P limitation for ma-574 rine biological production is less severe in the sensitivity experiment. This results in slightly 575 less reduced export fluxes of organic matter and  $CaCO_3$ , as compared to the standard 576 run (Fig. 41), translating into temporarily higher net ocean-sediment loss fluxes (Fig. 4m; 577 578 dashed vs solid lines). The higher loss of POC and  $CaCO_3$  causes DIC (and to a lesser extent ALK) and  $CO_2$  to decrease more in the sensitivity than standard setup. 579

<sup>580</sup> Differences in the isotopic perturbation can be readily understood from differences <sup>581</sup> in the burial-input fluxes and an even further reduced fractionation during marine pho-

to synthesis owing to the lower  $CO_2$  concentration (see Table 1 and Fig. 4e,m-o). The 582 majority of the carbon isotopic perturbation is again driven by the POC cycle (cf. Fig. 583 2c-d). The burial-input imbalance in the POC cycle is larger in the sensitivity experi-60/ ment with preferential phosphate leaching from sediments, as compared to the standard organic input step-change experiment. This leads to an amplified increase in the  $\delta^{13}C$ 586 signature of the reactive carbon pools (see the larger perturbation in  $\delta^{13}$ C in Fig. 4n,o). 587 At the same time, lower atmospheric  $CO_2$  and thus lower  $[CO_{2,aq}]$  lead to less fraction-588 ation during marine photosynthesis and an increase in  $\delta^{13}$ C of the POC burial flux. In 589 total, this leads to initially higher positive  $\delta^{13}$ C values in the atmosphere and ocean for 590 the sensitivity experiment with subsequent declines in differences (Fig. 4n; 6b,d). 591

592

#### 3.3 Scalability of Step-change Responses

Here, we address the linearity of responses in carbon and carbon isotopes to step-593 changes of different magnitudes (Fig. 6). In a linear system, the responses shown in Fig. 594 1 and 4 would be independent of the magnitude of the step-change. Then, the impact 595 of any evolution of changes in CaCO<sub>3</sub> or organic matter input could be exactly described 596 with the help of the responses or Green's functions shown in Fig. 1 and 4. While the car-597 bon cycle is known to be non-linear, near-linear behavior may still emerge within cer-598 tain limits of change. In Fig. 6 we compare the responses in  $CO_2$ , DIC, and their  $\delta^{13}C$ 599 signatures from eight step-change simulations where  $CaCO_3$  or organic matter input was 600 changed at nominal time t=0 over a wide range (-80% to +80%). All responses are nor-601 malized to a step increase in  $CaCO_3$  or decrease in organic matter input of 40% for com-602 parison with the experiments discussed above. The responses for steps within  $\pm 40\%$  are 603 visually highlighted and linked by blue (for  $CaCO_3$ ) and red (for organic matter) shad-604 ing. The responses for larger step changes are not further discussed. 605

Generally, linearity is higher for step-changes in the CaCO<sub>3</sub> input as compared to 606 organic matter input. Changes in DIC,  $\delta^{13}C_{atm}$ , and  $\delta^{13}C_{DIC}$  are near-linear in the case 607 of step-changes in CaCO<sub>3</sub> input, although non-linearities increase somewhat with time 608 by up to  $\pm 14-16\%$  relative to the overall change at 100 kyr (Fig. 6b,c,d). The largest 609 uncertainties are associated with the response in  $CO_2$ , where the mean response is -39 610 ppm, with a spread of  $\pm$  7 ppm ( $\pm$  18%) after 100 kyr and the range between +40% and 611 -40% CaCO<sub>3</sub> input. The response in CO<sub>2</sub> is larger for a reduction than an increase in 612  $CaCO_3$  input of equal magnitude (Fig. 6a). For step-changes in the organic input, near 613 linearity is found for changes in DIC (spread:  $\pm~7\%$  ) and, to a lesser extent, for atmo-614 spheric CO<sub>2</sub> ( $\pm 7$  ppm,  $\pm 14\%$ ),  $\delta^{13}C_{atm}$  ( $\pm 20\%$  relative to peak change), while the spread 615 covers 0.2 % (± 37% relative to peak change) for  $\delta^{13}C_{DIC}$  after 100 kyr and the range 616 between +40% and -40% organic matter input. 617

In summary, the assumption of a near-linear response appears reasonable for changes in organic matter and CaCO<sub>3</sub> input within  $\pm 40\%$  around the mean state. Thus, the response functions displayed in Fig. 1and 4 broadly characterize the response to changes in CaCO<sub>3</sub> and organic matter input, enabling the construction of a simple emulator to simulate responses to time-varying input in a cost-efficient and transparent way as done in section 4.2.

#### $_{624}$ 4 Discussion

State-of-the-art Earth System Models or Earth System Models of Intermediate Complexity that explicitly represent processes on a 3-dimensional spatial grid and sub-annual
time scales are computationally too expensive for simulations exceeding millions of years.
However, the characteristic spatio-temporal responses of such complex models may be
used to build cost-efficient substitute models (section 2.3) that account for the underlying spatio-temporal complexity of the involved processes. In the future, such models
may also include spatially explicit weathering or ice sheets. The Green's response func-

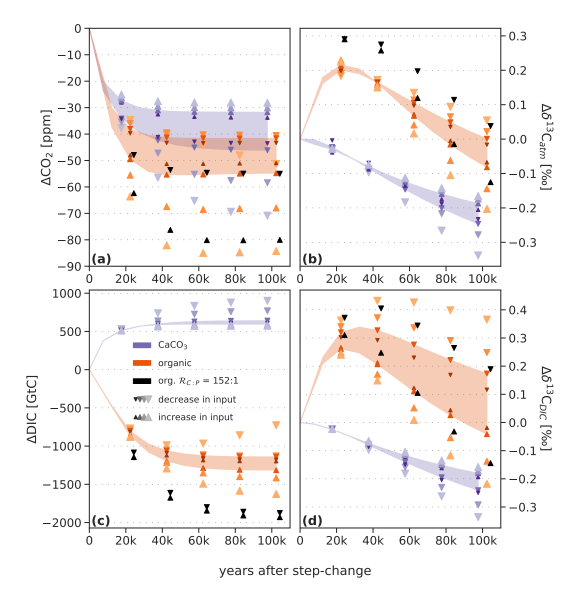


Figure 6. Normalized response functions. Change in (a)  $CO_2$ , (b)  $\delta^{13}C_{atm}$ , (c) DIC, and (d)  $\delta^{13}C_{DIC}$  to step-changes in the input of CaCO<sub>3</sub> (blue), organic material (red), and organic material with a different C:P ratio (black, see section 2.2) at different times. Step-changes range from -80 to 80% in 20% increments (size of the markers). Results are scaled to an increase of 40% in case of CaCO<sub>3</sub> and a decrease of 40% in case of organic material. Red and blue envelops indicate the range from +40% to -40% of changes in CaCO<sub>3</sub> and organic input. Triangles are slightly offset in x-direction for visibility and black triangles show results only for an in-/decrease of 40% in the organic input flux.

tions from the Bern3D Earth System Model of Intermediate Complexity for changes in
the net input of organic matter and CaCO<sub>3</sub> may also be viewed as a first step towards
bringing spatial scales to modelling the carbon cycle over periods of many millions of years.
In such long simulations, the atmosphere, ocean, land biosphere, and interactive sediments are usually taken as single reservoirs which differs from our approach, as we here
explicitly consider the transfer of carbon, alkalinity, and phosphorus between these reservoirs and within the ocean.

The response timescales and amplitudes to perturbations in the input from the litho-639 sphere are governed by the interplay between ocean circulation and air-sea exchange, nu-640 trient transport to the ocean surface, marine biological productivity and the export of 641 biogenic particles, and remineralization and redissolution of biogenic particles within the 642 water column and sediments. Spatial gradients within the ocean and ocean sediments 643 are important. For example, the removal rate of  $\delta^{13}$ C perturbations is significantly mod-644 ified by  $\delta^{13}$ C gradients within the ocean, influencing the isotopic composition of the burial 645 flux of organic and CaCO<sub>3</sub> particles in the lithosphere (Jeltsch-Thömmes & Joos, 2020). 646

The long timescales and the spatial complexity associated with burial-input im-647 balances and the carbon cycle is a challenge when investigating the glacial-interglacial 648 variations of the past few million years. Box models may not capture spatial gradients, 649 whereas glacial-interglacial simulations are computationally too expensive with weathering-650 sediment-enabled Earth System Models (Lacroix et al., 2020) and are even demanding 651 for Earth System Models of Intermediate Complexity (e.g. Menviel et al., 2012; Colbourn 652 et al., 2013; Ganopolski & Brovkin, 2017; Willeit et al., 2022). Cost-efficient substitute 653 models, representing faithfully the response of spatially-resolved models, provide an al-654 ternative. 655

In brief, evaluating the response to a step-like change reveals underlying system dynamics and spatio-temporal response characteristics. This is useful for model-model comparison and for understanding the model responses to a perturbation. Further, the response can be used to build a cost-efficient substitute model (or emulator) to explore different forcing histories over long time scales (see section 4.2).

