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Mantle rain towards the Earth's surface: a model for the internal cycle of water

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8 Abstract

9 The internal or deep water cycle controls the volume of the oceans at the surface of the Earth. The 10 advent of subduction 2-3 billion years ago initiated the transport of water back to the Earth's interior. 11 With one ocean mass injected into the deep mantle over the last 2-3 billion years, some mantle regions 12 must have become saturated and thus turned into a deep source of water. The mantle transition zone 13 (MTZ) between 410 and 660 km depths is unlikely to be a source of hydrous melt, because its minerals 14 can integrate several thousand ppm of water. On the contrary, the low-velocity layer (LVL) lying above 15 the 410 km-discontinuity is one such source. As proposed by the "Transition-Zone-Water-Filter Model", 16 the LVL is ubiquitously formed by the global uplift of the hydrous MTZ as a counter flow of subduction 17 of slabs into deeper regions. The seismic signature of the LVL is compatible with the presence of 0.5 to 18 1% melt. This melt is produced by dehydration melting during upwelling of the mantle transition zone 19 (MTZ) containing 2200(300) ppm wt H₂O, which corresponds to 0.6 ocean mass stored today in the 20 MTZ. Hydrous silicate melt can be gravitationally stable just above the 410 km discontinuity. We propose, here, that at the upper limit of the LVL it becomes buoyant, especially where the mantle is 21 22 particularly hot and/or hydrous. Once it becomes buoyant, the melt can percolate rapidly upwards 23 through the mantle. As a consequence, the olivine-bearing mantle (OBM) could be almost saturated in 24 water, due to the presence of upwelling hydrous melt. On its path, the melt may be responsible for the 25 seismic low-velocity zones at mantle depths of between 80 and 150 km. It could also be a source for 26 refertilisation of the lithospheric mantle. Based on this model, there should be ~1.0 oceanic mass (OM) 27 stored in the upper mantle today. Secular cooling of the mantle implies an increased capacity of the 28 OBM minerals to store water. The related decrease of oceans' mass at the Earth's surface is estimated to 29 $\sim 20\%$ per one billion years.

30

31 Introduction

Water is present at the surface of our planet, which made the development of life possible. Cosmochemical arguments indicate that the amount of initial water in the planet could have been equivalent to several oceanic masses ($1 \text{ OM} = 1.4.10^{21} \text{ kg H}_2\text{O}$) depending on the source (Marty, 2012; Piani et al., 2020). Regardless of the origin of water on Earth, the deep water cycle determines how much of the accreted water resides at the surface and how much is trapped in the planet's interior. In this article, we develop a model based on the water storage capacity of mantle minerals, which brings new constraints on the distribution of water in the Earth's mantle.

39 Modern plate tectonics has been operating for the last 2-3 billion years (Ga), during which water 40 has been continuously introduced into the mantle at subduction zones. This regassing flux is particularly 41 difficult to assess, since the different layers of a slab have heterogeneous water contents and the degree 42 of serpentinization can vary as a function of the geological setting. Recent estimates converge to 1 to 2 43 10^{12} kg.yr⁻¹ of water transported to the mantle at subduction zones, which equates to 0.7 to 1.4 times the 44 OM every Ga. Still, most of the water is thought to escape from the slab, percolate into the mantle wedge 45 and trigger arc volcanism. Only 30% to 40% of the initial slab water could be efficiently carried down to the deep mantle, resulting in a flux of 0.3 to 0.8 10¹² kg.yr⁻¹ H₂O or 0.2 to 0.5 OM/Ga (Rupke et al., 46

47 2004; van Keken et al., 2011). With increasing depth, the hydrous minerals that are stable at the Earth's 48 surface break down upon heating and their water content can eventually be incorporated as H point 49 defects in nominally anhydrous minerals (NAMs) (Ohtani et al., 2018; Padrón-Navarta and Hermann, 50 2017). This evolution is facilitated by an increasing H-solubility in NAMs with increasing pressure. At 51 depths greater than 410 km, the formation of wadsleyite and ringwoodite increases hydrogen solubility 52 by about one order of magnitude (Bolfan-Casanova et al., 2000; Ferot and Bolfan-Casanova, 2012; 53 Inoue, 1994; Ohtani et al., 2000). We note that a high water storage capacity does not necessarily imply 54 a high water content. Still, the discovery of the first terrestrial ringwoodite, which contains 1.2 wt% 55 H₂O, in a deep diamond, is a proof that water can reach the transition zone (Pearson et al., 2014).

56 Today, there is no consensus on the water content in the mantle transition zone (MTZ) with budgets 57 varying from 0.3 to 3 OM (corresponding to ~1000 ppm to 1 wt% H₂O). A number of arguments have 58 been proposed for or against high water contents based on different approaches: (i) elasticity 59 measurements related to observed velocity jumps at 410 km depth invoke water contents that vary 60 between 0.3 and 0.6 wt% H₂O (Buchen et al., 2018; Mao et al., 2008). (ii) Electrical conductivity yields 61 controversial results from negligible water contents (Yoshino et al., 2008) to 0.1-0.2 wt% (Huang et al., 62 2005). (iii) It has also been proposed that 1-2 wt% H₂O are necessary to explain the viscosity contrast 63 at 660 km depth, as inferred from the effect of water content on dislocation mobility in bridgmanite and 64 ringwoodite (Fei et al., 2017). Of course, the water content of the MTZ does not have to be 65 homogeneous. Hydrogen has a slow diffusion rate in the solid state (Demouchy, 2010; Hae et al., 2006), 66 which can, however, be counterbalanced by a long residence time of a slab in the MTZ (Fukao and 67 Obayashi, 2013).

68 A crucial geological setting for the internal cycle of water is the global low velocity layer (LVL) 69 above the 410 km discontinuity (Revenaugh and Sipkin, 1994; Tauzin et al., 2010; Vinnik and Farra, 70 2007). It is unanimously interpreted as being due to dehydration melting (Revenaugh and Jordan, 1991; 71 Song et al., 2004; Tauzin et al., 2010) and this is a strong indication that the transition zone is hydrous. 72 Melting triggered by dehydration of the rising mantle occurs if the water content in the MTZ exceeds 73 the water solubility of the olivine-bearing mantle (OBM). The LVL displays an average shear-wave 74 velocity drop ($\delta V_s/V_s$) of about 4-5%, as reported in previous works, compared to the overlying OBM. 75 Based on the experimental relationship between the Vs drop in hydrous peridotites and the amount of 76 melt produced, the melt content in the LVL was constrained between 0.5 to 1.0 vol. % (Freitas et al., 77 2017; Xiao et al., 2020). The Transition Zone Water Filter model (Bercovici and Karato, 2003) is used 78 as a mean of explaining the presence and fate of the melt in the LVL. It involves a continuous production 79 of melt in the LVL associated with a water cycle around the transition zone, functioning in a closed 80 system with only one input: water from the slab. The hypothesis is that, once formed atop the 410 km 81 discontinuity, the dense melt travels laterally in the LVL until it freezes in contact with the cold slab and 82 gets subducted (e.g. (Leahy and Bercovici, 2007)). The water would be released again after crossing the 83 660 km boundary and would diffuse up to the MTZ because water incorporation in bridgmanite is low 84 (Bolfan-Casanova et al., 2003; Liu et al., 2021). Seismic low velocity zones below the 660-km 85 discontinuity indicate inefficient penetration of water into the lower mantle (Durand et al., 2017; Liu et 86 al., 2016; Schmandt et al., 2014). However, it should be relatively difficult to subduct the LVL melt 87 because it is lighter than the MTZ (Matsukage et al., 2005; Sakamaki et al., 2006). In addition, the 88 sustainability of a closed deep water cycle with water entering from the slab is questionable: if 0.2 to 89 0.5 OM were to be injected into the deep mantle every Ga (see above), the water content in the MTZ 90 should have increased by 0.4-1.5 OM and hence the ocean level decreased of the same amount since the 91 beginning of plate tectonics 2-3 Ga ago. This is obviously too extreme and incompatible with previous 92 proposals of a 10-20% decrease in ocean mass since the Archean, based on hydrogen isotopes (Lecuyer 93 et al., 1998; Pope et al., 2012) and geodynamic modeling (Flament et al., 2013).

