

Inefficient water degassing inhibits ocean formation on rocky planets: An insight from self-consistent mantle degassing models

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ABSTRACT

A sufficient amount of water is required at the surface to develop water oceans. A significant fraction of water, however, remains in the mantle during magma ocean solidification, and thus the existence of water oceans is not guaranteed even for exoplanets located in the habitable zone. To discuss the likelihood of ocean formation, we built two models to predict the rate of mantle degassing during the magma ocean stage and the subsequent solid-state convection stage. We find that planets with low $\text{H}_2\text{O}/\text{CO}_2$ ratios would not have a sufficient amount of surface water to develop water oceans immediately after magma ocean solidification, and the majority of the water inventory would be retained in the mantle during their subsequent evolution regardless of planetary size. This is because oceanless planets are likely to operate under stagnant lid convection, and for such planets, dehydration stiffening of the depleted lithospheric mantle would limit the rate of mantle degassing. In contrast, a significant fraction of CO_2 would already be degassed during magma ocean solidification. With a strong greenhouse effect, all surface water would exist as vapor, and water oceans may be absent throughout planetary evolution. Volatile concentrations in the bulk silicate Earth are close to the threshold amount for ocean formation, so if Venus shared similar concentrations, small differences in solar radiation may explain the divergent evolutionary paths of Earth and Venus.

Keywords: habitability, magma ocean, ocean formation

1. INTRODUCTION

The continuing discovery of extrasolar terrestrial planets has increased the prospect of identifying habitable exoplanets. Earth-size exoplanets are starting to be found at a heliocentric distance where liquid water may be stable (Gillon et al. 2017; Gilbert et al. 2020), and one of the super-Earth size exoplanets identified in the habitable zone, K2-18b, has water vapor detected in its atmosphere (Benneke et al. 2019b; Tsiaras et al. 2019). With the upcoming James Webb Space Telescope, more detections of water vapor are expected on potentially habitable exoplanets (Gardner et al. 2006), and the possible presence of surface water makes such exoplanets a leading candidate to search for biosignatures (Schulze-Makuch et al. 2011; Madhusudhan et al. 2020). Given the importance of surface liquid water, one of the key questions is the likelihood of water ocean formation on an exoplanet with a water vapor atmosphere.

The presence of surface water has often been discussed in terms of the habitable zone (Kasting et al. 1993; Kopparapu et al. 2013; Kodama et al. 2019), which is the region where liquid water is stable at the temperature and pressure conditions of a planetary surface. The premise, however, is that the amount of surface water is large enough to develop oceans. Although a threshold value to form water oceans is $\sim 10^{16}$ kg for the present-day Earth and is thus well smaller than the current ocean mass (1.4×10^{21} kg), the threshold changes with the amount of greenhouse gases and net stellar radiation (Abe 1993a; Salvador et al. 2017). For example, during the early Hadean on Earth, an atmosphere of 100 bar CO_2 would have increased this threshold to ~ 0.1 ocean mass, so it is not guaranteed that the threshold amount of water would always exist on the surface of terrestrial planets. A significant fraction of the water inventory can be retained in the planetary interior, precluding the formation of water oceans.

The initial amount of surface water is determined by volatile partitioning during magma ocean solidification, whereas its subsequent evolution is governed by interaction with

mantle convection (Ito et al. 1983; Korenaga et al. 2017). Previous studies have often assumed that the mantle entirely degases during magma ocean solidification (Elkins-Tanton 2008; Lebrun et al. 2013; Salvador et al. 2017), but considering the rheological transition of a partially molten medium and its slow compaction rate, a significant amount of volatiles could be trapped in the mantle (Hier-Majumder and Hirschmann 2017). Furthermore, if water oceans are initially absent, the lack of surface water may have persisted for a geological timescale. The presence of surface water is critical in reducing the lithospheric strength and thus in triggering plate tectonics (Korenaga 2007, 2010a, 2020). Without oceans, the mantle would operate under stagnant lid convection, and mantle upwelling would be suppressed by a rigid lid covering the entire surface (Solomatov 1995). Furthermore, dehydration stiffening of the depleted lithospheric mantle, caused by loss of water upon melting, could further hinder mantle processing (Korenaga 2009b). Under such a mode of convection, mantle degassing becomes inefficient, and the amount of surface water could be limited over a substantial period of time.

In this paper, we investigate mantle degassing during and after the solidification of a magma ocean and discuss the likelihood of ocean formation on terrestrial planets. Initial conditions necessary for planets to develop water oceans are explored for various planetary sizes and volatile compositions. Previous studies discussing the impact of mantle degassing on habitability have mostly focused on the solid-state convection stage (Noack et al. 2017; Tosi et al. 2017; Vilella and Kaminski 2017; Dorn et al. 2018), but the atmosphere could be dominated by volatiles degassed during the preceding magma ocean. Moreover, volatile concentrations in the mantle affect viscosity and thus the thermal evolution of planets, so it is crucial to solve for the entire history of mantle degassing self-consistently. It is noted that we assume that planets are in an oxidized condition, and the atmospheric components considered here are limited to H_2O and CO_2 . The redox of the early Earth remains controversial, but recent studies suggest that planets larger than Mars-size would have an oxidized atmosphere (Hirschmann 2012; Deng et al. 2020).

2. DEGASSING DURING MAGMA OCEAN:

After the final giant impact and prior to the solidification of the mantle surface

We first consider mantle degassing during the early stage of magma ocean solidification. Here the focus is on the aftermath of the magma ocean produced by the final of multiple giant impacts, which are common in N -body simulations of planetary formation (e.g., Rubie et al. 2015; Quintana et al. 2016). The final giant impact was likely energetic enough to melt most of the mantle (Nakajima and Stevenson 2015), so the atmosphere existing prior to the impact would either have re-equilibrated with the magma ocean. The same is likely to be true for larger exoplanets because, with an impactor of 10% target mass, a typical impact velocity is high enough to produce substantial melting of the mantle (Stixrude 2014). Also, the amount of radiogenic Xe in the atmosphere suggests that a catastrophic atmospheric loss likely occurred around the time of the Moon-forming giant impact (Porcelli and Pepin 2000). Therefore, we assume that volatile partitioning would be reset by the final giant impact, and we estimate how the volatile budget would be distributed between the atmosphere and the mantle during the solidification of the final magma ocean.

The solidification of a magma ocean can be divided into two stages by its surface melt fraction (Figure 1). With a high melt fraction at the surface, the magma ocean produces a high convective heat flux, resulting in a surface temperature high enough to maintain a molten surface (Lebrun et al. 2013; Salvador et al. 2017). As the mantle cools down and melt fraction becomes lower than the critical value (~ 0.4), however, the partially molten mantle would undergo a rheological transition and start to behave as solid (Abe 1993b). Convective heat flux plummets, and the surface temperature would drop to < 500 K, which is well below the mantle solidus temperature (Lebrun et al. 2013; Salvador et al. 2017). Therefore, the surface layer would be completely solidified in a short timescale once the surface melt fraction reaches the critical value, even though the interior may still be

partially molten. Chemical equilibrium between the mantle and the hydrosphere is not maintained thereafter, and volatile exchange between the surface and the interior would instead be characterized by solid-state convection (Abe 1997). Although its contribution is expected to be small, some additional degassing processes may take place during the transition period from a magma ocean to solid-state convection and are discussed in Section 4.1.

In this section, we focus on volatile partitioning between the atmosphere and magma ocean while a partially molten surface is sustained, and the main goal here is to show the impact of the rheological transition on mantle degassing. We emphasize that the magma ocean considered here is produced by the final giant impact on the planet. Other degassing models have also examined the same period (e.g., Elkins-Tanton 2008; Lebrun et al. 2013; Salvador et al. 2017; Bower et al. 2019), and ours and these studies investigate a different evolutionary stage from accretionary magma ocean models (e.g., Matsui and Abe 1986; Zahnle et al. 1988). The latter type of model tracks the amount of added surface water as an impactor hits the planet and considers different degassing processes from the former. Also, accretionary magma ocean models do not consider large-scale melting at the final stage of accretion, so the mantle and atmosphere are never in equilibrium in terms of volatile partitioning. We assume that a giant impact most likely induces global-scale melting (Nakajima and Stevenson 2015) so that a substantial amount of volatiles would be partitioned into the mantle. Although both types of models aim to quantify surface water mass at the end of accretion, we caution that their validity depends on the assumed scale of mantle melting.

2.1 Volatile partitioning

When a magma ocean behaves rheologically as liquid, the atmosphere and the magma ocean are equilibrated in a short timescale, and volatile partitioning between the magma and the atmosphere would determine the atmospheric pressure (Elkins-Tanton 2008; Lebrun et al. 2013). Magma at the surface dissolves volatiles up to their solubility limits,

and volatiles in excess of saturation would reside in the atmosphere. As a magma ocean cools down, melt fraction decreases from the lowermost part of the mantle (e.g., Abe 1993b; Solomatov 2007), and the ‘melt-dominated’ layer shrinks and the ‘solid-dominated’ layer grows (Figure 1). The two layers are separated by the rheological transition and are defined by their melt fractions as $\phi \geq 0.4$ and $\phi < 0.4$, respectively. These layers were referred to as ‘soft’ and ‘hard’ magma oceans in Abe (1993a). We solve for volatile partitioning and mass balance at each time step as the magma ocean undergoes the rheological transition from the bottom upward.

Volatile concentration in the surface magma, x_i , corresponds to the solubility at its atmospheric partial pressure, p_i , for each volatile species i . The solubility measurements of CO₂ and H₂O are parameterized as follows (Blank and Brooker 1994; Lichtenberg et al. 2021):

$$p_{\text{H}_2\text{O}} = \left(\frac{x_{\text{H}_2\text{O}}}{6.8 \times 10^{-6}} \right)^{1.42}, \quad (1)$$

$$p_{\text{CO}_2} = \left(\frac{x_{\text{CO}_2}}{4.4 \times 10^{-10}} \right), \quad (2)$$

where pressure is shown in the unit of Pa and concentration is in wt%. Some models indicate higher solubility for both H₂O and CO₂ (Papale 1997; Gardner et al. 1999), but we adopt the least soluble model to prepare the most optimistic case for ocean formation. Regardless of the model, water is predicted to be more soluble in silicate melts than CO₂. The variables P_i and x_i can be solved using the following mass balance (Bower et al. 2019):

$$\phi x_i M_l + \frac{\mu_i}{\bar{\mu}} \frac{4\pi r_p^2}{g} p_i = m_i, \quad (3)$$

where ϕ is the melt fraction at the surface, M_l is the mass of the melt-dominated layer, μ_i is the molar mass of volatile i , $\bar{\mu}$ is the mean molar mass of the atmosphere, r_p is the planetary radius, and m_i is the total mass of volatile i in the atmosphere and the melt-dominated layer (Figure 1a). The surface melt fraction ϕ is determined based on the surface temperature, T_s , and a phase diagram of peridotite (Katz et al. 2003), and the surface temperature is calculated using on a 1-D atmospheric model of Nakajima et al. (1992) (see Appendix A.1 and A.2 for details).