#### 661

## 4.1 Uncertainties and Limitations

We apply an Earth system model of intermediate complexity. The use of the Bern3D 662 model enables us to conduct 600 kyr long simulations considering fluid dynamics and bio-663 geochemical processes in a spatially-resolved 3-dimensional setting. This is a progress 664 compared to box models or 2-dimensional ocean models, typically applied for such long 665 time scales. However, many processes are still represented by simplified parameteriza-666 tions and some of these, as, for example, the dissolution rate constants in the marine sed-667 iments or the remineralization of organic matter in the water column, rely on global uni-668 form parameter choices. 669

Another limitation relates to the stoichiometric ratios used in the model. The ques-670 tion of the C:P ratio in marine sediments and preferential P leaching to the ocean is dis-671 cussed in the literature (e.g., Ingall & Jahnke, 1994; Delaney, 1998; L. D. Anderson et 672 al., 2001). Under low oxygen conditions, for example, P is preferentially regenerated from 673 marine sediments, (e.g., Ingall & Jahnke, 1994). This could lead to a positive feedback 674 loop where more P is brought to the ocean surface enhancing primary production and 675 organic matter export, and, in turn, enhancing organic matter transport to and reminer-676 alization at depth, further depleting bottom water oxygen through consumption of oxy-677 678 gen during remineralization (e.g., Van Cappellen & Ingall, 1994; Wallmann, 2010; Palastanga et al., 2011; Niemeyer et al., 2016; Watson, 2016; Kemena et al., 2019). Release 679 of additional P from sediments could have a strong impact on  $\delta^{13}$ C (see e.g., Tschumi 680 et al., 2011, for a step-increase in the ocean's P inventory). In the current setup of the 681 Bern3D model, we assume constant Redfield ratios in the ocean and sediment. Further, 682

in the standard step-change experiment of the organic input flux, the same constant C:P 683 ratio of 117:1 is applied. To address this limitation, we conducted an additional exper-684 iment, where 30% of the P entering marine sediments in the model is immediately re-685 leased back to the ocean (see section 2.1). While the quantitative results differ, the qualitative response is similar for the experiment with and without this idealized enhanced 687 P leaching. Model formulations could be revised in the future for varying C:P ratios, pref-688 erential P leakage from sediments under low oxygen concentrations as done in the HAMOCC 689 and UVic models (e.g., Palastanga et al., 2011; Niemeyer et al., 2016; Kemena et al., 2019), 690 and variable stoichiometric ratios for biological production, e.g., Matsumoto et al. (2020). 691 Lacroix et al. (2020) further discuss the question of C:P ratios of riverine weathering in-692 put of organic material. While terrestrial dissolved organic matter exhibits very high C:P 693 ratios of up to 2583:1 (Meybeck, 1982; Compton et al., 2000), the ratio is uncertain in 694 particulate organic matter (POM), with a range of 56-499 (Meybeck, 1982; Ramirez & 695 Rose, 1992; Compton et al., 2000). In the experiments here, weathering of terrestrial dis-696 solved organic material is not considered and the input flux  $F_{org}$  is assumed to originate 697 from previously buried organic matter. In line with Lacroix et al. (2020), the C:P ra-698 tio of this  $F_{org}$  flux from the litosphere is the same as of oceanic POM. 699

For the evolution of  $\delta^{13}$ C, the organic matter cycle and fractionation during ma-700 rine photosynthesis plays an important role. Here, this fractionation is described using 701 the empirical relationship with dissolved  $CO_2$  by Freeman and Hayes (1992). Addition-702 ally, there are other environmental factors besides dissolved  $CO_2$  that might affect frac-703 tionation during marine photosynthesis (Goericke & Fry, 1994). Jahn et al. (2015) im-704 plemented three parameterizations in their ocean model. One where fractionation varies 705 with dissolved  $CO_2$  and two also considering phytoplankton growth rates. Overall, all 706 three parameterizations lead to the expected pattern of high  $\delta^{13}C_{DIC}$  in surface waters 707 and low  $\delta^{13}C_{DIC}$  values in water that has been out of contact with the atmosphere for 708 a long time. Liu et al. (2021) found good  $\delta^{13}$ C data-model agreement for discrimination 709 varying with dissolved  $CO_2$  according to Popp et al. (1989), while the agreement was 710 less favorable with a parameterization additionally varying with local phytoplankton growth 711 rates (Laws et al., 1995). These findings lend indirect justification for applying the pa-712 rameterization of Freeman and Hayes (1992), which is similar to that of Popp et al. (1989). 713 While the application of alternative parameterizations would somewhat affect the quan-714 titative findings of this study, the main messages would likely remain unchanged. 715

716 717

## 4.2 Scenarios for the past 800,000 years: Application of the Green's Function Substitute Model

We apply the Green's function emulator (Eq. 9; section 2.3) to easily investigate 718 different scenarios for changes in CaCO<sub>3</sub> weathering and organic matter input over the 719 past 800,000 years. We use the mean of the normalized responses to the +40 and -40%720 step-change in input for  $CaCO_3$  and organic matter, respectively. This emulator is an 721 approximation and non-linear interactions with carbon cycle changes other than those 722 related to changes in weathering input are not taken into account. The aim of this ex-723 ercise is not to provide conclusive answers how past changes in input fluxes affected at-724 mospheric  $CO_2$  and the carbon cycle, but rather to illustrate the usefulness of the Green's 725 function approach to explore different scenarios and hypotheses. 726

In the first scenario, we apply the 0-dimensional version of the RokGeM v0.9 model 727 (Colbourn et al., 2013) where  $CaCO_3$  input is a function of atmospheric  $CO_2$  (Bereiter 728 et al., 2015) and global mean surface air temperature (SAT). SAT evolution is computed 729 by linearly scaling the Antarctic temperature reconstruction of Jouzel et al. (2007)), as-730 suming a difference in SAT between preindustrial (PI) and Last Glacial Maximum (LGM) 731 of 5.5 K. RokGeM yields  $CaCO_3$  weathering to be about 40% lower at LGM than PI (Fig. 732 7, turquoise). This is in contrast to Börker et al. (2020) who suggest CaCO<sub>3</sub> weather-733 ing to be about 30% higher at LGM than PI. In our second scenario, we linearly scale 734

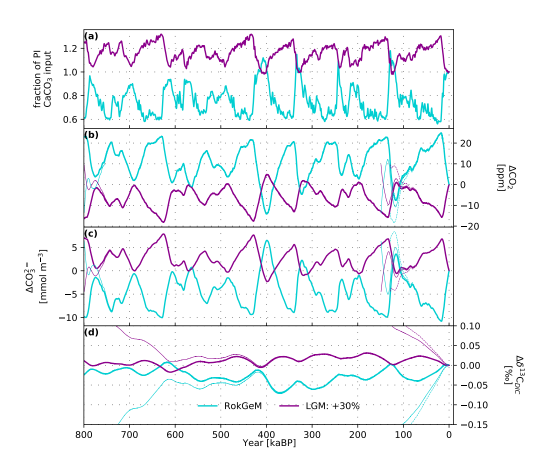


Figure 7. Two contrasting literature-based CaCO<sub>3</sub> weathering scenarios. (a) Prescribed changes in CaCO<sub>3</sub> weathering input (magenta: +30% in LGM, based on reconstructions of CaCO<sub>3</sub> weathering by Börker et al. (2020) and scaled with  $\delta^{18}$ O (Lisiecki & Raymo, 2005); Turquoise: as calculated with the Rock Geochemical Model RokGeM v0.9 (Colbourn et al., 2013), where maximum LGM cooling of 5.5 K is scaled with the Antarctic temperature reconstruction by Jouzel et al. (2007)). Changes (relative to PI) in (b) atmospheric CO<sub>2</sub>, (c) global mean  $CO_3^{2-}$ , and (d) global mean  $\delta^{13}C_{DIC}$  as emulated using the responses of the step-change experiment. The first glacial cycles was repeated twice before starting the standard run at 800 kyr (thick lines). Sensitivity simulations, illustrating the drift associated with a "old start", were started directly at at 800, 150, and 130 kaBP (thin lines); results show that initial conditions affect simulated CO<sub>2</sub> for several ten thousand years and  $\delta^{13}C$  for several hundred thousand years.

## <sup>735</sup> the $\delta^{18}$ O record of Lisiecki and Raymo (2005) to yield an LGM-PI difference of +30% <sup>736</sup> in CaCO<sub>3</sub> input.

The scenario based on Börker et al. (2020) yields LGM minima of up to  $\sim 15$  ppm 737 lower than at PI. In contrast, the scenario based on RokGem yields up to more than 20 738 ppm higher  $CO_2$  during glacials (Fig. 7a,b). Overall, differences between the two literature-739 based scenarios are up to 35 ppm or about a third of the glacial-interglacial  $CO_2$  am-740 plitude. Similarly, changes in  $CO_3^{2-}$  are opposite and differences between the two sce-741 narios are up to 18 mmol m<sup>-3</sup>. Generally,  $CO_2$  and  $CO_3^{2-}$  follow changes in weather-742 ing without much delay and capture some multi-millennial features of the forcing, while 743  $\delta^{13}C_{DIC}$  shows a strongly smoothed signal. Variations in  $\delta^{13}C_{DIC}$  are on the order < 0.05744 ‰ and the timing of maxima/minima does not directly mirror changes in the forcing 745 (Fig. 7d). 746

In the third scenario, we adopt the "shelf-weathering" hypothesis by Wallmann et 747 al. (2016), where input of organic-derived material to the ocean is elevated during pe-748 riods of low sea level. Taking their PI-LGM change at face value (see panel (h), Fig. 3 749 in Wallmann et al., 2016) translates into a  $\sim 10\%$  change of the total organic input in 750 the Bern3D model. This PI-LGM difference in forcing is then scaled with  $\delta^{18}$ O (Lisiecki 751 & Raymo, 2005) to yield a forcing history of the past 800 kyr (Fig. 8a). This yields  $CO_2$ 752 changes of up to  $\sim 10$  ppm with higher CO<sub>2</sub> during glacials than interglacials (Fig. 8b). 753 Changes in  $\delta^{13}C_{DIC}$  are on the order of 0.05 % with lower values during glacials. This 754 means that about 15% of the observed PI-LGM  $\delta^{13}C_{\text{DIC}}$  change (C. D. Peterson et al., 755 2014) could be attributed to changes in input from shelves. Contrary to changes in the 756  $CaCO_3$  weathering, the signal in  $\Delta \delta^{13}C_{DIC}$  is less smoothed and the timing reflects the 757 forcing history (Fig. 8a,d). This is likely a results from the initial strong perturbation 758 in  $\delta^{13}C_{\text{DIC}}$  resulting from changes in the input  $\delta^{13}C$  signature (see Fig. 3n, 5d-f and sec-759 tion 3.2). 760

761 762

## 4.3 How to Initialize Earth System Models: The Glacial-Interglacial Cold Start Problem

The long adjustment timescales of carbon and  $\delta^{13}$ C to perturbations in input-burial 763 imbalances pose a serious challenge for model initialisation. Initial conditions affect the 764 evolution of  $CO_2$  and  $\delta^{13}C$  over several ten thousand and hundred thousand years, re-765 spectively. In the standard setup of the emulator, the forcing history of the first glacial 766 cycle is repeated twice, before starting the simulation at the nominal year 800 kaBP. We 767 illustrate the impact and potential biases of such a "cold start" (Hasselmann et al., 1993) 768 by starting the emulator at 800 kaBP, the beginning of our scenarios, at 150 kaBP, i.e. 769 close to glacial maximum conditions, and at 130 kaBP, near the beginning of the last in-770 terglacial. 771

The initial trends in  $\text{CO}_2$  and  $\text{CO}_3^{2-}$  deviate strongly from standard results during many millennia (thin versus thick lines in Fig. 7) for all three sensitivity runs. This initial drift obscures the signals found in the standard run. For  $\delta^{13}$ C, deviations extend as expected over several hundred thousand years. Results for  $\delta^{13}$ C of the simulations started at 150 and 130 kaBP are useless and highly misleading. The simulated trend is about four times larger than the signal simulated in the standard setup.

