# Ferric iron evolution during crystallization of the Earth and Mars

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#### Abstract

Magma ocean crystallization models that track fO2 evolution can reproduce the D/H ratios of both the Earth and Mars without the need for exogenous processes. Fractional crystallization leads to compositional evolution of the bulk oxide components. Metal-saturated magma oceans have long been thought to contain negligible ferric iron oxide (Fe3+O1.5), but recent work suggests they may contain near-present-day Fe3+ concentrations. We model the fractional crystallization of Earth and Mars, including Fe2+ and Fe3+ as separate components. We use two independent equations of state (Deng, Armstrong EOS) to calculate Fe3+ partition coefficients for lower mantle minerals and compare results of fractional crystallization for different magma ocean configurations for both Earth and Mars. We calculate the oxygen fugacity (fO2) at the surface as the systems evolve and compare them to constraints on the fO2 of the last magma ocean atmosphere from D/H ratios. For Earth, we find that Fe3+ must behave incompatibly in the lower mantle to match the D/H constraint for whole mantle models, but shallow magma ocean models also provide reasonable matches to the constraints. For Mars, both EOSs produce near identical results but cannot match the D/H constraints on last fO2 unless the magma ocean begins with less than 50% of the predicted Fe3+. This model shows that Fe3+ partitioning has a measurable effect on magma ocean atmosphere redox state, which is not a static quantity but evolves throughout the magma ocean's lifetime. We highlight the need for additional experimental constraints on ferric iron partitioning.

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Ferric iron evolution during crystallization of the Earth and Mars 1 Laura Schaefer<sup>1</sup>, Kaveh Pahlevan<sup>2</sup>, Linda T. Elkins-Tanton<sup>3</sup> 2 3 <sup>1</sup> Department of Earth and Planetary Sciences, Stanford University 4 <sup>2</sup>Carl Sagan Center, SETI Institute, Mountain View, CA 5 <sup>3</sup>School of Earth and Space Exploration, Arizona State University, 6 Corresponding author: Laura Schaefer (lkschaef@stanford.edu) 7 **Key Points:** 8 We model mineral/melt partitioning during magma ocean crystallization using new Fe<sup>3+</sup> 9 • partition coefficients for lower mantle minerals. 10 We calculate oxygen fugacity  $(fO_2)$  of outgassing at the surface for Earth and Mars • 11 magma oceans and match to constraints from planetary D/H. 12

For Earth, only models using the Armstrong EOS match constraints, whereas Mars must
 start Fe<sup>3+</sup>-depleted to match constraints.

#### 16 Abstract

- 17 Magma ocean crystallization models that track *f*O<sub>2</sub> evolution can reproduce the D/H ratios of
- 18 both the Earth and Mars without the need for exogenous processes. Fractional crystallization
- 19 leads to compositional evolution of the bulk oxide components. Metal-saturated magma oceans
- have long been thought to contain negligible ferric iron oxide (Fe<sup>3+</sup>O<sub>1.5</sub>), but recent work
- suggests they may contain near-present-day Fe<sup>3+</sup> concentrations of planetary mantles. We model
- the fractional crystallization of Earth and Mars, including  $Fe^{2+}$  and  $Fe^{3+}$  as separate components.
- We use two independent equations of state (Deng, Armstrong EOS) to calculate  $Fe^{3+}$  partition
- 24 coefficients for lower mantle minerals and compare results of fractional crystallization for
- different magma ocean configurations for both Earth and Mars. We calculate the oxygen fugacity ( $fO_2$ ) at the surface as the systems evolve and compare them to constraints on the  $fO_2$  of the last
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- incompatibly in the lower mantle to match the D/H constraint for whole mantle models, but
- shallow magma ocean models also provide reasonable matches to the constraints. For Mars, both
- EOSs produce near identical results but cannot match the D/H constraints on last  $fO_2$  unless the
- magma ocean begins with less than 50% of the predicted  $Fe^{3+}$ . This model shows that  $Fe^{3+}$
- partitioning has a measurable effect on magma ocean atmosphere redox state, which is not a
- static quantity but evolves throughout the magma ocean's lifetime. We highlight the need for
- 34 additional experimental constraints on ferric iron partitioning.
- 35

#### 36 Plain Language Summary

- 37 During the formation of planets like Earth and Mars, many undergo a stage in which their rocky
- mantles are almost fully molten, called a magma ocean. The crystallization of different minerals
- 39 as the magma ocean cools leads to changes in the composition of the melt. The melt maintains
- 40 contact with the atmosphere throughout this stage, so changes in the melt's composition,
- 41 especially the availability of oxygen, can lead to changes in the atmosphere's composition. In
- 42 this paper, we use a model that looks at how the melt's composition changes, with a special
- focus on how different types (or species) of iron  $(Fe^{2+}, Fe^{3+})$  behave as minerals form. We use
- 44 two different models called equations of state (EOS) of how these iron species behave at high 45 pressures and compare how the melt's composition changes when we use each of them. For
- Earth, only one of the EOS models produces results that agree with information that we have
- 47 about the composition of the last atmosphere in contact with the magma ocean. For Mars, both
- 48 models produce similar results, but neither is successful in recreating the last atmosphere unless
- 49 the melt starts with much less of the  $Fe^{3+}$  species than previously predicted.

#### 50 **1. Introduction**

The oxidation state of Earth's present day upper mantle is close to the quartz-fayalite-magnetite 51 (QFM) buffer (Canil & O'Neill 1996; Frost & McCammon 2008), which results in oxidized 52 53 volcanic gases such as  $H_2O$  and  $CO_2$ . Measurements of redox proxies going back to at least 3900 Ma (Delano 2001), and possibly as far back as 4.3 Ga (Trail et al. 2011), are consistent with 54 present day values, indicating little secular change in mantle oxidation state through time, 55 although recent measurements suggest a slightly lower mantle oxidation state during the early 56 Archean of ~QFM-0.5/1 (Aulbach & Stagno 2016, Nicklas et al. 2019). However, models of 57 core formation suggest that the Earth's mantle would have been two log units below the iron-58 wüstite buffer (~IW-2) at the end of planet formation (Rubie et al. 2015, Badro et al. 2015). As 59 first suggested by Hirschmann (2012), recent measurements of ferric ( $Fe^{3+}$ ) iron in silicate melts 60 and first principles calculations indicate that high pressure conditions may stabilize ferric iron 61 over ferrous iron (Armstrong et al., 2019; Zhang et al., 2017, Deng et al. 2020, Kuwahara et al. 62 2023), unlike earlier lower pressure data (Kress & Carmichael 1991; O'Neill et al. 2006). Fe-63 bearing silicates equilibrated at high pressure then have non-negligible ferric iron contents, even 64 at reducing conditions. This suggests that once metal segregates during core formation, the 65 oxygen fugacity of a well-mixed mantle at low pressures – often characterized by its  $Fe^{+3}/Fe^{+2}$  – 66 would suddenly increase. Hirschmann (2022) recalibrated a silicate melt equation of state and 67 calculated the ferric iron contents of silicate melts under core-formation conditions for Earth and 68

- Mars and found  $Fe^{3+}/Fe_T = 0.04 0.10$  and 0.026 0.038, respectively, consistent with the
- 70 present day mantles of the two planets.
- 71

72 Chemical partitioning models of magma ocean evolution have also previously shown that

fractional crystallization will tend to increase the FeO abundance in the magma ocean during
 soldification (Elkins-Tanton et al., 2003, 2005). This results in a denser, iron-rich upper mantle,

soldification (Elkins-Tanton et al., 2003, 2005). This results in a denser, iron-rich upper mantle,
 which may cause a massive early mantle overturn event (Boukaré et al. 2018; Elkins-Tanton et

- al. 2003). Ferric iron is moderately incompatible during both partial melting and fractional
- crystallization (Canil & O'Neill 1996; Cottrell & Kelley 2011; Sorbadere et al. 2018) in the

vpper mantle and may therefore fractionate from ferrous iron during fractional crystallization of

the magma ocean, as first suggested by McCanta et al. (2009). Most previous fractional

80 crystallization models did not include ferric iron because it was thought to be present at

- vanishingly small amounts for oxygen fugacities below IW. Fractionation of ferric and ferrous
- 82 iron during magma ocean solidification would produce both vertical and temporal variations in
- redox state, which may influence mantle mineralogy as well as early outgassed atmospheric
- 84 compositions.
- 85

In this paper, we seek to determine the effect that ferric iron partitioning could have on the

oxygen fugacity of the last magma ocean atmospheres of Earth and Mars. Pahlevan et al. (2019,

88 2022) determine the oxygen fugacity of the last atmosphere equilibrated with a magma ocean by

- 89 determining the maximum amount of D/H fractionation that could have occurred for Earth and
- 90 Mars, using constraints on initial D/H of early hydrogen reservoirs. These models assume that all
- 91 hydrogen that escapes is carried to the upper atmosphere in the form of H<sub>2</sub>, and no hydrogen is
- 92 stripped from water vapor during the steam atmosphere stage. Then the relative abundances of
- $H_2$  and  $H_2O$  in the last magma ocean atmosphere constrain the oxygen fugacity through the
- 94 reaction:
- 95

 $H_2 + \frac{1}{2}O_2 = H_2O \tag{1}$ 

Assuming a chondritic source of early water on Earth, Pahlevan et al. (2019) find that only a 98 small amount of hydrogen escape is allowed, which requires the oxygen fugacity of the last 99 magma ocean atmosphere for Earth to be greater than IW + 1. Under these conditions, the 100 101 abundance of  $H_2$  is much lower than the abundance of  $H_2O$  in the gas phase equilibrated with the magma ocean. In contrast, there appears to be evidence for moderate to high escape of hydrogen 102 from early Mars, and Pahlevan et al. (2022) find that the D/H ratio would be consistent with a 103 last magma ocean atmosphere less than IW - 1. These constraints therefore provide a lower limit 104 on oxygen fugacity for Earth but an upper limit for the last magma ocean atmosphere of Mars. 105

106

107 In this paper, we present models for ferric iron partitioning in a crystallizing magma ocean on

Earth and Mars, considering both batch and fractional crystallization scenarios. Our model
 follows the evolution of the ferric iron as it partitions between mantle minerals and magma ocean

follows the evolution of the ferric iron as it partitions between mantle minerals and magma occ liquid. To do this, we must first determine ferric iron partition coefficients for all the

111 crystallizing minerals. Whereas measurements of ferric iron partition coefficients are available

for many of the upper mantle minerals (Mallmann & O'Neill 2009, Davis & Cottrell 2021,

113 Rudra & Hirschmann 2022), most of the lower mantle minerals have no such measurements. In

114 this paper, we therefore use the best available mineral constraints in conjuction with a silicate

melt model for the variation of  $Fe^{3+}/Fe^{2+}$  with oxygen fugacity in order to estimate the ferric iron

partition coefficients. Two silicate melt equations of state (EOS) extend up to high enough

pressure to use for these calculations (Armstrong et al. 2019, Deng et al. 2020). We find that

these two EOSs yield opposite partitioning behavior (compatible vs. incompatible) for several of

the lower mantle minerals, including bridgmanite. Our partitioning model therefore provides a

test of the two EOSs to determine which model is most consistent with the D/H constraints forEarth and Mars.

122

123 This paper is organized as follows. We first discuss the model below and describe the ferric iron

124 partition coefficients for the abundant mantle minerals (**Section 2**) using the two EOS

parameterizations for ferric iron oxide (Armstrong et al. 2019, Deng et al. 2020). We then

discuss what magma ocean conditions the D/H constraints of Pahlevan et al. (2019, 2022) may

127 correspond to. We then show results for the  $fO_2$  evolution at the surface for different initial ferric

iron abundances and different crystallization scenarios (Section 3) for both planets to determine

129 which model parameters best match the D/H constraints.