At the other end of the deep water cycle, mid-ocean ridges are the main recognized geological setting where water is transferred from the mantle to the crust (in this study, water outgassing by arc

- volcanism is not considered in the mantle budget because we only consider the flux entering the mantle
- beyond the formation of arcs, which is 0.3 to 0.8 10^{12} kg.yr⁻¹ H₂O, as proposed by van Keken et al.,
- 98 2011). The rate of basalt production is ~20 km³.yr⁻¹ (Cogne and Humler, 2006), corresponding to ~1.2
- 10^{11} kg.yr⁻¹ of water assuming 2000 ppm water in basalts (Michael, 1988). This is at least twice less
- water than the lowest estimate of the water flux that is efficiently injected into the deep mantle by subduction. The difference of at least $0.2-0.7 \ 10^{12} \text{ kg H}_2 \text{O.yr}^{-1}$ points out to the need of considering other
- 102 possible sources of outgassing.
 - 102 possible sources of outgassing.

103 In this article, the strategy is to use quantitative constraints arising from mineral physics data, such 104 as the water storage capacity of minerals, the effect of partial melting on seismic properties, the effect 105 of composition (FeO and H₂O) on melt density, to develop a new model of the internal cycle of water. 106 We explain how the LVL can play a major role in extracting water from the major reservoir constituted 107 by the MTZ. The amount of water transported by the melt counterbalances the incoming water flow 108 associated with the entry of slabs into the MTZ, resulting in a stable water balance for the MTZ. We 109 investigate the conditions that can make buoyant a water-bearing melt formed in the LVL. Finally, we 110 describe the possible implications of such "mantle rain" and compare them with various geological 111 observations.

112

113 Maximum water solubility of the olivine-bearing mantle

114 The upper mantle is constituted of nominally anhydrous minerals (NAMs), which can 115 accommodate water in their lattice in the form of OH defects. The major upper mantle phase, olivine, 116 can incorporate more than 1000 ppm by weight of H_2O at high upper mantle pressures. The maximum 117 water solubility in olivine is a function of pressure, temperature and oxygen fugacity. When the 118 solubility limit is overpassed, an H₂O-rich phase appears, which can be a water-rich fluid at pressures 119 lower than a couple of GPa or a water-bearing silicate melt at higher pressures (Kessel et al., 2015; Mibe 120 et al., 2007). The phase equilibrium between the H₂O-bearing melt and the H-saturated olivine yields 121 the following relations:

(2)

122
$$Mg_2SiO_4 + n H_2O \ll Mg_{1-n}H_{2n}SiO_4 + n MgO$$
 (1)

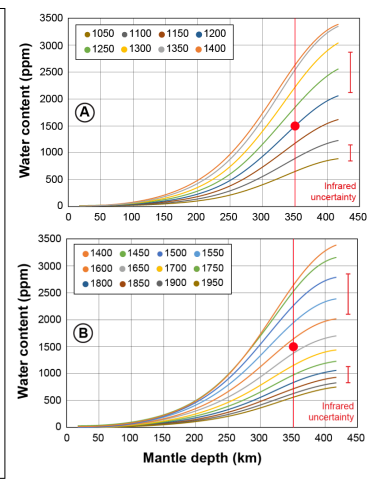
123
$$[OH]^{olivine} = f_{H2O}^n \exp(\Delta G^{Ol}/RT)$$

 f_{H2O} is the water fugacity (see details in Suppl. Material) and $\Delta G^{OI} (\Delta H^{OI} - T\Delta S^{OI} - P\Delta V^{OI})$ the Gibbs 124 125 free energy of the formation of H-defects in olivine, n is the exponent describing the incorporation 126 mechanism. A best fit to the available experimental data (see (Ferot and Bolfan-Casanova, 2012) and 127 references therein) yields the following values (assuming that H is substituting for Mg, hence n=1): (i) 128 formation enthalpy ΔH^{Ol} = 47 kJ/mol, which compares well with previous reports of 37.1 kJ/mol (Bali 129 et al., 2008) and 50 kJ/mol (Zhao et al., 2004), (ii) formation entropy ΔS^{Ol} increasing from 75 to 92 130 J/molK when pressure increases from 1 to 12 GPa, as described in (Bali et al., 2008; Smyth et al., 2006), 131 and (iii) constant activation volume at $\Delta V^{Ol}=10.6 \text{ cm}^3/\text{mol}$ (Kohlstedt et al., 1996; Mosenfelder et al., 132 2006). Note that the hypothesis of site substitution of H in Mg (n=1) or Si (n=2) site is necessary to 133 extract the thermodynamic parameters from the solubility data, however, as demonstrated by (Yang, 134 2016) both sets of $(n, \Delta V)$ for n=1 or n=2 equally describe the data. Then, the maximum water-solubility 135 limit in olivine can be calculated at all P-T conditions relevant to the upper mantle using Suppl-Eq. 1 136 (Suppl. Material) addressing the value of f_{H2O} in Eq. 2.

137 We can also calculate the H-content in the other mantle minerals coexisting with olivine under 138 water-saturation conditions, using the published partitioning coefficients of H between olivine (Ol), 139 enstatite (Opx), clinopyroxene (Cpx) and garnet (Gt). We adopt a H partition coefficients of 1 between 140 olivine and garnet $D^{water}_{ol/gt}$ (Ardia et al., 2012; Novella et al., 2014), a $D^{water}_{opxx/ol}$ decreasing from 2 to 141 0.9 between Opx and olivine with increasing pressure from 2.5 to 9 GPa (Ferot and Bolfan-Casanova, 2012) and a D^{water}_{cpx/opx} of 2 between Cpx and Opx (Demouchy et al., 2017). Finally, the well-known
modal fractions of each of these phases in a peridotite-type upper mantle (Irifune and Isshiki, 1998)
enables the calculation of the maximum water-solubility limit of the bulk olivine-bearing mantle (OBM)
at any depth, following a given adiabatic temperature gradient (Fig. 1).

146 For a present-day mantle potential temperature (T_P) of ~1650 K (Katsura et al., 2010), the H₂O 147 solubility limit of the OBM increases with depth from a few tens of ppm at the surface to ~1700 ppm 148 wt at the 410 km seismic discontinuity. The effect of temperature is major, with a solubility limit 149 decreasing by a factor of ~4 for T_P increasing from 1500 to 1950 K. It implies an increase of the water-150 storage capacity of the mantle with secular cooling that we will discuss later in more detail (see also 151 (Dong et al., 2021)). We note that this calculation does not include possible effects associated with an 152 increased water solubility limit in the first 100 km of the lithosphere (Green et al., 2014; Mierdel et al., 153 2007). In addition, the present model assumes conditions at which H_2O is the major hydrogen species 154 in the melt (Druzhbin et al., 2021).

Fig. 1: Water-saturation limit of a peridotite-type mantle. Colored curves present the water saturation limits of the mantle as a function of depth along an adiabatic temperature gradient prevailing for mantle potential temperatures (T_P) from (A)1050 to 1400 K and (B) 1400 to 1950 K (derived from (Ferot and Bolfan-Casanova, 2012)). A hydrous melt (or fluid) occurs at water contents above the saturation limit. At a given mantle depth, the saturation limit first increases with temperature up to T_P ~1400K and then decreases with further T increase. The latter effect is due to silicate dissolution in the melt and, therefore, a decrease in the activity of water in the hydrous silicate-melt at high temperature (see (Bali et al., 2008) and text). Uncertainty in the determination of the water content using infrared is +/-15% (Ferot and Bolfan-Casanova, 2012). Red dot corresponds to 1500 ppm H₂O in the mantle at 350 km depth (see text).