The initial adjustment after a cold start is an intrinsic property of the Earth system as implemented in the parent Bern3D model (see Figs 1 and 4) and inherited by the emulator. The results highlight that models used to simulate the last deglacial or glacialinterglacial cycles and featuring input and burial fluxes must be carefully initialised to avoid wrong results and interpretations.

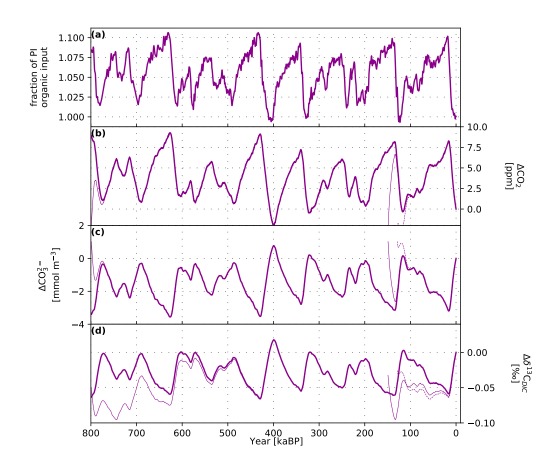


Figure 8. As Fig. 7 but for a scenario with varying input of organic-derived material (P:C:ALK=1:117:-17 and  $\delta^{13}$ C=-20.42 ‰), based on (Wallmann, 2014; Wallmann et al., 2016)

## 783 5 Summary and Conclusions

We conducted idealized simulations applying a step-change in the input fluxes of 784 DIC, ALK, P, and  $\delta^{13}$ C from the lithosphere to the ocean. We assumed step changes in 785 a conceptual input flux releasing the elements of former organic matter ( $F_{org}$ ; P:ALK:DIC=1:-786 17:-117 with  $\delta^{13}C=-20.42$  ‰) or from CaCO<sub>3</sub> weathering ( $F_{CaCO_3}$ ; ALK:DIC=2:1 with 787  $\delta^{13}C=2.88$  %). We analyzed the response in atmospheric CO<sub>2</sub>, marine parameters (DIC, 788 ALK,  $CO_3^{--}$ , pH, saturation horizon, export production), sedimentary and burial fluxes, 789 and in  $\delta^{13}$ C of various carbon reservoirs and fluxes in the Bern3D model. The two stan-790 dard simulations extend over 600,000 years and  $F_{CaCO_3}$  is increased by 40% and  $F_{org}$ 791 reduced by 40%, respectively. 792

Atmospheric CO<sub>2</sub> decreased exponentially by about 30 ppm in both simulations. The e-folding adjustment time scale for CO<sub>2</sub> and other carbon cycle parameters is ~7,500 years for the step change in  $F_{org}$  and ~9,900 years for the step in  $F_{CaCO_3}$ . The difference in time scales reflects differences in underlying processes. While sedimentary CaCO<sub>3</sub> compensation is the controlling process after a change in CaCO<sub>3</sub> weathering, a reduction in organic matter export and burial and the marine phosphorus inventory govern the carbon response after the reduction in  $F_{org}$  and P input to the ocean.

For the increase in  $F_{CaCO_3}$ ,  $\delta^{13}C$  of atmospheric CO<sub>2</sub>, DIC, and CaCO<sub>3</sub> decreases gradually with a time scale of ~57,300 years to stabilize 0.35‰ lower than before the step in  $F_{CaCO_3}$ . The adjustment in  $\delta^{13}C$  is controlled by burial-input imbalances in both the cycling of CaCO<sub>3</sub> and organic matter. The decrease in CO<sub>2</sub> in the surface ocean causes less fractionation during photosynthesis and an increase in  $\delta^{13}C$  of organic matter. In turn, a larger burial of  $\delta^{13}C$  is mediated by organic matter burial which is at the new equilibrium compensated by an opposite perturbation in the  $\delta^{13}C$  removal by CaCO<sub>3</sub> burial.

For the reduction in  $F_{org}$ ,  $\delta^{13}$ C of atmospheric CO<sub>2</sub>, DIC and CaCO<sub>3</sub> increases ini-808 tially by about 0.3%. This increase is in response to the reduced input of isotopically 809 light carbon and follows a time scale of about  $\sim 5,800$  years. At peaking, the reduced in-810 put is compensated by a reduced burial of organic matter. Afterward,  $\delta^{13}C$  decreases 811 with an adjustment time scale of  $\sim 110,000$  to stabilize only after several hundred thou-812 sand years. This decrease is, similar as for the change in  $CaCO_3$  weathering, related to 813 the positive shift in  $\delta^{13}$ C of marine organic matter due to lower CO<sub>2</sub>. The reduction in 814 the export and the remineralization fluxes of organic matter and their changes in iso-815 topic signature cause distinct spatial patterns within the ocean. The surface-to-thermocline 816 gradient is reduced by about 0.25%. 817

The application of a step change in input allows for probing the so-called Green's 818 function response. The normalized responses are used to build a simple, cost-efficient em-819 ulator. Simulating one million years takes a few seconds with the emulator, while it would 820 take about three months using Bern3D. The emulator is applied in a range of literature-821 based scenarios for the past 800,000 years to estimate carbon cycle changes to plausi-822 ble time-varying input fluxes from the lithosphere or shelves. These reveal differences 823 in simulated  $CO_2$  of up to a third of the reconstructed glacial-interglacial amplitude, and 824 differences in  $CO_3^{--}$ , and small (0.05 to 0.1 %), but non-negligible, changes in  $\delta^{13}C$ . 825 Our emulator, is probably most useful for periods with a close-to-modern continental con-826 figuration. However, Green's function emulators could also be valuable for studying Earth's 827 long history. 828

The long adjustment time scales of ten thousand years, and even a hundred thousand years for  $\delta^{13}$ C, pose a serious challenge for sediment-enabled model simulations. The model needs to be initialized carefully to avoid model drift obscuring the responses to any applied forcing, rendering results misleading and useless, as exemplified in coldstart simulations with the emulator. <sup>834</sup> Our results highlight the importance to consider weathering-burial (input-output) <sup>835</sup> imbalances and interactions between the atmosphere-ocean-land biosphere system and <sup>836</sup> reactive ocean sediments in modeling studies that investigate CO<sub>2</sub> and  $\delta^{13}$ C on multi-<sup>837</sup> millennial and longer timescales. A thorough understanding of the mechanisms affect-<sup>838</sup> ing atmospheric CO<sub>2</sub>,  $\delta^{13}$ C<sub>DIC</sub>, CO<sub>3</sub><sup>2-</sup> and other carbon cycle parameters will help in the <sup>839</sup> interpretation of paleo-records. Already small imbalances in the burial-input cycle can <sup>840</sup> have large effects on carbon and carbon isotopes in the Earth system and are thus likely <sup>841</sup> candidates to have contributed to the reconstructed glacial-interglacial variations.

#### Acknowledgments

The authors acknowledge funding from the Swiss National Science Foundation (No. 200020\_200511) 843 and the European Union's Horizon 2020 research and innovation program under grant 844 agreement No 820989 (project COMFORT, Our common future ocean in the Earth system-845 quantifying coupled cycles of carbon, oxygen, and nutrients for determining and achiev-846 ing safe operating spaces with respect to tipping points). The work reflects only the au-847 thors' view; the European Commission and their executive agency are not responsible 848 for any use that may be made of the information the work contains. Markus Adloff and 849 Fabrice Lacroix provided valuable feedback that helped to improve the manuscript. 850

Data used for the figures and necessary for construction of the emulator will be made available upon peer-review completion on zenodo.org and accessible via doi. Additional data is available upon request from the corresponding author. During peer-review, data can be accessed under the following share-link: https://cloud.climate.unibe.ch/s/ nfGTZRmkrs5SAyF

856

866

867

868

The authors declare that they have no conflict of interest.

