# Geophysical and geochemical controls on abiotic carbon cycling on Earth-like planets

Marc Neveu<sup>1</sup>, Taylor Louanne Che Bartlow<sup>2</sup>, Ryan Felton<sup>3</sup>, Shawn D<br/> Domagal-Goldman<sup>4</sup>, and Steven  $\mathrm{Desch}^5$ 

<sup>1</sup>Goddard Space Flight Center/University of Maryland <sup>2</sup>University of Maryland <sup>3</sup>Catholic University of America <sup>4</sup>National Aeronautics and Space Administration (NASA) <sup>5</sup>Arizona State University

February 27, 2023

## Abstract

We investigate how variations in a planet's size and the chemical (mineral) composition of its upper mantle and surface affect processes involved in the carbonate-silicate cycle, which is thought to have regulated the composition of Earth's atmosphere and its surface temperature over geologic time. We present models of geophysical and geochemical controls on these processes: outgassing, continental weathering, and seafloor weathering, and analyze sensitivities to planet size and composition. For Earth-like compositions, outgassing is maximized for planets of Earth's size. Smaller planets convect less vigorously; higher pressures inside larger planets hinder melting. For more felsic mantles, smaller planets (0.5-0.75 Earth mass) outgas more, whereas more mafic planets follow the size trend of Earth's composition. Planet size and composition can affect outgassing by two orders of magnitude, with variability driven by mass in the first 2.5 Gyr after formation and by composition past that time. In contrast, simulations spanning the diversity of surface composition or the patchiness of land as they are to surface temperature, with fluxes within a factor of five of Earth's. Seafloor weathering appears more sensitive to uncertainties in tectonic regime (occurrence, speed, and size of plates) than to seafloor composition. These results form a basis to interpret calculations of geological surface carbon fluxes to track atmospheric compositions, through time, of lifeless exo-Earths, providing a baseline against which the effect of biological activity may be distinguished with telescopic observations.

# Geophysical and geochemical controls on abiotic carbon cycling on Earth-like planets

M. Neveu<sup>1,2</sup>, T. Bartlow<sup>1</sup>, R. Felton<sup>3</sup>, S. Domagal-Goldman<sup>2</sup>, S. Desch<sup>4</sup>

<sup>1</sup>University of Maryland, College Park, 4296 Stadium Drive, College Park, MD 20742, USA
 <sup>2</sup>NASA Goddard Space Flight Center, 8800 Greenbelt Road, Greenbelt, MD 20771, USA
 <sup>3</sup>NASA Ames Research Center, Moffett Field, CA 94043, USA
 <sup>4</sup>Arizona State University, 781 E Terrace Mall, Tempe, AZ 85287

# **Key Points:**

3

# A new code enables modeling of carbon cycling on planets of size and composition that differ from Earth's. Outgassing tends to go down with geologic time, is maximal for planets around 1 Earth mass, and higher for more felsic upper mantles. Continental weathering is comparatively less sensitive to surface composition and patchiness, and insensitive to planet size.

Corresponding author: Marc Neveu, marc.f.neveu@nasa.gov

#### 15 Abstract

We investigate how variations in a planet's size and the chemical (mineral) composition 16 of its upper mantle and surface affect processes involved in the carbonate-silicate cycle, 17 which is thought to have regulated the composition of Earth's atmosphere and its sur-18 face temperature over geologic time. We present models of geophysical and geochemi-19 cal controls on these processes: outgassing, continental weathering, and seafloor weath-20 ering, and analyze sensitivities to planet size and composition. For Earth-like compo-21 sitions, outgassing is maximized for planets of Earth's size. Smaller planets convect less 22 vigorously; higher pressures inside larger planets hinder melting. For more felsic man-23 tles, smaller planets (0.5-0.75 Earth mass) outgas more, whereas more mafic planets fol-24 low the size trend of Earth's composition. Planet size and composition can affect out-25 gassing by two orders of magnitude, with variability driven by mass in the first 2.5 Gyr 26 after formation and by composition past that time. In contrast, simulations spanning 27 the diversity of surface compositions encountered in the inner solar system indicate that 28 continental weathering fluxes are about as sensitive to surface composition or the patch-29 iness of land as they are to surface temperature, with fluxes within a factor of five of Earth's. 30 Seafloor weathering appears more sensitive to uncertainties in tectonic regime (occur-31 rence, speed, and size of plates) than to seafloor composition. These results form a ba-32 sis to interpret calculations of geological surface carbon fluxes to track atmospheric com-33 positions, through time, of lifeless exo-Earths, providing a baseline against which the ef-34 fect of biological activity may be distinguished with telescopic observations. 35

36

# Plain Language Summary

Earth's surface pressure and temperature have been relatively stable over its his-37 tory. They are likely regulated by the greenhouse effect induced by carbon dioxide  $(CO_2)$ 38 in Earth's atmosphere.  $CO_2$  is thought to be outgassed from the mantle, e.g., through 39 volcanism.  $CO_2$  is removed from the atmosphere by chemical weathering (the reaction 40 of surface rocks with  $CO_2$ -bearing rainwater), and river transport of its products to the 41 ocean where carbonate rocks form. Carbonates are subducted to the mantle, complet-42 ing the carbonate-silicate cycle. Here, we develop a model of mantle convection, melt-43 ing, and chemical reactions between water and rock to investigate how this cycle might 44 vary on solid planets with different sizes and mineral compositions. We find that these 45 affect carbon outgassing more than weathering. This new ability to model carbon cy-46

cling on a diversity of solid planets with an ocean helps understand what these planets

<sup>48</sup> might look like without a biosphere. Deviations from that baseline could be used to search<sup>49</sup> for signs of life.

50 1 Introduction

Thousands of planets are now known to orbit stars other than our Sun (NASA Ex-51 oplanet Archive; Akeson et al. (2013)). Amidst the zoo of observed combinations of host-52 star types, star-planet distances, orbital eccentricities, and planet sizes, a handful of plan-53 ets bearing similarities with Earth have been identified (Schulze-Makuch et al., 2011; Barnes 54 et al., 2015; Kane et al., 2016; Gillon et al., 2017). These are primarily solid worlds for 55 which "exo-climate" models have predicted temperatures that might allow surface liq-56 uid water (Wordsworth et al., 2011). Such potentially habitable worlds show promise as 57 targets for detecting signs of life, e.g., via atmospheric spectral signatures (Krissansen-58 Totton et al., 2016; Grenfell, 2017; Schwieterman et al., 2018; Krissansen-Totton, Gar-59 land, et al., 2018; M. A. Thompson et al., 2022) in the coming decades (Kiang et al., 2018; 60 Fujii et al., 2018; Pidhorodetska et al., 2020; Lin et al., 2021; Mikal-Evans, 2022). 61

Confidently attributing atmospheric spectral features to biology demands a thor-62 ough understanding of the physics, chemistry, and geology of exoplanets (Krissansen-63 Totton & Catling, 2017; Catling et al., 2018; Lisse et al., 2020). To detect life on an ex-64 oplanet, we must first constrain the geochemical cycles on a lifeless world, so that we rec-65 ognize what is not attributable to abiotic processes. This requires the ability to model 66 the interplay among interior, surface, atmospheric, and orbital processes, which is an over-67 arching goal of the exoplanet community (Forget & Leconte, 2014; Lehmer et al., 2020; 68 Barnes et al., 2020; Kopparapu et al., 2020; Unterborn et al., 2021; Krissansen-Totton 69 et al., 2022). 70

Previous efforts to understand exoplanet geology and climate have so far focused
on tidal (Barnes et al., 2013; Pierrehumbert & Hammond, 2019; Blackledge et al., 2020;
Colose et al., 2021), orbital (Spiegel et al., 2009; Dressing et al., 2010; Kaspi & Showman, 2015; Lehmer et al., 2020; Vervoort et al., 2022), compositional (Bond et al., 2010;
Young et al., 2014; Unterborn et al., 2014, 2016; Shahar et al., 2019; Unterborn et al.,
2022), and land coverage controls (Abbot et al., 2012; Tziperman et al., 2012; Lewis et
al., 2018; Krissansen-Totton, Arney, & Catling, 2018; Madden & Kaltenegger, 2020). Foley

-3-

(2015) has shown that the negative climate feedbacks inherent in the long-term carbon
cycle are uninhibited by climate's effect on plate tectonics. Models of atmospheric chemistry and chemical evolution have assumed surface fluxes of gases from outgassing or weathering as lower boundary conditions (Hu et al., 2012; Domagal-Goldman et al., 2014; Felton et al., 2022), usually estimated from present Earth values (Edson et al., 2012; KrissansenTotton & Catling, 2017; Krissansen-Totton, Arney, & Catling, 2018; Lehmer et al., 2020).

Outgassing rates have been predicted to increase with planet size (Kite et al., 2009). 84 In addition, the oxidation state of a planet's environment influences the surface and at-85 mospheric speciation of key elements such as carbon. Carbon is a chief component of plan-86 etary atmospheres as  $CO_2$ , CO, or  $CH_4$ ; and an essential building block of life as we know 87 it (Orgel, 1998). As CO<sub>2</sub>, it is also a major control of Earth's climate via the carbonate-88 silicate cycle (Kasting, 1988). While some studies have probed the photochemistry of 89 reduced carbon-rich atmospheres on terrestrial planets (Haqq-Misra et al., 2008; Arney 90 et al., 2017; Felton et al., 2022), there is a need to investigate how the carbonate-silicate 91 cycle might differ on worlds with a different bulk composition (Bond et al., 2010; Young 92 et al., 2014; Unterborn et al., 2022). 93

To address this need, we have developed a model of geophysical and geochemical 94 controls on abiotic cycling of carbon, on planets of 0.5 to 2 Earth masses  $(M_{\oplus})$  with the 95 same insolation as Earth. We use this model to investigate how the carbonate-silicate 96 cycle varies depending on planet bulk composition, size, and age. The model incorpo-97 rates key features of carbon cycling (Rushby et al., 2018): (1) equilibrium between at-98 mospheric carbon and dissolved carbon in the ocean, (2) removal from the atmosphere 99 by dissolving in rainwater and reacting with silicate rocks (weathering), (3) carbon re-100 moval from the ocean by seafloor water-rock interaction (Sleep & Zahnle, 2001; Krissansen-101 Totton & Catling, 2017), and (4) release of atmospheric carbon by outgassing such as 102 volcanism or other venting. In this paper, we describe models for each of these processes 103 that are validated to modern Earth conditions, but are based on first principles of geo-104 physics and geochemistry rather than on scalings with modern-day Earth. This enables 105 a departure from the necessarily Earth-centric conditions beyond which Earth-based scal-106 ings no longer accurately approximate the underlying outgassing and weathering pro-107 cesses. The present models can thus be used to investigate planets whose size and chem-108 ical composition differ from Earth's. In a companion paper, we describe the results of 109 the integrated abiotic carbon cycling model. 110

-4-

#### manuscript submitted to JGR: Planets



Figure 1. Architecture of the box model used to compute a planet's atmospheric and ocean composition as a function of time, with key computed parameters and inputs. The two reservoirs of carbon are the mantle and {atmosphere+ocean}. Fluxes between these reservoirs are due to seafloor weathering and mantle outgassing. The seafloor weathering flux depends on ocean composition, which itself depends on material fluxes arising from continental weathering and on ocean equilibrium with the atmosphere.

This paper is organized as follows. In Section 2, we describe the models of continental weathering, seafloor weathering, and outgassing, and their ability to reproduce corresponding estimated carbon fluxes on modern Earth. In Section 3, continental weathering and outgassing fluxes are computed for a variety of planet sizes and upper mantle or surface compositions. Sensitivity to these properties and their implications are discussed in Section 4; in particular, an example computation of the full carbon cycle through time is presented. We provide major conclusions in Section 5.

## 118 **2** Model

The abiotic carbon cycling model (Fig. 1), determines net fluxes (in mol m<sup>-2</sup> s<sup>-1</sup>) of carbon species from the surface and interior to the atmosphere. We assume that these fluxes are primarily due to continental weathering, seafloor weathering, and outgassing, and that the atmosphere is equilibrated with the ocean into a single reservoir. This section focuses on describing models of each of these processes; description of their coupling in the overall model is provided in the companion paper. An implementation of the model in C language is freely available at https://github.com/MarcNeveu/ExoCcycleGeo. 126

# 2.1 Geodynamics and outgassing

Outgassing of a planet's interior to the atmosphere results from partial melting of 127 mantle material advected to shallow depths (Kite et al., 2009). Although localized vol-128 canism is a prominent form of outgassing, much of outgassing on a planet may occur from 129 more diffuse (e.g., hydrothermal) sources. Both sources are linked to internal convection, 130 melting, and subsequent conductive cooling of the crust. In this model, we do not dis-131 tinguish subaerial from subaqueous outgassing because we consider the atmosphere and 132 ocean as a single reservoir in which the atmosphere and ocean compositions are at equi-133 librium. 134

Our overall procedure for calculating outgassing fluxes is as follows. The thermal 135 evolution of a planet of a given size is computed by balancing accretional and radiogenic 136 heating with convective and conductive cooling (as a function of tectonic mode) to ob-137 tain temperature profiles as a function of depth at different time points in a planet's his-138 tory. Geodynamic model parameters and adopted values are described in Table 1. These 139 temperature profiles are used to determine the melt fraction as a function of depth for 140 rock of a specified composition, using the *alphaMELTS* command line implementation 141 (Smith & Asimow, 2005; R. N. Thompson et al., 2007; P. M. Antoshechkina & Asimow, 142 2010; P. M. Antoshechkina et al., 2010; P. Antoshechkina & Asimow, 2018) of the MELTS 143 software (Ghiorso & Sack, 1995; Asimow & Ghiorso, 1998). From melt fractions, using 144 the same simplifying assumption as Kite et al. (2009) that all melt reaches the surface 145 (or at least the near-surface crust), we obtain rates of melt extrusion (or intrusion). Fi-146 nally, we assume that the rate of carbon outgassing is directly proportional to that ex-147 trusion and intrusion rate, based on the assumed carbon content of the upper mantle ma-148 terial able to be outgassed. 149

150

#### 2.1.1 Planet structure

The user inputs a planet mass, core mass fraction, and three materials for an inner core, outer core, and mantle. From these parameters, an interior structure is generated for the planet using the approach described in Lorenzo et al. (2014). The planet is divided in 1-dimensional grid zones, assuming spherical symmetry. Given a first-guess radius, the gravitational acceleration is computed as a function of depth. The equation of hydrostatic equilibrium is then integrated to find the pressure at depth, assuming con**Table 1.** Geodynamic model parameters. The lack of dependency for this study of the assumed values for the geodynamical parameters described in this section on a planet's bulk composition (Table 1) is addressed in Section 4. <sup>*a*</sup> User-specified inputs via planet mass, core mass fraction, and inner core, outer core, and mantle materials (Lorenzo et al., 2014).

Parameter	Symbol	Value	Units	Notes and references
Planet Structure				
Planet radius $^{a}$	$r_p$		m	
Core-mantle boundary radius $^{a}$	$r_c$		m	
Grid zone radius	r		m	
Gravitational acceleration	g		${\rm m~s^{-2}}$	
Density <sup>a</sup>	ρ		${\rm kg}~{\rm m}^{-3}$	Lorenzo et al. (2014)
Pressure	P		Pa	$= \int_r^{r_p} \rho(r) \ g(r) \ dr$
Temperature	T		К	
Accretion				
Fraction of impactor energy deposited at depth	h	0.1		Canup et al. (2021)
Ratio of planetesimal velocity to escape velocity	χ	2.0		Canup et al. (2021)
Thermal Evolution				
Radiogenic power per unit mass	H		${\rm W~kg^{-1}}$	Nuclide heating rate $\times$
				mantle abundance (Table 2)
Radionuclide half-life	$t_{1/2}$		s	See Table 2
Mantle heat capacity	$C_p$	1295	$\rm J~kg^{-1}~K^{-1}$	Akaogi and Ito (1993)
Mantle thermal conductivity	k	4.18	$\mathrm{W}~\mathrm{m}^{-1}~\mathrm{K}^{-1}$	Turcotte and Schubert (2002)
Mantle thermal diffusivity	$\kappa$		$\mathrm{m}^2~\mathrm{s}^{-1}$	$= k/(\rho \ C_p)$
Mid-mantle temperature	$T_m$		К	From equation $(4)$
Surface temperature	$T_s$		К	
Mantle thermal expansivity	α	$3 \times 10^{-5}$	$K^{-1}$	Turcotte and Schubert (2002)
Radius of brittle-ductile transition	$r_{BDT}$		m	
Temperature in mantle adiabat	$T_{adiab}$		К	From equation $(5)$
Temperature in lithosphere	$T_{upbnd}$		К	From equation $(6)$
Depth of lithosphere	$z_{Lith}$		m	Where $T_{adiab} = T_{upbnd}$
Rayleigh number	Ra		—	
Critical Rayleigh number	$Ra_{cr}$	1707.762	—	Reid and Harris (1958)
Nusselt number scaling parameter	$\beta$	1/3	—	$Nu = (Ra/Ra_{cr})^{\beta}$
Convective velocity	$v_{conv}$		${\rm m~s^{-1}}$	From equation (7)
Convective timescale	$t_{conv}$		s	$= (r_{BDT} - r_c)/v_{conv}$

157	tinuity across core and mantle boundaries. Equations of state for each of the three ma-
158	terials are inverted to derive the densities as a function of pressure. Based on these den-
159	sities, the thickness of the layers is adjusted. The gravitational acceleration as a func-
160	tion of depth is then updated, and so on until a convergence threshold is reached on the
161	central density. Several equations of state are considered depending on the material: a
162	third-order Birch-Murnaghan equation of state (Lorenzo et al., 2014), a power law (Seager
163	et al., 2007), and for high-pressure phases of ice, a temperature-dependent equation (Choukroun
164	& Grasset, 2010). A database of planetary materials and equation-of-state parameters
165	is provided with the code and includes data from Seager et al. (2007) for $\alpha$ -(bcc)Fe, MgSiO <sub>3</sub> ,
166	(Mg,Fe)SiO <sub>3</sub> , high-pressure $H_2O$ , C (graphite), and SiC; Dubrovinsky et al. (2000) for
167	$\epsilon\text{-}\mathrm{Fe};$ W. W. Anderson and Ahrens (1994) for liquid Fe; Fei et al. (1995) for FeS IV; Sata
168	et al. (2010) for FeS VII; Duffy, Hemley, and Mao (1995) for MgO; Weidner et al. (1982)
169	for SiO <sub>2</sub> ; Wett and Ahrens (1983) for $Mg_2SiO_4$ ; Chung (1971) for (Mg,Fe) <sub>2</sub> SiO <sub>4</sub> ; Duffy,
170	Meade, et al. (1995) for $Mg(OH)_2$ ; Tyburczy et al. (1991) for serpentine; Nestola et al.
171	(2010) for antigorite; Auzende et al. (2006) for lizardite; Fortes et al. (2013) for $Na_2SO_4 \cdot 10H_2O$ ;
172	Gromnitskaya et al. (2013) for MgSO <sub>4</sub> ·7H <sub>2</sub> O; Choukroun and Grasset (2010) for liquid
173	$H_2O$ and its ices Ih, II, III, V, and VI; Frank et al. (2013) for $H_2O$ ice VII; Fortes et al.
174	(2003) for $NH_3 \cdot 2H_2O$ I; Griffiths et al. (2012) for $NH_3 \cdot 2H_2O$ IIa; and Ma et al. (2012)
175	for $NH_3 \cdot 2H_2O$ liquid. A typical grid comprises $10^4$ zones. This is much finer than required
176	for this structure calculation, but necessary to resolve the upper mantle finely enough
177	as to accurately compute melting as a function of depth. The oversampling of deeper
178	layers seldom affects the speed of computation because the planetary structure is com-
179	puted only once at the onset of a simulation.

180

# 2.1.2 Thermal evolution

181 182 183

184

185

Heat sources include both radiogenic heat and heat from accretion, the two sources responsible for nearly all the heat trapped inside the Earth. A lower fraction of radiogenic heat implies a lesser dependence of mantle temperature on planet size and, especially, age (Kite et al., 2009). We factor accretional heating into the planet's initial temperature at mid-mantle depth,  $T_{m0}$ , following Canup et al. (2021):

$$T_{m0} = T_{surf0} + 0.6 \times 4 \ h \ \chi \ G \ \pi \ \rho \ r_p^2 / (3C_p) \tag{1}$$

**Table 2.** Radionuclide properties. Abundances pertain to the terrestrial planets of the Solar System and are given at the present day, 4.57 Gyr after the birth of the Solar System. The default abundances are those of Turcotte and Schubert (2002).

Property	$^{40}$ K	<sup>232</sup> Th	$^{235}$ U	$^{238}\mathrm{U}$	Reference	Notes
Half-life (Gyr)	1.26	14.0	0.704	4.47		
Heating rate ( $\mu$ W per kg nuclide)	) 29.2	26.4	569	94.6		
Abundance	18.0	51.9	0.10	14.2	Lyubetskaya and Korenaga (2007)	Low end-member
(ppb by mass)	36.9	124.0	0.22	30.8	Turcotte and Schubert (2002)	Highest heating
	30.7	84.1	0.15	21.0	Ringwood (1991)	Intermediate heating
	22.8	62.6	0.12	17.2	Lyubetskaya and Korenaga (2007)	Low heating
	28.8	79.5	0.15	20.15	McDonough and Sun (1995)	Intermediate heating

186 Here,  $T_{surf0}$  is an initial surface temperature, h is the fraction of impactor kinetic energy retained by the planet,  $\chi$  is the ratio of the impactor velocity relative to the escape 187 velocity from the accreting planet, G is the gravitational constant,  $r_p$  is the planet's fi-188 nal radius, and  $C_p$  is its bulk-averaged heat capacity, taken to be that of the mantle. Vol-189 ume averaging leads to the factor of 0.6. The parameters  $\chi$  and especially h are not well 190 constrained. For the Earth, simulations of the Moon-forming impact suggest that the 191 temperature of silicates reached 3000 to 6000 K (Canup, 2004), which necessitates  $h\chi \approx$ 192 0.2. We thus adopt this value for the  $h\chi$  product, which corresponds to 20% energy re-193 tained for average impactor velocities equivalent to the Earth's escape velocity of 11.2 194  $km s^{-1}$ , and noting that basing such a value on the Moon-forming impact may not be 195 representative of other Earths that did not experience such an impact. The higher ac-196 cretional heating experienced by larger (higher-R) planets tends to cause early vigorous 197 convection; the heat thus eliminated leads to these planets having counter-intuitively lower 198 lithospheric heat fluxes than an Earth-sized planet in the first 0.5 to 1 Gyr after accre-199 tion. Past 2 Gyr, the imprint of accretional heating on thermal evolution vanishes and 200 thermal evolution is dominated by the balance of radiogenic heating and convective or 201 conductive cooling, with larger planets having a hotter mantle and higher heat fluxes (see 202 Fig. 5c in Section 3). 203

Because radiogenic elements tend to partition with silicates rather than metals, heat 204 production is assumed to take place solely in the mantle. Radionuclide abundances de-205 pend on the bulk composition of the accreted material; we allow for user selection from 206 different sets of literature values (Table 2). Canonically, the abundances of Turcotte and 207 Schubert (2002) are assumed (Table 2); the lack of dependency of these values on a planet's 208 bulk composition is discussed in Section 4. Short-lived radionuclides are ignored since 209 the terrestrial planets have been thought to form after many short-lived radionuclide half-210 lives (Walsh & Levison, 2019), although recent models implying formation within just 211 a few million years may make short-lived radionuclides relevant early in the planet's his-212 tory (Bhatia & Sahijpal, 2016, 2017; Saito & Kuramoto, 2018; Johansen et al., 2021). 213 The radiogenic power per unit mass H (W kg<sup>-1</sup>) is expressed as: 214

$$H = \sum_{i=1}^{4} H_0(i) e^{-ln(2) t/t_{1/2}}$$
(2)

where i = 1 - 4 are the four main long-lived radioisotopes (<sup>40</sup>K, <sup>232</sup>Th, <sup>235</sup>U, <sup>238</sup>U),  $H_0$  is the product of a radionuclide's specific heating rate with its mass fraction in mantle material at the time the first planet-building solids condense – 4.57 billion years (Gyr) ago for the Solar System –,  $t_{1/2}$  is its half-life, and t is time.

The thermal profile from the surface to the base of the crust is calculated as follows. First, a mid-mantle temperature  $T_m$  is calculated at each time step with the assumptions of Kite et al. (2009) (e.g., whole-mantle convection), except thermal equilibrium is not assumed:

$$\frac{dT_m}{dt} = H/C_p - \frac{k}{\rho C_p} Nu \left(T_m - T_{ref}\right) \left(\frac{2}{r_{BDT} - r_c}\right)^2 \tag{3}$$

with the Nusselt number (dimensionless ratio of total heat flux to conductive heat flux) given by  $Nu = (Ra/Ra_{cr})^{\beta}$ . The Rayleigh number Ra is itself given by:

$$Ra = \frac{g \alpha (T_m - T_{ref}) (r_{BDT} - r_c)^3}{\kappa \nu}$$
(4)

Here, k is the thermal conductivity,  $T_{ref}$  is the surface temperature  $T_s$  in the plate tectonics regime, or the temperature at the base of the lithosphere (see equations 5 and 6) in the stagnant lid regime,  $r_c$  is the radius of the core-mantle boundary,  $r_{BDT}$  is the radius of the brittle-ductile transition (BDT), g is the gravitational acceleration,  $\alpha$  is the thermal expansivity,  $\kappa = k/(\rho C_p)$  is the thermal diffusivity, and  $\nu$  is the temperaturedependent kinematic viscosity.  $T_m$ , g, and  $\nu$  are evaluated halfway between  $r_c$  and  $r_{BDT}$ .

The radius of brittle-ductile transition  $r_{BDT}$  is that at which the temperature  $T_{BDT}$ and pressure  $P_{BDT}$  are such that the brittle and ductile strengths of mantle material are equal. It is found by using the previous time step's  $P_{BDT}$  and solving for  $T_{BDT}$  using a combined binary and Newton-Raphson search (Press et al., 1992). The brittle strength is given by 0.85*P* at low pressure (P < 200 MPa) and 0.6*P* + 50 MPa at P > 200MPa (Byerlee, 1978). The ductile strength is the ratio of the viscosity (see below) to the time step.

 $Ra_{cr}\approx 10^3$  is the critical Raleigh number. We adopt  $Ra_{cr}=1707.762$  from Reid 238 and Harris (1958). We also adopt  $\beta = 1/3$ , relevant to an isoviscous convective layer 239 (Solomatov, 1995); the viscosity variation with pressure and temperature across the man-240 tle may warrant values closer to 1/4 (Moresi & Solomatov, 1995; Deschamps & Sotin, 241 2000). Since that viscosity variation is not well constrained, especially for mantle com-242 positions that differ from Earth's, it may make sense to vary  $\beta$  between 1/4 and 1/3; but 243 we find that varying planet mass provides a sufficient range of thermal evolutions for the 244 purposes of this study. 245

The viscosity of Earth's mantle is assumed non-Newtonian, although this is uncertain (Deschamps & Sotin, 2000). The viscosity is a parallel combination of those obtained with the diffusion and dislocation creep flow laws for olivine given in Table 3 of Korenaga and Karato (2008), with stress equated to lithostatic pressure and the choice of dry (default) or wet mechanism left as a user input.

Based on the mid-mantle temperature  $T_m$ , a temperature profile is computed with 251 radius r from the surface to the core. This profile is assumed to have two parts: an adi-252 abat section in the asthenosphere, where heat transfer is convective, and a conductive 253 section in the lithosphere, the upper and only thermal boundary layer since the fluid is 254 radiogenically heated from within (Turcotte & Schubert, 2002). This boundary layer (Fischer 255 et al., 2010) is formally distinguished from the brittle layer  $(r > r_{BDT})$ , but in prac-256 tice the two have similar thickness. The temperature profile is computed as  $T(r) = min[T_{adiab}(r), T_{upbnd}(r)]$ , 257 with: 258

$$T_{adiab}(r) = T_m - \alpha \ g \ T_m [r - 0.5(r_{BDT} + r_c)]/C_p \tag{5}$$

$$T_{upbnd}(r) = T_m + (T_s - T_m) \ erfc\left[\frac{r_{BDT} - r}{2}\sqrt{\frac{2 \ v_{conv}}{\kappa(r_{BDT} - r_c)}}\right]$$
(6)

Equation 5 is equation 4.254 of Turcotte and Schubert (2002), evaluated at the mid-mantle depth halfway between  $r_c$  and  $r_{BDT}$ ; see also Katsura et al. (2010). Equation 6 is equation 6.347 of Turcotte and Schubert (2002), evaluated at the lateral midpoint between upwelling and downwelling regions of a convective cell (whose width and height are equal). In equation (6), erfc is the complementary error function, and the convective velocity  $v_{conv}$  is computed as in equation 6.379 of Turcotte and Schubert (2002):

$$v_{conv} = 0.354 \ \kappa \ Ra^{1/2} / (r_{BDT} - r_c) \tag{7}$$

which is appropriate for a fluid heated from within. A corresponding convection timescale is  $t_{conv} = (r_{BDT} - r_c)/v_{conv}$ .

The lithospheric thickness  $z_{Lith}$  is the depth at which  $T_{adiab} = T_{upbnd}$ . At the first time step, the temperature profile is computed using the same equations (5) and (6), initializing  $r_{BDT} = r_p$  and  $v_{conv} = (r_p - 100 \text{ km} - r_c)/10 \text{ Myr}$ .

The sensitivity of temperature profiles to mid-mantle depth and rheology for a 1- $M_{\oplus}$  planet is shown in Fig. 2. A hotter mantle results in a thinner lithosphere, and so does a wet olivine rheology, whose lower resulting viscosities result in more vigorous convection. For the adopted olivine rheologies, mantle temperatures of 2400-2500 K result in a profile that fits constraints for the Earth (Fischer et al., 2010).

The tectonic mode is determined by comparing the yield stress, equated to the brittle (and ductile) strength at the BDT, with the convective drive stress  $\nu/t_{conv}$ . If the drive stress exceeds the yield stress, plate tectonics proceed; otherwise the planet is assumed to have a stagnant lid. This very rough determination requires viscosities above  $\sim 10^{27}$  Pa s for plate tectonics to take place, which is not achieved in our simulations. The tectonic mode only has a minor effect on equation (4) but can significantly affect seafloor weathering, as investigated in the companion paper.



Figure 2. Validation of the melting and outgassing code with present-day Earth. Geotherms (temperature-depth profiles; top x-axis) for a  $1-M_{\oplus}$  planet are overlain on constraints for Earth's geotherm from analyses of mantle peridotites and measurements of surface heat flux (Turcotte & Schubert, 2002; Fischer et al., 2010). Also shown are melt fraction-depth profiles (bottom x-axis).

#### 2.1.3 Outgassing carbon flux

283

282

The outgassing flux, in mol  $s^{-1}$ , is assumed to be given by:

$$F_{outgas} = \frac{M_{melt} f_{C,magma}}{M_C t_{conv}}$$
(8)

where  $M_{melt}$  is the total mass of melt integrated over depth,  $f_{C,magma}$  is the fraction of the melt mass that exsolves as carbon gas, and  $M_C$  is the molar mass of CO<sub>2</sub>.

To obtain  $M_{melt}$ , a mantle melt fraction is computed as a function of depth using 286 the *alphaMELTS* command line implementation of the *MELTS* code (Ghiorso & Sack, 287 1995; P. M. Antoshechkina & Asimow, 2010; P. M. Antoshechkina et al., 2010; P. An-288 toshechkina & Asimow, 2018). A specified oxide composition (see Table 4 in Section 3) 289 is provided to *alphaMELTS*, along with settings instructing computation of chemical equi-290 librium along the above-determined temperature profile, from the surface to the depth 291 at which P = 10 GPa. The MELTS model is valid for P < 3 GPa, and alphaMELTS 292 allows (potentially unreliable) calculations beyond this limit up to 4 GPa. Calculations 293 show that there can still be partial melting at 4 GPa. In this case, our model implemen-294 tation extrapolates linearly the melt fraction trend over the 5 gridpoints closest to 4 GPa, 295 as determined with *alphaMELTS*, to higher pressures until the melt fraction is zero. Based 296 on comparison with an analytical melting model for Earth's upper mantle composition 297 (McKenzie & Bickle, 1988), for which the melt fraction decreases to zero with depth (pres-298 sure) faster than linearly, this may slightly overestimate the amount of melt at high pres-299 sure. However, these amounts tend to be small relative to  $M_{melt}$ . 300

For the temperature profiles of Fig. 2, the rates of crustal generation from all melt,  $M_{melt}/(4\pi r_p^2 \rho_{melt} t_{conv})$  with the melt density  $\rho_{melt}$  obtained from *alphaMELTS*, are 30.1 ( $T_m = 2500$  K), 11 ( $T_m = 2450$  K), 1.23 ( $T_m = 2400$  K), and 47 m Myr<sup>-1</sup> ( $T_m = 2400$  K, wet rheology) compared to Earth's estimated 40 m Myr<sup>-1</sup> (Kite et al., 2009). These illustrate a trade between best reproducing Earth's temperature profile with depth (profiles with thicker lithosphere) and best reproducing its rate of crust generation (profiles with thinner lithosphere).

On modern Earth, the magma  $CO_2$  content  $f_{C,magma}$  has been variously reported to be 0.03–1 mass% (Hekinian et al., 2000; Jones et al., 2018) in mid-ocean ridge basalts, 0.1–0.65 mass% in ocean island basalts that also possibly approximate Archean Earth

-14-

erupted material (Gerlach & Graeber, 1985; A. Anderson, 1995; Hartley et al., 2014),

and 0.004-0.15 mass% in back-arc volcanism (Cioni, 2000; Saito et al., 2001; Wade et al.,

<sup>313</sup> 2006; Blundy et al., 2010). Adopting a canonical abundance of 0.2 mass%, which is within

- those ranges, yields an outgassing flux similar to that of modern Earth for a 1 Earth-
- mass, 4.57 Gyr-old planet whose upper mantle has a mid-ocean ridge basalt composi-
- $_{316}$  tion (see Fig. 5 in Section 3).

Outgassed carbon can be speciated into CO<sub>2</sub> and CH<sub>4</sub>, with the relative fraction 317 of methane  $f_{CH4}$  left as a user input and constant through a simulation. We approxi-318 mate  $f_{CH4} = 0$  for modern and Archean Earth cases, and  $f_{CH4} \ge 0.5$  for more metal-319 rich planetary mantles. For context, the redox state of Hadean Earth magmas was near 320 the fayalite-magnetite-quartz (FMQ) redox buffer (Trail et al., 2011), but a higher pro-321 portion of submarine (i.e., high-pressure) volcanism would have affected the composi-322 tion of the erupted gases (Gaillard et al., 2011), with low (atmospheric)-pressure out-323 gassing likely to be uniformly near FMQ-2 over a range of body sizes, assuming a sim-324 ilar solar-system-like bulk basaltic composition for rock-metal bodies (Gaillard & Scail-325 let, 2014). In this case, the prime carbon gas erupted is CO<sub>2</sub> (Gaillard & Scaillet, 2014). 326 For undifferentiated or more metal-rich objects, at 1 bar, the predominant carbon gases 327 are CH<sub>4</sub> below 750 K and CO above 750 K (Schaefer & Fegley, 2017). 328

329

# 2.2 Continental weathering

330

# 2.2.1 Geochemical kinetics

Weathering describes the physicochemical process which modifies rocky surface landforms by its physical and chemical interaction with fluids having solubilized atmospheric material. As fluids are usually richer in carbon than rock, weathering reactions tend to result in transfer of carbon from the fluid to the rock. A relevant reaction is (Urey, 1952):

$$CaSiO_3 + CO_2(aq) = CaCO_3 + SiO_2$$
  
wollastonite calcite silica (9)

where (aq) indicates  $CO_2$  in its aqueous form. The fluid-rock setting can be subaerial crust exposed to rain or running water (continental weathering), or suboceanic crust in contact with ocean water (seafloor weathering). **Table 3.** Key model parameters for calculating carbon fluxes. Fluxes in mol m<sup>-2</sup> s<sup>-1</sup> can be obtained by dividing  $F_i$  by  $4\pi r_p^2$ . <sup>*a*</sup> User-specified inputs. <sup>*b*</sup> User-specified input via river runoff rate.