156

157 H2O-saturated solidus of upper mantle

The solubility curves of Fig. 1 can be inverted into H_2O -saturated solidii (Fig. 2), considering that a melt appears when, at any P-T condition above the second critical point of water, the water solubility limit of the OBM is overpassed. The "500 ppm H_2O -saturated solidus", for instance, is the P-T profile at which a water content above 500 ppm produces an incipient amount of melt saturated in H_2O . Due to the formalism of Eq. 2 (see Suppl. Material), the calculated H_2O -saturated solidii present a complex shape in the P-T diagram. In particular, they do not fall parallel to the dry-solidus (here we use that from (Andrault et al., 2018), but the same observation would also stand for other works). Reasons for this are
intrinsic to the thermodynamical properties of the minerals and hydrous melt. If we consider, for
example, 1500 ppm of water in the mantle at 350 km depth (red dot in Fig. 1), such water content is
higher than the water saturation limit, and would therefore yield the formation of a aqueous fluid, at
temperatures below 1200 K (Fig. 1A) and hydrous melt above 1625 K (Fig. 1B). In contrast, the NAMs
integrate 1500 ppm in the temperature range between 1200 and 1625 K. This yields a loop shape H₂O-

170 saturated solidus in Fig. 2. We note that the effect of water on the

thermodynamic parameters of the solid and melt may also inducemantle melting at temperatures just below the dry-solidus (Aubaud

173et al., 2004). In this case, however, neither the melt nor the solid174residue would be saturated in H_2O . We do not report this effect in175Fig. 2.

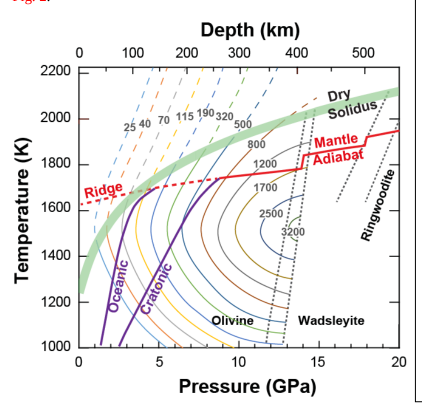


Fig. 2: The H₂O-saturated mantle solidii for water contents between 25 and 3200 ppm H_2O . The colored lines with associated water contents arise from the maximum water solubility displayed in Fig. 1. An H₂O-saturated melt appears on the left side of these curves (because water solubility decreases upon decompression) for H_2O contents equal or superior to the reported number. We also report (i) the dry-solidus of chondritic-type mantle (green thick line (Andrault et al., 2018)) and (ii) the adiabatic profile temperature corresponding to $T_P=1600$ -1650 K (Katsura et al., 2010). This figure does not consider the possible effect of a higher H₂O solubility limit in the lithosphere (Green et al., 2014; Mierdel et al., 2007; Sarafian et al., 2017). Uncertainty on water contents as in Fig. 1.

176

177 Continuous dehydration melting in the uprising mantle

178 A water content higher than the maximum water solubility induces the coexistence of water-179 saturated melt and solid residue. At a depth of 410 km, the OBM residue (produced by dehydration 180 melting) should contain about 1700(200) ppm-wt H₂O (Fig. 2). This is much higher than the water 181 content in the shallow source of mid ocean-ridge basalts (MORB), which contains between 50 and 250 182 ppm-wt H₂O at around 100 km depth (Michael, 1988). An appealing option to solve this discrepancy is 183 that, during upwelling, the mantle H₂O-content decreases along with the water storage capacity limit 184 (Ferot and Bolfan-Casanova, 2012) (Fig. 1). Hence, the mantle would remain partially molten at all 185 depths (Hirschmann et al., 2005), until, at shallow depths, it finally reaches the value invoked for the 186 MORB source. This seems incompatible with electrical conductivity measurements that do not favor a 187 partially molten upper mantle between 180 and 350 km depth (Freitas et al., 2019; Toffelmier and 188 Tyburczy, 2007). At least it implies that the fraction of melt should remain small. Based on the water-189 saturation profiles (Fig. 1), we calculate that the uprising by e.g. 10, 30 or 60 km from 410-km depth of 190 the water-saturated mantle residue would produce ~0.07, ~0.25 or ~0.58 % of melt (colored dots in Fig. 191 3), respectively, assuming that the hydrous melt contains 6.5 wt% of water (see Table 1 and text below). 192 The same calculation can be performed for the uplift of the mantle residue from any mantle depth. The 193 maximum amount of melt produced by an uplift of e.g. 30 km is ~0.37 wt%. This calculation shows that 194 the amount of melt can be very small at any mantle depth if the melt that is regularly produced by 195 dehydration melting is rapidly segregated from the uprising mantle. For a comparison, the seismic and 196 electrical-conductivity anomalies observed between 80 and 150 km depth are compatible with more than

197 0.2% (Chantel et al., 2016) and between 0.3 to 2.0 wt% (Ni et al.,

198 2011) of melt, respectively. At intermediate upper-mantle depths,

the melt is buoyant and can travel upwards toward the Earth'ssurface. This could make its presence undetectable to

201 geophysical measurements.

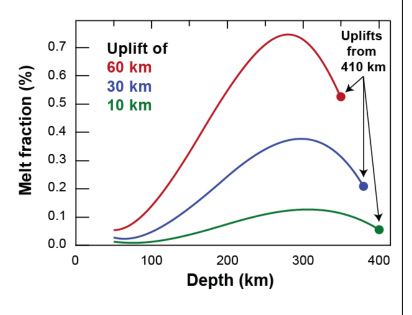


Fig. 3: Dehydration melting in a rising upper mantle: the adiabatic uplift (i.e. decompression) of a waterinduces residue saturated continuous dehydration melting, due to a decrease of the watersaturation limit with decreasing mantle depth (Fig. 1). Here, we report the fraction of melt that can be expected at a given depth when a water-saturated mantle residue comes from 10, 30 or 60 km deeper, without loss of water during its ascent. Uplift from the 410 km seismic discontinuity are the colored dots. The curved shape with maxima around 275-300 km depth is due to the complex shape of the maximum water solubility limit in the mantle (Fig. 1). Uncertainties are +/-15%, same as in Fig. 1.

202

203 Dehydration melting in the LVL: melt composition and phase diagram

204 The occurrence of a global low velocity layer (LVL) above the 410 km discontinuity (Revenaugh 205 and Sipkin, 1994; Tauzin et al., 2010; Vinnik and Farra, 2007) is unanimously interpreted as being due 206 to dehydration melting (Revenaugh and Jordan, 1991; Song et al., 2004; Tauzin et al., 2010). This is a 207 strong indication that the transition zone is hydrous. Melting triggered by dehydration of the rising 208 mantle occurs if the water content in the MTZ exceeds the water solubility of the olivine-bearing mantle 209 (OBM). The LVL displays an average shear-wave velocity drop ($\delta V_s/V_s$) between 4 and 5.6 %, as 210 reported in previous works, compared to the overlying OBM. Based on the experimental relationship 211 between the Vs drop in hydrous peridotite and the amount of melt produced, the melt content in the LVL 212 was constrained between 0.5 to 1.0 vol. % (Freitas et al., 2017; Xiao et al., 2020). Dihedral angles are 213 much lower (less than 10°) in the hydrous peridotite system at high-pressure (Freitas et al., 2017; 214 Yoshino et al., 2007) and the amount of melt necessary to explain $\delta Vs/Vs$ is predicted to decrease at 215 lower wetting angle (Hier-Majumder et al., 2014). Low wetting angles and low surface tension (Hier-216 Majumder et al., 2006; Yoshino et al., 2007) favor the fast mobility of the melt at grain boundaries. This 217 is compatible with another major seismic observation in the LVL; no major melt ponds but instead a 218 continuous global layer of partial melt that is not limited to regions associated with subduction or mantle 219 plumes (Tauzin et al., 2010).