130

## 131 2. Magma ocean model

132

133 2.1 Fractional Crystallization: phases and proportions

134 In this paper, we modify the magma ocean fractional crystallization model of Elkins-Tanton

135 (2008). The magma ocean crystallizes from a given depth (or pressure) upwards, following a

specified mineralization sequence determined by mineral phase stability along the mantle

137 solidus, with slightly different mineral proportions for Earth and Mars (Elkins-Tanton 2008,

138Table 2). We do not explicitly model the planetary thermal evolution timescales or volatile

139 evolution here, focusing instead on the detailed mineral/melt chemical evolution. We update the

140 Elkins-Tanton fractional crystallization model by modifying the internal structure calculation and

141 including ferric/ferrous iron partitioning between the silicate liquid and crystallizing phases. We

- 142 also explore the uncertainties in the ferric iron partition coefficients using Monte Carlo
- simulations. We discuss these updates in detail below.
- 144
- 145 2.2 Internal structure
- 146 We specify the core mass fraction of the planet and then the internal structure of the planet is
- 147 calculated for a liquid silicate mantle and isolated Fe core by solving the equations of internal
- structure with spherical symmetry and hydrostatic equilibrium:

149 
$$\frac{dm(r)}{dr} = 4\pi r^2 \rho(r)$$
(2)  
$$\frac{dm(r)}{dP(r)} = -Gm(r)\rho(r)$$

 $\frac{dr}{dr} = \frac{-Gm(r)\rho(r)}{r^2}$ (3)

As the planetary mantle solidifies, the density of the minerals in each solid layer is determined 151 and the radius of solidification is calculated, beginning from the core-mantle boundary. The 152 crystallization occurs in layers of 1 wt% of the mantle's mass. The depth of the magma ocean is 153 determined for the new magma ocean composition by determining the density of the silicate 154 liquid with depth. We compute the internal structure evolution for each of the magma ocean 155 scenarios (see Section 2.3) once, to determine the evolution of the radius of solidification, then 156 we use this calculated structure for each of the Monte Carlo simulations with different ferric iron 157 partition coefficients. Using this procedure, the density of the layers that are strongly ferric iron-158

- 159 enriched may be slightly underestimated.
- 160

161 2.3 Whole Mantle vs. Shallow Magma Ocean Models

In our models, we test different magma ocean configurations for both Earth and Mars (see 162 Figure 1), with either different initial depths of melting or a switch between crystallization style 163 from batch (or equilibrium) to fractional crystallization. Purely fractional crystallization 164 produces larger degrees of fractionation than batch crystallization, but the switch from batch to 165 fractional crystallization depends on crystal size within the magma ocean (Solomatov 2015), 166 with both mechanisms possible under the convective conditions of Earth's magma ocean. Xie et 167 al. (2020) use new measurements of silicate liquid viscosities to calculate crystal fractions and 168 find that fractional crystallization likely started when the base of Earth's magma ocean was 169 ~1000 km depth, with formation of a bridgmanite-rich layer at the top of the lower mantle during 170 batch crystallization. In the absence of good constraints on the Martian magma ocean, we will 171 172 adopt identical scenarios for both planets. We test whole mantle magma ocean models in which batch crystallization switches to fractional crystallization at 500 km and 1000 km, as shown in 173 Figure 1(*left*), compared to a fully fractional crystallization model. We also compare these 174 results to shallow magma ocean models, also starting at 500 or 1000 km, in which we assume 175 that the lower mantle does not melt and the starting compositions are the Bulk Silicate planet 176 compositions given in Table 1. The differences in results between whole mantle vs. shallow 177 magma oceans is driven entirely by the bulk composition at the initiation of fractional 178 crystallization. 179

For the whole mantle models, we adopt a simplified model of batch crystallization, inwhich we compute bulk partition coefficients for each of the melt oxide phases via:

183 
$$\overline{D} = \sum_{i} X_{i} D_{i}$$

184

182

(4)

where  $X_i$  is the mass fraction of mineral phase *i* in the solid,  $D_i$  is the partition coefficient between mineral *i* and the melt and  $\overline{D}$  is the bulk partition coefficient between solid and liquid. We then use the batch crystallization equation to calculate the composition of the liquid following crystallization of some mass fraction of the mantle.

- 189
- 190

$$\frac{C_{liq}}{C_{liq}^0} = \frac{1}{F(1-\overline{D})+\overline{D}} = \frac{1}{F+\overline{D}(1-F)}$$
(5)

191

where F is the degree of melting,  $C_{lig}$  is the mass fraction of an oxide in the melt after 192 crystallization,  $C_{lig}^{0}$  is the initial mass fraction of the oxide in the magma ocean. The conditions 193 for batch crystallization for each scenario are given in Table 2, including the pressure and 194 temperature at the base of the magma ocean layer (top of solid layer), the mass fraction of the 195 magma ocean relative to the total mantle, and the mineral mass fractions within the crystallized 196 197 layer, normalized to 1. We use these mineral fractions in equation 5 for calculating bulk partition coefficients for each model. We adopt the same starting pressure and temperature conditions for 198 the shallow magma ocean models. 199

200

In **Table S1**, we summarize the partition coefficients  $(D_i)$  for each of the melt oxide components 201 (SiO<sub>2</sub>, MgO, FeO, CaO, Al<sub>2</sub>O<sub>3</sub>) that we use in the batch crystallization model. Note that we do 202 not specifically calculate mineral phase compositions or apply stoichiometric corrections as done 203 during the fractional crystallization stage. We modify the exchange coefficients (e.g,  $K_D^{Fe/Mg}$  = 204  $D_{FeO}/D_{MaO}$ ) for the fractional crystallization stage to ensure consistency with the  $D_i$  used in the 205 calculation of the bulk partition coefficients (eqn 5). We summarize our new  $K_D$  values 206 compared to the Elkins-Tanton (2008) model in Table S2. Most values are in relatively close 207 208 agreement, and we find little change in the overall model behavior when using either set of values. 209

210

To take into account the uncertainties on the partition coefficients, we performed Monte Carlo simulations using 10<sup>5</sup> randomized draws of oxide partition coefficients, assuming uncorrelated normal distributions, to perform the batch crystallization calculations. These simulations produced mean oxide abundances that were reproducible within 0.01 wt%, which we use as the starting compositions for the fractional crystallization models. We report the mean values for each batch scenario in **Table 1**, along with the Bulk Silicate Earth (BSE) (McDonough 2003) and Bulk Silicate Mars (BSM) (Taylor 2013) compositions, which are also used as starting

218 conditions for the shallow magma ocean models.

- 219
- 220 2.4 Pressure of last equilibration

221

To relate the oxygen fugacity predicted by our fractional crystallization models to the D/H

constraints of Pahlevan et al. (2019, 2022), we must determine the conditions under which the

magma ocean and the atmosphere begin to evolve separately. Magma ocean thermal evolution

models suggest that the mantle evolves from a fully molten system to a partially molten "mush"

system, at which point a thick lid (> a few km) can form (Solomatov 2007, Lebrun et al. 2012,

Monteux et al. 2020). The lid inhibits direct magma ocean-atmosphere exchange, with subsequent transfer of volatiles from the interior to the atmosphere requiring physical advection

of material through the lid. This is often called the rheological transition. Some portion of the 229

- upper mantle remains molten at this stage, with additional volatiles trapped within the melt layer 230
- (Elkins-Tanton 2008, Hier-Majumder & Hirschmann 2017, Miyazaki & Korenaga 2022). We 231 characterize the separation of the atmosphere and remaining magma ocean by the pressure at the 232
- base of the partially molten layer when the thick lid forms, which we refer to as the *basal* 233
- pressure of last equilibration (BPLE). We show an example thermal profile that illustrates the 234
- intersection of solidus and magma ocean adiabat that determines the BPLE in Figure S4. 235
- Although our fractional crystallization models continue to lower pressures than the BPLE, we 236
- search for models which produce the allowed oxygen fugacity within the bounds of the BPLE 237
- that we calculate below. 238
- 239

We use a 1D magma ocean thermal evolution model (Schaefer et al. 2016, Lebrun et al. 2013) to 240

- constrain the BPLE for both Earth and Mars. Note that this model is a physical simulation that is 241 separate from the chemical model of fractional crystallization that we discuss throughout the rest 242
- of this paper. The thermal evolution model calculates the evolution of mantle and surface
- 243 temperatures along with volatile evolution within the coupled mantle-atmosphere system. We
- 244
- find that the BPLE is not sensitive to the volatile content or any parameters that determine the 245
- timing of solidification. Instead, we find that the BPLE depends most sensitively on the assumed 246
- solidus and liquidus curves, and especially the temperature difference between the solidus and 247 248 liquidus ( $T_{\text{liquidus}}$ - $T_{\text{solidus}}$  called liquidus offset hereafter) within the upper mantle layers, which
- determines melt fraction ( $\varphi = (T_{mantle} T_{solidus})/(T_{liquidus} T_{solidus})$ ) and dictates the evolution of the 249
- rheological front. We find that if the liquidus offset is reduced, the BPLE drops to lower 250
- pressures. 251

252

A conservative assumption of a dry peridotite solidus and liquidus (Pierru et al. 2022) for Earth 253 yields BPLE of 15.3 GPa. Pierru et al. (2022) find a liquidus offset of ~300 K in the upper 254 mantle, which increases to ~600 K at the core-mantle boundary. We find similar BPLE values 255 256 using Pierru et al (2022)'s solidus temperature with a constant liquidus offset of 300 K. However, the presence of volatiles such as water and CO<sub>2</sub> in the liquid will lower solidus and 257 liquidus temperatures (Katz et al. 2003, Medard & Grove 2008). If we reduce both the solidus 258 and liquidus temperatures equally by 200 and 400 K, we find BPLE for Earth of 14.6 and 14.0 259 GPa, respectively, a relatively small effect. If volatile effects do not equally reduce the solidus 260 and liquidus temperatures, these results may vary more greatly, with higher (lower) BPLE values 261 262 if the solidus-liquidus gap increases (decreases). An additional effect to consider is that fractional crystallization enriches the liquid in incompatible elements, which can produce an equal or 263 greater decrease in the solidus temperatures compared to volatile-rich models (Elkins-Tanton 264 265 2008). Elkins-Tanton (2008) determined solidus temperatures for Earth and Mars by calculating melting points for their fractionally crystallizing systems using the MELTs model, incorporating 266 both the incompatible element and volatile effects. We use their reported solidus temperatures 267 with liquidus offsets ranging from 300 to 600 K and find BPLE from 6 – 10 GPa. Note also that 268 we assume that the liquidus offset is constant with pressure, which is likely an oversimplification 269 270 but provides reasonable upper and lower bounds.

271

The solidus temperature of Mars is different from the peridotite solidus of the Earth due to higher 272

- FeO contents. Liquidus temperatures are not well-determined for Mars, but based on work by 273
- Longhi et al. (1992), we estimate a solidus-liquidus offset of between 300 400 K. Duncan et al. 274
- (2018) measured solidus temperatures for Mars-analog compositions. Using their solidus plus 275

liquidus offsets of 300 - 400 K, we find BPLE of 2.8 - 3.8 GPa for Mars. This range is much

lower than for the Earth due to the effect of the lower gravity of Mars on the temperature

structure with pressure. We again find minimal differences if both solidus and liquidus are

equally reduced by impurities. Elkins-Tanton (2008) also determined the solidus for a

fractionally crystallizing Mars. Using their solidus with 300 - 400 K liquidus offsets, we find

281 BPLE of 1.7 -2.5 GPa.

282

Based on these results, we adopt a range of 6 - 15 GPa for the BPLE for Earth, and 1.7 - 3.8

GPa for Mars. Miyazaki & Korenaga (2022) find a slightly lower but similar pressure of ~5 GPa for the base of the partially molten layer when the surface becomes solid for an Earth-mass planet. We emphasize that further investigation of the factors that control the pressure of last equilibration for magma ocean atmospheres, as well as the timing of separation of atmosphere and interior will be necessary to further our understanding of the early atmospheres of rocky planets.