The volatile mass m_i represents those that were not trapped in the solid-dominated layer existing in the deeper region (Figure 1). Taking into account the rheological transition of a partially molten medium, a significant amount of melt could be trapped in the pore space of the solid matrix, together with volatiles dissolved in the melt (Figure 1b). The percolation of pore melt is slower than the speed of magma ocean solidification (Hier-Majumder and Hirschmann 2017), and a newly formed rheologically-solid layer, which includes volatile-rich pore melt, would be quickly delivered to the deeper region by the Rayleigh-Taylor instability (Maurice et al. 2017; Miyazaki and Korenaga 2019a). The solid-dominated layer is much more viscous than the melt-dominated one, and its convection would be insignificant within the cooling timescale of the melt-dominated layer. However, because the solidus and liquidus curves of peridotite are superadiabatic (Figure S1), the density structure of the solid-dominated layer is gravitationally unstable after the rheological transition. For parameters used in Miyazaki and Korenaga (2019a), the entire lower mantle solidifies in ~ 4000 years, whereas the timescale for the Rayleigh-Taylor instability is less than a year just after the lower mantle undergoes the rheological transition. Volatiles trapped in the pore space of the solid matrix would thus be effectively sequestered in the deep mantle and be segregated from the remaining melt layer (Figure 1a, 1b).

During the instability-triggered downwelling, the melt in the pore space would solidify by adiabatic compression, and the melt would be supersaturated and exsolve volatiles. Yet, the saturation limit would be reached only when melt fraction becomes lower than ~ 0.002 . Here we consider an initial H_2O content below 0.1 wt%, and in such a case, concentrating volatiles by 500 times would still not result in exceeding saturation limits (Kawamoto and Holloway 1997). Therefore, the melt-solid mixture would be mostly solidified when volatiles in melt pockets finally start to be degassed, and exsolved volatiles would either diffuse into the surrounding nominally anhydrous minerals or be trapped in the solid matrix as fluid inclusions. Volatiles incorporated in the pore space of the solid matrix would thus be kept in the solid-dominated layer during the subsequent evolution, and we assume that the corresponding amount is subtracted from the volatile budget of the melt-

dominated layer. As the magma ocean cools down and the depth of the melt-dominated layer, d , decreases, the volatile budget m_i changes as:

$$m_i(d - \Delta d) = m_i(d) - \phi x_i \Delta M_l, \quad (4)$$

where ΔM_l is the mass of the magma ocean that underwent the rheological transition while d decreased by Δd . Considering that vigorous convection takes place within the melt-dominated layer, the bulk volatile concentration in a crystal-melt mixture is assumed to be constant ($= \phi x_i$) throughout the melt-dominated layer. Equations (1)-(4) are solved until the melt-dominated layer disappears ($M_l, d = 0$).

2.2 Degassing of the upper mantle by percolation

Once the entire mantle undergoes the rheological transition and the melt-dominated layer disappears, the evolution of a magma ocean is characterized by solid-state convection. Although percolation is slower than the Rayleigh-Taylor instability, it is faster than solid-state convection, so melt and dissolved volatiles in the pore space would escape towards the surface (Figure 1c). Volatiles included in the partial melt would be degassed to the surface, and their amount is proportional to the mass of partially molten layer when the melt-dominated layer disappears. We calculate its thickness from the adiabatic temperature profile with a critical melt fraction of 0.4 at the surface (Abe 1993b; Solomatov 2007). For simplicity, the mantle is assumed to be compositionally homogeneous and have a pyrolitic composition, and in such a case, the partial melt layer would extend to a depth of ~ 5 GPa (Appendix A.3). We note that, if the mantle experiences differentiation during magma ocean solidification, the thickness of the partial melt layer could decrease by half (Appendix A.4; Miyazaki and Korenaga 2022), and degassing by percolation could be smaller than presented in Figures 2-4. The mantle, however, would eventually be homogenized, and the long-term evolution of surface water would converge to results presented in Section 3.

2.3 Results

2.3.1. Degassing of Earth-size planets

Figure 2 shows a typical evolution of the atmospheric pressure for an Earth-size planet at 1 AU. A fraction of volatiles would be degassed from the beginning to maintain equilibrium with volatiles in the mantle, but further degassing is limited during the early stage of solidification because volatiles are trapped in the pore space of the solid-dominated layer and thus volatile concentrations remain mostly unchanged (Figure 2a). Here we assume initial volatile concentrations of 0.04 and 0.1 wt% for both H₂O and CO₂, and regardless of the initial concentration, >99% of the total H₂O budget is estimated to be retained in the mantle during the early stage of magma ocean solidification. On the other hand, because CO₂ is less soluble to silicate magma, ~70% of the total CO₂ would be released to the atmosphere (Figure 2a; Hier-Majumder and Hirschmann 2017), and such an atmosphere dominated by CO₂ would further suppress the degassing of lighter gases, including H₂O (Bower et al. 2019). Degassing resumes when the surface magma starts to solidify as volatiles concentrate in the remaining melt phase (Figure 2b) and a higher atmospheric pressure is required to maintain equilibrium between the atmosphere and the magma ocean. Degassing by percolation would also increase the volatile mass in the atmosphere, yet ~95% of H₂O would still be retained in the mantle.

2.3.2. Degassing of super-Earths

We repeat calculations in Section 2.3.1 for a range of initial volatile content and planetary mass. As expected, the pressure of degassed atmosphere increases with higher initial volatile concentrations, but a larger planetary mass does not necessarily result in a thicker atmosphere; the partial pressure for degassed H₂O atmosphere, $p_{\text{H}_2\text{O}}$, shows little dependence on planetary mass (Figure 3a). This is because the majority of H₂O is stored in the mantle, and the water concentration in the mantle changes little from the initial value. Therefore, the corresponding atmospheric pressure in equilibrium with the mantle reservoir is mostly the same regardless of planetary mass. In contrast, most of the CO₂ inventory is degassed to the atmosphere, and thus p_{CO_2} becomes higher for larger planets

(Figure 3b). The partial pressure is related to the mass of volatile i in the atmosphere, M_i , through:

$$p_i = \frac{M_i g}{4\pi r_p^2} \frac{\bar{\mu}}{\mu_i} \propto M_i, \quad (5)$$

where g is the gravitational acceleration and r_p is the planetary radius. The relation between g and r_p is calculated from the simplified interior model of Seager et al. (2007). Therefore, with the same volatile concentrations, M_{CO_2} increases with planetary mass (Figure 3b), whereas $M_{\text{H}_2\text{O}}$ remains the same for planets of any size (Figure 3a).

When the amount of surface H_2O is small, the entire surface water budget can exist as vapor in the atmosphere without forming water oceans. The threshold amount of H_2O for ocean formation is estimated using a 1-D radiative-convective model of Nakajima et al. (1992). The model predicts that the threshold amount increases with a thicker CO_2 atmosphere (Abe 1993a; Salvador et al. 2017) because a thicker atmosphere can contain more H_2O vapor, and its amount further increases under a higher surface temperature induced by a stronger greenhouse effect. Under the same initial volatile concentrations, therefore, larger planets are less likely to develop water oceans immediately after the solidification of the mantle surface (Figure 3a, 3c). For an initial CO_2 concentration of 0.01 wt%, the amount of degassed CO_2 (M_{CO_2}) increases from 5.1×10^{19} kg for a Mars-size planet to 1.1×10^{21} kg for a $10M_E$ super-Earth (Figure 3b), which would raise the amount of surface H_2O required to develop water oceans from $\sim 1.6 \times 10^{18}$ kg to $\sim 3.1 \times 10^{19}$ kg (Figures 3a). Nevertheless, the amount of surface water $M_{\text{H}_2\text{O}}$ does not change with planetary size, where $M_{\text{H}_2\text{O}}$ is $\sim 2 \times 10^{19}$ kg for any planets with an initial H_2O concentration of 0.01 wt%. Smaller planets are thus more likely to develop water oceans when the surface of the mantle solidifies. How the initial volatile concentration of terrestrial planets is characterized is still debated (Hirschmann et al. 2021; Li et al. 2021), but if terrestrial exoplanets have similar volatile concentrations to Earth, super-Earths may lack water oceans when the surface of the mantle solidifies (Figure 3c).

It is noted that the minimum surface H₂O mass to form water oceans in Figure 3a is associated with an uncertainty of a factor of ~2 because a 1-D gray atmospheric model is adopted here. Considering that the subsiding branch of the atmospheric circulation is undersaturated with water vapor, the average relative humidity should be smaller than 1 (e.g., Ishiwatari et al. 2002; Abe et al. 2011; Pierrehumbert and Ding 2016). It is thus assumed that the relative humidity is 0.7 throughout the troposphere in our model, but this value remains unconstrained for an atmosphere with an extreme CO₂ concentration and may range between 0.5 and 1. Also, CO₂ existing near the surface could be in a supercritical state when p_{CO_2} is larger than 74 bar. Although the temperature and pressure conditions considered here are distant from its critical point, physical properties including opacity may differ by some degree from the gas state. Implementing these factors, together with the wavelength dependence of the opacity, may modify our estimate of the minimum H₂O mass required for ocean formation by some factor. The overall trend of our results is unlikely to change, but a more detailed model would be needed to predict the actual threshold.

