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Solubility of water in peridotite liquids and the prevalence of steam atmospheres on rocky planets



Paolo A. Sossi^{a,*}, Peter M.E. Tollan^a, James Badro^b, Dan J. Bower^c

- ^a Institute of Geochemistry and Petrology, ETH Zürich, 8092 Zürich, Switzerland
- ^b Institut de Physique du Globe de Paris, Université de Paris, 75005 Paris, France
- ^c Center for Space and Habitability, Universität Bern, 3012 Bern, Switzerland

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ABSTRACT

Atmospheres are products of time-integrated mass exchange between the surface of a planet and its interior. On Earth and other planetary bodies, magma oceans likely marked significant atmosphere-forming events, during which both steam- and carbon-rich atmospheres may have been generated. However, the nature of Earth's early atmosphere, and those around other rocky planets, remains unclear for lack of constraints on the solubility of water in liquids of appropriate composition. Here we determine the solubility of water in 14 peridotite liquids, representative of Earth's mantle, synthesised in a laser-heated aerodynamic levitation furnace. We explore oxygen fugacities (fO_2) between -1.9 and +6.0 log units relative to the iron-wüstite buffer at constant temperature (2173 \pm 50 K) and total pressure (1 bar). The resulting fH2O ranged from nominally 0 to 0.027 bar and fH2 from 0 to 0.064 bar. Total H2O contents were determined by transmission FTIR spectroscopy of doubly-polished thick sections from the intensity of the absorption band at 3550 cm⁻¹ and applying the Beer-Lambert law. The mole fraction of water in the liquid is found to be proportional to $(fH_2O)^{0.5}$, attesting to its dissolution as OH. The data are fitted by a solubility coefficient of 524 \pm 16 ppmw/bar^{0.5}, given a molar absorption coefficient, ε_{3550} , of 6.3 \pm 0.3 m²/mol for basaltic glasses or 647 \pm 25 ppmw/bar^{0.5}, for a preliminary $\varepsilon_{3550} = 5.1 \pm 0.3$ m²/mol for peridotitic glasses. These solubility constants are roughly 10 - 25% lower than those for basaltic liquids at 1623 K and 1 bar. Higher temperatures result in lower water solubility in silicate melts, wholly offsetting the greater depolymerisation of peridotite melts that would otherwise increase H₂O solubility relative to basaltic liquids at constant temperature. Because the solubility of water remains high relative to that of CO₂, steam atmospheres are rare, although they may form under moderately oxidising conditions on telluric bodies, provided sufficiently high H/C ratios prevail.

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1. Introduction

Owing to the heat generated by planetary impacts (Melosh, 1990) and short-lived radioactive isotopes (Bhatia, 2021), magma ocean-forming events were likely commonplace during the main planet-building stage (up to \sim 100 Myr after CAI; Chao et al., 2021). Such events are significant because they likely result in the wholesale resetting of any pre-existing primary atmosphere captured from the solar nebula (Genda and Abe, 2005), and engender secondary atmosphere growth owing to the efficiency with which the molten planetary interior transports mass to the atmosphere. The Rayleigh number of a liquid terrestrial mantle, \sim 10 30 ,

yields mixing timescales of the order of days- to weeks (Solomatov, 2000), rapid with respect to atmospheric cooling timescales, meaning equilibrium between the interior and its atmosphere is readily attained. The corollary is that heat transfer is also efficient, meaning magma oceans are periodic, short-lived phenomena occurring over $\sim 10^3$ to 10^7 yr (Bower et al., 2019; Lebrun et al., 2013).

The nature of an atmosphere generated during a magma ocean stage on a given planet is key to establishing its long-term stability to photochemical destruction, condensation or escape, and whether it is amenable to the development of life (Catling and Kasting, 2017). The identities and partial pressures of gas species that comprise an atmosphere depend upon the *i*) bulk planetary endowment of atmosphere-forming elements (typically H, C, N, O and S), and *ii*) the solubility of these gases in the underlying magma ocean as a function of the intensive variables (pressure, temperature, composition) at the interface. The bulk silicate Earth

^{*} Corresponding author.

E-mail address: paolo.sossi@erdw.ethz.ch (P.A. Sossi).

(BSE) acts as a reference model for the estimation of abundances of atmosphere-forming elements (e.g., Hirschmann, 2018; Marty, 2012). Provided appropriate solubility laws are known, the composition of a model atmosphere in equilibrium with a magma ocean can be determined.

In this manner, Sossi et al. (2020) demonstrated that, at 2173 K, a CO-rich atmosphere would have formed around the Earth for oxygen fugacities (fO_2) near the iron-wüstite (IW) buffer. Such atmospheres would cool and partially condense to give rise to CO_2 -N₂-rich atmospheres at ambient temperatures. They highlighted that the type of atmosphere depends on fO_2 , which, at a given temperature and pressure, determines the ratios of fCO/fCO_2 and fH_2/fH_2O . Using a similar approach, Gaillard et al. (2022) highlighted that, should equilibrium between the magma ocean and atmosphere take place at very low fO_2 (Δ IW-3 or lower), then H_2 -rich atmospheres would form, whereas SO_2 -rich atmospheres could prevail under oxidising conditions ($>\Delta$ IW+2). However, the accuracy of these models is contingent upon the extrapolation of solubility laws of volatile species in molten peridotite from experimental data in molten basalt, and at much lower temperatures.

The determination of the solubilities of major volatiles in silicate melts has been largely confined to moderate- to highpressures relevant to the Earth's crust and upper mantle (Dalou et al., 2019; Dixon et al., 1995; Hirschmann et al., 2012), with only a handful at 1 bar (Newcombe et al., 2017). However, atmospheric pressures above magma oceans on rocky planets are likely to be below several hundred bar (e.g., Elkins-Tanton, 2008; Gaillard et al., 2022; Sossi et al., 2020). Moreover, the basaltic (and more evolved) compositions examined in these studies melt below \sim 1773 K (Gaillard et al., 2001; Moore et al., 1998). By contrast, the Earth's mantle and those of other rocky planets are broadly peridotitic, not basaltic, in composition (Unterborn et al., 2020) with liquidus temperatures of \sim 2023 K at 1 bar (Takahashi, 1986). The difficulty of achieving such temperatures experimentally, and in quenching peridotitic compositions into homogeneous glasses (cf. Green, 1975) has thus far prevented solubility measurements in magma ocean-relevant liquids.

Here we explore the occurrence of steam (H₂O-rich) atmospheres above magma oceans, a state often imposed in planetary cooling models (Abe and Matsui, 1985; Fegley et al., 2016; Lebrun et al., 2013). In order to do so, molten peridotite representative of Earth's mantle was equilibrated at 2173 K in a controlled atmosphere achieved by mixing Ar-CO₂-H₂ and O₂ gases in an aerodynamic laser-heated levitation furnace (ALLF) that permitted rapid quenching. The water contents and speciation in the resulting glasses were determined by Fourier-Transform InfraRed (FTIR) spectroscopy. Together, these data are used to devise a water solubility model for peridotite liquids and applied to understand atmospheres generated in equilibrium with magma oceans.

2. Methods

2.1. Experimental

The quenched peridotite liquids investigated in this study are those reported in Sossi et al. (2020). Briefly, a composition modelled on that of KLB-1 (Takahashi, 1986) was synthesised by mixing high-purity (>99.95% pure) powders of MgO, SiO₂, CaCO₃, Fe₂O₃ and Al₂O₃ together in the desired proportions. Following 24 h of decarbonation at 1273 K in air, ~20 mg pieces of pressed powder were placed in the ALLF (Institut de Physique du Globe de Paris, France). All samples were heated above their liquidus by a 125 W CO₂ laser (λ = 10.6 μ m) to 2173 \pm 50 K for 30 s in pure O₂ before being quenched and re-melted under the desired gas mixture at the same temperature, as recorded by optical pyrometry. Gas flow rates were set to a precision of \pm 2% relative

by Bronckhorst mass flow controllers delivering 92% Ar/8% H₂ and 92% Ar/8% CO₂ mixtures to the furnace nozzle, resulting in oxygen fugacities from Δ IW-1.9 to Δ IW+6.0. Gas flow rates were in the range 800 - 1000 sccm, resulting in nozzle velocities of 8 - 10 m/s (cf. Badro et al., 2021). Badro et al. (2021) and Young et al. (2022) showed that isotopic fractionation during evaporation in levitation furnaces results from the advective transport of the gas around the bead, resulting in partial pressures of evaporating species from the melt that are \sim 75% lower than their equilibrium partial pressures. For a flowing gas representing an infinite reservoir to impose fO_2 , fH_2 and fH_2O , the total pressure (P_T) around the sphere must be equal to 1 bar. At 2173 K, the temperatures are sufficiently high that the gas behaves ideally, such that the fugacity of species i, f_i , is equal to its partial pressure, p_i , calculated according to the equation $p_i = X_i P_T$, where X is the mole fraction. The fugacities at the experimental temperature and pressure were calculated by Gibbs Free Energy minimisation using FactSage. Samples were quenched below the glass transition by switching off the power to the laser, resulting in cooling rates of \sim 800 K/s. A summary of the experimental conditions can be found in Table 1. As reported by Sossi et al. (2020), the two most reduced samples experienced \sim 10% Fe loss due to evaporation with a concomitant decrease in Si/Mg by \sim 4% relative. The mean composition for the remaining peridotite glasses, by weight, is 46.53(26)% SiO₂, 4.37(4)% Al₂O₃, 8.44(29)% FeO^T (all Fe as FeO), 38.05(42)% MgO, and 2.06(5)% CaO. Both Na₂O and K_2O are <0.01%.

2.2. Fourier-Transform InfraRed spectroscopy

Water contents of peridotite glasses were determined by FTIR spectroscopy at the Institut für Geologie, Universität Bern, Switzerland (Tollan and Hermann, 2019). The glasses were mounted in epoxy and cut to a total thickness of $1(\pm 0.005)$ mm. The samples were then polished with diamond paste on both sides to a grade of 1 µm. Measurements were conducted in transmission mode under unpolarised light, using a Bruker Tensor II spectrometer with an SiC Globar infrared source and a KBr beamsplitter, coupled to a Bruker Hyperion 3000 microscope with a dry air-purged sample chamber. The infrared light transmitted through the sample was collected as a function of wavelength by a liquid N2-cooled mercury cadmium telluride (MCT) LN-D316 detector, with an integration area of $400 \times 15 \ \mu m$ per spot, 128 scans and 8 cm⁻¹ resolution over a spectral range from 6000 cm⁻¹ to 1000 cm⁻¹. For each sample, in-situ FTIR transects were performed across the diameter of the glasses, amounting to up to 10 spots/sample at a spacing of roughly 100 µm/point with the background determined every 20 points (Table S1, Fig. S1). The data were processed using the Bruker OPUS software, in which atmospheric absorbance from H₂O and CO₂ was corrected for using its built-in algorithm. Owing to the low absorption intensity of the 3550 cm⁻¹ feature used for total water quantification and the steeply curved intrinsic absorption of the peridotitic glass, a polynomial baseline correction with \sim 15 points was performed manually and independently three times, each \sim 6 months apart. The contribution of the 3550 cm⁻¹ band to the absorbance, was quantified by its peak intensity relative to the local baseline within the assigned spectral window, which ranged from 3750 \pm 50 cm⁻¹ to 3100 \pm 50 cm⁻¹ for all spectra.