289 290

#### 291 2.5 Ferric iron partitioning

Our chemical models of fractional crystallization allow the oxygen fugacity of the magma ocean 292 to evolve as a result of changes in the ferric to ferrous iron ratio  $(Fe^{3+}/Fe^{2+})$  in the silicate melt. 293 This ratio evolves as a result of differences in the partitioning behavior of the two iron oxide 294 species. To study this, we modify the model of Elkins-Tanton (2008) by adding ferric iron 295 partitioning between the silicate minerals and melt. The ferric iron partition coefficients used for 296 each mineral along with estimated uncertainties are given in Table 3. Models have shown that 297 the bulk FeO content of the melt phase will increase as the magma ocean crystallizes due to the 298 incompatibility of Fe relative to Mg in silicate minerals (Boukaré et al. 2018; Elkins-Tanton et 299 al. 2003). Fe/Mg partition coefficients have been updated from previous models (see Section 300 **2.3**) and are assumed to represent  $Fe^{2+}$  only. Ferric iron also behaves as a moderately 301 incompatible element in most mineral/melt systems (Canil and O'Neill, 1996). Bulk partition 302 303 coefficients for ferric oxide in peridotite/melt systems under upper mantle conditions are typically found to be between 0.1 - 0.3 (Canil & O'Neill 1996; Cottrell & Kelley 2011; 304 Sorbadere et al. 2018). Cottrell and Kelley (2011) calculated a bulk Fe<sub>2</sub>O<sub>3</sub> partition coefficient of 305 0.22 for fertile mantle composed of 55% olivine, 25% orthopyroxene, 18% clinopyroxene and 306 2% spinel using the partition coefficients of Mallmann and O'Neill (2009). The comparable 307 value of the bulk mantle/melt partition coefficient for FeO using the same mineralogy is 0.85, 308 indicating that ferrous iron behaves more compatibly than ferric iron. Therefore, our expectation 309

is that the  $Fe^{3+}/Fe^{2+}$  ratio of the magma ocean should increase as the magma ocean crystallizes,

although this depends in detail on the crystallizing phases and their proportions. In particular, the

ferric iron abundance and partitioning behavior in the lower mantle minerals are largely

unknown, but they have been shown to have a high capacity to hold ferric iron under current

- 314 mantle conditions in mineral physics experiments.
- 315

316 For most of the minerals in our model, we assume a constant mineral/melt partition coefficient

derived from experiments. It is likely that ferric iron partition coefficients depend on other

factors, including pressure, temperature, oxygen fugacity, melt fraction, electronic spin state

and/or bulk composition. However, there is insufficient experimental data to robustly constrain

these dependencies for any major mantle phase beyond the upper mantle. We consider the ferric

iron partition coefficients presented here to be the best current achievable estimate, but we

emphasize the need for additional experimental measurements in order to better constrain theevolution of magma ocean redox state.

324

In Section 3, we will explore how the uncertainty in the ferric iron partition coefficients may alter the results of the calculations and the implications for the evolution of the earliest

- atmospheres of Earth and Mars. Below, we briefly discuss the partition coefficients for the
- phases included in this model, and the typical  $Fe^{3+}$  abundance observed in mantle samples where
- available. Additional literature review of ferric iron partition coefficients is available in **Text S1**.
- 330
- 331 2.5.1 Upper mantle assemblage (ol, opx, cpx, sp)

We halt crystallization of the magma ocean model at 2 GPa, as accumulated compositional fractionation may alter low pressure crystallization sequences, and mass balance becomes difficult to achieve with our prescribed mineralogy. We therefore do not include plagioclase in the following discussion, which only crystallizes at pressures less than 2 GPa.

- 336
- 2.5.1.1 Olivine (ol)  $Fe^{3+}$  is highly incompatible in olivine, and typically has abundances below
- reporting standards in most upper mantle olivines. Many models of partitioning during partial

melting of the upper mantle assume that  $D_{Fe^{3+}}^{ol/melt} = 0$  (e.g. Canil et al. 1994, Rudra & Hirschmann 2022). Mallmann & O'Neill (2009) find a non-zero but very low partition

- Hirschmann 2022). Mallmann & O'Neill (2009) find a non-zero but very low partition coefficient of  $0.0626 \pm 0.0102$  for Fe<sup>3+</sup> into olivine. We adopt a partition coefficient of 0 for olivine, but calculations using the partition coefficient of Mallmann & O'Neill (2009) do not
- 343 produce significantly different results.
- 344
- 2.5.1.2 Spinel (*sp*) Spinel is a major host phase for ferric iron in the upper mantle. Although it is not an abundant mineral, ferric iron is very compatible in spinels. In lherzolite peridotites in the
- present day upper mantle, spinel has  $Fe^{3+}/\Sigma Fe$  of 15-34% (Woodland et al. 2006). Davis &
- Cottrell (2021) conducted peridotite partial melting experiments at 1.5 GPa and over a range of
- $fO_2$ . Using their data along with corrected data from Sorbadere et al. (2018), Davis & Cottrell
- (2021) find that  $D_{Fe^{3+}}^{sp/melt}$  increases with spinel Fe<sub>2</sub>O<sub>3</sub> concentration, independent of  $fO_2$ , and

351 decreases with temperature. We adopt their corrected  $D_{F\rho^{3+}}^{sp/melt}$  equation:

352 
$$ln D_{Fe_2O_3}^{sp/melt} = a \frac{10,000}{T} + b + c ln (C_{Fe_2O_3}^{sp} (wt\%))$$
 (6)  
353 where  $a = 0.87 \pm 0.07, b = -4.6 \pm 0.4$ , and  $c = 0.24 \pm 0.02$ .

- 354
- 2.5.1.3 *Clinopyroxene (cpx)- and Orthopyroxene (opx)* Ferric iron is more compatible in
- clinopyroxene than orthopyroxene, making it an important host phase in the upper mantle. In
- modern day lherzolite peridotites, opx and cpx have  $Fe^{3+}/\Sigma Fe = 2 9\%$  and 3 32%,
- respectively (Woodland et al. 2006). Rudra & Hirschmann (2022) used experiments on an
- and esite composition liquid to measure the  $Fe^{3+}$  partition coefficient for cpx at 1 2.5 GPa. They
- found that it depends on both the  $Al_2O_3$  and  $Fe_2O_3$  concentrations in the clinopyroxene, although
- the fit including  $Fe_2O_3$  gives only marginal improvement over fitting by  $Al_2O_3$  alone, which we prefer for simplicity. This relationship is given by the equation:
- 363

364 
$$D_{Fe^{3+}}^{cpx/melt} = (0.165 \pm 0.047) C_{Al_2O_3}^{cpx} (\text{wt\%}) + (-0.297 \pm 0.108)$$
 (7)  
365

However, this equation gives negative values for the partition coefficient when  $Al_2O_3$  is below 366

1.8 wt%, which is within the range of natural abundances for cpx in garnet peridotites 367

(Woodland 2009). Therefore, we refit this relationship using logarithmic values for both D and 368

 $Al_2O_3$  concentration. We include the three cpx/melt values from Davis and Cottrell (2021), 369

which are at higher Al<sub>2</sub>O<sub>3</sub> concentration, and derive: 370

371

372 
$$\ln D_{Fe^{3+}}^{cpx/melt} = (1.997 \pm 0.640) \ln C_{Al_2O_3}^{cpx} (\text{wt\%}) + (-3.987 \pm 0.965)$$
 (8)

373

We find an  $r^2 = 0.6027$ , compared with  $r^2 = 0.575$  for the equation given in Rudra & Hirschmann 374 (2022) (see Fig. S1).Our model finds  $Al_2O_3$  concentrations of 0.5 - 1.5 wt% in clinopyroxene, 375 with slightly wider ranges in whole mantle models (batch + fractional crystallization) versus 376 377 shallow mantle models (fractional crystallization only). High pressure clinopyroxenes are typically Mg-rich, becoming more Ca-rich as crystallization progresses. Because of the strong 378 dependence on Al<sub>2</sub>O<sub>3</sub> abundance, the cpx partition coefficients are much lower (~0.05) than 379 those determined by Mallmann & O'Neill (2009) of 0.45, which leads to slightly stronger 380 fractionation of ferric iron in the upper mantle. This effect is further propagated by linking the 381 clinopyroxene partition coefficient to those of orthopyroxene and garnet, as discussed below. 382 383

384 The ferric iron partition coefficient of orthopyroxene is poorly constrained by experimental data, especially regarding compositional dependencies. We follow Rudra & Hirschmann (2022) in 385 using the relationship: 386

387

 $D_{Fe^{3+}}^{opx/melt} = D_{Fe^{3+}}^{opx/cpx} \times D_{Fe^{3+}}^{cpx/melt}$ (9) 389

We fit data on ferric iron concentration in opx and cpx in peridotites to a log-normal distribution, 390 which gives  $D_{Fe^{3+}}^{opx/cpx} = 0.70^{+1.56}_{-0.64}$  (Woodland et al. 2006, Canil & O'Neill 1996, Malaspina et al. 391 2012, Nimis et al. 2015, Luth & Canil 1993) (see Fig. S2). We note that this is close to, but 392 somewhat lower than the values of 0.80 (Rudra & Hirschmann 2022, fit to normal distribution) 393 and 0.81 (Davis and Cottrell 2021, constrained by 3 experimental data points). Typical values of 394  $D_{Fe^{3+}}^{opx/melt}$  in our models range from 0.02 - 0.05. 395

396

2.5.2 *Garnet* (gt) 397

Garnet in upper mantle peridotites typically has  $Fe^{3+}/\Sigma Fe = 2 - 7\%$  (Canil & O'Neill 1996, 398 399 Malaspina et al. 2012, Nimis et al. 2015). As with other high pressure phases, little to no information is available on partitioning of ferric iron between silicate melt and garnet, although 400 data is available on partitioning of ferric iron between garnet and pyroxene phases in natural 401 samples (Canil & O'Neill 1996, Woodland et al. 2006, Woodland 2009, Nimis et al. 2015, 402 Aulbach et al. 2022). We therefore again adopt the parameterization of Rudra & Hirschmann 403 (2022), using the relationship: 404 405

406 
$$D_{Fe^{3+}}^{gt/melt} = D_{Fe^{3+}}^{gt/cpx} \times D_{Fe^{3+}}^{cpx/melt}$$
 (10)

407

We fit a log-normal distribution to data from natural peridotites, which yields  $D_{Fe^{3+}}^{gt/cpx} = 0.68^{+1.96}_{-0.51}$ 408 (Woodland 2009, Canil & O'Neill 1996, Malaspina et al. 2012, and Nimis et al. 2015) (see Fig. 409

- **S3**). Typical values of  $D_{Fe^{3+}}^{gt/melt}$  in our models range from 0.02 0.05. We note that this model 410
- does not capture the temperature dependence of gt/cpx partition coefficients noted by others 411
- (Canil and O'Neill 1996, Woodland 2009, Purwin et al. 2013). The garnet/cpx partition 412
- coefficient determined here is biased to sub-solidus temperatures. Because the garnet/cpx 413
- partition coefficient increases with temperature, using a constant value may underestimate the 414
- ferric iron partition coefficient that would be appropriate for a magma ocean system. We discuss 415
- the effect of uncertainties on the garnet partition coefficient further in Section 4.1. 416
- 417
- 418 2.5.3 Majorite (maj), Wadslevite (wad), Ringwoodite (ring)
- Ferric iron is more compatible in the high-pressure phases wadslevite and ringwoodite than in 419 the lower pressure polymorph olivine and becomes increasingly more compatible in the majoritic 420 421 form of garnet at high pressures. However, there are few studies that report measured ferric iron
- content of these minerals along with co-existing minerals like pyroxene or garnets and none with 422 melts. However, minimum ferric iron abundances in the minerals have been measured at metal 423
- saturation. O'Neill et al. (1993) found that the minimum  $Fe^{3+}/\Sigma Fe$  of majorite, wadslevite, and
- 424 ringwoodite coexisting with metallic iron was 0.07, 0.03 and 0.04, respectively. For magma 425
- ocean conditions at or below IW, we expect similar ferric iron contents in these minerals. We use 426
- the silicate melt model of Hirschmann (2022) for a melt that would hypothetically co-exist with 427
- the minerals at these conditions in order to estimate the ferric iron partition coefficient. We use 428
- 429 both the Deng and Armstrong EOS for ferric iron oxide (FeO<sub>1.5</sub>). We report our calculated
- average values in Table 3, with details about the calculation provided in Text S2 and Table S1. 430
- 431
- 2.5.4 Bridgmanite (bg) 432