Parameter	Symbol	Value	Units	Notes and references
Outgassing				
Outgassing flux	$F_{outgas}$		$ m mol~s^{-1}$	From equation $(8)$
Mass of mantle melt integrated over depth	$M_{melt}$		kg	From <i>alphaMELTS</i>
Magma carbon fraction $^{a}$	$f_{C,magma}$	0.002	$\rm kg \ kg^{-1}$	
Relative fraction of C outgassed as CH4 $^a$	$f_{CH4}$		$\rm kg \ kg^{-1}$	0 for Modern and Archean Earth
				$\geq 0.5$ for chondritic mantle
Continental weathering				
Continental weathering flux	$F_{contw}$		$ m mol~s^{-1}$	From equation (11)
Rainfall rate $^{b}$	p	0.7	${\rm m~yr^{-1}}$	Broecker and Peng (1982)
Fresh- and seawater density	ho	1000	${\rm kg}~{\rm m}^{-3}$	
A real fraction of planet covered by land $^{a}$	L		_	0.29 for Earth
Freshwater:rock mass ratio	$4\pi r_p^2 \ L \ p \ \rho/M_{rock}$	5000	${\rm kg \ kg^{-1}}$	Martin and Meybeck (1979)
Molalities of dissolved species in rivers	$m_{i,river}$		$ m mol~kg^{-1}$	From <i>PHREEQC</i> kinetic calculation
Continental weathering				
Seafloor weathering flux	$F_{seafw}$		$ m mol~s^{-1}$	From equation $(13)$
Mass of surface ocean $a$	$M_{ocean}$	$1.4\times 10^{21}$	kg	
Hydrothermal circulation timescale	$t_{circ}$	$10^{7}$	$\mathbf{yr}$	Mottl (1983); Kadko et al. $(1995)$
Length of plate boundaries	$L_{plate\_ridges}$		m	$1.5\times 2\pi r_p$ for modern Earth
where seafloor is created				$(4.5 - 7.5) \times 2\pi r_p$ for Archean Earth
Depth of fracturing below seafloor	$z_{crack}$	6	$\rm km$	Vance et al. $(2007)$
Molalities of dissolved species in the ocean	$m_{i,ocean}$		$ m mol~kg^{-1}$	From $PHREEQC$ and eq. (14)
River water mass flowed to ocean in a time step	$M_{river}$		kg	
Carbon incorporated in the crust	$\Delta C_{reac}$		mol	From equation $(16)$
Atmospheric evolution				
Net carbon flux	$F_C$		$ m mol~s^{-1}$	From equation $(17)$
${\rm Atmosphere+ocean}$ reservoir	$RC_{atm+oc}$		mol	
Mantle reservoir	$RC_{mantle}$		$\operatorname{mol}$	
Time step $^{a}$	$\Delta t$		s	Adapted based on $RC_i$ and $F_C$
Surface pressure $^{a}$	$P_{surf}$		Pa	Varies during simulation
Surface temperature $^{a}$	$T_{surf}$		Κ	Varies during simulation
Number of moles of gas in atmosphere	$n_{air}$		mol	
Atmospheric mixing ratios $a$	$x_i$		-	$i = CO_2, CH_4, N_2, O_2, H_2O;$
				vary during simulation
Atmospheric volume	$V_{atm}$		$m^3$	Held constant

In planetary-scale models of the carbonate-silicate cycle, continental weathering fluxes are usually assumed to vary with temperature as an Arrhenius law, with runoff (the amount of rainwater involved in weathering reactions and that eventually transports dissolved or particulate surface rock to the ocean), and with atmospheric CO<sub>2</sub> partial pressure. The variation with the latter two parameters is usually expressed as power law deviations from measured Earth values, as in equations 8a, 2, and 1, respectively, of Sleep and Zahnle (2001),

In natural systems, crustal materials are composed of a variety of minerals that all 345 react with carbon-bearing fluids at different rates. To capture the effect of surface com-346 position on reaction rates, new power laws would have to be determined for every sur-347 face composition, an impractical challenge. Instead, we opt to compute reaction rates 348 and corresponding continental weathering fluxes from first principles, based on the re-349 sults of simulations of water-rock interaction through time using the KINETICS function-350 ality of the PHREEQC software (Parkhurst & Appelo, 2013). PHREEQC database and 351 example input files are provided as supplementary material to this article in a Zenodo 352 repository, and data and input compositions are described below. 353

Chemical reactions, including those involved in continental weathering, proceed at 354 a rate that depends on numerous factors such as temperature, the extent of chemical dis-355 equilibrium, concentrations of species such as H<sup>+</sup> (pH), mineral surface properties (Pokrovsky 356 & Schott, 1999), and the action of catalysts. For the latter two processes, a thorough 357 understanding is lacking for a comprehensive set of relevant rock species. The depen-358 dence on temperature and composition is better understood, but experimental data have 359 been compiled only for a subset of species that are abundant at Earth's surface (Palandri 360 & Kharaka, 2004). As a result, the calculation of reaction rates is typically captured into 361 a simplified form, fitted to experimental measurements, such as (Palandri & Kharaka, 362 2004): 363

$$\frac{dm}{dt} = -\text{SA } A \ e^{-\frac{E}{R(T-T_0)}} \ a_{H^+}^n \ (1-\Omega^p)^q \tag{10}$$

Here, m represents the concentration (molality, in moles of species per kg of water solvent) and dm/dt its change over time, SA the surface area available for reaction, E an activation energy, R the gas constant, T the temperature (which influences reaction rates via an Arrhenius law), and  $T_0$  a reference temperature, usually 298.15 K.  $\Omega$  is a mea-

sure of chemical disequilibrium, equal to Q/K, where  $Q = \prod_i a_i^{\nu_i}$  is the product of the 368 species' thermodynamic activities  $a_i$ , scaled geometrically by their stoichiometric coef-369 ficient  $\nu_i$  (negative for reactants, positive for products), and K(T, P) is this product when 370 the reaction reaches equilibrium (G. M. Anderson, 2005). K is termed the reaction con-371 stant, but depends on temperature T and pressure P. The magnitude of  $\Omega$  hinges both 372 on the bulk initial compositions of the fluid (e.g., carbon content) and rock, and on how 373 far toward equilibrium the reaction has progressed. The constant A and the exponents 374 n, p, and q are determined experimentally; here, p and q are both approximated to 1 for 375 all species (Palandri & Kharaka, 2004). 376

Reaction constants K are obtained from the *core10.dat* thermodynamic database 377 described in Neveu et al. (2017), with data from references therein, distributed with PHREEQC 378 at https://wwwbrr.cr.usgs.gov/projects/GWC\_coupled/phreeqc. For this study, we have 379 augmented the database with kinetic rates for 56 solids compiled by Palandri and Kharaka 380 (2004) and references therein. These include quartz and silica, as well as mineral species 381 in the feldspar, olivine, garnet and other orthosilicates, cyclosilicate, pyrox-382 enoid, amphibole, mica, clay and other phyllosilicates, oxide, hydroxide, carbonate, sul-383 fate, sulfide, and halide groups. 384

Not all solids need be part of the reacting crust, but all are allowed to form in a 385 simulation. One exception is quartz, included in weathering simulations involving mod-386 ern Earth and the solar system surfaces of Table 5, but not allowed to form in those in-387 volving the surface composition of Archean Earth. This is because quartz is not reac-388 tive on relevant timescales (residence time of rain and river water on land) and as such 389 tends to make the system of differential equations (10) too stiff for *PHREEQC*'s imple-390 mentation of the CVODE solver (Cohen et al., 1996) to handle, preventing these simula-391 tions from proceeding. 392

The starting continental crust (Table 5) and rainwater fluid compositions are provided as inputs to *PHREEQC*. The composition of rainwater, including its carbon content, is set to be equilibrated with the atmospheric composition, assuming that the atmosphere is an infinite reservoir. Atmospheric  $N_2$  and its dissolved form are assumed to be inert.

A fluid:rock mass ratio of about 5000 is canonically assumed, although the diversity of terrains on Earth points to much variation from this global mean. This ratio is

-18-

based on a reference river runoff rate for Earth of 0.665 mm day<sup>-1</sup> (Edson et al., 2012), 400 which corresponds to a rainfall rate  $p = 0.7 \text{ m yr}^{-1}$  minus evaporation of 65% of the rain-401 water from rivers (Broecker & Peng, 1982; Martin & Meybeck, 1979). Multiplying by 402 land surface area  $4\pi r_p^2 L$ , with L the fraction of planet area covered by land (L = 0.29403 for Earth), and fluid density  $\rho\approx 1000~{\rm kg}~{\rm m}^{-3}$  yields a rainfall mass rate  $4\pi r_p^2 \;L\;p\;\rho=$ 404  $1.0 \times 10^{17}$  kg yr<sup>-1</sup>. The mass of parent rock affected by weathering at the continent sur-405 face drained by rivers to the ocean is estimated to be  $M_{rock} = 21.1 \times 10^{12} \text{ kg yr}^{-1}$  in 406 Table V, footnote 3 of Martin and Meybeck (1979). This yields a fluid:rock ratio  $4\pi r_p^2 L p \rho/M_{rock}$ 407 = 4918. The ratio is proportional to p but insensitive to  $4\pi r_p^2 L$  since  $M_{rock}$  too is pro-408 portional to this term. In our modern and Archean Earth simulations, the fluid:rock ra-409 tios are 5000 and 4944, respectively. 410

2.2.2 Estimating continental weathering carbon fluxes

411

The *PHREEQC* simulation yields fluid and mineral abundances as a function of 412 reaction time, assuming a chemically closed system (no removal of reactants or products). 413 In the full model (Fig. 1), all solutes are delivered to the ocean where they accumulate 414 and, once saturation is reached, form carbon-bearing minerals that upon subduction trans-415 fer carbon from the {atmosphere+ocean} reservoir back to the mantle reservoir (Sec-416 tion 2.3). This approach provides a level of chemical realism not achieved with treatment 417 of distinct continental and seafloor weathering fluxes of carbon, but does not single out 418 the continental contribution to the weathering flux and its dependency on surface com-419 position. 420

For the purposes of this study, we therefore calculate a continental weathering flux  $F_{contw}$ , in mol yr<sup>-1</sup>, of dissolved carbon that can be sequestered in minerals based on amounts (molalities)  $m_{i,river}$  of Ca, Mg, and Fe cations in mol per kg of runoff water after a reaction time equal to the mean residence time of rainwater on land, about 10-15 years on modern-day Earth (Begemann & Libby, 1957; Chahine, 1992). The flux is then:

$$F_{contw} = \sum_{i} j m_{i,river} 4\pi r_p^2 L p \rho$$
(11)

Here, *i* is a summation index on all combinations of aqueous Ca, Mg, Fe, arising from
the dissolution of silicate, carbonates, sulfates, and sulfides present in the system. The

stoichiometric integer j is equal to the balance of positive and negative charges among 429 the ions (other than  $H^+$  and  $OH^-$ ) arising from the dissolution of these minerals: +1, 430 +2, 0, and -2 for carbonates, silicates, sulfates or pyrrhotite, and pyrite, respectively. 431 Indeed, their dissolution yields a doubly charged cation plus, respectively,  $HCO_3^-$ , aque-432 ous  $SiO_2$ ,  $SO_4^{2-}$ , or 2  $SO_4^{2-}$  at the river and ocean pH and oxidation conditions of mod-433 ern Earth (Berner et al., 1983). The contribution of Na and K cations is neglected be-434 cause their bicarbonate salts are too soluble to sequester carbon. Each excess positive 435 charge can combine with  $HCO_3^-$  or  $1/2 SO_4^{2-}$  from rainwater. Twice the rainwater  $SO_4^{2-}$ 436 flux is thus removed from the total excess positive charges to obtain  $F_{contw}$ . 437

While this approach to quantifying continental weathering flux provides a basis for determining its sensitivity to surface composition, it is inadequate if the atmosphere, surface, and ocean compositions lead to a predominant dissolved carbon species with a different charge than bicarbonate, such as dissolved  $CO_2$  (carbonic acid, no charge) or carbonate  $CO_3^{2-}$  (doubly charged anion) at ocean pH lower than 5 or greater than 10 at Earth surface T and P, respectively, or dissolved organic carbon or  $CH_4$  (no charge) at reducing conditions. Those cases warrant use of the full model of Fig. 1.

445

#### 2.2.3 Validation: modern Earth

We validate and calibrate PHREEQC kinetic simulations of continental weather-446 ing at 288 K and 1 bar with estimates of river abundances of major rock-forming ele-447 ments (Martin & Meybeck, 1979) and corresponding continental weathering fluxes  $F_{contw}$ 448 for modern Earth (Berner et al., 1983; Lerman & Wu, 2006; Colbourn et al., 2015) at 449 reaction (residence) times of 10-15 years (Begemann & Libby, 1957; Chahine, 1992). Abun-450 dances of major rock-forming elements in rivers are obtained by dividing simulated mo-451 lalities  $m_{i,river}$  by 1 - 0.65 = 0.35 to account for 65% evaporation on land (Broecker 452 & Peng, 1982). 453

A felsic crust composition (see 'Earth (modern)' column of Table 5 in Section 3) is set by adjusting starting abundances of quartz; Na-, Ca-, and K-feldspars; phyllosilicates (annite, phlogopite, daphnite); and hornblende (anthophyllite and tremolite) to match mineral volume fractions from Nesbitt and Markovics (1997) and oxide mass fractions from Rudnick and Gao (2003) within 10%. To this, we add calcite and dolomite (which rapidly and fully dissolve) to obtain carbonate contributions of Ca and Mg cations

-20-



Figure 3. Validation of continental weathering model against modern Earth average river compositions and weathering fluxes. **a.** PHREEQC simulated molalities  $m_{i,river}$  of dissolved Na, Mg, Si, Ca, and Fe (solid lines) at 288 K and 1 bar, divided by (1 (0.65) to account for 65% evaporation, compared to measured averaged river abundances (dashed lines) at relevant residence times of 10 to 15 years (red window). b. Corresponding river pH. c. Resulting total net flux (thick black line), obtained from the  $m_{i,river}$  shown in (a) using equation (11), and contributions to this flux from various mineral groups (subsets of the terms summed in equation 11), compared to literature ranges (shaded bands and lines inside the red box). The total flux curve is corrected to exclude about 2.4 Tmol  $yr^{-1}$  of cation fluxes, which are assumed to react with the corresponding flux of sulfate from rainwater and sulfide weathering generated by the model (Berner et al., 1983). Because this approach does not attribute the total sulfate flux contribution to specific mineral groups, the curves showing mineral group contributions include this sulfate flux contribution and therefore slightly overestimate the contribution of each group to the  $CO_2$  drawdown flux. Another 2.4 Tmol yr<sup>-1</sup> of sulfate arises from weathering of sulfate minerals, chiefly gypsum, but the sulfate and calcium arising from it are not shown here as together they do not affect the charge balance. <sup>a</sup> Martin and Meybeck (1979). <sup>b</sup> Berner et al. (1983). <sup>c</sup> Begemann and Libby (1957). <sup>d</sup> Chahine (1992). <sup>e</sup> Lerman and Wu (2006). <sup>f</sup> Colbourn et al. (2015).

in the ranges given by equations 10 and 17 of Berner et al. (1983) and by Table S1 of

461 Colbourn et al. (2015) and references therein. The added amounts correspond to 5.3 vol%

of exposed land, within the 4.2 - 8.8 area% range estimated for the past 500 Myr on

Earth,  $(6.2 - 13) \times 10^6$  km<sup>2</sup> (Kiessling et al., 2003), although part of this area is under-

- water on continental shelves. We also add sulfate (gypsum) and sulfide (pyrite) at abun-
- dances that match their estimated modern contributions to river sulfate; see p. 645 of
- 466 Berner et al. (1983).

The liquid composition is set to be equilibrated with partial pressures of 0.78 atm 467 of  $N_2$ , 0.2 atm of  $O_2$ , and 4000  $\mu$ atm of  $CO_2$ . The  $CO_2$  partial pressure, within the sev-468 eral thousand  $\mu$ atm range measured for large rivers (Cole & Caraco, 2001), reflects su-469 persaturation relative to the atmosphere (pre-industrial  $pCO_2$  of 280  $\mu$ atm) due to bi-470 ological respiration of biogenic reduced carbon from soil (Rasilo et al., 2017; Lynch et 471 al., 2010). The liquid also comprises 8  $\mu$ mol kg<sup>-1</sup> of sulfate, corresponding to the esti-472 mated  $0.8 \text{ Tmol yr}^{-1}$  pre-industrial sulfur contribution from outgassing and atmospheric 473 cycling (Berner et al., 1983). 474

<sup>475</sup> *PHREEQC* calculations with this input reproduce both a set of average cation con-<sup>476</sup> centrations in Earth rivers (Fig. 3a) and reported ranges of the weathering flux, includ-<sup>477</sup> ing contributions from various mineral groups (largely Ca- and Mg- silicates and carbon-<sup>478</sup> ates) (Fig. 3c). The computed river water sulfur abundance (constant with time) of  $6.9 \times$ <sup>479</sup>  $10^{-5}$  mol kg<sup>-1</sup> S, which corresponds to 6.6 mg kg<sup>-1</sup> SO<sub>4</sub><sup>2-</sup>, is in the reported range of <sup>480</sup> 5.19 - 11.2 mg kg<sup>-1</sup> SO<sub>4</sub><sup>2-</sup> of Lerman et al. (2007) and references therein.

- A few differences stand out. First, in the simulation nearly all Fe is in minerals. 481 Measured dissolved river Fe molalities may be non-negligible only because of (biogenic) 482 organic complexation (Gibbs, 1975), and can only be matched in a simulation if precip-483 itation of Fe oxides is prevented. Second, the simulated river pH past 10 years of inter-484 action is at the high end of typical measured river values (Fig. 3b). Third, the Ca-carbonate 485 (calcite) contribution to weathering flux is lower than estimated by Berner et al. (1983). 486 However, the Mg-carbonate contribution matches their estimate and the total carbon-487 ate contribution is in the middle of the range reported in Table S1 of Colbourn et al. (2015). 488
- To convert dissolved abundances to weathering fluxes we have assumed L = 0.29, irrespective of whether this land is drained by rivers to the oceans (i.e., exorheic). Exorheic areas have been variously estimated as comprising half to two thirds of the con-

-22-

tinental area on modern Earth (Lerman et al., 2007). Excluding Greenland and Antarctica, on which short-term weathering is impeded by the ice cover, but including endorheic areas in the ratio of surface water reservoirs to riverine fluxes (Chahine, 1992) results in L = 0.26, thus lowering fluxes of Fig. 3c by 10%.

In this calculation, we altered the dissolution kinetics of feldspar along the solid 496 solution between Ca (anorthite, fastest dissolving) and Na (albite, slowest dissolving) 497 end-members in order to limit the rate of Ca dissolution. Otherwise, the additional Ca 498 in solution increases the pH, limiting dissolved Mg. We found that attributing 85% of 499 anorthite the kinetic law for labradorite (50-70% anorthite, 30-50% albite) and the other 500 15% the law for bytownite (70-90% anorthite, 10-30% albite) achieved a suitable balance 501 between dissolved Ca and dissolved Mg as shown in Fig. 3c. We did not change the dis-502 solution rate for albite, assumed to be the end-member's law, but instead controlled the 503 amount of Na in solution with NaCl (see Table 5 in Section 3). 504

When applying this model to other surface and atmospheric compositions in lifeless conditions, we restore  $pCO_2$  to the atmospheric value, scale the 8  $\mu$ mol kg<sup>-1</sup> of dissolved atmospheric sulfur by the ratio of  $pCO_2$  to Earth's modern  $pCO_2$ , and retain the 85% labradorite – 15% bytownite kinetic rate law for anorthite. Implicit in the second choice is the simplifying assumption that atmospheric CO<sub>2</sub> and SO<sub>2</sub> co-vary; the model does not explicitly track atmospheric SO<sub>2</sub>.

511

## 2.3 Seafloor weathering

Seafloor weathering describes the loss of carbon from the {atmosphere+ocean} reservoir to the mantle reservoir. If the planet does not experience plate tectonics, dissolved species delivered by rivers continue to accumulate until saturation is reached and minerals precipitate. If plate tectonics operate, ocean fluids can react with fresh (unreacted) new ocean crust exposed at the seafloor.

In either case, it is sensible to assume that unlike continental weathering reactions, ocean-seafloor reactions proceed to chemical equilibrium. If ocean crust is recycled, the recycling timescale depends, to first order, on the vigor of mantle convection. The current rates of production of oceanic crust are about 20 km<sup>3</sup> yr<sup>-1</sup>  $\approx 2 \times 10^{6}$  kg s<sup>-1</sup> on Earth (Cogné & Humler, 2006), with the mantle convecting on a timescale of  $10^{8}-10^{9}$ years. This timescale holds for the larger, younger, or compositionally different planets

in the simulations whose outgassing fluxes are shown in Fig. 5. This is slower than cir-523 culation of ocean fluids through fractures or porosity in this crust, which occurs on a timescale 524  $t_{circ} \sim 10^7$  years (Mottl, 1983; Kadko et al., 1995). It is also slower than chemical equi-525 librium between these fluids and crustal rock: a *PHREEQC* kinetic simulation of reac-526 tion between a fluid with modern-day seawater composition from Parkhurst and Appelo 527 (2013) (simplified from Nordstrom et al. (1979)) and mafic crust composition (see Ta-528 ble 5, 'Earth (Archean)' column, in Section 3) shows that chemical equilibrium is reached 529 on timescales of  $10^5 - 10^6$  years (Fig. 4). This equilibrium is calculated at the ocean 530 surface temperature (a lower bound), 405 bar (depth of about 4 km), and a water:rock 531 mass ratio of 1 (see below). 532

The reaction takes place at seafloor pressure and surface temperature, in the approximation of an isothermal ocean. Water-rock reactions at or below the seafloor can, but need not, take place at elevated temperatures relative to that of the ocean. The water:rock ratio by volume is determined by:

$$\frac{V_{fluid}}{V_{ocean\ crust}} = \frac{(M_{ocean}/\rho) \times \Delta t/t_{circ}}{\frac{1-L}{1-0.29}\ L_{plate\_ridges\ v_{conv}\ z_{crack}\ \Delta t}}$$
(12)

This ratio is independent of the time step  $\Delta t$ . Here,  $L_{plate\_ridges}$  is the length of 537 plate boundaries where new seafloor is created (this excludes subduction zones). On mod-538 ern Earth, this length is about 1.5 times Earth's circumference. We scale it for differ-539 ent mantle convective vigor as  $L_{plate\_ridges} = (Ra/2.3 \times 10^6)^{\beta} \times 1.5 \times 2\pi r_p$ , which is 540 consistent with a linear proportion to the Nusselt number and with a 3 to 5 times greater 541 ridge length in the Archean (Kadko et al., 1995).  $v_{conv}$  is the plate velocity, which can 542 simply be equated to the mantle convective velocity (Turcotte & Schubert, 2002), or ar-543 bitrarily decreased from that value to simulate more sluggish tectonics as may have been 544 the case at the onset of plate tectonics (Brown et al., 2020).  $z_{crack}$  is the depth into the 545 seafloor to which fluid can circulate, with estimates for the present day varying between 546 600 m (Johnson & Pruis, 2003) and 6 km (Vance et al., 2007) (see also Hasenclever et 547 al. (2014)), with deeper circulation appearing possible if plate velocity is very low (Tao 548 et al., 2020). We adopt  $z_{crack} = 6$  km. 549

The seafloor weathering flux, in moles per unit time, is the difference  $\Delta C_{reac}$  in the molality of aqueous carbon  $m_{C,ocean}$  in the ocean pre- vs. post-reaction (mol kg<sup>-1</sup>), multiplied by the ocean mass  $M_{ocean}$  and divided by the time step:



Figure 4. Abundances of minerals and solutes as a function of time (lines) in a kinetic simulation at 288 K and 405 bar of the interaction between seafloor rock and seawater at a mass ratio of 1. These abundances, compared to those computed for an equilibrium simulation (thick dots), indicate that equilibrium is approached at about  $10^5$  years, although some Al- and S-rich minerals such as, respectively, clinochlore and pyrite have not yet reacted to diaspore and pyrrhotite.

$$F_{seafw} = \Delta C_{reac} \ M_{ocean} / \Delta t \tag{13}$$

The pre-reaction ocean molality  $m_{i,ocean}$  of each element is an average of the ocean composition at the previous time step  $(m_{i-1,ocean})$  and that of riverine input (molalities  $m_{i,river}$ ), scaled by the relative masses of the ocean and river runoff over a timestep:

$$m_{i,ocean,t} = \frac{m_{i,ocean,t-1} \ M_{ocean} + m_{i,river} \ M_{river}}{M_{ocean} + M_{river}}.$$
(14)

 $M_{ocean}$  is a user input (default  $1.4 \times 10^{21}$  kg for modern Earth) and is assumed constant with time. The mass of river water  $M_{river}$  flowing into the ocean during a timestep is given by:

$$M_{river} = 4\pi r_p^2 \ L \ \frac{p}{1 - 0.65} \ \rho \ \Delta t \tag{15}$$

<sup>559</sup> Only a subset  $\Delta t/t_{circ}$  of the ocean reacts with the seafloor; the resulting fluid is <sup>560</sup> then mixed back with the rest of the ocean to update the ocean composition. Follow-<sup>561</sup> ing this mixing step, mineral precipitation is allowed in case the ocean has become su-<sup>562</sup> persaturated; the resulting fluid composition is the ocean composition post-reaction. This <sup>563</sup> allows determination of  $\Delta C_{reac}$  as:

$$\Delta C_{reac} = m_{C,ocean,post-reaction} - m_{C,ocean,pre-reaction} \tag{16}$$

where  $m_{C,ocean}$  refers to the total molality of dissolved carbon, summed over all dissolved carbon species (including but not limited to CO<sub>2</sub>, CH<sub>4</sub>, HCO<sub>3</sub><sup>-</sup>, and CO<sub>3</sub><sup>2-</sup>).

566 River-ocean and hydrothermal fluid-ocean mixing simulations are carried out us-

<sup>567</sup> ing *PHREEQC*'s MIX functionality, which allows consistent tracking of pH and redox con-

ditions through the mixing reactions. The pressure of reaction is adjusted using the REACTION\_PRESSURE

keyword. Fluid-rock equilibria are calculated using the EQUILIBRIUM\_PHASES function-

- ality, either in the presence of seafloor rock (assumed, e.g., to have the mafic Archean
- crust composition given in Section 3's Table 5) if plate tectonics occur, or without re-
- active rock otherwise. In the latter case, transfer of carbon from the ocean to minerals
- occurs if the ocean composition following riverine input is such that carbon-bearing min-

erals are supersaturated, as expected if the ocean is already in equilibrium with a carbon-

<sup>575</sup> bearing atmosphere prior to riverine input.

For an ocean saturated in carbon both before the addition of riverine input and after seafloor interaction and precipitation of any supersaturated minerals,  $\Delta C_{reac}$  is proportional to  $\Delta t$ . Therefore,  $F_{seafw}$  is independent of  $\Delta t$ , as confirmed with simulations involving seafloor weathering with various time steps (see companion paper).

580

#### 2.4 Net carbon flux and atmospheric evolution

The net flux  $F_C$  of carbon between the {atmosphere+ ocean} and mantle reservoirs is the sum of  $F_{outgas}$ , the outgassing flux given by equation (8), and  $F_{seafw}$  given by equation (13). The latter term accounts for contributions from continental weathering via the term  $\Delta C_{reac}$  (see equations 14 and 16), and as such represents the effects of both continental and seafloor weathering. If plate tectonics operate, this term is scaled by a fraction  $f_{arc}$  (default value 0.25 for Earth) that accounts for partial outgassing of subducted carbon via back-arc volcanism. Thus:

$$F_C = F_{outgas} + (1 - f_{arc}) F_{seafw}.$$
(17)

The {atmosphere+ocean} and mantle reservoirs of carbon, respectively  $RC_{atm+oc}$ and  $RC_{mantle}$  in mol, are respectively incremented or decremented at each timestep by  $F_C \Delta t$ . A maximum timestep is set as a user input (canonically, 1 Myr) and decreased if needed during a simulation to  $0.1 \times min(RC_{atm+oc}, RC_{mantle})/|F_C|$  in order to maintain numerical stability.

At the first time step,  $RC_{atm+oc}$  is initialized by specifying initial atmospheric par-593 tial pressures of  $CO_2$  and  $CH_4$  and using *PHREEQC* to determine the corresponding 594 (equilibrium)  $m_{C,ocean}$ . The initial  $RC_{mantle}$  is derived from the mantle mass (Section 595 2.1.1) and the canonical assumption that the mantle comprises 200 ppm C by mass. This 596 is in the inferred range of 20-1800 ppm (Dasgupta & Hirschmann, 2010) and implies de-597 pletion by a factor of 10 relative to the abundance of carbon in mantle melt of 0.2% (Sec-598 tion 2.1.3). Thus, there is canonically about  $10^3$  times more carbon in the mantle than 599 in the {atmosphere+ocean} reservoir (see Fig. 7 in Section 4). 600

The atmospheric surface pressure  $P_{surf}$  and mixing ratios  $x_{CO2} = pCO_2/P_{surf}$ and  $x_{CH4} = pCH_4/P_{surf}$  are updated at each time step based on  $F_C$  and  $f_{CH4}$ . This is done by equilibrating the atmospheric and ocean compositions using *PHREEQC* at surface temperature  $T_{surf}$  (fixed in this *PHREEQC* calculation) and pressure  $P_{surf}$  (modified by this *PHREEQC* calculation based on gas exchange with the ocean). Pre-equilibration conditions are set according to, respectively:

$$\Delta P_{surf} = F_C \ \Delta t \ R \ T_{surf} / V_{atm} \tag{18}$$

$$x_{CO2,t+1} = \frac{x_{CO2,t} \ n_{air} + (1 - f_{CH4}) F_C \Delta t}{n_{air} + F_C \Delta t}$$
(19)

$$x_{CH4,t+1} = \frac{x_{CH4,t} \ n_{air} + f_{CH4} F_C \Delta t}{n_{air} + F_C \Delta t} \tag{20}$$

where  $V_{atm}$  is the atmospheric volume and  $T_{surf}$  the surface temperature.

 $V_{atm}$  is held constant throughout the simulation, and computed at the first timestep as  $V_{atm} = n_{air} R T_{surf}/P_{surf}$ . The number of moles of air  $n_{air}$  is initiated as  $n_{air} =$  $P_{surf} \times 4\pi r_p^2/(g_{surf} M_{air})$ , with  $M_{air}$  the average atmospheric molar mass (calculated from the user-input atmospheric partial pressures of CO<sub>2</sub>, CH<sub>4</sub>, N<sub>2</sub>, O<sub>2</sub>, and H<sub>2</sub>O) and  $g_{surf}$  the surface gravity.  $n_{air}$  is incremented at each time step by  $F_C \Delta t$ .

The model is intended for coupling with models of atmospheric photochemistry and 613 radiative transfer (Kasting & Ackerman, 1986; Kopparapu et al., 2013; Arney et al., 2016; 614 Vidaurri et al., 2022). However, its implementation can be run independently, in which 615 case  $T_{surf}$  is computed by summing a term from insolation, which changes over geologic 616 timescales, and a term capturing the atmospheric greenhouse effect. Both terms are pa-617 rameterized as in Caldeira and Kasting (1992). This parameterization is valid for  $10^{-8}$ 618 bar  $< pCO_2 < 10^{-2}$  bar. For standalone simulations, we also roughly parameterize 619 the rainfall rate p as  $p \propto T_{surf}^{1.025}$  (with  $p = 0.7 \text{ m yr}^{-1}$  for  $T_{surf} = 288 \text{ K}$ ), reflecting 620 a roughly 2.5% increase in global mean precipitation on Earth per Kelvin of tempera-621 ture increase (Allen & Ingram, 2002; Trenberth et al., 2005; Pendergrass, 2020). An ex-622 ample such standalone simulation is provided in Section 4. 623

#### <sup>624</sup> 2.5 Neglected processes

The ocean mass is approximated as constant over time, even though on Earth it may have increased or decreased by a factor up to about 1.5 over geologic time (Harrison, 1999; Korenaga, 2008; Pope et al., 2012; Laneuville et al., 2018). We do not consider the potential sequestration of carbon species in ice caps or loss to space. Atmospheric speciation, photochemistry, and effects on surface temperature and runoff are not considered in this paper. This model does not differentiate between degassing of primordial vs. subducted carbon.

This model does not explicitly track outgassing chemistry as a function of source 632 depth, degassing transport, and outgassing pressure. While these aspects are thought 633 to impact the redox state of outgassed material, the extent to which they do is debated 634 (Burgisser & Scaillet, 2007; Dasgupta & Hirschmann, 2010). The  $fO_2$  of magmas and 635 associated abundances of gas species in equilibrium with the magma composition could 636 change up to 1.5-2 orders of magnitude upon ascent, due to degassing and pressure ef-637 fects on volatile exsolution (Burgisser & Scaillet, 2007). However, this variation could 638 go in either direction (toward a more oxidizing or reducing magma), depending on the 639 starting conditions, unless the volatiles are primarily H<sub>2</sub> and H<sub>2</sub>O, with little C and S 640 species. Moreover, the findings of Burgisser and Scaillet (2007) applied to Fe-poor (rhy-641 olitic) magmas, but more mafic magmas may provide added buffering capability, decreas-642 ing changes in magma redox during ascent. 643

#### 644 **3 Results**

In this section, we describe the sensitivity to planet size and composition of carbon fluxes arising from outgassing and weathering. The sensitivity of the carbon cycle and resulting atmospheric composition on these properties is investigated in the companion paper.

649

# 3.1 Geological controls on outgassing

The outgassing fluxes through time of rocky silicate planets with a surface water ocean and a surface temperature similar to Earth's are shown in Fig. 5a as a function of planet mass and upper mantle composition. Planet mass is varied between 0.5 and <sup>653</sup> 2  $M_{\oplus}$ , with the same planet-building materials and core mass fraction as in the mod-<sup>654</sup> ern Earth validation simulation shown in Fig. 2 and a dry olivine rheology.

Four upper mantle compositions are investigated: mid-ocean ridge basalt (Allan et al., 1989), chondritic (Sanloup et al., 1999), rhyolitic (Pamukcu et al., 2015), and felsic (Dolinschi, 2019) (Table 4). These oxide compositions were speciated into mineral compositions assuming an oxidation state corresponding to the FMQ buffer. Varying  $fO_2$ to several log units below this buffer did not significantly affect the propensity of these compositions to melt (or, therefore, the outgassing fluxes).

Simulations span the time interval between 0 and 5 Gyr after formation, but outgassing fluxes are computed only after 0.6 Gyr to allow for the initial geodynamic evolution to lead to a more steadily evolving mantle temperature and heat flux. The time step in these simulations is 10 Myr, which achieves numerical convergence since climate feedbacks are not tracked for this outgassing flux sensitivity analysis.

Outgassing fluxes span  $2 \times 10^4$  to  $6 \times 10^6$  mol s<sup>-1</sup>. They tend to be greater at 666 earlier times and for more felsic compositions. They decrease with increasing planet mass 667 if the mass is greater than  $1M_{\oplus}$ , but the mass at which outgassing fluxes peak is lower 668 for the two more felsic compositions:  $0.5 - 0.75 M_{\oplus}$  compared to  $1 M_{\oplus}$  for the basaltic 669 (Earth-like) upper mantle composition and  $1.5 M_\oplus$  for the ultramafic (chondritic) com-670 position. The outgassing fluxes for the  $1M_{\oplus}$  planets are usually within a few percent of 671 the peak values as a function of planet mass, irrespective of composition; whereas whether 672 the outgassing flux for 0.5-, 0.75-, or  $1.5-M_{\oplus}$  planets is close to this maximum depends 673 sensitively on composition. For example, for basaltic Earth-like or chondritic composi-674 tions, outgassing fluxes for 0.5- $M_{\oplus}$  planets are lower than those of 1- $M_{\oplus}$  planets by tens 675 of percent at any given time after formation. 676

The general decrease of outgassing fluxes with time is consistent with the global thermal evolution which tends toward lower mantle temperatures and heat fluxes as radiogenic heating decreases over time. The decrease spans up to two orders of magnitude between the first billion years after formation and the present day, but with considerable variability. It is less pronounced for higher-mass planets within the range investigated and for more felsic compositions. For 2  $M_{\oplus}$  planets, the flux first increases to peak at 1 to 2 Gyr after formation before decreasing to 5-Gyr values that are close to fluxes

Table 4. Simulated mantle compositions in mass% oxides: mid-ocean ridge basalt (MORB; Allan et al. (1989) as cited in Ghiorso and Sack (1995)), felsic (Dolinschi, 2019), rhyolitic (Pamukcu et al., 2015), and chondritic (Sanloup et al. (1999) as cited in Taylor (2013)). About 1 mass% or less of SiO<sub>2</sub> was either added or removed from the reported compositions in order for the total to reach 100%. These compositions are free of sub-percent mass fractions reported for Mn, Cr, and K oxides, as P. Antoshechkina and Asimow (2018) advised that including these as major oxides rather than trace elements in MELTS calculations can lead to unpredictable results. The rhyolitic composition's non-negligible K<sub>2</sub>O mass fraction (Pamukcu et al., 2015) was added as additional Na<sub>2</sub>O. Reported mass fractions of Al<sub>2</sub>O and Fe<sub>2</sub>O<sub>3</sub> were converted to mass fractions of Al<sub>2</sub>O<sub>3</sub> and FeO, respectively, because Al<sub>2</sub>O is not part of the MELTS chemical model and MELTS partitions Fe between Fe<sub>2</sub>O<sub>3</sub> and FeO during a calculation based on the chosen redox buffer (here, FMQ).

Oxide	MORB	Chondritic	Rhyolitic	Felsic
$SiO_2$	48.4	48.6	75.6	52.8
MgO	9.1	27.0	0.2	29.2
FeO	8.5	17.5	1.0	12.2
$Al_2O_3$	17.6	3.6	13.2	2.8
CaO	12.5	2.0	0.8	2.4
$\mathrm{TiO}_2$	1.0	0.1	0.2	-
Na <sub>2</sub> O	2.7	1.2	9.1	0.6
$P_2O_5$	0.1	-	-	-
$\rm H_2O$	0.2	-	-	-



Figure 5. a. Effect of planet mass and upper mantle composition on outgassing flux through time. Planet mass is varied between 0.5 and 2  $M_{\oplus}$  and shown as lighter to darker shade, respectively. Four upper mantle compositions are investigated: mid-ocean ridge basalt ("Earth-like", blue curves), chondritic (green curves), rhyolitic (yellow curves), and felsic (gray curves). The Earth symbol indicates the computed outgassing rate for modeled modern Earth conditions at which the model was validated (Fig. 2); the other three blue symbols indicate literature estimates of this value (Zhang & Zindler, 1993; Donnadieu et al., 2006; Burton et al., 2013). Gaps in curves or (especially for the rhyolitic composition) spuriously low values arise from lack of convergence of *MELTS* calculations at the full set of relevant shallow depths for those particular combinations of composition and mass- and age-dependent geotherm. **b.** Effect of planet mass (pressure-temperature profile) on the extent of melting of an upper mantle of basaltic (Earthlike) composition. **c.** Mid-mantle temperature through geologic time for planets of different masses, all with a basaltic upper mantle.

- computed at 0.6 Gyr. For felsic upper mantle compositions, the decrease in outgassing
  flux does not exceed 1 order of magnitude between 0.6 and 5 Gyr.
- At early times, mass has the largest effect on outgassing flux, but past 2 Gyr after formation, the outgassing fluxes become instead more strongly influenced by upper mantle composition. At 0.6 Gyr, the outgassing flux for  $\leq 1M_{\oplus}$  planets is greater than 10<sup>6</sup> mol s<sup>-1</sup> for all four upper mantle compositions simulated, whereas the outgassing flux for  $2M_{\oplus}$  planets is lower than 10<sup>6</sup> mol s<sup>-1</sup>. Instead, at 4 Gyr, the outgassing flux ranges corresponding to planet masses in the range  $0.5-2M_{\oplus}$  for each of the four compositions simulated no longer overlap: the flux ranges are about 8 - 12, 3 - 6, 0.8 -

 $_{693}$  2, and  $0.1 - 0.7 \times 10^5$  mol s<sup>-1</sup> respectively for the felsic, rhyolitic, basaltic, and chon-

<sup>694</sup> dritic mantle compositions.