## <sup>857</sup> References

- Anderson, L. A., & Sarmiento, J. L. (1994). Redfield ratios of remineralization
   determined by nutrient data analysis. *Global Biogeochemical Cycles*, 8(1), 65–
   80. Retrieved from https://agupubs.onlinelibrary.wiley.com/doi/abs/10
   .1029/93GB03318 doi: 10.1029/93GB03318
- Anderson, L. D., Delaney, M. L., & Faul, K. L. (2001). Carbon to phosphorus ratios
   in sediments: Implications for nutrient cycling. *Global Biogeochemical Cycles*,
   *15*(1), 65–79. Retrieved from http://doi.wiley.com/10.1029/2000GB001270
   doi: 10.1029/2000GB001270
  - Archer, D., Kheshgi, H., & Maier-Reimer, E. (1997). Multiple timescales for neutralization of fossil fuel CO<sub>2</sub>. Geophysical Research Letters, 24(4), 405–408. doi: 10.1029/97GL00168
- Archer, D., Kheshgi, H., & Maier-Reimer, E. (1998). Dynamics of fossil fuel CO<sub>2</sub>
   neutralization by marine CaCO3. *Global Biogeochemical Cycles*, 12(2), 259–
   276. doi: 10.1029/98GB00744
- 872Archer, D., & Maier-Reimer, E.(1994).Effect of deep-sea sedimentary calcite873preservation on atmospheric CO2 concentration.Nature, 367(6460), 260–263.874doi: 10.1038/367260a0
- Archer, D., Winguth, A., Lea, D., & Mahowald, N. (2000). What caused the glacial/interglacial atmospheric pCO<sub>2</sub> cycles? *Reviews of Geophysics*, 38(2), 159–189.
- 878Bastiaansen, R., Dijkstra, H. A., & Heydt, A. S. v. d.(2021).Projections879of the Transient State-Dependency of Climate Feedbacks.Geophysi-880cal Research Letters, 48(20), e2021GL094670.Retrieved from https://881agupubs.onlinelibrary.wiley.com/doi/abs/10.1029/2021GL094670
- (e2021GL094670 2021GL094670) doi: https://doi.org/10.1029/2021GL094670
- Battaglia, G., Steinacher, M., & Joos, F. (2016). A probabilistic assessment of cal cium carbonate export and dissolution in the modern ocean. *Biogeosciences*,
   13(9), 2823–2848. doi: 10.5194/bg-13-2823-2016
- Bereiter, B., Eggleston, S., Schmitt, J., Nehrbass-Ahles, C., Stocker, T. F., Fischer,
   H., ... Chappellaz, J. (2015). Revision of the EPICA Dome C CO<sub>2</sub> record
   from 800 to 600 kyr before present. *Geophysical Research Letters*, 42(2),
   542–549. doi: 10.1002/2014GL061957
- Bergman, N., Lenton, T., & Watson, A. (2004). COPSE: A new model of biogeochemical cycling over Phanerozoic time. American Journal of Science, 304(5), 397-437. doi: 10.2475/ajs.304.5.397
- Berner, R. A. (1990). Atmospheric Carbon Dioxide Levels Over Phanero-

894	zoic Time. Science, 249(4975), 1382-1386. Retrieved from https://
895	www.science.org/doi/abs/10.1126/science.249.4975.1382 doi:
896	10.1126/science.249.4975.1382
897	Berner, R. A. (2006). GEOCARBSULF: A combined model for Phanerozoic atmo-
898	spheric $O_2$ and $CO_2$ . Geochimica et Cosmochimica Acta, $70(23)$ , 5653-5664.
899	Retrieved from https://www.sciencedirect.com/science/article/pii/
900	S0016703706002031 (A Special Issue Dedicated to Robert A. Berner) doi:
901	https://doi.org/10.1016/j.gca.2005.11.032
902	Broecker, W. S. (1970). A boundary condition on the evolution of atmospheric
903	oxygen. Journal of Geophysical Research (1896-1977), 75(18), 3553-3557.
904	Retrieved from https://agupubs.onlinelibrary.wiley.com/doi/abs/
905	10.1029/JC075i018p03553 doi: https://doi.org/10.1029/JC075i018p03553
906	Broecker, W. S., & Peng, TH. (1987). The role of CaCO3 compensation in the
907	glacial to interglacial atmospheric CO <sub>2</sub> change. Global Biogeochemical Cycles,
908	1(1), 15-29. doi: $10.1029/GB001i001p00015$
909	Broecker, W. S., & Peng, TH. (1989). The cause of the glacial to interglacial atmo-
910	spheric CO <sub>2</sub> change: A polar alkalinity hypothesis. Global Biogeochemical Cy-
911	$cles,\ \Im(3),\ 215 ext{}239.$
912	Brovkin, V., Ganopolski, A., Archer, D., & Munhoven, G. (2012). Glacial CO <sub>2</sub> cy-
913	cle as a succession of key physical and biogeochemical processes. Climate of the
914	Past, 8(1), 251-264. doi: 10.5194/cp-8-251-2012
915	Börker, J., Hartmann, J., Amann, T., Romero-Mujalli, G., Moosdorf, N., & Jenkins,
916	C. (2020). Chemical Weathering of Loess and Its Contribution to Global
917	Alkalinity Fluxes to the Coastal Zone During the Last Glacial Maximum,
918	Mid-Holocene, and Present. Geochemistry, Geophysics, Geosystems, 21(7),
919	e2020GC008922. Retrieved from https://agupubs.onlinelibrary.wiley
920	.com/doi/abs/10.1029/2020GC008922 (e2020GC008922 2020GC008922) doi:
921	https://doi.org/10.1029/2020GC008922
922	Cartapanis, O., Bianchi, D., Jaccard, S., & Galbraith, E. D. (2016). Global pulses
923	of organic carbon burial in deep-sea sediments during glacial maxima. Nature
924	communications, 7, 10796. doi: 10.1038/ncomms10796
925	Cartapanis, O., Galbraith, E. D., Bianchi, D., & Jaccard, S. L. (2018). Carbon
926	burial in deep-sea sediment and implications for oceanic inventories of car-
927	bon and alkalinity over the last glacial cycle. Climate of the Past, $14(11)$ , 1810, 1850, doi: 10.5104/cp.14.1810, 2018
928	1819–1850. doi: $10.5194/cp-14-1819-2018$
929	Caves, J. K., Jost, A. B., Lau, K. V., & Maher, K. (2016). Cenozoic carbon cycle
930	imbalances and a variable weathering feedback. Earth and Planetary Science
931	Letters, 450, 152-163. doi: {10.1016/j.epsl.2016.06.035} Clark, P. U., Archer, D., Pollard, D., Blum, J. D., Rial, J. A., Brovkin, V., Roy,
932	M. (2006). The middle Pleistocene transition: characteristics, mechanisms, and
933	implications for long-term changes in atmospheric pCO <sub>2</sub> . Quaternary Science
934 935	<i>Reviews</i> , 25(23-24), 3150–3184. doi: 10.1016/j.quascirev.2006.07.008
	Colbourn, G., Ridgwell, A., & Lenton, T. M. (2013). The Rock Geochemical Model
936 937	(RokGeM) v0.9. Geoscientific Model Development, 6(5), 1543–1573. doi: 10
938	.5194/gmd-6-1543-2013
939	Colbourn, G., Ridgwell, A., & Lenton, T. M. (2015). The time scale of the silicate
940	weathering negative feedback on atmospheric $CO_2$ . Global Biogeochemical Cy-
941	cles, 29(5), 583-596. doi: 10.1002/2014GB005054
942	Compton, J., Mallinson, D., Glenn, C., Filippelli, G., Föllmi, K., Shields, G., &
943	Zanin, Y. (2000). Variations in the global phosphorus cycle. In <i>Marine</i>
944	Authigenesis: From Global to Microbial (p. 21-33). Wiley-Blackwell.
945	Delaney, M. L. (1998). Phosphorus accumulation in marine sediments and
946	the oceanic phosphorus cycle. Global Biogeochemical Cycles, 12(4), 563–
947	572. Retrieved from http://doi.wiley.com/10.1029/98GB02263 doi:
948	10.1029/98GB02263

949	Edwards, N. R., Willmott, A. J., & Killworth, P. D. (1998). On the Role of Topog-
950	raphy and Wind Stress on the Stability of the Thermohaline Circulation. Jour-
951	nal of Physical Oceanography, 28(5), 756–778. doi: 10.1175/1520-0485(1998)
952	$028\langle 0756:OTROTA\rangle 2.0.CO;2$
953	Eggleston, S., Schmitt, J., Bereiter, B., Schneider, R., & Fischer, H. (2016). Evolu-
954	tion of the stable carbon isotope composition of atmospheric $CO_2$ over the last
955	glacial cycle. Paleoceanography, 31(3), 434–452. doi: 10.1002/2015PA002874
956	Emerson, S., & Archer, D. (1992). Glacial carbonate dissolution cycles and atmo-
	spheric pCO <sub>2</sub> : A view from the ocean bottom. <i>Paleoceanography</i> , $7(3)$ , 319–
957	331. doi: 10.1029/92PA00773
958	Emerson, S., & Bender, M. (1981). Carbonate Fluxes at the Sediment-Water Inter-
959	face of the Deep-Sea - Calcium-Carbonate Preservation. Journal of Marine Re-
960	-
961	search, $39(1)$ , 139-162.
962	Feely, R. A., Sabine, C. L., Lee, K., Berelson, W., Kleypas, J., Fabry, V. J., &
963	Millero, F. J. (2004). Impact of Anthropogenic $CO_2$ on the CaCO3 System in
964	the Oceans. <i>Science</i> , <i>305</i> , 362–366.
965	Freeman, K. H., & Hayes, J. M. (1992). Fractionation of carbon isotopes by phy-
966	toplankton and estimates of ancient $CO_2$ levels. Global Biogeochemical Cycles,
967	6(2), 185-198. doi: 10.1029/92GB00190
968	Ganopolski, A., & Brovkin, V. (2017). Simulation of climate, ice sheets and $CO_2$
969	evolution during the last four glacial cycles with an Earth system model
970	of intermediate complexity. Climate of the Past, $13(12)$ , $1695-1716$ . doi:
971	10.5194/cp-13-1695-2017
972	Goericke, R., & Fry, B. (1994). Variations of marine plankton $\delta^{13}$ C with latitude,
973	temperature, and dissolved $CO_2$ in the world ocean. Global Biogeochemi-
974	cal Cycles, $\delta(1)$ , 85-90. Retrieved from https://agupubs.onlinelibrary
975	.wiley.com/doi/abs/10.1029/93GB03272 doi: https://doi.org/10.1029/
976	93GB03272
977	Goodwin, P., & Ridgwell, A. (2010). Ocean-atmosphere partitioning of anthro-
978	pogenic carbon dioxide on multimillennial timescales. Global Biogeochemical
979	Cycles, 24(2), GB2014,. Retrieved from http://doi.wiley.com/10.1029/
980	2008GB003449 doi: 10.1029/2008GB003449
981	Griffies, S. M. (1998). The Gent–McWilliams Skew Flux. Journal of Physi-
982	cal Oceanography, $28(5)$ , $831-841$ . doi: $10.1175/1520-0485(1998)028(0831)$ :
983	TGMSF $2.0.CO;2$
984	Hartmann, J., Moosdorf, N., Lauerwald, R., Hinderer, M., & West, A. J. (2014).
985	Global chemical weathering and associated P-release - The role of lithol-
986	ogy, temperature and soil properties. <i>Chemical Geology</i> , 363, 145-163. doi:
987	10.1016/j.chemgeo.2013.10.025
988	Hasselmann, K., Sausen, R., Maier-Reimer, E., & Voss, R. (1993). On the cold start
989	problem in transient simulations with coupled atmosphere-ocean models. <i>Cli</i> -
990	mate Dynamics, 9(2), 53-61. doi: 10.1007/BF00210008
991	Hayes, J. M., & Waldbauer, J. R. (2006). The carbon cycle and associated redox
992	processes through time. Philosophical Transactions of the Royal Society B -
993	Biological Sciences, 361(1470), 931-950. (Conference on Major Steps in Cell
994	Evolution - Palaeontological, Molecular and Cellular Evidence of their Timing
995	and Global Effects, Royal Soc, London, ENGLAND, SEP 26-27, 2005) doi:
996	10.1098/rstb.2006.1840
	Heinze, C., Maier-Reimer, E., Winguth, A. M. E., & Archer, D. (1999). A global
997 998	oceanic sediment model for long-term climate studies. Global Biogeochemical
998	Cycles, 13(1), 221-250. doi: 10.1029/98GB02812
1000	Hoogakker, B. A., Rohling, E. J., Palmer, M. R., Tyrrell, T., & Rothwell, R. G.
1000	(2006). Underlying causes for long-term global ocean $\delta$ 13C fluctuations over
1001	
	the last 1.20 WVC. Fath and Planelary Science Letters 248(1-2) 13-29 our
1003	the last 1.20 Myr. <i>Earth and Planetary Science Letters</i> , 248(1-2), 15–29. doi: 10.1016/j.epsl.2006.05.007

