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1 **Chlorine in wadsleyite and ringwoodite: an experimental study**

2 **Mathilde Roberge¹, H el ene Bureau¹, Nathalie Bolfan-Casanova², Caroline Raepsaet³,**

3 **Suzy Surble³, Hicham Khodja³, Anne-Line Auzende⁴, Patrick Cordier⁵, Guillaume**

4 **Fiquet¹.**

5
6 *¹Institut de Min eralogie, de Physique des Mat eriaux et de Cosmochimie (IMPMC), Sorbonne*
7 *Universit es – UPMC Univ. Paris 06, CNRS UMR 7590, Mus eum National d’Histoire Naturelle, IRD*
8 *UR 206, 4 place Jussieu, 75252 Paris Cedex 05, France*

9 *²Laboratoire Magmas et Volcans, Universit  Blaise Pascal, Clermont-Ferrand, France*

10 *³LEEL, NIMBE, CEA, CNRS, Universit  Paris-Saclay, CEA Saclay, 91191 Gif-sur-Yvette Cedex,*
11 *France*

12 *⁴Universit  Grenoble Alpes, ISTerre, UMR5275 F-38041 Grenoble cedex 9, France*

13 *⁵UMET Lille*

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20
21 **Corresponding author : H el ene Bureau**

22 **e-mail : helene.bureau@imPMC.upmc.fr**

23

Abstract

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We report concentrations of Chlorine (Cl) in synthetic wadsleyite (Wd) and ringwoodite (Rw) in the system $\text{NaCl}-(\text{Mg,Fe})_2\text{SiO}_4$ under hydrous and anhydrous conditions. Multi-anvil press experiments were performed under pressures (14-22 GPa) and temperatures (1100-1400°C) relevant to the transition zone (TZ: 410-670 km depth). Cl and H contents were measured using Particle Induced X-ray Emission (PIXE) and Elastic Recoil Detection Analysis (ERDA) respectively. Results show that Cl content in Rw and Wd is significantly higher than in other nominally anhydrous minerals from the upper mantle (olivine, pyroxene, garnet), with up to 490 ppm Cl in anhydrous Rw, and from 174 to 200 ppm Cl in hydrous Wd and up to 113 ppm Cl in hydrous Rw.

These results put constraints on the Cl budget of the deep Earth. Based on these results, we propose that the TZ may be a major repository for major halogen elements in the mantle, where Cl may be concentrated together with H_2O and F (see Roberge et al., 2015). Assuming a continuous supply by subduction and a water-rich TZ, we use the concentrations measured in Wd (174 ppm Cl) and in Rw (106 ppm Cl) and we obtain a maximum value for the Cl budget for the bulk silicate Earth (BSE) of 15.1×10^{22} g Cl, equivalent to 37 ppm Cl. This value is larger than the 17 ppm Cl proposed previously by McDonough and Sun (1995) and evidences that the Cl content of the mantle may be higher than previously thought. Comparison of the present results with the budget calculated for F (Roberge et al., 2015) shows that while both elements abundances are probably underestimated for the bulk silicate Earth, their relative abundances are preserved. The BSE is too rich in F with respect to heavy halogen elements to be compatible with a primordial origin from chondrites CI-like (carbonaceous chondrites CC) material only. We thus propose a combination of two processes to explain these relative abundances: a primordial contribution of different chondritic-like materials, including EC-like (enstatite chondrites), possibly followed by a distinct fractionation of F during the Earth differentiation due to its lithophile behaviour compared to Cl, Br and I.

54 **1. Introduction**

55

56 Halogen elements (fluorine F, chlorine Cl, bromine Br, iodine I) are minor volatiles compared
57 to hydrogen and carbon. Major halogens F and Cl have been mostly studied for their role in the
58 shallowest Earth's reservoirs: lithosphere, crust, atmosphere, and hydrosphere, mostly because
59 they are the most abundant halogens and because they have been shown to be strongly involved
60 in volcanic and igneous processes. Indeed, Cl is an important constituent of volcanic fumaroles
61 and plumes, and form individualized fluids such as brines and molten salts. These brines are
62 strongly involved in hydrothermal systems and in ore forming processes (see the reviews after
63 Pyle and Mather 2009; Shinohara 2009; Aiuppa et al., 2009). Cl is particularly used to trace
64 igneous processes and ore-forming processes, and to track magmas from their genesis to their
65 eruption Cl is known to affect magma properties (see Pyle and Mather, 2009), it is significantly
66 degassed from subaerial volcanic activity (e.g. Aiuppa, 2009), and it can impact the stratosphere
67 chemistry. Cl is enriched in sea water, it has been shown that oceanic subduction delivers fluids
68 to the mantle through serpentinites related processes (serpentinization and deserpentinization),
69 that are significantly Cl-rich (e.g. Ito et al., 1983; John et al., 2011; Kendrick et al., 2011),
70 making this last one an important constituent in mantle metasomatism processes, that enriches
71 the sources for arc magmatism (e.g. Tatsumi, 1989, Philippot et al., 1998; Scambelluri et al.,
72 2004). Indeed magmas from subduction zones are among the most enriched in Cl (Ito et al.,
73 1983; Straub and Layne, 2003; Kendrick et al., 2012).