The mass dependence of outgassing flux arises primarily from the influence of planet 695 mass on the geotherm (pressure-temperature profile) and its position relative to the solidus 696 for a given composition. For an Earth-like, basaltic upper mantle composition, planets 697 more massive than Earth have higher internal pressures and a shallower lithosphere-asthenosphere 698 boundary (Fig. 5b). This tends to limit the extent of mantle melting relative to a  $1-M_{\oplus}$ 699 planet. Planets less massive tend to be cooler because of their lower ratio of volume (ac-700 cretional and radiogenic heat production) to surface area (heat loss). This also limits the 701 extent of mantle melting (Fig. 5b). Thus, melting (and, proportionally, outgassing) is 702 generally maximized for  $1-M_{\oplus}$  planets. 703

The above analysis applies past 2 Gyr. At earlier times, outgassing is much more 704 suppressed on planets more massive than Earth owing to, counter-intuitively, relatively 705 low internal temperatures and heat fluxes (Fig. 5c). This is because convective heat trans-706 port is so efficient immediately after formation (due to the need to evacuate the high heat 707 of accretion) that mantle temperatures become uniform enough for the vigor of convec-708 tion and associated heat flux to drop below those of lower-mass planets. This results in 709 planets whose geotherm seldom crosses the solidus, and therefore in little melting and, 710 proportionally, outgassing. Later on, the heat flux increases back due to radiogenic heat-711 ing. 712

The compositional dependence of outgassing flux arises from the position of the upper mantle material's solidus curve relative to the geotherm of a planet of given mass and age. The more felsic compositions of Dolinschi (2019) and Pamukcu et al. (2015) have lower melting temperatures; therefore, the degree of melting for a given geotherm crossing their solidus is higher, leading to higher outgassing.

718

#### 3.2 Geological controls on continental weathering

The effect of surface composition on continental weathering fluxes of rocky silicate planets is shown in Fig. 6a. The diverse surface compositions simulated are reflective of those of inner solar system planets (Table 5). These simulations were carried out assuming a 20% CO<sub>2</sub>, 80% N<sub>2</sub>, 1-bar atmosphere; a 288 K surface temperature; and modern-Earth rainfall rate of 0.7 m yr<sup>-1</sup> (water:rock ratio  $\approx$  5000). As in Fig. 3c, weathering

-33-



Figure 6. a. Continental weathering flux as a function of planet surface composition and residence time of rainwater on land. This flux is expressed as the equivalent cation capacity able to combine with bicarbonate  $(\text{HCO}_3^-)$  to form mineral precipitates. For each curve, a thumbnail of the planetary surface and labels "1" and "2" indicate the surface composition based on Table 5. The jaggedness in the curves for the Mars- and Ceres-like surface compositions is due to *PHREEQC* having difficulties converging on equilibria involving the precipitation of insoluble oxidized iron minerals. **b.** Dependence of weathering flux as a function of residence time on temperature for surface compositions approximating those of Modern and Archean Earth (Table 5).

fluxes are expressed as the equivalent cation capacity able to combine with bicarbonate ( $HCO_3^-$ ) to form mineral precipitates.

As shown in Fig. 6a, surface composition has a moderate effect on continental weathering fluxes, which are all on the order of tens of Tmol yr<sup>-1</sup>. Weathering fluxes are higher if the surface is made of minerals able to rapidly dissolve, such as sulfates (Venus-like surface) and carbonates (Ceres-like surface). The resulting enrichment of river water in cations likely acts as a negative feedback onto the dissolution of other minerals as  $\Omega$  in equation (10) increases toward 1 (i.e.,  $Q \to K$ ), preventing weathering fluxes from varying by more than about 1 order of magnitude.

Continental weathering fluxes increase with increased continental residence time, because the longer water-rock interaction leads to more mineral dissolution not fully compensated by precipitation of supersaturated minerals. For a given rainfall rate, weathering should thus be more effective on less patchy land.

Continental weathering fluxes also generally increase with temperature (Fig. 6b), 737 as expected from the Arrhenius (exponential) dependence of reaction kinetic rates on 738 the inverse of temperature. Despite this general trend, the relationship is more complex 739 and depends, e.g., on residence time of water on land. For the modern Earth case, if the 740 residence time exceeds 50 years (i.e., for continental masses more consolidated than to-741 day), the weathering flux is lower at higher temperatures because mineral formation re-742 actions that consume dissolve cations proceed faster owing both to faster kinetics and 743 to the faster accumulation of river cations (from prior mineral dissolution) to reach min-744 eral saturation. Thus, especially on less patchy land, weathering fluxes may instead be 745 controlled mainly by (proportional to) land area and rainfall. 746

Expressing weathering flux as the equivalent cation capacity able to combine with 747 bicarbonate  $(HCO_3^-)$  to form mineral precipitates requires making the implicit assump-748 tion that oxidized carbon is present as bicarbonate, i.e., that the ocean pH is roughly 749 between 6 (below which dissolved CO<sub>2</sub>, i.e., carbonic acid, dominates) and 10 (above which 750 doubly charged carbonate dominates). At pH > 10, the oxidized carbon cation trapping 751 capacity would be halved to conserve charge balance; at pH < 6, the trapping capacity 752 of cations would be zero since carbonic acid is neutral. In carbon cycling evolution sim-753 ulations that combine outgassing, weathering, and ocean-atmosphere equilibrium, the 754

-35-
ocean pH tends to be between 6 and 9 (see Fig. 7 in Section 4 and companion paper),

<sup>756</sup> making bicarbonate trapping capacity a reasonable means of evaluating weathering flux.

## 757 4 Discussion

Carbon is cycled between the atmosphere, surface, and interior of a planet by outgassing, weathering, and subduction. The balance between fluxes associated with these processes determines the amount of carbon accumulated in the atmosphere-surface and interior reservoirs. The above results enable us to understand how a planet's size and composition affects the intrinsic strength of these fluxes, i.e., of carbon cycle feedbacks. To do so, we have evaluated the magnitude of outgassing and continental weathering fluxes; for the latter, in the specific conditions of Earth's modern-day atmosphere.

765

766

# 4.1 Effect of planet size and composition on continental weathering and outgassing

Our simulations suggest that the efficiency of outgassing and subduction decreases 767 with mass for planets between 1 and 2 Earth masses. This contradicts the findings of 768 Kite et al. (2009) but is in line with more recent work accounting for the effects of high 769 pressure increasing mantle viscosity and thus decreasing its propensity to convect (Stamenković 770 et al., 2012). Here, planet mass is found to influence outgassing in other ways: at lower 771 masses, lower mantle temperatures decrease the degree of melting. At higher masses, higher 772 pressures decrease the degree of melting. We have assumed, as did Kite et al. (2009), that 773 all melt produced reaches the surface, but planet mass may also have an impact on the 774 degree to which intrusive volcanism does not lead to outgassing, proportionally decreas-775 ing outgassing compared to the values shown in Fig. 5a. 776

We have neglected the dependence on bulk composition of the thermophysical prop-777 erties and radionuclide abundances that affect a planet's geodynamic evolution (Tables 778 1 and 2). More matic compositions may lead to more vigorous convection and faster-cooling 779 planets that may outgas comparatively more early on and less at later ages than the flux 780 changes shown in Fig. 5a. The linkage between bulk compositions and radionuclide abun-781 dances is not straightforward. The internal distribution of radionuclides is unclear, be-782 cause elements Th and U tend to be incorporated as impurities in minerals (Oelkers et 783 al., 2009; Neveu et al., 2017) and because K can be exchanged between planet-building 784

-36-

materials such as silicates and volatile ices through aqueous alteration (Kirk & Steven-785 son, 1987; Engel et al., 1994; Castillo-Rogez & Lunine, 2010; Neveu et al., 2017). More-786 over, the starting abundances of radionuclides are difficult to predict from first princi-787 ples. The dominant heat source is <sup>40</sup>K, but Earth is rather depleted (relative to chon-788 drites) in moderately volatile elements, including K, by unknown mechanisms (Desch et 789 al., 2020; Unterborn et al., 2022). The solar system is probably not unusual in its abun-790 dances of short-lived radionuclides such as <sup>26</sup>Al, but these would be even more variable 791 (Desch et al., 2022). Thus, radiogenic heating may best be left as a freely varied param-792 eter, with the simulations shown here providing a solar system-centric baseline. 793

More mafic compositions tend to have a higher solidus temperature for a given pressure, and therefore are less prone to generating melt (i.e., outgassing). Although this was not tracked in our simulations, more mafic (reducing) compositions are generally expected to result in more reducing outgassing. The detailed carbon gas speciation into relative proportions of  $CO_2$  and  $CH_4$  depends on effects such as redox changes during ascent (Burgisser & Scaillet, 2007) and planet surface pressure (gravity and atmospheric density) (Gaillard & Scaillet, 2014), which are not considered here (Section 2.5).

Surface gravity is not expected to affect the continental weathering flux via the runoff rate to which this flux is proportional. Runoff rate is in turn proportional to river flow velocity, which depends on surface gravity as v = 2gh. Since declivity (a planet's relief) h is inversely proportional to surface gravity, the effects of gravity cancel out.

A planet's surface composition does affect the continental weathering flux to an ex-805 tent comparable to the effect of residence time of rainwater on land (Fig. 6a). However, 806 the effect of composition is much more pronounced on outgassing, for which it induces 807 order-of-magnitude differences (Fig. 5a). Time-evolution simulations in which the at-808 mospheric composition varies as a function of outgassing and (continental+seafloor) weath-809 ering show that in practice, the combined weathering flux varies to match the outgassing 810 forcing so as to keep the sum of atmospheric and ocean carbon roughly constant within 811 a factor of a few. An example such simulation, in which it is assumed that seafloor weath-812 ering proceeds as described in Section 2.3, is shown in Fig. 7. A broader set of such sim-813 ulations for different planet sizes and compositions is presented in the companion pa-814 per. 815

-37-



Figure 7. Time evolution of a. carbon reservoirs, b. carbon fluxes, c. surface temperature and pressure, d. atmospheric composition, e. pH, and f. ocean composition on a  $1-M_{\oplus}$  planet with an upper mantle composition comparable to that of mid-ocean ridge basalts (Table 4) and a surface composition comparable to that of Archean Earth (Table 5). The combined (continental + seafloor) weathering flux varies to match the outgassing forcing, maintaining roughly constant amounts of carbon in the {atmosphere+ocean} reservoir. Changes in the partitioning of carbon between the atmosphere and ocean arise due to secular changes in ocean chemistry (in this case, increasing pH). The ocean composition is initiated to the present-day composition, and the atmospheric composition is initiated to 0.5 bar total pressure of 80% N<sub>2</sub>-20% CO<sub>2</sub>.

Overall, planet mass and composition are expected to affect the carbon cycle as 816 follows, assuming the planet receives insolation similar to Earth's. First, the magnitude 817 of injection rates (fluxes) of carbon into the {atmosphere+ocean} reservoir varies by or-818 ders of magnitude (Fig. 5a) and is expected to generally decrease with geologic time. The 819 added carbon, dissolved in rainwater and in the ocean, reacts with land and seafloor rock. 820 The former yields an influx of carbon and cations into the ocean at a rate that is not very 821 sensitive to surface composition or residence time (continental patchiness) (Fig. 6) and 822 insensitive to planet mass (surface gravity), but which is proportional to rainfall rate (it-823 self weakly increasing with temperature via the atmospheric  $H_2O$  content; Section 2.4) 824 and land area. Continental patchiness affects evaporation, i.e., water:rock ratio, but sim-825 ulations with water:rock ratios varied by a factor of a few do not yield major differences 826 in weathering fluxes, especially since the effects of water:rock ratio and residence time 827 offset each other (reactions with more concentrated reactants proceed faster). The finite 828 solubility of carbon minerals drives precipitation and incorporation of carbon into the 829 seafloor at rates that should match the rate of outgassing (Fig. 7). Variations in the con-830 tinental weathering input of cations should drive changes in ocean composition on ge-831 ological timescales that may affect the partitioning of carbon between the ocean and at-832 mosphere due to the dependence of carbon speciation on, e.g., pH. Detailed investiga-833 tion of these effects is the topic of the companion paper. 834

835

### 4.2 The Earth through time

While Earth's mean surface temperature has likely remained relatively constant at a few degrees above freezing throughout much of its evolution due to the carbonatesilicate cycle (Sleep & Zahnle, 2001), its land composition has become more felsic, its land less patchy, and its land coverage has increased (Korenaga, 2018). Its atmosphere has become less carbon-rich, with  $pCO_2$  changing from about 0.1–1 bar to a few  $10^{-4}$ bar (Catling & Zahnle, 2020).

This change in surface composition has acted to decrease continental weathering fluxes (Fig. 8) due to the lesser disequilibrium with Earth's atmosphere. Had Earth's composition remained the same, the continental weathering flux today would be closer to 100 Tmol yr<sup>-1</sup> of carbon than the estimated 20–25 Tmol yr<sup>-1</sup> (see Fig. 3c). However, the higher patchiness and lower areal fraction of land on Archean Earth implies a lower flux: about  $55 \times \frac{L}{0.29}$  Tmol/year for a residence time of 0.1 rather than 10 years.



Figure 8. Effect of land coverage and patchiness on continental weathering flux for a  $1-M_{\oplus}$ planet. Here, we have assumed a primitive mantle composition for the Earth's crust, which may be adequate for the Archean, and an Archean-like atmospheric composition of 0.5 bar N<sub>2</sub> and 0.5 bar CO<sub>2</sub>. The inset near 10 years residence time shows the same fluxes for modern Earth, as reproduced from Fig. 3.

This matches the modern-day weathering flux for a land fraction of about 15%, implying that the continental weathering flux need not have changed significantly over time.

The outgassing flux was likely an order of magnitude higher than today (Fig. 5a; 850 see also Sleep and Zahnle (2001) and Krissansen-Totton, Arney, and Catling (2018)), im-851 plying a greater forcing that decreased with time. This flux was balanced by a greater 852 seafloor weathering flux arising from a combination of higher convective velocity (Laneuville 853 et al., 2018) and 3-5 times greater ridge length (Kadko et al., 1995) in equation (12) (lower 854 water:rock ratio), even though, given the above, the accumulation rate of reactants from 855 river delivery to the ocean was roughly the same as today. The gradual increase with 856 time in the water:rock ratio of seafloor reactions, decrease in outgassing forcing, and neg-857 ative feedback of the cycle likely all acted to maintain carbon levels in the atmosphere 858 until carbon became significantly sequestered in biomass. 859

# 5 Conclusions

We have introduced a new, open-access model of geological carbon fluxes, whose implementation leverages existing, widely used geochemical codes of solid-melt equilib-

-40-

ria for silicate rocks (*MELTS*) and of equilibria and kinetics of water-rock interactions (*PHREEQC*). Coupled with a simple numerical computation of global thermal evolution, this model enables investigation of the effects of planet size (mass) and composition on carbon cycling through geologic time. Its applicable size range (0.5 to 2 Earth masses) is limited by the fidelity of the geodynamic model; the applicable range of composition is limited by those that can be handled by *MELTS* and *PHREEQC*.

We find that these planetary properties can influence outgassing fluxes of carbon 869 by as much as two orders of magnitude, with 0.5–1- $M_{\oplus}$  planets with felsic upper man-870 tles outgassing the most. In contrast, continental weathering fluxes are independent of 871 planet size and relatively insensitive to surface composition in terms of disequilibrium 872 with an Earth-like atmosphere; the variation in fluxes is at most a factor of a few. Con-873 tinental patchiness (residence time of rainwater) has an effect of similar magnitude, with 874 greater residence times generally leading to higher fluxes by allowing kinetically limited 875 reactions to proceed further. Surface composition and land patchiness thus likely have 876 lesser effects on continental weathering than the fraction of a planet's surface covered 877 by land. These continental weathering fluxes likely affect the rate of change of oceanic 878 chemical composition, which governs the partitioning of carbon between ocean and at-879 mosphere. The resulting evolutions of atmosphere and ocean compositions on Earth-like 880 planets of various sizes and solid compositions are the focus of a companion paper. 881

- 882 Open Research Section
- The data supporting this paper's conclusions can be obtained by running the ExoCcycleGeo code (version 22.4) with the crust and upper mantle composition templates freely available from https://github.com/MarcNeveu/ExoCcycleGeo. The software dependencies can be downloaded at the following webpages:

887

888

*PHREEQC*: https://wwwbrr.cr.usgs.gov/projects/GWC\_coupled/phreeqc;

- alphaMELTS: https://magmasource.caltech.edu/alphamelts; MELTS itself is available available
- able at https://melts.ofm-research.org.
- The data underlying Fig. 3 to 8 are archived in a Zenodo repository at https://zenodo.org/deposit/7640681 (size: 377.5 MBytes). The subset of data underlying Fig. 3, 4, 6, and 8 were obtained by running *PHREEQC* alone. Those underlying Fig. 2 were obtained by hard-coding a value for the mid-mantle temperature in *ExoCcycleGeo* and running a single time step.

-41-

### 894 Acknowledgments

- This work was funded by NASA Exoplanet Research Program award NNX17AB95G. We
- thank the reviewers of the proposal that led to this award for providing valuable feed-

<sup>897</sup> back and guidance that helped improve this research.

### 898 References

- Abbot, D. S., Cowan, N. B., & Ciesla, F. J. (2012). Indication of insensitivity of
   planetary weathering behavior and habitable zone to surface land fraction. *The Astrophysical Journal*, 756(2), 178. doi: 10.1088/0004-637X/756/2/178
- Akaogi, M., & Ito, E. (1993). Heat capacity of MgSiO<sub>3</sub> perovskite. *Geophysical Research Letters*, 20(2), 105–108. doi: 10.1029/92GL02655
- Akeson, R., Chen, X., Ciardi, D., Crane, M., Good, J., Harbut, M., ... Zhang, A.
   (2013). The NASA Exoplanet Archive: data and tools for exoplanet research.
   *Publications of the Astronomical Society of the Pacific*, 125(930), 989. doi:
   10.1086/672273
- Allan, J. F., Batiza, R., Perfit, M. R., Fornari, D. J., & Sack, R. O. (1989). Petrology of lavas from the lamont seamount chain and adjacent East Pacific Rise,
  10 N. Journal of Petrology, 30(5), 1245–1298. doi: 10.1093/petrology/ 30.5.1245
- Allen, M. R., & Ingram, W. J. (2002). Constraints on future changes in climate and
  the hydrologic cycle. *Nature*, 419(6903), 228–232.
- Anderson, A. (1995). CO<sub>2</sub> and the eruptibility of picrite and komatiite. *Lithos*, 34(1-3), 19–25. doi: 10.1016/0024-4937(95)90005-5
- Anderson, G. M. (2005). Thermodynamics of Natural Systems. Cambridge Univer sity Press.
- Anderson, W. W., & Ahrens, T. J. (1994). An equation of state for liquid iron
   and implications for the Earth's core. Journal of Geophysical Research: Solid
   Earth, 99(B3), 4273–4284. doi: 10.1029/93JB03158
- Antoshechkina, P., & Asimow, P. (2018). Alphamelts software manual a text
   driven interface for MELTS, pMELTS & pHMELTS (Tech. Rep.). California
   Institute of Technology.
- Antoshechkina, P. M., & Asimow, P. D. (2010). Adiabat\_1ph 3.0 and the MAGMA website: educational and research tools for studying the petrology and geo-

926	chemistry of plate margins. In AGU Fall Meeting Abstracts (Vol. 2010, pp.
927	ED41B-0644).
928	Antoshechkina, P. M., Asimow, P. D., Hauri, E. H., & Luffi, P. I. (2010). Effect of
929	water on mantle melting and magma differentiation, as modeled using Adia-
930	bat_1ph 3.0. In AGU Fall Meeting Abstracts (Vol. 2010, pp. V53C–2264).
931	Arney, G. N., Domagal-Goldman, S. D., Meadows, V. S., Wolf, E. T., Schwieterman,
932	E., Charnay, B., Trainer, M. G. (2016). The pale orange dot: the spectrum
933	and habitability of hazy Archean Earth. Astrobiology, $16(11)$ , 873–899. doi:
934	10.1089/ast.2015.1422
935	Arney, G. N., Meadows, V. S., Domagal-Goldman, S. D., Deming, D., Robinson,
936	T. D., Tovar, G., Schwieterman, E. (2017). Pale orange dots: the impact of
937	organic haze on the habitability and detectability of Earthlike exoplanets. $\ The$
938	Astrophysical Journal, 836(1), 49. doi: 10.3847/1538-4357/836/1/49
939	Asimow, P. D., & Ghiorso, M. S. (1998). Algorithmic modifications extending
940	MELTS to calculate subsolidus phase relations. American Mineralogist, $83(9-$
941	10), 1127–1132. doi: 10.2138/am-1998-9-1022
942	Auzende, AL., Pellenq, RM., Devouard, B., Baronnet, A., & Grauby, O. (2006).
943	Atomistic calculations of structural and elastic properties of serpentine miner-
944	als: the case of lizardite. Physics and Chemistry of Minerals, 33(4), 266–275.
945	doi: $10.1007/s00269-006-0078-x$
946	Barnes, R., Luger, R., Deitrick, R., Driscoll, P., Quinn, T. R., Fleming, D. P.,
947	Armstrong, J. (2020). VPLanet: the virtual planet simulator. Publica-
948	tions of the Astronomical Society of the Pacific, $132(1008)$ , $024502$ . doi:
949	10.1088/1538-3873/ab3ce8
950	Barnes, R., Meadows, V. S., & Evans, N. (2015). Comparative habitability of tran-
951	siting exoplanets. The Astrophysical Journal, $814(2)$ , 91. doi: 10.1088/0004
952	-637X/814/2/91
953	Barnes, R., Mullins, K., Goldblatt, C., Meadows, V. S., Kasting, J. F., & Heller,
954	R. $(2013)$ . Tidal Venuses: triggering a climate catastrophe via tidal heating.
955	Astrobiology, $13(3)$ , 225–250. doi: 10.1089/ast.2012.0851
956	Begemann, F., & Libby, W. F. (1957). Continental water balance, ground water
957	inventory and storage times, surface ocean mixing rates and world-wide wa-
958	ter circulation patterns from cosmic-ray and bomb tritium. Geochimica et

959	Cosmochimica Acta, $12(4)$ , 277–296. doi: $10.1016/0016-7037(57)90040-6$
960	Berner, R. A., Lasaga, A. C., & Garrels, R. M. (1983). The carbonate-silicate
961	geochemical cycle and its effect on atmospheric carbon dioxide over the
962	past 100 million years. American Journal of Science, 283, 641–683. doi:
963	10.2475/ajs.283.7.641
964	Bhatia, G. K., & Sahijpal, S. (2016). The early thermal evolution of Mars. Meteorit-
965	ics and Planetary Science, $51(1)$ , 138-154. doi: 10.1111/maps.12573
966	Bhatia, G. K., & Sahijpal, S. $(2017)$ . Did $^{26}$ Al and impact-induced heating differ-
967	entiate Mercury? Meteoritics and Planetary Science, 52(2), 295-319. doi: 10
968	.1111/maps.12789
969	Blackledge, B., Green, J., Barnes, R., & Way, M. J. (2020). Tides on other Earths:
970	Implications for exoplanet and palaeo-tidal simulations. Geophysical Research
971	$Letters,\ 47(12),\ e2019GL085746.\ doi:\ https://doi.org/10.1029/2019GL085746$
972	Blundy, J., Cashman, K. V., Rust, A., & Witham, F. $(2010)$ . A case for CO <sub>2</sub> -rich
973	arc magmas. Earth and Planetary Science Letters, 290(3-4), 289–301. doi: 10
974	.1016/j.epsl.2009.12.013
975	Bond, J. C., O'Brien, D. P., & Lauretta, D. S. (2010). The compositional diversity
976	of extrasolar terrestrial planets. I. In situ simulations. The Astrophysical Jour-
977	nal, 715(2), 1050. doi: 10.1088/0004-637X/715/2/1050
978	Broecker, W. S., & Peng, T. H. (1982). Tracers in the Sea. Lamont-Doherty Geolog-
979	ical Observatory, Palisades, NY.
980	Brown, M., Johnson, T., & Gardiner, N. J. (2020). Plate tectonics and the archean
981	earth. Annual Review of Earth and Planetary Sciences, 48, 291–320. doi: 10
982	.1146/annurev-earth-081619-052705
983	Burgisser, A., & Scaillet, B. (2007). Redox evolution of a degassing magma rising to
984	the surface. Nature, $445(7124)$ , 194–197. doi: 10.1038/nature05509
985	Burton, M. R., Sawyer, G. M., & Granieri, D. (2013). Deep carbon emissions from
986	volcanoes. Reviews in Mineralogy and Geochemistry, $75(1)$ , $323-354$ . doi: 10
987	$.2138/\mathrm{rmg}.2013.75.11$
988	Byerlee, J. (1978). Friction of rocks. In Rock Friction and Earthquake Prediction
989	(pp. 615–626). Springer.
990	Caldeira, K., & Kasting, J. F. (1992). The life span of the biosphere revisited. Na-
991	ture, $360(6406)$ , 721–723. doi: 10.1038/360721a0

992	Canup, R. M. (2004). Simulations of a late lunar-forming impact. <i>Icarus</i> , 168(2),
993	433–456. doi: 10.1016/j.icarus.2003.09.028
994	Canup, R. M., Kratter, K. M., & Neveu, M. (2021). On the origin of the Pluto sys-
995	tem. In The Pluto System after New Horizons (p. 475-506). University of Ari-
996	zona Press.
997	Castillo-Rogez, J. C., & Lunine, J. I. (2010). Evolution of Titan's rocky core con-
998	strained by Cassini observations. Geophysical Research Letters, $37(20)$ . doi: 10
999	.1029/2010GL044398
1000	Catling, D. C., Kiang, N. Y., Robinson, T. D., Rushby, A. J., Del Genio, A., et al.
1001	(2018). Exoplanet biosignatures: a framework for their assessment. Astrobiol-
1002	ogy. doi: 10.1089/ast.2017.1737
1003	Catling, D. C., & Zahnle, K. J. (2020). The Archean atmosphere. Science Advances,
1004	$\boldsymbol{6}(9),$ eaax1420. doi: 10.1126/sciadv.aax1420
1005	Chahine, M. T. (1992). The hydrological cycle and its influence on climate. Nature,
1006	359(6394), 373-380.doi: 10.1038/359373a0
1007	Choukroun, M., & Grasset, O. (2010). Thermodynamic data and modeling of
1008	the water and ammonia-water phase diagrams up to $2.2$ GPa for plane-
1009	tary geophysics. The Journal of Chemical Physics, 133(14), 144502. doi:
1010	10.1063/1.3487520
1011	Chung, D. (1971). Elasticity and equations of state of olivines in the $Mg_2SiO_4$ -
1012	$Fe_2SiO_4$ system. Geophysical Journal International, 25(5), 511–538. doi: 10
1013	.1111/j.1365-246X.1971.tb02201.x
1014	Cioni, R. (2000). Volatile content and degassing processes in the AD 79 magma
1015	chamber at Vesuvius (Italy). Contributions to Mineralogy and Petrology,
1016	140(1), 40-54. doi: 10.1007/s004100000167
1017	Cogné, JP., & Humler, E. (2006). Trends and rhythms in global seafloor gen-
1018	eration rate. Geochemistry, Geophysics, Geosystems, $7(3)$ . doi: 10.1029/
1019	2005GC001148
1020	Cohen, S. D., Hindmarsh, A. C., & Dubois, P. F. (1996). CVODE, a stiff/nonstiff
1021	ODE solver in C. Computers in Physics, $10(2)$ , 138–143. doi: 10.1063/
1022	1.4822377
1023	Colbourn, G., Ridgwell, A., & Lenton, T. (2015). The time scale of the silicate
1024	weathering negative feedback on atmospheric CO <sub>2</sub> . Global Biogeochemical Cy-

1025	cles, 29(5), 583–596. doi: 10.1002/2014GB005054
1026	Cole, J. J., & Caraco, N. F. (2001). Carbon in catchments: connecting terrestrial
1027	carbon losses with aquatic metabolism. Marine and Freshwater Research,
1028	52(1), 101-110. doi: $10.1071/MF00084$
1029	Colose, C. M., Haqq-Misra, J., Wolf, E. T., Del Genio, A. D., Barnes, R., Way,
1030	M. J., & Ruedy, R. (2021). Effects of spin–orbit resonances and tidal heating
1031	on the inner edge of the habitable zone. The Astrophysical Journal, $921(1)$ ,
1032	25. doi: 10.3847/1538-4357/ac135c
1033	Dasgupta, R., & Hirschmann, M. M. (2010). The deep carbon cycle and melting in
1034	Earth's interior. Earth and Planetary Science Letters, 298(1-2), 1–13. doi: 10
1035	.1016/j.epsl.2010.06.039
1036	De Sanctis, M. C., Ammannito, E., Raponi, A., Marchi, S., McCord, T. B., Mc-
1037	Sween, H., Russell, C. T. (2015). Ammoniated phyllosilicates with a likely
1038	outer solar system origin on (1) Ceres. Nature, 528(7581), 241–244. doi:
1039	10.1038/nature16172
1040	Desch, S. J., Abbot, D., Krijt, S., Unterborn, C., Morard, G., & Hartnett, H. E.
1041	(2020). The volatile content of rocky planets. In <i>Planetary Diversity: Rocky</i>
1042	planet processes and their observational signatures (pp. $6.1$ – $6.40$ ). IOP.
1043	Desch, S. J., Young, E. D., Dunham, E. T., Fujimoto, Y., & Dunlap, D. R. (2022).
1044	Short-Lived Radionuclides in Meteorites and the Sun's Birth Environment.
1045	$arXiv\ e\text{-}prints.$ doi: 10.48550/arXiv.2203.11169
1046	Deschamps, F., & Sotin, C. (2000). Inversion of two-dimensional numerical
1047	convection experiments for a fluid with a strongly temperature-dependent
1048	viscosity. Geophysical Journal International, 143(1), 204–218. doi:
1049	10.1046/j.1365-246x.2000.00228.x
1050	Des Marais, D. J., Jakosky, B. M., & Hynek, B. M. (2007). Astrobiological impli-
1051	cations of Mars' surface composition and properties. In The Martian Surface-
1052	Composition, Mineralogy, and Physical Properties (pp. 599–624). Cambridge
1053	University Press.
1054	Dolinschi, J. D. (2019). Mineralogy of the silicon-rich mantle: Implications for Mars
1055	and exoplanets. <i>PhD dissertation</i> . Retrieved from https://keep.lib.asu
1056	.edu/_flysystem/fedora/c7/218100/Dolinschi_asu_0010N_19331.pdf
1057	Domagal-Goldman, S. D., Segura, A., Claire, M. W., Robinson, T. D., & Meadows,

-46-

1058	V. S. (2014). Abiotic ozone and oxygen in atmospheres similar to prebiotic
1059	Earth. The Astrophysical Journal, $792(2)$ , 90. doi: $10.1088/0004-637X/792/2/2/2$
1060	90
1061	Donnadieu, Y., Goddéris, Y., Pierrehumbert, R., Dromart, G., Fluteau, F., & Jacob,
1062	R. (2006). A GEOCLIM simulation of climatic and biogeochemical conse-
1063	quences of Pangea breakup. $Geochemistry, Geophysics, Geosystems, 7(11).$
1064	doi: 10.1029/2006GC001278
1065	Dressing, C. D., Spiegel, D. S., Scharf, C. A., Menou, K., & Raymond, S. N. (2010).
1066	Habitable climates: the influence of eccentricity. The Astrophysical Journal,
1067	721(2), 1295. doi: 10.1088/0004-637X/721/2/1295
1068	Dubrovinsky, L. S., Saxena, S. K., Tutti, F., Rekhi, S., & LeBehan, T. (2000).
1069	In situ X-ray study of thermal expansion and phase transition of iron
1070	at multimegabar pressure. Physical Review Letters, $84(8)$ , 1720. doi:
1071	10.1103/PhysRevLett.84.1720
1072	Duffy, T. S., Hemley, R. J., & Mao, HK. (1995). Equation of state and shear
1073	strength at multimegabar pressures: Magnesium oxide to 227 GPa. $Physical$
1074	Review Letters, 74(8), 1371. doi: 10.1103/PhysRevLett.74.1371
1075	Duffy, T. S., Meade, C., Fei, Y., Mao, HK., & Hemley, R. J. (1995). High-pressure
1076	phase transition in brucite, Mg (OH) <sub>2</sub> . American Mineralogist, $80(3-4)$ , 222–
1077	230. doi: 10.2138/am-1995-3-403
1078	Dyar, M. D., Helbert, J., Cooper, R. F., Sklute, E. C., Maturilli, A., Mueller, N. T.,
1079	Smrekar, S. E. (2021). Surface weathering on Venus: Constraints from
1080	kinetic, spectroscopic, and geochemical data. <i>Icarus</i> , 358, 114139. doi:
1081	10.1016/j.icarus.2020.114139
1082	Edson, A. R., Kasting, J. F., Pollard, D., Lee, S., & Bannon, P. R. (2012). The
1083	carbonate-silicate cycle and $\mathrm{CO}_2/\mathrm{climate}$ feedbacks on tidally locked terrestrial
1084	planets. Astrobiology, $12(6)$ , 562–571. doi: 10.1089/ast.2011.0762
1085	Engel, S., Lunine, J. I., & Norton, D. L. (1994). Silicate interactions with ammonia-
1086	water fluids on early Titan. Journal of Geophysical Research: Planets, $99(E2)$ ,
1087	3745–3752. doi: 10.1029/93JE03433
1088	Fei, Y., Prewitt, C. T., Mao, Hk., & Bertka, C. M. (1995). Structure and density of
1089	FeS at high pressure and high temperature and the internal structure of Mars.
1090	Science, $268(5219)$ , 1892–1894. doi: 10.1126/science.268.5219.1892

1091	Felton, R. C., Bastelberger, S. T., Mandt, K. E., Luspay-Kuti, A., Fauchez, T. J.,
1092	& Domagal-Goldman, S. D. (2022). The role of atmospheric exchange in
1093	false-positive biosignature detection. Journal of Geophysical Research: Planets,
1094	127(3), e2021JE006853. doi: 10.1029/2021JE006853
1095	Fischer, K. M., Ford, H. A., Abt, D. L., & Rychert, C. A. (2010). The lithosphere-
1096	asthenosphere boundary. Annual Review of Earth and Planetary Sciences, 38,
1097	551–575. doi: 10.1146/annurev-earth-040809-152438
1098	Foley, B. J. (2015). The role of plate tectonic–climate coupling and exposed land
1099	area in the development of habitable climates on rocky planets. The Astro-
1100	physical Journal, 812(1), 36. doi: 10.1088/0004-637X/812/1/36
1101	Forget, F., & Leconte, J. (2014). Possible climates on terrestrial exoplanets. <i>Phil.</i>
1102	Trans. R. Soc. A, 372(2014), 20130084. doi: 10.1098/rsta.2013.0084
1103	Fortes, A. D., Brand, H., Vočadlo, L., Lindsay-Scott, A., Fernandez-Alonso,
1104	F., & Wood, I. (2013). $P-V-T$ equation of state of synthetic mirabilite
1105	$(Na_2SO_4 \cdot 10D_2O)$ determined by powder neutron diffraction. Journal of Ap-
1106	plied Crystallography, $46(2)$ , 448–460. doi: 10.1107/S0021889813001362
1107	Fortes, A. D., Wood, I. G., Brodholt, J. P., & Vočadlo, L. (2003). The structure,
1108	ordering and equation of state of ammonia dihydrate ( $NH_3 \cdot 2H_2O$ ). Icarus,
1109	162(1), 59-73. doi: 10.1016/S0019-1035(02)00073-8
1110	Frank, M. R., Aarestad, E., Scott, H. P., & Prakapenka, V. B. (2013). A comparison
1111	of ice VII formed in the $H_2O$ , NaCl- $H_2O$ , and $CH_3OH-H_2O$ systems: Implica-
1112	tions for $H_2O$ -rich planets. Physics of the Earth and Planetary Interiors, 215,
1113	12–20. doi: 10.1016/j.pepi.2012.10.010
1114	Fujii, Y., Angerhausen, D., Deitrick, R., Domagal-Goldman, S., Grenfell, J. L., Hori,
1115	Y., Stevenson, K. B. (2018). Exoplanet biosignatures: observational
1116	prospects. Astrobiology, $18(6)$ , 739–778. doi: 10.1089/ast.2017.1733
1117	Gaillard, F., & Scaillet, B. (2014). A theoretical framework for volcanic degassing
1118	chemistry in a comparative planetology perspective and implications for plan-
1119	etary atmospheres. Earth and Planetary Science Letters, 403, 307–316. doi:
1120	10.1016/j.epsl.2014.07.009
1121	Gaillard, F., Scaillet, B., & Arndt, N. T. (2011). Atmospheric oxygenation caused by
1122	a change in volcanic degassing pressure. Nature, $478(7368),229.$ doi: 10.1038/
1123	nature10460