Boujibar et al. (2016) report Fe<sup>3+</sup>/Fe<sub>T</sub> measurements of bridgmanite specimens under partial 433 melting conditions at 25 GPa and IW -2, with values ranging from 14 - 25%. They also report 434 degree of melting and the bulk oxide abundances of both minerals and the quenched melt phase. 435 We use the measured  $fO_2$  and melt phase abundances to calculate the Fe<sup>3+</sup>/Fe<sub>T</sub> ratio in the silicate 436 melt using the model of Hirschmann (2022) and both the Deng and Armstrong EOS for ferric 437 iron oxide (FeO<sub>1.5</sub>). Additional details are given in the Text S3 and Table S2. We find an 438 average value of  $D_{Fe^{3+}}^{bg/melt}$  of 2.07 ± 0.50 with the Deng EOS and 0.17 ± 0.03 with the 439 Armstrong EOS. The two equations of state predict radically different partitioning behavior of 440 ferric iron in bridgmanite. More recently, Kuwahara & Nakada (2023) report ferric iron partition 441 coefficients between silicate melt and bridgmanite at 23-27 GPa, with values of  $D_{Fe^{3+}}^{bg/melt} =$ 442 0.43 - 0.51. These experiments were at  $fO_2$  conditions slightly below the Re-ReO<sub>2</sub> buffer, with 443 calculated values of IW+1.0 to +1.5 (note that we calculate slightly higher  $fO_2$  than reported in 444 the paper, assuming pure Re metal foil, as alloy compositions were not reported). These 445 experiments, as well as work by Kuwahara et al. (2023), suggest closer agreement with the 446 Armstrong EOS, although Kuwahara et al. adopt different values for the first pressure derivative 447 of the bulk modulus  $\kappa$ ' of both FeO and FeO<sub>1.5</sub> of 4 and 1.4, compared to 8 and 1.3 from 448 Armstrong et al. . In our model below, we prefer our calculated partition coefficients for 449 bridgmanite rather than the more recent measurements by Kuwahara et al. for three reasons: 1) 450 the Boujibar et al. (2016) experiments occur at oxygen fugacity conditions more consistent with 451 predicted magma ocean conditions post-core-formation (e.g. IW-2, Rubie et al. 2015), 2) they 452 allow us to explore a more extreme range of partitioning behaviors in the lower mantle, and 3) to 453 conduct a consistent test of a single EOS's predictions for ferric iron partitioning behavior. We 454

455 explore how the outcomes of our models change when using the partition coefficient of

456 Kuwahara et al. in Section 4.1. There is a clear need for more partitioning experiments at a wider

457 range of experimental conditions in order to better constrain deep mantle behavior.

458

#### 459 2.5.5 Magnesiowüstite (mw)

Magnesiowüstite (or ferropericlase) contains a relatively small amount of ferric iron, ranging 460 from ~0 to 0.1 times the ferrous iron abundance (McCammon et al. 2004a) when in equilibrium 461 with metallic iron. At higher oxygen fugacities, periclase can incorporate more significant 462 amounts of ferric iron. Otsuka et al. (2013) show that the ferric iron abundance in periclase 463 depends strongly on composition (Fe#), oxygen fugacity, and pressure. They find that ferric iron 464 abundances can be large for Fe#  $> \sim 0.2$  at relatively high oxygen fugacities. However, ferric 465 iron abundances are likely negligible (< 1%) for all reasonable Fe#s at pressures greater than ~ 466 24GPa. Measurements of natural ferropericlase inclusions in diamonds from the lower mantle 467 find Fe<sup>3+</sup>/ $\Sigma$ Fe  $\lesssim$  7% (Kaminsky et al. 2015; McCammon et al. 1997, 2004b), consistent with the 468 experimental results. For our calculations, we therefore neglect ferric iron in magnesiowüstite 469 under the assumption that the lower mantle will begin crystallizing at a relatively low ( $\langle IW \rangle$ )  $fO_2$ .

470 471

#### 472 *2.5.6 Testing uncertainties*

473 We report estimated uncertainties for the ferric iron partition coefficients in **Table 3**. To explore

- how these uncertainties would affect the redox evolution of the crystallizing magma ocean, we
- 475 perform Monte Carlo simulations with 1000 random draws from normal distributions (log-
- normal for opx/cpx and gt/cpx) for each of the ferric iron partition coefficients. For
- clinopyroxene and spinel, we draw for each of the fit coefficients from equations (6) and (8). We
  assume all uncertainties are uncorrelated. We found that 100 draws was sufficient to produce
- 478 assume an uncertainties are uncorrelated. We found that foo draws was sufficient to produce 479 stable mean values for  $Fe^{3+}/Fe_T$  and  $fO_2$  but that the ranges of these calculated values were
- substantially smaller than when using 1000 simulations. In order to more accurately estimate the uncertainty on the calculated  $fO_2$ , we used the larger number of simulations. Simulations beyond 1000 did not substantially change means or increase calculated ranges but did substantially
- 483 increase computation time.
- 484

#### 485 2.6 Initial $Fe^{3+}$ abundance

- 486 Hirschmann (2022) calculated ferric iron contents of magma oceans for Earth and Mars at core-
- forming conditions, using different pressure-temperature- $fO_2$  from literature models of trace
- element behavior during core-formation. He found ranges of  $Fe^{3+}/Fe_T = 0.034 0.10$  for Earth
- and 0.026 0.038 for Mars, depending on mantle FeO<sub>T</sub> abundance. We use these as our starting
- range of ferric iron abundance in our magma ocean models. Hirschmann (2022) modified his calculated ferric iron abundance due to  $Cr^{2+}$  oxidation during sub-solidus reactions with Fe<sup>3+</sup> in
- 491 calculated ferric iron abundance due to  $Cr^{2+}$  oxidation during sub-solidus reactions with Fe<sup>3+</sup> in 492 the upper mantle. We neglect this effect here as we are largely focused on the liquid silicate
- 492 the upper manue. We neglect this effect here as we are largely focused on the inquid sincate 493 behavior. We start with nominal abundances of  $Fe^{3+}/Fe_T = 0.10$  for Earth and 0.02 for Mars.
- More recently, Kuwahara et al. (2023) experimentally found a  $Fe^{3+}/Fe_T = 0.2 0.5$  for peridotite
- under metal-saturated conditions at pressures of 23 28 GPa. While these numbers are
- 496 challenging to explain from a mantle redox evolution perspective (Hirschmann 2023), they are
- 497 consistent with a modified Armstrong et al. (2019) EOS for  $FeO_{1.5}$ . We will therefore explore a
- 498 range of initial  $Fe^{3+}/Fe_T$  up to 0.20.
- 499
- 500 2.7 Interstitial Liquids

- 501 Previous magma ocean models have noted that in the crystallization of magma chambers, some
- 502 liquid is often trapped by the crystals that settle to the bottom of the magma chamber. Elkins-
- Tanton et al. (2005) adopted an interstitial liquid volume fraction in each solid layer of 1%.
- Recent models by Hier-Majumder & Hirschmann (2017) & Miyazaki & Korenga (2022) looking
- at the two-phase dynamics of magma ocean crystallization have suggested that the trapped liquid
- fraction may be much higher, especially in the upper mantle, and increase as the rate of solidification increases. Hier-Majumder & Hirschmann (2017) find trapped liquid fractions of
- 20-30 vol% in the upper mantle. This has implications for trapped volatile fraction but will also
- modify the results of our redox-partitioning model explored here. We find the effect to be minor
- 510 in comparison to uncertainties on the ferric iron partition coefficients. We use a nominal value of
- 511 20% in our fractional crystallization models but discuss results with as little as 1%. Effects of
- 512 interstitial fluid on compositional fractionation are minimal for pressures higher than 5 GPa.
- 513

## 514 **3 Results**

- 515 In the following section, we discuss the oxygen fugacity evolution of the magma oceans of Earth
- and Mars, with a focus on matching the constraints for the D/H ratios of the last magma ocean
- atmospheres at the BPLE. We show results for different initial  $Fe^{3+}$  abundances and consider the
- different magma ocean depths and crystallization scenarios discussed in Section 2.6 and Figure
- **1**. The evolution of the oxide abundances within the solid and liquid phases largely matches the
- results of previous work (Elkins-Tanton et al. 2003, 2005, Elkins-Tanton 2008), with increasing
- 521 concentration of incompatible oxides such as FeO, CaO, and Al<sub>2</sub>O<sub>3</sub> and decreasing 522 concentrations of compatible oxides including MgO and SiO<sub>2</sub> in the late-stage liquid. We
- provide a brief description of these results in **Supplemental Text S4** but focus here on the new results related to  $Fe^{3+}$  partitioning and oxygen fugacity.
- 524 525

## 526 *3.1 Fe<sup>3+</sup> Evolution of the Terrestrial Magma Ocean*

- The Deng and Armstrong EOSs produce opposite ferric iron partitioning behavior in the lower mantle, with the Deng EOS yielding compatible behavior ( $D_{Fe^{3+}}^{min/melt} > 1$ ) and the Armstrong EOS yielding incompatible behavior ( $D_{Fe^{3+}}^{min/melt} < 1$ ) (see **Table 3**). For the Deng EOS, ferric iron concentrations therefore decrease during crystallization of the lower mantle and transition zone, whereas ferric iron concentrations increase for the Armstrong EOS. Within the upper mantle, ferric iron partition coefficients are identical and Fe<sup>3+</sup> concentrations generally increase due to bulk partition coefficients being much less than 1.
- 535

The oxygen fugacity of a silicate melt can be calculated using the ratio of  $Fe^{3+}$  to  $Fe^{2+}$  (Kress & Carmichael, 1991, O'Neill et al. 2006, Jayasuriya et al. 2004, Zhang et al. 2017, Armstrong et al.