74 During subduction, the interaction between seawater and rocks produces secondary minerals
75 containing significant amounts of Cl (Ito et al., 1983; Pagé et al., 2016). It has thus been
76 proposed that the subduction of oceanic lithospheric material at convergent plate boundaries
77 would drive an annual global flux of $2.9\text{--}22 \times 10^{12}$ g Cl to the Earth's interior (John et al., 2011).
78 It has also been proposed that a part of the subducted Cl would reach high depths in the mantle
79 (>200 km) and would possibly enrich the sources for Ocean Island Basalts (Kendrick et al.,
80 2015; Joachim et al., 2015). Cl is significantly present in ultrahigh pressure metamorphic rocks
81 (Scambelluri et al., 2004; Ottolini and Fèvre, 2008; Pagé et al., 2016), in kimberlites
82 (Kamenetsky et al., 2004), and in inclusions of saline brines in diamonds (Weiss et al., 2014).
83 Recent studies show that Cl is present in olivine, pyroxene, and garnet: the highest contents of
84 Cl are measured in minerals formed by metamorphic dehydration of serpentine, olivines and
85 pyroxenes (up to 400 ppm Cl, Scambelluri et al., 2004; Ottolini and Fèvre, 2008). In natural
86 upper mantle nominally anhydrous minerals, the Cl contents are very low: up to 6.3 ppm in

87 olivine (Beyer et al., 2012). Partitioning experiments performed for natural compositions
88 however demonstrate the capability of upper mantle minerals to be the Cl carriers: up to 148
89 ppm Cl in orthopyroxene, 17 ppm Cl in clinopyroxene, 13 ppm in garnet, 5 ppm in plagioclase,
90 up to 170 ppm in olivine (Dalou et al., 2012). To our knowledge, there is currently no evidence
91 about the presence of Cl in the transition zone and in the lower mantle. Recently we have
92 suggested that a strong link connects the water global cycle and that of fluorine at depth
93 (Crepisson et al., 2014; Roberge et al., 2015). We have proposed that the transition zone can be
94 a major reservoir for fluorine (Roberge et al., 2015). It could be similar for Cl.
95 This study aims at constraining the possible deep storage and cycling of Cl through the
96 determination of Cl potential contents in wadsleyite (Wd) and ringwoodite (Rw), the major
97 minerals of the transition zone (TZ). We use these data to discuss the Cl content of the bulk
98 silicate Earth.

99

100 **2. Materials and methods**

101

102 **2.1. Starting materials an experimental strategy**

103 The starting bulk composition was olivine Fo₉₀ with a slight excess of silica ((Mg+Fe)/Si atomic
104 ratio =1.75), obtained with a mixture of (1) oxide powders of MgO, SiO₂, FeO, or (2) natural
105 Fo₉₀ olivine with SiO₂ powder. 5 wt% of Cl was added to the mixture as crushed NaCl powder.
106 NaCl was chosen as the source of Cl because it is enriched in sea water and in subducted oceanic
107 floor. For experiments under hydrous conditions, 2wt% of water was added as brucite
108 Mg(OH)₂, an amount close to the expected water solubility in wadsleyite and ringwoodite in
109 the transition zone. Samples were synthesized in multi-anvil presses at LMV Clermont-Ferrand
110 (France) and at the Bayerisches Geoinstitut of Bayreuth (Germany) following the procedure
111 described in Frost et al. (2001) and Demouchy et al. (2005). Experiments were performed from
112 14 to 22 GPa and from 1100°C to 1400°C during 0.5 to 9 hours, (Table 1). The samples were
113 enclosed in Re, Pt or Au-Pd capsules. Temperatures were monitored with W3Re/W25Re
114 thermocouples located at the top of the capsules. After the run, the experiments were quenched
115 by switching off the electric power before decompression.