2.4 Discussion: early Earth and early Venus

The amount of volatiles in the bulk silicate Earth (BSE) could have been just above the threshold for stabilizing water oceans at the point of magma ocean solidification. Although the volatile content in the terrestrial mantle remains controversial, recent estimates suggest that the mantle and the hydrosphere in total contain 1.2-3 ocean mass of H₂O (Hirschmann and Dasgupta 2009; Korenaga et al. 2017) and $4.8\text{--}26 \times 10^{20}$ kg of CO₂, which translates to initial concentrations of 0.04-0.1 wt% and 0.01-0.07 wt%, respectively, in the mantle. The estimated volatile concentrations lie just inside the boundary between the regimes of ocean formation and dry surface (Figure 3c), and if BSE had a lower-bound H₂O and an upper-bound CO₂ concentration, water oceans may have been absent during the early stage of evolution. As we discuss in Section 3, the rate of mantle degassing decreases substantially after the mantle surface solidifies, and thus the surface would continue to lack water oceans if oceans are absent at the time of surface solidification. The presence of surface water in the Hadean has been suggested from zircon records

(e.g., Mojzsis et al. 2001; Wilde et al. 2001), so water oceans were likely present on the surface of Earth from the beginning of its evolution. A combination of a low H_2O and a high CO_2 concentration in BSE (>0.05 wt%) may be ruled out if water oceans indeed existed on the surface of the Hadean Earth.

The early Venus, on the other hand, may have lacked water oceans because a larger amount of surface H_2O is required for planets closer to the central star (Figure 4a). Assuming the same albedo, the net solar radiation for the early Venus was 330 W m^{-2} , which is two times higher than the value for the early Earth. To balance stronger solar radiation, the overall thermal structure, including the planetary surface, becomes hotter, thus allowing more H_2O to exist as water vapor in the atmosphere (Nakajima et al. 1992). The minimum amount of H_2O to form oceans doubles for a M_{CO_2} of 10^{21} kg and increases by a factor of 5 for $M_{\text{CO}_2} \sim 10^{20}$ kg. As a result, for a planet receiving 330 W m^{-2} of stellar radiation, water oceans would be absent for a wider range of initial volatile concentrations. If Venus had similar volatile concentrations to Earth, our result suggests that Venus would develop water oceans only for a combination of a high H_2O and a low CO_2 concentration (Figure 4b). Because the threshold for ocean formation lies within the estimated volatile concentrations in BSE, the location of planets and thus the amount of net solar radiation may have decided the fate of two similar planets (see Section 3.3 for further discussions). Mars, on the other hand, likely developed water oceans immediately after magma ocean solidification because of its small size (Figure 3c) and its distance from the Sun.

Volatile concentrations required to develop water oceans is much higher in our model than estimated in Salvador et al. (2017) because we take into account volatiles that are trapped in the pore space of the melt-solid mixture. Models of magma ocean solidification have often neglected this effect and assumed that volatiles, behaving as incompatible elements, concentrate in the residual melt layer as the magma ocean solidifies from the bottom to the top (Elkins-Tanton 2008; Hamano et al. 2013; Lebrun et al. 2013; Salvador et al. 2017). In such a scenario, the concentration of volatiles eventually exceeds saturation levels and results in efficient degassing. Volatiles, however, would not be efficiently degassed because the compaction of melt-solid mixture is slow (Hier-Majumder

and Hirschmann 2017) and because the Rayleigh-Taylor instability efficiently delivers volatile-rich pore melt to the deeper mantle. Therefore, the remaining melt layer near the surface experiences little change in volatile concentrations, and the mantle is likely to remain hydrated (Section 2.1, Figure 2a).

3. DEGASSING DURING SOLID-STATE CONVECTION:

After the solidification of the mantle surface

Once the surface of the mantle solidifies, mantle degassing during the subsequent evolution would be characterized by solid-state convection. Although partial melt may still exist in the interior, the planetary surface would be covered by a solidified lid, and the mode of volatile exchange would be similar to what is seen on the present-day Venus and Mars (Noack et al. 2017; Dorn et al. 2018). The solid lid prohibits the mantle and the hydrosphere to reach equilibrium, and instead the evolution of the atmosphere becomes controlled by a balance between ingassing to and degassing from the mantle. The rate of ingassing and degassing differs appreciably depending on the mode of solid-state convection, so whether the mantle operates under plate tectonics or stagnant lid convection would control the atmospheric evolution of terrestrial planets. Here, plate tectonics refers to a mode that allows the continuous recycling of the uppermost layer, whereas in stagnant lid convection, a rigid immobile shell covers the surface, and convective motion is limited to the interior region beneath the shell (Solomatov 1995).

The key to triggering plate tectonics is the weakening of oceanic lithosphere (Moresi and Solomatov 1998), and several mechanisms have been proposed, including grain-size reduction (e.g., Kameyama et al. 1997; Bercovici and Ricard 2012) and thermal cracking (Korenaga 2007). Comparison between different mechanisms are discussed in several reviews (Bercovici et al. 2015; Korenaga 2020), but the weakening of the strongest part of the lithosphere requires the reduction of friction coefficient by some means, for which thermal cracking is so far the only mechanism that is consistent with our understanding of rock mechanics. Positive feedback between thermal cracking and serpentinization

could potentially hydrate the lithosphere deeply (Korenaga 2017), with trapped water reducing lithostatic-hydrostatic pressure difference and thus effective friction coefficient. The presence of surface water is likely the key to lowering the yield strength of the lithosphere, and although the mechanism is different, the importance of surface water in triggering plate tectonics has been suggested in other studies as well (Regenauer-Lieb et al. 2001; Gerya et al. 2008). Indeed, other oceanless planets, including Venus, Mars, and Moon, are generally considered to be operating under the mode of stagnant lid convection. Sporadic subduction may be triggered by large meteorite impacts during the early stage of planetary evolution (Hansen 2007; O'Neill et al. 2017), but its effect is likely to be ephemeral, and the mode of convection would revert to stagnant lid without water oceans.

The absence of surface water thus leads to stagnant lid convection, which limits both ingassing and outgassing compared to plate tectonics (Kite et al. 2009). Under solid-state convection, the degassing of the mantle is triggered when the mantle material undergoes partial melting during its upwelling (Fraeman and Korenaga 2010; O'Rourke and Korenaga 2012), but a rigid immobile shell would limit the extent of such upwelling motion. With the same convective velocity, the processing of the mantle would be significantly slower for stagnant lid convection than plate tectonics, under which the mantle material is brought up all the way to the surface. Oceanless planets would therefore have a smaller supply of H₂O from the mantle than those with oceans, implying that an oceanless state may persist for the geological time.

The lack of surface recycling also prevents the transport of volatiles from the hydrosphere to the mantle, whereas the sequestration of volatiles is possible under plate tectonics (Sleep and Zahnle 2001; Miyazaki and Korenaga 2022). Volatiles, in particular CO₂, can be stored in oceanic crust as carbonate minerals (e.g., Alt and Teagle 1999), which could then be delivered to the interior by subduction. The majority of atmospheric CO₂ existed after magma ocean solidification on Earth should have been transported to the mantle by the end of Hadean (Catling and Zahnle 2020), and a rapid removal of atmospheric CO₂ is also expected for other terrestrial planets with plate tectonics. On the other hand, the

removal of atmospheric CO_2 would be inhibited for stagnant lid convection, and rather CO_2 would continue to accumulate in the atmosphere by mantle degassing. The threshold amount of H_2O to develop water oceans thus continues to increase with time for such planets, which would be another factor that further delays the formation of water oceans.

Here, we estimate how long it takes for initially oceanless dry planets to develop water oceans after solid-state convection starts, by modeling the thermal evolution of terrestrial planets. Previous studies discussing habitability in the context of stagnant lid convection have focused on the degassing of CO_2 because the availability of greenhouse gas on the surface was considered to regulate the outer edge of the habitable zone (Noack et al. 2017; Tosi et al. 2017; Vilella and Kaminski 2017; Dorn et al. 2018). However, they have not taken into account the massive CO_2 atmosphere released during magma ocean (Figure 2a), and with its effect included, the amount of greenhouse effect should not be a limiting factor for habitability for planets with stagnant lid convection. Because the focus of this paper is the formation of water oceans on terrestrial planets, we do not model the evolution after water oceans form, but its impact on habitability is discussed in Section 4.2.

3.1. Methods: Mantle degassing under stagnant lid convection

The degassing rate of terrestrial planets that initially lack water oceans is calculated based on a heat flow scaling of stagnant lid convection (Korenaga 2009b). This scaling has been applied to the evolution of Mars, Venus, and super-Venus planets (Fraeman and Korenaga 2010; O'Rourke and Korenaga 2012), where they modeled the thermal evolution of crust, mantle, and core self-consistently by incorporating the influence of mantle processing on rheology and radiogenic heating. Our theoretical formulation follows Fraeman and Korenaga (2010) and O'Rourke and Korenaga (2012), and readers are referred to these studies for details. We summarize the key equations of our model in the following, focusing on the difference from previous models.

Our primary goal is to estimate the processing rate of the mantle to track the atmospheric evolution over time: melting by adiabatic decompression generates new crust and depleted lithospheric mantle (DLM), and volatiles are degassed during crust formation. A wide range of initial conditions were explored in O'Rourke and Korenaga (2012), but sub-solidus convection likely starts immediately after the surface of the mantle solidifies and degassing by percolation takes place (Section 2.2). We thus adopt the thermal structure and volatile concentrations predicted in Section 2 as initial conditions to model the history of mantle degassing in a self-consistent manner.

3.1.1. Scaling of stagnant lid convection

The vigor of convection is characterized by the Nusselt number: convective heat flux normalized by conductive heat flux. The Nusselt number, Nu , is calculated through a local stability analysis of the top thermal boundary layer (Korenaga 2009b), and we consider a condition where a bottom fraction of the boundary layer becomes gravitationally unstable. The Nusselt number is a function of the following variables; the mantle potential temperature, T_m , the temperature difference across the thermal boundary layer, ΔT , the mantle viscosity at the base of the thermal boundary layer, η_m , viscosity increase by dehydration, $\Delta\eta_w$, and the thickness of the depleted lithospheric mantle, h_{DLM} . The temperatures T_m and ΔT are solved from the energy balance of the mantle, and η_m is calculated as a function of potential temperature T_m and volatile concentrations in the mantle (Section 3.1.2). The evolution of h_{DLM} is controlled by the combination of mantle processing, delamination, and rewetting (Section 3.1.3).