The composition of the water-bearing LVL-melt can be studied experimentally by reproducing an
 equilibrium texture between peridotite and 0.5 to 1 wt% melt. Analyses of thin veins of melt
 corresponding to ~2% partial melting indicate ~21.2 wt% FeO (recalculated after (Freitas et al., 2017)).
 However, the composition of liquids originating from low degrees of partial melting can be biased by

- 224 crystallization upon quenching, inducing an enrichment of incompatible elements in the residual melt.
- 225 Alternative methods have been developed in order to determine the composition of incipient melts 226
- accurately, such as the iterative sandwich method (Dasgupta and Hirschmann, 2007), which yields a 227 melt FeO-content of ~13.4 wt% for the LVL melt (Condamine and Frost, 2018). On the other hand,
- 228 density measurements show variations of +21 and -16.5 kg/m³ for every additional mol% FeO and wt%
- 229 H₂O, respectively (Matsukage et al., 2005), in agreement with thermodynamical modeling (Freitas et
- 230 al., 2017). As a result, a melt with an FeO-content of 11.6 wt% (used in (Matsukage et al., 2005)) or
- 231 13.4 wt% (ref. (Condamine and Frost, 2018), Table 1) would be neutrally buoyant at 350 km, at the top
- 232 of the LVL, for H₂O contents of ~6 or ~7.1 wt% H₂O, respectively, in good agreement with previous
- 233 work (Matsukage et al., 2005; Sakamaki et al., 2006). For modelling purposes, we use an average LVL-
- 234 melt with reference FeO and H_2O contents of 12.5(1.0) and 6.5(1.0) wt %, respectively. We note that
- 235 such FeO-content in the LVL of a peridotite-type mantle implies a solid-melt FeO partition coefficient
- 236 $D_{\text{FeO}} \sim 0.65$, which is compatible with $D_{\text{Fe}} \sim 0.56$ proposed previously (Mibe et al., 2006).

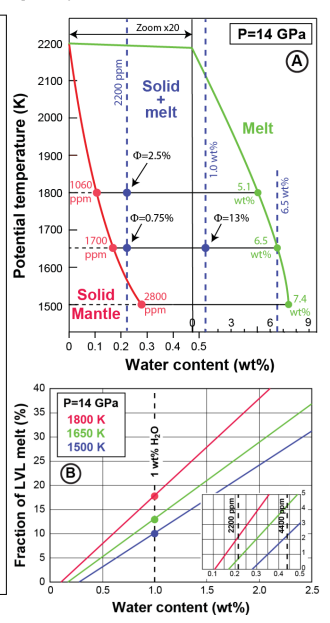
Composition of LVL	melts	SiO2	TiO2	AI2O3	FeO	MgO	CaO	Na2O	H20	Total
Matsukage et al.	meas. (wt%	35.5		3.3	11.6	30.4	14.1		6.0	100.9
(2005)	norm. (wt%	35.2		3.3	11.5	30.1	14.0		5.9	100.0
	mol%	27.8		1.5	7.6	35.5	11.8		15.7	100.0
Condamine and	meas. (wt%	34.9	0.6	0.7	12.9	27.0	12.5	0.45	7.1	96.1
Frost (2018)	norm. (wt%	36.3	0.6	0.7	13.4	28.1	13.0	0.47	7.4	100.0
	mol%	28.1	0.4	0.3	8.7	32.4	10.8	0.35	19.0	100.0
Freitas et al. (2017)	meas. (wt%	29.7		2.6	21.2	2.6	9.0	0.07	13.6	78.8
F=2%	norm. (wt%	37.8		3.2	27.0	3.3	11.4	0.09	17.3	100.0
	mol%	27.6		1.4	16.4	3.6	8.9	0.06	42.0	100.0
Preferred values	wt%				12.5				6.5	

Table 1: Composition of the LVL melt: (Matsukage et al., 2005) reported the neutral buoyancy in the LVL of a melt containing 11.6 wt% FeO and 6.0 wt% H₂O. The concentration in major elements of the LVL-melt was further refined using the iterative sandwich method (Condamine and Frost, 2018). They refined a melt Fe-content of \sim 13.4 wt%, which should be neutrally buoyant in the LVL for an H₂O content of ~7.1 wt% (Freitas et al., 2017; Matsukage et al., 2005). In a later work, incipient melting was induced by dehydration melting while measuring the sound wave velocities under conditions relevant to the LVL (Freitas et al., 2017). In these experiments, however, crystallization of some olivine could have occurred upon quenching, making the primary melt composition difficult to retrieve. We calculate a solid-melt FeO partitioning coefficient D_{FeO}~0.65 between the LVL melt and the solid mantle residue.

237 We now draw the melting diagram of the hydrous mantle at 14 GPa (Fig. 4A). The solidus is the water storage capacity based on mineral properties (Fig. 1). At temperatures of 1500, 1650 and 1800 K, the maximum H_2O -content in the mantle is 2800(300), 1700(200) and 1060(100) K, respectively. For a mantle potential temperature (T_P) of 1650 K, the hydrous melt that is in equilibrium with the watersaturated mantle should contain ~6.5 wt% H₂O (Table 1). Using mass conservation we calculate the melt fraction in the LVL as a function of the bulk water content (Fig. 4B). A bulk-mantle water content of 2200 ppm (left vertical dashed line), 0.5 or 1 wt% (middle vertical dashed line) leads to the formation of 0.75, 5.2 or 13 wt% of melt in the LVL. The latter values are well above the 0.5-1% of melt required to account for low LVL velocities (Freitas et al., 2017), clearly indicating that the bulk water content of the LVL, and that of the MTZ as well, must be well below 1 wt%. One may argue that higher MTZ water-content could be possible, if the LVL-melt returns quickly to the MTZ (Leahy and Bercovici, 248 2007). However, (i) the LVL-melt is lighter than the MTZ (Fig. 5), with therefore no reason to sink 249 through the 410-km discontinuity, and (ii) return of hydrous LVL-melt into the MTZ would indefinitely 250 increase its water content, which does not seem sustainable. A higher water content in the MTZ is only possible if the water-content in the LVL-melt has been underestimated in previous studies (Table 1).
We will discuss the fate of the LVL-melt later in this article.

Finally, we draw the liquidus of this phase diagram passing through the melt composition at T_P=1650 K (~6.5 wt%) and assuming that it has the same, but inversed, curvature than that of the solidus (Fig. 5a). The resulting diagram suggests a slight decrease of the water content in the hydrous melt, from e.g. 7.4 to 5.1, when the potential temperature increases from 1500 to 1800 K, respectively. This indicates that for 2200 pm of water in the LVL, the melt fraction, Φ , increases from 0, ~0.75 and ~2.3 %, when temperature is 1500, 1650 and 1800 K, respectively.

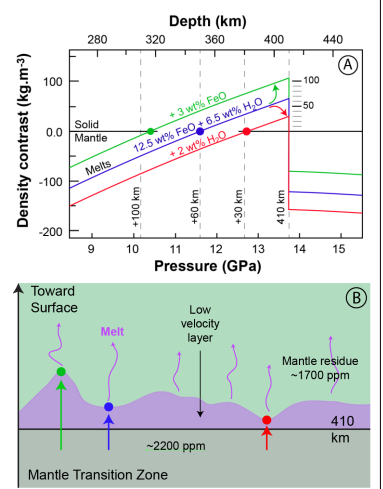
Fig. 4: (A) Phase diagram relevant to dehydration melting of the mantle: above the 410 km depth discontinuity (~14 GPa), exceeding the maximum water storage capacity of peridotite induces dehydration melting. At a mantle potential temperature (T_P) of 1650 K, the solid mantle can store a maximum of ~1700 ppm H_2O (Fig. 1) and the melt contains ~6.5 wt% H₂O (Table 1). It results that a bulk water content of 2200 ppm (left vertical dashed line) leads to the formation of 0.75% of melt, in agreement with seismic observations of the low-velocity layer (Freitas et al., 2017). Thus, the melting diagram is mainly built using constraints from previous reports. Alternatively, a bulk water content of 1 wt% (middle vertical dashed line) would yield 13% melt. At T_P=1800 K, a bulk water content of 2200 ppm would produce 2.5% of melt containing 5.1 wt% H₂O. This diagram shows that mantle heterogeneities of temperature and water-content affect significantly the degree of partial melting and the melt composition. Uncertainty on water contents is same as in Fig. 1. (B) Fraction of LVL-melt: based on the solidus and liquidus profiles, we calculate, from Fig. 4A, the fraction of melt that must be present in the LVL based on bulk mantle water content and for T_P of 1500, 1650 and 1800 K. Inset is a zoom of the melt fraction at low water contents. Uncertainty on the melt fraction is estimated to $\sim 25\%$.