116

117 **2.2. Sample characterization**

118 The mineral assemblages were recovered, embedded in epoxy and mirror polished on one side.
119 They were analyzed with Raman spectroscopy and scanning electron microscopy at first. The

120 textures were investigated with a Zeiss Ultra 55 field emission scanning electron microscope
121 (SEM) equipped with an Energy Dispersive X-Ray spectroscopy (EDX) system. Major
122 elements compositions of wadsleyites and ringwoodites were subsequently measured by
123 Electron Probe Micro Analysis (EPMA) on CAMECA-SX100 at CAMPARIS facility (UPMC,
124 France). For these quantitative analyses, we used an acceleration voltage of 15 kV and a beam
125 current of 10 nA with an 15 μm defocused beam.

126 A thin section of ringwoodite sample was prepared with a Focus Ion Beam (FIB), using a
127 gallium beam with a FEI Strata DB 235 at IEMN (Lille France). A transmission Electron
128 Microscopy (TEM) study was performed on this FIB section at UMET, Lille (France) with a
129 FEI Tecnai G2-20 twin operating at 200 kV.

130 Chlorine and hydrogen contents were measured using ion beam analysis, Particle Induced X-
131 Ray Emission (PIXE) for Cl and Elastic Recoil Detection Analysis (ERDA) for H at the nuclear
132 microprobe of the Laboratoire d'Etude des Eléments Légers (LEEL), CEA Saclay. Cl
133 concentrations of the mineral phases Wd and Rw were obtained using incident beams of 3×3
134 μm^2 mapped on large areas (50×50 to $200 \times 20 \mu\text{m}^2$). The procedure is detailed in (Bureau et al.,
135 2010; 2015). We used different incident beams (H^+ and $^4\text{He}^+$ from 1.7 to 3 MeV) in order to
136 control the depth of investigation in the samples. With mineral phases a few tens of μm in size
137 embedded in a quenched Cl-Si-rich glass (present at grain boundaries), compromise had to be
138 found between energy and detection in order to avoid any chlorine contribution from glassy
139 phase. Indeed the depths investigated by the ion beams depend both on the nature of the incident
140 ion (H^+ , $^4\text{He}^+$) and on the energy of the beam, (*i.e.* the highest the energy is the deepest is the
141 investigation). For a San Carlos olivine, the depth of analysis for a proton beam is 50 μm at 3
142 MeV and 25 μm at 1.7 MeV. For one given energy, this depth is reduced when the incident ion
143 is heavier than H^+ , at 2 MeV the investigated depth of $^4\text{He}^+$ is 6 μm .

144 Hydrogen (*i.e.* water) contents of Rw and Wd were measured using ERDA. We used a 3 MeV
145 $^4\text{He}^+$ beam, following the protocol described in Raepsaet et al. (2008), Bureau et al. (2009) and
146 Withers et al. (2012). Simultaneous Cl analysis were also performed by PIXE during ERDA.
147 We scanned the beam on selected areas of the sample (from 30×30 to $150 \times 100 \mu\text{m}^2$). Durations
148 of analysis were chosen from 1800 to 7200 seconds. ERDA and PIXE were associated to
149 simultaneous Rutherford Backscattering Spectrometry (RBS) measurements used to monitor
150 the cumulated charge delivered to the sample during the acquisition (see Bureau et al., 2009).
151 They also provided information on the matrix chemical composition of the samples. For all ion
152 beam analysis, the software RISMIN (Daudin et al., 2003) was used to process the data by
153 selecting the areas of interest in the chemical maps (see Fig. 1). It was particularly useful for

154 the detection of NaCl-rich glasses (grain boundaries, cracks or surface contamination), which
155 would affect H and Cl contents. Once the areas were selected, ERDA spectra were processed
156 by using SIMNRA (Mayer et al., 1997) and PIXE spectra were processed by using the
157 GUPIXWIN software (Campbell et al., 2000). Analysis were cross-checked against: NIST
158 SRM610 glass (Rocholl et al., 1997), KE12 (pantellerite lava from Kenya, Metrich and
159 Rutherford, 1992), EtC3 (Cl-Br-I-bearing NaAlSi₃O₈ glass, Bureau et al., 2000). The sensitivity
160 (i.e. detection limit) with respect to Cl was of a few tens of ppm (30-40 at maximum) depending
161 on the conditions (beam energy, grain size, size of the selected areas).