2019, Sossi et al. 2020, Deng et al., 2020). Here we use the recently recalibrated equation of

Hirschmann (2022) to calculate the oxygen fugacity at the surface of the magma ocean, which is
 relevant for the outgassed atmosphere. This equation depends on the pressure, temperature and

- 541 oxide abundances of the silicate liquid as follows:
- 542

$$\log\left(\frac{X_{FeO_{1.5}}}{X_{FeO}}\right) = a\log f_{O_2} + b + \frac{c}{T} - \frac{\Delta C_p}{R\ln 10} \left[1 - \frac{T_0}{T} - \ln\left(\frac{T}{T_0}\right)\right] - \frac{\int_{P_0}^{P} \Delta V dP}{RT\ln 10} + \frac{1}{T} \left[Y_1 X_{SiO_2} + Y_2 X_{TiO_2} + Y_3 X_{MgO} + Y_4 X_{CaO} + Y_5 X_{NaO_{0.5}} + Y_6 X_{KO_{0.5}} + Y_7 X_{PO_{2.5}} + Y_8 X_{SiO_2} X_{AlO_{1.5}} + Y_9 X_{SiO_2} X_{MgO}\right]$$
(11)

where parameter values are given in **Table 4**, T is temperature in K, P is pressure in GPa, and  $X_i$ 543 are mole fractions of the oxides in the silicate melt. We calculate the temperature at the surface 544 of the magma ocean by assuming a one-phase magma ocean adiabatic profile anchored by the 545 solidus temperature at the base of the magma ocean. We assume a pressure of 1 bar at the 546 547 surface, so that the  $\Delta V dP$  term is negligible. Models suggest that magma oceans may have atmospheres of 100 – 1000 bars (e.g. Hamano et al. 2013, Lebrun et al. 2013), but this produces 548 a very minor pressure effect that we neglect here. We also utilize the evolving oxide mole 549 fractions determined with the fractional crystallization model (see Figure S8) in equation (11). 550 This calculation assumes that the composition of the silicate liquid remains homogeneous 551 throughout the liquid layer due to rapid convection and therefore that the relative concentrations 552 of  $Fe^{3+}$  and  $Fe^{2+}$  and the other oxides are the same at both the surface and the base of the magma 553 ocean. Note that we prefer the relationship between  $Fe^{3+}/Fe^{2+}$  and  $fO_2$  from Hirschmann (2022) 554 rather than Sossi et al. (2020) because of the dependence on the oxide mole fractions in the 555 silicate melt, which deviate from the peridotite composition measured by Sossi et al. after 556 fractional crystallization. 557

558

Figure 2 shows the evolution of the oxygen fugacity of outgassing for the whole mantle magma 559 ocean models, with either fully fractional crystallization (*right*, 'whole Earth') or a switch from 560 batch to fractional crystallization occurring at 500 (*left*) or 1000 km (*middle*). The base of the 561 magma ocean evolves from right to left in these figures, with time from start of crystallization 562 following in the same direction, although note that we do not specifically model the timing of 563 solidification. For these calculations, we use a nominal initial  $Fe^{3+}/Fe_T = 0.10$ . This value is the 564 upper limit of Fe<sup>3+</sup>/Fe<sub>T</sub> determined for core-formation conditions by Hirschmann (2022). In 565 Figure 2, we plot all 1000 draws of different ferric iron partition coefficient values in grey and 566 the average of the 1000 simulations in black. The calculations shown in the top row use partition 567 coefficients for the lower mantle and transition zone calculated with the Deng EOS, whereas the 568 bottom row uses those derived with the Armstrong EOS. The pink horizontal shaded region 569 570 shows the allowed  $fO_2$  of the last magma ocean atmosphere from D/H constraints (Pahlevan et al. 2019) and the grav vertical shaded region shows our constraints for the BPLEBPLE from 571 572 Section 2.4; the overlap between pink and gray areas represent valid solutions.

573

574 The oxygen fugacity evolution closely follows the ferric iron content of the magma ocean.

575 Therefore the Deng EOS models produce a decreasing  $fO_2$  as pressures drop from 120 GPa to 18

576 GPa (**Fig. 2**, *right* and *middle*) due to crystallization of bridgmanite, majorite, ringwoodite and 577 wadsleyite, whereas the Armstrong EOS produces increasing  $fO_2$  across the same region. The

578 Armstrong EOS produces a smaller spread in solutions because of the smaller absolute value and

therefore smaller uncertainty on the bridgmanite partition coefficient compared to the Deng EOS

(note that we treat both uncertainties as a normal distribution, truncated at zero). For both EOS

models, the 500 km magma ocean shows increasing  $fO_2$  as the magma ocean crystallizes up to

pressures of ~5 GPa and a relatively narrow range of possible solutions, but note that the Armstrong EOS model starts at much higher  $fO_2$  because of the incompatible behavior of Fe<sup>3+</sup>

during the batch crystallization stage. In contrast, the  $fO_2$  of both the 1000 km and whole Earth

585 models using Deng EOS partition coefficients initially decrease before increasing at pressures

586 less than ~20 GPa. This is due to the compatible behavior of ferric iron in the transition zone and

less than ~20 GPa. This is due to the compatible behavior of ferric iron in the transition zone and lower mantle minerals, unlike the upper mantle. At pressures lower than 5 GPa,  $Fe^{3+}$  in the liquid

and  $fO_2$  of outgassing decreases due to a) increase in  $D_{Fe^{3+}}^{cpx/melt}$  because of increasing Al<sub>2</sub>O<sub>3</sub>

contents in the cpx phase, and b) the eventual crystallization of spinel, which has a large

- 590  $D_{Fe^{3+}}^{sp/melt}$  value.
- 591

Only 7 of the 3000 realizations (all for the whole Earth fractional model) of Deng EOS models 592 shown in Fig. 2(right) produce results that agree with the combined D/H and BPLE bounds. In 593 594 contrast, all 3000 realizations of the Armstrong EOS models produce valid solutions. The successful Deng EOS models have an average bridgmanite partition coefficient of 1.00 in 595 comparison to the average for all of the whole Earth Deng EOS models of 1.99. Taken at face 596 value, the D/H constraints suggest that the Armstrong EOS produces more realistic ferric iron 597 partition coefficients, and therefore that ferric iron behaved incompatibly throughout the entire 598 magma ocean. 599

600

However, it is possible that the initial ferric iron content of the magma ocean was either higher or lower than our nominal value and that the Deng EOS could produce results consistent with D/H

with different initial conditions. **Figure 3** shows only the mean oxygen fugacities for the three

different whole mantle configurations with different starting  $Fe^{3+}/Fe_T$  from 0.04 up to 0.20. For the Deng EOS models, initial  $Fe^{3+}/Fe_T$  of 0.04 and 0.14 do not produce valid solutions to the

the Deng EOS models, initial  $Fe^{3+}/Fe_T$  of 0.04 and 0.14 do not produce valid solutions to the D/H constraints. For the initial  $Fe^{3+}/Fe_T = 0.20$ , we find that 993, 47, and 249 realizations out of

1000 match the 500 km, 1000 km and whole Earth models, respectively (see Table S5). In
 contrast, effectively all realizations of all Armstrong EOS models match the D/H constraints

609 within the BPLE bounds. The Armstrong EOS model only fails if  $Fe^{3+}/Fe_T$  is less than 0.04.

610 However, we note that for  $Fe^{3+}/Fe_T > 0.10$ , the Armstrong EOS models predict  $fO_2$  greater than

the present day upper mantle, which are likely too high to be valid. Recent experimental work by Kuwahara et al. (2023) suggests that initial mantle  $Fe^{3+}/Fe_T$  was 0.2-0.50 and favors the

Kuwahara et al. (2023) suggests that initial mantle  $Fe^{3+}/Fe_T$  was 0.2-0.50 and favors the Armstrong EOS over the Deng EOS, although with modified κ' for both FeO and FeO<sub>1.5</sub>. We

find these conclusions contradictory with the extreme redox evolution of the models using the

615 Armstrong EOS developed here.

616

To find whether there are any configurations under which the Deng EOS predicts valid solutions, 617 we additionally consider shallow magma oceans, by beginning the 500 km and 1000 km models 618 with a BSE composition (see **Fig.1** (*right*)). Shallow magma oceans may be generated by smaller 619 620 impactors in which only a portion of the mantle is melted. Alternatively, some models suggest that a magma ocean may solidify from the middle of the mantle, with a basal magma ocean 621 crystallizing much more slowly than the upper mantle portion of the magma ocean (Labrosse et 622 al. 2007). Our shallow magma ocean models could then be interpreted as simulating the upper 623 mantle portion of such a layered crystallization model. By using BSE as the starting composition, 624 we assume that the mantle is homogenized prior to settling of crystals to the starting depth (either 625 626 500 or 1000 km), followed by upward fractional crystallization. We plot the averages of these realizations as dashed lines in Figure 3. These realizations are much more successful at 627 matching the D/H constraint for the Deng EOS because the BSE has a higher starting  $Fe^{3+}/Fe_T$ 628 than the batch crystallization models. For the 500 km partial magma ocean, 0, 989 and 1000 629 realizations match for initial  $Fe^{3+}/Fe_T = 0.04, 0.10$ , and 0.14. For the 1000 km magma ocean, 0, 630 620 and 976 realizations are successful for the same initial Fe<sup>3+</sup>/Fe<sub>T</sub>. Bulk partition coefficients 631 632 are similar (see Figure S5) between the whole mantle and shallow magma ocean models, with outcomes differing due to the differences in ferric iron content of the magma ocean when 633

634 fractional crystallization begins.

We also run simulations with different amounts of interstitial liquid, as discussed in Section 2.7. 636

Our nominal simulations use 20% interstitial fluid. We neglect the compositional evolution of 637 the interstitial fluid as it crystallizes in situ. Smaller amounts of interstitial fluid generate slightly 638

more compositional fractionation, with differences becoming significant at pressures less than 5 639

GPa. These models produce steeper increases of  $Fe^{3+}$  in the liquid and therefore higher oxygen 640

- fugacities for outgassing. However, since the BPLEBPLE for Earth is at pressures higher than 5 641
- GPa, interstitial fluid abundance does not affect our conclusions regarding the oxygen fugacity at 642
- the time that the atmosphere and magma ocean begin to evolve separately. 643 644
- 3.2 Fe<sup>3+</sup> Evolution of a Martian Magma Ocean 645
- 646 Our nominal model for Mars used initial  $Fe^{3+}/Fe_T = 0.02$ , slightly lower than the range of values 647 calculated for core formation conditions by Hirschmann (2022) of 0.026 – 0.038 (see Figure 648 **S6**). The D/H constraint for Mars (Pahlevan et al. 2022) is a maximum  $fO_2$  of IW-1 for the last 649
- magma ocean atmosphere. All of the whole mantle magma ocean models produce monotonically 650
- increasing values for  $Fe^{3+}/Fe_T$  and  $fO_2$  of outgassing with decreasing pressure, due to 651
- incompatible Fe<sup>3+</sup> behavior throughout most of the Martian mantle. Because there is only a small 652
- portion of the Martian mantle containing high pressure phases, the Deng and Armstrong EOS 653
- models produce nearly identical results within ~0.5 log units of  $fO_2$  of each other, unlike the 654
- Earth models, although the Deng models are modestly more successful, especially for the whole 655 mantle fractional crystallization model. For calculations using our nominal ferric iron abundance, 656
- most models produce oxygen fugacities of IW to IW+1 at 2 GPa. Only 265, 89, and 875 out of 657
- 658 1000 simulations are successful for the three Deng EOS models, but 0, 2, and 171 for the
- Armstrong models. The successful simulations have bulk ferric iron partition coefficients 659
- roughly an order of magnitude higher than the mean in the 2.5-15 GPa region, where olivine, 660
- orthopyroxene, clinopyroxene and garnet all form. Reducing the initial ferric iron content of the 661 planet produced more successful results for both EOS models. 662
- 663
- In **Figure 4**, we show simulations for a lower initial  $Fe^{3+}/Fe_T = 0.01$  for the three whole mantle 664 magma ocean models, which provides a better fit to the constraints. This initial ferric iron 665 content is substantially lower than predicted values but produces 100% successful outcomes for 666 all three whole mantle batch crystallization models for both EOSs. Successful models have 667 oxygen fugacities lower than IW-1 somewhere within the predicted BPLEBPLE range, although 668
- not necessarily at the lowest pressure. In Figure 5, we compare mean values for 1000 669
- simulations for the whole mantle and shallow magma ocean models with different initial
- 670  $Fe^{3+}/Fe_T$ . 671
- 672
- 673 The shallow magma oceans, which start from the Bulk Silicate Mars (BSM) composition (see dashed lines for 500 and 1000 km models), diverge more strongly for the 500 km models at low 674
- pressure, and more so for the Armstrong EOS than the Deng EOS. As for the Earth, these models 675 simulate either shallower magma oceans, or ones that produce a neutrally buoyant cumulate pile 676
- at midmantle depth that isolates the upper mantle from the lower mantle. For the 1000 km 677
- models, evolution is nearly identical to the whole mantle crystallization case, with the  $Fe^{3+}/Fe_T =$ 678
- 0.01 model producing 100% successful simulations, whereas with  $Fe^{3+}/Fe_T = 0.02$ , there are only 679
- ~1% successful simulations for both EOSs. In comparison, the 0.02 model for the 500 km partial 680
- magma ocean produces 100% successful simulations for both EOSs. Changes in interstitial fluid 681