259

260 Melt-Mantle density contrast

The fate of the LVL-melt is primarily controlled by the melt-mantle density contrast ($\Delta \rho = \rho_{melt}$ p_{mantle}). A melt-mantle density crossover (that is where both densities of solid and melt become equal), occurs around the 410 km depth corresponding to ~14 GPa (Matsukage et al., 2005; Sakamaki et al., 2006) (Fig. 5a), due to (i) the higher compressibility of the melt, compared to the solid mantle, (ii) the absence of a major phase transition in mantle minerals above the 410 km discontinuity, and (iii) the 266 preference of FeO for the melt. Based on the equations of 267 state and the dependence of the melt density on its FeO and 268 H₂O contents (Jing and Karato, 2012; Matsukage et al., 269 2005), we recalculate the density contrast as a function of depth for various possible melt compositions (Fig. 5a). As 270 271 expected, there is a maximum in the density contrast just 272 above the 410 km discontinuity that then decreases with 273 decreasing depth. In addition, we calculate that small 274 variations in the FeO (+/- 3 wt%) and H₂O (+/- 2 wt%) 275 contents can significantly affect the mantle depth at which 276 neutral melt-mantle buoyancy is achieved.



277 Thus, variations in melt composition can create 278 variations of the thickness of the LVL (ΔZ_{LVL}). Indeed, the 279 top of the LVL is the point of neutral melt-mantle density 280 (red, blue and green dots in Fig. 5a) and further uprising of 281 the mantle beyond this limit yields a buoyant melt, which upward migration is forced by gravity. A melt composition 282 283 enriched in iron, by e.g. 3 wt%, yields a denser melt that is 284 gravitationally stable across a broader region (from 410 to 285 \sim 320 km depth, up to the green dot in Fig. 5a), thus implying 286 an LVL which is thicker, by e.g. +40 km, compared to the 287 mean mantle, which melt is gravitationally stable up to ~ 60 288 km above the 410 discontinuity (blue dot in Fig. 5a). In

Fig. 5: (A) Melt-Mantle density cross-over. Because melts have a higher compressibility compared to solid mantle, the density the difference between melt and solid increases with increasing depth, and the melt eventually becomes denser than the solid between the neutral buoyancy depth and the 410 km depth, at the olivine-wadsleyite phase transition. This depth interval (ΔZ_{LVL}) defines the thickness of the LVL. Neutral melt-mantle buoyancy is expected at the upper limit of the LVL, which is seismically observed 60 km in average above the 410 km discontinuity (blue dot). For a mantle of peridotite composition, the LVLmelt should contain 12.5 wt% FeO and 6.5 wt% H₂O (blue line, Table The melt density changes 1). significantly as a function of its FeO (green line) and H₂O (red line) contents (Matsukage et al., 2005; Sakamaki et al., 2006). Thus, the thickness of the region where the melt is denser (within ΔZ_{LVL}), can vary laterally inside the mantle due to heterogeneities in temperature and composition (see text). Depending on their composition, melts can be buoyant (less dense) at depths shallower than the green, blue or red dot. For comparison, density of the OBM at 410 km depth is ~3500 kg.m⁻³. Uncertainties are detailed in previous works (Matsukage et al., 2005; Sakamaki et al., 2006) (B) The "mantle-rain" model proposes that the LVL-melts rise upward through the mantle once they reach the upper limit of the LVL. Melts migration occurs vertically and laterally along grain boundaries. The observed regional variations of the LVLthickness can be explained by mantle heterogeneities in FeO, H₂O contents and temperature, which induce LVLmelts that are relatively denser (green dot) or lighter (red dot), compared to the typical LVL-melt (blue dot, Table 1). Still, any melt composition becomes buoyant above a given mantle depths (Fig. 5a).

contrast, a melt enriched in H_2O , by e.g. 2 wt%, yields a lighter melt that is gravitationally stable up to 30 km only above the 410 km discontinuity (red dot in Fig. 5a). Chemical variations of the melt can be induced by mantle heterogeneities in temperature, water content or even fertility. They can induce variations of the LVL thickness from ~30 to ~100 km that are compatible with seismological observations (Revenaugh and Sipkin, 1994; Tauzin et al., 2010; Vinnik and Farra, 2007).

Knowing the melting diagram of hydrous peridotite (Fig.4a), the manner in which the variation in mantle composition and temperature affect the density of the LVL melt can be assessed by constraining the melt fraction (Fig.4b). Indeed, given that the 410 km discontinuity is well explained globally by a peridotite (or pyrolite) mantle, there is no reason at this stage to invoke other components such as eclogite or sediments.

299 (i) Since water is an incompatible compound, a local increase of the water content in the MTZ 300 would induce a higher degree of partial melting in the LVL (Fig. 4b). We determine that twice more 301 water in the MTZ than the value of 2200 ppm would produce ~6 times more LVL-melt (~4.4 wt%, see 302 inset of Fig. 4b). Such melt would contain the same amount of H₂O (fixed at the liquidus, Fig. 4a), but 303 a lower FeO-content (reduced by ~1% compared to 6.5 wt%; Table 1), according to mass conservation 304 $[FeO]_{Melt} = [FeO]_{Mantle} / (\Phi + (1 - \Phi)D_{FeO}), where \Phi is the melt fraction and using D_{FeO} ~ 0.65, the solid-melt$ 305 FeO partition coefficient. As a result, such melt would be $\sim 0.2\%$ lighter (+21 kg/m³ for every additional 306 mol% FeO (Matsukage et al., 2005)), compared to the melt formed from a mantle containing 2200 ppm 307 H₂O. Similarly, a bulk 1 wt% H₂O would produce 13 wt% of LVL-melt (see Fig. 4b) with ~5% less FeO 308 and a density lowered by ~1%. The mantle residue also loses some Fe with increasing Φ , however, to a 309 much lesser extent. Altogether, bulk water contents of 4400 ppm or 1 wt% imply a decrease of ~6.3 and 310 \sim 36 kg.m⁻³, respectively, from a melt-mantle density contrast of \sim 65 kg.m⁻³ at the base of the LVL for a 311 water content of 2200 ppm H₂O (Fig 5a). This would yield a LVL up to ~30 km thinner, compared to 312 the average mantle.

313 (ii) The mantle could also present variations in its fertility, implying significant heterogeneities in 314 FeO-content, which incompatibility and molecular weight have important consequences on melt density, 315 as we have seen. A reduction by 10% (from e.g. 8.0 to 7.2 wt% FeO) in the FeO content of the mantle 316 would result in a stronger decrease of the melt FeO content, compared to that of the residue (due to 317 $D_{Fe}^{Solid/Melt}=0.65$), and a decrease of the melt-mantle density contrast by 0.8% or ~28 kg.m⁻³ (Matsukage 318 et al., 2005). This would yield a LVL ~30 km thicker, compared to the average mantle.

(iii) A locally higher mantle temperature would increase Φ (Fig. 4b). Such LVL-melt would be depleted in both H₂O and FeO, compared to mean mantle conditions as incompatible elements get diluted with increasing melt fraction. It is difficult to tell which of the two opposite effects (increasing FeO content or increasing H₂O content) dominates the melt-mantle density contrast. The imprecise shape of the liquidus in Fig. 4a yields a large uncertainty on Φ as a function of temperature. However, the effect of bulk water content could be significant, as illustrated by the change from 0.75 to 2.5 wt% of melt when T_P increases from 1650 to 1800 K, for a bulk H₂O-content of 2200 ppm (Fig. 4b).

(iv) Finally, the presence of carbonate in the LVL melt cannot be discarded. Carbonates are subducted to great depths and are known to react to form diamond in the transition zone (Pearson et al., 2014; Shirey et al., 2021). If present in the melt, C in the form of $CO_3^{2^2}$ is expected to increase the melt buoyancy relative to the surrounding mantle. Still, carbonates may more likely be present at depths shallower than ~150 km where the oxygen fugacity is higher (Stagno et al., 2013).

- Overall, mantle heterogeneities in FeO and H₂O contents and possibly also temperature and CO₂content appear to have a significant effect on the melt buoyancy, especially in the shallow part of the LVL where the melt-mantle density contrast is small (Fig. 5a). We acknowledge important uncertainties
- in the amplitude of the melt-density dependence on FeO, H_2O , CO_2 contents and temperature. However,

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5 the trends described above are qualitatively correct and, therefore, mantle heterogeneities could be a major controlling parameter of the LVL-thickness.