162

163

164 **3. Results**

165

166 **3.1. Synthesized samples**

167

168 Ten samples were synthesized (see Table 1). Among these samples, two are volatile-free (no
169 H₂O and no Cl), two are anhydrous (Cl-enriched but no H₂O), and six are Cl-H₂O-enriched.
170 Recovered mineralogical assemblages are described in Table 1. They correspond to the mineral
171 assemblages expected for the pressure and temperature conditions of the transition zone. The
172 minerals are in equilibrium with a saline and silica-rich melts mostly present at grain
173 boundaries. SEM shows that wadsleyite and ringwoodite phases exhibit subhedral or euhedral
174 shapes and various crystal sizes (Fig. 1 and 2). In the runs performed at 14 GPa, we observe
175 enstatite in agreement with literature (Inoue 1994; Inoue et al. 1995; Bolfan-Casanova 2000).
176 For runs performed at 22 GPa, stishovite (~10 μm in size) is found in equilibrium with
177 ringwoodite (50-300 μm in size Fig. 2). For all experiments, crystals are embedded in a Si-Na-
178 Cl-rich interstitial glass at grain boundaries which corresponds to the residual melt. No
179 particular textural differences or abrupt increase in the relative proportion amount of quenched
180 glasses are noted between experiments performed at different temperatures. Optical observation
181 and SEM measurements show that the crystals are pure, inclusion-free ringwoodites and
182 wadsleyites (no visible inclusions). Furthermore, a 100 nm thickness lamella of ringwoodite
183 (sample #S5553) was studied by transmission electron microscopy TEM (Figure 3). No Cl-rich
184 or NaCl-rich micro or nano-inclusions have been observed.

185 EMPA analyses show that the Cl-bearing wadsleyites and ringwoodites exhibit a Mg# ratio
186 ($Mg\# = Mg/(Mg+Fe)$ atomic ratio) ranging from 0.90 to 0.92 (Table 1 and 2), similar to the
187 Earth's mantle mineral assemblage.

188

189 **3.2. Chemical Characterization**

190

191 PIXE and ERDA mapping of large areas of the samples show that there is no chemical zoning
192 with respect to Cl and H in the Rw and Wd crystals. About 5 to 20 % in volume of NaCl-rich
193 glasses are present in grain boundaries (Fig. 1). Enstatite is present at 5-10% volume in all the
194 samples containing wadsleyite and stishovite is present at about 5-15% volume in all the
195 samples containing ringwoodite. A comparison between the different Cl contents obtained from
196 each analytical conditions (3 and 1.7 MeV, H^+ and He^+ beams) shows that detection limits and
197 uncertainties for Cl are dependent on the energy and ion source used (Table 2). The analyses
198 performed with a 3 MeV proton beam are the most sensitive to Cl contents but also to potential
199 contamination (Cl-rich glasses located at grain boundary) because the depth of investigation is
200 of about 50 μm . This depth is reduced to 25 μm for a beam of 1.7 MeV, and to a few μm for a
201 $^4He^+$ beam of 3 MeV, but these two last conditions are much less sensitive. As the Cl contents
202 determined at 3 MeV were systematically higher than the measurements performed at 1.7 MeV
203 with a beam of H^+ or at 2 MeV with a beam of $^4He^+$, we have only considered the measurements
204 performed at 1.7 MeV with a beam of H^+ and at 2 and 3 MeV with a beam of $^4He^+$. The lowest
205 Cl content analysed was of 60 ppm, a value higher than the detection limit. Two samples could
206 not be analysed for Cl due to the very small size of the crystals.

207 Results show that Cl contents are significant, and range from 60 ± 60 to 200 ± 48 ppm Cl for
208 Wd and from 99 ± 12 to 490 ± 33 ppm Cl for Rw. Anhydrous Rw contain more Cl than hydrous
209 Rw, with a maximum of 490 ppm Cl. Unfortunately, we failed analysing Cl in anhydrous Wd
210 because crystals were too small. Cl contents of hydrous Wd are ranging from 60 ± 60 to $200 \pm$
211 48 ppm Cl with corresponding water contents ranging from 917 ± 15 to 1659 ± 14 ppm H_2O .
212 Cl contents of hydrous Rw are ranging from 99 ± 12 to 490 ± 33 ppm Cl for water contents
213 from 932 ± 14 to 4766 ± 13 ppm H_2O .

214 No relationship is observed between the temperature and both Cl and H_2O contents (i.e. OH
215 contents expressed as water equivalent concentrations, see Fig. 4 and 5). A slight decrease of
216 Cl content with pressure may be suggested for hydrous samples. The incorporation of Cl in
217 anhydrous Rw is higher than in hydrous Rw (Fig. 4 and 5). No significant dependency is found
218 between the Cl content and Mg#, FeO, SiO_2 or $(Mg+Fe)/Si$ atomic ratio.