The Beer-Lambert law states that the total abundance of water dissolved in the glass, $C(H_2O)$, is related to the measured total absorbance of the IR-active band at 3550 cm⁻¹ (I_{3550}) produced by OH- and molecular H_2O stretching modes (Stolper, 1982), its molar mass $M(H_2O)$ (in kg/mol), the path length of transmitted light through the absorbing medium (d in m) and its density (ρ in kg/m³), by a proportionality constant, the molar absorption cross section for that band (ε_{3550}), with units of m^2/mol .

$$C(H_2O)(ppm) = 10^6 \frac{I_{3550}M(H_2O)}{d\rho\varepsilon_{3550}}$$
 (1)

Here, $M(H_2O) = 0.018015$ kg/mol, $d = 10^{-3}(\pm 5 \times 10^{-6})$ m, and $ho = 2991 \pm 11 \text{ kg/m}^3$ at the glass transition temperature, $T_{\rm g} =$ 1012 ± 6 K (Dingwell et al., 2004), calculated using thermal expansivities and partial molar volumes from (Lange and Carmichael, 1987) and compressibilities from (Kress and Carmichael, 1991), assuming $Fe^{3+}/\Sigma Fe = 0$ and anhydrous compositions for all samples. The calculated density extrapolated to 2173 K is in excellent agreement with synchrotron X-ray absorption measurements of peridotitic liquids, $\rho = 2700 \text{ kg/m}^3$ (Sakamaki et al., 2010). The molar absorption cross section has been determined for the 3550 cm⁻¹ band in a wide range of silicate compositions, from basaltic to rhyolitic (Shishkina et al., 2014; Stolper, 1982) yet that for peridotitic compositions is unknown. Stolper (1982) derived a value of 6.71 \pm 0.67 m²/mol for ε_{3550} for basaltic- to rhyolitic glasses, while Shishkina et al. (2014) determined $\varepsilon_{3550} = 5.92 \pm 0.40 \text{ m}^2/\text{mol for}$ the alkali basaltic and basanitic compositions, suggesting little dependence of ε_{3550} on melt composition over this range. On the other hand, Mercier et al. (2010) concluded on the basis of SIMS and FTIR analyses of basanitic glasses that increasing non-bridging oxygen over tetrahedral oxygens (NBO/T) in the melt leads to a decrease in ε_{3550} to roughly 4.5 m²/mol at NBO/T \sim 1. The water contents of the peridotite glasses were too low to permit accurate independent determination of C(H2O) by Karl-Fischer Titration or by Elastic Recoil Detection Analysis (ERDA), which has a detection limit near ~50 - 100 ppm (Bureau et al., 2009; Withers et al., 2012). However, thanks to the recent development of a novel rapid-quench multi-anvil technique, Bondar et al. (2021) were able to synthesise a set of hydrous peridotitic glasses with MgO contents up to 43.5 wt.% and water contents ranging from 0 to 5 wt.% H₂O (Bondar et al., 2022). On the basis of the hydrous peridotitic glass with the lowest water content, Bondar (2022) obtained a preliminary molar absorption coefficient for ε_{3550} of 5.1 \pm 0.3 m²/mol. For completeness, we present H₂O concentrations calculated according to a mean value $\varepsilon_{3550} = 6.3 \pm 0.3 \text{ m}^2/\text{mol}$ (n=8) for basaltic glasses, and the preliminary value of 5.1 \pm 0.3 m²/mol for peridotitic glasses. The uncertainties on these quantities are propagated to those on $C(H_2O)$, reported as 1σ .

3. Results

A summary of the results is shown in Table 1 and raw, unnormalised spectra for each sample are presented as Extended Data. Fig. 1 shows the calculated gas fugacities of the major species present in the gas mixes used in the experiments at a constant temperature of 2173 K. Owing to the binary mixture of CO_2 -H₂ (diluted with 92% Ar), CO_2 is the dominant gas species for oxygen fugacities 1.5 log units above the IW buffer, while fH_2 predominates below IW. In the interval between IW and Δ IW+1.5, H₂O is the prevailing gas species in the C-O-H system. The calculated fH_2O at 2173 K ranges from nominally 0 bar (Per-1, -2 and -5) to 0.027 bar (Per-10), while fH_2 extends up to 0.064 bar (Per-12) (Table 1).

The only H-bearing absorption band observed in the glasses is that located at 3550 cm⁻¹. Hence, we find no evidence for the dissolution of molecular H_2 at 4130 cm⁻¹. Within a sample, the normalised absorbance of the 3550 cm⁻¹ band varies by <7% relative, with an average of $2.8 \pm 1.7\%$, indicating water is homogeneously distributed in the glasses (Table S1). Moreover, I_{3550} scales with fH_2O , reaching a maximum at \sim 0.5 log units above the iron-wüstite buffer (Fig. 2). The corresponding water contents of peridotite glasses as determined by eq. (1) range from 32.5 ± 2.3 ppmw for $\varepsilon_{3550} = 6.3$ m²/mol or 40.2 ± 3.2 ppmw for $\varepsilon_{3550} = 5.1$ m²/mol (Per-5) to 137.4 ± 7.3 ppmw ($\varepsilon_{3550} = 6.3$ m²/mol) or

Table 1 Experimental conditions, calculated gas fugacities, measured infrared absorbance intensity of the 3550 cm⁻¹ band and calculated H₂O contents

Temperature Time ΔIW^a fH_2	ΔIW^a fH_2	fH_2			fH_20	# of	Fit 1	Fit 2	Fit 3	Mean	1α	Mean	1α	Corrected Mean	Corrected Mean
	(bar)			9	ar)	points	Absorbance	Absorbance	Absorbance	$\varepsilon = 6.3 \text{ m}^2/\text{mol}$		$\varepsilon = 5.1 \text{ m}^2/\text{mol}$		$\varepsilon = 6.3 \text{ m}^2/\text{mol}$	$\varepsilon = 5.1 \text{ m}^2/\text{mol}$
										(ppmw)		(ppmw)		(ppmw)	(bpmw)
2173 ± 21 30 5.97 0 0	30 5.97 0 0	5.97 0 0	0 0	0		3	0.036	0.037	0.038	35.4	2.0	43.7	3.0	2.9	3.5
0	0	0	0 0	0		7	0.039	0.040	0.039	37.6	2.0	46.5	3.1	5.1	6.3
± 29 32 3.42 1.28E-05	3.42 1.28E-05	1.28E-05		0	.00071	7	0.053	0.054	0.050	20.0	3.2	61.8	4.6	17.5	21.7
\pm 59 60 3.46 9.88E-07	3.46 9.88E-07	9.88E-07		5.	74E-05	7	0.041	0.040	0.039	38.2	2.2	47.2	3.2	5.7	7.1
± 25 36 3.46 5.17E-08	3.46 5.17E-08	5.17E-08		3.0)2E-06	9	0.036	0.033	0.033	32.5	2.3	40.2	3.2	0.0	0.0
\pm 23 25 3.37 2.77E-05	3.37 2.77E-05	2.77E-05		0.0	315	8	0.053	0.055	0.057	52.6	3.2	65.0	4.7	20.1	24.8
\pm 13 27 3.23 8.41E-05	3.23 8.41E-05	8.41E-05		0.0	38	7	690.0	0.070	0.072	67.2	3.8	83.1	9.6	34.7	42.9
2197 ± 13 31 2.94 0.00024 0.00	2.94 0.00024	0.00024		0.00	9/	8	0.088	0.090	0.094	86.7	5.2	107.1	9.7	54.2	6.99
\pm 15 30 2.03 0.0016	2.03 0.0016	0.0016		0.0	18	6	0.115	0.120	0.118	112.5	6.3	139.0	9.4	80.0	8.86
± 21 30 0.64 0.012	0.64 0.012	0.012		0.0	27	6	0.141	0.142	0.143	135.8	7.1	167.7	10.9	103.3	127.5
± 16 28 -0.77 0.041	-0.77 0.041	0.041		0.0	018	10	0.145	0.142	0.144	137.4	7.3	169.7	11.1	104.8	129.5
0.064	0.064	0.064		0	8200	10	0.133	0.132	0.138	128.4	7.3	158.6	10.8	95.9	118.5
\pm 13 10 3.23 8.41E-05	3.23 8.41E-05	8.41E-05		0	0.0038	8	990.0	990.0	0.068	63.7	3.5	78.7	5.2	31.2	38.6
2148 ± 41 120 3.23 8.41E-05 0.	3.23 8.41E-05	8.41E-05		0	0038	8	0.064	0.074	0.071	9.99	9.9	82.3	2.6	34.1	42.1
				l											

IW buffer calculated according to the revised expression of Hirschmann (2021) and hence the delta values shown here are 0.5 log units lower than reported in Sossi et al. (2020)

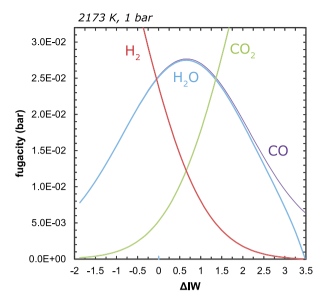


Fig. 1. Equilibrium speciation of the gas phase as calculated in *FactSage* for the CO_2 : H_2 mixtures used in the experiments, from 0.11 to pure CO_2 , in 92% Ar, at 2173 K and 1 bar, as a function of fO_2 relative to the IW buffer (Δ IW) of Hirschmann (2021).

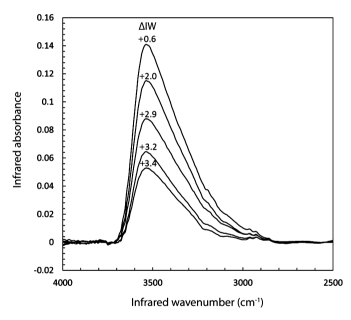


Fig. 2. Background-normalised FTIR spectra (see section 2.2) showing the intensity of the OH stretching absorbance band (\sim 3550 cm $^{-1}$) for glasses over a range of fO_2 , from Δ IW+0.6 to +3.4 (where IW = oxygen fugacity defined by the iron-wüstite buffer at 2173 K).

 169.7 ± 11.1 ppmw ($\varepsilon_{3550} = 6.3 \text{ m}^2/\text{mol}$) (Per-11). By examining the water contents determined amongst three glasses synthesised at the same experimental conditions (Per-TS1, Per-7, Per-TS2) but for different times (10 s, 30 s, and 120 s, respectively), equilibrium distribution of H_2O between the liquid and the surrounding gas phase is achieved in under 10 s. The H_2O content in each of the three samples overlaps within uncertainty, with a relative standard deviation of 5.6%, on par with that observed for a single sample across the three different fitting schemes. Based on these replicate experimental samples and baseline fitting procedures, the total precision on $C(H_2O)$ is estimated to be $\pm 5\%$ relative (Table 1).