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338 Fate of hydrous-melt in the upper mantle

339 Overall, we expect the production of water-rich melts to be widespread in the OBM: not only the 340 LVL could be a continuous source of buoyant melts, but the uprising residual mantle is expected to 341 dehydrate progressively on its way toward the surface (Fig. 3). While the vertical mobility of the melt 342 is certainly triggered by a melt being more buoyant with increasing elevation from the 410 km 343 discontinuity (Fig. 5a), lateral migration is also facilitated by several parameters: (i) a low dihedral angle 344 in the hydrous peridotite system at high pressure (Yoshino et al., 2007) (ii) significant grain to grain 345 surface energy between silicate minerals, ~4 J/m² for olivine (Swain and Atkinson, 1978), though 346 unfortunately poorly constrained in the presence of majoritic garnet, the second major phase at high 347 upper-mantle depths and (iii) a major relaxation of the surface energy under hydrous conditions (de 348 Leeuw et al., 2000). Large-distance melt propagation has already been proposed for the LVL-melt above 349 the 410 km discontinuity (Leahy and Bercovici, 2007). Widespread circulation of metasomatic agents 350 is also recognized in the cratonic lithosphere (O'Reilly and Griffin, 2013). We argue that a similar 351 process also occurs within the convecting mantle and it is possible that chemical reactions favor mantle 352 refertilization. As observed in nature, the percolation of hydrous melt into a mantle region that is not 353 saturated in water would induce hydration of NAMs, drying out the melt and inducting its solidification 354 (Denis et al., 2015; Doucet et al., 2014; Le Roux et al., 2007). Over geological time, the travelling melts 355 can resolve at least partially the heterogeneity in water content between the different mantle regions. In 356 this way, all mantle regions (even those that do not undergo dehydration melting upon uprising) could 357 achieve a water content close to the saturation limit. For further modeling of the total water content in 358 the upper mantle, we consider that the minerals of the OBM carry an amount of water that is close to 359 the water saturation limit (Fig. 1).

360 On its way through the OBM, the buoyant hydrous melt eventually meets a critical depth where it 361 decomposes into a mixture of volatile-bearing fluid and silicate-melt (Kessel et al., 2015; Mibe et al., 362 2007). The appearance of a fluid (Gaillard et al., 2008; Hirschmann, 2010) as well as the local 363 accumulation of a viscous silicate-melt (Chantel et al., 2016; Priestley and McKenzie, 2013) could 364 produce the low velocity zones (LVZ) detected by seismic and electrical conductivity measurements. 365 The range of depths between 80 and 150 km where LVZs are detected (e.g. (Debayle and Ricard, 2012; 366 Kawakatsu et al., 2009)) is compatible with the P-T conditions, around 3.8 GPa and 1273 K, at which 367 the second critical end point was reported in the peridotite-H₂O system (Mibe et al., 2007). It is therefore 368 appealing to make a causal link between the "mantle rain" and the LVZs. A major observation that needs 369 to be taken into account is that the LVZ seismic and electrical-conductivity anomalies become weaker 370 and deeper with increasing plate age (e.g. (Debayle et al., 2020)), while we expect ubiquitous upwelling 371 of the melt from the LVL. A greater impact of the melt in young geological settings could arise from 372 two major factors: (i) less hydrous melt reaches old compared to young lithospheres. The mantle 373 underneath mid-ocean ridges is uprising and, therefore, the mantle residue is expected to undergo 374 continuous dehydration melting on its path (Fig. 3). The hydrous-melt produced this way is added to the 375 melt coming from the LVL (left side of Fig. 6). Alternatively, the mantle underneath an old lithosphere 376 should be either stagnant (in depth) or sinking. Thus, no melt is produced by dehydration-melting upon 377 decompression. Now if we consider that the regions where the mantle is sinking, one can expect that it 378 would react and stop the LVL-melt at mid upper-mantle depth due to mantle rehydration (right side of 379 Fig. 6). (ii) Young oceanic crust presents a much steeper temperature gradient in the first 100-150 km 380 depth, compared to the continental crust. By comparing these temperature profiles and the mantle 381 solidus, it appears that the formation of a water-saturated melt evolves progressively from ~150 ppm at 382 100-150 km depth, to \sim 700 ppm H₂O at 250-300 km depth, for young oceanic to old continental settings, 383 respectively (Fig. 2). This denotes a larger impact of water in the young lithosphere.

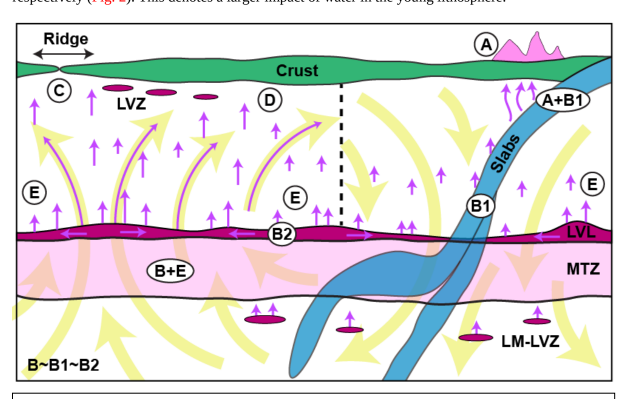


Fig. 6: Internal cycle of water and the mantle rain model. The mantle convects upward or downward in left and right parts of the diagram, respectively. Upward movement should dominate, though, in order to balance the inflow of slab (blue) into the deep mantle. Upon uprising of the water-rich MTZ material (light pink) and crossing of the 410 km mineralogical discontinuity, the mantle becomes water-saturated and dehydration melting produces the LVL-melt (dark pink layer). In addition, the uprising of the mantle residue continuously produces a water-bearing melt (curved pink arrows, Fig. 3). Above the upper limit of the LVL, the melt is buoyant and rises toward the surface (linear pink arrows, Fig. 5). Migration of the hydrous melt should take place in all directions, due to low surface tension; for clarity, lateral melt movements are not represented in this figure. except in the LVL. In mantle regions that are sinking, chemical reactions with the travelling hydrous melt is expected to hydrate mantle NAMs. Similar reactions occurring in the lithospheric mantle (point D) can produce mantle refertilisation and eventually intraplate volcanism. (A+B1) is the inflow of water at subduction zones; ~1-2 10¹² kg.yr⁻¹ but only (B1) penetrates the deep mantle; 0.3-0.8 10¹² kg.yr⁻¹. (B2) is the water contribution to produce the LVL-melt: 0.5(0.2) 10¹² kg.yr⁻¹. (C) is the outflow of water at the ocean ridges; ~0.1 10^{12} kg.yr⁻¹. (B2) should roughly equal (C+D), with (D) the water flux into the lithosphere: 0.2-0.7 10^{12} kg.yr⁻¹. While (A) to (D) are water fluxes, (E) is the steady water content in the solid upper mantle. Considering that the upper mantle is almost saturated in water yields a total water content of 8.5(0.8) 10^{20} kg H₂O or 0.6(0.1) mass of the oceans. Calculations are detailed in the main text and captions of other figures. This cycle assumes a constant amount of water in the lower mantle and in the core. We note that some dehydration melting may occur below the 660 km and induce low seismic-velocity zones (LM-LVZ).

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The transfer of melts and fluids enriched in incompatible and volatile elements to the lithospheric 385 mantle may contribute to its refertilisation. In addition to well identified, hydrous metasomatism in 386 mantle xenoliths from the cratonic mantle (Doucet et al., 2014; Le Roux et al., 2007), the presence of a 387 few percent of pyroxenite is often invoked to explain the trace element and isotopic composition of mid-388 ocean ridge basalts (MORBs) originating from the oceanic mantle (Lambart et al., 2009). The hydrous 389 melts originating from the transition zone could also help explaining the occurrence of less well-known 390 magmatism. For example, the presence of residual fluids under-plating the crust could trigger the 391 392

formation of petit-spot volcanoes (Machida et al., 2017). Hydrous plumes coming from the MTZ have also recently been invoked to explain continental magmatism such as continental flood basalts beneath

393 China (Kuritani et al., 2019; Wang et al., 2015), and beneath the Turkish-Iranian plateau

394 (Soltanmohammadi et al., 2018). It is also notable that only a few identified mantle plumes (Liu et al.,

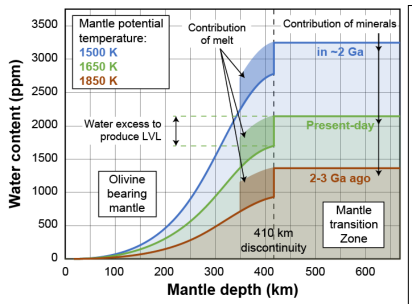
395 2018; Vinnik and Farra, 2007; Zhang et al., 2018) come from great depth (Courtillot et al., 2003).