1124	Gerlach, T. M., & Graeber, E. J. (1985). Volatile budget of Kilauea volcano. Nature,
1125	313(6000), 273-277.doi: 10.1038/313273a0
1126	Ghiorso, M. S., & Sack, R. O. (1995). Chemical mass transfer in magmatic pro-
1127	cesses IV. A revised and internally consistent thermodynamic model for the
1128	interpolation and extrapolation of liquid-solid equilibria in magmatic systems
1129	at elevated temperatures and pressures. Contributions to Mineralogy and
1130	Petrology, 119(2), 197-212.doi: 10.1007/BF00307281
1131	Gibbs, R. (1975). Assessing Potential Ocean Pollutants. National Academy of Sci-
1132	ences.
1133	Gillon, M., Triaud, A. H., Demory, BO., Jehin, E., Agol, E., Deck, K. M.,
1134	Queloz, D. (2017). Seven temperate terrestrial planets around the nearby
1135	ultracool dwarf star TRAPPIST-1. Nature, $542(7642)$ , $456-460$ . doi:
1136	10.1038/nature21360
1137	Grenfell, J. L. (2017). A review of exoplanetary biosignatures. Physics Reports, 713,
1138	1-17. doi: 10.1016/j.physrep.2017.08.003
1139	Griffiths, G. I., Fortes, A. D., Pickard, C. J., & Needs, R. (2012). Crystal structure
1140	of ammonia dihydrate II. The Journal of Chemical Physics, 136(17), 174512.
1141	doi: 10.1063/1.4707930
1142	Gromnitskaya, E. L., Yagafarov, O. F., Lyapin, A. G., Brazhkin, V. V., Wood, I. G.,
1143	Tucker, M. G., & Fortes, A. D. (2013). The high-pressure phase diagram
1144	of synthetic epsomite (MgSO <sub>4</sub> ·7H <sub>2</sub> O and MgSO <sub>4</sub> ·7D <sub>2</sub> O) from ultrasonic and
1145	neutron powder diffraction measurements. Physics and Chemistry of Minerals,
1146	40(3), 271-285. doi: 10.1007/s00269-013-0567-7
1147	Haqq-Misra, J. D., Domagal-Goldman, S. D., Kasting, P. J., & Kasting, J. F.
1148	(2008). A revised, hazy methane greenhouse for the Archean Earth. $Astro-$
1149	biology, 8(6), 1127-1137.doi: 10.1089/ast.2007.0197
1150	Harrison, C. $(1999)$ . Constraints on ocean volume change since the Archean. Geo-
1151	physical Research Letters, $26(13)$ , 1913–1916. doi: 10.1029/1999GL900425
1152	Hartley, M. E., Maclennan, J., Edmonds, M., & Thordarson, T. (2014). Reconstruct-
1153	ing the deep $CO_2$ degassing behaviour of large basaltic fissure eruptions. Earth
1154	and Planetary Science Letters, 393, 120–131. doi: 10.1016/j.epsl.2014.02.031
1155	Hasenclever, J., Theissen-Krah, S., Rüpke, L. H., Morgan, J. P., Iyer, K., Petersen,
1156	S., & Devey, C. W. (2014). Hybrid shallow on-axis and deep off-axis hy-

-49-

1157	drothermal circulation at fast-spreading ridges. Nature, 508(7497), 508–512.
1158	doi: 10.1038/nature13174
1159	Hekinian, R., Pineau, F., Shilobreeva, S., Bideau, D., Gracia, E., & Javoy, M.
1160	(2000). Deep sea explosive activity on the Mid-Atlantic Ridge near 34
1161	50' N: magma composition, vesicularity and volatile content. Journal
1162	of Volcanology and Geothermal Research, 98(1-4), 49–77. doi: 10.1016/
1163	S0377-0273(99)00190-0
1164	Hofmann, A. W. (1988). Chemical differentiation of the Earth: the relationship be-
1165	tween mantle, continental crust, and oceanic crust. Earth and Planetary Sci-
1166	ence Letters, $90(3)$ , 297–314. doi: 10.1016/0012-821X(88)90132-X
1167	Hu, R., Seager, S., & Bains, W. (2012). Photochemistry in terrestrial exoplanet at-
1168	mospheres. I. Photochemistry model and benchmark cases. The Astrophysical
1169	Journal, 761(2), 166. doi: 10.1088/0004-637X/761/2/166
1170	Johansen, A., Ronnet, T., Bizzarro, M., Schiller, M., Lambrechts, M., Nordlund,
1171	Å., & Lammer, H. (2021). A pebble accretion model for the formation of the
1172	terrestrial planets in the Solar System. Science Advances, $7(8)$ , eabc0444. doi:
1173	10.1126/sciadv.abc0444
1174	Johnson, H. P., & Pruis, M. J. (2003). Fluxes of fluid and heat from the oceanic
1175	crustal reservoir. Earth and Planetary Science Letters, 216(4), 565–574. doi:
1176	10.1016/S0012-821X(03)00545-4
1177	Jones, M., Soule, S., Gonnermann, H., Le Roux, V., & Clague, D. (2018). Magma
1178	ascent and lava flow emplacement rates during the 2011 Axial Seamount erup-
1179	tion based on $CO_2$ degassing. Earth and Planetary Science Letters, 494,
1180	32-41. doi: 10.1016/j.epsl.2018.04.044
1181	Kadko, D., Baross, J., & Alt, J. (1995). The magnitude and global implications of
1182	hydrothermal flux. Geophysical Monograph-American Geophysical Union, 91,
1183	446–466. doi: $10.1029/GM091p0446$
1184	Kane, S. R., Hill, M. L., Kasting, J. F., Kopparapu, R. K., Quintana, E. V., Barclay,
1185	T., Torres, G. (2016). A catalog of Kepler habitable zone exoplanet candi-
1186	dates. The Astrophysical Journal, $830(1)$ , 1. doi: $10.3847/0004-637X/830/1/$
1187	1
1188	Kaspi, Y., & Showman, A. P. (2015). Atmospheric dynamics of terrestrial exoplan-
1189	ets over a wide range of orbital and atmospheric parameters. The Astrophysical

Journal, 804 (1), 60. doi: 10.1088/0004-637X/804/1/60
 Kasting, J. F. (1988). Runaway and moist greenhouse atmospheres and the evolution of Earth and Venus. *Icarus*, 74 (3), 472–494. doi: 10.1016/0019-1035(88)
 90116-9

- Kasting, J. F., & Ackerman, T. P. (1986). Climatic consequences of very high carbon dioxide levels in the Earth's early atmosphere. *Science*, 234 (4782), 1383–
  1385. doi: 10.1126/science.11539665
- Katsura, T., Yoneda, A., Yamazaki, D., Yoshino, T., & Ito, E. (2010). Adiabatic
   temperature profile in the mantle. *Physics of the Earth and Planetary Interi-* ors, 183(1-2), 212–218. doi: 10.1016/j.pepi.2010.07.001
- <sup>1200</sup> Kiang, N. Y., Domagal-Goldman, S., Parenteau, M. N., Catling, D. C., Fujii, Y.,
- Meadows, V. S., ... Walker, S. I. (2018). Exoplanet biosignatures: at the dawn of a new era of planetary observations. *Astrobiology*, 18(6), 619–629. doi: 10.1089/ast.2018.1862
- Kiessling, W., Flügel, E., & Golonka, J. (2003). Patterns of Phanerozoic carbonate platform sedimentation. Lethaia, 36(3), 195–225. doi: 10.1080/
  00241160310004648
- Kirk, R. L., & Stevenson, D. J. (1987). Thermal evolution of a differentiated
   Ganymede and implications for surface features. *Icarus*, 69(1), 91–134. doi:
   10.1016/0019-1035(87)90009-1
- Kite, E. S., Manga, M., & Gaidos, E. (2009). Geodynamics and rate of volcanism on
   massive Earth-like planets. *The Astrophysical Journal*, 700(2), 1732. doi: 10
   .1088/0004-637X/700/2/1732
- <sup>1213</sup> Kopparapu, R. K., Ramirez, R., Kasting, J. F., Eymet, V., Robinson, T. D., Ma-
- hadevan, S., ... Deshpande, R. (2013). Habitable zones around mainsequence stars: New estimates. *The Astrophysical Journal*, 765, 131. doi:
  10.1088/0004-637X/765/2/131
- Kopparapu, R. K., Wolf, E. T., & Meadows, V. S. (2020). Characterizing exoplanet
   habitability. In *Planetary Astrobiology* (p. 449). University of Arizona Press.
- Korenaga, J. (2008). Urey ratio and the structure and evolution of Earth's mantle.
   *Reviews of Geophysics*, 46(2). doi: 10.1029/2007RG000241
- Korenaga, J. (2018). Crustal evolution and mantle dynamics through Earth history.
   *Philosophical Transactions of the Royal Society A: Mathematical, Physical and*

1223	Engineering Sciences, 376 (2132), 20170408. doi: 10.1098/rsta.2017.0408
1224	Korenaga, J., & Karato, SI. (2008). A new analysis of experimental data on olivine
1225	rheology. Journal of Geophysical Research: Solid Earth, 113(B2). doi: 10
1226	.1029/2007 JB005100
1227	Krissansen-Totton, J., Arney, G. N., & Catling, D. C. (2018). Constraining the
1228	climate and ocean pH of the early Earth with a geological carbon cycle model.
1229	Proceedings of the National Academy of Sciences, 115(16), 4105–4110. doi:
1230	10.1073/pnas.1721296115
1231	Krissansen-Totton, J., Bergsman, D. S., & Catling, D. C. (2016). On detecting
1232	biospheres from chemical thermodynamic disequilibrium in planetary atmo-
1233	spheres. Astrobiology, $16(1)$ , 39–67. doi: 10.1089/ast.2015.1327
1234	Krissansen-Totton, J., & Catling, D. C. (2017). Constraining climate sensitivity and
1235	continental versus seafloor weathering using an inverse geological carbon cycle
1236	model. Nature Communications, $\mathcal{S}(1)$ , 1–15. doi: 10.1038/ncomms15423
1237	Krissansen-Totton, J., Garland, R., Irwin, P., & Catling, D. C. (2018). Detectability
1238	of biosignatures in an oxic atmospheres with the James Webb Space Telescope:
1239	A TRAPPIST-1e case study. The Astronomical Journal, 156(3), 114. doi:
1240	10.3847/1538- $3881/aad564$
1241	Krissansen-Totton, J., Thompson, M., Galloway, M. L., & Fortney, J. J. (2022).
1242	Understanding planetary context to enable life detection on exoplanets and
1243	test the Copernican principle. Nature Astronomy, $6(2)$ , 189–198. doi:
1244	10.1038/s41550-021-01579-7
1245	Laneuville, M., Kameya, M., & Cleaves, H. J. (2018). Earth without life: A systems
1246	model of a global abiotic nitrogen cycle. Astrobiology, 18, 897–914. doi: 10
1247	.1089/ast.2017.1700
1248	Lehmer, O. R., Catling, D. C., & Krissansen-Totton, J. (2020). Carbonate-
1249	silicate cycle predictions of Earth-like planetary climates and testing the
1250	habitable zone concept. Nature Communications, $11(1)$ , 1–10. doi:
1251	10.1038/s41467-020-19896-2
1252	Lerman, A., & Wu, L. (2006). $CO_2$ and sulfuric acid controls of weathering and river
1253	water composition. Journal of Geochemical Exploration, 88(1-3), 427–430. doi:
1254	10.1016/j.gexplo.2005.08.100
1255	Lerman, A., Wu, L., & Mackenzie, F. T. (2007). CO <sub>2</sub> and H <sub>2</sub> SO <sub>4</sub> consump-

-52-

manuscript submitted to JGR: Planets

1256	tion in weathering and material transport to the ocean, and their role in
1257	the global carbon balance. Marine Chemistry, $106(1-2)$ , $326-350$ . doi:
1258	10.1016/j.marchem.2006.04.004
1259	Lewis, N. T., Lambert, F. H., Boutle, I. A., Mayne, N. J., Manners, J., & Acreman,
1260	D. M. (2018). The influence of a substellar continent on the climate of a
1261	tidally locked exoplanet. The Astrophysical Journal, 854(2), 171.
1262	Lin, Z., MacDonald, R. J., Kaltenegger, L., & Wilson, D. J. (2021). Differentiat-
1263	ing modern and prebiotic Earth scenarios for TRAPPIST-1e: high-resolution
1264	transmission spectra and predictions for JWST. Monthly Notices of the Royal
1265	Astronomical Society, $505(3)$ , $3562-3578$ . doi: $10.1093/mnras/stab1486$
1266	Lisse, C. M., Desch, S. J., Unterborn, C. T., Kane, S. R., Young, P. R., Hartnett,
1267	H. E., Izenberg, N. R. (2020). A geologically robust procedure for ob-
1268	serving rocky exoplanets to ensure that detection of atmospheric oxygen is a
1269	modern Earth-like biosignature. The Astrophysical Journal Letters, $898(1)$ ,
1270	L17. doi: 10.3847/2041-8213/ab9b91
1271	Lorenzo, A., Desch, S. J., & Shim, SH. (2014). On the lower radius limit of exo-
1272	planets. In Lunar and Planetary Science Conference (p. 1636).
1273	Lynch, J. K., Beatty, C. M., Seidel, M. P., Jungst, L. J., & DeGrandpre, M. D.
1274	(2010). Controls of riverine $CO_2$ over an annual cycle determined using di-
1275	rect, high temporal resolution $pCO_2$ measurements. Journal of Geophysical
1276	Research: Biogeosciences, $115(G3)$ . doi: $10.1029/2009JG001132$
1277	Lyubetskaya, T., & Korenaga, J. (2007). Chemical composition of Earth's prim-
1278	itive mantle and its variance: 1. Method and results. Journal of Geophysical
1279	Research: Solid Earth, 112(B3). doi: 10.1029/2005JB004223
1280	Ma, C., Wu, X., Huang, F., Zhou, Q., Li, F., & Cui, Q. (2012). The acoustic ve-
1281	locity, refractive index, and equation of state of liquid ammonia dihydrate
1282	under high pressure and high temperature. The Journal of Chemical Physics,
1283	137(10), 104504. doi: 10.1063/1.4751944
1284	Madden, J., & Kaltenegger, L. (2020). How surfaces shape the climate of habitable
1285	exoplanets. Monthly Notices of the Royal Astronomical Society, 495(1), 1–11.
1286	doi: 10.1093/mnras/staa387
1287	Martin, JM., & Meybeck, M. (1979). Elemental mass-balance of material carried
1288	by major world rivers. Marine Chemistry, 7(3), 173–206. doi: 10.1016/0304

1289	-4203(79)90039-2
1290	McDonough, W. F., & Sun, SS. (1995). The composition of the Earth. Chemical
1291	Geology, 120(3-4), 223–253. doi: 10.1016/0009-2541(94)00140-4
1292	McKenzie, D. A. N., & Bickle, M. J. (1988). The volume and composition of melt
1293	generated by extension of the lithosphere. Journal of Petrology, $29(3)$ , $625-$
1294	679. doi: $10.1093/\text{petrology}/29.3.625$
1295	McLennan, S. M., & Grotzinger, J. P. (2008). The sedimentary rock cycle of Mars.
1296	In The Martian Surface–Composition, Mineralogy, and Physical Properties
1297	(pp. 541–577). Cambridge University Press.
1298	Mikal-Evans, T. $(2022)$ . Detecting the proposed $CH_4$ – $CO_2$ biosignature pair with
1299	the James Webb Space Telescope: TRAPPIST-1e and the effect of cloud/haze.
1300	Monthly Notices of the Royal Astronomical Society, $510(1)$ , $980-991$ . doi:
1301	10.1093/mnras/stab3383
1302	Ming, D. W., Morris, R. V., & Clark, B. C. (2008). Aqueous alteration on Mars. In
1303	$The \ Martian \ Surface-Composition, \ Mineralogy, \ and \ Physical \ Properties \ (pp.$
1304	519–540). Cambridge University Press.
1305	Moresi, LN., & Solomatov, V. S. (1995). Numerical investigation of 2D convection
1306	with extremely large viscosity variations. Physics of Fluids, $\gamma(9)$ , 2154–2162.
1307	doi: 10.1063/1.868465
1308	Mottl, M. J. (1983). Metabasalts, axial hot springs, and the structure of hydrother-
1309	mal systems at mid-ocean ridges. Geological Society of America Bulletin,
1310	94(2), 161–180. doi: https://doi.org/10.1130/0016-7606(1983)94\%3C161:
1311	$MAHSAT \ 3E2.0.CO;2$
1312	Nesbitt, H. W., & Markovics, G. (1997). Weathering of granodioritic crust, long-
1313	term storage of elements in weathering profiles, and petrogenesis of siliciclas-
1314	tic sediments. Geochimica et Cosmochimica Acta, 61(8), 1653–1670. doi:
1315	10.1016/S0016-7037(97)00031-8
1316	Nestola, F., Angel, R. J., Zhao, J., Garrido, C. J., Sánchez-Vizcaíno, V. L., Capitani,
1317	G., & Mellini, M. (2010). Antigorite equation of state and anomalous softening
1318	at 6 GPa: an in situ single-crystal x-ray diffraction study. Contributions to
1319	Mineralogy and Petrology, $160(1)$ , $33-43$ . doi: $10.1007/s00410-009-0463-9$
1320	Neveu, M., Desch, S. J., & Castillo-Rogez, J. C. (2017). Aqueous geochemistry in
1321	icy world interiors: Equilibrium fluid, rock, and gas compositions, and fate

-54-

1322	of antifreezes and radionuclides. Geochimica et Cosmochimica Acta, 212,
1323	324–371. doi: 10.1016/j.gca.2017.06.023
1324	Nordstrom, D. K., Plummer, L. N., Wigley, T. M. L., Wolery, T. J., Ball, J. W.,
1325	Jenne, E. A., Thrailkill, J. (1979). A comparison of computer-
1326	ized chemical models for equilibrium calculations in aqueous systems. In
1327	ACS Symposium Series (Vol. 93, p. 857-892). ACS Publications. doi:
1328	10.1021/bk-1979-0093.ch038
1329	Oelkers, E., Bénézeth, P., & Pokrovski, G. S. (2009). Thermodynamic databases for
1330	water-rock interaction. Reviews in Mineralogy and Geochemistry, $70(1)$ , 1–46.
1331	doi: 10.2138/rmg.2009.70.1
1332	Orgel, L. E. (1998). The origin of life–a review of facts and speculations. Trends in
1333	Biochemical Sciences, 23(12), 491–495. doi: 10.1016/s0968-0004(98)01300-0
1334	Palandri, J. L., & Kharaka, Y. K. (2004). A compilation of rate parameters of
1335	water-mineral interaction kinetics for application to geochemical modeling
1336	(Tech. Rep.). US Geological Survey. Menlo Park. CA.
1337	Pamukcu, A. S., Gualda, G. A. R., Ghiorso, M. S., Miller, C. F., & McCracken,
1338	R. G. (2015). Phase-equilibrium geobarometers for silicic rocks based on
1339	rhyolite-MELTS—Part 3: Application to the Peach Spring tuff (Arizona–
1340	California–Nevada, USA). Contributions to Mineralogy and Petrology, 169(3),
1341	33. doi: 10.1007/s00410-015-1122-y
1342	Papike, J. J., Ryder, G., & Shearer, C. K. (1998). Lunar samples. In J. J. Papike
1343	(Ed.), Planetary Materials (Vol. 36). Walter de Gruyter GmbH & Co KG.
1344	Parkhurst, D. L., & Appelo, C. A. J. (2013). Description of input and examples
1345	for PHREEQC version $3-A$ computer program for speciation, batch-reaction,
1346	one-dimensional transport, and inverse geochemical calculations (Tech. Rep.).
1347	US Geological Survey, Reston, VA.
1348	Pendergrass, A. G. (2020). The global-mean precipitation response to $CO_2$ -
1349	induced warming in CMIP6 models. $Geophysical Research Letters, 47(17),$
1350	e2020GL089964. doi: $10.1029/2020GL089964$
1351	Pidhorodetska, D., Fauchez, T. J., Villanueva, G. L., Domagal-Goldman, S. D.,
1352	& Kopparapu, R. K. (2020). Detectability of molecular signatures on
1353	TRAPPIST-1e through transmission spectroscopy simulated for future space-
1354	based observatories. The Astrophysical Journal Letters, 898(2), L33. doi:

1355	10.3847/2041- $8213/aba4a1$
1356	Pierrehumbert, R. T., & Hammond, M. (2019). Atmospheric circulation of tide-
1357	locked exoplanets. Annual Review of Fluid Mechanics, 51, 275–303. doi: 10
1358	.1146/annurev-fluid-010518-040516
1359	Pokrovsky, O. S., & Schott, J. (1999). Processes at the magnesium-bearing
1360	carbonates/solution interface. II. Kinetics and mechanism of magnesite
1361	dissolution. Geochimica et Cosmochimica Acta, 63(6), 881–897. doi:
1362	10.1016/S0016-7037(99)00013-7
1363	Pope, E. C., Bird, D. K., & Rosing, M. T. (2012). Isotope composition and vol-
1364	ume of Earth's early oceans. Proceedings of the National Academy of Sciences,
1365	109(12), 4371-4376.doi: 10.1073/pnas.1115705109
1366	Press, W. H., Teukolsky, S. A., Vetterling, W. T., & Flannery, B. P. (1992). Numer-
1367	ical Recipes in C, 2nd edition. Cambridge University, New York.
1368	Rasilo, T., Hutchins, R. H. S., Ruiz-González, C., & Del Giorgio, P. A. (2017).
1369	Transport and transformation of soil-derived $CO_2$ , $CH_4$ and DOC sustain $CO_2$
1370	supersaturation in small boreal streams. Science of the Total Environment,
1371	579, 902–912. doi: 10.1016/j.scitotenv.2016.10.187
1372	Reid, W. H., & Harris, D. L. (1958). Some further results on the Bénard problem.
1373	The Physics of Fluids, $1(2)$ , 102–110. doi: 10.1063/1.1705871
1374	Ringwood, A. E. (1991). Phase transformations and their bearing on the consti-
1375	tution and dynamics of the mantle. Geochimica et Cosmochimica Acta, $55(8)$ ,
1376	2083–2110. doi: 10.1016/0016-7037(91)90090-R
1377	Rudnick, R. L., & Gao, S. (2003). Composition of the continental crust. In <i>Treatise</i>
1378	on Geochemistry (Vol. 3, p. 659). Elsevier.
1379	Rushby, A. J., Johnson, M., Mills, B. J., Watson, A. J., & Claire, M. W. (2018).
1380	Long-term planetary habitability and the carbonate-silicate cycle. Astrobiol-
1381	ogy, 18(5), 469-480.doi: 10.1089/ast.2017.1693
1382	Saito, G., Kazahaya, K., Shinohara, H., Stimac, J., & Kawanabe, Y. (2001). Varia-
1383	tion of volatile concentration in a magma system of Satsuma-Iwojima volcano
1384	deduced from melt inclusion analyses. Journal of Volcanology and Geothermal
1385	Research, $108(1-4)$ , 11–31. doi: 10.1016/S0377-0273(00)00276-6
1386	Saito, H., & Kuramoto, K. (2018). Formation of a hybrid-type proto-atmosphere on
1387	Mars accreting in the solar nebula. Monthly Notices of the Royal Astronomical

1388	Society, $475(1)$ , 1274-1287. doi: 10.1093/mnras/stx3176
1389	Sanloup, C., Jambon, A., & Gillet, P. (1999). A simple chondritic model of Mars.
1390	Physics of the Earth and Planetary Interiors, 112(1-2), 43–54. doi: 10.1016/
1391	S0031-9201(98)00175-7
1392	Sata, N., Hirose, K., Shen, G., Nakajima, Y., Ohishi, Y., & Hirao, N. (2010). Com-
1393	pression of FeSi, Fe3C, $Fe_{0.95}O$ , and FeS under the core pressures and impli-
1394	cation for light element in the Earth's core. Journal of Geophysical Research:
1395	Solid Earth, $115(B9)$ . doi: $10.1029/2009JB006975$
1396	Schaefer, L., & Fegley, B. (2017). Redox states of initial atmospheres outgassed on
1397	rocky planets and planetesimals. The Astrophysical Journal, $843(2)$ , 120. doi:
1398	10.3847/1538-4357/aa784f
1399	Schulze-Makuch, D., Méndez, A., Fairén, A. G., Von Paris, P., Turse, C., Boyer, G.,
1400	$\dots$ Irwin, L. N. (2011). A two-tiered approach to assessing the habitability of
1401	exoplanets. Astrobiology, $11(10)$ , 1041–1052. doi: 10.1089/ast.2010.0592
1402	Schwieterman, E. W., Kiang, N. Y., Parenteau, M. N., Harman, C. E., DasSarma,
1403	S., Fisher, T. M., Lyons, T. W. (2018). Exoplanet biosignatures: a re-
1404	view of remotely detectable signs of life. Astrobiology, $18(6)$ , $663-708$ . doi:
1405	10.1089/ast.2017.1729
1406	Seager, S., Kuchner, M., Hier-Majumder, C., & Militzer, B. (2007). Mass-radius re-
1407	lationships for solid exoplanets. The Astrophysical Journal, $669(2)$ , 1279. doi:
1408	10.1086/521346
1409	Shahar, A., Driscoll, P., Weinberger, A., & Cody, G. (2019). What makes a planet
1410	habitable? Science, $364(6439)$ , $434-435$ . doi: 10.1126/science.aaw4326
1411	Sleep, N. H., & Zahnle, K. (2001). Carbon dioxide cycling and implications for
1412	climate on ancient Earth. Journal of Geophysical Research: Planets, $106(E1)$ ,
1413	1373–1399. doi: $10.1029/2000$ JE001247
1414	Smith, P. M., & Asimow, P. D. (2005). Adiabat_1ph: A new public front-end to the
1415	MELTS, pMELTS, and pHMELTS models. Geochemistry, Geophysics, Geosys-
1416	tems, $6(2)$ . doi: 10.1029/2004GC000816
1417	Solomatov, V. S. (1995). Scaling of temperature-and stress-dependent viscosity con-
1418	vection. Physics of Fluids, 7(2), 266–274. doi: 10.1063/1.868624
1419	Spiegel, D. S., Menou, K., & Scharf, C. A. (2009). Habitable climates: the influence
1420	of obliquity. The Astrophysical Journal, 691(1), 596. doi: 10.1088/0004-637X/

691/1/596

1421

1422	Stamenković, V., Noack, L., Breuer, D., & Spohn, T. (2012). The influence of									
1423	pressure-dependent viscosity on the thermal evolution of super-Earths. The									
1424	Astrophysical Journal, 748(1), 41. doi: 10.1088/0004-637X/748/1/41									
1425	Tao, C., Sevfried, W., Lowell, R., Liu, Y., Liang, J., Guo, Z.,, Li, W. (2020).									
1426	Deep high-temperature hydrothermal circulation in a detachment faulting sys-									
1427	tem on the ultra-slow spreading ridge. Nature Communications, 11(1), 1–9.									
1428	doi: 10.1038/s41467-020-15062-w									
1429	Taylor, G. J. (2013). The bulk composition of Mars. <i>Geochemistry</i> , 73(4), 401–420.									
1430	doi: 10.1016/j.chemer.2013.09.006									
1431	Thompson, M. A., Krissansen-Totton, J., Wogan, N., Telus, M., & Fortney, J. J.									
1432	(2022). The case and context for atmospheric methane as an exoplanet									
1433	biosignature. Proceedings of the National Academy of Sciences, 119(14),									
1434	e2117933119. doi: 10.1073/pnas.2117933119									
1435	Thompson, R. N., Riches, A. J. V., Antoshechkina, P. M., Pearson, D. G., Nowell,									
1436	G. M., Ottley, C. J., Niku-Paavola, V. (2007). Origin of CFB magmatism:									
1437	multi-tiered intracrustal picrite–rhyolite magmatic plumbing at Spitzkoppe,									
1438	western Namibia, during early Cretaceous Etendeka magmatism. Journal of									
1439	Petrology, 48(6), 1119-1154.doi: 10.1093/petrology/egm012									
1440	Tosca, N. J., McLennan, S. M., Lindsley, D. H., & Schoonen, M. A. (2004).									
1441	Acid-sulfate weathering of synthetic Martian basalt: The acid fog model									
1442	revisited. Journal of Geophysical Research: Planets, 109(E5). doi:									
1443	10.1029/2003JE002218									
1444	Trail, D., Watson, E. B., & Tailby, N. D. (2011). The oxidation state of Hadean									
1445	magmas and implications for early Earth's atmosphere. Nature, $480(7375)$ , 79.									
1446	doi: 10.1038/nature10655									
1447	Treiman, A. H., & Schwenzer, S. P. (2009). Basalt–atmosphere interaction on									
1448	Venus: Preliminary results on weathering of minerals and bulk rock. In $Venus$									
1449	Geochemistry: Progress, Prospects, and New Missions (p. abstract $#2011$ ).									
1450	Trenberth, K. E., Fasullo, J., & Smith, L. (2005). Trends and variability in column-									
1451	integrated atmospheric water vapor. Climate dynamics, $24(7)$ , 741–758. doi:									
1452	10.1007/s00382-005-0017-4									

-58-

1454	Tyburczy, J. A., Duffy, T. S., Ahrens, T. J., & Lange, M. A. (1991). Shock wave
1455	equation of state of serpentine to 150 GPa: Implications for the occurrence
1456	of water in the Earth's lower mantle. Journal of Geophysical Research: Solid
1457	Earth, $96(B11)$ , 18011–18027. doi: 10.1029/91JB01573
1458	Tziperman, E., Abbot, D. S., Ashkenazy, Y., Gildor, H., Pollard, D., Schoof, C. G.,

- <sup>1459</sup> & Schrag, D. P. (2012). Continental constriction and oceanic ice-cover thick <sup>1460</sup> ness in a Snowball-Earth scenario. Journal of Geophysical Research: Oceans,
   <sup>1461</sup> 117(C5). doi: 10.1029/2011JC007730
- Unterborn, C. T., Byrne, P. K., Anbar, A. D., Arney, G. N., Brain, D., Desch, S. J.,
  ... Way, M. J. (2021). Exogeoscience and its role in characterizing exoplanet
  habitability and the detectability of life. Bulletin of the American Astronomical Society. doi: 10.3847/25c2cfeb.5209dd13
- Unterborn, C. T., Dismukes, E. E., & Panero, W. R. (2016). Scaling the Earth: a
   sensitivity analysis of terrestrial exoplanetary interior models. *The Astrophysi- cal Journal*, 819(1), 32. doi: 10.3847/0004-637X/819/1/32
- <sup>1469</sup> Unterborn, C. T., Foley, B. J., Desch, S. J., Young, P. A., Vance, G., Chiffelle,
- <sup>1470</sup> L., & Kane, S. R. (2022). Mantle degassing lifetimes through galac-
- tic time and the maximum age stagnant-lid rocky exoplanets can support
- temperate climates. *The Astrophysical Journal Letters*, 930(1), L6. doi: 1473 10.3847/2041-8213/ac6596
- Unterborn, C. T., Kabbes, J. E., Pigott, J. S., Reaman, D. M., & Panero, W. R.
  (2014). The role of carbon in extrasolar planetary geodynamics and habitability. *The Astrophysical Journal*, 793(2), 124. doi: 10.1088/0004-637X/793/2/
  124
- <sup>1478</sup> Urey, H. C. (1952). The Planets: their Origin and Development. Yale Univ. Press.
- Vance, S., Harnmeijer, J., Kimura, J., Hussmann, H., DeMartin, B., & Brown, J. M.
  (2007). Hydrothermal systems in small ocean planets. Astrobiology, 7(6),
  987–1005. doi: 10.1089/ast.2007.0075
- Vervoort, P., Horner, J., Kane, S. R., Turner, S. K., & Gilmore, J. B. (2022). System
   architecture and planetary obliquity: Implications for long-term habitability.
   The Astronomical Journal, 164 (4), 130. doi: 10.3847/1538-3881/ac87fd
- Vidaurri, M. R., Bastelberger, S. T., Wolf, E. T., Domagal-Goldman, S., & Koppa-
- rapu, R. K. (2022). The outer edge of the Venus zone around main-sequence

1487	stars. The Planetary Science Journal, $3(6)$ , 137. doi: 10.3847/PSJ/ac68e2
1488	Wade, J. A., Plank, T., Melson, W. G., Soto, G. J., & Hauri, E. H. (2006). The
1489	volatile content of magmas from Arenal volcano, Costa Rica. Journal of Vol-
1490	canology and Geothermal Research, 157(1-3), 94–120. doi: 10.1016/j.jvolgeores
1491	.2006.03.045
1492	Walsh, K. J., & Levison, H. F. (2019). Planetesimals to terrestrial planets: colli-
1493	sional evolution amidst a dissipating gas disk. Icarus, 329, 88–100. doi: 10
1494	.1016/j.icarus.2019.03.031
1495	Weider, S. Z. (2019). Petrology and geochemistry of Mercury. Oxford Research En-
1496	cyclopedia of Planetary Science, 127. doi: 10.1093/acrefore/9780190647926.013
1497	.127
1498	Weidner, D. J., Bass, J. D., Ringwood, A. E., & Sinclair, W. (1982). The single-
1499	crystal elastic moduli of stishovite. Journal of Geophysical Research: Solid
1500	Earth, $87(B6)$ , $4740-4746$ . doi: 10.1029/JB087iB06p04740
1501	Wett, J. P., & Ahrens, T. J. (1983). Shock compression of single-crystal forsterite.
1502	Journal of Geophysical Research: Solid Earth, 88(B11), 9500–9512. doi:
1503	10.1029/JB088iB11p09500
1504	Wordsworth, R. D., Forget, F., Selsis, F., Millour, E., Charnay, B., & Madeleine,
1505	JB. (2011). Gliese 581d is the first discovered terrestrial-mass exoplanet
1506	in the habitable zone. The Astrophysical Journal Letters, $733(2)$ , L48. doi:
1507	10.1088/2041-8205/733/2/L48
1508	Young, P. A., Desch, S. J., Anbar, A. D., Barnes, R., Hinkel, N. R., Kopparapu,
1509	R., Truitt, A. (2014). Astrobiological stoichiometry. Astrobiology, $14(7)$ ,
1510	603-626. doi: $10.1089/ast.2014.1143$
1511	Zhang, Y., & Zindler, A. (1993). Distribution and evolution of carbon and nitro-
1512	gen in Earth. Earth and Planetary Science Letters, 117(3-4), 331–345. doi: 10
1513	.1016/0012- $821X(93)90088$ -Q

Table 5. Starting crustal compositions in mol% for continental weathering simulations. Horizontal lines delineate broad mineral groups: pyroxenes and olivines, feldspars, cyclosilicates, silica polymorphs, hornblende, secondary minerals, carbonates, sulfates, sulfides, and halides. Notes and references: <sup>a</sup> Sedimentary. <sup>b</sup> Hydrothermal. <sup>c</sup> Matches both oxide mass fractions from Rudnick and Gao (2003) and mineral volume fractions from Nesbitt and Markovics (1997) within 10% prior to the addition of carbonates, sulfates, sulfates, and halite. The kinetic parameters of annite and daphnite are assumed to be the same as those of phlogopite and clinochlore, respectively, from Palandri and Kharaka (2004).<sup>d</sup> Matches primitive mantle oxide mass fractions of Hofmann (1988) within 4% prior to addition of pyrite and halite. <sup>e</sup> From Table 3 (average Mercury Southern hemisphere composition) of Weider (2019), matching both mineral and elemental mass fractions, and attributing all hyperstheme, olivine, and sulfide respectively as enstatite, forsterite, and pyrite to match the high Mg/Fe and S/Fe ratios.  $^{f}$  Results of modeled interaction between basalt and Venus' atmosphere (Treiman & Schwenzer, 2009; Dyar et al., 2021).  $^{g}$  Highland composition (Table 5.1 of Papike et al. (1998)) with feldspar composition from their Table A5.6, olivine composition from the Fe/(Fe+Mg) ratios of their Table A5.9, and Ca- and Mg-rich pyroxene compositions (attributed to 50% diopside / 25% enstatite / 25% wollastonite in case 1; 100% diopside in case 2) from columns 17 and 19 of their Table A5.4 due to the lack of kinetic data for Fe-pyroxene (ferrosilite) in Palandri and Kharaka (2004).  $^{h}$  From the Martian upper crust experimental mineralogy used by Tosca et al. (2004) as reported in Table 24.3 of McLennan and Grotzinger (2008), with plagioclase attributed to albite based on the lack of anorthite in a Meridiani Planum outcrop (Table 23.3 of Ming et al. (2008)), oxides attributed to hematite, and addition of jarosite and anhydrite at abundances bracketed by the lack of sulfates in the Tosca et al. (2004) composition and the tens of mass% of sulfates in Table 24.3 of McLennan and Grotzinger (2008) and Table 26.2 of Des Marais et al. (2007). <sup>i</sup> Compositions for Fig. 4b and 5d of De Sanctis et al. (2015), assuming their optically dark component is magnetite.

Mineral	Formula	$\begin{array}{c} \operatorname{Earth}^{c} \\ (\operatorname{modern}) \end{array}$	$\begin{array}{c} \operatorname{Earth}^{d} \\ (\operatorname{Archean}) \end{array}$	Mercury	Venus $1^f$ (reduced)	Venus $2^f$ (oxidized)	Lunar 1 <sup>9</sup>	Lunar 2 <sup>9</sup>	<sup><i>I</i></sup> Mars <sup><i>h</i></sup>	Ceres 1	$i$ Ceres $2^i$
Fayalite Forsterite Enstatite Wollastonite Diopside	$\begin{array}{c} \mathrm{Fe_2SiO_4}\\ \mathrm{Mg_2SiO_4}\\ \mathrm{MgSiO_3}\\ \mathrm{CaSiO_3}\\ \mathrm{CaMg(SiO_3)_2} \end{array}$	- - - -	7.5 53.6 27.2 3.1	$\frac{8.5}{51.1}$	- 38.8 -	12.2	$2.0 \\ 4.9 \\ 9.5 \\ 9.5 \\ 18.9$	2.4 5.3 - 32.4	2.0 10.7 23.8 -	- - - -	
Anorthite Albite K-feldspar	$\begin{array}{c} { m CaAl_2Si_2O_8} \\ { m NaAlSi_3O_8} \\ { m KAlSi_3O_8} \end{array}$	$^{8.4}_{15.6}_{5.0}$	4.7 1.5	$11.7 \\ 17.9 \\ 0.4$	11.8	11.2	$16.5 \\ 0.9 \\ 0.1$	$17.7 \\ 0.9 \\ 0.2$	$12.3 \\ 0.4$	- - -	- - -
Cordierite	$\mathrm{Mg_2Al_4Si_5O_{18}}$	-	-	-	7.6	7.2	-	-	-	-	-
$\begin{array}{c} \label{eq:Quartz} \\ \mbox{Amorph. SiO}_2 \end{array}$	${siO_2} {SiO_2}$	59.7	-	-	12.2	31.1	37.8	41.2	39.2	-	-
Anthophyllite Tremolite	$\substack{ Mg_{7}Si_{8}O_{22}(OH)_{2}\\ Ca_{2}Mg_{5}Si_{8}O_{22}(OH)_{2} }$	$\begin{array}{c} 0.21 \\ 0.43 \end{array}$	-	-	-	-	-	-	-	-	-
Annite Phlogopite Daphnite Lizardite Magnetite Hematite	$\begin{array}{c} {\rm KAlFe_3Si_3O_{10}(OH)_2}\\ {\rm KAlMg_3Si_3O_{10}(OH)_2}\\ {\rm Fe_5AlAlSi_3O_{10}(OH)_8}\\ {\rm Mg_3Si_2O_5(OH)_4}\\ {\rm Fe_3O_4}\\ {\rm Fe_2O_3} \end{array}$	1.0 1.8 1.4 - -	- - - - -		- - - -	- - 9.9		- - - - -	- - - 7.1	1.6 - 3.6 57.1	2.2 - 4.4 72.3
	${caCO_3 \atop CaMg(CO_3)_2 \atop CaMg(CO_3)_2 \atop MgCO_3}$	4.8 1.7 -	6 	1	- - -	- - -	- - -	- - - -	- - -	- 37.7	- 21.2
Gypsum Anhydrite Jarosite	$\begin{array}{c} {\rm CaSO_4\cdot 2H_2O}\\ {\rm CaSO_4}\\ {\rm KFe_3(SO_4)_2(OH)_6} \end{array}$	0.89	- - -	- - -	29.6	28.4	- - -	- - -	$2.8 \\ 1.8$	- - -	- - -
Pyrite	$\mathrm{FeS}_2$	0.17	0.21	6.1	-	-	-	-	-	-	-
Halite	NaCl	3.8	2.3	-	-	-	-	-	-	-	-

# Geophysical and geochemical controls on abiotic carbon cycling on Earth-like planets

M. Neveu<sup>1,2</sup>, T. Bartlow<sup>1</sup>, R. Felton<sup>3</sup>, S. Domagal-Goldman<sup>2</sup>, S. Desch<sup>4</sup>

<sup>1</sup>University of Maryland, College Park, 4296 Stadium Drive, College Park, MD 20742, USA
 <sup>2</sup>NASA Goddard Space Flight Center, 8800 Greenbelt Road, Greenbelt, MD 20771, USA
 <sup>3</sup>NASA Ames Research Center, Moffett Field, CA 94043, USA
 <sup>4</sup>Arizona State University, 781 E Terrace Mall, Tempe, AZ 85287

# **Key Points:**

3

# A new code enables modeling of carbon cycling on planets of size and composition that differ from Earth's. Outgassing tends to go down with geologic time, is maximal for planets around 1 Earth mass, and higher for more felsic upper mantles. Continental weathering is comparatively less sensitive to surface composition and patchiness, and insensitive to planet size.