When $C(H_2O)$ is plotted against fH_2O (Fig. 3), it is observed that dissolved water contents do not tend to nil at $fH_2O = 0$ as expected thermodynamically, but rather reach a constant value

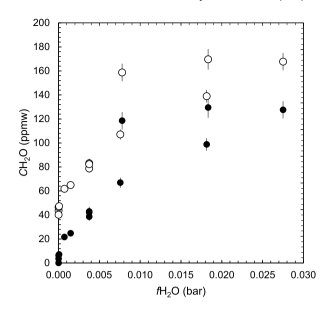


Fig. 3. Water contents of the glasses (in ppmw using $\varepsilon_{3550} = 5.1 \text{ m}^2/\text{mol}$) expressed as a function of its fugacity in the gas phase at 2173 K. White points indicate uncorrected data, showing the water contents remain at \sim 40 ppmw for nominal $fH_2O = 0$, and black points show the data corrected according to eq. (2).

of \sim 32 - 40 ppmw. As the $C(H_2O)$ must formally be equal to zero when $fH_2O=0$, the $C(H_2O)$ contents of all samples are renormalised to those of samples with $fH_2O=0$;

$$C(H_2O)_{norm} = C(H_2O)_{meas} - C(H_2O)_0$$
 (2)

Here we adopt the lowest $C(H_2O)$ value measured in the glasses, 32.5 ppmw ($\varepsilon_{3550} = 6.3 \text{ m}^2/\text{mol}$) or 40.2 ppmw ($\varepsilon_{3550} = 5.1$ m^2/mol), as the normalising value, $C(H_2O)_0$. Alternatively, the non-negligible absorbance for samples with nominal fH2O of zero could result from a constant fH2O background during glass synthesis. However, when excluding these samples on a plot of $\log f H_2 O$ vs. $\log C H_2 O$, the data define a slope of ~ 0.33 , well below the value of 0.5 expected for dissolution of OH in the melt. This explanation is therefore untenable, as it implies a gas-liquid reaction that does not relate to any physical dissolution stoichiometry. Moreover, H₂O contents of the ALPHAGAZ O₂ and Ar-CO₂ mixtures used in the experiments were <3 ppmw, which would translate into ~1 ppmw dissolved in the melt using pre-existing calibrations (Newcombe et al., 2017). As the experiments were not performed within a chamber, it cannot be excluded that the melt bead interacted with small quantities of air during the experiment, as levitation sometimes becomes unstable during laser power and/or gas fluctuations. Therefore, the remnant absorption at 3550 cm⁻¹ in melts synthesised under $fH_2O = 0$ is attributed to background contributions from either the environment or nonequilibrium water in the glass. That this water likely resides in the glass is supported by the fact that the measurements were performed in transmission mode, traversing the sample, and that the spectral peak shape and position are similar to those glasses that contain higher H₂O contents. It is noteworthy that Newcombe et al. (2019) also observed 6 - 22 ppmw H₂O dissolved in glasses equilibrated in nominally dry CO-CO2 gas mixtures, which they attributed to kinetic inhibition of H2O degassing. The corrected values in Table 1 will be used for the remainder of the study.

The stoichiometry of the dissolution of water in silicate melts can be inferred from the dependence of CH_2O on fH_2O . The dependence of water content in a liquid in equilibrium with a gas phase can be described:

$$C(H_2O) = \alpha [f(H_2O)]^{\beta}$$
 (3)

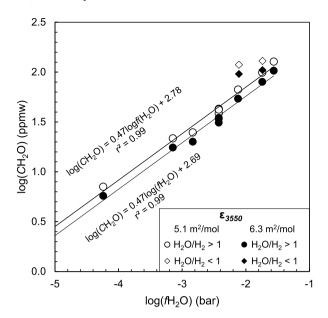


Fig. 4. Fit of equation (4) to the water contents (in ppmw) determined in the glass and the corresponding water fugacities in the gas phase. Water contents are calculated with $\varepsilon_{3550} = 5.1 \text{ m}^2/\text{mol}$ (white) and with $\varepsilon_{3550} = 6.3 \text{ m}^2/\text{mol}$ (black). Points shown in diamonds correspond to samples synthesised under high fH_2 and were not included in the regression. Samples that were synthesised at $fH_2O = 0$ (Per-1, Per-2, Per-5) were excluded from the plot.

Where α is the solubility coefficient and β the stoichiometric coefficient of the reaction; Henry's Law is recovered for $\beta = 1$. The values of constants α and β are determined empirically by taking the logarithm of both variables (Fig. 4);

$$\log C(H_2O) = \beta \log[f(H_2O)] + \log \alpha \tag{4}$$

A least-squares regression of the data (n=9) to eq. (4) yields $\beta=0.47\pm0.02$ and, for a peridotite glass molar absorption coefficient $(\varepsilon_{3550}=5.1~\text{m}^2/\text{mol})$, $\log\alpha$ of $2.78\pm0.05~(\alpha=605^{+68}_{-60}~\text{ppmw/bar}^{0.47})$ while for $\varepsilon_{3550}=6.3~\text{m}^2/\text{mol}$, $\log\alpha=2.69\pm0.07~(\alpha=490^{+68}_{-60}~\text{ppmw/bar}^{0.47})$ (Fig. 4). Two samples synthesised at the lowest fO_2 and highest fH_2 (Per-11 and Per-12) plot to markedly higher CH_2O than those predicted by eq. (4) (see Fig. 4) and were excluded from the fitting procedure, along with samples synthesised at $fH_2O=0$ (Per-1, Per-2 and Per-5). The accuracy of equation (4) is contingent on the assumption that the quantity of H in the glasses depends exclusively on fH_2O ; one that will be discussed below.

4. Discussion

4.1. Mechanisms and equilibria of H dissolution

The value of β obtained from the fit to the experimental data is approximately 0.5, a solubility dependence known as Sieverts' Law (Sieverts, 1929). This implies dissociation of the dissolved species in the liquid phase, a mechanism that was well-established in earlier experiments of water solubility in silicate melts in equilibrium with a gas or fluid phase (Dixon et al., 1995; Shishkina et al., 2010; Stolper, 1982; Tomlinson, 1956). More recently, similar results were obtained by Newcombe et al. (2017), who also observed a dependence of H_2O content of lunar- and anorthite-diopside liquids equilibrated in a 1 bar furnace with CO_2 - H_2 gas mixtures on $f(H_2O)^{0.5}$. Taken together, these solubility relationships are strong evidence for the dissolution of water into silicate melts by

$$H_2O(g) = H_2O(l)$$
 (5)

and the homogeneous equilibrium in the melt for its dissociation into hydroxyl groups

$$H_2O(l) + O(l) = 2OH(l)$$
 (6)

Where, at equilibrium, eq. (5) and (6) combine to yield:

$$K_{(7)} = \frac{a \, (\text{OH})^2}{f \, (\text{H}_2\text{O}) \, a \, (\text{O})}.$$
 (7)

Here, a refers to the activity and states that the mole fraction of OH dissolved in the melt should be proportional to $(f \, H_2 \, O)^{0.5}$. However, as shown in Fig. 4, the $H_2 \, O$ contents of samples Per-11 and Per-12 do not abide by this theoretical expectation. This mismatch implies the contribution of a second equilibrium;

$$H_2(g) = H_2(l)$$
 (8)

because H_2 is the predominant H-bearing species in the gas phase in equilibrium with Per-11 and Per-12 (Table 1). Given that OH is expected to be the prevailing species at low $C(H_2O)$ (Newcombe et al., 2017) and no molecular hydrogen absorption band was observed in the spectra, a dissociation reaction equivalent to that of eq. (6) can be written for H_2 ,

$$H_2(l) + 2O(l) = 2OH(l)$$
 (9)

leading to:

$$K_{(10)} = \frac{a \, (\text{OH})^2}{f \, (\text{H}_2) \, a \, (\text{O})^2} \tag{10}$$

Reaction (10) illustrates how the presence of $H_2(g)$ contributes to the dissolution of H as OH in peridotite liquids. The dataset (excluding samples Per-1, -2 and -5) is therefore fit by least squares regression to equation (11) that accounts for the contributions of both fH_2 and fH_2O in the atmosphere, given a $\beta_{H_2O}=\beta_{H_2}=0.5$ dependence (eq. (7), (10)) and varying α_{H_2O} and α_{H_2} . This yields an expression for the solubility of H_2O in peridotite liquids at 2173 K and 1 bar total pressure:

$$C(H_2O)(ppm) = (\alpha_{H_2O})f(H_2O)^{0.5} + (\alpha_{H_2})f(H_2)^{0.5}$$
 (11)

For $\varepsilon_{3550}=6.3~\text{m}^2/\text{mol}$, we find $\alpha_{\text{H}_2\text{O}}=524\pm16~\text{ppmw/bar}^{0.5}$ and $\alpha_{\text{H}_2}=183\pm14~\text{ppmw/bar}^{0.5}$ whereas with $\varepsilon_{3550}=5.1~\text{m}^2/\text{mol}$, these values are $\alpha_{\text{H}_2\text{O}}=647\pm25~\text{ppmw/bar}^{0.5}$ and $\alpha_{\text{H}_2}=225\pm13~\text{ppmw/bar}^{0.5}$. The goodness of fit with respect to the data is shown in Fig. 5. The solubility constant for H₂O, is roughly 10% (5.1 m²/mol) to 25% (6.3 m²/mol) lower than that observed for a lunar green glass (LG) and anorthite-diopside eutectic (ADeu) melts equilibrated at 1623 K (Newcombe et al., 2017). Unlike Newcombe et al. (2017), who are unable to detect any increase in OH solubility in the presence of H₂(g) in LG and ADeu melts, peridotite liquids in equilibrium with H₂(g) are able to dissolve OH with a solubility constant a factor of \sim 3 lower than that for H₂O(g).

Comparison with models based on pre-existing data for H_2O solubility should only be considered qualitative because the experiments fall well outside their temperature calibration range. Nevertheless, for completeness, the model of Moore et al. (1998) (973 < T [K] < 1473) underpredicts the H_2O contents by a factor $\sim\!\!3$, while that of Iacono-Marziano et al. (2012) (1373 < T [K] < 1673) overpredicts them by approximately the same factor (Table S2). Both serve to show that current models are inadequate for extrapolation beyond their calibration ranges, and applicability of such models to magma ocean-like conditions and compositions is not recommended.

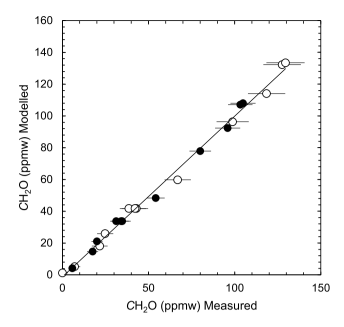


Fig. 5. Comparison between the measured water content of silicate melts determined in this work (H₂O, Measured) and the modelled value (H₂O, Modelled) determined using eq. (11) for $\varepsilon_{3550} = 5.1 \text{ m}^2/\text{mol}$ (white) and $\varepsilon_{3550} = 6.3 \text{ m}^2/\text{mol}$ (black). The fits are H₂O (ppmw, modelled) = 1.01 × H₂O (ppmw, measured) – 1.25, $r^2 = 0.995$ and H₂O (ppmw, modelled) = 1.02 × H₂O (ppmw, measured) – 1.6, $r^2 = 0.994$, respectively.