1004	Hooss, G., Voss, R., Hasselmann, K., Maier-Reimer, E., & Joos, F. (2001). A
1005	nonlinear impulse response model of the coupled carbon cycle-climate system
1006	(NICCS). Climate Dynamics, 18(3-4), 189–202. doi: 10.1007/s003820100170
1007	Huybers, P., & Langmuir, C. (2009). Feedback between deglaciation, volcanism, and
1008	atmospheric CO <sub>2</sub> . Earth and Planetary Science Letters, $286(3-4)$ , 479–491.
1009	doi: 10.1016/j.epsl.2009.07.014
1010	Ingall, E., & Jahnke, R. (1994). Evidence for enhanced phosphorus regenera-
1011	tion from marine sediments overlain by oxygen depleted waters. Geochim-
1012	ica et Cosmochimica Acta, 58(11), 2571–2575. Retrieved from https://
1013	linkinghub.elsevier.com/retrieve/pii/0016703794900337 doi:
1014	10.1016/0016-7037(94)90033-7
1015	Isson, T. T., Planavsky, N. J., Coogan, L. A., Stewart, E. M., Ague, J. J., Bolton,
1016	E. W., Kump, L. R. (2020). Evolution of the Global Carbon Cycle and
1017	Climate Regulation on Earth. Global Biogeochemical Cycles, $34(2)$ . doi:
1018	10.1029/2018GB006061
1019	Jahn, A., Lindsay, K., Giraud, X., Gruber, N., Otto-Bliesner, B. L., Liu, Z., &
1020	Brady, E. C. (2015). Carbon isotopes in the ocean model of the Community
1021	Earth System Model (CESM1). Geoscientific Model Development, 8(8), 2419– 2424. Betriaud from https://cmd.companyi.cug.org/onticleg/8/2410/2015/
1022	2434. Retrieved from https://gmd.copernicus.org/articles/8/2419/2015/ doi: 10.5104/gmd.8.2410.2015
1023	doi: 10.5194/gmd-8-2419-2015 Jeltsch-Thömmes, A., Battaglia, G., Cartapanis, O., Jaccard, S. L., & Joos, F.
1024	(2019). Low terrestrial carbon storage at the Last Glacial Maximum: con-
1025	straints from multi-proxy data. Climate of the Past, 15(2), 849–879. doi:
1026 1027	10.5194/cp-15-849-2019
	Jeltsch-Thömmes, A., & Joos, F. (2020). Modeling the evolution of pulse-like per-
1028 1029	turbations in atmospheric carbon and carbon isotopes: the role of weathering-
1029	sedimentation imbalances. Climate of the Past, $16(2)$ , $423-451$ . Retrieved from
1030	https://www.clim-past.net/16/423/2020/ doi: 10.5194/cp-16-423-2020
1032	Joos, F., & Bruno, M. (1996). Pulse response functions are cost-efficient tools
1032	to model the link between carbon emissions, atmospheric $CO_2$ and global
1034	warming. Physics and Chemistry of The Earth, 21 (5-6), 471–476. doi:
1035	10.1016/S0079-1946(97)81144-5
1036	Joos, F., Prentice, I. C., Sitch, S., Meyer, R., Hooss, G., Plattner, Gk., Hassel-
1037	mann, K. (2001). Global warming feedbacks on terrestrial carbon uptake under
1038	the Intergovernmental Panel on Climate Change (IPCC) Emission Scenarios.
1039	Global Biogeochemical Cycles, 15(4), 891–907. doi: 10.1029/2000GB001375
1040	Joos, F., Roth, R., Fuglestvedt, J. S., Peters, G. P., Enting, I. G., Von Bloh, W.,
1041	Weaver, A. J. (2013). Carbon dioxide and climate impulse response
1042	functions for the computation of greenhouse gas metrics: A multi-model
1043	analysis. Atmospheric Chemistry and Physics, $13(5)$ , $2793-2825$ . doi:
1044	10.5194/acp-13-2793-2013
1045	Jouzel, J., Masson-Delmotte, V., Cattani, O., Dreyfus, G., Falourd, S., Hoffmann,
1046	G., Wolff, E. W. (2007). Orbital and Millennial Antarctic Climate Vari-
1047	ability over the Past 800,000 Years. Science, 317(5839), 793–796. Retrieved
1048	from https://www.sciencemag.org/lookup/doi/10.1126/science.1141038
1049	doi: 10.1126/science.1141038
1050	Kalnay, E., Kanamitsu, M., Kistler, R., Collins, W., Deaven, D., Gandin, L.,
1051	Joseph, D. (1996). The ncep/ncar 40-year reanalysis project. Bulletin of the American Maternalogical Society $27(2)$ , $427$ , $472$ , Detriving from
1052	of the American Meteorological Society, 77(3), 437 - 472. Retrieved from
1053	https://journals.ametsoc.org/view/journals/bams/77/3/1520-0477
1054	_1996_077_0437_tnyrp_2_0_co_2.xml doi: 10.1175/1520-0477(1996)077(0437: TNYRP)2.0.CO;2
1055	Kasting, J. F. (2019). The Goldilocks Planet? How Silicate Weathering Maintains
1056	Earth "Just Right". Elements, 15(4), 235-240. doi: 10.2138/gselements.15.4
1057 1058	.235