DLM lacks volatiles, so the layer is more viscous than the source mantle. Thermal convection is suppressed with such effect of dehydration stiffening (van Thienen 2007), and its effect is stronger for a higher mantle potential temperature because of a thicker DLM. For a potential temperature of 1600 °C, the thermal boundary layer becomes thicker by a factor of ~5 than what is predicted in a scaling without dehydration (Korenaga 2009b).

Therefore, hotter mantle does not indicate more efficient thermal convection, and considering the effect of dehydration stiffening is crucial for estimating Nu . The thickness of the thermal boundary layer (TBL) can be calculated as $h_{\text{TBL}} = h_{\text{cm}}/Nu$, where h_{cm} is the depth of the convective mantle.

3.1.2. Mantle rheology

Viscosity η_m is described as a function of mantle potential temperature T_m and water content, c_w :

$$\eta_m = \begin{cases} A \exp\left(-\frac{E}{RT_m}\right) (\Delta\eta_w)^{1-c_w/c_0} & (c_w \leq c_0), \\ A \exp\left(-\frac{E}{RT_m}\right) \left(\frac{c_0}{c_w}\right) & (c_w > c_0), \end{cases} \quad (6)$$

where E is an activation energy (300 kJ mol⁻¹) and R is the universal gas constant. Viscosity is assumed to decrease linearly with the water concentration c_w (Mei and Kohlstedt 2000; Jain et al. 2019), but an exponential relation is adopted below a cutoff value, $c_0 = 50$ ppm, to prevent viscosity from reaching infinity. CO₂ might also reduce mantle viscosity, but because it is mostly degassed before the solidification of the mantle surface (Figure 2a), its effect is not considered. Viscosity contrast between dry and wet mantle of $c_w = c_0$ is assumed to be $\Delta\eta_w = 125$ (Hirth and Kohlstedt 1996; Mei and Kohlstedt 2000), and we adjust the preexponential constant, A , so that viscosity is 10¹⁹ Pa s at 1350 °C and $c_w = 0.04$ wt% as suggested from rock mechanics and the possibility of small-scale convection beneath oceanic lithosphere (Hirth and Kohlstedt 1996; Davaille and Jaupart 1994; Dumoulin et al. 1999). Initial volatile concentrations in the mantle are chosen from Figures 3c and 4b so that water oceans are absent at the beginning of solid-state convection stage.

3.1.3. Mantle processing and crust formation

The processing of the mantle is assumed to start at a depth where the temperature exceeds a dry solidus. A wet mantle has a lower solidus temperature (e.g., Kawamoto

and Holloway 1997), but the mantle would not be completely dehydrated until the melt fraction becomes sufficiently large, which becomes possible after crossing the dry solidus (e.g., Hirth and Kohlstedt 1996). The initial depth of mantle processing can thus be approximated by the dry solidus, and we parameterize the initial pressure of melting, P_i [GPa], as

$$P_i = \frac{T_m - 1423 \text{ K}}{100}. \quad (7)$$

The melting is assumed to stop where it reaches the base of the thermal boundary layer at $P_f = \rho_L g(h_{\text{TBL}} + h_c)$, where ρ_L is the lithosphere density and h_c is crustal thickness. The thickness of melting zone, h_p , can be described as

$$h_p = \frac{P_i - P_f}{\rho_L g}, \quad (8)$$

and assuming that downwelling is much more localized than upwelling, the volumetric rate of mantle processing is given by

$$\dot{V}_{\text{proc}} = \frac{2h_p u_{\text{conv}}}{h_{cm}} 4\pi r_p^2. \quad (9)$$

We adopt the scaling of Solomatov and Moresi (2000) for the average convective velocity beneath the stagnant lid:

$$u_{\text{conv}} = 0.38 \frac{\kappa}{h_{cm}} \left(\frac{Ra}{\theta} \right)^{\frac{1}{2}}, \quad (10)$$

where κ is the thermal diffusivity, Ra is the internal Rayleigh number, and θ is the Frank-Kamenetskii parameter. Convective velocity is influenced by the viscosity contrast across the stagnant lid and thus is a function of θ , which describes the degree of temperature dependency of viscosity:

$$\theta = \frac{E\Delta T}{RT_m^2}. \quad (11)$$

With a typical activation energy of $E = 300 \text{ kJ mol}^{-1}$ (Section 3.1.2), θ is ~ 14 for a mantle temperature of 1600°C , corresponding to a state immediately after magma ocean solidification.

Finally, we can calculate the volumetric melt productivity by multiplying \dot{V}_{proc} by the average melt fraction in the partial melt layer, $\bar{\phi}$, which is estimated based on the melt productivity by adiabatic decompression ($d\phi/dP = 0.1 \text{ GPa}^{-1}$). The growth of crust is equivalent to the volumetric melt productivity, and the volume of DLM increases by

$$\dot{V}_{\text{DLM}} = (1 - \bar{\phi})\dot{V}_{\text{proc}}. \quad (12)$$

Volatiles included in the processed mantle are released to the mantle, and the rate of degassing is proportional to \dot{V}_{proc} as well. Volatiles are highly incompatible at the depths where melting occurs, so we consider that the processed mantle becomes entirely dry after melting. It is noted that, in some previous studies, some large fraction of volatiles (90% of volatiles with 10% melting) was assumed to remain in the depleted mantle after melting (Noack et al. 2017; Dorn et al. 2018), which could underestimate the efficiency of degassing a factor of ~ 10 .

Crustal thickness continues to grow by mantle processing, but the thickness of the depleted lithospheric mantle is reduced by delamination or rehydration from the underlying mantle. When the thermal boundary layer is thinner than DLM ($h_{\text{DLM}} > h_{\text{TBL}}$), DLM would be eroded by convection (Figure 5), so we assume that such a delaminated fraction of DLM becomes mixed with the source mantle. Volatile concentrations in the mantle would be diluted as a result of mixing, and we adjust the concentrations accordingly (Fraeman and Korenaga 2010). Also, DLM is continuously rewetted by hydrogen diffusion from the underlying mantle. At each time step, a diffusion length $\Delta h_{\text{DLM}} = \sqrt{D_{\text{diff}}\Delta t}$ is calculated, and we assume that the lower Δh_{DLM} of DLM is reincorporated into the source mantle reservoir. The diffusion coefficient is taken from a parameterization given in Korenaga (2009):

$$D_{\text{diff}} = 6 \times 10^{-5} \text{ m}^2\text{s}^{-1} \times (-0.0027 + 2.19 \times 10^{-6}(T_m - 273)). \quad (13)$$

Diffusive rewetting is not considered when DLM thickness is reduced by delamination.

The initial thicknesses of crust h_c and depleted lithospheric mantle h_{DLM} are inherited from the last stage of magma ocean solidification. As discussed in Section 2.2, the uppermost partially molten layer becomes depleted in volatiles through melt escape by

percolation when the surface of the mantle solidifies (Figure 1d). The bottom pressure of the initially processed mantle is set to ~5 GPa, which is equivalent to ~170 km for Earth-size planets and ~80 km for $5M_E$ super-Earths.

3.1.4. Thermal evolution of the mantle

The thermal evolution of the convecting mantle is controlled by a balance of radiogenic heating, heating from the core, convective heat flux of the mantle, and latent heat of mantle melting:

$$\rho_m C_m V_{cm} \gamma_m \frac{dT_m}{dt} = \frac{4}{3} \pi (r_m^3 - r_c^3) Q_m + 4\pi (r_c^2 F_c - r_m^2 F_m) - \rho_L f_m L_m, \quad (14)$$

where ρ_m is the average density of mantle, C_m is the specific heat of mantle, V_{cm} is the volume of convective mantle, r_m and r_c are the radii of the mantle and core, Q_m is radiogenic heat production per unit volume, and L_m is the latent heat of mantle melting per unit mass. The constant γ_m is adopted to convert potential temperature to the average temperature of the mantle. The convective heat flux of the mantle, F_m , can be calculated from Nu (Section 3.1.1), and heat flux from the core, F_c , is derived using the scaling of Stevenson et al. (1983).

The convecting mantle represents the whole mantle for planets smaller than Earth-size, but the deeper region of super-Earths may be too viscous for convection because its major constituting mineral, post-perovskite, is expected to have viscosity higher by a few orders of magnitude than the terrestrial mantle (Tackley et al. 2013). We thus assume that only the region shallower than $P_{ppv} = 200$ GPa participates in thermal convection, and the value of V_{cm} is adjusted accordingly. Also, heat flux supplied to the convecting mantle would be limited to those from radiogenic decay because an extreme viscosity suppresses the cooling of the deeper mantle (Tackley et al. 2013; Dorn et al. 2018) and the core. With a post-perovskite viscosity of 10^{24} Pa s, which is $\sim 10^5$ higher than the that of the typical upper mantle, a local stability analysis of the bottom boundary layer (Stevenson et al. 1983) suggests that core cooling would be smaller by a factor of ~ 50 for

super-Earths. The presence of post-perovskite should thus lead to the efficient cooling of the shallower mantle while maintaining a hot deep interior, and such an effect is incorporated in our parameterized convection model.

3.2. Results

3.2.1. Evolution of Earth-size planets

Figure 6 shows the thermal evolution and degassing history of a typical Earth-size planet, which illustrates that mantle degassing continues but gradually decreases throughout its evolution (Figure 6a). The rate of mantle degassing can be understood from change in the thickness of the depleted lithospheric mantle h_{DLM} because mantle processing is reflected in the growth of DLM. During the first ~ 2 Gyr, the depth where mantle melting starts becomes deeper with time (Figure 6b) as mantle temperature increases by radioactive elements. Therefore, the net growth of DLM is observed, resulting in efficient degassing during this period (Figure 6c). In the subsequent stage, however, mantle temperature decreases with time, and the mantle is newly processed only as DLM delaminates or is rewetted by hydrogen diffusion. The growth of DLM by mantle melting and its delamination by convective erosion are occurring concurrently, and the two competing processes maintain h_{DLM} comparable to h_{TBL} (Figure 6a). As a result, degassing is less efficient than in the first 2 Gyr even though the temperature is higher, and its rate further slows down as the mantle cools down and convective velocity decreases (Figure 6c, 6d). It is noted that the rate of degassing experiences temporal fluctuation as a result of convective delamination. The bottom part of DLM becomes unstable when a sufficient fraction of DLM is rewetted by hydrogen diffusion. The resulting delamination of DLM allows mantle upwelling to reach a shallower depth, briefly triggering a larger degree of new melting.