- 682 produce larger increases in oxygen fugacity as pressure decreases, so there are fewer successful
- realizations for each magma ocean model. However, mean values for  $Fe^{3+}/Fe_T = 0.01$  with both 20% and 1% intersticial fluid are successful for all three magma access denths
- 20% and 1% interstitial fluid are successful for all three magma ocean depths.
- Based on the D/H constraints within the BPLE, the initial ferric iron content of Mars may have 686 been lower than calculated from core-formation conditions. This could be achievable through 687 less efficient equilibration between the mantle and core or continuous accretion of metal-bearing 688 building blocks rather than a single-stage core formation scenario that sets the magma ocean 689 redox state. Alternatively, a shallow, 500 km magma ocean with higher initial  $Fe^{3+}/Fe_T$  would 690 also match the observed D/H constraints. Another possible solution is that our calculated BPLE 691 is too low and the separation between atmosphere and magma ocean occurs at higher pressures, 692 where oxygen fugacities are lower. This would be possible if the liquidus offset was smaller than 693 the 300-400 K that we assumed, or if the solidus were steeper than the Elkins-Tanton (2008) 694 model. 695
- 695 696
- Bulk ferric iron partition coefficients are shown in **Figure S7**. The ferric iron partition
- coefficient does not change significantly for different initial Fe<sup>3+</sup>/Fe<sub>T</sub>, and bulk partition
- coefficients are generally the same for all of the whole mantle (batch + fractional crystallization)
- models for a given mineralogical layer. Partition coefficients are greater than 1 at pressures
- above 15 Gpa for the Deng EOS models, where wadsleyite, ringwoodite and majorite form, but
- slightly below 1 in this pressure region for the Armstrong EOS models. In the upper mantle, the
   2.5-15 Gpa layer contains olivine, orthopyroxene, clinopyroxene and garnet. Partition
- coefficients are largely determined by  $Al_2O_3$  content in clinopyroxenes, which is low. A spike in
- the bulk partition coefficient at low pressure occurs where spinel begins to form. Ferric iron is
- compatible in spinel, but the bulk partition coefficient remains less than 1 because of the
   relatively small phase proportion of spinel.
- 709 **4. Discussion**
- 710

## 711 *4.1 Ferric iron partition coefficient uncertainties*

- Ferric iron mineral-melt partition coefficients for several critical mineral phases remain mostly
  unmeasured. We have estimated those quantities here to the extent possible, but a high degree of
- uncertainty remains. In particular, garnet plays a key role in the redox budget of the upper
- mantle, but there have been no direct measurements of ferric iron mineral-melt partition
- coefficients. There is only one direct measurement of the ferric iron partition coefficient
- 718 (Kuwahara and Nakada 2023) of bridgmanite and no direct measurements of ferric iron partition
- coefficients for any of the other lower mantle minerals (majoritic garnet, wadsleyite,
- ringwoodite). In addition, the pressure, temperature and compositional dependencies of ferric
- iron partition coefficients would be valuable to improve our understanding of the redox evolution
- 722 of the Earth's mantle.
- 723
- **Figures 2 & 4** show the effect of the current estimated uncertainties of the current ferric iron
- partition coefficient estimates on the oxygen fugacity calculations. Both the 1000 km and whole
- mantle models for the Deng EOS have much larger uncertainties on the  $fO_2$  evolution than for the 500 km model. This is because the uncertainties on the ferric iron partition coefficients of the
- 127 the 500 km model. This is because the uncertainties on the ferric non partition coefficients (
   108 lower mantle minerals allow both incompatible and compatible behaviors, which produce a

- vider spread of behaviors. In contrast, ferric iron partition coefficients of the upper mantle are
- very small and absolute uncertainties are therefore also much smaller than the lower mantle,
- producing a narrower range of outcomes. However, this likely does not capture the full
- uncertainties of partitioning because we do not include the uncertainty of the major oxide
- partitioning, which effects cpx, and therefore opx and gt, partition coefficients.
- The Armstrong EOS produces nearly compatible behavior for ringwoodite, wadsleyite, and 735 majorite, whereas the Deng EOS predicts strongly compatible behavior for these minerals. The 736 two EOS predict more contrasting behavior for bridgmanite (~2 vs. ~0.2), but the calculated 737 uncertainty for the Armstrong EOS led us to only explore a narrow range of strongly 738 incompatible states, as opposed to the moderately incompatible behavior shown by Kuwahara & 739 Nakada (2023). To explore how our model results would be affected if these minerals (bg, ring, 740 maj, wad, as well as gt) had ferric iron partition coefficients that produced different compatibility 741 behavior, we run simulations of the whole Earth magma ocean with lower values of ferric iron 742 743 partition coefficients (0.50  $\pm$  0.50 for bg (based on Kuwahara & Nakada 2023), 0.50  $\pm$  0.50 for ring and maj,  $0.25\pm0.25$  for wad) for all of the lower mantle minerals and a higher value for 744 garnet (1.75  $\pm$  0.75). We run each of these changes individually to determine which partition 745 coefficient has the strongest effect on the  $fO_2$  evolution (Figure 6a), otherwise using the Deng 746 EOS ferric iron partition coefficients. We find minimal effects of changing the partition 747 coefficients of ringwoodite, wadsleyite, majorite and garnet. However, bridgmanite has a very 748 strong effect on the  $fO_2$  evolution. Within the BPLE region, average  $fO_2$  are 3 log units higher 749 than the nominal model, approximately the same as the Armstrong EOS results, and match the 750 D/H constraints on  $fO_2$  very well. This effect is due to the very large volume fraction which 751 bridgmanite occupies in the whole Earth model (~70 vol% of whole mantle), whereas all of the 752 transition zone minerals combined occupy only ~8 vol%. Garnet has similarly low volume 753 fraction (~1.5 vol%) and therefore has only a minor effect. Therefore this test confirms that the 754 primary difference between the Deng and Armstrong EOS results is due to the radically different 755 ferric iron partition coefficients these two EOS predict for bridgmanite. 756
- 757

We vary the same ferric iron partition coefficients for the whole Mars model, shown in **Fig. 6b**. 758 759 We find a much smaller effect on the magma ocean oxygen fugacity for Mars. Increasing the garnet partition coefficient causes a drop in the final oxygen fugacity of ~0.5 log units, whereas 760 decreasing all of the lower mantle mineral partition coefficients together increases the final 761 oxygen fugacity ~0.5 log units. Garnet occupies ~6 vol% of the Martian mantle, whereas the 762 763 combined lower mantle minerals (ringwoodite, majorite, wadsleyite) occupy ~ 21%. Note that variations for Mars are much smaller than for Earth because there is no bridgmanite present. 764 Overall, we find that the largest improvements in the Earth and Mars models would come from 765 more constraints on the ferric iron partition coefficient of bridgmanite and garnet under 766 conditions appropriate to a magma ocean. 767

768769 *4.1.1 Bridgmanite* 

Since bridgmanite's mineral-melt partition coefficient has the largest effect on the Earth model
automage it is worth discussing in more detail. The shundance of ferrie ince in hidemonite

- outcomes, it is worth discussing in more detail. The abundance of ferric iron in bridgmanite
- under current mantle conditions has been shown to depend on Al content, oxygen fugacity,
- temperature, and total Fe content (Huang et al. 2021). In addition, ferric iron content seems to vary with depth in the modern mantle (Kurnosov et al. 2017, Mashino et al. 2020), likely due to

changes in substitution mechanisms at different pressures (Shim et al. 2017), which may be 776 caused by changes in the spin state of  $Fe^{3+}$  on the B site (Lin et al. 2013). Shim et al. (2017) find 777 that the  $Fe^{3+}/Fe_T$  of bridgmanite may drop to low values (~0.10) between 1,000 and 1,700 km, 778 779 with higher values (>0.50) at both higher and lower pressures. Our nominal whole Earth model  $(Fe^{3+}/Fe_T = 0.10)$ , which assumes both constant ferric iron partition coefficient as well as Fe/Mg 780 exchange coefficient, produces bridgmanite with  $FeO_T = 4 - 6$  wt% (Deng EOS) or 3-6% 781 (Armstrong EOS) increasing at lower pressures, whereas  $(Fe^{3+}/Fe_T)_{hg}$  decreases from ~0.40 at the 782 core mantle boundary to ~0.10 at 22 GPa for the Deng EOS, but only changes from 0.05 - 0.06783 for the Armstrong EOS. The variations with depth are due to the enrichment of the melt with 784 FeO and relative depletion/enrichment of  $FeO_{1.5}$ . Although we can artificially impose a decrease 785 786 in the ferric iron partition coefficient for the depths suggested by Shim et al. (2017), we find that doing so produces a minimal change in the oxygen fugacity evolution. For the Armstrong EOS, 787 we find maximum values of  $(Fe^{3+}/Fe_T)_{bg}$  of 0.11 when the initial bulk  $(Fe^{3+}/Fe_T) = 0.20$ . 788 Achieving higher values of (Fe<sup>3+</sup>/Fe<sub>T</sub>)<sub>bg</sub> of 0.50 would require initial bulk (Fe<sup>3+</sup>/Fe<sub>T</sub>) up to or 789 greater than 0.50, which would predict final outgassed atmospheres with  $fO_2 = IW+7 - IW+9$ , 790 well above present day upper mantle oxidation state (QFM ~IW+4) and high enough that 791 792 molecular  $O_2$  gas would be significant in the outgassed atmosphere. We find this an implausible result, given evidence that the Archean mantle was 1 to 2 log units more reduced than the present 793 day (Aulbach & Stagno 2016, Nicklas et al. 2019, Gao et al. 2022, Stagno & Aulbach 2021). 794 795 However, note that the late accretion of small reduced planetesimals throughout the Hadean and late Archean may have delivered sufficient metal to reduce the upper mantle to the observed 796 Archean values (Pahlevan et al. 2019, Kuwahara & Nakada 2023). 797

798

The thermodynamic model of Huang et al. (2021) shows that at a given pressure,  $Fe^{3+}/Fe_T$  in

bridgmanite increases with decreasing temperature. Therefore, as the lower mantle cools below the solidus,  $Fe^{3+}$  should become more stabilized in bridgmanite, and additional

disproportionation is likely to occur to increase the  $Fe^{3+}/Fe_T$  of the lower mantle. This is further

supported by evidence that a metallic phase could be present within the modern day mantle

804 (Frost et al. 2004, Rohrbach et al. 2007, Rohrbach & Schmidt 2011, Zhang et al. 2016).

Therefore, we should not expect our magma ocean model to predict present-day  $Fe^{3+}$  abundances in bridgmanite. Pressure or temperature-dependence of the ferric iron partition coefficient or

later thermochemical modification or mixing may account for the differences between our

predictions for the magma ocean vs. present day mantle. However, a better understanding of how

809 much disproportionation has occurred over the planet's lifetime is necessary to permit better

modeling of the redox evolution during both the magma ocean stage and subsequent evolution.

811

812 *4.2 Redox effects not included* 

813

In this paper, we have explored only the crystal-liquid chemistry of a dry, volatile-free magma ocean, and so these models are strictly only applicable to volatile-depleted planets. However, it is likely that the final magma oceans of both the Earth and Mars contained nearly their full complement of planetary volatiles. Recent experimental works have shown that all of the major volatile elements (H, C, N, S) can be highly soluble in core-forming alloy fluids under various conditions (Hirschmann 2016, Dalou et al. 2019, Grewal et al. 2019). Following the segregation

of the core, volatiles within the magma ocean will partition between the atmosphere and silicate

liquid based on their solubilities to form the earliest atmospheres (e.g. Elkins-Tanton 2008).