1059	Kemena, T. P., Landolfi, A., Oschlies, A., Wallmann, K., & Dale, A. W. (2019).
1060	Ocean phosphorus inventory: large uncertainties in future projections
1061	on millennial timescales and their consequences for ocean deoxygena-
1062	tion. Earth System Dynamics, 10(3), 539–553. Retrieved from https://
1063	www.earth-syst-dynam.net/10/539/2019/ doi: $10.5194/esd-10-539-2019$
1064	Komar, N., & Zeebe, R. E. (2021). Reconciling atmospheric cojsub¿2j/sub¿,
1065	weathering, and calcite compensation depth across the cenozoic. Science
1066	Advances, 7(4), eabd4876. Retrieved from https://www.science.org/doi/
1067	abs/10.1126/sciadv.abd4876 doi: 10.1126/sciadv.abd4876
1068	Kump, L. R., & Alley, R. B. (1994). Global chemical weathering on glacial time
1069	scales. Material Fluxes on the Surface of the Earth, 46–60.
1070	Köhler, P., & Munhoven, G. (2020). Late Pleistocene Carbon Cycle Re-
1071	visited by Considering Solid Earth Processes. Paleoceanography and
1072	Paleoclimatology, 35(12), e2020PA004020. Retrieved from https://
1073	agupubs.onlinelibrary.wiley.com/doi/abs/10.1029/2020PA004020
1074	(e2020PA004020 10.1029/2020PA004020) doi: https://doi.org/10.1029/
1075	2020PA004020
1076	Lacroix, F., Ilyina, T., & Hartmann, J. (2020). Oceanic CO <sub>2</sub> outgassing and bio-
1077	logical production hotspots induced by pre-industrial river loads of nutrients
1078	and carbon in a global modeling approach. Biogeosciences, $17(1)$ , 55-88. doi:
1079	10.5194/bg-17-55-2020
1080	Lacroix, F., Ilyina, T., Laruelle, G. G., & Regnier, P. (2021). Reconstructing
1081	the Preindustrial Coastal Carbon Cycle Through a Global Ocean Circula-
1082	tion Model: Was the Global Continental Shelf Already Both Autotrophic
1083	and a $CO_2$ Sink? Global Biogeochemical Cycles, $35(2)$ , e2020GB006603.
	Retrieved from https://agupubs.onlinelibrary.wiley.com/doi/
1084	abs/10.1029/2020GB006603 (e2020GB006603 2020GB006603) doi:
1085	https://doi.org/10.1029/2020GB006603
1086	Laws, E. A., Popp, B. N., Bidigare, R. R., Kennicutt, M. C., & Macko, S. A.
1087	(1995). Dependence of phytoplankton carbon isotopic composition on
1088	growth rate and $[CO_{2,aq}]$ : Theoretical considerations and experimental re-
1089	sults. Geochimica et Cosmochimica Acta, $59(6)$ , 1131-1138. Retrieved from
1090	https://www.sciencedirect.com/science/article/pii/0016703795000304
1091	doi: https://doi.org/10.1016/0016-7037(95)00030-4
1092	
1093	
1094	over the last 3 Myr. <i>Paleoceanography</i> , 29(2), 71-86. Retrieved from https://
1095	agupubs.onlinelibrary.wiley.com/doi/abs/10.1002/2013PA002505 doi: https://doi.org/10.1002/2013PA002505
1096	
1097	Lisiecki, L. E., & Raymo, M. E. (2005). A Pliocene-Pleistocene stack of 57 glob-
1098	ally distributed benchic d18O records. <i>Paleoceanography</i> , $20(1)$ , 1–17. doi: 10
1099	.1029/2004PA001071
1100	Liu, B., Six, K. D., & Ilyina, T. (2021). Incorporating the stable carbon iso-
1101	tope <sup>13</sup> C in the ocean biogeochemical component of the Max Planck In-
1102	stitute Earth System Model. $Biogeosciences, 18(14), 4389-4429.$ Re-
1103	trieved from https://bg.copernicus.org/articles/18/4389/2021/ doi:
1104	10.5194/bg-18-4389-2021
1105	Lourantou, A., Chappellaz, J., Barnola, J. M., Masson-Delmotte, V., & Raynaud, D.
1106	(2010). Changes in atmospheric $CO_2$ and its carbon isotopic ratio during the
1107	penultimate deglaciation. Quaternary Science Reviews, 29(17-18), 1983–1992.
1108	doi: 10.1016/j.quascirev.2010.05.002
1109	Lüthi, D., Le Floch, M., Bereiter, B., Blunier, T., Barnola, J. M., Siegenthaler,
1110	U., Stocker, T. F. (2008). High-resolution carbon dioxide concentration
1111	record $650,000-800,000$ years before present. Nature, $453(7193)$ , $379-382$ . doi:
1112	10.1038/nature06949
	Maier-Reimer, E., & Hasselmann, K. (1987). Transport and storage of $CO_2$ in the

1114 1115	ocean - an inorganic ocean-circulation carbon cycle model. Climate Dynamics, $\mathcal{Z}(2)$ , 63–90. doi: 10.1007/BF01054491
1116	Marcott, S. a., Bauska, T. K., Buizert, C., Steig, E. J., Rosen, J. L., Cuffey,
1117	K. M., Brook, E. J. (2014). Centennial-scale changes in the global
1118	carbon cycle during the last deglaciation. Nature, 514 (7524), 616–9. doi:
1119	10.1038/nature13799
1120	Matsumoto, K., Rickaby, R., & Tanioka, T. (2020). Carbon Export Buffering and
1121	CO <sub>2</sub> Drawdown by Flexible Phytoplankton C:N:P Under Glacial Conditions.
1122	Paleoceanography and Paleoclimatology, 35(7), e2019PA003823. Retrieved
1123	from https://agupubs.onlinelibrary.wiley.com/doi/abs/10.1029/
1124	2019PA003823 (e2019PA003823 2019PA003823) doi: https://doi.org/10.1029/
1125	2019PA003823
1126	Menviel, L., Joos, F., & Ritz, S. P. (2012). Simulating atmospheric CO <sub>2</sub> , 13C and
1127	the marine carbon cycle during the Last Glacial-Interglacial cycle: Possible
1128	role for a deepening of the mean remineralization depth and an increase in
1129	the oceanic nutrient inventory. Quaternary Science Reviews, 56, 46–68. doi:
1130	10.1016/j.quascirev.2012.09.012
1131	Metzler, H., Müller, M., & Sierra, C. A. (2018). Transit-time and age distri-
1132	butions for nonlinear time-dependent compartmental systems. Proceed-
1133	ings of the National Academy of Sciences, 115(6), 1150-1155. Retrieved
1134	from https://www.pnas.org/doi/abs/10.1073/pnas.1705296115 doi:
1135	10.1073/pnas.1705296115
1136	Meybeck, M. (1982). Carbon, Nitrogen, ans Phosphorous Transport by World
1137	Rivers. American Journal of Science, 282(4), 401-450. doi: 10.2475/
1138	ajs.282.4.401
1139	Milliman, J. D., & Droxler, A. W. (1996). Neritic and pelagic carbonate sedimenta-
1140	tion in the marine environment: Ignorance is not bliss. <i>Geologische Rundschau</i> ,
1141	85(3), 496-504, doi: 10.1007/BF02369004
1141 1142	85(3), 496–504. doi: 10.1007/BF02369004 Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., & Lenton, T. M.
1142	Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., & Lenton, T. M.
1142 1143	Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., & Lenton, T. M. (2017). Elevated CO <sub>2</sub> degassing rates prevented the return of Snow-
1142 1143 1144	Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., & Lenton, T. M.(2017).Elevated CO2 degassing rates prevented the return of Snow- ball Earth during the Phanerozoic.Nature Communications, 8.doi:
1142 1143 1144 1145	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> </ul>
1142 1143 1144	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Re-</li> </ul>
1142 1143 1144 1145 1146 1147	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211-223. doi: 10.1016/0077-7579(86)90043-8</li> </ul>
1142 1143 1144 1145 1146 1147 1148	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water</li> </ul>
1142 1143 1144 1145 1146 1147	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479–5499. doi:</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211-223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479-5499. doi: 10.1175/JCLI3911.1</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211-223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479-5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008).</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211-223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479-5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange for-</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479–5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange formulation and ocean transport strength. Global Biogeochemical Cycles, 22(3),</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1154 1155	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20 (2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19 (21), 5479–5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange formulation and ocean transport strength. Global Biogeochemical Cycles, 22 (3), 1–14. doi: 10.1029/2007GB003065</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1154 1155	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479–5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange formulation and ocean transport strength. Global Biogeochemical Cycles, 22(3), 1–14. doi: 10.1029/2007GB003065</li> <li>Munhoven, G. (2002). Glacial - Interglacial changes of continental weather-</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1154 1155	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snow- ball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Re- search, 20(2-3), 211-223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three- Dimensional Ocean Model. Journal of Climate, 19(21), 5479-5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange for- mulation and ocean transport strength. Global Biogeochemical Cycles, 22(3), 1-14. doi: 10.1029/2007GB003065</li> <li>Munhoven, G. (2002). Glacial - Interglacial changes of continental weather- ing: Estimates of the related CO<sub>2</sub> and HCO3 - flux variations and their</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1154 1155 1156	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479–5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange formulation and ocean transport strength. Global Biogeochemical Cycles, 22(3), 1–14. doi: 10.1029/2007GB003065</li> <li>Munhoven, G. (2002). Glacial - Interglacial changes of continental weathering: Estimates of the related CO<sub>2</sub> and HCO3 - flux variations and their uncertainties. Global and Planetary Change, 33(1-2), 155–176. doi:</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1154 1155 1156 1157	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snow- ball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Re- search, 20(2-3), 211-223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three- Dimensional Ocean Model. Journal of Climate, 19(21), 5479-5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange for- mulation and ocean transport strength. Global Biogeochemical Cycles, 22(3), 1-14. doi: 10.1029/2007GB003065</li> <li>Munhoven, G. (2002). Glacial - Interglacial changes of continental weather- ing: Estimates of the related CO<sub>2</sub> and HCO3 - flux variations and their uncertainties. Global and Planetary Change, 33(1-2), 155-176. doi: 10.1016/S0921-8181(02)00068-1</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1155 1156 1157 1158 1159 1159	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snow- ball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Re- search, 20(2-3), 211-223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three- Dimensional Ocean Model. Journal of Climate, 19(21), 5479-5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange for- mulation and ocean transport strength. Global Biogeochemical Cycles, 22(3), 1-14. doi: 10.1029/2007GB003065</li> <li>Munhoven, G. (2002). Glacial - Interglacial changes of continental weather- ing: Estimates of the related CO<sub>2</sub> and HCO3 - flux variations and their uncertainties. Global and Planetary Change, 33(1-2), 155-176. doi: 10.1016/S0921-8181(02)00068-1</li> <li>Munhoven, G., &amp; François, L. M. (1996). Glacial-interglacial variability of atmo-</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1154 1155 1156 1157	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211-223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479-5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange formulation and ocean transport strength. Global Biogeochemical Cycles, 22(3), 1-14. doi: 10.1029/2007GB003065</li> <li>Munhoven, G. (2002). Glacial - Interglacial changes of continental weathering: Estimates of the related CO<sub>2</sub> and HCO3 - flux variations and their uncertainties. Global and Planetary Change, 33(1-2), 155-176. doi: 10.1016/S0921-8181(02)00068-1</li> <li>Munhoven, G., &amp; François, L. M. (1996). Glacial-interglacial variability of atmospheric CO<sub>2</sub> due to changing continental silicate rock weathering: A model</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1154 1155 1156 1157 1158 1159 1160	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479–5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange formulation and ocean transport strength. Global Biogeochemical Cycles, 22(3), 1–14. doi: 10.1029/2007GB003065</li> <li>Munhoven, G. (2002). Glacial - Interglacial changes of continental weathering: Estimates of the related CO<sub>2</sub> and HCO3 - flux variations and their uncertainties. Global and Planetary Change, 33(1-2), 155–176. doi: 10.1016/S0921-8181(02)00068-1</li> <li>Munhoven, G., &amp; François, L. M. (1996). Glacial-interglacial variability of atmospheric CO<sub>2</sub> due to changing continental silicate rock weathering: A model study. Journal of Geophysical Research: Atmospheres, 101(D16), 21423-21437.</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1154 1155 1156 1157 1158 1159 1160 1161	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479–5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange formulation and ocean transport strength. Global Biogeochemical Cycles, 22(3), 1–14. doi: 10.1029/2007GB003065</li> <li>Munhoven, G. (2002). Glacial - Interglacial changes of continental weathering: Estimates of the related CO<sub>2</sub> and HCO3 - flux variations and their uncertainties. Global and Planetary Change, 33(1-2), 155–176. doi: 10.1016/S0921-8181(02)00068-1</li> <li>Munhoven, G., &amp; François, L. M. (1996). Glacial-interglacial variability of atmospheric CO<sub>2</sub> due to changing continental silicate rock weathering: A model study. Journal of Geophysical Research: Atmospheres, 101(D16), 21423-21437. Retrieved from https://agupubs.onlinelibrary.wiley.com/doi/abs/</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1154 1155 1155 1156 1157 1158 1159 1160 1161	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479–5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange formulation and ocean transport strength. Global Biogeochemical Cycles, 22(3), 1–14. doi: 10.1029/2007GB003065</li> <li>Munhoven, G. (2002). Glacial - Interglacial changes of continental weathering: Estimates of the related CO<sub>2</sub> and HCO3 - flux variations and their uncertainties. Global and Planetary Change, 33(1-2), 155–176. doi: 10.1016/S0921-8181(02)00068-1</li> <li>Munhoven, G., &amp; François, L. M. (1996). Glacial-interglacial variability of atmospheric CO<sub>2</sub> due to changing continental silicate rock weathering: A model study. Journal of Geophysical Research: Atmospheres, 101(D16), 21423-21437. Retrieved from https://agupubs.onlinelibrary.wiley.com/doi/abs/ 10.1029/96JD01842 doi: https://doi.org/10.1029/96JD01842</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1154 1155 1156 1157 1158 1159 1160 1161 1162 1163	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479–5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange formulation and ocean transport strength. Global Biogeochemical Cycles, 22(3), 1–14. doi: 10.1029/2007GB003065</li> <li>Munhoven, G. (2002). Glacial - Interglacial changes of continental weathering: Estimates of the related CO<sub>2</sub> and HCO3 - flux variations and their uncertainties. Global and Planetary Change, 33(1-2), 155–176. doi: 10.1016/S0921-8181(02)00068-1</li> <li>Munhoven, G., &amp; François, L. M. (1996). Glacial-interglacial variability of atmospheric CO<sub>2</sub> due to changing continental silicate rock weathering: A model study. Journal of Geophysical Research: Atmospheres, 101(D16), 21423-21437. Retrieved from https://agupubs.onlinelibrary.wiley.com/doi/abs/</li> </ul>
1142 1143 1144 1145 1146 1147 1148 1149 1150 1151 1152 1153 1154 1155 1156 1157 1158 1159 1160 1161 1162 1163	<ul> <li>Mills, B. J. W., Scotese, C. R., Walding, N. G., Shields, G. A., &amp; Lenton, T. M. (2017). Elevated CO<sub>2</sub> degassing rates prevented the return of Snowball Earth during the Phanerozoic. Nature Communications, 8. doi: 10.1038/s41467-017-01456-w</li> <li>Mook, W. G. (1986). 13C in atmospheric CO<sub>2</sub>. Netherlands Journal of Sea Research, 20(2-3), 211–223. doi: 10.1016/0077-7579(86)90043-8</li> <li>Müller, S. A., Joos, F., Edwards, N. R., &amp; Stocker, T. F. (2006). Water Mass Distribution and Ventilation Time Scales in a Cost-Efficient, Three-Dimensional Ocean Model. Journal of Climate, 19(21), 5479–5499. doi: 10.1175/JCLI3911.1</li> <li>Müller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., &amp; Stocker, T. F. (2008). Modeled natural and excess radiocarbon: Sensitivities to the gas exchange formulation and ocean transport strength. Global Biogeochemical Cycles, 22(3), 1–14. doi: 10.1029/2007GB003065</li> <li>Munhoven, G. (2002). Glacial - Interglacial changes of continental weathering: Estimates of the related CO<sub>2</sub> and HCO3 - flux variations and their uncertainties. Global and Planetary Change, 33(1-2), 155–176. doi: 10.1016/S0921-8181(02)00068-1</li> <li>Munhoven, G., &amp; François, L. M. (1996). Glacial-interglacial variability of atmospheric CO<sub>2</sub> due to changing continental silicate rock weathering: A model study. Journal of Geophysical Research: Atmospheres, 101(D16), 21423-21437. Retrieved from https://agupubs.onlinelibrary.wiley.com/doi/abs/ 10.1029/96JD01842 doi: https://doi.org/10.1029/96JD01842</li> <li>Najjar, R. G., &amp; Orr, J. C. (1999). Biotic-HOWTO. Internal OCMIP (Tech. Rep.).</li> </ul>