4.2. Thermodynamic constraints on the solubility of water at low fH_2O

Quantifying the influence of temperature and composition on the solubility of water in silicate melts requires evaluation of eq. (7). The lack of thermodynamic data for individual melt species restricts estimates of the equilibrium constant, $K_{(7)}$, to semi-empirical means (Hirschmann et al., 2012). The quantity of OH dissolved is given by:

$$X (OH) = \frac{\sqrt{K_{(7)} f(H_2 O) a(O)}}{\gamma (OH)}$$
 (12)

Where X refers to the mole fraction and γ to the activity coefficient. The activity coefficient of OH in silicate melts is not known, and is therefore assumed to be unity. Iacono-Marziano et al. (2012) considered that the activity of oxygen in silicate melts, O, is related to the ratio of Non-Bridging Oxygens (NBO) to total oxygen (Σ O). Here we define:

$$a(0) = \frac{\text{NBO}}{\Sigma O},\tag{13}$$

where NBO and O are calculated following Iacono-Marziano et al. (2012). Given that $f \, H_2 O = (X(OH)/\alpha_{XH_2O})^2$, (cf. eq. (3), where X denotes α calculated by mole fraction of H_2O in the melt) and $a = X\gamma$, one can rewrite eq. (7) as:

$$K_{(7)} = \frac{(\gamma \text{ (OH) } \alpha_{\text{XH}_2\text{O}})^2}{a(\text{O})}$$
 (14)

This analysis illustrates that, all else being equal, α is expected to correlate positively with aO for a given pressure and temperature. That is, liquids with a greater proportion of Non-Bridging Oxygens should be able to dissolve more water at a given fH_2O , temperature and pressure. The peridotite liquids investigated here have aO=0.43, while the LG and ADeu compositions of Newcombe et al. (2017) have lower NBO/ ΣO , 0.35 and 0.28, respectively, reflecting their more polymerised melt structure. Because α is expected to increase with $(aO)^{0.5}$, which is opposite to observations, we

conclude that composition plays only a subordinate role in determining the solubility constant of eq. (7) (*cf.* Shishkina et al., 2010; lacono-Marziano et al., 2012), though systematic experimentation is required to further investigate compositional dependence at 1 bar total pressure.

By the same logic, to understand the effect of melt composition on the dissolution of OH derived from H_2 , a reaction equivalent to eq. (14) may be written using $K_{(10)}$:

$$K_{(10)} = \frac{(\gamma \text{ (OH) } \alpha_{XH_2})^2}{a(0)^2}.$$
 (15)

Equation (15) illustrates that the solubility of OH in melts in equilibrium with $H_2(g)$ should increase proportional to aO. As a result, relative to the LG and ADeu compositions (Newcombe et al., 2017) with aO of 0.35 and 0.28, respectively, a 25 – 50% increase in OH solubility in peridotite liquid, with aO of 0.43, is expected. This increase is qualitatively compatible with the observation of a contribution of OH dissolution from the H_2 gas in peridotite liquids (cf. Fig. 4, eq. (11)) but not in the experiments of Newcombe et al. (2017). It is therefore likely that highly depolymerised melt compositions could incorporate large proportions of OH in their structure, even in the absence of H_2O in the gas phase.

The other key variable, temperature, is poorly constrained because there are very few studies that report results of a wide experimental temperature series for a given composition at fixed $f\,H_2O$ or total pressure. Here we evaluate the dependence of water solubility on temperature at low total pressures and low $f\,H_2O$ such that the only dissolved species is OH (i.e., where eq. (7) applies). In evaluating $K_{(7)}$ from eq. (14), the value of α_{H_2O} is calculated on a molar basis, where $\alpha_{X(H_2O)} = \alpha_{H_2O} \times (5.55 \times 10^{-6})$, $\gamma(OH)$ is assumed to be unity, and aO is calculated on the basis of melt composition (eq. (13)). The values of $K_{(7)}$ are calculated in this manner for five silicate melt compositions (Table 2). These values are converted to a Gibbs Free Energy change of reaction (7) by:

$$\Delta G_{(7)} = -RT ln K_{(7)} \tag{16}$$

The resultant $\Delta G_{(7)}$ is plotted as a function of temperature in Fig. 6.

The enthalpy and entropy changes associated with the reaction are given by the intercept and slope, respectively, of Fig. 6. This is the $2^{\rm nd}$ Law Method, which implicitly assumes that heat capacity changes over the temperature range are sufficiently small so as to be considered constant. Thus, the values of $\Delta H = -54552 \pm 12500$ J/mol (-32160 ± 12500 J/mol for $\varepsilon_{3550} = 5.1$ m²/mol) and $\Delta S = 115.2 \pm 6.9$ J/mol.K (101.4 ± 6.9 J/mol for $\varepsilon_{3550} = 5.1$ m²/mol) correspond to the mean temperature, 1800 K, over which the curve is fit. That the uncertainties on both quantities are relatively large reflects the three data points at 1 bar total pressure used to fit the relationship. However, using all data in Table 2, overlapping estimates are obtained, whereby $\Delta H = -61618 \pm 8044$ J/mol and $\Delta S = 116.1 \pm 5.6$ J/mol.K ($\Delta H = -53867 \pm 7540$ J/mol and $\Delta S = 110.3 \pm 5.3$ J/mol.K for $\varepsilon_{3550} = 5.1$ m²/mol).

When translated to H_2O dissolved in the silicate melt, a change in temperature from 1200 K to 2200 K results in a \sim 2.5-fold decrease in water content of the melt. Notably, the model of Moore et al. (1998) predicts a similar temperature dependence (a factor \sim 1.7 decrease over the same range), while the temperature term in the expression of lacono-Marziano et al. (2012) is so small so as to be insignificant. The amount of H_2O dissolved flattens out at high fH_2O , the implications of which are examined in the following sections.

Table 2Compilation of experiments performed assessing water solubility in silicate melts at low total pressures over a range of temperatures and compositions. NB. The value of $\alpha_{\rm H_{2}O}$ is calculated on the basis of mole fraction rather than ppmw.

Reference	Composition	Temperature	α	fH_2O	# of experiments
		(K)	(bar ^{-0.5})	(bar)	
This work ($\varepsilon_{3550} = 6.3 \text{ m}^2/\text{mol}$)	Peridotite	2173	2.91×10^{-3}	$5.7 \times 10^{-5} - 0.027$	14
This work ($\varepsilon_{3550} = 5.1 \text{ m}^2/\text{mol}$)	Peridotite	2173	3.59×10^{-3}	$5.7 \times 10^{-5} - 0.027$	14
Newcombe et al. (2017)	Anorthite-Diopside eutectic	1623	4.22×10^{-3}	$9.8 \times 10^{-3} - 0.32$	14
Newcombe et al. (2017)	Lunar Green Glass	1623	4.04×10^{-3}	$9.8 \times 10^{-3} - 0.32$	11
Dixon et al. (1995)	Mid-Ocean Ridge Basalt	1473	5.36×10^{-3}	17 - 709	14
Hamilton and Oxtoby (1986)	NaAlSi ₃ O ₈	1123 - 1573	$7.59 \times 10^{-3} - 9.91 \times 10^{-3}$	1685 - 2160	13

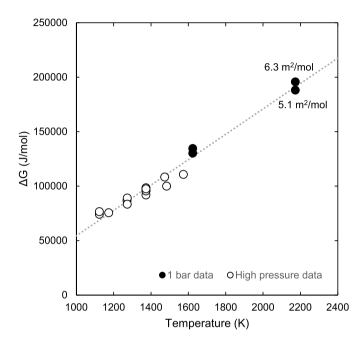


Fig. 6. The Gibbs Free Energy change of reaction (7) plotted as a function of temperature. Black = 1 bar data from this study and Newcombe et al. (2017); White = higher pressure data from Dixon et al. (1995) and Hamilton and Oxtoby (1986). These latter authors only provide total pressure, therefore, assuming a pure $\rm H_2O$ fluid in their experiments, total pressure was converted to $\rm fH_2O$ using an equation of state (Duan and Zhang, 2006). Grey dotted line = best fit to all data.

5. Implications

5.1. Effect of planetary mass on the extent of water retention

The formation of planetary bodies in the early Solar System proceeds by accretion of material through impacts, which become progressively more energetic as the masses of the bodies involved increases. Moreover, as these processes occurred within the first several million years following the collapse of the molecular cloud that led to the protoplanetary disk, short-lived radioactive isotopes, most notably, ²⁶Al with a half-life of 0.7 Myr, were still active (*e.g.*, Bhatia, 2021). These provide sources of heat that can lead to extensive melting of the precursor rocky bodies of the Earth and the other terrestrial planets (Chao et al., 2021).

Records of these bodies are preserved in achondritic meteorites, such as eucrites from the asteroid IV-Vesta. These samples, which are broadly basaltic in composition, contain nominally anhydrous minerals, namely clinopyroxenes, that preserve low H_2O contents (\sim 5 – 10 ppmw; Sarafian et al., 2017, 2019; Stephant et al., 2021) with respect to their terrestrial counterparts (50 – 700 ppmw; Warren and Hauri, 2014). The uniformly low H_2O contents (<20 ppmw) are a general feature of clinopyroxenes from achondritic mafic meteorites from small telluric bodies, including angrites and ureilites (Peterson et al., 2022; Sarafian et al., 2017).

This observation, together with their negative- to chondritic δD ratios (McCubbin and Barnes, 2019), suggests that i) H originates from the accretion of the parent body and ii) the H preserved is not residual from partial evaporative loss, which would produce positive δD with respect to chondrites. Here we examine these constraints in the context of magma ocean outgassing on planetary bodies.

The partial pressure, p, exerted by a given gas species in a single-species atmosphere (see Bower et al., 2019 for a multispecies atmosphere), i, at a planetary surface is given by:

$$p_i = \frac{gM_i^a}{4\pi r^2},\tag{17}$$

where M is the mass of i in the atmosphere, a, and r is the radius of the body. We model scenarios as a function of planetary radius in which the bulk water content is fixed (1000 ppmw), and can reside either in the magma ocean (assumed equivalent to the 70% the total mass of the body, i.e., comprised of a silicate mantle with a 30% core by mass) or in the atmosphere. This is expressed in the mass balance constraint:

$$M_i^T = M_i^a + M_i^m \tag{18a}$$

where T and m refer to the total mass and mantle mass of i, respectively. Substituting equation (17) into equation (18a), and given that $M_i^T = M_T C_{i,T}$, where $C_{i,T}$ is the concentration (by mass) of i, one obtains:

$$M_T C_{i,T} = p_i \frac{4\pi r^2}{g} + M_m C_{i,m}$$
 (18b)

The bulk mass of the body, M_T , is calculated by solving the static structure equations that also yield gravity and internal pressure as a function of radius, r (Valencia et al., 2007; Appendix A). The density of the mantle with depth is calculated assuming compression under adiabatic conditions, according to the Adams-Williamson equation, which accounts for the compressibility of typical mantle peridotite (Jackson, 1998). Finally, the quantity $C_{i,m}$ is eliminated by substitution of eq. (3), connecting the mantle abundance of H_2O to its partial pressure (note ideality in the gas phase is assumed, and p is interchangeable with f, the fugacity):

$$M_T C_{H_2O,T} = f H_2 O \frac{4\pi r^2}{g} + M_m \alpha_{H_2O} (f H_2O)^{0.5}$$
 (19)

Equation (19) is solved numerically, by varying f H₂O such that the right-hand side equals the left-hand side.