Corresponding author: Marc Neveu, marc.f.neveu@nasa.gov

### 15 Abstract

We investigate how variations in a planet's size and the chemical (mineral) composition 16 of its upper mantle and surface affect processes involved in the carbonate-silicate cycle, 17 which is thought to have regulated the composition of Earth's atmosphere and its sur-18 face temperature over geologic time. We present models of geophysical and geochemi-19 cal controls on these processes: outgassing, continental weathering, and seafloor weath-20 ering, and analyze sensitivities to planet size and composition. For Earth-like compo-21 sitions, outgassing is maximized for planets of Earth's size. Smaller planets convect less 22 vigorously; higher pressures inside larger planets hinder melting. For more felsic man-23 tles, smaller planets (0.5-0.75 Earth mass) outgas more, whereas more mafic planets fol-24 low the size trend of Earth's composition. Planet size and composition can affect out-25 gassing by two orders of magnitude, with variability driven by mass in the first 2.5 Gyr 26 after formation and by composition past that time. In contrast, simulations spanning 27 the diversity of surface compositions encountered in the inner solar system indicate that 28 continental weathering fluxes are about as sensitive to surface composition or the patch-29 iness of land as they are to surface temperature, with fluxes within a factor of five of Earth's. 30 Seafloor weathering appears more sensitive to uncertainties in tectonic regime (occur-31 rence, speed, and size of plates) than to seafloor composition. These results form a ba-32 sis to interpret calculations of geological surface carbon fluxes to track atmospheric com-33 positions, through time, of lifeless exo-Earths, providing a baseline against which the ef-34 fect of biological activity may be distinguished with telescopic observations. 35

36

# Plain Language Summary

Earth's surface pressure and temperature have been relatively stable over its his-37 tory. They are likely regulated by the greenhouse effect induced by carbon dioxide  $(CO_2)$ 38 in Earth's atmosphere.  $CO_2$  is thought to be outgassed from the mantle, e.g., through 39 volcanism.  $CO_2$  is removed from the atmosphere by chemical weathering (the reaction 40 of surface rocks with  $CO_2$ -bearing rainwater), and river transport of its products to the 41 ocean where carbonate rocks form. Carbonates are subducted to the mantle, complet-42 ing the carbonate-silicate cycle. Here, we develop a model of mantle convection, melt-43 ing, and chemical reactions between water and rock to investigate how this cycle might 44 vary on solid planets with different sizes and mineral compositions. We find that these 45 affect carbon outgassing more than weathering. This new ability to model carbon cy-46

cling on a diversity of solid planets with an ocean helps understand what these planets

<sup>48</sup> might look like without a biosphere. Deviations from that baseline could be used to search<sup>49</sup> for signs of life.

50 1 Introduction

Thousands of planets are now known to orbit stars other than our Sun (NASA Ex-51 oplanet Archive; Akeson et al. (2013)). Amidst the zoo of observed combinations of host-52 star types, star-planet distances, orbital eccentricities, and planet sizes, a handful of plan-53 ets bearing similarities with Earth have been identified (Schulze-Makuch et al., 2011; Barnes 54 et al., 2015; Kane et al., 2016; Gillon et al., 2017). These are primarily solid worlds for 55 which "exo-climate" models have predicted temperatures that might allow surface liq-56 uid water (Wordsworth et al., 2011). Such potentially habitable worlds show promise as 57 targets for detecting signs of life, e.g., via atmospheric spectral signatures (Krissansen-58 Totton et al., 2016; Grenfell, 2017; Schwieterman et al., 2018; Krissansen-Totton, Gar-59 land, et al., 2018; M. A. Thompson et al., 2022) in the coming decades (Kiang et al., 2018; 60 Fujii et al., 2018; Pidhorodetska et al., 2020; Lin et al., 2021; Mikal-Evans, 2022). 61

Confidently attributing atmospheric spectral features to biology demands a thor-62 ough understanding of the physics, chemistry, and geology of exoplanets (Krissansen-63 Totton & Catling, 2017; Catling et al., 2018; Lisse et al., 2020). To detect life on an ex-64 oplanet, we must first constrain the geochemical cycles on a lifeless world, so that we rec-65 ognize what is not attributable to abiotic processes. This requires the ability to model 66 the interplay among interior, surface, atmospheric, and orbital processes, which is an over-67 arching goal of the exoplanet community (Forget & Leconte, 2014; Lehmer et al., 2020; 68 Barnes et al., 2020; Kopparapu et al., 2020; Unterborn et al., 2021; Krissansen-Totton 69 et al., 2022). 70

Previous efforts to understand exoplanet geology and climate have so far focused
on tidal (Barnes et al., 2013; Pierrehumbert & Hammond, 2019; Blackledge et al., 2020;
Colose et al., 2021), orbital (Spiegel et al., 2009; Dressing et al., 2010; Kaspi & Showman, 2015; Lehmer et al., 2020; Vervoort et al., 2022), compositional (Bond et al., 2010;
Young et al., 2014; Unterborn et al., 2014, 2016; Shahar et al., 2019; Unterborn et al.,
2022), and land coverage controls (Abbot et al., 2012; Tziperman et al., 2012; Lewis et
al., 2018; Krissansen-Totton, Arney, & Catling, 2018; Madden & Kaltenegger, 2020). Foley

-3-

(2015) has shown that the negative climate feedbacks inherent in the long-term carbon
cycle are uninhibited by climate's effect on plate tectonics. Models of atmospheric chemistry and chemical evolution have assumed surface fluxes of gases from outgassing or weathering as lower boundary conditions (Hu et al., 2012; Domagal-Goldman et al., 2014; Felton et al., 2022), usually estimated from present Earth values (Edson et al., 2012; KrissansenTotton & Catling, 2017; Krissansen-Totton, Arney, & Catling, 2018; Lehmer et al., 2020).

Outgassing rates have been predicted to increase with planet size (Kite et al., 2009). 84 In addition, the oxidation state of a planet's environment influences the surface and at-85 mospheric speciation of key elements such as carbon. Carbon is a chief component of plan-86 etary atmospheres as  $CO_2$ , CO, or  $CH_4$ ; and an essential building block of life as we know 87 it (Orgel, 1998). As CO<sub>2</sub>, it is also a major control of Earth's climate via the carbonate-88 silicate cycle (Kasting, 1988). While some studies have probed the photochemistry of 89 reduced carbon-rich atmospheres on terrestrial planets (Haqq-Misra et al., 2008; Arney 90 et al., 2017; Felton et al., 2022), there is a need to investigate how the carbonate-silicate 91 cycle might differ on worlds with a different bulk composition (Bond et al., 2010; Young 92 et al., 2014; Unterborn et al., 2022). 93

To address this need, we have developed a model of geophysical and geochemical 94 controls on abiotic cycling of carbon, on planets of 0.5 to 2 Earth masses  $(M_{\oplus})$  with the 95 same insolation as Earth. We use this model to investigate how the carbonate-silicate 96 cycle varies depending on planet bulk composition, size, and age. The model incorpo-97 rates key features of carbon cycling (Rushby et al., 2018): (1) equilibrium between at-98 mospheric carbon and dissolved carbon in the ocean, (2) removal from the atmosphere 99 by dissolving in rainwater and reacting with silicate rocks (weathering), (3) carbon re-100 moval from the ocean by seafloor water-rock interaction (Sleep & Zahnle, 2001; Krissansen-101 Totton & Catling, 2017), and (4) release of atmospheric carbon by outgassing such as 102 volcanism or other venting. In this paper, we describe models for each of these processes 103 that are validated to modern Earth conditions, but are based on first principles of geo-104 physics and geochemistry rather than on scalings with modern-day Earth. This enables 105 a departure from the necessarily Earth-centric conditions beyond which Earth-based scal-106 ings no longer accurately approximate the underlying outgassing and weathering pro-107 cesses. The present models can thus be used to investigate planets whose size and chem-108 ical composition differ from Earth's. In a companion paper, we describe the results of 109 the integrated abiotic carbon cycling model. 110

-4-

#### manuscript submitted to JGR: Planets



Figure 1. Architecture of the box model used to compute a planet's atmospheric and ocean composition as a function of time, with key computed parameters and inputs. The two reservoirs of carbon are the mantle and {atmosphere+ocean}. Fluxes between these reservoirs are due to seafloor weathering and mantle outgassing. The seafloor weathering flux depends on ocean composition, which itself depends on material fluxes arising from continental weathering and on ocean equilibrium with the atmosphere.

This paper is organized as follows. In Section 2, we describe the models of continental weathering, seafloor weathering, and outgassing, and their ability to reproduce corresponding estimated carbon fluxes on modern Earth. In Section 3, continental weathering and outgassing fluxes are computed for a variety of planet sizes and upper mantle or surface compositions. Sensitivity to these properties and their implications are discussed in Section 4; in particular, an example computation of the full carbon cycle through time is presented. We provide major conclusions in Section 5.

## 118 **2** Model

The abiotic carbon cycling model (Fig. 1), determines net fluxes (in mol m<sup>-2</sup> s<sup>-1</sup>) of carbon species from the surface and interior to the atmosphere. We assume that these fluxes are primarily due to continental weathering, seafloor weathering, and outgassing, and that the atmosphere is equilibrated with the ocean into a single reservoir. This section focuses on describing models of each of these processes; description of their coupling in the overall model is provided in the companion paper. An implementation of the model in C language is freely available at https://github.com/MarcNeveu/ExoCcycleGeo. 126

# 2.1 Geodynamics and outgassing

Outgassing of a planet's interior to the atmosphere results from partial melting of 127 mantle material advected to shallow depths (Kite et al., 2009). Although localized vol-128 canism is a prominent form of outgassing, much of outgassing on a planet may occur from 129 more diffuse (e.g., hydrothermal) sources. Both sources are linked to internal convection, 130 melting, and subsequent conductive cooling of the crust. In this model, we do not dis-131 tinguish subaerial from subaqueous outgassing because we consider the atmosphere and 132 ocean as a single reservoir in which the atmosphere and ocean compositions are at equi-133 librium. 134

Our overall procedure for calculating outgassing fluxes is as follows. The thermal 135 evolution of a planet of a given size is computed by balancing accretional and radiogenic 136 heating with convective and conductive cooling (as a function of tectonic mode) to ob-137 tain temperature profiles as a function of depth at different time points in a planet's his-138 tory. Geodynamic model parameters and adopted values are described in Table 1. These 139 temperature profiles are used to determine the melt fraction as a function of depth for 140 rock of a specified composition, using the *alphaMELTS* command line implementation 141 (Smith & Asimow, 2005; R. N. Thompson et al., 2007; P. M. Antoshechkina & Asimow, 142 2010; P. M. Antoshechkina et al., 2010; P. Antoshechkina & Asimow, 2018) of the MELTS 143 software (Ghiorso & Sack, 1995; Asimow & Ghiorso, 1998). From melt fractions, using 144 the same simplifying assumption as Kite et al. (2009) that all melt reaches the surface 145 (or at least the near-surface crust), we obtain rates of melt extrusion (or intrusion). Fi-146 nally, we assume that the rate of carbon outgassing is directly proportional to that ex-147 trusion and intrusion rate, based on the assumed carbon content of the upper mantle ma-148 terial able to be outgassed. 149

150

### 2.1.1 Planet structure

The user inputs a planet mass, core mass fraction, and three materials for an inner core, outer core, and mantle. From these parameters, an interior structure is generated for the planet using the approach described in Lorenzo et al. (2014). The planet is divided in 1-dimensional grid zones, assuming spherical symmetry. Given a first-guess radius, the gravitational acceleration is computed as a function of depth. The equation of hydrostatic equilibrium is then integrated to find the pressure at depth, assuming con**Table 1.** Geodynamic model parameters. The lack of dependency for this study of the assumed values for the geodynamical parameters described in this section on a planet's bulk composition (Table 1) is addressed in Section 4. <sup>*a*</sup> User-specified inputs via planet mass, core mass fraction, and inner core, outer core, and mantle materials (Lorenzo et al., 2014).

Parameter	Symbol	Value	Units	Notes and references
Planet Structure				
Planet radius $^{a}$	$r_p$		m	
Core-mantle boundary radius $^{a}$	$r_c$		m	
Grid zone radius	r		m	
Gravitational acceleration	g		${\rm m~s^{-2}}$	
Density <sup>a</sup>	ρ		${\rm kg}~{\rm m}^{-3}$	Lorenzo et al. (2014)
Pressure	P		Pa	$= \int_r^{r_p} \rho(r) \ g(r) \ dr$
Temperature	T		К	
Accretion				
Fraction of impactor energy deposited at depth	h	0.1		Canup et al. (2021)
Ratio of planetesimal velocity to escape velocity	χ	2.0		Canup et al. (2021)
Thermal Evolution				
Radiogenic power per unit mass	H		${\rm W~kg^{-1}}$	Nuclide heating rate $\times$
				mantle abundance (Table 2)
Radionuclide half-life	$t_{1/2}$		s	See Table 2
Mantle heat capacity	$C_p$	1295	$\rm J~kg^{-1}~K^{-1}$	Akaogi and Ito (1993)
Mantle thermal conductivity	k	4.18	$\mathrm{W}~\mathrm{m}^{-1}~\mathrm{K}^{-1}$	Turcotte and Schubert (2002)
Mantle thermal diffusivity	$\kappa$		$\mathrm{m}^2~\mathrm{s}^{-1}$	$= k/(\rho \ C_p)$
Mid-mantle temperature	$T_m$		К	From equation $(4)$
Surface temperature	$T_s$		К	
Mantle thermal expansivity	α	$3 \times 10^{-5}$	$K^{-1}$	Turcotte and Schubert (2002)
Radius of brittle-ductile transition	$r_{BDT}$		m	
Temperature in mantle adiabat	$T_{adiab}$		К	From equation $(5)$
Temperature in lithosphere	$T_{upbnd}$		К	From equation $(6)$
Depth of lithosphere	$z_{Lith}$		m	Where $T_{adiab} = T_{upbnd}$
Rayleigh number	Ra		—	
Critical Rayleigh number	$Ra_{cr}$	1707.762	—	Reid and Harris (1958)
Nusselt number scaling parameter	$\beta$	1/3	—	$Nu = (Ra/Ra_{cr})^{\beta}$
Convective velocity	$v_{conv}$		${\rm m~s^{-1}}$	From equation (7)
Convective timescale	$t_{conv}$		s	$= (r_{BDT} - r_c)/v_{conv}$

157	tinuity across core and mantle boundaries. Equations of state for each of the three ma-
158	terials are inverted to derive the densities as a function of pressure. Based on these den-
159	sities, the thickness of the layers is adjusted. The gravitational acceleration as a func-
160	tion of depth is then updated, and so on until a convergence threshold is reached on the
161	central density. Several equations of state are considered depending on the material: a
162	third-order Birch-Murnaghan equation of state (Lorenzo et al., 2014), a power law (Seager
163	et al., 2007), and for high-pressure phases of ice, a temperature-dependent equation (Choukroun
164	& Grasset, 2010). A database of planetary materials and equation-of-state parameters
165	is provided with the code and includes data from Seager et al. (2007) for $\alpha$ -(bcc)Fe, MgSiO <sub>3</sub> ,
166	(Mg,Fe)SiO <sub>3</sub> , high-pressure $H_2O$ , C (graphite), and SiC; Dubrovinsky et al. (2000) for
167	$\epsilon\text{-}\mathrm{Fe};$ W. W. Anderson and Ahrens (1994) for liquid Fe; Fei et al. (1995) for FeS IV; Sata
168	et al. (2010) for FeS VII; Duffy, Hemley, and Mao (1995) for MgO; Weidner et al. (1982)
169	for SiO <sub>2</sub> ; Wett and Ahrens (1983) for $Mg_2SiO_4$ ; Chung (1971) for (Mg,Fe) <sub>2</sub> SiO <sub>4</sub> ; Duffy,
170	Meade, et al. (1995) for $Mg(OH)_2$ ; Tyburczy et al. (1991) for serpentine; Nestola et al.
171	(2010) for antigorite; Auzende et al. (2006) for lizardite; Fortes et al. (2013) for $Na_2SO_4 \cdot 10H_2O$ ;
172	Gromnitskaya et al. (2013) for MgSO <sub>4</sub> ·7H <sub>2</sub> O; Choukroun and Grasset (2010) for liquid
173	$H_2O$ and its ices Ih, II, III, V, and VI; Frank et al. (2013) for $H_2O$ ice VII; Fortes et al.
174	(2003) for $NH_3 \cdot 2H_2O$ I; Griffiths et al. (2012) for $NH_3 \cdot 2H_2O$ IIa; and Ma et al. (2012)
175	for $NH_3 \cdot 2H_2O$ liquid. A typical grid comprises $10^4$ zones. This is much finer than required
176	for this structure calculation, but necessary to resolve the upper mantle finely enough
177	as to accurately compute melting as a function of depth. The oversampling of deeper
178	layers seldom affects the speed of computation because the planetary structure is com-
179	puted only once at the onset of a simulation.

180

# 2.1.2 Thermal evolution

181 182 183

184

185

Heat sources include both radiogenic heat and heat from accretion, the two sources responsible for nearly all the heat trapped inside the Earth. A lower fraction of radiogenic heat implies a lesser dependence of mantle temperature on planet size and, especially, age (Kite et al., 2009). We factor accretional heating into the planet's initial temperature at mid-mantle depth,  $T_{m0}$ , following Canup et al. (2021):

$$T_{m0} = T_{surf0} + 0.6 \times 4 \ h \ \chi \ G \ \pi \ \rho \ r_p^2 / (3C_p) \tag{1}$$

**Table 2.** Radionuclide properties. Abundances pertain to the terrestrial planets of the Solar System and are given at the present day, 4.57 Gyr after the birth of the Solar System. The default abundances are those of Turcotte and Schubert (2002).

Property	$^{40}$ K	<sup>232</sup> Th	$^{235}$ U	$^{238}\mathrm{U}$	Reference	Notes
Half-life (Gyr)	1.26	14.0	0.704	4.47		
Heating rate ( $\mu$ W per kg nuclide)	) 29.2	26.4	569	94.6		
Abundance	18.0	51.9	0.10	14.2	Lyubetskaya and Korenaga (2007)	Low end-member
(ppb by mass)	36.9	124.0	0.22	30.8	Turcotte and Schubert (2002)	Highest heating
	30.7	84.1	0.15	21.0	Ringwood (1991)	Intermediate heating
	22.8	62.6	0.12	17.2	Lyubetskaya and Korenaga (2007)	Low heating
	28.8	79.5	0.15	20.15	McDonough and Sun (1995)	Intermediate heating

186 Here,  $T_{surf0}$  is an initial surface temperature, h is the fraction of impactor kinetic energy retained by the planet,  $\chi$  is the ratio of the impactor velocity relative to the escape 187 velocity from the accreting planet, G is the gravitational constant,  $r_p$  is the planet's fi-188 nal radius, and  $C_p$  is its bulk-averaged heat capacity, taken to be that of the mantle. Vol-189 ume averaging leads to the factor of 0.6. The parameters  $\chi$  and especially h are not well 190 constrained. For the Earth, simulations of the Moon-forming impact suggest that the 191 temperature of silicates reached 3000 to 6000 K (Canup, 2004), which necessitates  $h\chi \approx$ 192 0.2. We thus adopt this value for the  $h\chi$  product, which corresponds to 20% energy re-193 tained for average impactor velocities equivalent to the Earth's escape velocity of 11.2 194  $km s^{-1}$ , and noting that basing such a value on the Moon-forming impact may not be 195 representative of other Earths that did not experience such an impact. The higher ac-196 cretional heating experienced by larger (higher-R) planets tends to cause early vigorous 197 convection; the heat thus eliminated leads to these planets having counter-intuitively lower 198 lithospheric heat fluxes than an Earth-sized planet in the first 0.5 to 1 Gyr after accre-199 tion. Past 2 Gyr, the imprint of accretional heating on thermal evolution vanishes and 200 thermal evolution is dominated by the balance of radiogenic heating and convective or 201 conductive cooling, with larger planets having a hotter mantle and higher heat fluxes (see 202 Fig. 5c in Section 3). 203

Because radiogenic elements tend to partition with silicates rather than metals, heat 204 production is assumed to take place solely in the mantle. Radionuclide abundances de-205 pend on the bulk composition of the accreted material; we allow for user selection from 206 different sets of literature values (Table 2). Canonically, the abundances of Turcotte and 207 Schubert (2002) are assumed (Table 2); the lack of dependency of these values on a planet's 208 bulk composition is discussed in Section 4. Short-lived radionuclides are ignored since 209 the terrestrial planets have been thought to form after many short-lived radionuclide half-210 lives (Walsh & Levison, 2019), although recent models implying formation within just 211 a few million years may make short-lived radionuclides relevant early in the planet's his-212 tory (Bhatia & Sahijpal, 2016, 2017; Saito & Kuramoto, 2018; Johansen et al., 2021). 213 The radiogenic power per unit mass H (W kg<sup>-1</sup>) is expressed as: 214

$$H = \sum_{i=1}^{4} H_0(i) e^{-ln(2) t/t_{1/2}}$$
(2)

where i = 1 - 4 are the four main long-lived radioisotopes (<sup>40</sup>K, <sup>232</sup>Th, <sup>235</sup>U, <sup>238</sup>U),  $H_0$  is the product of a radionuclide's specific heating rate with its mass fraction in mantle material at the time the first planet-building solids condense – 4.57 billion years (Gyr) ago for the Solar System –,  $t_{1/2}$  is its half-life, and t is time.

The thermal profile from the surface to the base of the crust is calculated as follows. First, a mid-mantle temperature  $T_m$  is calculated at each time step with the assumptions of Kite et al. (2009) (e.g., whole-mantle convection), except thermal equilibrium is not assumed:

$$\frac{dT_m}{dt} = H/C_p - \frac{k}{\rho C_p} Nu \left(T_m - T_{ref}\right) \left(\frac{2}{r_{BDT} - r_c}\right)^2 \tag{3}$$

with the Nusselt number (dimensionless ratio of total heat flux to conductive heat flux) given by  $Nu = (Ra/Ra_{cr})^{\beta}$ . The Rayleigh number Ra is itself given by:

$$Ra = \frac{g \alpha (T_m - T_{ref}) (r_{BDT} - r_c)^3}{\kappa \nu}$$
(4)

Here, k is the thermal conductivity,  $T_{ref}$  is the surface temperature  $T_s$  in the plate tectonics regime, or the temperature at the base of the lithosphere (see equations 5 and 6) in the stagnant lid regime,  $r_c$  is the radius of the core-mantle boundary,  $r_{BDT}$  is the ra-
dius of the brittle-ductile transition (BDT), g is the gravitational acceleration,  $\alpha$  is the thermal expansivity,  $\kappa = k/(\rho C_p)$  is the thermal diffusivity, and  $\nu$  is the temperaturedependent kinematic viscosity.  $T_m$ , g, and  $\nu$  are evaluated halfway between  $r_c$  and  $r_{BDT}$ .

The radius of brittle-ductile transition  $r_{BDT}$  is that at which the temperature  $T_{BDT}$ and pressure  $P_{BDT}$  are such that the brittle and ductile strengths of mantle material are equal. It is found by using the previous time step's  $P_{BDT}$  and solving for  $T_{BDT}$  using a combined binary and Newton-Raphson search (Press et al., 1992). The brittle strength is given by 0.85*P* at low pressure (P < 200 MPa) and 0.6*P* + 50 MPa at P > 200MPa (Byerlee, 1978). The ductile strength is the ratio of the viscosity (see below) to the time step.

 $Ra_{cr}\approx 10^3$  is the critical Raleigh number. We adopt  $Ra_{cr}=1707.762$  from Reid 238 and Harris (1958). We also adopt  $\beta = 1/3$ , relevant to an isoviscous convective layer 239 (Solomatov, 1995); the viscosity variation with pressure and temperature across the man-240 tle may warrant values closer to 1/4 (Moresi & Solomatov, 1995; Deschamps & Sotin, 241 2000). Since that viscosity variation is not well constrained, especially for mantle com-242 positions that differ from Earth's, it may make sense to vary  $\beta$  between 1/4 and 1/3; but 243 we find that varying planet mass provides a sufficient range of thermal evolutions for the 244 purposes of this study. 245

The viscosity of Earth's mantle is assumed non-Newtonian, although this is uncertain (Deschamps & Sotin, 2000). The viscosity is a parallel combination of those obtained with the diffusion and dislocation creep flow laws for olivine given in Table 3 of Korenaga and Karato (2008), with stress equated to lithostatic pressure and the choice of dry (default) or wet mechanism left as a user input.

Based on the mid-mantle temperature  $T_m$ , a temperature profile is computed with 251 radius r from the surface to the core. This profile is assumed to have two parts: an adi-252 abat section in the asthenosphere, where heat transfer is convective, and a conductive 253 section in the lithosphere, the upper and only thermal boundary layer since the fluid is 254 radiogenically heated from within (Turcotte & Schubert, 2002). This boundary layer (Fischer 255 et al., 2010) is formally distinguished from the brittle layer  $(r > r_{BDT})$ , but in prac-256 tice the two have similar thickness. The temperature profile is computed as  $T(r) = min[T_{adiab}(r), T_{upbnd}(r)]$ , 257 with: 258

$$T_{adiab}(r) = T_m - \alpha \ g \ T_m [r - 0.5(r_{BDT} + r_c)]/C_p \tag{5}$$

$$T_{upbnd}(r) = T_m + (T_s - T_m) \ erfc\left[\frac{r_{BDT} - r}{2}\sqrt{\frac{2 \ v_{conv}}{\kappa(r_{BDT} - r_c)}}\right]$$
(6)

Equation 5 is equation 4.254 of Turcotte and Schubert (2002), evaluated at the mid-mantle depth halfway between  $r_c$  and  $r_{BDT}$ ; see also Katsura et al. (2010). Equation 6 is equation 6.347 of Turcotte and Schubert (2002), evaluated at the lateral midpoint between upwelling and downwelling regions of a convective cell (whose width and height are equal). In equation (6), erfc is the complementary error function, and the convective velocity  $v_{conv}$  is computed as in equation 6.379 of Turcotte and Schubert (2002):

$$v_{conv} = 0.354 \ \kappa \ Ra^{1/2} / (r_{BDT} - r_c) \tag{7}$$

which is appropriate for a fluid heated from within. A corresponding convection timescale is  $t_{conv} = (r_{BDT} - r_c)/v_{conv}$ .

The lithospheric thickness  $z_{Lith}$  is the depth at which  $T_{adiab} = T_{upbnd}$ . At the first time step, the temperature profile is computed using the same equations (5) and (6), initializing  $r_{BDT} = r_p$  and  $v_{conv} = (r_p - 100 \text{ km} - r_c)/10 \text{ Myr}$ .

The sensitivity of temperature profiles to mid-mantle depth and rheology for a 1- $M_{\oplus}$  planet is shown in Fig. 2. A hotter mantle results in a thinner lithosphere, and so does a wet olivine rheology, whose lower resulting viscosities result in more vigorous convection. For the adopted olivine rheologies, mantle temperatures of 2400-2500 K result in a profile that fits constraints for the Earth (Fischer et al., 2010).

The tectonic mode is determined by comparing the yield stress, equated to the brittle (and ductile) strength at the BDT, with the convective drive stress  $\nu/t_{conv}$ . If the drive stress exceeds the yield stress, plate tectonics proceed; otherwise the planet is assumed to have a stagnant lid. This very rough determination requires viscosities above  $\sim 10^{27}$  Pa s for plate tectonics to take place, which is not achieved in our simulations. The tectonic mode only has a minor effect on equation (4) but can significantly affect seafloor weathering, as investigated in the companion paper.



Figure 2. Validation of the melting and outgassing code with present-day Earth. Geotherms (temperature-depth profiles; top x-axis) for a  $1-M_{\oplus}$  planet are overlain on constraints for Earth's geotherm from analyses of mantle peridotites and measurements of surface heat flux (Turcotte & Schubert, 2002; Fischer et al., 2010). Also shown are melt fraction-depth profiles (bottom x-axis).

#### 2.1.3 Outgassing carbon flux

283

282

The outgassing flux, in mol  $s^{-1}$ , is assumed to be given by:

$$F_{outgas} = \frac{M_{melt} f_{C,magma}}{M_C t_{conv}}$$
(8)

where  $M_{melt}$  is the total mass of melt integrated over depth,  $f_{C,magma}$  is the fraction of the melt mass that exsolves as carbon gas, and  $M_C$  is the molar mass of CO<sub>2</sub>.

To obtain  $M_{melt}$ , a mantle melt fraction is computed as a function of depth using 286 the *alphaMELTS* command line implementation of the *MELTS* code (Ghiorso & Sack, 287 1995; P. M. Antoshechkina & Asimow, 2010; P. M. Antoshechkina et al., 2010; P. An-288 toshechkina & Asimow, 2018). A specified oxide composition (see Table 4 in Section 3) 289 is provided to *alphaMELTS*, along with settings instructing computation of chemical equi-290 librium along the above-determined temperature profile, from the surface to the depth 291 at which P = 10 GPa. The MELTS model is valid for P < 3 GPa, and alphaMELTS 292 allows (potentially unreliable) calculations beyond this limit up to 4 GPa. Calculations 293 show that there can still be partial melting at 4 GPa. In this case, our model implemen-294 tation extrapolates linearly the melt fraction trend over the 5 gridpoints closest to 4 GPa, 295 as determined with *alphaMELTS*, to higher pressures until the melt fraction is zero. Based 296 on comparison with an analytical melting model for Earth's upper mantle composition 297 (McKenzie & Bickle, 1988), for which the melt fraction decreases to zero with depth (pres-298 sure) faster than linearly, this may slightly overestimate the amount of melt at high pres-299 sure. However, these amounts tend to be small relative to  $M_{melt}$ . 300

For the temperature profiles of Fig. 2, the rates of crustal generation from all melt,  $M_{melt}/(4\pi r_p^2 \rho_{melt} t_{conv})$  with the melt density  $\rho_{melt}$  obtained from *alphaMELTS*, are 30.1 ( $T_m = 2500$  K), 11 ( $T_m = 2450$  K), 1.23 ( $T_m = 2400$  K), and 47 m Myr<sup>-1</sup> ( $T_m = 2400$  K, wet rheology) compared to Earth's estimated 40 m Myr<sup>-1</sup> (Kite et al., 2009). These illustrate a trade between best reproducing Earth's temperature profile with depth (profiles with thicker lithosphere) and best reproducing its rate of crust generation (profiles with thinner lithosphere).

On modern Earth, the magma  $CO_2$  content  $f_{C,magma}$  has been variously reported to be 0.03–1 mass% (Hekinian et al., 2000; Jones et al., 2018) in mid-ocean ridge basalts, 0.1–0.65 mass% in ocean island basalts that also possibly approximate Archean Earth

-14-

erupted material (Gerlach & Graeber, 1985; A. Anderson, 1995; Hartley et al., 2014),

and 0.004-0.15 mass% in back-arc volcanism (Cioni, 2000; Saito et al., 2001; Wade et al.,

<sup>313</sup> 2006; Blundy et al., 2010). Adopting a canonical abundance of 0.2 mass%, which is within

- those ranges, yields an outgassing flux similar to that of modern Earth for a 1 Earth-
- mass, 4.57 Gyr-old planet whose upper mantle has a mid-ocean ridge basalt composi-
- $_{316}$  tion (see Fig. 5 in Section 3).

Outgassed carbon can be speciated into CO<sub>2</sub> and CH<sub>4</sub>, with the relative fraction 317 of methane  $f_{CH4}$  left as a user input and constant through a simulation. We approxi-318 mate  $f_{CH4} = 0$  for modern and Archean Earth cases, and  $f_{CH4} \ge 0.5$  for more metal-319 rich planetary mantles. For context, the redox state of Hadean Earth magmas was near 320 the fayalite-magnetite-quartz (FMQ) redox buffer (Trail et al., 2011), but a higher pro-321 portion of submarine (i.e., high-pressure) volcanism would have affected the composi-322 tion of the erupted gases (Gaillard et al., 2011), with low (atmospheric)-pressure out-323 gassing likely to be uniformly near FMQ-2 over a range of body sizes, assuming a sim-324 ilar solar-system-like bulk basaltic composition for rock-metal bodies (Gaillard & Scail-325 let, 2014). In this case, the prime carbon gas erupted is CO<sub>2</sub> (Gaillard & Scaillet, 2014). 326 For undifferentiated or more metal-rich objects, at 1 bar, the predominant carbon gases 327 are CH<sub>4</sub> below 750 K and CO above 750 K (Schaefer & Fegley, 2017). 328

329

# 2.2 Continental weathering

330

## 2.2.1 Geochemical kinetics

Weathering describes the physicochemical process which modifies rocky surface landforms by its physical and chemical interaction with fluids having solubilized atmospheric material. As fluids are usually richer in carbon than rock, weathering reactions tend to result in transfer of carbon from the fluid to the rock. A relevant reaction is (Urey, 1952):

$$CaSiO_3 + CO_2(aq) = CaCO_3 + SiO_2$$
  
wollastonite calcite silica (9)

where (aq) indicates  $CO_2$  in its aqueous form. The fluid-rock setting can be subaerial crust exposed to rain or running water (continental weathering), or suboceanic crust in contact with ocean water (seafloor weathering). **Table 3.** Key model parameters for calculating carbon fluxes. Fluxes in mol m<sup>-2</sup> s<sup>-1</sup> can be obtained by dividing  $F_i$  by  $4\pi r_p^2$ . <sup>*a*</sup> User-specified inputs. <sup>*b*</sup> User-specified input via river runoff rate.