1169	40,000 yr. Nature, 295(5846), 220–223. doi: 10.1038/295220a0
1170	Niemeyer, D., Kemena, T. P., Meissner, K. J., & Oschlies, A. (2016). A model
1171	study of warming-induced phosphorus-oxygen feedbacks in open-ocean oxygen
1172	minimum zones on millennial timescales. Earth System Dynamics Discussions,
1173	1-18. doi: $10.5194/esd-2016-50$
1174	Oliver, K. I. C., Hoogakker, B. A. A., Crowhurst, S., Henderson, G. M., Rickaby,
1175	R. E. M., Edwards, N. R., & Elderfield, H. (2010). A synthesis of marine sed-
1176	iment core d13C data over the last 150 000 years. Climate of the Past, $6(5)$ ,
1177	645–673. doi: 10.5194/cp-6-645-2010
1178	Orr, J. C., & Epitalon, J. M. (2015). Improved routines to model the ocean carbon-
1179	ate system: Mocsy 2.0. Geoscientific Model Development, 8(3), 485–499. doi:
1180	10.5194/gmd-8-485-2015
1181	Palastanga, V., Slomp, C. P., & Heinze, C. (2011). Long-term controls on ocean
1182	phosphorus and oxygen in a global biogeochemical model. Global Biogeochemi-
1183	cal Cycles, 25(3), 1–19. doi: 10.1029/2010GB003827
1184	Parekh, P., Joos, F., & Müller, S. A. (2008). A modeling assessment of the inter-
1185	play between aeolian iron fluxes and iron-binding ligands in controlling carbon
1186	dioxide fluctuations during Antarctic warm events. Paleoceanography, 23(4),
1187	PA4202. doi: 10.1029/2007PA001531
1188	Paulmier, A., Kriest, I., & Oschlies, A. (2009). Stoichiometries of remineralisation
1189	and denitrification in global biogeochemical ocean models. <i>Biogeosciences</i> ,
1105	6(5), 923-935. Retrieved from https://bg.copernicus.org/articles/6/
1190	923/2009/ doi: 10.5194/bg-6-923-2009
1192	Peterson, C. D., & Lisiecki, L. E. (2018). Deglacial carbon cycle changes observed in
1192	a compilation of 127 benthic d13C time series (20-6 ka). Climate of the Past,
	14(8), 1229-1252. doi: $10.5194/cp-14-1229-2018$
1194	Peterson, C. D., Lisiecki, L. E., & Stern, J. V. (2014). Deglacial whole-ocean d13C
1195	change estimated from 480 benthic foraminiferal records. <i>Paleoceanography</i> ,
1196	29, 549–563. doi: 10.1002/2013PA002552.Received
1197	Peterson, L. C., & Prell, W. L. (1985). Carbonate Preservation and Rates of
1198	Climatic Change: An 800 kyr Record from the Indian Ocean. In <i>The Car</i> -
1199	bon Cycle and Atmospheric $CO_2$ : Natural Variations Archean to Present
1200	(p. 251-269). American Geophysical Union (AGU). Retrieved from
1201	https://agupubs.onlinelibrary.wiley.com/doi/abs/10.1029/GM032p0251
1202	doi: https://doi.org/10.1029/GM032p0251
1203	Popp, B. N., Takigiku, R., Hayes, J. M., Louda, J. W., & Baker, E. W. (1989).
1204	The post-Paleozoic chronology and mechanism of 13 C depletion in primary
1205	marine organic matter. American Journal of Science, 289(4), 436–454.
1206	Retrieved from https://www.ajsonline.org/content/289/4/436 doi:
1207	10.2475/ajs.289.4.436
1208	Ramirez, A. J., & Rose, A. W. (1992). Analytical geochemistry of organic phos-
1209	phorus and its correlation with organic carbon in marine and fluvial sed-
1210	iments and soils. <i>American Journal of Science</i> , 292(6), 421–454. Re-
1211	trieved from https://www.ajsonline.org/content/292/6/421 doi:
1212	10.2475/ajs.292.6.421
1213	Raymo, M. E., Oppo, D. W., & Curry, W. (1997). The mid-Pleistocene climate
1214	transition: A deep sea carbon isotopic perspective. Paleoceanography, 12(4),
1215	546–559. doi: 10.1029/97PA01019
1216	
1217	Regnier, P., Friedlingstein, P., Ciais, P., Mackenzie, F. T., Gruber, N., Janssens, L.A. Thullner, M. (2012) Anthropogonia porturbation of the car
1218	I. A., Thullner, M. (2013). Anthropogenic perturbation of the car- bon fluxes from land to ocean. <i>Nature Geoscience</i> , 6(8), 597-607. doi:
1219	bon fluxes from land to ocean. Nature Geoscience, $6(8)$ , 597-607. doi: 10.1038/NGEO1830
1220	
1221	Ridgwell, A., & Zeebe, R. (2005). The role of the global carbonate cycle in the regu-
1222	lation and evolution of the Earth system. Earth and Planetary Science Letters,
1223	234(3-4), 299-315. doi: 10.1016/j.epsl.2005.03.006