Solutions to eq. (19) are represented graphically in Fig. 7. For a constant H_2O content of 1000 ppmw, partial pressures increase markedly above bodies with surface gravity up to $\sim 2 \text{ m/s}^2$, corresponding to radii of $\sim 2000 \text{ km}$. Over the same interval, the fraction of water outgassed decreases drastically, until the partial

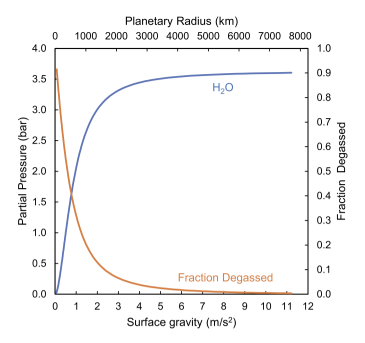


Fig. 7. The partial pressure of H_2O (blue line) in a single-species atmosphere and the corresponding fraction of total water in the atmosphere (orange line) above a planetary surface as a function of its surface gravity and radius. While the absolute partial pressure increases with increasing planetary mass, the relative fraction of water present in the atmosphere (as opposed to dissolved in the liquid mantle) decreases. The calculations assume a fixed total water concentration (1000 ppmw as H_2O) for all planetary masses, a bulk density calculated by static structure equations assuming adiabatic compression (Appendix A), an α_{H_2O} of 524 ppmw/bar^{0.5} (use of $\alpha_{H_2O} = 647$ ppmw/bar^{0.5} would result in even lower pressures), a temperature of 2173 K and that the mass of the mantle is equal to 0.7 times the mass of the planet (i.e., an Earth-like core mass fraction). (For interpretation of the colours in the figure(s), the reader is referred to the web version of this article.)

pressure reaches an asymptote at roughly 3.5 bar for more massive planets.

As a result, the H_2O budget can be lost more readily from smaller planetary bodies with respect to larger ones, despite lower partial pressures in their atmospheres, because a greater fraction of the water resides in the atmosphere. For example, magma oceans on bodies with the surface gravity of Vesta (0.25 m/s^2) would be expected to have $\sim \! 50\%$ of their total H_2O present in the atmosphere, compared with 0.5% for a body with the mass of Earth (Fig. 7). All else being equal, smaller bodies will tend to outgas atmospheres whose composition closely reflects that of their bulk planetary inventory, while species in the atmospheres of larger bodies will be moderated by their solubilities.

The foregoing analysis presumes all H is present as H₂O, and that the fO_2 is independent of planetary size. It is likely, however, that a significant fraction (~50%) of H existed as H₂ in the atmospheres of small telluric bodies, given that the fO_2 recorded in basaltic achondrites is near the IW buffer (Wadhwa, 2008). Accordingly, because H₂ is less soluble than is H₂O in silicate liquid (Hirschmann et al., 2012), the lower oxygen fugacities of small rocky bodies relative to Earth's present-day upper mantle (Δ IW+3.5) make H loss from the former more pronounced. Therefore, the escape of H₂O from small planetary bodies is not only favoured by their lower escape velocities, but also by their lower fO₂ and the solubility behaviour of H₂O that results in proportionally greater degassing extents. These controls provide a physical explanation for the observation that all achondrites from small telluric bodies (Sarafian et al., 2019; Peterson et al., 2022) are strongly depleted in water with respect to the Earth in a manner inversely proportional to their radii (McCubbin and Barnes, 2019).

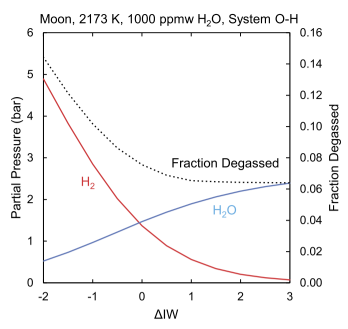


Fig. 8. Partial pressures of H-bearing species in the system O-H in an hypothetical primitive lunar atmosphere at 2173 K as a function of oxygen fugacity expressed relative to the iron-wüstite buffer (Δ IW). Red = H₂, Blue = H₂O. The initial concentration of H₂O in the Moon is assumed to be that of Earth's mantle (1000 ppmw = 7.33 \times 10¹⁹ kg), its radius 1737 km and surface gravity = 1.62 m/s². An $\alpha_{\rm H_2O}$ of 524 ppmw/bar^{0.5} is used (use of $\alpha_{\rm H_2O}$ = 647 ppmw/bar^{0.5} would result in even lower pressures).

5.2. Water in the primitive atmosphere of the Moon

While similarities in the chemical and isotopic composition of refractory and major rock-forming elements in the Earth and Moon, such as Cr, Ti and O, have been used to argue for formation of the Moon directly from, or by complete mixing with Earth-like material (Lock et al., 2020; Pahlevan and Stevenson, 2007), this concordance breaks down for elements more volatile than Li, which are systematically depleted in the Moon (Tartèse et al., 2021). This depletion extends to hydrogen, whose abundance in the lunar mantle has been estimated through measurement of pyroclastic glasses (Saal et al., 2008) as well as from melt inclusions in mare basalts (Chen et al., 2015; Ni et al., 2019), to fall in the range 3 – 84 ppmw $\rm H_2O$ (McCubbin and Barnes, 2019 and references therein). Though the range is wide, there is little doubt that the Moon is depleted in H with respect to Earth's mantle by a factor of 10 - 1000.

Nakajima and Stevenson (2018) argued that, in an H_2O -dominated protolunar atmosphere, H loss would be limited by its sluggish diffusion through the atmosphere with a mean molecular mass greater than that of H_2 . However, these authors considered only H_2O -rich atmospheres ($fH_2O/fH_2>100$) which implicitly assumes oxidising conditions (at 2000 K, the fH_2O/fH_2 ratio reaches $\sim\!100$ at the Fayalite-Magnetite-Quartz, FMQ buffer). Depending on the temperature and fO_2 at which the atmosphere equilibrated with the magma, fH_2/fH_2O could vary by orders of magnitude, potentially aiding the escape of H and other gases in the atmosphere.

Beginning with the premise that the Moon was derived from the Earth's mantle (Ringwood and Kesson, 1977), the initial concentration of H_2O in the lunar mantle is expected to have been that of the present-day Earth's mantle, that is, roughly 1000 ppmw (Marty, 2012; Hirschmann, 2018). The formation of a primitive lunar atmosphere in equilibrium with a magma ocean at 2173 K in the simple system O-H as a function of fO_2 (Fig. 8) is simulated using eq. (11) for H_2O and the data of Hirschmann et al. (2012)

for the dissolution of H_2 . For this purpose, the homogeneous gas phase equilibrium $H_2 + 1/2O_2 = H_2O$ is described by $\log K = 13152/T - 3.039$ (Chase, 1998) and is used to provide self-consistent solutions for fH_2 and fH_2O . An equation analogous to eq. (19) is also solved for H_2 .

The results indicate that i) the total pressure of the sum of H-bearing species never exceeds \sim 6 bar, ii) the dominant H-bearing species is H₂ below IW, and iii) the fraction of H degassed increases from 0.06 at Δ IW+3 to 0.15 at Δ IW-2, owing to the lower solubility of molecular hydrogen with respect to water.

As the partial pressures of gas species derived from silicates (e.g., Mg(g), SiO(g)) are relatively low even at high temperatures (10^{-3}) bar at 2000 K, Charnoz et al., 2021), it is likely that the Moon never had a thick atmosphere (fewer than tens of bars; see also Needham and Kring, 2017 that estimate mare volcanism produces only 0.01 bar). Moreover, loss of H above a protolunar magma ocean would have been favoured under reducing conditions as not only does increasing fH_2/fH_2O lower the mean molecular mass of the atmosphere from 17.5 at Δ IW+3 to 3.5 at Δ IW-2, but a factor \sim 2 higher fraction of the budget of H resides in the atmosphere. Stated otherwise, should the Moon have had an initial H2O budget similar to that of Earth, and an oxidised magma ocean, then its water depletion could likely only have arisen owing to H loss in the protolunar disk, prior to its accretion. Hence, consideration of the fO_2 of the protolunar disk and above its magma ocean are paramount to assessing whether the scenario envisaged by Nakajima and Stevenson (2018) is applicable (above $\sim \Delta IW+0.5$), and therefore the ease with which the Moon dehydrated during its formation.

5.3. Prevalence of steam atmospheres around rocky bodies

A steam atmosphere is defined as one dominated by water vapour with respect to other gaseous species that may occur in planetary atmospheres at high temperatures (CO, CO₂, H₂). Such atmospheres are thought to have been relevant to the formation of the Earth in a magma ocean state (e.g., Abe and Matsui, 1985; Fegley et al., 2016; Lebrun et al., 2013), to the terrestrial planets more generally (Lammer et al., 2020), as well as for exoplanets (Turbet et al., 2020). Indeed, the dissolution of water into silicate has been proposed to account, in part, for the 'radius gap' observed between super-Earths and sub-Neptunes (Zeng et al., 2021). However, whether steam atmospheres form is contingent upon the abundance and solubility of H with respect to other atmophile elements, of which the most abundant is C.

As demonstrated above, the fraction of H outgassed is not only dependent on intensive variables such as temperature and oxygen fugacity, but also extensive variables, namely, its abundance and planetary mass. To assess the prevalence and occurrence of steam atmospheres above magma oceans on Earth-like planets, a Monte Carlo simulation with 10,000 runs is performed across three variables that are uniformly distributed, the molar C/H ratio of bulk planet $\{C/H \in \mathbb{R} \mid 0.1 < C/H < 1\}$, the abundance of H in ocean masses, $N_{Ocean} \{ N_{Ocean} \in \mathbb{R} \mid 1 < N_{Ocean} < 10 \}$ and the fO_2 at the magma ocean atmosphere surface $\{\Delta IW \in \mathbb{R} \mid -4 < \Delta IW < 4\}$. The partial pressures of the species CO, CO₂, H₂ and H₂O are calculated in these model atmospheres at a constant temperature, 2173 K and planetary mass, that of the Earth (1 M_E, where 0.68 of this is its mantle) in a self-consistent manner according to mass balance constraints (eq. (19)) and homogeneous gas phase equilibria (see Bower et al., 2022, for a complete description).