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397 Water budget in the present-day deep mantle

398 In mantle regions subjected to continuous upwelling, melt is continuously formed in the LVL, while 399 a partially depleted, but still water saturated, residue proceeds upwards (Bercovici and Karato, 2003). 400 On a global scale, an incoming flow of ~2.5 10¹¹ m³.yr⁻¹ of subducted slab material (Wen and Anderson, 401 1995) through the MTZ implies an average uplift of ~ 0.6 mm.yr⁻¹ across the whole 410 km discontinuity. 402 In order to induce dehydration melting, the water content in the transition zone needs to be higher than 403 the maximum storage capacity of the OBM, which is 1700(200) ppm H₂O at 410 km for a mantle 404 potential temperature of 1650 K (Fig. 1, see details in Suppl. Material). In addition, some water is carried 405 by 0.5 to 1% of LVL melt containing 6.5(1) wt% H₂O. This represents an additional 300 to 700 ppm 406 (thus, 500(200) ppm) H₂O, yielding a total mantle content of 2200(300) ppm H₂O (Fig. 4) to explain the 407 seismic velocity drop in the LVL. We now consider that the average water content in the MTZ is 408 comparable to that in the LVL (2200(300) ppm), which is only ~30% higher than the maximum 409 solubility limit of the mantle (1700(200) ppm) in the LVL. This assumption implies that there is no 410 accumulation of melt in the LVL, but instead continuous melt formation by dehydration melting and 411 escape by Mantle Rain of a same amount of LVL-melt. Here, we evaluate what are the consequences of 412 this assumption: (i) if the water content of the mantle uprising from the MTZ would be lower, between 413 1700(200) and 2200(300) ppm water, then there would still be dehydration melting. There would be less 414 LVL-melt produced for a same amount of uprising MTZ material. This would lower the fluxes of LVL-415 melt out of the LVL and loss by uprising. (ii) Alternatively, more than 2200(300) ppm water in the MTZ 416 would produce a fraction of LVL-melt higher than expected from the comparison between experiments 417 and seismic data. This could be accommodated by the Mantle Rain model with a greater flux of uprising 418 LVL-melt. Still, it should be compensated by a higher influx of water in the MTZ, which does not seem 419 to be available (see text above and Fig. 6). It makes it unreasonable a MTZ water content significantly 420 higher than 2200(300) ppm. In summary, slight variations of the MTZ water content away from the 421 average value of 2200(300) ppm does not significantly change either the Mantle Rain model or the upper 422 mantle water balance estimate that we develop below. We note that this value is far below the level of 423 water saturation in MTZ, because wadsleyite can include up to 5 times more hydrogen than olivine at 424 these P-T conditions (Ferot and Bolfan-Casanova, 2012). Based on the average value of 2200(300) ppm, 425 the MTZ should contain $8.5(0.8) \ 10^{20}$ kg of water, corresponding to 0.6(0.1) OM.

426 The continuous rise of the water-bearing MTZ material by ~ 0.6 mm.yr⁻¹ corresponds to a total of ~2.0(0.3) 10^{12} kg.yr⁻¹ H₂O crossing upwards the 410 km discontinuity. This value is significantly larger 427 than the 0.3 to 0.8 10^{12} kg.yr⁻¹ H₂O injected in the deep mantle by subduction ((van Keken et al., 2011); 428 429 B1 in Fig. 6). Instead, it is the flux of water producing the LVL-melt, about 0.5(0.2) 10¹² kg.yr⁻¹ (B2 in 430 Fig. 6), that is well compatible with the flux of water entering the MTZ by subduction. This indicates 431 that the production of LVL-melt is primarily triggered by the amount of deeply subducted water. On the 432 other hand, the origin of the water present in NAMs uprising to the LVL cannot come from the subducted 433 water. In the framework of global convection, NAMs are expected to cross the 410 km boundary in both 434 directions and, for this reason, their contribution to the water transfer through the 410 km boundary is 435 balanced. By analogy, the situation is comparable to the presence of incompatible elements in peridotite; 436 they do not come from the subduction of MORBs. The water content of NAMs sinking through the 410 437 km discontinuity is unknown. However, it could be close to the NAMs saturation limit due to reaction 438 with hydrous-melt present in the upper mantle (see above). In addition, they cross the water-saturated 439 LVL and could react there with the water-bearing melt. For this reason, we consider that the water440 content of NAMs in the upper mantle as a constant (contribution E in Fig. 6).



441 Now we calculate the total amount of water present in the 442 upper mantle (Fig. 7). An OBM close to its water saturation limit 443 yields an integrated value of its water content of $\sim 4.3(0.4) \ 10^{20} \ \text{kg}$ 444 or ~0.31(0.03) OM. Added to the 0.6(0.1) OM in the MTZ (see 445 above), we conclude that a water-equivalent of $\sim 1.0(0.1)$ OM of 446 water currently resides in the mantle above the 660 km 447 discontinuity. We acknowledge that this calculation does not 448 include possible effects associated with an increase in the water 449 solubility limit in the lithospheric mantle and the crust (Green et al., 450 2014; Mierdel et al., 2007; Sarafian et al., 2017). For estimating a 451 water budget over the entire mantle, the amount of water residing 452 in the lower mantle should also be considered: different 453 experimental works based on high-quality single-crystal of

Fig. 7: Water content in the upper mantle as a function of mantle potential temperature. At a depth shallower than 410 km, the H₂O content of the mantle follows the water-saturation profiles (Fig. 1). On the other hand, the water content in the mantle transition zone can be estimated as being the sum of the solubility limit in the mantle above 410 km and the content needed water to produce 0.5 to 1 wt% of hydrous-melt in the lowvelocity layer, in agreement with seismic data. The mean thickness of the LVL is ~60 km, but depends on mantle heterogeneities in H₂O and FeO contents, and temperature (see text). The same calculation is performed for mantle potential temperatures of 1850 K (red: 2-3 Ga ago), 1650 K (green: ~present-day) and 1500 K (blue: in the future). Uncertainty on water contents is same as in Fig. 1.

bridgmanite find a radically different water storage capacity from ~100 ppm (Bolfan-Casanova et al., 2003) to ~1000 ppm (Fu et al., 2019), which translates into maxima of ~0.3 to ~3 OM in the lower mantle. Ferropericlase and CaSiO₃-perovskite may also integrate minor amounts of H. In addition, some water may be carried below the 660 km discontinuity in dense hydrous phases that remain stable in lowtemperature slabs (Nishi et al., 2014; Pamato et al., 2015). Still, partition coefficients greater than 10 between most NAMs and bridgmanite favor the storage of water in the MTZ, compared to the lower mantle (Bolfan-Casanova et al., 2003) and thus we consider that the lower mantle should be fairly dry.

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462 Water budget of the mantle over the Earth's history

463 We evaluate the amount of water that has been stored in the mantle over the course of the Earth's 464 history. Analyses of ancient non-arc basalts and komatiites suggest a rate of mantle cooling of ~100 465 K/Ga (e.g. (Herzberg et al., 2010)). A higher mantle temperature in the past implies a lower water storage 466 capacity for NAMs (Fig. 1) and thus less water stored in the upper mantle (Fig. 7). The amount of water 467 stored in the MTZ in the past can be estimated from the maximum water storage capacity of the mantle 468 just above the 410 km discontinuity making following assumptions: (i) the LVL has persisted since the 469 establishment of slab subduction, due to the parental link between subducted water and formation of the 470 LVL (see above). (ii) LVL properties, e.g. its dependence on mantle temperature and water content,

471 have remained the same over time, which enables the use of the melting diagram (Fig. 4). It implies that 472 the excess water in the MTZ, compared to the OBM, is regulated by the LVL (Fig. 7). In other words, 473 we assume that although the deep water cycle has evolved over the last 2-3 Ga, a stable equilibrium 474 between the flow of water into the MTZ via subduction and the flow of water out the MTZ via melt 475 formed at the LVL was established early on and maintained over geological times. We thus obtain water 476 contents of 930(100) or 2800(300) ppm for the OBM at 410 km depth from Fig. 1, and 1400(200) or 477 3300(500) ppm for the MTZ, for T_P varying from the highest values of 1850 K (in the past) to 1500 K 478 (in the future), respectively (Fig. 7).