822 Carbon and sulfur may exsolve as separate solid/liquid species, including carbides, sulfides,

graphite, and diamonds (Grewal et al. 2020, Gaillard et al. 2022). The compositions of secondary 823 phases will depend strongly on the redox. Redox reactions between volatile elements and Fe-824 bearing species may also play a role in both the overall redox balance and the final composition 825 of the system (Gaillard et al. 2014, 2022) as well as long term evolution (Hirschmann 2023). For 826 instance, Genda & Ikoma (2008) and more recently Young et al. (2023) suggest that all of 827 Earth's water could be produced by redox reactions between solar nebular H<sub>2</sub> gas with FeO-828 bearing silicate liquids within the magma ocean. Similar models have also been applied to 829 exoplanets (Kite & Schaefer 2021, Schlichting & Young 2022). While these volatile-driven 830 redox mechanisms are likely important for the overall redox evolution, incorporating them into 831 the model presented here would require a more complete thermodynamic model for reactions 832 833 between volatiles and iron-bearing species in both the melt and each mineral species throughout the full pressure range of the Earth's mantle, which is beyond the scope of this paper. 834 835 We note that additional redox evolution is likely to happen in the crystallizing mushy upper 836 mantle layers, including post-solidus evolution due to  $Cr^{2+}$  oxidation by  $Fe^{3+}$  reduction 837 (Hirschmann 2022), which we do not model here. Other redox sensitive elements such as Ti and 838 V are also likely to play a role in the redox evolution as well, but are relatively low in abundance 839 and are likely to produce more second-order effect. Given the large uncertainties in the behavior 840 of ferric iron alone, the additional complexity of including these minor elements is not warranted 841 at this time. 842 843 Sub-solidus reactions as well as crystallization of trapped interstitial melts are also likely to 844 modify the Fe<sup>3+</sup>/Fe<sub>T</sub> of the mantle past the point where our models halt, so we do not expect our 845

models to produce a direct match to the present day mantle. In addition, post-magma ocean
mantle overturn and solid state convection will redistribute ferric and ferrous iron throughout the
modern day mantle. Recent models suggest that, chondritic D/H values notwithstanding, solid
state convection may initiate within the solidified cumulate layer below a liquid layer, even
before full solidification, which may erase some of the density stratification generated by magma
ocean crystallization (Maurice et al. 2017, Ballmer et al. 2017, Boukare et al. 2018, Morison et
al. 2019, Bolrao et al. 2021, Sharp et al. 2013).

853

Because many of our models predict oxygen fugacities that are well below IW, many of our 854 models may be metal-saturated, especially for low initial Fe<sup>3+</sup>/Fe<sub>T</sub> values for the Earth models. 855 While the initial Fe<sup>3+</sup>/Fe<sub>T</sub> values are set by metal-silicate partitioning (Hirschmann 2022) under 856 different conditions, removal of Fe<sup>3+</sup> from the liquid by mineral/melt partitioning forces the 857 liquid phase into below metal saturation. Therefore additional Fe<sup>3+</sup> and metal production may 858 occur during the magma ocean crystallization. We choose not to model this process here in the 859 interest of minimizing the number of free parameters in the model, but in future work, we will 860 explore how much additional disproportionation may occur as a result of crystallization. 861

862

*4.4 Atmosphere evolution beyond the rheological transition* 

864
865 Using the thermal evolution model from Section 2.4, we examine the rate and timing of
866 outgassing that occurs following the lid formation to estimate how quickly post-magma ocean
867 outgassing will modify the last magma-ocean-equilibrated atmosphere. Unlike the BPLE, the
868 timing of outgassing is sensitive to the absolute values of the solidus and liquidus temperatures,
869 not just their offset. For dry solidus temperatures (Pierru et al. 2022), solid state outgassing for

the Earth can increase the atmospheric mass by 10% in 2-14 Myr, depending on initial volatile 870 budget. However, if the solidus is depressed by 200 K, then it takes 200-300 Myr to see a similar 871 increase in atmospheric mass. Bower et al. (2019) similarly find that outgassing pauses for ~100 872 Myr following solidification of the surface for an Earth-mass planet using a mixing-length 873 thermal evolution model. The long pause before outgassing resumes suggests that the last magma 874 ocean atmosphere remains unmodified by outgassing for a sufficient length of time for D/H 875 fractionation to occur, such that the D/H constraint (Pahlevan et al. 2019) for Earth is a true 876 probe of the oxygen fugacity of the magma ocean. We note that the Sun is most active and 877 capable of driving atmospheric escape for the first 100 Myr after formation, so this lag in 878 outgassing may distinctly separate a last magma ocean atmosphere from outgassing occurring in 879 880 the mush mode. Recent work by Miyazaki & Korenaga (2022) suggests that upward melt transport via percolation may enhance early outgassing. However, Monteux et al. (2020) showed 881 that the remaining melt layers may take 500 - 800 Myr to fully crystallize. We stress the need for 882 additional dynamical magma ocean models to further assess the behavior of volatile transport 883 during this time period. 884

885

For Mars, we find continued outgassing can increase the atmospheric mass by 10% within  $\sim 0.5$ 886

Myr when using the dry solidus (Duncan et al. 2018), but may take as much as 150-170 Myrs 887 when the dry solidus is depressed by 200 K. Given the fast formation time of Mars (< 10 Myr, 888 Dauphas & Pourmand 2011), solid-state outgassing may initiate well within the early active 889 period of the Sun. Therefore, the lag in outgassing fluxes after surface solidification may not 890 robustly separate the last magma ocean atmosphere from subsolidus outgassing, so it is possible 891 that the D/H record for Mars may probe an atmosphere combined of both last magma ocean 892 atmosphere and early sub-solidus outgassing.

893 894

#### **5** Conclusions 895

896

In this paper, we show that it is possible to match the the D/H ratios of both the Earth and Mars 897

through fractional crystallization of a magma ocean without the need for exogenous processes. 898

899 We explore the redox evolution of magma oceans due to the different crystallization behaviors of ferric and ferrous iron. 900

901

We also constrain the pressure at the base of the magma ocean at the time of last atmosphere-902 mantle equilibration (BPLE). Magma ocean thermal evolution models predict a spread of 903 pressures that depends sensitively on the gap between solidus and liquidus temperatures. 904 905 Fractional crystallization and volatile trapping in melt will both act to lower solidus and liquidus temperatures, although it remains unclear if the effect on both is equal. More experimental data 906 on residual liquid compositions, including both solidus and liquidus temperatures, are needed to 907 better constrain the behavior of the late-stage magma ocean-atmosphere system, and subsequent 908 outgassing. Further exploration of the dynamics of crystallization and melt sequestration during 909 the late stage magma ocean is also needed to better understand what constitutes the last magma 910 911 ocean atmosphere.

912

Hirschmann (2022) has calculated from first principles what the initial ferric iron content of 913

- Earth (Fe<sup>3+</sup>/Fe<sub>T</sub> = 0.034 0.10) and Mars (Fe<sup>3+</sup>/Fe<sub>T</sub> = 0.026 0.038) should be at the time of 914
- core formation using new data and equations of state for FeO and FeO<sub>1.5</sub>. The range of ferric iron 915
- abundances span observed values for the present day. However, no whole mantle model for the 916

Earth with these ferric iron contents match the last magma ocean atmosphere oxygen fugacity,

- based on measurements of D/H ratios (Pahlevan et al. 2019) when using lower mantle ferric iron
- partition coefficients calculated with the Deng EOS. The best fitting model for the Earth with the
- Deng EOS requires shallow magma ocean conditions, starting from a Bulk Silicate Earth composition with initial  $Fe^{3+}/Fe_T$  above ~0.06. A partial magma ocean could be explained by
- either an impact smaller than the moon-forming impact, or by the formation of a buoyant layer of
- minerals at mid-mantle depths that leads to separate evolution of an upper and lower (basal)
- magma ocean. However, a likelier explanation for the Earth is that the Armstrong EOS provides
- a more accurate description of lower mantle ferric iron partitioning, which is strongly
- 926 incompatible. This is consistent with new bridgmanite partition coefficient measurements by
- 927 Kuwahara and Nakada (2023). We find that all of our tested Earth magma ocean scenarios with
- the Armstrong EOS except the lowest initial  $Fe^{3+}/Fe_T$  are consistent with the D/H constraints.
- 929

For Mars, best fit models have initial ferric iron abundances ~2.5x lower than predicted values,

but partial magma oceans produce somewhat better matches to the D/H constraint. Because the

- minerals wadsleyite, ringwoodite, and majorite do not make up a significant portion of the
- Martian mantle, there is little difference between the Deng and Armstrong EOS models for Mars.
- If the magma ocean and atmosphere are separated when the magma ocean still has a greater
- depth than our predicted BPLE, then it would be possible to match the D/H constraint with larger
- initial ferric iron abundances more consistent with predictions.
- 937

We emphasize the need for more experimental data on ferric iron partition coefficients and melting temperatures of diverse compositions. Experimental synthesis of bridgmanite and

transition zone minerals suggests that ferric iron could behave more compatibly in the lower

- mantle than the upper mantle, but unfortunately, there is little data on mineral/melt partition
- coefficients for lower mantle minerals and the two available equations of state for ferric iron
- oxide predict opposite partitioning behavior for these minerals. Some key upper mantle minerals
- 944 (garnet) are likewise missing direct measurements of ferric iron partitioning with a co-existing

945 melt. In particular, measurements of the ferric iron partition coefficients of bridgmanite and

- garnet will produce the biggest improvements in redox evolution models.
- 947

There remain many uncertainties in the redox evolution of magma oceans on rocky planets. We 948 949 show that the most likely behavior for solar system bodies is an overall net increase in oxygen fugacity of outgassing, but this behavior does not necessarily hold true for all planets, depending 950 on the bulk composition and minerals that form at different depths. Because we find that ferric 951 iron must behave incompatibly in the lower mantle in order to match the Earth, we find that 952 larger planets such as super-Earths, which have larger lower mantle volumes, will likely produce 953 higher final oxygen fugacities than a smaller planet of the same initial composition. Therefore, 954 955 on average, larger planets should produce more oxidized initial atmospheres than smaller planets. 956

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- 962

#### 963 **Open Research**

- The magma ocean crystallization simulation code developed in this article is available at the
- Stanford Digital Repository (Schaefer, 2023a, https://doi.org/10.25740/yg376bh0915) along with
- all necessary input files to run the simulations shown in Figures 2-5. Matlab data files of all
- simulations are available in the archive.
- The magma ocean thermal evolution model used in Section 2.7 and 4.1 is a modified version of
- the model described by Schaefer et al. (2016). The code used in this paper is archived at the
- Stanford Digital Repository (Schaefer, 2023b, https://doi.org/10.25740/rk050tc3031).
- The literature data used to calculate ferric iron partition coefficients and bulk oxide partition
- coefficients are provided in the supplemental material as an .xlsx file.
- 973
- 974
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**Table 1.** Composition of residual magma ocean liquid following batch crystallization. Starting compositions are the Bulk Silicate Earth (BSE) (McDonough 2003) and Bulk Silicate Mars (BSM) (Taylor 2013). We use lower mantle ferric iron partition coefficients calculated from both the Deng (D) and Armstrong (A) EOSs for the batch calculations.