At 2173 K, only the four major gas species in the system C-O-H shown in Fig. 9 are stable (cf. Sossi et al., 2020; Bower et al., 2022). A striking feature of the generated compositions is the mutual exclusivity of CO_2 -H₂-rich atmospheres (Fig. 9). Within the explored parameter space, a maximum of roughly 75 bar H₂ and

75 bar CO₂ can coexist at intermediate fO₂ near the IW buffer (10 Ocean masses, C/H = 1), at which the major gas species is CO(\sim 200 to 400 bar). On average, CO is the most abundant species expected in the range of modelled atmospheres, despite the fact that bulk C/H ratios are <1. This is a product of its low solubility with respect to H₂O and CO₂, and the fact that, at 2173 K, chemical equilibrium dictates that $CO/CO_2 = 1$ at IW+1.5 (with CO predominant below this value) as compared to IW for $H_2/H_2O =$ 1. As a result, there is a wide parameter space of CO-H₂-rich atmospheres that occurs for all C/H ratios and ocean masses below the IW buffer. Considering that the formation of iron-rich cores on rocky planets in our Solar System sets the fO_2 to between Δ IW-1 and Δ IW-4 (e.g., Frost et al., 2008), this condition may be commonplace among rocky planets. Nevertheless, the fO_2 set by core formation may not reflect that at the surface of the magma ocean, owing to the volume changes of redox equilibria, notably of iron (Hirschmann, 2012), such that higher fO_2 conditions at the interface are possible (Deng et al., 2020; Sossi et al., 2020).

Steam atmospheres sensu stricto, in which $fH_2O/(fH_2O +$ $fH_2 + fCO + fCO_2$) > 0.5, do not form; the maximum value of the index achieved in the simulations is 0.34, and only 0.72% have values >0.25. Water, as the most soluble species amongst the four considered, reaches a maximum fugacity of only 37 bar (as compared with \sim 700 bar for both CO₂ and H₂, and \sim 500 bar for CO), meaning steam atmospheres are, on average, far more tenuous than the equivalent carbon-rich atmospheres produced at a given C/H ratio. Part of this behaviour derives from the dependence of dissolved H₂O on $(fH_2O)^{0.5}$, one that holds until ~1000 bar fH_2O (e.g., Berndt et al., 2002; Dixon et al., 1995). At pressures higher than this, significant quantities of H₂O can more readily enter the atmosphere, as $K_{(6)}$ shifts to the left and XH_2O becomes proportional to fH_2O . Care should therefore be taken in applying eq. (11) to magma oceans of planets with large g and/or water budgets ($fH_2O > 1000$ bar), in which water dissolution would tend to a value of $\beta = 1$. Importantly, steam atmospheres for the scenarios considered are never composed entirely of steam; they contain significant quantities of H2, CO2 and CO. Indeed, the formation of steam atmospheres requires a bulk planet with >10 ocean masses, C/H ratios <0.1 and oxygen fugacities at the atmosphere-magma interface just above the IW buffer. This is because, although fH_2O scales with $(fO_2)^{0.5}$, so too does the fugacity of the less soluble CO_2 , meaning fCO_2 quickly surpasses fH_2O above IW.

As a result, steam atmospheres are likely to be uncommon on Earth-sized rocky planets, lest they have molar C/H ratios significantly lower than that of Earth (\sim 1). In the solar neighbourhood, the C/H ratios of most G-type stars vary by ± 0.5 dex units about the solar ratio (Hinkel et al., 2014). If this variability is indicative of that found in any potential orbiting rocky planets, then only a minority of stars, about 53 of 3819 G-type stars with a C/H ratio \sim 10 times lower than that of the Sun, would be expected to host rocky planets that harbour steam atmospheres.

The foregoing assessment neglects other atmophile elements (notably S and N), which would further dilute the H_2O fraction in the atmosphere, particularly at high fO_2 (> Δ IW+2) at which SO_2 becomes a significant component (Gaillard et al., 2022). Moreover, only high temperature atmospheres (2173 K) in equilibrium with an Earth's mantle-sized magma ocean are considered. However, a variety of post-magma ocean processes have the potential to modify the constitution of the magma-generated atmosphere (Bower et al., 2022; Gaillard et al., 2022; Hier-Majumder and Hirschmann, 2017; Sossi et al., 2020). Notably, homogeneous gas phase equilibria, precipitation of graphite and other condensed phases, and their subsequent separation from the atmosphere during cooling can promote the production of H_2O by increasing fO_2 and decreasing the C/H ratio, or its consumption through condensation (Sossi et al., 2020). Even prior to its outgassing, dynamical pro-

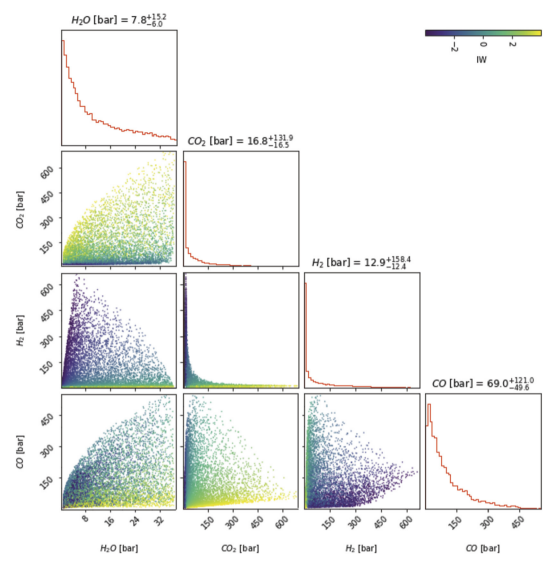


Fig. 9. Corner plot of the partial pressures of the four gas species present in the atmospheres of planets with 1 Earth mass (M_E) and a fully-molten mantle of 0.68 M_E at constant temperature (2173 K), generated according to a Monte-Carlo simulation by variation of three parameters – C/H ratio, ocean masses of H, and oxygen fugacity (see text for details). The colours of the points are coded according to the fO_2 value, reported with respect to the iron-wüstite buffer (Δ IW). Equivalent plots in which the points are colour-coded by C/H ratio and ocean masses are shown in the Supplementary Information. Histograms are constructed with 50 bins, and the title shows the 0.5 quantile of the data, with lower and upper bounds that pertain to the 0.16 and 0.84 quantiles, respectively. An α_{H_2O} of 524 ppmw/bar^{0.5} is used (use of $\alpha_{H_2O} = 647$ ppmw/bar^{0.5} would result in even lower pressures of H_2O).

cesses in the magma ocean, such as melt trapping (Hier-Majumder and Hirschmann, 2017) and dynamical lock-up (Bower et al., 2022) can prevent wholesale outgassing of H_2O , while fractional crystallisation prolongs its outgassing by maintaining contact between the melt and the atmosphere (Bower et al., 2022). Therefore, whether steam atmospheres around planets eventually arise during cooling and crystallisation depends upon i) efficient liberation of dissolved water and/or ii) chemical equilibration of the atmosphere and its components upon cooling.

6. Conclusions

The generation and prevalence of steam atmospheres around rocky planets is an oft-invoked, yet poorly understood phenomenon. This is largely owing to the scarcity of solubility data for compositions, and at pressures and temperatures, relevant to the formation of atmospheres from molten planetary mantles. We demonstrate, through experimental determination of H_2O contents in peridotite liquids at 2173 K using Fourier-Transform InfraRed spectroscopy, that the dissolved content of H_2O is proportional to

the square-root of the fugacity of H_2O with which it is in equilibrium. This dependence confirms its incorporation as hydroxyl groups (OH) in silicate melts at the low total fH_2O (up to 0.027 bar) of the experiments. Moreover, fH_2 promotes dissolution of OH at low fO_2 below the iron-wüstite buffer. At a given fH_2O , temperature exerts the primary control on water solubility in silicate liquids, and the water contents of peridotite melts at 2173 K are 10 - 25% lower than for basaltic melts at 1623 K at equivalent fH_2O , offsetting the effect of lower melt polymerisation, which is expected to increase H_2O solubility. Peridotite liquids are able to dissolve $H_2(g)$ as OH in their structure, with a coefficient 1/3 that for dissolution of $H_2O(g)$, a property linked to their highly depolymerised nature.

We demonstrate that a greater fraction of the total water budget of a given planetary body is outgassed for smaller with respect to larger bodies, owing to the square-root dependence of fH_2O on dissolved H_2O . This property, together with the reducing fO_2 of basaltic achondrites compared to terrestrial basalts, provides an explanation for the systematically low H_2O contents of nominally anhydrous minerals in achondritic meteorites derived from the crusts

and mantles of small telluric bodies with respect to their terrestrial counterparts. Assuming an initially Earth-like H budget, one such small body, the Moon, likely only ever harboured a tenuous atmosphere, with less than several bars of H as H_2 (\sim 6 bar, below IW) or H_2O (\sim 2 bar, above IW). The fO_2 that prevailed in the protolunar atmosphere affects its mean molar mass, and is therefore key to determining whether H is retained as H₂O or more readily escapes as H₂. A survey of the possible chemical variability among rocky (exo)planets shows that the fO_2 at the atmosphere-magma interface, together with the C/H ratio and number of oceans in the bulk planet, control the likelihood of steam atmosphere formation at magmatic temperatures. We show, through Monte-Carlo simulation, that the high solubility of H₂O with respect to other gases in the C-O-H system limits the extent of its atmospheric outgassing. Therefore, steam atmospheres are rare outcomes of magma oceanderived secondary atmospheres, and require low C/H ratios, and high H budgets and oxygen fugacities to prevail.

CRediT authorship contribution statement

Paolo A. Sossi: Conceptualization, Formal analysis, Investigation, Writing – original draft, Writing – review & editing. **Peter M.E. Tollan:** Formal analysis, Investigation, Writing – review & editing. **James Badro:** Investigation, Resources, Writing – review & editing. **Dan J. Bower:** Formal analysis, Writing – review & editing.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Raw data is available as csv files in the supplementary information.

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Appendix A

We solve the interior static structure equations (e.g., Valencia et al., 2007) to determine the structure and bulk properties of differentiated planetesimals and planets with radii ranging from around 10 km to 7700 km. To this end, we first prescribe the bulk planetary mass and a fixed core mass fraction of 30% across all radii, where the density of the iron core is assumed to be constant (pressure-independent) at 10000 kg/m^3 . This approach enables determination of the radius of the core-mantle boundary and the mass of the mantle. The remaining step is to determine the radius of the planet, r_0 , given the mass of the mantle and core, the

radius of the core-mantle boundary, and an equation of state for the silicate mantle. To capture the density change $(d\rho)$ of peridotite with radius (dr) we assume adiabatic self-compression (the Adams-Williamson equation),

$$\frac{d\rho}{dr} = -\frac{\rho (r)^2 g(r)}{K},\tag{A.1}$$

where ρ is the density, g is the acceleration due to gravity, and K is the adiabatic bulk modulus of the mantle. Here we employ a value of K=264 GPa determined for bridgmanite (Jackson, 1998) to approximate that of peridotite. The remaining static structure equations are:

$$\frac{dg}{dr} = 4\pi G\rho(r) - \frac{2Gm(r)}{r^3},\tag{A.2}$$

$$\frac{dm}{dr} = 4\pi r^2 \rho (r), \qquad (A.3)$$

$$\frac{dP}{dr} = -\rho(r)g(r). \tag{A.4}$$

In the above equations, G is the universal gravitational constant, m is the mass contained within r and P is the pressure. To solve the system of equations we prescribe boundary conditions at the planetary radius (r_0) and integrate to the core-mantle boundary:

$$g(r_0) = Gm_0/r_0^2$$
, $m(r_0) = m_0$, $P(r_0) = 0$, $\rho(r_0) = \rho_0$ (A.5)

We adopt a surface density ρ_0 of 3300 kg/m³ for peridotite (Lee, 2003). Using Newton's method we then solve the structure equations for the planetary radius (r_0) that results in a silicate mantle that constitutes 70% of the prescribed bulk planetary mass. For the size range of planetary bodies we consider, this provides a self-consistent mantle & bulk planetary mass, planetary radius, and surface gravity to use in eqns. (17) and (18).