Parameter	Symbol	Value	Units	Notes and references
Outgassing				
Outgassing flux	$F_{outgas}$		$ m mol~s^{-1}$	From equation $(8)$
Mass of mantle melt integrated over depth	$M_{melt}$		kg	From <i>alphaMELTS</i>
Magma carbon fraction $^{a}$	$f_{C,magma}$	0.002	$\rm kg \ kg^{-1}$	
Relative fraction of C outgassed as CH4 $^a$	$f_{CH4}$		$\rm kg \ kg^{-1}$	0 for Modern and Archean Earth
				$\geq 0.5$ for chondritic mantle
Continental weathering				
Continental weathering flux	$F_{contw}$		$ m mol~s^{-1}$	From equation (11)
Rainfall rate $^{b}$	p	0.7	${\rm m~yr^{-1}}$	Broecker and Peng (1982)
Fresh- and seawater density	ho	1000	${\rm kg}~{\rm m}^{-3}$	
A real fraction of planet covered by land $^{a}$	L		_	0.29 for Earth
Freshwater:rock mass ratio	$4\pi r_p^2 \ L \ p \ \rho/M_{rock}$	5000	${\rm kg \ kg^{-1}}$	Martin and Meybeck (1979)
Molalities of dissolved species in rivers	$m_{i,river}$		$ m mol~kg^{-1}$	From <i>PHREEQC</i> kinetic calculation
Continental weathering				
Seafloor weathering flux	$F_{seafw}$		$ m mol~s^{-1}$	From equation $(13)$
Mass of surface ocean $a$	$M_{ocean}$	$1.4\times 10^{21}$	kg	
Hydrothermal circulation timescale	$t_{circ}$	$10^{7}$	$\mathbf{yr}$	Mottl (1983); Kadko et al. $(1995)$
Length of plate boundaries	$L_{plate\_ridges}$		m	$1.5\times 2\pi r_p$ for modern Earth
where seafloor is created				$(4.5 - 7.5) \times 2\pi r_p$ for Archean Earth
Depth of fracturing below seafloor	$z_{crack}$	6	$\rm km$	Vance et al. $(2007)$
Molalities of dissolved species in the ocean	$m_{i,ocean}$		$ m mol~kg^{-1}$	From $PHREEQC$ and eq. (14)
River water mass flowed to ocean in a time step	$M_{river}$		kg	
Carbon incorporated in the crust	$\Delta C_{reac}$		mol	From equation $(16)$
Atmospheric evolution				
Net carbon flux	$F_C$		$ m mol~s^{-1}$	From equation $(17)$
${\rm Atmosphere+ocean}$ reservoir	$RC_{atm+oc}$		mol	
Mantle reservoir	$RC_{mantle}$		$\operatorname{mol}$	
Time step $^{a}$	$\Delta t$		s	Adapted based on $RC_i$ and $F_C$
Surface pressure $^{a}$	$P_{surf}$		Pa	Varies during simulation
Surface temperature $^{a}$	$T_{surf}$		Κ	Varies during simulation
Number of moles of gas in atmosphere	$n_{air}$		mol	
Atmospheric mixing ratios $a$	$x_i$		-	$i = CO_2, CH_4, N_2, O_2, H_2O;$
				vary during simulation
Atmospheric volume	$V_{atm}$		$m^3$	Held constant

In planetary-scale models of the carbonate-silicate cycle, continental weathering fluxes are usually assumed to vary with temperature as an Arrhenius law, with runoff (the amount of rainwater involved in weathering reactions and that eventually transports dissolved or particulate surface rock to the ocean), and with atmospheric CO<sub>2</sub> partial pressure. The variation with the latter two parameters is usually expressed as power law deviations from measured Earth values, as in equations 8a, 2, and 1, respectively, of Sleep and Zahnle (2001),

In natural systems, crustal materials are composed of a variety of minerals that all 345 react with carbon-bearing fluids at different rates. To capture the effect of surface com-346 position on reaction rates, new power laws would have to be determined for every sur-347 face composition, an impractical challenge. Instead, we opt to compute reaction rates 348 and corresponding continental weathering fluxes from first principles, based on the re-349 sults of simulations of water-rock interaction through time using the KINETICS function-350 ality of the PHREEQC software (Parkhurst & Appelo, 2013). PHREEQC database and 351 example input files are provided as supplementary material to this article in a Zenodo 352 repository, and data and input compositions are described below. 353

Chemical reactions, including those involved in continental weathering, proceed at 354 a rate that depends on numerous factors such as temperature, the extent of chemical dis-355 equilibrium, concentrations of species such as H<sup>+</sup> (pH), mineral surface properties (Pokrovsky 356 & Schott, 1999), and the action of catalysts. For the latter two processes, a thorough 357 understanding is lacking for a comprehensive set of relevant rock species. The depen-358 dence on temperature and composition is better understood, but experimental data have 359 been compiled only for a subset of species that are abundant at Earth's surface (Palandri 360 & Kharaka, 2004). As a result, the calculation of reaction rates is typically captured into 361 a simplified form, fitted to experimental measurements, such as (Palandri & Kharaka, 362 2004): 363

$$\frac{dm}{dt} = -\text{SA } A \ e^{-\frac{E}{R(T-T_0)}} \ a_{H^+}^n \ (1-\Omega^p)^q \tag{10}$$

Here, m represents the concentration (molality, in moles of species per kg of water solvent) and dm/dt its change over time, SA the surface area available for reaction, E an activation energy, R the gas constant, T the temperature (which influences reaction rates via an Arrhenius law), and  $T_0$  a reference temperature, usually 298.15 K.  $\Omega$  is a mea-

sure of chemical disequilibrium, equal to Q/K, where  $Q = \prod_i a_i^{\nu_i}$  is the product of the 368 species' thermodynamic activities  $a_i$ , scaled geometrically by their stoichiometric coef-369 ficient  $\nu_i$  (negative for reactants, positive for products), and K(T, P) is this product when 370 the reaction reaches equilibrium (G. M. Anderson, 2005). K is termed the reaction con-371 stant, but depends on temperature T and pressure P. The magnitude of  $\Omega$  hinges both 372 on the bulk initial compositions of the fluid (e.g., carbon content) and rock, and on how 373 far toward equilibrium the reaction has progressed. The constant A and the exponents 374 n, p, and q are determined experimentally; here, p and q are both approximated to 1 for 375 all species (Palandri & Kharaka, 2004). 376

Reaction constants K are obtained from the *core10.dat* thermodynamic database 377 described in Neveu et al. (2017), with data from references therein, distributed with PHREEQC 378 at https://wwwbrr.cr.usgs.gov/projects/GWC\_coupled/phreeqc. For this study, we have 379 augmented the database with kinetic rates for 56 solids compiled by Palandri and Kharaka 380 (2004) and references therein. These include quartz and silica, as well as mineral species 381 in the feldspar, olivine, garnet and other orthosilicates, cyclosilicate, pyrox-382 enoid, amphibole, mica, clay and other phyllosilicates, oxide, hydroxide, carbonate, sul-383 fate, sulfide, and halide groups. 384

Not all solids need be part of the reacting crust, but all are allowed to form in a 385 simulation. One exception is quartz, included in weathering simulations involving mod-386 ern Earth and the solar system surfaces of Table 5, but not allowed to form in those in-387 volving the surface composition of Archean Earth. This is because quartz is not reac-388 tive on relevant timescales (residence time of rain and river water on land) and as such 389 tends to make the system of differential equations (10) too stiff for *PHREEQC*'s imple-390 mentation of the CVODE solver (Cohen et al., 1996) to handle, preventing these simula-391 tions from proceeding. 392

The starting continental crust (Table 5) and rainwater fluid compositions are provided as inputs to *PHREEQC*. The composition of rainwater, including its carbon content, is set to be equilibrated with the atmospheric composition, assuming that the atmosphere is an infinite reservoir. Atmospheric  $N_2$  and its dissolved form are assumed to be inert.

A fluid:rock mass ratio of about 5000 is canonically assumed, although the diversity of terrains on Earth points to much variation from this global mean. This ratio is

-18-

based on a reference river runoff rate for Earth of 0.665 mm day<sup>-1</sup> (Edson et al., 2012), 400 which corresponds to a rainfall rate  $p = 0.7 \text{ m yr}^{-1}$  minus evaporation of 65% of the rain-401 water from rivers (Broecker & Peng, 1982; Martin & Meybeck, 1979). Multiplying by 402 land surface area  $4\pi r_p^2 L$ , with L the fraction of planet area covered by land (L = 0.29403 for Earth), and fluid density  $\rho\approx 1000~{\rm kg}~{\rm m}^{-3}$  yields a rainfall mass rate  $4\pi r_p^2 \;L\;p\;\rho=$ 404  $1.0 \times 10^{17}$  kg yr<sup>-1</sup>. The mass of parent rock affected by weathering at the continent sur-405 face drained by rivers to the ocean is estimated to be  $M_{rock} = 21.1 \times 10^{12} \text{ kg yr}^{-1}$  in 406 Table V, footnote 3 of Martin and Meybeck (1979). This yields a fluid:rock ratio  $4\pi r_p^2 L p \rho/M_{rock}$ 407 = 4918. The ratio is proportional to p but insensitive to  $4\pi r_p^2 L$  since  $M_{rock}$  too is pro-408 portional to this term. In our modern and Archean Earth simulations, the fluid:rock ra-409 tios are 5000 and 4944, respectively. 410

2.2.2 Estimating continental weathering carbon fluxes

411

The *PHREEQC* simulation yields fluid and mineral abundances as a function of 412 reaction time, assuming a chemically closed system (no removal of reactants or products). 413 In the full model (Fig. 1), all solutes are delivered to the ocean where they accumulate 414 and, once saturation is reached, form carbon-bearing minerals that upon subduction trans-415 fer carbon from the {atmosphere+ocean} reservoir back to the mantle reservoir (Sec-416 tion 2.3). This approach provides a level of chemical realism not achieved with treatment 417 of distinct continental and seafloor weathering fluxes of carbon, but does not single out 418 the continental contribution to the weathering flux and its dependency on surface com-419 position. 420

For the purposes of this study, we therefore calculate a continental weathering flux  $F_{contw}$ , in mol yr<sup>-1</sup>, of dissolved carbon that can be sequestered in minerals based on amounts (molalities)  $m_{i,river}$  of Ca, Mg, and Fe cations in mol per kg of runoff water after a reaction time equal to the mean residence time of rainwater on land, about 10-15 years on modern-day Earth (Begemann & Libby, 1957; Chahine, 1992). The flux is then:

$$F_{contw} = \sum_{i} j m_{i,river} 4\pi r_p^2 L p \rho$$
(11)

Here, *i* is a summation index on all combinations of aqueous Ca, Mg, Fe, arising from
the dissolution of silicate, carbonates, sulfates, and sulfides present in the system. The

stoichiometric integer j is equal to the balance of positive and negative charges among 429 the ions (other than  $H^+$  and  $OH^-$ ) arising from the dissolution of these minerals: +1, 430 +2, 0, and -2 for carbonates, silicates, sulfates or pyrrhotite, and pyrite, respectively. 431 Indeed, their dissolution yields a doubly charged cation plus, respectively,  $HCO_3^-$ , aque-432 ous  $SiO_2$ ,  $SO_4^{2-}$ , or 2  $SO_4^{2-}$  at the river and ocean pH and oxidation conditions of mod-433 ern Earth (Berner et al., 1983). The contribution of Na and K cations is neglected be-434 cause their bicarbonate salts are too soluble to sequester carbon. Each excess positive 435 charge can combine with  $HCO_3^-$  or  $1/2 SO_4^{2-}$  from rainwater. Twice the rainwater  $SO_4^{2-}$ 436 flux is thus removed from the total excess positive charges to obtain  $F_{contw}$ . 437

While this approach to quantifying continental weathering flux provides a basis for determining its sensitivity to surface composition, it is inadequate if the atmosphere, surface, and ocean compositions lead to a predominant dissolved carbon species with a different charge than bicarbonate, such as dissolved  $CO_2$  (carbonic acid, no charge) or carbonate  $CO_3^{2-}$  (doubly charged anion) at ocean pH lower than 5 or greater than 10 at Earth surface T and P, respectively, or dissolved organic carbon or  $CH_4$  (no charge) at reducing conditions. Those cases warrant use of the full model of Fig. 1.

445

#### 2.2.3 Validation: modern Earth

We validate and calibrate PHREEQC kinetic simulations of continental weather-446 ing at 288 K and 1 bar with estimates of river abundances of major rock-forming ele-447 ments (Martin & Meybeck, 1979) and corresponding continental weathering fluxes  $F_{contw}$ 448 for modern Earth (Berner et al., 1983; Lerman & Wu, 2006; Colbourn et al., 2015) at 449 reaction (residence) times of 10-15 years (Begemann & Libby, 1957; Chahine, 1992). Abun-450 dances of major rock-forming elements in rivers are obtained by dividing simulated mo-451 lalities  $m_{i,river}$  by 1 - 0.65 = 0.35 to account for 65% evaporation on land (Broecker 452 & Peng, 1982). 453

A felsic crust composition (see 'Earth (modern)' column of Table 5 in Section 3) is set by adjusting starting abundances of quartz; Na-, Ca-, and K-feldspars; phyllosilicates (annite, phlogopite, daphnite); and hornblende (anthophyllite and tremolite) to match mineral volume fractions from Nesbitt and Markovics (1997) and oxide mass fractions from Rudnick and Gao (2003) within 10%. To this, we add calcite and dolomite (which rapidly and fully dissolve) to obtain carbonate contributions of Ca and Mg cations

-20-



Figure 3. Validation of continental weathering model against modern Earth average river compositions and weathering fluxes. **a.** PHREEQC simulated molalities  $m_{i,river}$  of dissolved Na, Mg, Si, Ca, and Fe (solid lines) at 288 K and 1 bar, divided by (1 (0.65) to account for 65% evaporation, compared to measured averaged river abundances (dashed lines) at relevant residence times of 10 to 15 years (red window). b. Corresponding river pH. c. Resulting total net flux (thick black line), obtained from the  $m_{i,river}$  shown in (a) using equation (11), and contributions to this flux from various mineral groups (subsets of the terms summed in equation 11), compared to literature ranges (shaded bands and lines inside the red box). The total flux curve is corrected to exclude about 2.4 Tmol  $yr^{-1}$  of cation fluxes, which are assumed to react with the corresponding flux of sulfate from rainwater and sulfide weathering generated by the model (Berner et al., 1983). Because this approach does not attribute the total sulfate flux contribution to specific mineral groups, the curves showing mineral group contributions include this sulfate flux contribution and therefore slightly overestimate the contribution of each group to the  $CO_2$  drawdown flux. Another 2.4 Tmol yr<sup>-1</sup> of sulfate arises from weathering of sulfate minerals, chiefly gypsum, but the sulfate and calcium arising from it are not shown here as together they do not affect the charge balance. <sup>a</sup> Martin and Meybeck (1979). <sup>b</sup> Berner et al. (1983). <sup>c</sup> Begemann and Libby (1957). <sup>d</sup> Chahine (1992). <sup>e</sup> Lerman and Wu (2006). <sup>f</sup> Colbourn et al. (2015).

in the ranges given by equations 10 and 17 of Berner et al. (1983) and by Table S1 of

461 Colbourn et al. (2015) and references therein. The added amounts correspond to 5.3 vol%

of exposed land, within the 4.2 - 8.8 area% range estimated for the past 500 Myr on

Earth,  $(6.2 - 13) \times 10^6$  km<sup>2</sup> (Kiessling et al., 2003), although part of this area is under-

- water on continental shelves. We also add sulfate (gypsum) and sulfide (pyrite) at abun-
- dances that match their estimated modern contributions to river sulfate; see p. 645 of
- 466 Berner et al. (1983).

The liquid composition is set to be equilibrated with partial pressures of 0.78 atm 467 of  $N_2$ , 0.2 atm of  $O_2$ , and 4000  $\mu$ atm of  $CO_2$ . The  $CO_2$  partial pressure, within the sev-468 eral thousand  $\mu$ atm range measured for large rivers (Cole & Caraco, 2001), reflects su-469 persaturation relative to the atmosphere (pre-industrial  $pCO_2$  of 280  $\mu$ atm) due to bi-470 ological respiration of biogenic reduced carbon from soil (Rasilo et al., 2017; Lynch et 471 al., 2010). The liquid also comprises 8  $\mu$ mol kg<sup>-1</sup> of sulfate, corresponding to the esti-472 mated  $0.8 \text{ Tmol yr}^{-1}$  pre-industrial sulfur contribution from outgassing and atmospheric 473 cycling (Berner et al., 1983). 474

<sup>475</sup> *PHREEQC* calculations with this input reproduce both a set of average cation con-<sup>476</sup> centrations in Earth rivers (Fig. 3a) and reported ranges of the weathering flux, includ-<sup>477</sup> ing contributions from various mineral groups (largely Ca- and Mg- silicates and carbon-<sup>478</sup> ates) (Fig. 3c). The computed river water sulfur abundance (constant with time) of  $6.9 \times$ <sup>479</sup>  $10^{-5}$  mol kg<sup>-1</sup> S, which corresponds to 6.6 mg kg<sup>-1</sup> SO<sub>4</sub><sup>2-</sup>, is in the reported range of <sup>480</sup> 5.19 - 11.2 mg kg<sup>-1</sup> SO<sub>4</sub><sup>2-</sup> of Lerman et al. (2007) and references therein.

- A few differences stand out. First, in the simulation nearly all Fe is in minerals. 481 Measured dissolved river Fe molalities may be non-negligible only because of (biogenic) 482 organic complexation (Gibbs, 1975), and can only be matched in a simulation if precip-483 itation of Fe oxides is prevented. Second, the simulated river pH past 10 years of inter-484 action is at the high end of typical measured river values (Fig. 3b). Third, the Ca-carbonate 485 (calcite) contribution to weathering flux is lower than estimated by Berner et al. (1983). 486 However, the Mg-carbonate contribution matches their estimate and the total carbon-487 ate contribution is in the middle of the range reported in Table S1 of Colbourn et al. (2015). 488
- To convert dissolved abundances to weathering fluxes we have assumed L = 0.29, irrespective of whether this land is drained by rivers to the oceans (i.e., exorheic). Exorheic areas have been variously estimated as comprising half to two thirds of the con-

-22-

tinental area on modern Earth (Lerman et al., 2007). Excluding Greenland and Antarctica, on which short-term weathering is impeded by the ice cover, but including endorheic areas in the ratio of surface water reservoirs to riverine fluxes (Chahine, 1992) results in L = 0.26, thus lowering fluxes of Fig. 3c by 10%.

In this calculation, we altered the dissolution kinetics of feldspar along the solid 496 solution between Ca (anorthite, fastest dissolving) and Na (albite, slowest dissolving) 497 end-members in order to limit the rate of Ca dissolution. Otherwise, the additional Ca 498 in solution increases the pH, limiting dissolved Mg. We found that attributing 85% of 499 anorthite the kinetic law for labradorite (50-70% anorthite, 30-50% albite) and the other 500 15% the law for bytownite (70-90% anorthite, 10-30% albite) achieved a suitable balance 501 between dissolved Ca and dissolved Mg as shown in Fig. 3c. We did not change the dis-502 solution rate for albite, assumed to be the end-member's law, but instead controlled the 503 amount of Na in solution with NaCl (see Table 5 in Section 3). 504

When applying this model to other surface and atmospheric compositions in lifeless conditions, we restore  $pCO_2$  to the atmospheric value, scale the 8  $\mu$ mol kg<sup>-1</sup> of dissolved atmospheric sulfur by the ratio of  $pCO_2$  to Earth's modern  $pCO_2$ , and retain the 85% labradorite – 15% bytownite kinetic rate law for anorthite. Implicit in the second choice is the simplifying assumption that atmospheric CO<sub>2</sub> and SO<sub>2</sub> co-vary; the model does not explicitly track atmospheric SO<sub>2</sub>.

511

## 2.3 Seafloor weathering

Seafloor weathering describes the loss of carbon from the {atmosphere+ocean} reservoir to the mantle reservoir. If the planet does not experience plate tectonics, dissolved species delivered by rivers continue to accumulate until saturation is reached and minerals precipitate. If plate tectonics operate, ocean fluids can react with fresh (unreacted) new ocean crust exposed at the seafloor.

In either case, it is sensible to assume that unlike continental weathering reactions, ocean-seafloor reactions proceed to chemical equilibrium. If ocean crust is recycled, the recycling timescale depends, to first order, on the vigor of mantle convection. The current rates of production of oceanic crust are about 20 km<sup>3</sup> yr<sup>-1</sup>  $\approx 2 \times 10^{6}$  kg s<sup>-1</sup> on Earth (Cogné & Humler, 2006), with the mantle convecting on a timescale of  $10^{8}-10^{9}$ years. This timescale holds for the larger, younger, or compositionally different planets

in the simulations whose outgassing fluxes are shown in Fig. 5. This is slower than cir-523 culation of ocean fluids through fractures or porosity in this crust, which occurs on a timescale 524  $t_{circ} \sim 10^7$  years (Mottl, 1983; Kadko et al., 1995). It is also slower than chemical equi-525 librium between these fluids and crustal rock: a *PHREEQC* kinetic simulation of reac-526 tion between a fluid with modern-day seawater composition from Parkhurst and Appelo 527 (2013) (simplified from Nordstrom et al. (1979)) and mafic crust composition (see Ta-528 ble 5, 'Earth (Archean)' column, in Section 3) shows that chemical equilibrium is reached 529 on timescales of  $10^5 - 10^6$  years (Fig. 4). This equilibrium is calculated at the ocean 530 surface temperature (a lower bound), 405 bar (depth of about 4 km), and a water:rock 531 mass ratio of 1 (see below). 532

The reaction takes place at seafloor pressure and surface temperature, in the approximation of an isothermal ocean. Water-rock reactions at or below the seafloor can, but need not, take place at elevated temperatures relative to that of the ocean. The water:rock ratio by volume is determined by:

$$\frac{V_{fluid}}{V_{ocean\ crust}} = \frac{(M_{ocean}/\rho) \times \Delta t/t_{circ}}{\frac{1-L}{1-0.29}\ L_{plate\_ridges\ v_{conv}\ z_{crack}\ \Delta t}}$$
(12)

This ratio is independent of the time step  $\Delta t$ . Here,  $L_{plate\_ridges}$  is the length of 537 plate boundaries where new seafloor is created (this excludes subduction zones). On mod-538 ern Earth, this length is about 1.5 times Earth's circumference. We scale it for differ-539 ent mantle convective vigor as  $L_{plate\_ridges} = (Ra/2.3 \times 10^6)^{\beta} \times 1.5 \times 2\pi r_p$ , which is 540 consistent with a linear proportion to the Nusselt number and with a 3 to 5 times greater 541 ridge length in the Archean (Kadko et al., 1995).  $v_{conv}$  is the plate velocity, which can 542 simply be equated to the mantle convective velocity (Turcotte & Schubert, 2002), or ar-543 bitrarily decreased from that value to simulate more sluggish tectonics as may have been 544 the case at the onset of plate tectonics (Brown et al., 2020).  $z_{crack}$  is the depth into the 545 seafloor to which fluid can circulate, with estimates for the present day varying between 546 600 m (Johnson & Pruis, 2003) and 6 km (Vance et al., 2007) (see also Hasenclever et 547 al. (2014)), with deeper circulation appearing possible if plate velocity is very low (Tao 548 et al., 2020). We adopt  $z_{crack} = 6$  km. 549

The seafloor weathering flux, in moles per unit time, is the difference  $\Delta C_{reac}$  in the molality of aqueous carbon  $m_{C,ocean}$  in the ocean pre- vs. post-reaction (mol kg<sup>-1</sup>), multiplied by the ocean mass  $M_{ocean}$  and divided by the time step:



Figure 4. Abundances of minerals and solutes as a function of time (lines) in a kinetic simulation at 288 K and 405 bar of the interaction between seafloor rock and seawater at a mass ratio of 1. These abundances, compared to those computed for an equilibrium simulation (thick dots), indicate that equilibrium is approached at about  $10^5$  years, although some Al- and S-rich minerals such as, respectively, clinochlore and pyrite have not yet reacted to diaspore and pyrrhotite.

$$F_{seafw} = \Delta C_{reac} \ M_{ocean} / \Delta t \tag{13}$$

The pre-reaction ocean molality  $m_{i,ocean}$  of each element is an average of the ocean composition at the previous time step  $(m_{i-1,ocean})$  and that of riverine input (molalities  $m_{i,river}$ ), scaled by the relative masses of the ocean and river runoff over a timestep:

$$m_{i,ocean,t} = \frac{m_{i,ocean,t-1} \ M_{ocean} + m_{i,river} \ M_{river}}{M_{ocean} + M_{river}}.$$
(14)

 $M_{ocean}$  is a user input (default  $1.4 \times 10^{21}$  kg for modern Earth) and is assumed constant with time. The mass of river water  $M_{river}$  flowing into the ocean during a timestep is given by:

$$M_{river} = 4\pi r_p^2 \ L \ \frac{p}{1 - 0.65} \ \rho \ \Delta t \tag{15}$$

<sup>559</sup> Only a subset  $\Delta t/t_{circ}$  of the ocean reacts with the seafloor; the resulting fluid is <sup>560</sup> then mixed back with the rest of the ocean to update the ocean composition. Follow-<sup>561</sup> ing this mixing step, mineral precipitation is allowed in case the ocean has become su-<sup>562</sup> persaturated; the resulting fluid composition is the ocean composition post-reaction. This <sup>563</sup> allows determination of  $\Delta C_{reac}$  as:

$$\Delta C_{reac} = m_{C,ocean,post-reaction} - m_{C,ocean,pre-reaction} \tag{16}$$

where  $m_{C,ocean}$  refers to the total molality of dissolved carbon, summed over all dissolved carbon species (including but not limited to CO<sub>2</sub>, CH<sub>4</sub>, HCO<sub>3</sub><sup>-</sup>, and CO<sub>3</sub><sup>2-</sup>).

566 River-ocean and hydrothermal fluid-ocean mixing simulations are carried out us-

<sup>567</sup> ing *PHREEQC*'s MIX functionality, which allows consistent tracking of pH and redox con-

ditions through the mixing reactions. The pressure of reaction is adjusted using the REACTION\_PRESSURE

keyword. Fluid-rock equilibria are calculated using the EQUILIBRIUM\_PHASES function-

- ality, either in the presence of seafloor rock (assumed, e.g., to have the mafic Archean
- crust composition given in Section 3's Table 5) if plate tectonics occur, or without re-
- active rock otherwise. In the latter case, transfer of carbon from the ocean to minerals
- occurs if the ocean composition following riverine input is such that carbon-bearing min-

erals are supersaturated, as expected if the ocean is already in equilibrium with a carbon-

<sup>575</sup> bearing atmosphere prior to riverine input.

For an ocean saturated in carbon both before the addition of riverine input and after seafloor interaction and precipitation of any supersaturated minerals,  $\Delta C_{reac}$  is proportional to  $\Delta t$ . Therefore,  $F_{seafw}$  is independent of  $\Delta t$ , as confirmed with simulations involving seafloor weathering with various time steps (see companion paper).

580

## 2.4 Net carbon flux and atmospheric evolution

The net flux  $F_C$  of carbon between the {atmosphere+ ocean} and mantle reservoirs is the sum of  $F_{outgas}$ , the outgassing flux given by equation (8), and  $F_{seafw}$  given by equation (13). The latter term accounts for contributions from continental weathering via the term  $\Delta C_{reac}$  (see equations 14 and 16), and as such represents the effects of both continental and seafloor weathering. If plate tectonics operate, this term is scaled by a fraction  $f_{arc}$  (default value 0.25 for Earth) that accounts for partial outgassing of subducted carbon via back-arc volcanism. Thus:

$$F_C = F_{outgas} + (1 - f_{arc}) F_{seafw}.$$
(17)

The {atmosphere+ocean} and mantle reservoirs of carbon, respectively  $RC_{atm+oc}$ and  $RC_{mantle}$  in mol, are respectively incremented or decremented at each timestep by  $F_C \Delta t$ . A maximum timestep is set as a user input (canonically, 1 Myr) and decreased if needed during a simulation to  $0.1 \times min(RC_{atm+oc}, RC_{mantle})/|F_C|$  in order to maintain numerical stability.

At the first time step,  $RC_{atm+oc}$  is initialized by specifying initial atmospheric par-593 tial pressures of  $CO_2$  and  $CH_4$  and using *PHREEQC* to determine the corresponding 594 (equilibrium)  $m_{C,ocean}$ . The initial  $RC_{mantle}$  is derived from the mantle mass (Section 595 2.1.1) and the canonical assumption that the mantle comprises 200 ppm C by mass. This 596 is in the inferred range of 20-1800 ppm (Dasgupta & Hirschmann, 2010) and implies de-597 pletion by a factor of 10 relative to the abundance of carbon in mantle melt of 0.2% (Sec-598 tion 2.1.3). Thus, there is canonically about  $10^3$  times more carbon in the mantle than 599 in the {atmosphere+ocean} reservoir (see Fig. 7 in Section 4). 600

The atmospheric surface pressure  $P_{surf}$  and mixing ratios  $x_{CO2} = pCO_2/P_{surf}$ and  $x_{CH4} = pCH_4/P_{surf}$  are updated at each time step based on  $F_C$  and  $f_{CH4}$ . This is done by equilibrating the atmospheric and ocean compositions using *PHREEQC* at surface temperature  $T_{surf}$  (fixed in this *PHREEQC* calculation) and pressure  $P_{surf}$  (modified by this *PHREEQC* calculation based on gas exchange with the ocean). Pre-equilibration conditions are set according to, respectively:

$$\Delta P_{surf} = F_C \ \Delta t \ R \ T_{surf} / V_{atm} \tag{18}$$

$$x_{CO2,t+1} = \frac{x_{CO2,t} \ n_{air} + (1 - f_{CH4}) F_C \Delta t}{n_{air} + F_C \Delta t}$$
(19)

$$x_{CH4,t+1} = \frac{x_{CH4,t} \ n_{air} + f_{CH4} F_C \Delta t}{n_{air} + F_C \Delta t} \tag{20}$$

where  $V_{atm}$  is the atmospheric volume and  $T_{surf}$  the surface temperature.

 $V_{atm}$  is held constant throughout the simulation, and computed at the first timestep as  $V_{atm} = n_{air} R T_{surf}/P_{surf}$ . The number of moles of air  $n_{air}$  is initiated as  $n_{air} =$  $P_{surf} \times 4\pi r_p^2/(g_{surf} M_{air})$ , with  $M_{air}$  the average atmospheric molar mass (calculated from the user-input atmospheric partial pressures of CO<sub>2</sub>, CH<sub>4</sub>, N<sub>2</sub>, O<sub>2</sub>, and H<sub>2</sub>O) and  $g_{surf}$  the surface gravity.  $n_{air}$  is incremented at each time step by  $F_C \Delta t$ .

The model is intended for coupling with models of atmospheric photochemistry and 613 radiative transfer (Kasting & Ackerman, 1986; Kopparapu et al., 2013; Arney et al., 2016; 614 Vidaurri et al., 2022). However, its implementation can be run independently, in which 615 case  $T_{surf}$  is computed by summing a term from insolation, which changes over geologic 616 timescales, and a term capturing the atmospheric greenhouse effect. Both terms are pa-617 rameterized as in Caldeira and Kasting (1992). This parameterization is valid for  $10^{-8}$ 618 bar  $< pCO_2 < 10^{-2}$  bar. For standalone simulations, we also roughly parameterize 619 the rainfall rate p as  $p \propto T_{surf}^{1.025}$  (with  $p = 0.7 \text{ m yr}^{-1}$  for  $T_{surf} = 288 \text{ K}$ ), reflecting 620 a roughly 2.5% increase in global mean precipitation on Earth per Kelvin of tempera-621 ture increase (Allen & Ingram, 2002; Trenberth et al., 2005; Pendergrass, 2020). An ex-622 ample such standalone simulation is provided in Section 4. 623

#### <sup>624</sup> 2.5 Neglected processes

The ocean mass is approximated as constant over time, even though on Earth it may have increased or decreased by a factor up to about 1.5 over geologic time (Harrison, 1999; Korenaga, 2008; Pope et al., 2012; Laneuville et al., 2018). We do not consider the potential sequestration of carbon species in ice caps or loss to space. Atmospheric speciation, photochemistry, and effects on surface temperature and runoff are not considered in this paper. This model does not differentiate between degassing of primordial vs. subducted carbon.

This model does not explicitly track outgassing chemistry as a function of source 632 depth, degassing transport, and outgassing pressure. While these aspects are thought 633 to impact the redox state of outgassed material, the extent to which they do is debated 634 (Burgisser & Scaillet, 2007; Dasgupta & Hirschmann, 2010). The  $fO_2$  of magmas and 635 associated abundances of gas species in equilibrium with the magma composition could 636 change up to 1.5-2 orders of magnitude upon ascent, due to degassing and pressure ef-637 fects on volatile exsolution (Burgisser & Scaillet, 2007). However, this variation could 638 go in either direction (toward a more oxidizing or reducing magma), depending on the 639 starting conditions, unless the volatiles are primarily H<sub>2</sub> and H<sub>2</sub>O, with little C and S 640 species. Moreover, the findings of Burgisser and Scaillet (2007) applied to Fe-poor (rhy-641 olitic) magmas, but more mafic magmas may provide added buffering capability, decreas-642 ing changes in magma redox during ascent. 643

#### 644 **3 Results**

In this section, we describe the sensitivity to planet size and composition of carbon fluxes arising from outgassing and weathering. The sensitivity of the carbon cycle and resulting atmospheric composition on these properties is investigated in the companion paper.

649

## 3.1 Geological controls on outgassing

The outgassing fluxes through time of rocky silicate planets with a surface water ocean and a surface temperature similar to Earth's are shown in Fig. 5a as a function of planet mass and upper mantle composition. Planet mass is varied between 0.5 and  $_{653}$  2  $M_{\oplus}$ , with the same planet-building materials and core mass fraction as in the modern Earth validation simulation shown in Fig. 2 and a dry olivine rheology.

Four upper mantle compositions are investigated: mid-ocean ridge basalt (Allan et al., 1989), chondritic (Sanloup et al., 1999), rhyolitic (Pamukcu et al., 2015), and felsic (Dolinschi, 2019) (Table 4). These oxide compositions were speciated into mineral compositions assuming an oxidation state corresponding to the FMQ buffer. Varying  $fO_2$ to several log units below this buffer did not significantly affect the propensity of these compositions to melt (or, therefore, the outgassing fluxes).

Simulations span the time interval between 0 and 5 Gyr after formation, but outgassing fluxes are computed only after 0.6 Gyr to allow for the initial geodynamic evolution to lead to a more steadily evolving mantle temperature and heat flux. The time step in these simulations is 10 Myr, which achieves numerical convergence since climate feedbacks are not tracked for this outgassing flux sensitivity analysis.

Outgassing fluxes span  $2 \times 10^4$  to  $6 \times 10^6$  mol s<sup>-1</sup>. They tend to be greater at 666 earlier times and for more felsic compositions. They decrease with increasing planet mass 667 if the mass is greater than  $1M_{\oplus}$ , but the mass at which outgassing fluxes peak is lower 668 for the two more felsic compositions:  $0.5 - 0.75 M_{\oplus}$  compared to  $1 M_{\oplus}$  for the basaltic 669 (Earth-like) upper mantle composition and  $1.5 M_\oplus$  for the ultramafic (chondritic) com-670 position. The outgassing fluxes for the  $1M_{\oplus}$  planets are usually within a few percent of 671 the peak values as a function of planet mass, irrespective of composition; whereas whether 672 the outgassing flux for 0.5-, 0.75-, or  $1.5-M_{\oplus}$  planets is close to this maximum depends 673 sensitively on composition. For example, for basaltic Earth-like or chondritic composi-674 tions, outgassing fluxes for 0.5- $M_{\oplus}$  planets are lower than those of 1- $M_{\oplus}$  planets by tens 675 of percent at any given time after formation. 676

The general decrease of outgassing fluxes with time is consistent with the global thermal evolution which tends toward lower mantle temperatures and heat fluxes as radiogenic heating decreases over time. The decrease spans up to two orders of magnitude between the first billion years after formation and the present day, but with considerable variability. It is less pronounced for higher-mass planets within the range investigated and for more felsic compositions. For 2  $M_{\oplus}$  planets, the flux first increases to peak at 1 to 2 Gyr after formation before decreasing to 5-Gyr values that are close to fluxes

Table 4. Simulated mantle compositions in mass% oxides: mid-ocean ridge basalt (MORB; Allan et al. (1989) as cited in Ghiorso and Sack (1995)), felsic (Dolinschi, 2019), rhyolitic (Pamukcu et al., 2015), and chondritic (Sanloup et al. (1999) as cited in Taylor (2013)). About 1 mass% or less of SiO<sub>2</sub> was either added or removed from the reported compositions in order for the total to reach 100%. These compositions are free of sub-percent mass fractions reported for Mn, Cr, and K oxides, as P. Antoshechkina and Asimow (2018) advised that including these as major oxides rather than trace elements in MELTS calculations can lead to unpredictable results. The rhyolitic composition's non-negligible K<sub>2</sub>O mass fraction (Pamukcu et al., 2015) was added as additional Na<sub>2</sub>O. Reported mass fractions of Al<sub>2</sub>O and Fe<sub>2</sub>O<sub>3</sub> were converted to mass fractions of Al<sub>2</sub>O<sub>3</sub> and FeO, respectively, because Al<sub>2</sub>O is not part of the MELTS chemical model and MELTS partitions Fe between Fe<sub>2</sub>O<sub>3</sub> and FeO during a calculation based on the chosen redox buffer (here, FMQ).

Oxide	MORB	Chondritic	Rhyolitic	Felsic
$SiO_2$	48.4	48.6	75.6	52.8
MgO	9.1	27.0	0.2	29.2
FeO	8.5	17.5	1.0	12.2
$Al_2O_3$	17.6	3.6	13.2	2.8
CaO	12.5	2.0	0.8	2.4
$\mathrm{TiO}_2$	1.0	0.1	0.2	-
Na <sub>2</sub> O	2.7	1.2	9.1	0.6
$P_2O_5$	0.1	-	-	-
$\rm H_2O$	0.2	-	-	-



Figure 5. a. Effect of planet mass and upper mantle composition on outgassing flux through time. Planet mass is varied between 0.5 and 2  $M_{\oplus}$  and shown as lighter to darker shade, respectively. Four upper mantle compositions are investigated: mid-ocean ridge basalt ("Earth-like", blue curves), chondritic (green curves), rhyolitic (yellow curves), and felsic (gray curves). The Earth symbol indicates the computed outgassing rate for modeled modern Earth conditions at which the model was validated (Fig. 2); the other three blue symbols indicate literature estimates of this value (Zhang & Zindler, 1993; Donnadieu et al., 2006; Burton et al., 2013). Gaps in curves or (especially for the rhyolitic composition) spuriously low values arise from lack of convergence of *MELTS* calculations at the full set of relevant shallow depths for those particular combinations of composition and mass- and age-dependent geotherm. **b.** Effect of planet mass (pressure-temperature profile) on the extent of melting of an upper mantle of basaltic (Earthlike) composition. **c.** Mid-mantle temperature through geologic time for planets of different masses, all with a basaltic upper mantle.

- computed at 0.6 Gyr. For felsic upper mantle compositions, the decrease in outgassing
  flux does not exceed 1 order of magnitude between 0.6 and 5 Gyr.
- At early times, mass has the largest effect on outgassing flux, but past 2 Gyr after formation, the outgassing fluxes become instead more strongly influenced by upper mantle composition. At 0.6 Gyr, the outgassing flux for  $\leq 1M_{\oplus}$  planets is greater than 10<sup>6</sup> mol s<sup>-1</sup> for all four upper mantle compositions simulated, whereas the outgassing flux for  $2M_{\oplus}$  planets is lower than 10<sup>6</sup> mol s<sup>-1</sup>. Instead, at 4 Gyr, the outgassing flux ranges corresponding to planet masses in the range  $0.5-2M_{\oplus}$  for each of the four compositions simulated no longer overlap: the flux ranges are about 8 - 12, 3 - 6, 0.8 -

 $_{693}$  2, and  $0.1 - 0.7 \times 10^5$  mol s<sup>-1</sup> respectively for the felsic, rhyolitic, basaltic, and chon-

<sup>694</sup> dritic mantle compositions.

The mass dependence of outgassing flux arises primarily from the influence of planet 695 mass on the geotherm (pressure-temperature profile) and its position relative to the solidus 696 for a given composition. For an Earth-like, basaltic upper mantle composition, planets 697 more massive than Earth have higher internal pressures and a shallower lithosphere-asthenosphere 698 boundary (Fig. 5b). This tends to limit the extent of mantle melting relative to a  $1-M_{\oplus}$ 699 planet. Planets less massive tend to be cooler because of their lower ratio of volume (ac-700 cretional and radiogenic heat production) to surface area (heat loss). This also limits the 701 extent of mantle melting (Fig. 5b). Thus, melting (and, proportionally, outgassing) is 702 generally maximized for 1- $M_{\oplus}$  planets. 703

The above analysis applies past 2 Gyr. At earlier times, outgassing is much more 704 suppressed on planets more massive than Earth owing to, counter-intuitively, relatively 705 low internal temperatures and heat fluxes (Fig. 5c). This is because convective heat trans-706 port is so efficient immediately after formation (due to the need to evacuate the high heat 707 of accretion) that mantle temperatures become uniform enough for the vigor of convec-708 tion and associated heat flux to drop below those of lower-mass planets. This results in 709 planets whose geotherm seldom crosses the solidus, and therefore in little melting and, 710 proportionally, outgassing. Later on, the heat flux increases back due to radiogenic heat-711 ing. 712

The compositional dependence of outgassing flux arises from the position of the upper mantle material's solidus curve relative to the geotherm of a planet of given mass and age. The more felsic compositions of Dolinschi (2019) and Pamukcu et al. (2015) have lower melting temperatures; therefore, the degree of melting for a given geotherm crossing their solidus is higher, leading to higher outgassing.