We ran the model for different initial water concentrations, and the results show that, under the same $\text{H}_2\text{O}/\text{CO}_2$ ratio, the timing of water ocean formation depends weakly on the H_2O content ($x_{\text{H}_2\text{O}}$; Figure 6a). This is a result of two competing processes. Because

of a lower viscosity, convective velocity and thus the rate of mantle processing are higher for a wetter mantle: A mantle with a $\times 10$ higher H_2O concentration has a >10 times higher H_2O degassing rate. Yet, the amount of surface H_2O necessary to stabilize oceans also becomes increasingly larger: An order of magnitude larger CO_2 concentration would require a surface H_2O amount larger by 40 times (Figure 3a). These two effects nearly cancel out, and thus an oceanless world persists for a similar duration as long as the $\text{H}_2\text{O}/\text{CO}_2$ ratio remains the same. Plate tectonics would operate once the threshold is reached (Korenaga 2020), and mantle degassing would become more efficient afterwards. A wet surface is thus expected to be maintained during the subsequent evolution.

For an Earth-size planet, the atmosphere contains $\sim 5\%$ of the total H_2O budget immediately after the solidification of the mantle surface (Figures 2a), and an additional $\sim 6\%$ would be degassed in the next 1 Gyr during solid-state convection (Figure 6a). These estimates correspond to initial H_2O concentrations between 0.01 and 0.1 wt%. For an $\text{H}_2\text{O}/\text{CO}_2$ ratio of 0.67, $\sim 10\%$ of the total H_2O budget needs to reside at the surface to stabilize water oceans, and such a condition is satisfied only after 0.5-1 Gyr of evolution (Figure 6a). Because degassing slows down with time, and a planet with a smaller $\text{H}_2\text{O}/\text{CO}_2$ ratio requires an increasingly longer duration to form water oceans. The $\text{H}_2\text{O}/\text{CO}_2$ ratio therefore largely characterizes when oceans emerge on terrestrial planets.

We note that a thick depleted lithospheric mantle acts as a limiting factor for heat transport and thus mantle degassing (Figure 7). In a conventional scaling of stagnant lid convection, which does not consider dehydration stiffening, a higher mantle temperature leads to a thinner thermal boundary layer and thus a higher convective heat flux to promote cooling (the case of $\Delta\eta = 1$; Solomatov and Moresi 2000). Assuming such a scaling, the mantle is predicted to be efficiently processed from the beginning, crustal thickness would rapidly grow (Figure 7b), and water oceans would form within the first 100 Myr (Figure 7a). Neglecting dehydration stiffening thus severely overestimates the rate of mantle degassing. In reality, the presence of a dry DLM impedes convection, and the thermal boundary layer becomes thicker at higher mantle potential temperatures (Figure 6a, 6c). The Nusselt number Nu is inversely proportional to h_{TBL} , so a hotter mantle actually

results in lower convective heat flux (Korenaga 2009b). Because of inefficient cooling, excess radiogenic heat production heats up the mantle during the first ~ 1.5 Gyr (Figure 6c), and mantle temperature starts to decrease only after this period. The depth where mantle melting starts then becomes shallower with time, DLM thins, and with a weaker influence of dehydration stiffening, h_{TBL} starts to decrease (Figure 6a). A larger viscosity contrast between dry and wet mantle $\Delta\eta_w$ further delays the rate of cooling and degassing (Figure 7c), and thus better constraining the dependence of viscosity on water content would be important to accurately predict the timing of ocean formation. Some experimental studies suggest that the water dependence of mantle viscosity can be more drastic than assumed here (Faul and Jackson 2007).

3.2.2. Evolution of super-Earths

For larger planets, the formation of water oceans is predicted only under a restricted range of initial conditions, and even when oceans are expected to emerge, the formation would take longer than Earth-size planets (Figure 8a). As discussed in Section 2.3, more CO_2 is released during the magma ocean stage on larger planets (Figure 3b), so a greater amount of surface H_2O is required to stabilize oceans (Figure 3a). Yet, the amount of water residing at the surface is nearly independent of planetary mass at the beginning of the evolution (Figure 3a), and the amount of degassed volatiles during solid-state convection increases little with planetary size (Figure 9). Under the same initial volatile concentrations, larger planets thus may not have a sufficient amount of surface H_2O to stabilize oceans.

Larger planets have a higher rate of degassing, but mantle processing ceases earlier during solid-state convection. This is seen in Figure 9a, which compares the evolution of surface water for Earth-size and $5M_E$ super-Earth planets: the total amount of degassed water increases faster during the first 1-2 Gyr for a $5 M_E$ super-Earth, yet no degassing is observed during the subsequent period. The scaling for mantle processing can be derived from Equation (7):

$$V_{\text{proc}} \sim f_m \Delta t \sim \frac{h_p u_{\text{conv}} r_p^2}{h_{cm}} \Delta t. \quad (15)$$

The scaling for whole mantle convection ($h_{cm} = h_m$) has been provided in O'Rourke and Korenaga (2012) as $V_{\text{proc}} \sim M_p^{0.24} \Delta t$, and a similar scaling for super-Earths can be derived considering the effect of viscous post-perovskite in the deeper mantle: h_{cm} decreases to $P_{ppv}/(\rho_m g)$, and u_{conv} to $Ra^{1/2}/h_{cm} \sim (h_{cm} g)^{1/2} \sim 1$. Using the interior model of Valencia et al. (2006) ($r_p \sim M_p^{0.26}$) and $h_p \sim g^{-1}$, V_{proc} follows $r_p^2 \Delta t \sim M_p^{0.52} \Delta t$, and degassing is in general more efficient for larger planets during the first 1 Gyr of continuous degassing (Figure 9). The duration of degassing, Δt , however, shortens with increasing planetary size. Mantle processing takes space when the initial depth of melting, $h_i = P_i/(\rho_L g)$, is greater than the base of the thermal boundary layer h_{TBL} , but larger planets are less likely to meet such a condition. When super-Earths of different sizes under the same thermal state are considered, the initial depth of melting scales with g^{-1} , whereas the thickness of the thermal boundary layer is scaled as

$$h_{\text{TBL}} = \frac{h_{cm}}{Nu} \sim \frac{h_{cm}}{Ra^{1/3}} \sim g^{-1/3}. \quad (16)$$

With increasing planetary mass, the initial melting depth becomes shallower at a faster rate than TBL, and activating mantle processing becomes more difficult. Consequently, mantle processing ceases earlier for larger planets, and even though the rate of mantle degassing is higher during the early stage, the total amount of degassed volatiles throughout their evolution does not change appreciably with planetary size (Figure 9b). We note that the limited processing of the mantle for larger planets is in a broad agreement with previous studies of 2-D mantle convection models (Dorn et al. 2018). A larger fraction of the water budget is thus expected to remain in the mantle for super-Earths without degassing to the surface, precluding the formation of water oceans.

3.3 Discussion: Hydrogen escape and long-term habitability

Hydrogen escape can be important for the timing of ocean formation when the initial H_2O concentration in the mantle ($x_{\text{H}_2\text{O}}$) is small and the planet is close to the star. The

efficiency of hydrogen escape depends on the mixing ratio of water vapor at the top of the atmosphere, and diffusion is the rate-controlling process when the mixing ratio is below ~2-3%, assuming a similar EUV flux to the present-day Sun and a heating efficiency of 30% (Watson et al. 1981). Although the luminosity of younger stars may be higher than the present-day Sun, hydrogen escape from a planet without a significant envelope would always be limited by diffusion, and thus a larger flux size would not affect the rate of water loss. The 1-D radiative-convective model (Appendix A.2. Nakajima et al. 1992) predicts that the mixing ratio is smaller than 1% for conditions considered in this study (Figure 10), and thus hydrogen escape would be limited by diffusion (e.g., Catling and Kasting 2017). For a $5M_E$ super-Earth with a water vapor mixing ratio of 0.003, the total amount of hydrogen escape during a 1 Gyr period could be as large as 4×10^{18} kg, which is equivalent to 4×10^{20} kg of H_2O . This amount is comparable to the total amount of degassed water for $x_{H_2O} < 0.04$ wt% (Figure 9a), and thus ocean formation would be delayed, or in some case precluded, compared to what is predicted in Figures 6a and 8a. Diffusion-limited flux is linearly proportional to the mixing ratio, so under a weaker net stellar radiation, the amount of water lost by hydrogen escape is negligible because the atmosphere contains a smaller amount of water vapor (Figure 10b).

We calculate mantle degassing under solid-state convection and estimate the likelihood and timing of ocean formation for various initial volatile concentrations, considering the loss of water by hydrogen escape. With a stronger net stellar radiation and thus a higher water mixing ratio, hydrogen escape becomes significant and prohibits the formation of ocean formation at a small initial H_2O concentration (x_{H_2O}). On early Venus, for example, if oceans are absent at the time of magma ocean solidification, degassing during solid-state convection does not supply a sufficient amount of water to stabilize oceans, and the surface would lack oceans throughout its evolution under $x_{H_2O} < 0.08$ wt%, equivalent to a total water inventory of ~2.4 ocean mass (Figure 11a). Although the threshold x_{H_2O} is higher for larger planets (Figure 11c), the general trend remains the same. On the other hand, if x_{CO_2} is small and thus water oceans are formed from the beginning, the mantle would operate under plate tectonics, and efficient mantle degassing would maintain oceans during the subsequent period. Therefore, for planets receiving $>240 \text{ W m}^{-2}$ of

radiation with a small initial H₂O content, whether water oceans exist immediately after magma ocean solidification characterizes the long-term existence of water oceans and thus the habitability of planets (Figure 11).