219 The water contents determined in Wd and Rw from this study (from 917 to 4766 ppm wt. H₂O)
220 are about one order of magnitude lower than the previously observed solubility values from the
221 literature (Bolfan-Casanova et al., 2000, Inoue et al., 1995, Demouchy et al., 2005) that reached
222 3 to 2.2 wt. % respectively, at similar temperatures. In regard to the low Cl concentrations, we
223 may exclude a competition between OH⁻ and Cl⁻ to enter in Wd and Rw structures. The
224 significant reduction of OH incorporation cannot be compensated by an exchange between OH⁻
225 and Cl⁻. Similar low water contents have been found in NaF-doped wadsleyite and ringwoodite
226 (Roberge et al., 2015). One sample: #40_Cl, has been synthesized together with sample #40_F
227 from this previous study. This suggests that the same process could lower water content in Rw
228 and Wd for both F and Cl-bearing samples.

229 We propose the sodium salts NaCl as NaF to be responsible of this behaviour (see Roberge et
230 al., 2015). A coupled incorporation mechanism of Na⁺ and H⁺ in both F and Cl-rich Wd and
231 Rw may be responsible of the low water content rather than a competitive effect between
232 halogen elements and hydroxyl groups (Roberge et al., 2015). Stalder et al. (2008) have
233 reported a strong decrease of hydrogen incorporation in pure synthetic enstatite at 2.5 GPa,
234 1150-1400°C, and shown to be a function of the NaCl added in the starting materials. The low
235 concentration of OH in enstatite is explained by a reduction of water activity in the fluid phase
236 (melt or aqueous fluid) due to the presence of dissolved salt. In our experiments, the low water
237 and possibly Cl content of the samples may be due to partitioning reaction between Rw and Wd
238 and the Si-water and NaCl-rich melt (up to 20% in volume) in equilibrium with the minerals
239 during the experiments. If true, the Cl concentrations cannot be considered as solubilities. It
240 would also explain why the water contents are so low (4766 ppm wt% at maximum) compared
241 to the initial amount of water (about 2 wt.%). Additional partitioning experiments involving a
242 Mg-rich peridotitic melt would be necessary to validate this hypothesis. As a first assumption,
243 we will consider that the Cl contents measured in Wd and Rw are corresponding to the
244 maximum contents expected in Wd and Rw in a transition zone where interstitial melt may be
245 present (e.g. Toffelmier et Tyburczy, 2007; Schmandt et al. 2014).

246

247 **3.3 Cl versus F in ringwoodite and wadsleyite**

248

249 A comparison between Cl contents (this study) and comparable F contents in Rw and Wd
250 (Roberge et al., 2015) at conditions relevant to the transition zone is presented in figure 6. All
251 the samples are comparable as we have used alternatively NaF or NaCl salts as starting
252 materials, and the same pressure and temperature conditions. We observe that Cl contents are

253 lower than F contents by a factor of 3 in anhydrous ringwoodites. For hydrous Rw and Wd it is
254 more variable, as Cl contents are ranging from 60 ± 60 to 200 ± 48 ppm when F contents vary
255 from 186 ± 19 to 850 ± 85 ppm. The difference between F and Cl solubility's in such chemical
256 systems may reflect the difference in ionic size between F^- and Cl^- . It is likely F and OH share
257 the same sites in olivine (Ol), wadsleyite and ringwoodite (Crépisson et al., 2014; Roberge et
258 al., 2015), but this seems more unlikely for Cl which is larger than F and OH (Cl^- 181 nm, OH^-
259 140 nm). Fluorine incorporation in Ol, Wd, Rd is shown to be related to water incorporation
260 (Crépisson et al., 2014, Roberge et al., 2015), but this is unclear for Cl. Indeed, similar Cl
261 contents are observed in two samples of Rw with different water contents (Fig. 4). Further
262 studies (i.e. FTIR spectroscopy) would be necessary to investigate into this direction. It is
263 difficult to determine where Cl is incorporated in the crystal structure. The presence of OH
264 defects in wadsleyite (Wd) and ringwoodite (Rw) have been investigated by coupling
265 experimental observations to the modelling of infrared spectra (e.g, Blanchard et al. 2009;
266 Blanchard et al., 2013) and provided insights into the crystal chemistry of OH in these minerals.
267 The incorporation of Cl in Rw and Wd could be indirectly determined by studying the spectral
268 signature of hydroxyl defects in NAMs, as for F-bearing hydrous olivines (Crépisson et al.,
269 2014). As a first hypothesis, we suggest that in Wd and Rw, Cl, F and OH may occupy similar
270 sites in the lattice, possibly mostly in association with Mg vacancies (as shown by Blanchard
271 et al., 2013).