479 The water storage capacity in the entire upper mantle (above 660 km depth) is found to increase

480 from ~0.6(0.1) to ~1.4(0.2) OM from 2-3 Ga ago (T_P ~1850 K) to a 481 point in the future when T_P will achieve 1500 K (Fig. 8). This 482 corresponds to a net decrease of the water elevation of the oceans by 483 ~900 m in the last 2-3 Ga, for a mean elevation today of 4.5 km. These 484 trends correspond to a decrease of the oceanic mass of ~20% per Ga, in 485 agreement with previous findings of a loss of several 10s % of the OM 486 over geological time, based on hydrogen isotopes (Lecuyer et al., 1998; 487 Pope et al., 2012) and geodynamic modeling (Flament et al., 2013). The 488 mantle storage capacity will continue to increase to a maximum value 489 of ~1.6 OM, when T_P may eventually decrease to 1400 K in several Ga 490 from now.

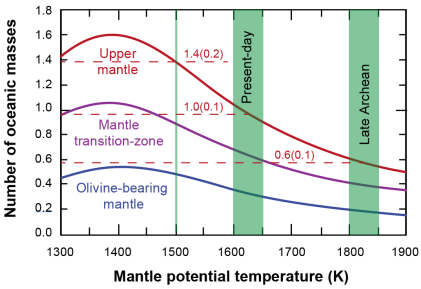


Fig. 8: Water content of upper mantle reservoirs throughout Earth's history. Water contents in the entire upper mantle (red), constituted of the OBM (blue) and of the MTZ (purple), calculated from the integration of contents water as a function of depth (Fig. 7) for various T_P. With present-day T_P ~1650 K, we expect ~1.0(0.1) ocean mass (OM) trapped in the mantle above 660 km (central red dashed line), with ~ 0.4 and ~ 0.6 OM in OBM and MTZ, respectively). The water storage capacity in the upper mantle increases with secular cooling at a rate of ~20% of the ocean mass per billion years. Uncertainty is +/-15% similar to that of Fig. 1.

491 Finally, we investigate the critical amount of water needed to produce a planet like the Earth in 492 terms of the surface ratio between oceans and emerged-continents. For the Earth, the present-day ~2.0 493 OM (1.0 OM residing at the surface and 1.0 OM in the mantle) stored above the 660 km discontinuity 494 ensures the existence of an ocean at the surface of the planet regardless of its age (i.e. its mantle 495 temperature). During the Archean, the mantle was hotter and thus its water storage capacity was lower, 496 ~0.6 OM, implying a volume of Archean oceans ~1.3 times that of today's. This agrees with inferences, 497 based on the ⁸⁷Sr/⁸⁶Sr isotopic composition of marine carbonates, that there was little (less than 4%) 498 emerged land 2.5 Ga ago (Flament et al., 2008). With cooling to the present-day potential temperature 499 of 1650 K, the hydrosphere would have been entirely buried into the solid mantle if the total terrestrial 500 water content had been less than ~ 1.0 OM (Fig. 8). In the future, the water storage capacity of the upper 501 mantle is expected to further increase with secular cooling, possibly reaching a maximum of ~ 1.6 OM 502 for a mantle potential temperature of 1400 K. This would leave only 0.4 OM at the Earth's surface. 503 Therefore, variations of the total amount of water in the mantle of a planet with the size of the Earth

- 504 may easily result in no emerged land early in the history, or the absence of a surface ocean a few Ga after the planet's formation, just because of the interplay with the mantle.
- 506

507 Conclusion

508 In this article, we develop the "mantle rain" model and its implications. The model is based on the 509 mechanism of dehydration melting taking place above the 410 km discontinuity and producing a water-510 saturated melt and solid residue, which together form the LVL observed by seismology. Starting from 511 the well-accepted concept of a neutral melt-mantle buoyancy at the upper limit of the LVL, we develop 512 the idea that the LVL is a source for upwelling melts. With such mechanism, it becomes possible to 513 balance the incoming (by subduction) and outgoing (by rise of LVL-melt) water flows for the MTZ. 514 Within the upper mantle, a water-bearing melt is expected to travel several orders of magnitude faster 515 than the mantle residue stirred by convection movements. The melt could travel relatively fast laterally 516 and vertically, favoring homogenization of the water content, mantle refertilisation, seismic anomalies 517 such as the LVZ, intraplate volcanism, etc.

518 The occurrence of dehydration melting in the LVL also offers a clue to estimate the average water 519 content in the MTZ. It should be higher than the maximum water-storage capacity of the OBM at 410 520 km depth, 1700(200) ppm, as determined experimentally, and should also account for the continuous 521 production of the water-bearing LVL-melt: thus, it should be ~2200(300) ppm H₂O for Tp of 1650 K. 522 Based on this value and with the idea that the OBM should be almost saturated in water, we find that ~1 523 ocean mass of water is stored in the entire upper mantle today. The same calculation can be performed 524 for mantle potential temperatures higher (in the past) and lower (in the future) than that of 1600-1650 K 525 prevailing today. With an increasing water storage capacity of the mantle as a function of decreasing 526 mantle temperature, the mass of the oceans at the Earth's surface is found to decrease by 20% every 527 billion years, when considering a rate of mantle cooling of ~100 K/Ga.

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529 Supplementary Material:

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531 Water fugacity in the upper mantle

We calculate the water fugacity (f_{H2O}) of pure water based on the previous equation of state (EoS) of H₂O (Pitzer and Sterner, 1994). However, geological fluids/melts are not pure water, at high pressures above ~3 GPa, as they dissolve a significant amount of silicate at high mantle temperatures (Massuyeau et al., 2015; Mibe et al., 2007). It makes the f_{H2O} significantly lower than that of pure H₂O. To take this effect into account, we apply a correction to the f_{H2O} of pure water based on a previous experimental report of the water content of olivine up to 12 GPa and 1673 K (Suppl. Fig. 1) (Bali et al., 2008):

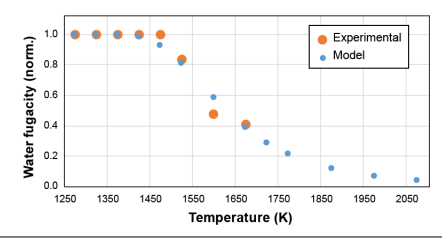
$$f_{H2O} = f_{H2O}^{\text{Pure-water}} * [1 - A_1 \exp(-\Delta G_1 / RT)]$$
(Suppl-Eq. 1)

with ΔG_1 related to the energy of dissolution of the silicate component in the fluid. ΔG_1 can vary with several parameters, in particular temperature and fluid composition. In order to reproduce the ratio of real water fugacity over that of the pure fluid reported previously (Suppl. Fig. 1; (Bali et al., 2008)), we adopted a ΔG_1 value that decreases exponentially with increasing temperature:

 $\Delta G_1 = A_2 \exp(-\Delta G_2/RT)$

(Suppl-Eq. 2)

with A₁=1, A₂=2.50 10⁻² and Δ G₂=-140 kJ/mol. For convenience, we use this formalism for further modeling of *f*_{*H2O*} in supercritical fluids or hydrous melts at upper mantle pressures and temperatures.



Suppl. Fig. 1: Temperature dependence of water fugacity (i.e. the correction to apply to the f_{H20} of pure water). Experimental data obtained between 3 and 12 GPa (Bali et al., 2008) are fitted using a double exponential function (Suppl-Eq. 1 and 2). The model reproduces well the amplitude of the f_{H20} decrease induced by a change of the melt composition (from water-rich to silicate-rich) with increasing the temperature above ~1500 K.

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 - 560 561

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