Ritz, S., Stocker, T. F., & Joos, F. (2011).A Coupled Dynamical Ocean-Energy 1224 Balance Atmosphere Model for Paleoclimate Studies. Journal of Climate, 24, 1225 349-375. doi: 10.1175/2010JCLI3351.1 1226 Roth, R., & Joos, F. (2012). Model limits on the role of volcanic carbon emissions in 1227 regulating glacial-interglacial CO<sub>2</sub>variations. Earth and Planetary Science Let-1228 ters, 329-330, 141-149. doi: 10.1016/j.epsl.2012.02.019 1229 Roth, R., Ritz, S. P., & Joos, F. (2014). Burial-nutrient feedbacks amplify the sensi-1230 tivity of atmospheric carbon dioxide to changes in organic matter remineralisa-1231 tion. Earth System Dynamics, 5(2), 321-343. doi: 10.5194/esd-5-321-2014 1232 Rugenstein, J. K. C., Ibarra, D. E., & von Blanckenburg, F. (2019). Neogene cooling 1233 driven by land surface reactivity rather than increased weathering fluxes. Na-1234 ture, 571(7763), 99+. doi:  $\{10.1038/s41586-019-1332-y\}$ 1235 Sarmiento, J. L., & Gruber, N. (2006). Ocean Biogeochemical Dynamics. Princeton: 1236 Princeton University Press. 1237 Sarmiento, J. L., Orr, J. C., & Siegenthaler, U. (1992). A perturbation simulation of 1238 CO<sub>2</sub> uptake in an ocean general circulation model. Journal of Geophysical Re-1239 search: Oceans, 97(C3), 3621-3645. doi: 10.1029/91JC02849 1240 Schmitt, J., Schneider, R., Elsig, J., Leuenberger, D., Lourantou, a., Chappellaz, J., 1241 ... Fischer, H. (2012). Carbon Isotope Constraints on the Deglacial  $CO_2$  Rise 1242 from Ice Cores. Science, 336(6082), 711-714. doi: 10.1126/science.1217161 1243 Schmittner, A., Gruber, N., Mix, A. C., Key, R. M., Tagliabue, A., & Westberry, 1244 Τ. (2013).Biology and air–sea gas exchange controls on the distribution of 1245 carbon isotope ratios (d13C) in the ocean. Biogeosciences, 10, 5793-5816. doi: 1246 10.5194/bg-10-5793-2013 1247 Schneider, R., Schmitt, J., Köhler, P., Joos, F., & Fischer, H. (2013). A reconstruc-1248 tion of atmospheric carbon dioxide and its stable carbon isotopic composition 1249 from the penultimate glacial maximum to the last glacial inception. Climate of 1250 the Past, 9(6), 2507–2523. doi: 10.5194/cp-9-2507-2013 1251 Schrag, D. P., Higgins, J. A., Macdonald, F. A., & Johnston, D. T. (2013).Au-1252 thigenic Carbonate and the History of the Global Carbon Cycle. Science. 1253 Retrieved from https://www.science.org/doi/abs/ 339(6119), 540-543.1254 10.1126/science.1229578 doi: 10.1126/science.1229578 1255 Siegenthaler, U., & Muennich, K. O. (1981). Carbon-13/carbon-12 fractionation dur-1256 ing carbon dioxide transfer from air to sea. In B. Bolin (Ed.), Scope 16: Car-1257 bon cycle modelling (pp. 249–257). Chichester: Wiley. 1258 Siegenthaler, U., & Oeschger, H. (1987).Biospheric CO<sub>2</sub> emissions dur-1259 ing the past 200 years reconstructed by deconvolution of ice core data. 1260 Tellus B: Chemical and Physical Meteorology, 39(1-2), 140–154. doi: 1261 10.3402/tellusb.v39i1-2.15331 1262 Siegenthaler, U., Stocker, T. F., Monnin, E., Lüthi, D., Schwander, J., Stauffer, B., ... Jouzel, J. (2005). Stable Carbon Cycle-Climate Relationship During the 1264 Late Pleistocene. Science, 310, 1313–1317. 1265 Sigman, D. M., & Boyle, E. A. (2000).Glacial/Interglacial Variations In Atmo-1266 spheric Carbon Dioxide. Nature, 407(6806), 859–869. doi: 10.1038/35038000 1267 Sigman, D. M., Hain, M. P., & Haug, G. H. (2010).The polar ocean and glacial 1268 cycles in atmospheric  $CO_2$  concentration. Nature, 466(7302), 47–55. doi: 10 .1038/nature09149 Sigman, D. M., McCorkle, D. C., & Martin, W. R. (1998). The calcite lysocline as a 1271 constraint on glacial/interglacial low-latitude production changes. Global Bio-1272 geochemical Cycles, 12(3), 409–427. doi: 10.1029/98GB01184 1273 Stolper, D. A., Bender, M. L., Dreyfus, G. B., Yan, Y., & Higgins, J. A. (2016).1274 A Pleistocene ice core record of atmospheric O<sub>2</sub> concentrations. Science, 1275 353(6306), 1427–1430. doi: 10.1126/science.aaf5445 1276 Strassmann, K. M., & Joos, F. (2018). The Bern Simple Climate Model (BernSCM) 1277 v1.0: an extensible and fully documented open-source re-implementation of the 1278

1279	Bern reduced-form model for global carbon cycle-climate simulations. Geosci-
1280	entific Model Development, 11(5), 1887-1908. doi: 10.5194/gmd-11-1887-2018
1281	Suchet, P., & Probst, J. (1995). A global-model for present day atmospheric soil
1282	$CO_2$ consumption by chemical erosion of continental rocks (GEM-CO2). Tellus
1283	Series B - Chemical and Physical Meteorology, 47(1-2), 273-280. (4th Atmo-
1284	spheric CO(2) International Conference, CARQUEIRANNE, FRANCE, SEP
1285	13-17, 1993) doi: 10.1034/j.1600-0889.47.issue1.23.x
1286	Thompson, M., & Randerson, J. (1999). Impulse response functions of terrestrial
1287	carbon cycle models: method and application. Global Change Biology, $5(4)$ ,
1288	371-394. doi: 10.1046/j.1365-2486.1999.00235.x
1289	Tréguer, P. J., Sutton, J. N., Brzezinski, M., Charette, M. A., Devries, T.,
1290	Dutkiewicz, S., Rouxel, O. (2021). Reviews and syntheses: The biogeo-
1291	chemical cycle of silicon in the modern ocean. Biogeosciences, 18(4), 1269–
1292	1289. Retrieved from https://bg.copernicus.org/articles/18/1269/2021/
1293	doi: 10.5194/bg-18-1269-2021
1294	Tschumi, T., Joos, F., Gehlen, M., & Heinze, C. (2011). Deep ocean ventilation, car-
1295	bon isotopes, marine sedimentation and the deglacial $CO_2$ rise. Climate of the
1296	Past, 7(3), 771-800. doi: 10.5194/cp-7-771-2011
1297	Van Cappellen, P., & Ingall, E. D. (1994). Benthic phosphorus regeneration, net
1298	primary production, and ocean anoxia: A model of the coupled marine bio-
1299	geochemical cycles of carbon and phosphorus. Paleoceanography, 9(5), 677–
1300	692. Retrieved from http://doi.wiley.com/10.1029/94PA01455 doi:
1301	10.1029/94PA01455
1302	Walker, J. C. G., Hays, P. B., & Kasting, J. F. (1981). A negative feedback mecha-
1303	nism for the long-term stabilization of Earth's surface temperature. Journal of
1304	Geophysical Research: Oceans, 86(C10), 9776-9782. Retrieved from https://
1305	agupubs.onlinelibrary.wiley.com/doi/abs/10.1029/JC086iC10p09776
1306	doi: https://doi.org/10.1029/JC086iC10p09776
1307	Wallmann, K. (2010). Phosphorus imbalance in the global ocean? Global Bio-
1308	geochemical Cycles, 24(4), GB4030. Retrieved from http://doi.wiley.com/
1309	10.1029/2009GB003643 doi: 10.1029/2009GB003643
1310	Wallmann, K. (2014). Is late Quaternary climate change governed by self-sustained
1311	oscillations in atmospheric CO <sub>2</sub> ? Geochimica et Cosmochimica Acta, 132, 413–
1312	439. doi: 10.1016/j.gca.2013.10.046
1313	Wallmann, K., Schneider, B., & Sarnthein, M. (2016). Effects of eustatic sea-level
1314	change, ocean dynamics, and nutrient utilization on atmospheric $pco_2$ and
1315	seawater composition over the last 130 000 years: a model study. Climate
1316	of the Past, 12(2), 339-375. Retrieved from https://cp.copernicus.org/
1317	articles/12/339/2016/ doi: 10.5194/cp-12-339-2016
1318	Wanninkhof, R. (2014). Relationship between wind speed and gas exchange over the
1319	ocean revisited. Limnology and Oceanography: Methods, $12(6)$ , $351-362$ . doi:
1320	10.4319/lom.2014.12.351
1321	Watson, B. A. J. (2016). Oceans on the edge of anoxia. Science, $354(6319)$ , $1529-$
1322	1530.
1323	Willeit, M., Ganopolski, A., Calov, R., & Brovkin, V. (2019). Mid-Pleistocene tran-
1324	sition in glacial cycles explained by declining $CO_2$ and regolith removal. Sci-
1325	ence Advances, $5(4)$ . doi: 10.1126/sciadv.aav7337
1326	Willeit, M., Ganopolski, A., Robinson, A., & Edwards, N. R. (2022). The Earth
1327	system model CLIMBER-X v1.0 – Part 1: Climate model description and
1328	validation. Geoscientific Model Development, 15(14), 5905–5948. Re-
1329	trieved from https://gmd.copernicus.org/articles/15/5905/2022/ doi:
1330	10.5194/gmd-15-5905-2022