This could provide an explanation for why Venus and Earth had divergent evolutionary paths: Venus may have continued to lack oceans throughout its history because of the comparable rates of mantle degassing and hydrogen escape, whereas Earth has had water oceans and plate tectonics since the solidification of the mantle surface. The net solar radiation received by early Venus is considered to be close to the tropospheric radiation limit (Hamano et al. 2013), but even if Venus was not in a runaway greenhouse state, water oceans are not guaranteed on early Venus. Limited mantle degassing under stagnant lid convection also implies that the Venusian mantle could still be wet, and that the amount of hydrogen escape and thus its by-product oxygen could be smaller than previously predicted. Such a small amount of oxygen could be consumed by the oxidation of surface rocks without invoking other removal mechanisms (e.g., Kurosawa 2015), given that the Venusian surface is likely to have continually been resurfaced by magmatism even in the mode of stagnant lid convection (O'Rourke and Korenaga 2015).

Hydrogen escape has little influence on ocean formation when the stellar radiation is weaker and/or a mantle contains a greater amount of H₂O (Figure 11b, 11d). In such cases, planets with the same H₂O/CO₂ ratio would develop water oceans on a similar timescale regardless of the absolute value of initial H₂O concentrations as discussed in Section 3.2.1. For planets resembling early-Earth, oceans would exist from the beginning for H₂O/CO₂>1.5, and the formation timescale becomes increasingly longer as the H₂O/CO₂ ratio decreases because degassing slows down as the mantle cools (Figures 6c, 8c; Section 3.2.2). Degassing eventually discontinues, and thus the surface would never acquire a sufficient amount of water to stabilize oceans for H₂O/CO₂ ratios lower than 0.4. Such threshold ratios for ocean formation increase with planetary mass and net stellar radiation. These values are within the estimate for the bulk silicate Earth (2.4 ± 1.0 ; Hirschmann and Dasgupta 2009), so if terrestrial exoplanets have similar volatile concentrations to Earth, the presence of water oceans is not guaranteed, especially for

super-Earths. Planets may have a different fate with a slight difference in their volatile concentrations, and the habitability of planets would depend largely on their size and the $\text{H}_2\text{O}/\text{CO}_2$ ratio of the mantle, in addition to the distance from the central star.

Although its effect is not considered in our study, the rate of hydrogen escape during the magma ocean phase is much higher than discussed here. A hot surface allows most of the surface water budget to exist as vapor, increasing the mixing ratio at the top of the atmosphere. Because a molten surface only persists for less than 0.1 Myr in a period after the final giant impact, water loss by hydrogen escape is negligible in the scope of Section 2. Hydrogen escape during the accretionary phase, however, could lead to a significant water loss (Zahnle et al. 1988) and would be important when constraining the water budget of terrestrial planets from the perspective of the chemical composition of planetary building blocks.

4. DISCUSSION and CONCLUSIONS

4.1. Additional processes to promote mantle degassing

During the transition from a magma ocean to solid-state convection, we assumed that the solid-dominated layer quickly solidified by the Rayleigh-Taylor instability within the layer. As discussed in Section 2.1, the density structure of the solid-dominated layer is gravitationally unstable, so a newly-solidified cold material would continuously be mixed with the pre-existing solid-dominated layer (Maurice et al. 2017). Because the cooling of the surface melt-dominated layer is much faster than the solid-dominated one, the thermal evolution of the solid-dominated layer would mostly be governed by the cold downwelling of newly-solidified materials, which is likely to be more efficient than cooling by solid state convection (Lebrun et al. 2013; Zahnle et al. 2015). Its effect, however, remains unquantified because its modeling requires the implementation of solidification by adiabatic compression. Viscosity also changes drastically as partially molten materials solidify during the downwelling. Fully incorporating these effects would require some numerical efforts, so we cannot rule out the possibility that the solid-dominated layer

remains superadiabatic at the time of surface solidification. In such a case, remaining heat may be released by episodic eruptions, and the amount of surface water could be slightly higher than predicted in Section 2.

Mantle degassing may also be promoted by tidal heating. It has been suggested that tidal heating inhibits the cooling of a solid-dominated magma ocean (Zahnle et al. 2015), and if the cooling timescale becomes long enough, the percolation of pore melt may transport a significant amount of trapped volatiles to the surface. Quantifying the timescale of mantle cooling, however, is not straightforward. As discussed in the previous paragraph, the cooling of the solid-dominated layer is mostly controlled by the overlying melt-dominated layer, and any heat added by tidal dissipation could efficiently be transported upwards by the Rayleigh-Taylor instability within the layer. A simple 1-D model would be insufficient to address this complex transition period, and further investigation is warranted. Also, the presence of satellites is not always guaranteed, and even if a satellite exists, its size is not necessarily going to be as large as the Earth's moon. The results shown here would thus serve as a baseline understanding on volatiles degassing, and we plan to address additional processes in future studies.

4.2. Plate tectonics and the removal of thick CO₂ atmosphere

Plate tectonics maintains a habitable atmospheric composition over geological time (Berner 2004), so the presence of surface water, a key to plate tectonics, is crucial for the long-term habitability of the planet. Its role is particularly important during the early stage of evolution because the majority of the CO₂ budget is degassed during magma ocean solidification. When $p_{\text{CO}_2} > 100$ bar, its greenhouse effect would maintain a surface temperature over 100 °C (Abe and Matsui 1988; Abe 1993a), which may prevent life forms similar to those that exist on Earth from emerging. The removal of such a thick CO₂ atmosphere likely requires rapid plate motion (Sleep and Zahnle 2001; Zahnle et al. 2007), so although this study has focused on the formation of water oceans, stabilizing water oceans is only one of the requirements for creating a habitable environment.

The efficiency of CO₂ removal from the atmosphere depends on plate velocity, and when the mantle has a pyrolitic composition, plate velocity could be slower for a hotter mantle than in the present-day (Korenaga 2006, 2010b). Miyazaki and Korenaga (2022) suggests that a chemically heterogeneous mantle resulting from magma oceans solidification (Appendix A.4) would develop a thinner depleted lithospheric mantle and thus allows faster plate velocity than a pyrolitic mantle. Whether such chemical differentiation occurs in a solidifying magma ocean is a critical factor in developing a habitable environment, although it depends on several unconstrained physical properties and is subjected to further investigation (Solomatov and Stevenson 1993; Solomatov et al. 1993; Xie et al. 2020).

Plate velocity also depends strongly on the degree of mantle hydration (Korenaga 2010b), and the idea of a wet mantle after magma ocean solidification is also consistent with the removal of the thick CO₂ atmosphere. Some previous models of magma ocean degassing have often assumed that the mantle would be entirely dehydrated during solidification (Elkins-Tanton 2008; Hamano et al. 2013; Lebrun et al. 2013; Salvador et al. 2017), but a dry mantle would have a slower plate motion than in the present-day, despite the hotter mantle temperature in the past (Korenaga 2006). If so, the removal of CO₂ over >100 bar would become highly inefficient (Miyazaki and Korenaga 2022), which fails to explain the emergence of the moderate climate by the early Archean. The wet mantle and thus rapid plate motion also agree with recent geochemical modeling, which suggests the efficient recycling of the surface material in the Hadean (Rosas and Korenaga 2018; Hyung and Jacobsen 2020). Studies on magma ocean have often suffered from a lack of observational constraints, but we may be able to obtain new insights into this period by taking into account the aftermath of a magma ocean.

4.3. Conclusions

We estimated the likelihood of ocean formation and its timing for planets with various volatile concentrations by solving for the evolution of mantle degassing during and after magma ocean solidification. Considering the rheological transition of a partially molten medium, a magma ocean is predicted to retain a significant fraction of the H_2O inventory, whereas CO_2 , which is much less soluble to magma, would mostly reside in the atmosphere after the solidification. When the $\text{H}_2\text{O}/\text{CO}_2$ ratio is lower than a threshold, surface water exists entirely as water vapor as a result of a strong greenhouse effect, and water oceans would be absent on the surface. Larger planets are less likely to develop water oceans because more greenhouse gases are emitted to the atmosphere during the solidification process.

If water oceans do not form at the beginning of solid-state convection, planets most likely operate under stagnant lid convection, severely limiting mantle degassing during the subsequent evolution. The rate of mantle degassing decreases and eventually reaches zero as the mantle cools and the thermal boundary layer grows, so the majority of H_2O is expected to be retained in the mantle throughout its evolution. The total amount of degassed water is nearly independent of planetary mass under the same volatile concentrations, but because a CO_2 atmosphere is thicker and thus a larger amount of surface water is necessary to form oceans for larger planets, the threshold $\text{H}_2\text{O}/\text{CO}_2$ ratio for ocean formation increases with planetary size. The threshold also increases with net stellar radiation as a larger amount of water vapor is contained in the atmosphere. In addition, water loss by hydrogen escape may preclude ocean formation for planets receiving strong radiation and containing a small amount of H_2O . These would not be the case once water oceans form at the surface because the mode of mantle convection would likely change to plate tectonics, promoting mantle degassing during the subsequent evolution.

If the volatile contents of rocky exoplanets are similar to those of the bulk silicate Earth, mantle degassing during magma ocean would not supply a sufficient amount of H_2O when planets are larger and/or receive stronger radiation than Earth. Exoplanets could be richer in volatiles than Earth, and indeed, exoplanets of larger size are known to have an

atmosphere of smaller metallicity (Benneke et al. 2019a). Yet, the redox evolution of rocky planets is not well understood, and if hydrogen existed in a reduced form, a high H/C ratio may not necessarily result in the formation of water oceans. An increasing number of observations for the exoplanet atmosphere is expected with upcoming missions, but the detection of water vapor does not necessarily indicate the presence of water oceans. In addition to the habitable zone, planetary mass and the $\text{H}_2\text{O}/\text{CO}_2$ ratio are also important parameters in predicting the likelihood of oceans on exoplanets. Because the threshold $\text{H}_2\text{O}/\text{CO}_2$ ratio for ocean formation is lower for smaller planets receiving weaker radiation, such planets are a more promising candidate to search for potential life, although they may be more difficult to detect and observe the atmosphere.