	Earth				Mars					
Oxide	500	) km	1000	) km	BSE	500	km	1000	) km	BSM
(wt%)	D	А	D	А		D	А	D	А	
FeO	12.7	12.5	11.2	11.0	7.82	24.2	24.2	21.4	21.3	18.1
MgO	31.8	31.2	34.0	33.6	38.3	17.8	17.7	25.2	25.2	31.2
SiO <sub>2</sub>	40.1	39.4	42.2	41.7	45.5	49.3	49.3	46.7	46.6	44.6
CaO	10.5	10.3	7.45	7.35	3.58	4.60	4.60	3.27	3.27	2.48
Al <sub>2</sub> O <sub>3</sub>	4.33	4.25	4.58	4.52	4.49	$3.58^{\dagger}$	3.57†	3.09†	3.09†	3.15
FeO <sub>1.5</sub>	$0.53^{*}$	$2.42^{*}$	$0.60^{*}$	$1.87^{*}$	$0.36^{*}$	$0.54^{**}$	$0.67^{**}$	$0.41^{**}$	$0.48^{**}$	$0.41^{**}$
Fe <sup>3+</sup> /Fe <sub>T</sub>	0.037	0.17	0.049	0.15	0.10	0.020	0.025	0.017	0.020	0.02

\*reported for initial  $Fe^{3+}/Fe_T = 0.10$ , \*\*Reported for initial  $Fe^{3+}/Fe_T = 0.02$ , <sup>†</sup>assumes  $Al_2O_3 = 1.2$ wt% in cpx, based on full fractional crystallization model.

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**Table 2**: Parameters for models that switch from batch to fractional crystallization at different depths, including the mass fraction of magma ocean, the pressure and depth at the base of the magma ocean when it transitions from batch to fractional, and the mineral fractions within the solid layer crystallized during the batch step.

	Ea	arth	Mars		
	500 km	1000 km	500 km	1000 km	
F <sub>mo</sub>	0.20	0.39	0.36	0.665	
P <sub>trans</sub> (GPa)	18	37	6.20	12.3	
z <sub>trans</sub> (km)	535	1005	508	1002	
T <sub>trans</sub> (K)	2160	2850	1930	2070	
Mineral fractions in batch	Bg 0.90	Bg 0.95	Maj 0.15	Maj 0.31	
crystallized layer	Mw 0.05	Mw 0.05	Ring 0.08	Ring 0.16	
	Maj 0.03		Wad 0.06	Wad 0.12	
	Ring 0.02		Cpx 0.14	Cpx 0.13	
			Opx 0.07	Opx 0.03	
			Gt 0.10	Gt 0.05	
			Ol 0.40	Ol 0.20	

**Table 3.** Ferric iron partition coefficients for the minerals included in our magma ocean model (see also Elkins-Tanton 2008). Partition coefficients for wadsleyite, majorite, ringwoodite, and bridgmanite are calculated using either the Deng or Armstrong (*italic*) EOS for ferric iron in silicate melts (see **Text S2** and **S3** for details). Abbreviations: sp – spinel, ol – olivine, cpx – clinopyroxene, opx – orthopyroxene, gt – garnet, wad – wadsleyite, maj – majorite, ring – ringwoodite, bg – bridgmanite, mw – magnesiowüstite

phase	$D_{Fe^{3+}}^{mineral/melt}$	Ref.
Sp	$exp[(0.87 \pm 0.07) \frac{10,000}{T} + (-4.6 \pm 0.4) + (0.24 \pm 0.02) ln \left(C_{Fe_2O_3}^{sp}(wt\%)\right)]$	1
Ol	0	
Срх	$\exp\left[(1.997 \pm 0.640) \ln C_{Al_2O_2}^{cpx} + (-3.987 \pm 0.965)\right]$	2
Opx	$(0.70^{+1.56}_{-0.64})D^{cpx/melt}_{Fe^{3+}}$	3-7
Gt	$(0.68^{+1.95}_{-0.51})D^{cpx/melt}_{Fe^{3+}}$	3-6
Wad	1.08±0.49	8
	$0.98{\pm}0.44$	
Maj	1.95 ±0.69	8
	0.97 ±0.34	
Ring	$2.13 \pm 1.17$	8
	0.97±0.53	
Bg	$2.07 \pm 0.50$	9
-	0.17±0.03	
Mw	0	

1 – Davis & Cottrell (2021), 2 – refit from Rudra & Hirschmann (2022), 3 - Woodland et al. 2006, 4-Canil & O'Neill 1996, 5 - Malaspina et al. 2012, 6- Nimis et al. 2015, 7-Luth & Canil 1993, 8 – calculated from data in O'Neill et al. 1993, 9 – calculated from data in Boujibar et al. 2016

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<sup>A</sup> FeO				
Reproduced from Hirschmann (2022), Table 2.				
Parameter	Value	Units		
а	0.1917			
b	-1.961			
С	4158.1	Κ		
$\Delta C_p$	33.25	J/K		
$T_0$	1673.15	Κ		
У1	-520.46	Κ		
<i>y</i> 2	-185.37	Κ		
<i>уз</i>	494.39	Κ		
<i>Y</i> 4	1938.34	Κ		
<i>y</i> 5	2888.48	Κ		
<u>у</u> 6	3473.68	Κ		
<i>Y</i> 7	-4473.6	Κ		
<i>y</i> 8	-1245.09	Κ		
<u>y</u> 9	-1156.86	K		

Table 4. Fit param	neters for calculating fO2 from
$\frac{X_{Fe0_{1.5}}}{X_{Fe0}}$ in silicate me	elts from equation (12).



1401 Figure 1. Diagram of model setup showing the differences between the whole mantle models 1402 and shallow magma ocean models. All models begin with a BSE or BSM composition (see 1403 Table 1). In the whole mantle models (*left*), we assume the magma ocean initially occupies the 1404 whole mantle, with either pure fractional crystallization or a mix of batch and fractional 1405 crystallization, with the switch occurring once the base of the magma ocean is above R<sub>trans</sub>. We 1406 model batch crystallization as a single stage to find the compositions from which to initialize the 1407 fractional crystallization model (see **Table 3**). **Table 2** gives the conditions (P, T, z) at R<sub>trans</sub>. We 1408 adopt the whole mantle magma ocean configurations as our nominal models and show the 1409 1410 evolution during the fractional crystallization stages in Fig. 2 and Fig. 4, as well as the solid lines in Fig. 3 and Fig. 5. In the shallow magma ocean models (*right*), we assume that only a portion 1411 of the mantle melts ( $R_{mo} = R_p - 1000$  km, or  $R_p - 500$  km) and solidifies entirely through 1412 1413 fractional crystallization. The physical conditions at R<sub>mo</sub> for the shallow models are the same as 1414 for Rtrans given in Table 2. The shallow magma ocean models are shown as dashed lines in Fig. 3 1415 and Fig. 5.



Figure 2. Evolution of oxygen fugacity for the fractional crystallization stages of whole mantle 1420 magma ocean models of Earth (see Fig. 1 (*left*)). The oxygen fugacity is calculated at the surface 1421 of the Earth's magma ocean and is plotted as a function of the pressure at the base of the magma 1422 1423 ocean at a given time. Therefore, the x-axis represents time evolution of the magma ocean, with time from the beginning of the magma ocean increasing from right to left. These calculations 1424 start with a nominal initial  $Fe^{3+}/Fe_T = 0.10$  and use ferric iron partition coefficients calculated 1425 from either the Deng EOS (top row) or Armstrong EOS (bottom row). Oxygen fugacity is shown 1426 1427 relative to the iron-wüstite buffer (Hirschmann 2021). The starting conditions for the three models are given in Table 1, assuming batch crystallization of the lower mantle for the 500 and 1428 1429 1000 km models. Grey lines are individual simulations with randomly drawn ferric iron partition coefficients, the black line is the average of all 1000 simulations. Pink horizontal shaded region 1430 1431 highlights fO<sub>2</sub> permitted by D/H constraint (Pahlevan et al. 2019), vertical grey shaded region 1432 highlights basal pressure of last equilibration (BPLEBPLE). 1433





1439 Figure 3. Mean oxygen fugacities at the surface of the Earth's magma ocean for different initial 1440  $Fe^{3+}/Fe_T$  (blue: 0.04, red: 0.10, yellow: 0.14, purple: 0.20). The oxygen fugacity is calculated at the surface of the Earth's magma ocean and is plotted as a function of the pressure at the base of 1441 the magma ocean at a given time. Therefore, the x-axis represents time evolution of the magma 1442 1443 ocean, with time from the beginning of the magma ocean increasing from right to left. Ferric iron partition coefficients were determined with the Deng EOS (top) or the Armstrong EOS (bottom). 1444 All lines represent the mean of 1000 simulations. Solid lines show the fractional crystallization 1445 evolution of whole mantle magma ocean models with interstitial liquid = 20% (see Fig. 1 (*left*)). 1446 Dashed lines are for the shallow magma ocean models (see Fig. 1 (right)) starting at either 500 1447 km or 1000 km (interstitial = 20%). Table S5 gives the number of successful simulations for 1448 1449 each model configuration. Pink horizontal shaded region highlights fO2 permitted by D/H constraint (Pahlevan et al. 2019), vertical grey shaded region highlights basal pressure of last 1450 equilibration (BPLEBPLE). 1451 1452





Figure 4. Evolution of oxygen fugacity for the fractional crystallization stages of whole mantle 1458 magma ocean models of Mars (see **Fig. 1** (*left*)). The oxygen fugacity is calculated at the surface 1459 of Mars' magma ocean and is plotted as a function of the pressure at the base of the magma 1460 ocean at a given time. Therefore, the x-axis represents time evolution of the magma ocean, with 1461 time from the beginning of the magma ocean increasing from right to left. These calculations 1462 start with a nominal initial  $Fe^{3+}/Fe_T = 0.01$  and use ferric iron partition coefficients calculated 1463 from either the Deng EOS (top row) or Armstrong EOS (bottom row). Oxygen fugacity is shown 1464 relative to the iron-wüstite buffer (Hirschmann 2021). The starting conditions for the three 1465 models are given in Table 1, assuming batch crystallization of the lower mantle for the 500 and 1466 1467 1000 km models. Grey lines are individual simulations with randomly drawn ferric iron partition coefficients, the black line is the average of all 1000 simulations. Pink horizontal shaded region 1468 highlights  $fO_2$  permitted by D/H constraint (Pahlevan et al. 2022), vertical grey shaded region 1469 1470 highlights pressure of last equilibration (BPLEBPLE). 1471



Figure 5. Mean oxygen fugacities at the surface of Mars' magma ocean for different initial 1475 Fe<sup>3+</sup>/Fe<sub>T</sub> (blue: 0.01, red: 0.02). The oxygen fugacity is calculated at the surface of Mars' magma 1476 ocean and is plotted as a function of the pressure at the base of the magma ocean at a given time. 1477 Therefore, the x-axis represents time evolution of the magma ocean, with time from the 1478 1479 beginning of the magma ocean increasing from right to left. Ferric iron partition coefficients were determined with the Deng EOS (top) or the Armstrong EOS (bottom). All lines represent 1480 the mean of 1000 simulations. Solid lines show the fractional crystallization evolution of whole 1481 mantle magma ocean models with interstitial liquid = 20% (see Fig. 1 (*left*)). Dashed lines are 1482 for the shallow magma ocean models (see Fig. 1 (right)) starting at either 500 km or 1000 km 1483 (interstitial = 20%). Table S6 gives the number of successful simulations for each model 1484 1485 configuration. Pink horizontal shaded region highlights fO<sub>2</sub> permitted by D/H constraint (Pahlevan et al. 2022), vertical grey shaded region highlights basal pressure of last equilibration 1486 (BPLEBPLE). 1487





- 1490 oxygen fugacity at the surface of the magma ocean. *Left*: whole Earth magma ocean with
- 1491  $(Fe^{3+}/Fe_T)_{init} = 0.10$ , *right*: whole Mars magma ocean with  $(Fe^{3+}/Fe_T)_{init} = 0.01$ . Bold lines show
- nominal mean values from **Figures 1 & 3**. We reduce the partition coefficients of
- 1493 ringwoodite(18 22 GPa), wadsleyite (15-18 GPa), majorite (15 22 GPa), bridgmanite (>22
- 1494 GPa) individually, and all together (l.m. = lower mantle) to  $0.5\pm0.5$ . We also increase the garnet
- 1495 (2.5 15 GPa) partition coefficient to  $1.75 \pm 0.75$ .