Appendix B. Supplementary material

Supplementary material related to this article can be found online at https://doi.org/10.1016/j.epsl.2022.117894.

References

Abe, Y., Matsui, T., 1985. The formation of an impact-generated $\rm H_2O$ atmosphere and its implications for the early thermal history of the Earth. J. Geophys. Res. 90, C545. https://doi.org/10.1029/jb090is02p0c545.

Badro, J., Sossi, P.A., Deng, Z., Borensztajn, S., Wehr, N., Ryerson, F.J., 2021. Experimental investigation of elemental and isotopic evaporation processes by laser heating in an aerodynamic levitation furnace. C. R. Géosci. 353 (1), 101–114.

Berndt, J., Liebske, C., Holtz, F., Friese, M., Nowak, M., Ziegenbein, D., Hurkuck, W., Koepke, J., 2002. A combined rapid-quench and H₂-membrane setup for internally heated pressure vessels: description and application for water solubility in basaltic melts. Am. Mineral. 87, 1717–1726.

Bhatia, G.K., 2021. Early thermal evolution of the embryos of Earth: role of ²⁶Al and impact-generated steam atmosphere. Planet. Space Sci. 207, 105335. https://doi.org/10.1016/j.pss.2021.105335.

Bondar, D., 2022. Water Partition Coefficients Between Upper Mantle Minerals and Melts as a Function of P/T and CO₂ Content Using a Rapid Quench Cell in Multi-Anvil Experiments. Bayerisches GeoInstitut.

Bondar, D., Fei, H., Withers, A.C., Ishii, T., Chanyshev, A., Katsura, T., 2021. A simplified rapid-quench multi-anvil technique. Rev. Sci. Instrum. 92. https://doi.org/10.1063/5.0062525

Bondar, D., Zandonà, A., Withers, A.C., Fei, H., Genova, D. Di, Miyajima, N., Katsura, T., 2022. Rapid-quenching of high-pressure depolymerized hydrous silicate (peridotitic) glasses. J. Non-Cryst. Solids 578. https://doi.org/10.1016/j.jnoncrysol. 2021.121347.

Bower, D.J., Hakim, K., Sossi, P.A., Sanan, P., 2022. Retention of water in terrestrial magma oceans and carbon-rich early atmospheres. Planet. Sci. J. 3, 93.

Bower, D.J., Kitzmann, D., Wolf, A.S., Sanan, P., Dorn, C., Oza, A.V., 2019. Linking the evolution of terrestrial interiors and an early outgassed atmosphere to astrophysical observations. Astron. Astrophys. 631, A103.

- Bureau, H., Raepsaet, C., Khodja, H., Carraro, A., Aubaud, C., 2009. Determination of hydrogen content in geological samples using elastic recoil detection analysis (ERDA). Geochim. Cosmochim. Acta 73, 3311–3322. https://doi.org/10.1016/j.gca. 2009.03.009.
- Catling, D.C., Kasting, J.F., 2017. Atmospheric Evolution on Inhabited and Lifeless Worlds. Cambridge, UK.
- Chao, K.-H., DeGraffenried, R., Lach, M., Nelson, W., Truax, K., Gaidos, E., 2021. Lava worlds: from early Earth to exoplanets. Geochemistry 81, 125735.
- Charnoz, S., Sossi, P.A., Lee, Y.N., Siebert, J., Hyodo, R., Allibert, L., Pignatale, F.C., Landeau, M., Oza, A.V., Moynier, F., 2021. Tidal pull of the Earth strips the proto-Moon of its volatiles. Icarus 364, 114451. https://doi.org/10.1016/j.icarus.2021. 114451
- Chase, M.W., 1998. NIST-JANAF Thermochemical Tables, 4th edition. American Chemical Society, Washington, D.C.
- Chen, Y., Zhang, Y., Liu, Y., Guan, Y., Eiler, J., Stolper, E.M., 2015. Water, fluorine, and sulfur concentrations in the lunar mantle. Earth Planet. Sci. Lett. 427, 37–46. https://doi.org/10.1016/j.epsl.2015.06.046.
- Dalou, C., Hirschmann, M.M., Jacobsen, S.D., Le Losq, C., 2019. Raman spectroscopy study of C-O-H-N speciation in reduced basaltic glasses: implications for reduced planetary mantles. Geochim. Cosmochim. Acta 265, 32–47. https://doi. org/10.1016/j.gca.2019.08.029.
- Deng, J., Du, Z., Karki, B.B., Ghosh, D.B., Lee, K.K.M., 2020. A magma ocean origin to divergent redox evolutions of rocky planetary bodies and early atmospheres. Nat. Commun. 11, 1–7. https://doi.org/10.1038/s41467-020-15757-0.
- Dingwell, D.B., Courtial, P., Giordano, D., Nichols, A.R.L., 2004. Viscosity of peridotite liquid. Earth Planet. Sci. Lett. 226, 127–138. https://doi.org/10.1016/j.epsl.2004. 07.017
- Dixon, J.E., Stolper, E.M., Holloway, J.R., 1995. An experimental study of water and carbon dioxide solubilities in mid-ocean ridge basaltic liquids. Part I: calibrations and solubility models. J. Petrol. 36, 1607–1631. https://doi.org/10.1093/ oxfordjournals.petrology.a037268.
- Duan, Z., Zhang, Z., 2006. Equation of state of the H₂O, CO₂, and H₂O-CO₂ systems up to 10 GPa and 2573.15 K: molecular dynamics simulations with ab initio potential surface. Geochim. Cosmochim. Acta 70, 2311–2324. https://doi.org/10.1016/i.gca.2006.02.009.
- Elkins-Tanton, L.T., 2008. Linked magma ocean solidification and atmospheric growth for Earth and Mars. Earth Planet. Sci. Lett. 271, 181–191. https://doi.org/10.1016/j.epsl.2008.03.062.
- Fegley, B., Jacobson, N.S., Williams, K.B., Plane, J.M.C., Schaefer, L., Lodders, K., 2016. Solubility of rock in steam atmospheres of planets. Astrophys. J. 824, 103.
- Frost, D.J., Mann, U., Asahara, Y., Rubie, D.C., 2008. The redox state of the mantle during and just after core formation. Philos. Trans. A. Math. Phys. Eng. Sci. 366, 4315–4337. https://doi.org/10.1098/rsta.2008.0147.
- Gaillard, F., Bernadou, F., Roskosz, M., Bouhifd, M.A., Marrocchi, Y., Iacono-Marziano, G., Moreira, M., Scaillet, B., Rogerie, G., 2022. Redox controls during magma ocean degassing. Earth Planet. Sci. Lett. 577, 117255. https://doi.org/10.1016/j.epsl.2021.117255.
- Gaillard, F., Scaillet, B., Pichavant, M., Beny, J., 2001. The effect of water and fO_2 on the ferric ferrous ratio of silicic melts. Chem. Geol. 174, 255–273.
- Genda, H., Abe, Y., 2005. Enhanced atmospheric loss on protoplanets at the giant impact phase in the presence of oceans. Nature 433, 842–844.
- Green, D.H., 1975. Genesis of Archean peridotitic magmas and constraints on Archean geothermal gradients and tectonics. Geology 3, 15–18. https://doi.org/10.1130/0091-7613(1975)3<15.
- Hamilton, D.L., Oxtoby, S., 1986. Solubility of water in albite-melt determined by the weight-loss method. J. Geol. 94, 626–630.
- Hier-Majumder, S., Hirschmann, M.M., 2017. The origin of volatiles in the Earth's mantle. Geochem. Geophys. Geosyst. 18, 3078–3092. https://doi.org/10.1002/2017GC006937.
- Hinkel, N.R., Timmes, F.X., Young, P.A., Pagano, M.D., Turnbull, M.C., 2014. Stellar abundances in the solar neighborhood: the hypatia catalog. Astron. J. 148. https://doi.org/10.1088/0004-6256/148/3/54.
- Hirschmann, M.M., 2021. Iron-wüstite revisited: a revised calibration accounting for variable stoichiometry and the effects of pressure. Geochim. Cosmochim. Acta 313, 74–84. https://doi.org/10.1016/j.gca.2021.08.039.
- Hirschmann, M.M., 2018. Comparative deep Earth volatile cycles: the case for C recycling from exosphere/mantle fractionation of major (H₂O, C, N) volatiles and from H₂O/Ce, CO₂/Ba, and CO₂/Nb exosphere ratios. Earth Planet. Sci. Lett. 502, 262–273. https://doi.org/10.1016/j.epsl.2018.08.023.
- Hirschmann, M.M., 2012. Magma ocean influence on early atmosphere mass and composition. Earth Planet. Sci. Lett. 341–344, 48–57.
- Hirschmann, M.M., Withers, A.C., Ardia, P., Foley, N.T., 2012. Solubility of molecular hydrogen in silicate melts and consequences for volatile evolution of terrestrial planets. Earth Planet. Sci. Lett. 345–348, 38–48. https://doi.org/10.1016/j.epsl. 2012.06.031
- Iacono-Marziano, G., Morizet, Y., Le Trong, E., Gaillard, F., 2012. New experimental data and semi-empirical parameterization of H₂O-CO₂ solubility in mafic melts. Geochim. Cosmochim. Acta 97, 1–23. https://doi.org/10.1016/j.gca.2012.08.035.
- Jackson, I., 1998. Elasticity, composition and temperature of the Earth's lower mantle: a reappraisal. Geophys. J. Int. 134, 291–311. https://doi.org/10.1046/j.1365-246X.1998.00560.x.