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APPENDIX: METHOD DETAILS

A.1. The thermal structure of the melt-dominated layer

Mantle potential temperature, T_m , is calculated to be consistent with the depth of the melt-dominated layer d . The magma ocean is considered to be adiabatic as a result of rapid convection, and T_m is obtained by integrating the adiabatic temperature gradient from the

bottom of the melt-dominated layer to the surface (Figure S1). The adiabatic gradient is described as

$$\left(\frac{dT}{dP}\right)_s = \frac{\alpha' T}{\rho c_p'}, \quad (A1)$$

where thermal expansivity, α' , and specific heat, c_p' , include the effect of phase transition. They are obtained by differentiating the Gibbs free energy of peridotite described in Miyazaki and Korenaga (2019b), and this thermodynamic model is also used to determine other thermodynamic properties, such as density and melt fraction at each depth. The relation between T_m and d for an Earth-size planet is shown in Figure 2a.

The surface temperature T_s is calculated using a 1-D atmospheric model (Appendix A.2) and depends on the atmospheric mass and composition, calculated from Equations (1-3), and the net outgoing infrared radiation ($F_{\uparrow}(\tau = 0)$ in Appendix A.2). The outgoing infrared radiation is the sum of absorbed stellar radiation and convective heat flux, F_{conv} , given by (Zahnle et al. 1988; Solomatov 2007):

$$F_{\text{conv}} = 0.089 \frac{k(T_m - T_s)}{d} Ra^{1/3}, \quad (A2)$$

where k is thermal conductivity and Ra is the Rayleigh number of the melt-dominated layer. Equation (A2) assumes that heat flux from the solid-dominated layer can be ignored because of a much higher viscosity than the melt-dominated one. The Rayleigh number is defined as

$$Ra = \frac{\alpha \rho g (T_m - T_s) d^3}{\kappa \eta_l}, \quad (A3)$$

where α is thermal expansivity, κ is thermal diffusivity, and η_l is the viscosity of the melt-dominated layer. The viscosity η_l does not change significantly until magma experiences rheological transition and thus is set to a constant of 1000 Pa s. The net stellar radiation would be $\sim 167 \text{ W m}^{-2}$ for the early Earth and $\sim 319 \text{ W m}^{-2}$ for the early Venus with an albedo of 0.3 and a solar radiation 30% weaker than the present day.

A.2. Atmospheric model

The surface temperature and the total amount water vapor in the atmosphere are estimated using a 1-D radiative-convective model based on Nakajima et al. (1992). We consider two layers within the atmosphere: stratosphere and troposphere, where the temperature is controlled by different heat transport mechanism. Whereas radiative equilibrium governs the stratosphere, the thermal structure in the troposphere is characterized by convective heat transport from the bottom. The atmosphere is assumed to be plane-parallel and is transparent to stellar radiation, but it is opaque to infrared radiation regardless of the wavelength. Although this assumption of gray atmosphere may underestimate the surface temperature during the earlier stage of magma ocean, the calculated temperatures are similar for the gray and nongray cases for surface temperatures below 200 °C (Salvador et al. 2017). For simplicity, the effects of clouds are neglected in our model.

The thermal profile of the stratosphere can be written as a function of optical thickness, τ :

$$\sigma_B T(\tau)^4 = \frac{1}{2} F_{\text{net}} \left(\frac{3}{2} \tau + 1 \right), \quad (A4)$$

where σ_B is the Stefan-Boltzmann constant and F_{net} is the net infrared flux emitted from the top of the atmosphere. Upward, F_{\uparrow} , and downward radiation fluxes, F_{\downarrow} , are also calculated as

$$F_{\uparrow}(\tau) = \frac{1}{2} F_{\text{net}} \left(\frac{3}{2} \tau + 2 \right), \quad (A5)$$

$$F_{\downarrow}(\tau) = \frac{1}{2} F_{\text{net}} \left(\frac{3}{2} \tau \right). \quad (A6)$$

Optical thickness τ is defined so that it increases towards the Earth's surface:

$$\tau(z) = - \int_{\infty}^z \kappa \rho_g dz. \quad (A7)$$

Opacity, κ , depends on the atmospheric concentrations of greenhouse gases, and for the atmosphere which has a molar fraction of x_i of gas species i , κ is given by

$$\kappa = \frac{1}{\mu} \sum_i \kappa_i x_i \mu_i, \quad (A8)$$

where $\bar{\mu}$ is the mean molar mass of gas, κ_i and μ_i are the opacity and molar mass, respectively, of species i . For an infrared absorption coefficient, we use 10^{-2} for H_2O , 1.3×10^{-4} for CO_2 , and 0 for N_2 (Abe and Matsui 1985). All species are assumed to be transparent to stellar radiation, and thus the downward radiation F_{\downarrow} is set to 0 at $\tau = 0$. The composition of the atmosphere is considered to be uniform in the stratosphere.

In the troposphere, the thermal structure is controlled by the moist adiabatic lapse rate, which is described as

$$\frac{dT}{dP} = \frac{\bar{\mu}}{\rho_g C_p} \frac{1 + \frac{qL}{RT}}{1 + \frac{qL^2}{C_p RT^2}}, \quad (\text{A9})$$

where ρ_g is the mean gas density, C_p is heat capacity, q is water mixing ratio, and L is the latent heat of the water. Water mixing ratio q is set to be saturated, which is likely in a CO_2 dominated atmosphere, and is obtained from the water phase diagram. Although the moist adiabatic lapse rate is derived assuming a completely saturated atmosphere, we assume that relative humidity in the troposphere is fixed at a value smaller than 1 (e.g., Nakajima et al. 1992).

The tropopause, a boundary between the stratosphere and troposphere, is determined as a height that the radiation energy is balanced. The upward radiation flux emitted from the troposphere can be calculated as

$$F_{\uparrow}(\tau) = -\sigma_B T^4 + \int_{\tau_b}^{\tau} e^{-(\tau' - \tau)} \frac{d}{d\tau'} (\sigma_B T(\tau')^4) d\tau', \quad (\text{A10})$$

where τ_b is the optical depth at the bottom of the atmosphere. We assume that the upward flux at the ground is the blackbody radiation of the ground temperature ($F_{\uparrow, \text{surf}} = -\sigma_B T_{\text{surf}}^4$). At the optical depth of the tropopause, τ_p , the values of $F_{\uparrow}(\tau_p)$ calculated from Equations (A5) and (A10) should be identical to satisfy the energy balance. We search for the profiles of temperature and water vapor content that agree with energy conservation. The temperature and the water vapor mixing ratio are smoothly connected at the tropopause.

A.3. The initial thickness of depleted lithospheric mantle

The thickness of the partial melt layer when the melt-dominated layer disappears (Figure 5) is estimated based on an adiabatic temperature profile with a critical melt fraction of 0.4 at the surface. The timescale for the Rayleigh-Taylor instability is predicted to be faster than the rate of solidification (Maurice et al. 2017; Miyazaki and Korenaga 2019a), and thus the upper mantle is likely to have an adiabatic thermal gradient when the surface of the mantle solidifies. The partial melt layer would experience degassing by percolation, so the bottom pressure of the partially melt layer also becomes the initial base pressure of depleted lithospheric mantle (Section 3.1.2).

The adiabatic gradient is expressed as (Mckenzie 1984):

$$\left(\frac{dT}{dP}\right)_s = \phi \frac{\alpha_l T}{\rho_l c_{p,l}} + (1 - \phi) \frac{\alpha_s T}{\rho_s c_{p,s}} - \frac{T \Delta S}{c_p} \left(\frac{d\phi}{dP}\right)_s, \quad (A11)$$

where c_p is specific heat per unit mass and ΔS is entropy change upon melting. Subscripts l and s denote the melt and liquid phases, respectively. Values adopted for thermal expansivity are 3.0 and $4.6 \times 10^{-5} \text{ K}^{-1}$ for α_s and α_l , and, for density, ρ_l and ρ_s are taken to be 2900 and 3300 kg m^{-3} , respectively. We adopt 1000 $\text{J kg}^{-1} \text{ K}^{-1}$ for c_p and 300 $\text{J kg}^{-1} \text{ K}^{-1}$ for ΔS . Melt fraction at a given temperature and pressure is calculated using the Rhyolite-MELTS model (Gualda et al. 2012), and the term $d\phi/dP$ in Equation (A11) is calculated so that changes in T and ϕ are consistent with thermodynamics.

A.4. Mantle composition after magma ocean solidification

The depth of the partial melt layer and the thickness of DML depend on the mantle composition after magma ocean solidification, which could be different from the present-day mantle as a result of differentiation (Elkins-Tanton et al. 2003; Maurice et al. 2017; Miyazaki and Korenaga 2019a). We assume a homogeneous pyrolitic mantle in this study, but if the mantle experiences differentiation during magma ocean, Miyazaki and Korenaga

(2019a) suggests that the mantle would develop a chemically heterogeneous structure, consisting of high-Mg# pyroxenite and Fe-rich blobs. The settling and sedimentation of a liquidus mineral (Mg-rich bridgmanite; de Koker et al. 2013) at the base of the melt-dominated layer produces a lower mantle comprised of high-Mg# and SiO₂-rich materials, whereas the upper mantle is left with residual Fe-rich dense materials. This structure is gravitationally unstable and is subject to small-scale Rayleigh-Taylor instabilities with a length scale shorter than 100 km. Fe-rich blobs may break down by some degree during the downwelling, but because of high solid viscosity, the two components are unlikely to be completely homogenized in a short period. At the end of solidification, Fe-rich materials would thus have existed as small blobs embedded in a high-Mg# pyroxenite matrix.

The Rhyolite-MELTS model suggests that such a pyroxenite would be mostly dehydrated at ~2 GPa under a mantle potential temperature of 1600 °C (see Extended Data Figure 1 in Miyazaki and Korenaga 2022 for details). On the other hand, a homogeneous pyrolytic mantle becomes dehydrated by ~5 GPa under the same mantle potential temperature. This is because the high-Mg# pyroxenite has lower FeO and CaO contents and lacks low-solidus minerals including clinopyroxene and plagioclase. Decompressional melting would thus start at a shallower depth for the chemically heterogeneous mantle, and the amount of volatiles degassed by percolation (Figures 2-4) is expected to be smaller than that of a pyrolitic mantle. The long-term evolution of surface water (Figures 6-9), however, would not be significantly different between these two cases because the chemically heterogeneous mantle should be homogenized after some time. The solidus temperature would decrease as high-Mg# pyroxenite and Fe-rich blobs are mixed, and thus the amount of surface water would quickly increase and eventually become similar to the case of a pyrolitic mantle.