272 To summarize our major results: (1) the concentration of Cl in Wd and Rw is significant and
273 can amount 490 ppm; (2) the concentration of Cl in Wd and Rw is 3 times smaller than
274 compared to the concentration of F in the same phases synthesized in similar conditions; (3)
275 anhydrous Rw are richer in both Cl and F than hydrous Rw; (4) the presence of water is not a
276 pre-requisite to store Cl and F in these minerals. When present, water content is small, Cl and
277 F contents may not be considered as solubility values for Wd and Rw, but would rather reflect
278 partitioning reactions with an interstitial melt.

279

280

281 **4. Discussion**

282

283 **4.1 Cl storage in the mantle Transition Zone's wadsleyite and ringwoodite phases**

284

285 It is now recognized that the TZ is hydrated at least locally as shown by recent studies: (1) the
286 significant water content of a hydrous ringwoodite trapped in a natural diamond exhumed from

287 the transition zone (Pearson et al., 2014), (2) geophysical data (e.g. Huang et al., 2005). We
288 consider hydrous Wd and Rw in the following sections: according to our measurements,
289 ringwoodite contains 60 ± 60 ppm Cl and 917 ± 15 ppm H₂O; wadsleyite can take up to $200 \pm$
290 48 ppm Cl and 1115 ± 14 ppm H₂O. Such significant amounts are not surprising if we compare
291 with the incorporation of fluorine in Wd and Rw (665 to 1045 ppm F, up to 956 ppm H₂O in
292 Wd and 186 to 1235 ppm F, up to 1404 ppm H₂O in Rw, see Roberge et al., 2015). Furthermore,
293 the range of water contents measured in Cl-rich Rw and Wd are consistent with what is
294 predicted for the transition zone i.e. 1000-2000 ppm (Huang et al., 2005) and 2000 ppm – 2
295 wt.% (Bercovici and Karato, 2003).

296 We assume, that no significant amounts of Cl are dissolved in the other minor mineral phases
297 of the TZ. We use an average pyrolitic composition with modal abundances of 60%wt.
298 wadsleyite from 410 to 520 km and 60%wt. ringwoodite from 520 to 660 km with addition of
299 40%wt. of garnet and clinopyroxene from 410 to 660 km (Ringwood, 1975), and with the Cl
300 contents of hydrous Fo₉₀ wadsleyite and ringwoodite (174 ppm and 106 ppm respectively).
301 An upper bound for Cl storage in the transition zone would be of 3×10^{22} g Cl, corresponding to
302 80 ppm Cl. Wd and the Rw could be major carriers of the deep Cl, and should be thus taken
303 into account for the Earth budget of this element.

304 In order to explain a chemical contrast between low water contents in the upper mantle with
305 about 142 ppm water (Saal et al., 2002) and assuming a water-rich transition zone (having 0.2-
306 2% wt% water), Bercovici and Karato (2003) have proposed the transition zone water-filter
307 model. In this model, the downwelling fluxes driven by slabs into the transition zone and deeper
308 lower mantle trigger a passive upwelling flow. At 410 km, this upwelling flow would rise out
309 of the high-water-solubility transition zone with the transition of Wd to olivine together with a
310 water release, as olivine has a lower solubility with respect to water than Wd (Bolfan-Casanova
311 et al., 2000). Such a phase transition would then favour the formation of a melt layer overlying
312 the transition zone, having a density higher than the solid phase. This layer would act as a filter,
313 and would maintain a chemical contrast for trace and volatile elements between a depleted
314 upper mantle and an enriched upper transition zone (Bercovici and Karato, 2003), it would
315 permit a significant storage of Cl, F and H₂O in the transition zone.