718

## 3.2 Geological controls on continental weathering

The effect of surface composition on continental weathering fluxes of rocky silicate planets is shown in Fig. 6a. The diverse surface compositions simulated are reflective of those of inner solar system planets (Table 5). These simulations were carried out assuming a 20% CO<sub>2</sub>, 80% N<sub>2</sub>, 1-bar atmosphere; a 288 K surface temperature; and modern-Earth rainfall rate of 0.7 m yr<sup>-1</sup> (water:rock ratio  $\approx$  5000). As in Fig. 3c, weathering

-33-



Figure 6. a. Continental weathering flux as a function of planet surface composition and residence time of rainwater on land. This flux is expressed as the equivalent cation capacity able to combine with bicarbonate  $(\text{HCO}_3^-)$  to form mineral precipitates. For each curve, a thumbnail of the planetary surface and labels "1" and "2" indicate the surface composition based on Table 5. The jaggedness in the curves for the Mars- and Ceres-like surface compositions is due to *PHREEQC* having difficulties converging on equilibria involving the precipitation of insoluble oxidized iron minerals. **b.** Dependence of weathering flux as a function of residence time on temperature for surface compositions approximating those of Modern and Archean Earth (Table 5).

fluxes are expressed as the equivalent cation capacity able to combine with bicarbonate ( $HCO_3^-$ ) to form mineral precipitates.

As shown in Fig. 6a, surface composition has a moderate effect on continental weathering fluxes, which are all on the order of tens of Tmol yr<sup>-1</sup>. Weathering fluxes are higher if the surface is made of minerals able to rapidly dissolve, such as sulfates (Venus-like surface) and carbonates (Ceres-like surface). The resulting enrichment of river water in cations likely acts as a negative feedback onto the dissolution of other minerals as  $\Omega$  in equation (10) increases toward 1 (i.e.,  $Q \to K$ ), preventing weathering fluxes from varying by more than about 1 order of magnitude.

Continental weathering fluxes increase with increased continental residence time, because the longer water-rock interaction leads to more mineral dissolution not fully compensated by precipitation of supersaturated minerals. For a given rainfall rate, weathering should thus be more effective on less patchy land.

Continental weathering fluxes also generally increase with temperature (Fig. 6b), 737 as expected from the Arrhenius (exponential) dependence of reaction kinetic rates on 738 the inverse of temperature. Despite this general trend, the relationship is more complex 739 and depends, e.g., on residence time of water on land. For the modern Earth case, if the 740 residence time exceeds 50 years (i.e., for continental masses more consolidated than to-741 day), the weathering flux is lower at higher temperatures because mineral formation re-742 actions that consume dissolve cations proceed faster owing both to faster kinetics and 743 to the faster accumulation of river cations (from prior mineral dissolution) to reach min-744 eral saturation. Thus, especially on less patchy land, weathering fluxes may instead be 745 controlled mainly by (proportional to) land area and rainfall. 746

Expressing weathering flux as the equivalent cation capacity able to combine with 747 bicarbonate  $(HCO_3^-)$  to form mineral precipitates requires making the implicit assump-748 tion that oxidized carbon is present as bicarbonate, i.e., that the ocean pH is roughly 749 between 6 (below which dissolved CO<sub>2</sub>, i.e., carbonic acid, dominates) and 10 (above which 750 doubly charged carbonate dominates). At pH > 10, the oxidized carbon cation trapping 751 capacity would be halved to conserve charge balance; at pH < 6, the trapping capacity 752 of cations would be zero since carbonic acid is neutral. In carbon cycling evolution sim-753 ulations that combine outgassing, weathering, and ocean-atmosphere equilibrium, the 754

-35-

ocean pH tends to be between 6 and 9 (see Fig. 7 in Section 4 and companion paper),

<sup>756</sup> making bicarbonate trapping capacity a reasonable means of evaluating weathering flux.

## 757 4 Discussion

Carbon is cycled between the atmosphere, surface, and interior of a planet by outgassing, weathering, and subduction. The balance between fluxes associated with these processes determines the amount of carbon accumulated in the atmosphere-surface and interior reservoirs. The above results enable us to understand how a planet's size and composition affects the intrinsic strength of these fluxes, i.e., of carbon cycle feedbacks. To do so, we have evaluated the magnitude of outgassing and continental weathering fluxes; for the latter, in the specific conditions of Earth's modern-day atmosphere.

765

766

# 4.1 Effect of planet size and composition on continental weathering and outgassing

Our simulations suggest that the efficiency of outgassing and subduction decreases 767 with mass for planets between 1 and 2 Earth masses. This contradicts the findings of 768 Kite et al. (2009) but is in line with more recent work accounting for the effects of high 769 pressure increasing mantle viscosity and thus decreasing its propensity to convect (Stamenković 770 et al., 2012). Here, planet mass is found to influence outgassing in other ways: at lower 771 masses, lower mantle temperatures decrease the degree of melting. At higher masses, higher 772 pressures decrease the degree of melting. We have assumed, as did Kite et al. (2009), that 773 all melt produced reaches the surface, but planet mass may also have an impact on the 774 degree to which intrusive volcanism does not lead to outgassing, proportionally decreas-775 ing outgassing compared to the values shown in Fig. 5a. 776

We have neglected the dependence on bulk composition of the thermophysical prop-777 erties and radionuclide abundances that affect a planet's geodynamic evolution (Tables 778 1 and 2). More matic compositions may lead to more vigorous convection and faster-cooling 779 planets that may outgas comparatively more early on and less at later ages than the flux 780 changes shown in Fig. 5a. The linkage between bulk compositions and radionuclide abun-781 dances is not straightforward. The internal distribution of radionuclides is unclear, be-782 cause elements Th and U tend to be incorporated as impurities in minerals (Oelkers et 783 al., 2009; Neveu et al., 2017) and because K can be exchanged between planet-building 784

-36-

materials such as silicates and volatile ices through aqueous alteration (Kirk & Steven-785 son, 1987; Engel et al., 1994; Castillo-Rogez & Lunine, 2010; Neveu et al., 2017). More-786 over, the starting abundances of radionuclides are difficult to predict from first princi-787 ples. The dominant heat source is <sup>40</sup>K, but Earth is rather depleted (relative to chon-788 drites) in moderately volatile elements, including K, by unknown mechanisms (Desch et 789 al., 2020; Unterborn et al., 2022). The solar system is probably not unusual in its abun-790 dances of short-lived radionuclides such as <sup>26</sup>Al, but these would be even more variable 791 (Desch et al., 2022). Thus, radiogenic heating may best be left as a freely varied param-792 eter, with the simulations shown here providing a solar system-centric baseline. 793

More mafic compositions tend to have a higher solidus temperature for a given pressure, and therefore are less prone to generating melt (i.e., outgassing). Although this was not tracked in our simulations, more mafic (reducing) compositions are generally expected to result in more reducing outgassing. The detailed carbon gas speciation into relative proportions of  $CO_2$  and  $CH_4$  depends on effects such as redox changes during ascent (Burgisser & Scaillet, 2007) and planet surface pressure (gravity and atmospheric density) (Gaillard & Scaillet, 2014), which are not considered here (Section 2.5).

Surface gravity is not expected to affect the continental weathering flux via the runoff rate to which this flux is proportional. Runoff rate is in turn proportional to river flow velocity, which depends on surface gravity as v = 2gh. Since declivity (a planet's relief) h is inversely proportional to surface gravity, the effects of gravity cancel out.

A planet's surface composition does affect the continental weathering flux to an ex-805 tent comparable to the effect of residence time of rainwater on land (Fig. 6a). However, 806 the effect of composition is much more pronounced on outgassing, for which it induces 807 order-of-magnitude differences (Fig. 5a). Time-evolution simulations in which the at-808 mospheric composition varies as a function of outgassing and (continental+seafloor) weath-809 ering show that in practice, the combined weathering flux varies to match the outgassing 810 forcing so as to keep the sum of atmospheric and ocean carbon roughly constant within 811 a factor of a few. An example such simulation, in which it is assumed that seafloor weath-812 ering proceeds as described in Section 2.3, is shown in Fig. 7. A broader set of such sim-813 ulations for different planet sizes and compositions is presented in the companion pa-814 per. 815

-37-



Figure 7. Time evolution of a. carbon reservoirs, b. carbon fluxes, c. surface temperature and pressure, d. atmospheric composition, e. pH, and f. ocean composition on a  $1-M_{\oplus}$  planet with an upper mantle composition comparable to that of mid-ocean ridge basalts (Table 4) and a surface composition comparable to that of Archean Earth (Table 5). The combined (continental + seafloor) weathering flux varies to match the outgassing forcing, maintaining roughly constant amounts of carbon in the {atmosphere+ocean} reservoir. Changes in the partitioning of carbon between the atmosphere and ocean arise due to secular changes in ocean chemistry (in this case, increasing pH). The ocean composition is initiated to the present-day composition, and the atmospheric composition is initiated to 0.5 bar total pressure of 80% N<sub>2</sub>-20% CO<sub>2</sub>.

Overall, planet mass and composition are expected to affect the carbon cycle as 816 follows, assuming the planet receives insolation similar to Earth's. First, the magnitude 817 of injection rates (fluxes) of carbon into the {atmosphere+ocean} reservoir varies by or-818 ders of magnitude (Fig. 5a) and is expected to generally decrease with geologic time. The 819 added carbon, dissolved in rainwater and in the ocean, reacts with land and seafloor rock. 820 The former yields an influx of carbon and cations into the ocean at a rate that is not very 821 sensitive to surface composition or residence time (continental patchiness) (Fig. 6) and 822 insensitive to planet mass (surface gravity), but which is proportional to rainfall rate (it-823 self weakly increasing with temperature via the atmospheric  $H_2O$  content; Section 2.4) 824 and land area. Continental patchiness affects evaporation, i.e., water:rock ratio, but sim-825 ulations with water:rock ratios varied by a factor of a few do not yield major differences 826 in weathering fluxes, especially since the effects of water:rock ratio and residence time 827 offset each other (reactions with more concentrated reactants proceed faster). The finite 828 solubility of carbon minerals drives precipitation and incorporation of carbon into the 829 seafloor at rates that should match the rate of outgassing (Fig. 7). Variations in the con-830 tinental weathering input of cations should drive changes in ocean composition on ge-831 ological timescales that may affect the partitioning of carbon between the ocean and at-832 mosphere due to the dependence of carbon speciation on, e.g., pH. Detailed investiga-833 tion of these effects is the topic of the companion paper. 834

835

#### 4.2 The Earth through time

While Earth's mean surface temperature has likely remained relatively constant at a few degrees above freezing throughout much of its evolution due to the carbonatesilicate cycle (Sleep & Zahnle, 2001), its land composition has become more felsic, its land less patchy, and its land coverage has increased (Korenaga, 2018). Its atmosphere has become less carbon-rich, with  $pCO_2$  changing from about 0.1–1 bar to a few  $10^{-4}$ bar (Catling & Zahnle, 2020).

This change in surface composition has acted to decrease continental weathering fluxes (Fig. 8) due to the lesser disequilibrium with Earth's atmosphere. Had Earth's composition remained the same, the continental weathering flux today would be closer to 100 Tmol yr<sup>-1</sup> of carbon than the estimated 20–25 Tmol yr<sup>-1</sup> (see Fig. 3c). However, the higher patchiness and lower areal fraction of land on Archean Earth implies a lower flux: about  $55 \times \frac{L}{0.29}$  Tmol/year for a residence time of 0.1 rather than 10 years.



Figure 8. Effect of land coverage and patchiness on continental weathering flux for a  $1-M_{\oplus}$ planet. Here, we have assumed a primitive mantle composition for the Earth's crust, which may be adequate for the Archean, and an Archean-like atmospheric composition of 0.5 bar N<sub>2</sub> and 0.5 bar CO<sub>2</sub>. The inset near 10 years residence time shows the same fluxes for modern Earth, as reproduced from Fig. 3.

This matches the modern-day weathering flux for a land fraction of about 15%, implying that the continental weathering flux need not have changed significantly over time.

The outgassing flux was likely an order of magnitude higher than today (Fig. 5a; 850 see also Sleep and Zahnle (2001) and Krissansen-Totton, Arney, and Catling (2018)), im-851 plying a greater forcing that decreased with time. This flux was balanced by a greater 852 seafloor weathering flux arising from a combination of higher convective velocity (Laneuville 853 et al., 2018) and 3-5 times greater ridge length (Kadko et al., 1995) in equation (12) (lower 854 water:rock ratio), even though, given the above, the accumulation rate of reactants from 855 river delivery to the ocean was roughly the same as today. The gradual increase with 856 time in the water:rock ratio of seafloor reactions, decrease in outgassing forcing, and neg-857 ative feedback of the cycle likely all acted to maintain carbon levels in the atmosphere 858 until carbon became significantly sequestered in biomass. 859

## 5 Conclusions

We have introduced a new, open-access model of geological carbon fluxes, whose implementation leverages existing, widely used geochemical codes of solid-melt equilib-

-40-

ria for silicate rocks (*MELTS*) and of equilibria and kinetics of water-rock interactions (*PHREEQC*). Coupled with a simple numerical computation of global thermal evolution, this model enables investigation of the effects of planet size (mass) and composition on carbon cycling through geologic time. Its applicable size range (0.5 to 2 Earth masses) is limited by the fidelity of the geodynamic model; the applicable range of composition is limited by those that can be handled by *MELTS* and *PHREEQC*.

We find that these planetary properties can influence outgassing fluxes of carbon 869 by as much as two orders of magnitude, with 0.5–1- $M_{\oplus}$  planets with felsic upper man-870 tles outgassing the most. In contrast, continental weathering fluxes are independent of 871 planet size and relatively insensitive to surface composition in terms of disequilibrium 872 with an Earth-like atmosphere; the variation in fluxes is at most a factor of a few. Con-873 tinental patchiness (residence time of rainwater) has an effect of similar magnitude, with 874 greater residence times generally leading to higher fluxes by allowing kinetically limited 875 reactions to proceed further. Surface composition and land patchiness thus likely have 876 lesser effects on continental weathering than the fraction of a planet's surface covered 877 by land. These continental weathering fluxes likely affect the rate of change of oceanic 878 chemical composition, which governs the partitioning of carbon between ocean and at-879 mosphere. The resulting evolutions of atmosphere and ocean compositions on Earth-like 880 planets of various sizes and solid compositions are the focus of a companion paper. 881

- 882 Open Research Section
- The data supporting this paper's conclusions can be obtained by running the ExoCcycleGeo code (version 22.4) with the crust and upper mantle composition templates freely available from https://github.com/MarcNeveu/ExoCcycleGeo. The software dependencies can be downloaded at the following webpages:

887

888

*PHREEQC*: https://wwwbrr.cr.usgs.gov/projects/GWC\_coupled/phreeqc;

- alphaMELTS: https://magmasource.caltech.edu/alphamelts; MELTS itself is available available
- able at https://melts.ofm-research.org.
- The data underlying Fig. 3 to 8 are archived in a Zenodo repository at https://zenodo.org/deposit/7640681 (size: 377.5 MBytes). The subset of data underlying Fig. 3, 4, 6, and 8 were obtained by running *PHREEQC* alone. Those underlying Fig. 2 were obtained by hard-coding a value for the mid-mantle temperature in *ExoCcycleGeo* and running a single time step.

-41-

### 894 Acknowledgments

- This work was funded by NASA Exoplanet Research Program award NNX17AB95G. We
- thank the reviewers of the proposal that led to this award for providing valuable feed-

<sup>897</sup> back and guidance that helped improve this research.

#### 898 References

- Abbot, D. S., Cowan, N. B., & Ciesla, F. J. (2012). Indication of insensitivity of
   planetary weathering behavior and habitable zone to surface land fraction. *The Astrophysical Journal*, 756(2), 178. doi: 10.1088/0004-637X/756/2/178
- Akaogi, M., & Ito, E. (1993). Heat capacity of MgSiO<sub>3</sub> perovskite. *Geophysical Research Letters*, 20(2), 105–108. doi: 10.1029/92GL02655
- Akeson, R., Chen, X., Ciardi, D., Crane, M., Good, J., Harbut, M., ... Zhang, A.
   (2013). The NASA Exoplanet Archive: data and tools for exoplanet research.
   *Publications of the Astronomical Society of the Pacific*, 125(930), 989. doi:
   10.1086/672273
- Allan, J. F., Batiza, R., Perfit, M. R., Fornari, D. J., & Sack, R. O. (1989). Petrology of lavas from the lamont seamount chain and adjacent East Pacific Rise,
  10 N. Journal of Petrology, 30(5), 1245–1298. doi: 10.1093/petrology/ 30.5.1245
- Allen, M. R., & Ingram, W. J. (2002). Constraints on future changes in climate and
  the hydrologic cycle. *Nature*, 419(6903), 228–232.
- Anderson, A. (1995). CO<sub>2</sub> and the eruptibility of picrite and komatiite. *Lithos*, 34(1-3), 19–25. doi: 10.1016/0024-4937(95)90005-5
- Anderson, G. M. (2005). Thermodynamics of Natural Systems. Cambridge Univer sity Press.
- Anderson, W. W., & Ahrens, T. J. (1994). An equation of state for liquid iron
   and implications for the Earth's core. Journal of Geophysical Research: Solid
   Earth, 99(B3), 4273–4284. doi: 10.1029/93JB03158
- Antoshechkina, P., & Asimow, P. (2018). Alphamelts software manual a text
   driven interface for MELTS, pMELTS & pHMELTS (Tech. Rep.). California
   Institute of Technology.
- Antoshechkina, P. M., & Asimow, P. D. (2010). Adiabat\_1ph 3.0 and the MAGMA website: educational and research tools for studying the petrology and geo-

926	chemistry of plate margins. In AGU Fall Meeting Abstracts (Vol. 2010, pp.
927	ED41B-0644).
928	Antoshechkina, P. M., Asimow, P. D., Hauri, E. H., & Luffi, P. I. (2010). Effect of
929	water on mantle melting and magma differentiation, as modeled using Adia-
930	bat_1ph 3.0. In AGU Fall Meeting Abstracts (Vol. 2010, pp. V53C–2264).
931	Arney, G. N., Domagal-Goldman, S. D., Meadows, V. S., Wolf, E. T., Schwieterman,
932	E., Charnay, B., Trainer, M. G. (2016). The pale orange dot: the spectrum
933	and habitability of hazy Archean Earth. Astrobiology, $16(11)$ , 873–899. doi:
934	10.1089/ast.2015.1422
935	Arney, G. N., Meadows, V. S., Domagal-Goldman, S. D., Deming, D., Robinson,
936	T. D., Tovar, G., Schwieterman, E. (2017). Pale orange dots: the impact of
937	organic haze on the habitability and detectability of Earthlike exoplanets. $\ The$
938	Astrophysical Journal, 836(1), 49. doi: 10.3847/1538-4357/836/1/49
939	Asimow, P. D., & Ghiorso, M. S. (1998). Algorithmic modifications extending
940	MELTS to calculate subsolidus phase relations. American Mineralogist, $83(9-$
941	10), 1127–1132. doi: 10.2138/am-1998-9-1022
942	Auzende, AL., Pellenq, RM., Devouard, B., Baronnet, A., & Grauby, O. (2006).
943	Atomistic calculations of structural and elastic properties of serpentine miner-
944	als: the case of lizardite. Physics and Chemistry of Minerals, 33(4), 266–275.
945	doi: 10.1007/s00269-006-0078-x
946	Barnes, R., Luger, R., Deitrick, R., Driscoll, P., Quinn, T. R., Fleming, D. P.,
947	Armstrong, J. (2020). VPLanet: the virtual planet simulator. <i>Publica</i> -
948	tions of the Astronomical Society of the Pacific, $132(1008)$ , $024502$ . doi:
949	10.1088/1538-3873/ab3ce8
950	Barnes, R., Meadows, V. S., & Evans, N. (2015). Comparative habitability of tran-
951	siting exoplanets. The Astrophysical Journal, $814(2)$ , 91. doi: 10.1088/0004
952	-637X/814/2/91
953	Barnes, R., Mullins, K., Goldblatt, C., Meadows, V. S., Kasting, J. F., & Heller,
954	R. $(2013)$ . Tidal Venuses: triggering a climate catastrophe via tidal heating.
955	Astrobiology, $13(3)$ , 225–250. doi: 10.1089/ast.2012.0851
956	Begemann, F., & Libby, W. F. (1957). Continental water balance, ground water
957	inventory and storage times, surface ocean mixing rates and world-wide wa-
958	ter circulation patterns from cosmic-ray and bomb tritium. Geochimica et

959	Cosmochimica Acta, $12(4)$ , 277–296. doi: $10.1016/0016-7037(57)90040-6$
960	Berner, R. A., Lasaga, A. C., & Garrels, R. M. (1983). The carbonate-silicate
961	geochemical cycle and its effect on atmospheric carbon dioxide over the
962	past 100 million years. American Journal of Science, 283, 641–683. doi:
963	10.2475/ajs.283.7.641
964	Bhatia, G. K., & Sahijpal, S. (2016). The early thermal evolution of Mars. Meteorit-
965	ics and Planetary Science, $51(1)$ , 138-154. doi: 10.1111/maps.12573
966	Bhatia, G. K., & Sahijpal, S. $(2017)$ . Did $^{26}$ Al and impact-induced heating differ-
967	entiate Mercury? Meteoritics and Planetary Science, 52(2), 295-319. doi: 10
968	.1111/maps.12789
969	Blackledge, B., Green, J., Barnes, R., & Way, M. J. (2020). Tides on other Earths:
970	Implications for exoplanet and palaeo-tidal simulations. Geophysical Research
971	$Letters,\ 47(12),\ e2019GL085746.\ doi:\ https://doi.org/10.1029/2019GL085746$
972	Blundy, J., Cashman, K. V., Rust, A., & Witham, F. $(2010)$ . A case for CO <sub>2</sub> -rich
973	arc magmas. Earth and Planetary Science Letters, 290(3-4), 289–301. doi: 10
974	.1016/j.epsl.2009.12.013
975	Bond, J. C., O'Brien, D. P., & Lauretta, D. S. (2010). The compositional diversity
976	of extrasolar terrestrial planets. I. In situ simulations. The Astrophysical Jour-
977	nal, 715(2), 1050. doi: 10.1088/0004-637X/715/2/1050
978	Broecker, W. S., & Peng, T. H. (1982). Tracers in the Sea. Lamont-Doherty Geolog-
979	ical Observatory, Palisades, NY.
980	Brown, M., Johnson, T., & Gardiner, N. J. (2020). Plate tectonics and the archean
981	earth. Annual Review of Earth and Planetary Sciences, 48, 291–320. doi: 10
982	.1146/annurev-earth-081619-052705
983	Burgisser, A., & Scaillet, B. (2007). Redox evolution of a degassing magma rising to
984	the surface. Nature, $445(7124)$ , 194–197. doi: 10.1038/nature05509
985	Burton, M. R., Sawyer, G. M., & Granieri, D. (2013). Deep carbon emissions from
986	volcanoes. Reviews in Mineralogy and Geochemistry, $75(1)$ , $323-354$ . doi: 10
987	$.2138/\mathrm{rmg}.2013.75.11$
988	Byerlee, J. (1978). Friction of rocks. In Rock Friction and Earthquake Prediction
989	(pp. 615–626). Springer.
990	Caldeira, K., & Kasting, J. F. (1992). The life span of the biosphere revisited. Na-
991	ture, $360(6406)$ , 721–723. doi: 10.1038/360721a0

992	Canup, R. M. (2004). Simulations of a late lunar-forming impact. <i>Icarus</i> , 168(2),
993	433–456. doi: 10.1016/j.icarus.2003.09.028
994	Canup, R. M., Kratter, K. M., & Neveu, M. (2021). On the origin of the Pluto sys-
995	tem. In The Pluto System after New Horizons (p. 475-506). University of Ari-
996	zona Press.
997	Castillo-Rogez, J. C., & Lunine, J. I. (2010). Evolution of Titan's rocky core con-
998	strained by Cassini observations. Geophysical Research Letters, $37(20)$ . doi: 10
999	.1029/2010GL044398
1000	Catling, D. C., Kiang, N. Y., Robinson, T. D., Rushby, A. J., Del Genio, A., et al.
1001	(2018). Exoplanet biosignatures: a framework for their assessment. Astrobiol-
1002	ogy. doi: 10.1089/ast.2017.1737
1003	Catling, D. C., & Zahnle, K. J. (2020). The Archean atmosphere. Science Advances,
1004	$\boldsymbol{6}(9),$ eaax 1420. doi: 10.1126/sciadv.aax 1420
1005	Chahine, M. T. (1992). The hydrological cycle and its influence on climate. Nature,
1006	359(6394), 373-380.doi: 10.1038/359373a0
1007	Choukroun, M., & Grasset, O. (2010). Thermodynamic data and modeling of
1008	the water and ammonia-water phase diagrams up to $2.2$ GPa for plane-
1009	tary geophysics. The Journal of Chemical Physics, 133(14), 144502. doi:
1010	10.1063/1.3487520
1011	Chung, D. (1971). Elasticity and equations of state of olivines in the $Mg_2SiO_4$ -
1012	$Fe_2SiO_4$ system. Geophysical Journal International, $25(5)$ , 511–538. doi: 10
1013	.1111/j.1365-246X.1971.tb02201.x
1014	Cioni, R. (2000). Volatile content and degassing processes in the AD 79 magma
1015	chamber at Vesuvius (Italy). Contributions to Mineralogy and Petrology,
1016	140(1), 40-54. doi: 10.1007/s004100000167
1017	Cogné, JP., & Humler, E. (2006). Trends and rhythms in global seafloor gen-
1018	eration rate. Geochemistry, Geophysics, Geosystems, $7(3)$ . doi: 10.1029/
1019	2005GC001148
1020	Cohen, S. D., Hindmarsh, A. C., & Dubois, P. F. (1996). CVODE, a stiff/nonstiff
1021	ODE solver in C. Computers in Physics, $10(2)$ , 138–143. doi: 10.1063/
1022	1.4822377
1023	Colbourn, G., Ridgwell, A., & Lenton, T. (2015). The time scale of the silicate
1024	weathering negative feedback on atmospheric CO <sub>2</sub> . Global Biogeochemical Cy-

1025	cles, 29(5), 583–596. doi: 10.1002/2014GB005054
1026	Cole, J. J., & Caraco, N. F. (2001). Carbon in catchments: connecting terrestrial
1027	carbon losses with aquatic metabolism. Marine and Freshwater Research,
1028	52(1), 101-110. doi: $10.1071/MF00084$
1029	Colose, C. M., Haqq-Misra, J., Wolf, E. T., Del Genio, A. D., Barnes, R., Way,
1030	M. J., & Ruedy, R. (2021). Effects of spin–orbit resonances and tidal heating
1031	on the inner edge of the habitable zone. The Astrophysical Journal, $921(1)$ ,
1032	25. doi: 10.3847/1538-4357/ac135c
1033	Dasgupta, R., & Hirschmann, M. M. (2010). The deep carbon cycle and melting in
1034	Earth's interior. Earth and Planetary Science Letters, 298(1-2), 1–13. doi: 10
1035	.1016/j.epsl.2010.06.039
1036	De Sanctis, M. C., Ammannito, E., Raponi, A., Marchi, S., McCord, T. B., Mc-
1037	Sween, H., Russell, C. T. (2015). Ammoniated phyllosilicates with a likely
1038	outer solar system origin on (1) Ceres. Nature, 528(7581), 241–244. doi:
1039	10.1038/nature16172
1040	Desch, S. J., Abbot, D., Krijt, S., Unterborn, C., Morard, G., & Hartnett, H. E.
1041	(2020). The volatile content of rocky planets. In <i>Planetary Diversity: Rocky</i>
1042	planet processes and their observational signatures (pp. $6.1$ – $6.40$ ). IOP.
1043	Desch, S. J., Young, E. D., Dunham, E. T., Fujimoto, Y., & Dunlap, D. R. (2022).
1044	Short-Lived Radionuclides in Meteorites and the Sun's Birth Environment.
1045	$arXiv\ e\text{-}prints.$ doi: 10.48550/arXiv.2203.11169
1046	Deschamps, F., & Sotin, C. (2000). Inversion of two-dimensional numerical
1047	convection experiments for a fluid with a strongly temperature-dependent
1048	viscosity. Geophysical Journal International, 143(1), 204–218. doi:
1049	10.1046/j.1365-246x.2000.00228.x
1050	Des Marais, D. J., Jakosky, B. M., & Hynek, B. M. (2007). Astrobiological impli-
1051	cations of Mars' surface composition and properties. In The Martian Surface-
1052	Composition, Mineralogy, and Physical Properties (pp. 599–624). Cambridge
1053	University Press.
1054	Dolinschi, J. D. (2019). Mineralogy of the silicon-rich mantle: Implications for Mars
1055	and exoplanets. <i>PhD dissertation</i> . Retrieved from https://keep.lib.asu
1056	.edu/_flysystem/fedora/c7/218100/Dolinschi_asu_0010N_19331.pdf
1057	Domagal-Goldman, S. D., Segura, A., Claire, M. W., Robinson, T. D., & Meadows,

-46-
1058	V. S. (2014). Abiotic ozone and oxygen in atmospheres similar to prebiotic
1059	Earth. The Astrophysical Journal, $792(2)$ , 90. doi: $10.1088/0004-637X/792/2/2/2$
1060	90
1061	Donnadieu, Y., Goddéris, Y., Pierrehumbert, R., Dromart, G., Fluteau, F., & Jacob,
1062	R. (2006). A GEOCLIM simulation of climatic and biogeochemical conse-
1063	quences of Pangea breakup. $Geochemistry, Geophysics, Geosystems, 7(11).$
1064	doi: 10.1029/2006GC001278
1065	Dressing, C. D., Spiegel, D. S., Scharf, C. A., Menou, K., & Raymond, S. N. (2010).
1066	Habitable climates: the influence of eccentricity. The Astrophysical Journal,
1067	721(2), 1295. doi: 10.1088/0004-637X/721/2/1295
1068	Dubrovinsky, L. S., Saxena, S. K., Tutti, F., Rekhi, S., & LeBehan, T. (2000).
1069	In situ X-ray study of thermal expansion and phase transition of iron
1070	at multimegabar pressure. Physical Review Letters, $84(8)$ , 1720. doi:
1071	10.1103/PhysRevLett.84.1720
1072	Duffy, T. S., Hemley, R. J., & Mao, HK. (1995). Equation of state and shear
1073	strength at multimegabar pressures: Magnesium oxide to 227 GPa. $Physical$
1074	Review Letters, 74(8), 1371. doi: 10.1103/PhysRevLett.74.1371
1075	Duffy, T. S., Meade, C., Fei, Y., Mao, HK., & Hemley, R. J. (1995). High-pressure
1076	phase transition in brucite, Mg (OH) <sub>2</sub> . American Mineralogist, $80(3-4)$ , 222–
1077	230. doi: 10.2138/am-1995-3-403
1078	Dyar, M. D., Helbert, J., Cooper, R. F., Sklute, E. C., Maturilli, A., Mueller, N. T.,
1079	Smrekar, S. E. (2021). Surface weathering on Venus: Constraints from
1080	kinetic, spectroscopic, and geochemical data. <i>Icarus</i> , 358, 114139. doi:
1081	10.1016/j.icarus.2020.114139
1082	Edson, A. R., Kasting, J. F., Pollard, D., Lee, S., & Bannon, P. R. (2012). The
1083	carbonate-silicate cycle and $\mathrm{CO}_2/\mathrm{climate}$ feedbacks on tidally locked terrestrial
1084	planets. Astrobiology, $12(6)$ , 562–571. doi: 10.1089/ast.2011.0762
1085	Engel, S., Lunine, J. I., & Norton, D. L. (1994). Silicate interactions with ammonia-
1086	water fluids on early Titan. Journal of Geophysical Research: Planets, $99(E2)$ ,
1087	3745–3752. doi: 10.1029/93JE03433
1088	Fei, Y., Prewitt, C. T., Mao, Hk., & Bertka, C. M. (1995). Structure and density of
1089	FeS at high pressure and high temperature and the internal structure of Mars.
1090	Science, $268(5219)$ , 1892–1894. doi: 10.1126/science.268.5219.1892

1091	Felton, R. C., Bastelberger, S. T., Mandt, K. E., Luspay-Kuti, A., Fauchez, T. J.,
1092	& Domagal-Goldman, S. D. (2022). The role of atmospheric exchange in
1093	false-positive biosignature detection. Journal of Geophysical Research: Planets,
1094	127(3), e2021JE006853. doi: 10.1029/2021JE006853
1095	Fischer, K. M., Ford, H. A., Abt, D. L., & Rychert, C. A. (2010). The lithosphere-
1096	asthenosphere boundary. Annual Review of Earth and Planetary Sciences, 38,
1097	551–575. doi: 10.1146/annurev-earth-040809-152438
1098	Foley, B. J. (2015). The role of plate tectonic–climate coupling and exposed land
1099	area in the development of habitable climates on rocky planets. The Astro-
1100	physical Journal, 812(1), 36. doi: 10.1088/0004-637X/812/1/36
1101	Forget, F., & Leconte, J. (2014). Possible climates on terrestrial exoplanets. <i>Phil.</i>
1102	Trans. R. Soc. A, 372(2014), 20130084. doi: 10.1098/rsta.2013.0084
1103	Fortes, A. D., Brand, H., Vočadlo, L., Lindsay-Scott, A., Fernandez-Alonso,
1104	F., & Wood, I. (2013). $P-V-T$ equation of state of synthetic mirabilite
1105	$(Na_2SO_4 \cdot 10D_2O)$ determined by powder neutron diffraction. Journal of Ap-
1106	plied Crystallography, $46(2)$ , 448–460. doi: 10.1107/S0021889813001362
1107	Fortes, A. D., Wood, I. G., Brodholt, J. P., & Vočadlo, L. (2003). The structure,
1108	ordering and equation of state of ammonia dihydrate ( $NH_3 \cdot 2H_2O$ ). Icarus,
1109	162(1), 59-73. doi: 10.1016/S0019-1035(02)00073-8
1110	Frank, M. R., Aarestad, E., Scott, H. P., & Prakapenka, V. B. (2013). A comparison
1111	of ice VII formed in the $H_2O$ , NaCl- $H_2O$ , and $CH_3OH-H_2O$ systems: Implica-
1112	tions for $H_2O$ -rich planets. Physics of the Earth and Planetary Interiors, 215,
1113	12–20. doi: 10.1016/j.pepi.2012.10.010
1114	Fujii, Y., Angerhausen, D., Deitrick, R., Domagal-Goldman, S., Grenfell, J. L., Hori,
1115	Y., Stevenson, K. B. (2018). Exoplanet biosignatures: observational
1116	prospects. Astrobiology, $18(6)$ , 739–778. doi: 10.1089/ast.2017.1733
1117	Gaillard, F., & Scaillet, B. (2014). A theoretical framework for volcanic degassing
1118	chemistry in a comparative planetology perspective and implications for plan-
1119	etary atmospheres. Earth and Planetary Science Letters, 403, 307–316. doi:
1120	10.1016/j.epsl.2014.07.009
1121	Gaillard, F., Scaillet, B., & Arndt, N. T. (2011). Atmospheric oxygenation caused by
1122	a change in volcanic degassing pressure. Nature, $478(7368),229.$ doi: 10.1038/
1123	nature10460

1124	Gerlach, T. M., & Graeber, E. J. (1985). Volatile budget of Kilauea volcano. Nature,
1125	313(6000), 273-277.doi: 10.1038/313273a0
1126	Ghiorso, M. S., & Sack, R. O. (1995). Chemical mass transfer in magmatic pro-
1127	cesses IV. A revised and internally consistent thermodynamic model for the
1128	interpolation and extrapolation of liquid-solid equilibria in magmatic systems
1129	at elevated temperatures and pressures. Contributions to Mineralogy and
1130	Petrology, 119(2), 197-212.doi: 10.1007/BF00307281
1131	Gibbs, R. (1975). Assessing Potential Ocean Pollutants. National Academy of Sci-
1132	ences.
1133	Gillon, M., Triaud, A. H., Demory, BO., Jehin, E., Agol, E., Deck, K. M.,
1134	Queloz, D. (2017). Seven temperate terrestrial planets around the nearby
1135	ultracool dwarf star TRAPPIST-1. Nature, $542(7642)$ , $456-460$ . doi:
1136	10.1038/nature21360
1137	Grenfell, J. L. (2017). A review of exoplanetary biosignatures. Physics Reports, 713,
1138	1-17. doi: 10.1016/j.physrep.2017.08.003
1139	Griffiths, G. I., Fortes, A. D., Pickard, C. J., & Needs, R. (2012). Crystal structure
1140	of ammonia dihydrate II. The Journal of Chemical Physics, 136(17), 174512.
1141	doi: 10.1063/1.4707930
1142	Gromnitskaya, E. L., Yagafarov, O. F., Lyapin, A. G., Brazhkin, V. V., Wood, I. G.,
1143	Tucker, M. G., & Fortes, A. D. (2013). The high-pressure phase diagram
1144	of synthetic epsomite (MgSO <sub>4</sub> ·7H <sub>2</sub> O and MgSO <sub>4</sub> ·7D <sub>2</sub> O) from ultrasonic and
1145	neutron powder diffraction measurements. Physics and Chemistry of Minerals,
1146	40(3), 271-285. doi: 10.1007/s00269-013-0567-7
1147	Haqq-Misra, J. D., Domagal-Goldman, S. D., Kasting, P. J., & Kasting, J. F.
1148	(2008). A revised, hazy methane greenhouse for the Archean Earth. $Astro-$
1149	biology, 8(6), 1127-1137.doi: 10.1089/ast.2007.0197
1150	Harrison, C. $(1999)$ . Constraints on ocean volume change since the Archean. Geo-
1151	physical Research Letters, $26(13)$ , 1913–1916. doi: 10.1029/1999GL900425
1152	Hartley, M. E., Maclennan, J., Edmonds, M., & Thordarson, T. (2014). Reconstruct-
1153	ing the deep $CO_2$ degassing behaviour of large basaltic fissure eruptions. Earth
1154	and Planetary Science Letters, 393, 120–131. doi: 10.1016/j.epsl.2014.02.031
1155	Hasenclever, J., Theissen-Krah, S., Rüpke, L. H., Morgan, J. P., Iyer, K., Petersen,
1156	S., & Devey, C. W. (2014). Hybrid shallow on-axis and deep off-axis hy-