- Kress, V.C., Carmichael, I.S.E., 1991. The compressibility of silicate liquids containing Fe₂O₃ and the effect of composition, temperature, oxygen fugacity and pressure on their redox states. Contrib. Mineral. Petrol. 108, 82–92. https://doi.org/10. 1007/BF00307328.
- Lammer, H., Scherf, M., Kurokawa, H., Ueno, Y., Burger, C., Maindl, T., Johnstone, C.P., Leizinger, M., Benedikt, M., Fossati, L., Kislyakova, K.G., Marty, B., Avice, G., Fegley, B., Odert, P., 2020. Loss and fractionation of noble gas isotopes and moderately volatile elements from planetary embryos and early Venus, Earth and Mars. Space Sci. Rev. 216. https://doi.org/10.1007/s11214-020-00701-x.
- Lange, R.A., Carmichael, I.S.E., 1987. Densities of Na₂O-K₂O-CaO-MgO-FeO-Fe₂O₃-Al₂O₃-TiO₂-SiO₂ liquids: new measurements and derived partial molar properties. Geochim. Cosmochim. Acta 51, 2931–2946. https://doi.org/10.1016/0016-7037(87)90368-1
- Lebrun, T., Massol, H., Chassefière, E., Davaille, A., Marcq, E., Sarda, P., Leblanc, F., Brandeis, G., 2013. Thermal evolution of an early magma ocean in interaction with the atmosphere. J. Geophys. Res. E Planets 118, 1155–1176. https://doi.org/10.1002/jgre.20068.
- Lee, C.T.A., 2003. Compositional variation of density and seismic velocities in natural peridotites at STP conditions: implications for seismic imaging of compositional heterogeneities in the upper mantle. J. Geophys. Res., Solid Earth 108 (B9). https://doi.org/10.1029/2003JB002413.
- Lock, S.J., Bermingham, K.R., Parai, R., Boyet, M., 2020. Geochemical constraints on the origin of the moon and preservation of ancient terrestrial heterogeneities. Space Sci. Rev. 216. https://doi.org/10.1007/s11214-020-00729-z.
- Marty, B., 2012. The origins and concentrations of water, carbon, nitrogen and noble gases on Earth. Earth Planet. Sci. Lett. 313–314, 56–66. https://doi.org/10.1016/j.epsl.2011.10.040.
- McCubbin, F.M., Barnes, J.J., 2019. Origin and abundances of H₂O in the terrestrial planets, Moon, and asteroids. Earth Planet. Sci. Lett. 526, 115771. https://doi.org/10.1016/j.epsl.2019.115771.
- Melosh, H.J., 1990. Giant impacts and the thermal state of the early Earth. In: Newsom, H.E., Jones, J.H. (Eds.), Origin of the Earth. Oxford University, New York, pp. 69–83.
- Mercier, M., Muro, A. Di, Métrich, N., Giordano, D., Belhadj, O., Mandeville, C.W., 2010. Spectroscopic analysis (FTIR, Raman) of water in mafic and intermediate glasses and glass inclusions. Geochim. Cosmochim. Acta 74, 5641–5656. https:// doi.org/10.1016/j.gca.2010.06.020.
- Moore, G., Vennemann, T., Carmichael, I.S.E., 1998. An empirical model for the solubility of H₂O in magmas to 3 kilobars. Am. Mineral. 83, 36–42.
- Nakajima, M., Stevenson, D.J., 2018. Inefficient volatile loss from the Moon-forming disk: reconciling the giant impact hypothesis and a wet Moon. Earth Planet. Sci. Lett. 487, 117–126. https://doi.org/10.1016/j.epsl.2018.01.026.
- Needham, D.H., Kring, D.A., 2017. Lunar volcanism produced a transient atmosphere around the ancient Moon. Earth Planet. Sci. Lett. 478, 175–178. https://doi.org/10.1016/j.epsl.2017.09.002.
- Newcombe, M.E., Beckett, J.R., Baker, M.B., Newman, S., Guan, Y., Eiler, J.M., Stolper, E.M., 2019. Effects of pH₂O, pH₂ and fO₂ on the diffusion of H-bearing species in lunar basaltic liquid and an iron-free basaltic analog at 1 atm. Geochim. Cosmochim. Acta 259, 316–343. https://doi.org/10.1016/j.gca.2019.05.033.
- Newcombe, M.E., Brett, A., Beckett, J.R., Baker, M.B., Newman, S., Guan, Y., Eiler, J.M., Stolper, E.M., 2017. Solubility of water in lunar basalt at low pH₂O. Geochim. Cosmochim. Acta 200, 330–352. https://doi.org/10.1016/j.gca.2016.12.026.
- Ni, P., Zhang, Y., Chen, S., Gagnon, J., 2019. A melt inclusion study on volatile abundances in the lunar mantle. Geochim. Cosmochim. Acta 249, 17–41. https://doi.org/10.1016/j.gca.2018.12.034.
- Pahlevan, K., Stevenson, D.J., 2007. Equilibration in the aftermath of the lunarforming giant impact. Earth Planet. Sci. Lett. 262, 438–449. https://doi.org/10. 1016/j.epsl.2007.07.055.
- Peterson, L., Newcombe, M.E., Alexander, C., Wang, J., Sarafian, A., Bischof, A., Nielsen, S., 2022. The water content of the ureilite parent body. Geochim. Cosmochim. Acta. https://doi.org/10.1016/j.gca.2022.10.036.
- Ringwood, A.E., Kesson, S.E., 1977. Basaltic magmatism and the bulk composition of the moon. Moon 16, 425–464. https://doi.org/10.1007/BF00577902.
- Saal, A.E., Hauri, E.H., Cascio, M. Lo, Van Orman, J.A., Rutherford, M.C., Cooper, R.F., 2008. Volatile content of lunar volcanic glasses and the presence of water in the Moon's interior. Nature 454, 192–195. https://doi.org/10.1038/nature07047.
- Sakamaki, T., Ohtani, E., Urakawa, S., Suzuki, A., Katayama, Y., 2010. Density of dry peridotite magma at high pressure using an X-ray absorption method. Am. Mineral. 95, 144–147. https://doi.org/10.2138/am.2010.3143.
- Sarafian, A.R., John, T., Roszjar, J., Whitehouse, M.J., 2017. Chlorine and hydrogen degassing in Vesta's magma ocean. Earth Planet. Sci. Lett. 459, 311–319. https:// doi.org/10.1016/j.epsl.2016.10.029.
- Sarafian, A.R., Nielsen, S.G., Marschall, H.R., Gaetani, G.A., Righter, K., Berger, E.L., 2019. The water and fluorine content of 4 Vesta. Geochim. Cosmochim. Acta 266, 568–581. https://doi.org/10.1016/j.gca.2019.08.023.
- Shishkina, T.A., Botcharnikov, R.E., Holtz, F., Almeev, R.R., Jazwa, A.M., Jakubiak, A.A., 2014. Compositional and pressure effects on the solubility of H₂O and CO₂ in mafic melts. Chem. Geol. 388, 112–129. https://doi.org/10.1016/j.chemgeo.2014. 09.001.

- Shishkina, T.A., Botcharnikov, R.E., Holtz, F., Almeev, R.R., Portnyagin, M.V., 2010. Solubility of H₂O- and CO₂-bearing fluids in tholeitic basalts at pressures up to 500 MPa. Chem. Geol. 277, 115–125.
- Sieverts, A., 1929. Die aufnahme von gasen durch metalle. Z. Met.kd. 21, 37–46. Solomatov, V.S., 2000. Fluid dynamics of a terrestrial magma ocean. In: Canup. R.M.,
- Solomatov, V.S., 2000. Fluid dynamics of a terrestrial magma ocean. In: Canup, R.M Righter, K. (Eds.), Origin of the Earth and Moon. Tucson, pp. 323–338.
- Sossi, P.A., Burnham, A.D., Badro, J., Lanzirotti, A., Newville, M., O'Neill, H.S.C., 2020. Redox state of Earth's magma ocean and its Venus-like early atmosphere. Sci. Adv. 6, 1–9. https://doi.org/10.1126/sciadv.abd1387.
- Stephant, A., Wadhwa, M., Hervig, R., Bose, M., Zhao, X., Barrett, T.J., Anand, M., Franchi, I.A., 2021. A deuterium-poor water reservoir in the asteroid 4 Vesta and the inner solar system. Geochim. Cosmochim. Acta 297, 203–219. https://doi.org/10.1016/j.gca.2021.01.004.
- Stolper, E., 1982. Water in silicate glasses: an infrared spectroscopic study. Contrib. Mineral. Petrol. 81, 1–17.
- Takahashi, E., 1986. Melting of a dry peridotite KLB-1 up to 14 GPa: implications on the origin of peridotitic upper mantle. J. Geophys. Res. 91, 9367–9382.
- Tartèse, R., Sossi, P.A., Moynier, F., 2021. Conditions and extent of volatile loss from the Moon during formation of the Procellarum basin. Proc. Natl. Acad. Sci. USA 118, 1–8. https://doi.org/10.1073/pnas.2023023118.
- Tollan, P., Hermann, J., 2019. Arc magmas oxidized by water dissociation and hydrogen incorporation in orthopyroxene. Nat. Geosci. 12, 667–671. https://doi.org/10.1038/s41561-019-0411-x.
- Tomlinson, J.W., 1956. A note on the solubility of water in a molten sodium silicate. J. Soc. Glass Technol. 40, 25–31.
- Turbet, M., Bolmont, E., Bourrier, V., Demory, B.O., Leconte, J., Owen, J., Wolf, E.T., 2020. A review of possible planetary atmospheres in the TRAPPIST-1 system. Space Sci. Rev. 216, 100. https://doi.org/10.1007/s11214-020-00719-1.

- Unterborn, C.T., Schaefer, L., Krijt, S., 2020. The composition of rocky planets. In: Tasker, E.J., Unterborn, C.T., Laneuville, M., Fujii, Y., Desch, S.J., Hartnett, H.E. (Eds.), Planetary Diversity. AAS-IOP, Bristol, UK, pp. 1–52.
- Valencia, D., Sasselov, D.D., O'Connell, R.J., 2007. Radius and structure models of the first super-Earth planet. Astrophys. J. 656, 545–551. https://doi.org/10.1086/509800
- Wadhwa, M., 2008. Redox conditions on small bodies, the Moon and Mars. Rev. Mineral. Geochem. 68, 493–510. https://doi.org/10.2138/rmg.2008.68.17.
- Warren, J.M., Hauri, E.H., 2014. Pyroxenes as tracers of mantle water variations. J. Geophys. Res., Solid Earth 119, 1851–1881. https://doi.org/10.1002/2013JB010328.Received.
- Withers, A.C., Bureau, H., Raepsaet, C., Hirschmann, M.M., 2012. Calibration of infrared spectroscopy by elastic recoil detection analysis of H in synthetic olivine. Chem. Geol. 334, 92–98. https://doi.org/10.1016/j.chem.
- Young, E.D., Macris, C.A., Tang, H., Hogan, A.A., Shollenberger, Q.R., 2022. Isotope velocimetry: experimental and theoretical demonstration of the potential importance of gas flow for isotope fractionation during evaporation of protoplanetary material. Earth Planet. Sci. Lett. 589, 117575. https://doi.org/10.1016/j.epsl.2022. 117575
- Zeng, L., Jacobsen, S.B., Hyung, E., Levi, A., Nava, C., Kirk, J., Piaulet, C., Lacedelli, G., Sasselov, D.D., Petaev, M.I., Stewart, S.T., Alam, M.K., López-Morales, M., Damasso, M., Latham, D.W., 2021. New perspectives on the exoplanet radius gap from a mathematica tool and visualized water equation of state. Astrophys. J. 923, 247. https://doi.org/10.3847/1538-4357/ac3137.