316

317

318 **4.2 How much Cl in the Bulk Silicate Earth?**

319

320 Based on the average Cl contents in Mid Ocean Ridge Basalts, McDonough and Sun (1995)
321 have estimated an abundance of 17 ppm Cl for the bulk silicate Earth. We have calculated the
322 maximum Cl storage of the bulk silicate Earth, by adding the Cl contribution for each reservoir.
323 Results are summarized in the figure 7, for Cl (this study) and for F (Roberge et al., 2015).
324 For the shallow reservoirs we assume the Cl crust contribution (ocean, evaporite, brine and
325 crustal rock abundances) to be 5.8×10^{22} g (Sharp et al., 2013); we have used the value of 1 ppm
326 Cl from Saal et al., (2002) to represent the depleted shallow upper mantle. For the transition
327 zone, we assumed a hydrous TZ enriched in Cl, due to a continuous supply by subduction, we
328 obtained a maximum storage of 3×10^{22} g of Cl, corresponding to the average concentration of
329 80 ppm. For the lower mantle we used the value of 21 ppm Cl calculated from Ocean Island
330 Basalts compositions by Kovalenko et al. (2006), assumed to be representative of the primitive
331 mantle. The resulting Bulk silicate Earth content for chlorine would then be of 15×10^{22} g,
332 corresponding to a value of 37 ppm Cl, with a contribution of 20% from the transition zone
333 (Figure 7). No significant change of the F/Cl ratio is observed (0.68, McDonough and Sun,
334 1995, 0.62, this study).

335 This estimate of 37 ppm Cl in the bulk silicate Earth is twice the previous one of 17 ppm
336 (McDonough and Sun, 1995), it shows that the Earth's budget of Cl and F could be largely
337 underestimated.

338

339

340 **4.5 What about the origin of chlorine and fluorine of the Earth?**

341

342 The origin of the volatile elements for the Earth is still highly debated today, but most of the
343 models (e.g. Albarède, 2009; Javoy et al., 2010; Marty et al., 2012) propose that they originated
344 from a cosmochemical reservoir that also sourced the parent bodies of primitive meteorites.
345 The absence of correct condensation temperatures (Lodders, 2003) for both Cl and F makes
346 difficult the calculation of the predicted abundances for Cl and F in an Earth accreted from any
347 chondritic materials. For water, a chondritic origin is preferred because the terrestrial and
348 chondritic D/H ratios are similar (see Marty and Yokochi, 2006 and references therein). Among
349 the primitive meteorites, carbonaceous chondrites CC are the most popular candidates to reflect
350 the Earth's volatile source. Marty (2012) shows that a later contribution of $2 \pm 1\%$ of
351 carbonaceous chondrite from the CI species to a dry proto-Earth would be consistent with the
352 Earth atmophile elements (C, H, N, noble gases) and is in agreement with heavy halogen
353 elements (Cl, Br, I) contents determined by McDonough and Sun (1995). However, according

354 to Marty (2012), a contribution of 17% of Cl would be necessary to explain the F abundance
355 from McDonough and Sun (1995). A greater halogen content such as the one presented here
356 cannot be easily reconciled with an Earth only built by homogeneous accretion of Cl
357 carbonaceous chondrite material. Alternatively, enstatite chondrites (EC) are F-enriched (i.e.
358 $F/Cl = 2.77$ to 1.16 , Rubin and Choi, 2009), with a heavy halogen element content similar to
359 those of Cl (Rubin and Choi, 2009). EC were already proposed to be the building material for
360 the Earth (e.g. Javoy, 1995), based on the isotopic signature of oxygen and nitrogen. However,
361 EC are water-poor, the Earth's Si isotopic compositions cannot be explained from a core-
362 formation scenario from a BSE having a EC's composition, and the Mg/Si ratio of the upper
363 mantle differs too much from the EC's. It is also proposed that the Earth was built from the
364 mixing of a number of chondritic like end-members having different volatile compositions
365 (Marty and Yokochi, 2006), including EC. The mixing of different volatile-rich late veneers is
366 however not sufficient to explain the depletion of Cl (and also Br and I) compared to F of the
367 BSE, which may reflect processes that have occurred during differentiation, after volatile
368 acquisition (primitive or from a late veneer). Fluorine has a lithophile character compared to
369 Cl, Br, I (McDonough and Sun, 1995). F content is greater than Cl content in both Wd and Rw.
370 During the magma ocean crystallization, F could have been stored in mantle minerals whereas
371 the heavier halogen elements may have been partially lost from the magma by devolatilization,
372 a loss possibly due to their hydrophilic affinity (Sharp and Draper, 2013; Bureau et al., 2015).
373 As long as the actual Cl and F budget of the Earth will remain un-determined, the origin of
374 volatile elements of the Earth will remain an open question.