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Figure 1. Schematic illustration of the upper mantle and atmosphere during the solidification of magma ocean. (a) Magma ocean undergoes the rheological transition from the bottom, forming a solid-dominated layer. Melt would be trapped in pore spaces with volatiles, and thus volatiles would not concentrate in the melt-dominated layer. Variables used in Equation (3) are labeled in corresponding reservoirs. (b) Magma ocean continues to solidify, and the thickness of the solid-dominated layer grows. Rapid convection and the Rayleigh-Taylor instability would efficiently cool the melt- and solid-dominated layers, respectively, both creating an adiabatic thermal profile. Because of the rapid onset of the instability, volatiles would efficiently be sequestered in the deeper region. (c) When the melt-dominated layer disappears, convective heat flux plummets, and the surface temperature drops below the mantle solidus. When sufficient amount of water is degassed, oceans may form. Because melt percolation is faster than solid-state convection, percolation contributes to additional degassing. (d) Erupted melt forms a crust at the top, leaving a dry depleted layer below the crust. Because the deeper region is completely solidified, percolation would not occur, and the most of the mantle would remain hydrated.

Figure 2. The evolution of (a) volatiles in the atmosphere and (b) the thermal state as a function of the melt-dominated layer depth. As solidification proceeds, the layer thickness decreases, so the system evolves from left to right. (a) The ratio of degassed volatiles to the total volatiles budget for H₂O (solid) and CO₂ (dashed). Initial mantle concentrations of 0.04 wt% (red) and 0.1 wt% (yellow) are tested for both volatiles. For H₂O, the two values correspond to 1.2 and 3 ocean masses, respectively. (b) The mantle potential temperature (black solid) and the surface temperature (colored solid, left axis) with the melt fraction of magma ocean at the surface (dashed, right axis). The snapshots of the mantle structure are shown in Figure 1, with triangles indicating the corresponding panels.

Figure 3. (a, b) Atmospheric mass of (a) H_2O and (b) CO_2 when the surface of the magma ocean solidified, as a function of the initial volatile concentration. Colors denote different planetary sizes: Mars- (red), Earth-size (M_E , yellow), and $10M_E$ super-Earths (cyan). The value of $M_{\text{H}_2\text{O}}$ is similar among planets of different size sizes, and all lines are very close to each other. In (a), the minimum amount of H_2O necessary to stabilize water oceans is also plotted, assuming that the $\text{H}_2\text{O}/\text{CO}_2$ mass ratio is 1. Mars- and Earth-size planets would have sufficient amount of water to form oceans, whereas $10M_E$ super-Earths would lack oceans immediately after the solidification of the mantle surface. (c) Conditions necessary for water ocean formation as a function of initial H_2O and CO_2 concentrations. Colored lines separate the two regimes: the formation (upper left) and absence of water oceans (lower right) when the mantle surface solidified. With the same volatile concentrations, smaller planets are more likely to develop water oceans. Shaded rectangles represent the estimated volatile concentrations in the bulk silicate Earth (BSE) from Hirschmann and Dasgupta (2009) and Korenaga et al. (2017).

Figure 4. (a) The minimum amount of surface H_2O required to form water oceans as a function of M_{CO_2} . Results are calculated for different levels of net stellar radiation: 165 W m^{-2} (solid, early Earth) and 330 W m^{-2} (dashed, early Venus). Colors denote different planetary sizes: Earth-size (M_E , yellow) and $10 M_E$ super-Earth (cyan). (b) The same as Figure 3c but with different levels of net stellar radiation. Colored lines separate the two regimes: the formation (upper left) and absence of water oceans (lower right) when the planetary surface solidified. Planets further away from the central star are more likely to develop water oceans, but planetary size plays a larger role in characterizing ocean formation. Shaded rectangles represent the estimated volatile concentrations in the bulk silicate Earth (BSE) from Hirschmann and Dasgupta (2009) and Korenaga et al. (2017).

Figure 5. Schematic illustration of the structure of terrestrial planets, and parameters described in Section 3.1 are labeled. When the depleted lithospheric mantle layer is thicker than the thermal boundary layer, the excess layer would delaminate and be mixed with the source mantle. The values of h_{DML} shown in the subsequent figures are those after delamination. An enlarged view of the melting zone is also shown as an inset on the right.

Figure 6. The thermal evolution and degassing history of Earth-size planets under stagnant lid convection. (a) Total amount of degassed water, (b) thicknesses of crust (dashed) and depleted lithospheric mantle (solid), (c) convective velocity beneath the lid (solid) and the rate of water degassing averaged over 50 Myr (dashed, light-colored), and (d) mantle potential temperature are shown for three different initial H_2O concentrations: 0.01 (red), 0.04 (yellow), and 0.1 wt% (green), corresponding to a total H_2O mass of 0.3, 1.2, and 3 ocean masses, respectively. Initial conditions are adopted from the final results of the magma ocean degassing model described in Section 2. Markers in (a) show the time when a threshold amount of surface H_2O is degassed to stabilize water oceans. The $\text{H}_2\text{O}/\text{CO}_2$ ratios of 0.66 (star) and net stellar radiation of 330 W m^{-2} are adopted here. With an H_2O concentration of 0.04 wt%, p_{CO_2} corresponds to ~ 330 bar. Once water oceans form at the surface, the mode of convection likely changes to plate tectonics, so modeling results after ocean formation would underestimate convective heat flux and the rate of mantle degassing. The gray line in (b) describes the thickness of thermal boundary layer. As a result of convective delamination, h_{DLM} becomes almost identical to h_{TBL} throughout the planetary evolution.

Figure 7. The same as Figure 6 but for an initial H_2O concentration of 0.04 wt% with two different values for viscosity contrast between dry and wet mantle: $\Delta\eta_w = 1$ (cyan) and 125 (yellow). A larger viscosity contrast results in slower degassing, and thus an oceanless surface would be maintained for a longer time. The case with $\Delta\eta_w = 1$ does not exhibit any dehydration stiffening, and thus convection and mantle degassing are more efficient than the other two cases, although it is an unrealistic assumption. While

depleted lithospheric mantle is actively eroded by thermal convection and its thickness is kept thin, crust grows rapidly to the point that, after 700 Myr, it becomes thick enough to halt mantle processing (shown in (c)). Markers in (a) show the time when a threshold amount of surface H₂O is degassed to stabilize water oceans.

Figure 8. The thermal evolution and degassing history of $5M_E$ super-Earths under stagnant lid convection. (a) Total amount of degassed water, (b) thicknesses of crust (dashed), depleted lithospheric mantle (solid), and thermal boundary layer (gray) (c) convective velocity beneath the lid (solid) and the rate of water degassing averaged over 50 Myr (dashed, light-colored), and (d) mantle potential temperature are shown for three different initial H₂O concentrations: 0.01 (red), 0.04 (yellow), and 0.1 wt% (green). Markers in (a) show the time when a threshold amount of surface H₂O is degassed to stabilize water oceans, where the H₂O/CO₂ ratio of 1.1 (triangle) and net stellar radiation of 330 W m⁻² are assumed. For a H₂O concentration of 0.04 wt%, H₂O/CO₂=1.1 corresponds to $p_{\text{CO}_2} \sim 500$ bar, and the total H₂O inventory is 6 ocean masses because the mantle mass of a $5M_E$ super-Earth is ~ 5 times that of Earth.

Figure 9. (a) The total amount of degassed water for Earth-size (solid) and $5M_E$ super-Earth planets (dotted) with three different initial H₂O concentrations in the mantle $x_{\text{H}_2\text{O}}$: 0.01 (red), 0.04 (yellow), and 0.1 wt% (green). Mantle degassing is faster during the first 1 Gyr but ceases after ~ 2 Gyr for a $5M_E$ super-Earth. (b) The amount of degassed water for different planetary sizes with H₂O concentrations of 0.01 (red) and 0.04 wt% (yellow). The bars show the amount of degassed water during three different periods: before the solidification of a magma ocean (gray), before 1 Gyr (light red/yellow) and 5 Gyr of solid-state convection (dark red/yellow). The minimum amount of surface H₂O for ocean formation is also plotted for initial CO₂ concentrations x_{CO_2} of 0.02 (black), 0.04 (dark gray), and 0.07 wt% (light gray).

Figure 10. The mixing ratio of H₂O vapor at the top of the atmosphere as a function of (a) the initial CO₂ concentration in the mantle and (b) as of net stellar radiation. A planet with water oceans is assumed, and an oceanless one would have a smaller mixing ratio than shown here. Colors indicate planetary mass: Earth-size (red), $2M_E$ (yellow), and $5M_E$ super-Earth (green). (a) Solid and dashed lines represent net stellar radiation of 330 and 165 W m⁻², respectively. Dotted gray line describes a critical ratio above which hydrogen escape is limited by the magnitude of EUV flux, and for the range of stellar radiation considered here, hydrogen escape is limited by diffusion. (b) An atmosphere containing 1 bar of N₂ and 100 bar of CO₂ is assumed here. Water loss by diffusion-limited hydrogen escape is estimated as well for an Earth-size planet using a binary diffusion coefficient of $b_{H_2} = 1.46 \times 10^{21} \text{ m}^{-1} \text{ s}^{-1}$ (Catling and Kasting 2017). Water is assumed to readily convert to hydrogen at the top of the atmosphere, and thus the values should be treated as an upper bound estimate for water loss.

Figure 11. Contours showing the timing of ocean formation for Earth-size (top) and $5M_E$ super-Earth planets (bottom). Results are calculated for three different levels of net stellar radiation: 330 (left) and 165 W m⁻² (right). Water oceans form immediately after magma ocean solidification and exist throughout the evolution of planets in regions left of black lines, whereas oceans would be absent for longer than 5 Gyr under conditions shaded in light gray. Rectangles represent the estimated volatile concentrations in the bulk silicate Earth (BSE) Hirschmann and Dasgupta (2009) and Korenaga et al. (2017).

Figure S1. (a) Adiabatic temperature profiles with potential temperatures of 2000 K (red), 2500 K (orange), and 3000 K (green) for pyrolitic composition. The solidus and liquidus of pyrolite (dashed), along with temperature where melt fraction is 40% (dotted), are also shown for comparison. (b) Melt fraction of the adiabatic temperature profiles. Dashed line describes the critical melt fraction for the rheological transition.