-49-

1157	drothermal circulation at fast-spreading ridges. Nature, 508(7497), 508–512.
1158	doi: 10.1038/nature13174
1159	Hekinian, R., Pineau, F., Shilobreeva, S., Bideau, D., Gracia, E., & Javoy, M.
1160	(2000). Deep sea explosive activity on the Mid-Atlantic Ridge near 34
1161	50' N: magma composition, vesicularity and volatile content. Journal
1162	of Volcanology and Geothermal Research, 98(1-4), 49–77. doi: 10.1016/
1163	S0377-0273(99)00190-0
1164	Hofmann, A. W. (1988). Chemical differentiation of the Earth: the relationship be-
1165	tween mantle, continental crust, and oceanic crust. Earth and Planetary Sci-
1166	ence Letters, $90(3)$ , 297–314. doi: 10.1016/0012-821X(88)90132-X
1167	Hu, R., Seager, S., & Bains, W. (2012). Photochemistry in terrestrial exoplanet at-
1168	mospheres. I. Photochemistry model and benchmark cases. The Astrophysical
1169	Journal, 761(2), 166. doi: 10.1088/0004-637X/761/2/166
1170	Johansen, A., Ronnet, T., Bizzarro, M., Schiller, M., Lambrechts, M., Nordlund,
1171	Å., & Lammer, H. (2021). A pebble accretion model for the formation of the
1172	terrestrial planets in the Solar System. Science Advances, $7(8)$ , eabc0444. doi:
1173	10.1126/sciadv.abc0444
1174	Johnson, H. P., & Pruis, M. J. (2003). Fluxes of fluid and heat from the oceanic
1175	crustal reservoir. Earth and Planetary Science Letters, 216(4), 565–574. doi:
1176	10.1016/S0012-821X(03)00545-4
1177	Jones, M., Soule, S., Gonnermann, H., Le Roux, V., & Clague, D. (2018). Magma
1178	ascent and lava flow emplacement rates during the 2011 Axial Seamount erup-
1179	tion based on $CO_2$ degassing. Earth and Planetary Science Letters, 494,
1180	32-41. doi: 10.1016/j.epsl.2018.04.044
1181	Kadko, D., Baross, J., & Alt, J. (1995). The magnitude and global implications of
1182	hydrothermal flux. Geophysical Monograph-American Geophysical Union, 91,
1183	446–466. doi: $10.1029/GM091p0446$
1184	Kane, S. R., Hill, M. L., Kasting, J. F., Kopparapu, R. K., Quintana, E. V., Barclay,
1185	T., Torres, G. (2016). A catalog of Kepler habitable zone exoplanet candi-
1186	dates. The Astrophysical Journal, $830(1)$ , 1. doi: $10.3847/0004-637X/830/1/$
1187	1
1188	Kaspi, Y., & Showman, A. P. (2015). Atmospheric dynamics of terrestrial exoplan-
1189	ets over a wide range of orbital and atmospheric parameters. The Astrophysical

Journal, 804 (1), 60. doi: 10.1088/0004-637X/804/1/60
 Kasting, J. F. (1988). Runaway and moist greenhouse atmospheres and the evolution of Earth and Venus. *Icarus*, 74 (3), 472–494. doi: 10.1016/0019-1035(88)
 90116-9

- Kasting, J. F., & Ackerman, T. P. (1986). Climatic consequences of very high carbon dioxide levels in the Earth's early atmosphere. *Science*, 234 (4782), 1383–
  1385. doi: 10.1126/science.11539665
- Katsura, T., Yoneda, A., Yamazaki, D., Yoshino, T., & Ito, E. (2010). Adiabatic
   temperature profile in the mantle. *Physics of the Earth and Planetary Interi-* ors, 183(1-2), 212–218. doi: 10.1016/j.pepi.2010.07.001
- <sup>1200</sup> Kiang, N. Y., Domagal-Goldman, S., Parenteau, M. N., Catling, D. C., Fujii, Y.,
- Meadows, V. S., ... Walker, S. I. (2018). Exoplanet biosignatures: at the dawn of a new era of planetary observations. *Astrobiology*, 18(6), 619–629. doi: 10.1089/ast.2018.1862
- Kiessling, W., Flügel, E., & Golonka, J. (2003). Patterns of Phanerozoic carbonate platform sedimentation. Lethaia, 36(3), 195–225. doi: 10.1080/
  00241160310004648
- Kirk, R. L., & Stevenson, D. J. (1987). Thermal evolution of a differentiated
   Ganymede and implications for surface features. *Icarus*, 69(1), 91–134. doi:
   10.1016/0019-1035(87)90009-1
- Kite, E. S., Manga, M., & Gaidos, E. (2009). Geodynamics and rate of volcanism on
   massive Earth-like planets. *The Astrophysical Journal*, 700(2), 1732. doi: 10
   .1088/0004-637X/700/2/1732
- <sup>1213</sup> Kopparapu, R. K., Ramirez, R., Kasting, J. F., Eymet, V., Robinson, T. D., Ma-
- hadevan, S., ... Deshpande, R. (2013). Habitable zones around mainsequence stars: New estimates. *The Astrophysical Journal*, 765, 131. doi:
  10.1088/0004-637X/765/2/131
- Kopparapu, R. K., Wolf, E. T., & Meadows, V. S. (2020). Characterizing exoplanet
   habitability. In *Planetary Astrobiology* (p. 449). University of Arizona Press.
- Korenaga, J. (2008). Urey ratio and the structure and evolution of Earth's mantle.
   *Reviews of Geophysics*, 46(2). doi: 10.1029/2007RG000241
- Korenaga, J. (2018). Crustal evolution and mantle dynamics through Earth history.
   *Philosophical Transactions of the Royal Society A: Mathematical, Physical and*

1223	Engineering Sciences, 376 (2132), 20170408. doi: 10.1098/rsta.2017.0408
1224	Korenaga, J., & Karato, SI. (2008). A new analysis of experimental data on olivine
1225	rheology. Journal of Geophysical Research: Solid Earth, 113(B2). doi: 10
1226	.1029/2007 JB005100
1227	Krissansen-Totton, J., Arney, G. N., & Catling, D. C. (2018). Constraining the
1228	climate and ocean pH of the early Earth with a geological carbon cycle model.
1229	Proceedings of the National Academy of Sciences, 115(16), 4105–4110. doi:
1230	10.1073/pnas.1721296115
1231	Krissansen-Totton, J., Bergsman, D. S., & Catling, D. C. (2016). On detecting
1232	biospheres from chemical thermodynamic disequilibrium in planetary atmo-
1233	spheres. Astrobiology, $16(1)$ , 39–67. doi: 10.1089/ast.2015.1327
1234	Krissansen-Totton, J., & Catling, D. C. (2017). Constraining climate sensitivity and
1235	continental versus seafloor weathering using an inverse geological carbon cycle
1236	model. Nature Communications, $\mathcal{S}(1)$ , 1–15. doi: 10.1038/ncomms15423
1237	Krissansen-Totton, J., Garland, R., Irwin, P., & Catling, D. C. (2018). Detectability
1238	of biosignatures in an oxic atmospheres with the James Webb Space Telescope:
1239	A TRAPPIST-1e case study. The Astronomical Journal, 156(3), 114. doi:
1240	10.3847/1538- $3881/aad564$
1241	Krissansen-Totton, J., Thompson, M., Galloway, M. L., & Fortney, J. J. (2022).
1242	Understanding planetary context to enable life detection on exoplanets and
1243	test the Copernican principle. Nature Astronomy, $6(2)$ , 189–198. doi:
1244	10.1038/s41550-021-01579-7
1245	Laneuville, M., Kameya, M., & Cleaves, H. J. (2018). Earth without life: A systems
1246	model of a global abiotic nitrogen cycle. Astrobiology, 18, 897–914. doi: 10
1247	.1089/ast.2017.1700
1248	Lehmer, O. R., Catling, D. C., & Krissansen-Totton, J. (2020). Carbonate-
1249	silicate cycle predictions of Earth-like planetary climates and testing the
1250	habitable zone concept. Nature Communications, $11(1)$ , 1–10. doi:
1251	10.1038/s41467-020-19896-2
1252	Lerman, A., & Wu, L. (2006). $CO_2$ and sulfuric acid controls of weathering and river
1253	water composition. Journal of Geochemical Exploration, 88(1-3), 427–430. doi:
1254	10.1016/j.gexplo.2005.08.100
1255	Lerman, A., Wu, L., & Mackenzie, F. T. (2007). CO <sub>2</sub> and H <sub>2</sub> SO <sub>4</sub> consump-

-52-

manuscript submitted to JGR: Planets

1256	tion in weathering and material transport to the ocean, and their role in
1257	the global carbon balance. Marine Chemistry, $106(1-2)$ , $326-350$ . doi:
1258	10.1016/j.marchem.2006.04.004
1259	Lewis, N. T., Lambert, F. H., Boutle, I. A., Mayne, N. J., Manners, J., & Acreman,
1260	D. M. (2018). The influence of a substellar continent on the climate of a
1261	tidally locked exoplanet. The Astrophysical Journal, 854(2), 171.
1262	Lin, Z., MacDonald, R. J., Kaltenegger, L., & Wilson, D. J. (2021). Differentiat-
1263	ing modern and prebiotic Earth scenarios for TRAPPIST-1e: high-resolution
1264	transmission spectra and predictions for JWST. Monthly Notices of the Royal
1265	Astronomical Society, $505(3)$ , $3562-3578$ . doi: $10.1093/mnras/stab1486$
1266	Lisse, C. M., Desch, S. J., Unterborn, C. T., Kane, S. R., Young, P. R., Hartnett,
1267	H. E., Izenberg, N. R. (2020). A geologically robust procedure for ob-
1268	serving rocky exoplanets to ensure that detection of atmospheric oxygen is a
1269	modern Earth-like biosignature. The Astrophysical Journal Letters, $898(1)$ ,
1270	L17. doi: 10.3847/2041-8213/ab9b91
1271	Lorenzo, A., Desch, S. J., & Shim, SH. (2014). On the lower radius limit of exo-
1272	planets. In Lunar and Planetary Science Conference (p. 1636).
1273	Lynch, J. K., Beatty, C. M., Seidel, M. P., Jungst, L. J., & DeGrandpre, M. D.
1274	(2010). Controls of riverine $CO_2$ over an annual cycle determined using di-
1275	rect, high temporal resolution $pCO_2$ measurements. Journal of Geophysical
1276	Research: Biogeosciences, $115(G3)$ . doi: $10.1029/2009JG001132$
1277	Lyubetskaya, T., & Korenaga, J. (2007). Chemical composition of Earth's prim-
1278	itive mantle and its variance: 1. Method and results. Journal of Geophysical
1279	Research: Solid Earth, 112(B3). doi: 10.1029/2005JB004223
1280	Ma, C., Wu, X., Huang, F., Zhou, Q., Li, F., & Cui, Q. (2012). The acoustic ve-
1281	locity, refractive index, and equation of state of liquid ammonia dihydrate
1282	under high pressure and high temperature. The Journal of Chemical Physics,
1283	137(10), 104504. doi: $10.1063/1.4751944$
1284	Madden, J., & Kaltenegger, L. (2020). How surfaces shape the climate of habitable
1285	exoplanets. Monthly Notices of the Royal Astronomical Society, 495(1), 1–11.
1286	doi: 10.1093/mnras/staa387
1287	Martin, JM., & Meybeck, M. (1979). Elemental mass-balance of material carried
1288	by major world rivers. Marine Chemistry, 7(3), 173–206. doi: 10.1016/0304

1289	-4203(79)90039-2
1290	McDonough, W. F., & Sun, SS. (1995). The composition of the Earth. Chemical
1291	Geology, 120(3-4), 223–253. doi: 10.1016/0009-2541(94)00140-4
1292	McKenzie, D. A. N., & Bickle, M. J. (1988). The volume and composition of melt
1293	generated by extension of the lithosphere. Journal of Petrology, $29(3)$ , $625-$
1294	679. doi: $10.1093/\text{petrology}/29.3.625$
1295	McLennan, S. M., & Grotzinger, J. P. (2008). The sedimentary rock cycle of Mars.
1296	In The Martian Surface–Composition, Mineralogy, and Physical Properties
1297	(pp. 541–577). Cambridge University Press.
1298	Mikal-Evans, T. $(2022)$ . Detecting the proposed $CH_4$ – $CO_2$ biosignature pair with
1299	the James Webb Space Telescope: TRAPPIST-1e and the effect of cloud/haze.
1300	Monthly Notices of the Royal Astronomical Society, $510(1)$ , $980-991$ . doi:
1301	10.1093/mnras/stab3383
1302	Ming, D. W., Morris, R. V., & Clark, B. C. (2008). Aqueous alteration on Mars. In
1303	$The \ Martian \ Surface-Composition, \ Mineralogy, \ and \ Physical \ Properties \ (pp.$
1304	519–540). Cambridge University Press.
1305	Moresi, LN., & Solomatov, V. S. (1995). Numerical investigation of 2D convection
1306	with extremely large viscosity variations. Physics of Fluids, $\gamma(9)$ , 2154–2162.
1307	doi: 10.1063/1.868465
1308	Mottl, M. J. (1983). Metabasalts, axial hot springs, and the structure of hydrother-
1309	mal systems at mid-ocean ridges. Geological Society of America Bulletin,
1310	94(2), 161–180. doi: https://doi.org/10.1130/0016-7606(1983)94\%3C161:
1311	$MAHSAT \ 3E2.0.CO;2$
1312	Nesbitt, H. W., & Markovics, G. (1997). Weathering of granodioritic crust, long-
1313	term storage of elements in weathering profiles, and petrogenesis of siliciclas-
1314	tic sediments. Geochimica et Cosmochimica Acta, 61(8), 1653–1670. doi:
1315	10.1016/S0016-7037(97)00031-8
1316	Nestola, F., Angel, R. J., Zhao, J., Garrido, C. J., Sánchez-Vizcaíno, V. L., Capitani,
1317	G., & Mellini, M. (2010). Antigorite equation of state and anomalous softening
1318	at 6 GPa: an in situ single-crystal x-ray diffraction study. Contributions to
1319	Mineralogy and Petrology, $160(1)$ , $33-43$ . doi: $10.1007/s00410-009-0463-9$
1320	Neveu, M., Desch, S. J., & Castillo-Rogez, J. C. (2017). Aqueous geochemistry in
1321	icy world interiors: Equilibrium fluid, rock, and gas compositions, and fate

-54-

1322	of antifreezes and radionuclides. Geochimica et Cosmochimica Acta, 212,
1323	324–371. doi: 10.1016/j.gca.2017.06.023
1324	Nordstrom, D. K., Plummer, L. N., Wigley, T. M. L., Wolery, T. J., Ball, J. W.,
1325	Jenne, E. A., Thrailkill, J. (1979). A comparison of computer-
1326	ized chemical models for equilibrium calculations in aqueous systems. In
1327	ACS Symposium Series (Vol. 93, p. 857-892). ACS Publications. doi:
1328	10.1021/bk-1979-0093.ch038
1329	Oelkers, E., Bénézeth, P., & Pokrovski, G. S. (2009). Thermodynamic databases for
1330	water-rock interaction. Reviews in Mineralogy and Geochemistry, $70(1)$ , 1–46.
1331	doi: 10.2138/rmg.2009.70.1
1332	Orgel, L. E. (1998). The origin of life–a review of facts and speculations. Trends in
1333	Biochemical Sciences, 23(12), 491–495. doi: 10.1016/s0968-0004(98)01300-0
1334	Palandri, J. L., & Kharaka, Y. K. (2004). A compilation of rate parameters of
1335	water-mineral interaction kinetics for application to geochemical modeling
1336	(Tech. Rep.). US Geological Survey. Menlo Park. CA.
1337	Pamukcu, A. S., Gualda, G. A. R., Ghiorso, M. S., Miller, C. F., & McCracken,
1338	R. G. (2015). Phase-equilibrium geobarometers for silicic rocks based on
1339	rhyolite-MELTS—Part 3: Application to the Peach Spring tuff (Arizona–
1340	California–Nevada, USA). Contributions to Mineralogy and Petrology, 169(3),
1341	33. doi: 10.1007/s00410-015-1122-y
1342	Papike, J. J., Ryder, G., & Shearer, C. K. (1998). Lunar samples. In J. J. Papike
1343	(Ed.), Planetary Materials (Vol. 36). Walter de Gruyter GmbH & Co KG.
1344	Parkhurst, D. L., & Appelo, C. A. J. (2013). Description of input and examples
1345	for PHREEQC version $3-A$ computer program for speciation, batch-reaction,
1346	one-dimensional transport, and inverse geochemical calculations (Tech. Rep.).
1347	US Geological Survey, Reston, VA.
1348	Pendergrass, A. G. (2020). The global-mean precipitation response to $CO_2$ -
1349	induced warming in CMIP6 models. $Geophysical Research Letters, 47(17),$
1350	e2020GL089964. doi: $10.1029/2020GL089964$
1351	Pidhorodetska, D., Fauchez, T. J., Villanueva, G. L., Domagal-Goldman, S. D.,
1352	& Kopparapu, R. K. (2020). Detectability of molecular signatures on
1353	TRAPPIST-1e through transmission spectroscopy simulated for future space-
1354	based observatories. The Astrophysical Journal Letters, 898(2), L33. doi:

1355	10.3847/2041- $8213/aba4a1$
1356	Pierrehumbert, R. T., & Hammond, M. (2019). Atmospheric circulation of tide-
1357	locked exoplanets. Annual Review of Fluid Mechanics, 51, 275–303. doi: 10
1358	.1146/annurev-fluid-010518-040516
1359	Pokrovsky, O. S., & Schott, J. (1999). Processes at the magnesium-bearing
1360	carbonates/solution interface. II. Kinetics and mechanism of magnesite
1361	dissolution. Geochimica et Cosmochimica Acta, 63(6), 881–897. doi:
1362	10.1016/S0016-7037(99)00013-7
1363	Pope, E. C., Bird, D. K., & Rosing, M. T. (2012). Isotope composition and vol-
1364	ume of Earth's early oceans. Proceedings of the National Academy of Sciences,
1365	109(12), 4371-4376.doi: 10.1073/pnas.1115705109
1366	Press, W. H., Teukolsky, S. A., Vetterling, W. T., & Flannery, B. P. (1992). Numer-
1367	ical Recipes in C, 2nd edition. Cambridge University, New York.
1368	Rasilo, T., Hutchins, R. H. S., Ruiz-González, C., & Del Giorgio, P. A. (2017).
1369	Transport and transformation of soil-derived $CO_2$ , $CH_4$ and DOC sustain $CO_2$
1370	supersaturation in small boreal streams. Science of the Total Environment,
1371	579, 902–912. doi: 10.1016/j.scitotenv.2016.10.187
1372	Reid, W. H., & Harris, D. L. (1958). Some further results on the Bénard problem.
1373	The Physics of Fluids, $1(2)$ , 102–110. doi: 10.1063/1.1705871
1374	Ringwood, A. E. (1991). Phase transformations and their bearing on the consti-
1375	tution and dynamics of the mantle. Geochimica et Cosmochimica Acta, $55(8)$ ,
1376	2083–2110. doi: 10.1016/0016-7037(91)90090-R
1377	Rudnick, R. L., & Gao, S. (2003). Composition of the continental crust. In <i>Treatise</i>
1378	on Geochemistry (Vol. 3, p. 659). Elsevier.
1379	Rushby, A. J., Johnson, M., Mills, B. J., Watson, A. J., & Claire, M. W. (2018).
1380	Long-term planetary habitability and the carbonate-silicate cycle. Astrobiol-
1381	ogy, 18(5), 469-480.doi: 10.1089/ast.2017.1693
1382	Saito, G., Kazahaya, K., Shinohara, H., Stimac, J., & Kawanabe, Y. (2001). Varia-
1383	tion of volatile concentration in a magma system of Satsuma-Iwojima volcano
1384	deduced from melt inclusion analyses. Journal of Volcanology and Geothermal
1385	Research, $108(1-4)$ , 11–31. doi: 10.1016/S0377-0273(00)00276-6
1386	Saito, H., & Kuramoto, K. (2018). Formation of a hybrid-type proto-atmosphere on
1387	Mars accreting in the solar nebula. Monthly Notices of the Royal Astronomical

1388	Society, $475(1)$ , 1274-1287. doi: 10.1093/mnras/stx3176
1389	Sanloup, C., Jambon, A., & Gillet, P. (1999). A simple chondritic model of Mars.
1390	Physics of the Earth and Planetary Interiors, 112(1-2), 43–54. doi: 10.1016/
1391	S0031-9201(98)00175-7
1392	Sata, N., Hirose, K., Shen, G., Nakajima, Y., Ohishi, Y., & Hirao, N. (2010). Com-
1393	pression of FeSi, Fe3C, $Fe_{0.95}O$ , and FeS under the core pressures and impli-
1394	cation for light element in the Earth's core. Journal of Geophysical Research:
1395	Solid Earth, $115(B9)$ . doi: $10.1029/2009JB006975$
1396	Schaefer, L., & Fegley, B. (2017). Redox states of initial atmospheres outgassed on
1397	rocky planets and planetesimals. The Astrophysical Journal, $843(2)$ , 120. doi:
1398	10.3847/1538-4357/aa784f
1399	Schulze-Makuch, D., Méndez, A., Fairén, A. G., Von Paris, P., Turse, C., Boyer, G.,
1400	$\dots$ Irwin, L. N. (2011). A two-tiered approach to assessing the habitability of
1401	exoplanets. Astrobiology, $11(10)$ , 1041–1052. doi: 10.1089/ast.2010.0592
1402	Schwieterman, E. W., Kiang, N. Y., Parenteau, M. N., Harman, C. E., DasSarma,
1403	S., Fisher, T. M., Lyons, T. W. (2018). Exoplanet biosignatures: a re-
1404	view of remotely detectable signs of life. Astrobiology, $18(6)$ , $663-708$ . doi:
1405	10.1089/ast.2017.1729
1406	Seager, S., Kuchner, M., Hier-Majumder, C., & Militzer, B. (2007). Mass-radius re-
1407	lationships for solid exoplanets. The Astrophysical Journal, $669(2)$ , 1279. doi:
1408	10.1086/521346
1409	Shahar, A., Driscoll, P., Weinberger, A., & Cody, G. (2019). What makes a planet
1410	habitable? Science, $364(6439)$ , $434-435$ . doi: 10.1126/science.aaw4326
1411	Sleep, N. H., & Zahnle, K. (2001). Carbon dioxide cycling and implications for
1412	climate on ancient Earth. Journal of Geophysical Research: Planets, $106(E1)$ ,
1413	1373–1399. doi: $10.1029/2000$ JE001247
1414	Smith, P. M., & Asimow, P. D. (2005). Adiabat_1ph: A new public front-end to the
1415	MELTS, pMELTS, and pHMELTS models. Geochemistry, Geophysics, Geosys-
1416	tems, $6(2)$ . doi: 10.1029/2004GC000816
1417	Solomatov, V. S. (1995). Scaling of temperature-and stress-dependent viscosity con-
1418	vection. Physics of Fluids, 7(2), 266–274. doi: 10.1063/1.868624
1419	Spiegel, D. S., Menou, K., & Scharf, C. A. (2009). Habitable climates: the influence
1420	of obliquity. The Astrophysical Journal, 691(1), 596. doi: 10.1088/0004-637X/

691/1/596

1421

1422	Stamenković, V., Noack, L., Breuer, D., & Spohn, T. (2012). The influence of
1423	pressure-dependent viscosity on the thermal evolution of super-Earths. The
1424	Astrophysical Journal, 748(1), 41. doi: 10.1088/0004-637X/748/1/41
1425	Tao, C., Sevfried, W., Lowell, R., Liu, Y., Liang, J., Guo, Z.,, Li, W. (2020).
1426	Deep high-temperature hydrothermal circulation in a detachment faulting sys-
1427	tem on the ultra-slow spreading ridge. Nature Communications, 11(1), 1–9.
1428	doi: 10.1038/s41467-020-15062-w
1429	Taylor, G. J. (2013). The bulk composition of Mars. <i>Geochemistry</i> , 73(4), 401–420.
1430	doi: 10.1016/j.chemer.2013.09.006
1431	Thompson, M. A., Krissansen-Totton, J., Wogan, N., Telus, M., & Fortney, J. J.
1432	(2022). The case and context for atmospheric methane as an exoplanet
1433	biosignature. Proceedings of the National Academy of Sciences, 119(14),
1434	e2117933119. doi: 10.1073/pnas.2117933119
1435	Thompson, R. N., Riches, A. J. V., Antoshechkina, P. M., Pearson, D. G., Nowell,
1436	G. M., Ottley, C. J., Niku-Paavola, V. (2007). Origin of CFB magmatism:
1437	multi-tiered intracrustal picrite–rhyolite magmatic plumbing at Spitzkoppe,
1438	western Namibia, during early Cretaceous Etendeka magmatism. Journal of
1439	Petrology, 48(6), 1119-1154.doi: 10.1093/petrology/egm012
1440	Tosca, N. J., McLennan, S. M., Lindsley, D. H., & Schoonen, M. A. (2004).
1441	Acid-sulfate weathering of synthetic Martian basalt: The acid fog model
1442	revisited. Journal of Geophysical Research: Planets, 109(E5). doi:
1443	10.1029/2003JE002218
1444	Trail, D., Watson, E. B., & Tailby, N. D. (2011). The oxidation state of Hadean
1445	magmas and implications for early Earth's atmosphere. Nature, $480(7375)$ , 79.
1446	doi: 10.1038/nature10655
1447	Treiman, A. H., & Schwenzer, S. P. (2009). Basalt–atmosphere interaction on
1448	Venus: Preliminary results on weathering of minerals and bulk rock. In $Venus$
1449	Geochemistry: Progress, Prospects, and New Missions (p. abstract $#2011$ ).
1450	Trenberth, K. E., Fasullo, J., & Smith, L. (2005). Trends and variability in column-
1451	integrated atmospheric water vapor. Climate dynamics, $24(7)$ , 741–758. doi:
1452	10.1007/s00382-005-0017-4
	,

-58-

1454	Tyburczy, J. A., Duffy, T. S., Ahrens, T. J., & Lange, M. A. (1991). Shock wave
1455	equation of state of serpentine to 150 GPa: Implications for the occurrence
1456	of water in the Earth's lower mantle. Journal of Geophysical Research: Solid
1457	Earth, $96(B11)$ , 18011–18027. doi: 10.1029/91JB01573
1458	Tziperman, E., Abbot, D. S., Ashkenazy, Y., Gildor, H., Pollard, D., Schoof, C. G.,

- <sup>1459</sup> & Schrag, D. P. (2012). Continental constriction and oceanic ice-cover thick <sup>1460</sup> ness in a Snowball-Earth scenario. Journal of Geophysical Research: Oceans,
   <sup>1461</sup> 117(C5). doi: 10.1029/2011JC007730
- Unterborn, C. T., Byrne, P. K., Anbar, A. D., Arney, G. N., Brain, D., Desch, S. J.,
  ... Way, M. J. (2021). Exogeoscience and its role in characterizing exoplanet
  habitability and the detectability of life. Bulletin of the American Astronomical Society. doi: 10.3847/25c2cfeb.5209dd13
- Unterborn, C. T., Dismukes, E. E., & Panero, W. R. (2016). Scaling the Earth: a
   sensitivity analysis of terrestrial exoplanetary interior models. *The Astrophysi- cal Journal*, 819(1), 32. doi: 10.3847/0004-637X/819/1/32
- <sup>1469</sup> Unterborn, C. T., Foley, B. J., Desch, S. J., Young, P. A., Vance, G., Chiffelle,
- <sup>1470</sup> L., & Kane, S. R. (2022). Mantle degassing lifetimes through galac-
- tic time and the maximum age stagnant-lid rocky exoplanets can support
- temperate climates. The Astrophysical Journal Letters, 930(1), L6. doi: 1473 10.3847/2041-8213/ac6596
- Unterborn, C. T., Kabbes, J. E., Pigott, J. S., Reaman, D. M., & Panero, W. R.
  (2014). The role of carbon in extrasolar planetary geodynamics and habitability. *The Astrophysical Journal*, 793(2), 124. doi: 10.1088/0004-637X/793/2/
  124
- <sup>1478</sup> Urey, H. C. (1952). The Planets: their Origin and Development. Yale Univ. Press.
- Vance, S., Harnmeijer, J., Kimura, J., Hussmann, H., DeMartin, B., & Brown, J. M.
  (2007). Hydrothermal systems in small ocean planets. Astrobiology, 7(6),
  987–1005. doi: 10.1089/ast.2007.0075
- Vervoort, P., Horner, J., Kane, S. R., Turner, S. K., & Gilmore, J. B. (2022). System
   architecture and planetary obliquity: Implications for long-term habitability.
   The Astronomical Journal, 164 (4), 130. doi: 10.3847/1538-3881/ac87fd
- Vidaurri, M. R., Bastelberger, S. T., Wolf, E. T., Domagal-Goldman, S., & Koppa-
- rapu, R. K. (2022). The outer edge of the Venus zone around main-sequence

1487	stars. The Planetary Science Journal, $3(6)$ , 137. doi: 10.3847/PSJ/ac68e2									
1488	Wade, J. A., Plank, T., Melson, W. G., Soto, G. J., & Hauri, E. H. (2006). The									
1489	volatile content of magmas from Arenal volcano, Costa Rica. Journal of Vol-									
1490	canology and Geothermal Research, 157(1-3), 94–120. doi: 10.1016/j.jvolgeores									
1491	.2006.03.045									
1492	Walsh, K. J., & Levison, H. F. (2019). Planetesimals to terrestrial planets: colli-									
1493	sional evolution amidst a dissipating gas disk. <i>Icarus</i> , 329, 88–100. doi: 10									
1494	.1016/j.icarus.2019.03.031									
1495	Weider, S. Z. (2019). Petrology and geochemistry of Mercury. Oxford Research En-									
1496	cyclopedia of Planetary Science, 127. doi: 10.1093/acrefore/9780190647926.013									
1497	.127									
1498	Weidner, D. J., Bass, J. D., Ringwood, A. E., & Sinclair, W. (1982). The single-									
1499	crystal elastic moduli of stishovite. Journal of Geophysical Research: Solid									
1500	Earth, $87(B6)$ , $4740-4746$ . doi: 10.1029/JB087iB06p04740									
1501	Wett, J. P., & Ahrens, T. J. (1983). Shock compression of single-crystal forsterite.									
1502	Journal of Geophysical Research: Solid Earth, 88(B11), 9500–9512. doi:									
1503	10.1029/JB088iB11p09500									
1504	Wordsworth, R. D., Forget, F., Selsis, F., Millour, E., Charnay, B., & Madeleine,									
1505	JB. (2011). Gliese 581d is the first discovered terrestrial-mass exoplanet									
1506	in the habitable zone. The Astrophysical Journal Letters, $733(2)$ , L48. doi:									
1507	10.1088/2041-8205/733/2/L48									
1508	Young, P. A., Desch, S. J., Anbar, A. D., Barnes, R., Hinkel, N. R., Kopparapu,									
1509	R., Truitt, A. (2014). Astrobiological stoichiometry. Astrobiology, $14(7)$ ,									
1510	603-626. doi: $10.1089/ast.2014.1143$									
1511	Zhang, Y., & Zindler, A. (1993). Distribution and evolution of carbon and nitro-									
1512	gen in Earth. Earth and Planetary Science Letters, 117(3-4), 331–345. doi: 10									
1513	.1016/0012- $821X(93)90088$ -Q									

Table 5. Starting crustal compositions in mol% for continental weathering simulations. Horizontal lines delineate broad mineral groups: pyroxenes and olivines, feldspars, cyclosilicates, silica polymorphs, hornblende, secondary minerals, carbonates, sulfates, sulfides, and halides. Notes and references: <sup>a</sup> Sedimentary. <sup>b</sup> Hydrothermal. <sup>c</sup> Matches both oxide mass fractions from Rudnick and Gao (2003) and mineral volume fractions from Nesbitt and Markovics (1997) within 10% prior to the addition of carbonates, sulfates, sulfates, and halite. The kinetic parameters of annite and daphnite are assumed to be the same as those of phlogopite and clinochlore, respectively, from Palandri and Kharaka (2004).<sup>d</sup> Matches primitive mantle oxide mass fractions of Hofmann (1988) within 4% prior to addition of pyrite and halite. <sup>e</sup> From Table 3 (average Mercury Southern hemisphere composition) of Weider (2019), matching both mineral and elemental mass fractions, and attributing all hyperstheme, olivine, and sulfide respectively as enstatite, forsterite, and pyrite to match the high Mg/Fe and S/Fe ratios.  $^{f}$  Results of modeled interaction between basalt and Venus' atmosphere (Treiman & Schwenzer, 2009; Dyar et al., 2021).  $^{g}$  Highland composition (Table 5.1 of Papike et al. (1998)) with feldspar composition from their Table A5.6, olivine composition from the Fe/(Fe+Mg) ratios of their Table A5.9, and Ca- and Mg-rich pyroxene compositions (attributed to 50% diopside / 25% enstatite / 25% wollastonite in case 1; 100% diopside in case 2) from columns 17 and 19 of their Table A5.4 due to the lack of kinetic data for Fe-pyroxene (ferrosilite) in Palandri and Kharaka (2004).  $^{h}$  From the Martian upper crust experimental mineralogy used by Tosca et al. (2004) as reported in Table 24.3 of McLennan and Grotzinger (2008), with plagioclase attributed to albite based on the lack of anorthite in a Meridiani Planum outcrop (Table 23.3 of Ming et al. (2008)), oxides attributed to hematite, and addition of jarosite and anhydrite at abundances bracketed by the lack of sulfates in the Tosca et al. (2004) composition and the tens of mass% of sulfates in Table 24.3 of McLennan and Grotzinger (2008) and Table 26.2 of Des Marais et al. (2007). <sup>i</sup> Compositions for Fig. 4b and 5d of De Sanctis et al. (2015), assuming their optically dark component is magnetite.

Mineral	Formula	$\begin{array}{c} \operatorname{Earth}^{c} \\ (\operatorname{modern}) \end{array}$	$\begin{array}{c} \operatorname{Earth}^{d} \\ (\operatorname{Archean}) \end{array}$	Mercury	<sup>e</sup> Venus $1^f$ (reduced)	Venus $2^f$ (oxidized)	Lunar 1 <sup>9</sup>	Lunar 2 <sup>9</sup>	<sup><i>I</i></sup> Mars <sup><i>h</i></sup>	Ceres 1	$i$ Ceres $2^i$
Fayalite Forsterite Enstatite Wollastonite Diopside	$\begin{array}{c} \mathrm{Fe_2SiO_4}\\ \mathrm{Mg_2SiO_4}\\ \mathrm{MgSiO_3}\\ \mathrm{CaSiO_3}\\ \mathrm{CaMg(SiO_3)_2} \end{array}$	- - - -	7.5 53.6 27.2 3.1	$\frac{8.5}{51.1}$	- 38.8 -	12.2	$2.0 \\ 4.9 \\ 9.5 \\ 9.5 \\ 18.9$	2.4 5.3 - 32.4	2.0 10.7 23.8 -	- - - -	
Anorthite Albite K-feldspar	$\begin{array}{c} { m CaAl_2Si_2O_8} \\ { m NaAlSi_3O_8} \\ { m KAlSi_3O_8} \end{array}$	$^{8.4}_{15.6}_{5.0}$	4.7 1.5	$11.7 \\ 17.9 \\ 0.4$	11.8	11.2	$16.5 \\ 0.9 \\ 0.1$	$17.7 \\ 0.9 \\ 0.2$	$12.3 \\ 0.4$	- - -	- - -
Cordierite	$\mathrm{Mg_2Al_4Si_5O_{18}}$	-	-	-	7.6	7.2	-	-	-	-	-
$\begin{array}{c} \label{eq:Quartz} \\ \mbox{Amorph. SiO}_2 \end{array}$	${{ m SiO}_2} m SiO_2$	59.7	-	-	12.2	31.1	37.8	41.2	39.2	-	-
Anthophyllite Tremolite	$\substack{ Mg_{7}Si_{8}O_{22}(OH)_{2}\\ Ca_{2}Mg_{5}Si_{8}O_{22}(OH)_{2} }$	$\begin{array}{c} 0.21 \\ 0.43 \end{array}$	-	-	-	-	-	-	-	-	-
Annite Phlogopite Daphnite Lizardite Magnetite Hematite	$\begin{array}{c} {\rm KAlFe_3Si_3O_{10}(OH)_2}\\ {\rm KAlMg_3Si_3O_{10}(OH)_2}\\ {\rm Fe_5AlAlSi_3O_{10}(OH)_8}\\ {\rm Mg_3Si_2O_5(OH)_4}\\ {\rm Fe_3O_4}\\ {\rm Fe_2O_3} \end{array}$	1.0 1.8 1.4 - -	- - - - -		- - - -	- - 9.9		- - - - -	- - - 7.1	1.6 - 3.6 57.1	2.2 - 4.4 72.3
	${caCO_3 \atop CaMg(CO_3)_2 \atop CaMg(CO_3)_2 \atop MgCO_3}$	4.8 1.7 -	6 	1	- - -	- - -		- - - -	- - - -	- 37.7	- 21.2
Gypsum Anhydrite Jarosite	$\begin{array}{c} {\rm CaSO_4\cdot 2H_2O}\\ {\rm CaSO_4}\\ {\rm KFe_3(SO_4)_2(OH)_6} \end{array}$	0.89	- - -	- - -	29.6	28.4	- - -	- - -	$2.8 \\ 1.8$	- - -	- - -
Pyrite	$\mathrm{FeS}_2$	0.17	0.21	6.1	-	-	-	-	-	-	-
Halite	NaCl	3.8	2.3	-	-	-	-	-	-	-	-