375

376

377 **CONCLUSION**

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379 The experimental determination of the Cl concentration in hydrous wadsleyite and ringwoodite
380 shows that they can be major carriers for Cl in the deep Earth. The transition zone can thus be
381 a deep significant repository for Cl. Cl contents are lower by a factor of 3 than those observed
382 by Roberge et al. (2015) for F. For both Cl and F, the presence of water lowers their
383 concentrations in Rw and Wd, but this storage capacity remains significant. It is thus likely that
384 both halogen elements are stored with water in the Earth's transition zone with a maximum
385 content of 80 ppm Cl.

386 Nominally anhydrous minerals are the main Cl carriers in the mantle at least down to 660 km
387 depth. Although it has been shown that F is strongly associated to water through incorporation

388 mechanisms in the NAMs, Cl incorporation in silicate minerals lattice is not so clear and would
389 deserve to be further studied. Assuming a continuous supply from subduction recycling, we
390 calculate a maximum BSE content for chlorine of 15×10^{22} g Cl, corresponding to 37 ppm Cl is
391 proposed. The corresponding global budget for Cl is twice the value proposed by McDonough
392 and Sun (1995). In the light of these experimental results for both Cl and F, we propose that the
393 modern relative abundances of Cl and F may result from the combination of two processes
394 during the Earth's accretion: a mixing contribution of CC and EC chondritic-like materials
395 (required to generate the F enrichment compared to Cl, Br, I); possibly followed by a
396 fractionation of F from other halogen elements during the Earth's differentiation, whereas most
397 of the F being stored in mantle minerals when part of the heavy halogens would have been
398 preferentially partitioned in the magma ocean, degassed and lost.

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404

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563 **Figure captions**

564

565 **Figure 1:**

566 Sample #S5553 (1100°C and 20 GPa): hydrous Cl-bearing ringwoodite synthesized at 1100°C and 20
567 GPa.

568 A. PIXE spectrum of one Rw crystal Cl and Fe Kalpha and Kbeta rays respectively at 2.62 – 2.81 and
569 6.40 – 7.05 KeV are labelled on the spectrum.

570 B. SEM photography of sample #S5553. Ringwoodite (in light grey), the largest grain sizes are of more
571 than 70 µm; stishovite is not present on this section, a Cl-rich melt is visible in grain boundaries. The
572 white square is corresponding to the total map analysed with PIXE.

573 C. PIXE elemental map for iron.

574 D. PIXE elemental map for Cl.

575 A pure ringwoodite crystal is selected using the combination of the SEM photography with the elemental
576 compositions (e.g. iron C and chlorine D) in order to avoid any chemical contamination from interstitial
577 melt. This crystal is represented by the black shape in the SEM photography, and by the white shape in
578 the chemical maps C and D. This area of interest (one ringwoodite crystal) is used to generate the PIXE
579 spectrum presented in A.

580

581 **Figure 2:**

582 SEM photography of A) S5551 (anhydrous Cl-bearing ringwoodite synthesized at 1100°C and 20 GPa).
583 Rw ringwoodite (in light grey), the grain sizes are of about 15 µm; St stishovite crystals are in dark grey
584 with 6 µm medium size, intergranular glass is dark B) H3697 (hydrous Cl-bearing ringwoodite
585 synthesized at 1400°C and 22 GPa), Rw ringwoodite (in light grey), the grain sizes are of about 20 µm;
586 St stishovite crystals are in dark grey, intergranular glass is dark.

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589 **Figure 3**

590 TEM analysis of a ringwoodite thin section of about 100 nm thickness (prepared by focused ion beam)
591 for transmission electron microscopy study, from sample #S5553 (20 GPa, 1100°C, 240 min.)
592 containing 99 ppm wt. Cl and 4766 ppm wt. H₂O.

593 A: Scanning Transmission Electron Microscopy (STEM) bright field micrograph of the ringwoodite
594 showing the only heterogeneities found in the sample.

595 B and C: EDX analysis of Na and Cl contents within the green rectangle in A (1800 x 1100 nm²), they
596 show that heterogeneities are not related to any Cl-rich or NaCl-rich inclusions, and emphasize the
597 uniform distribution.

598

599 **Figure 4:**

600 Cl content in ppm wt. versus temperature in °C. No significant effect of the temperature on the Cl content
601 is seen.

602 Legend: white squares: hydrous wadsleyite, white triangles: hydrous ringwoodite. black triangle:
603 anhydrous ringwoodite.

604

605 **Figure 5:**

606 Cl content in ppm wt. versus water contents in ppm wt. The highest Cl concentration is corresponding
607 to anhydrous ringwoodite. Cl content seems to slightly decrease with water content, but we notice that
608 for a same Cl content of about 100 ppm wt., the water content is ranging from 932 to 4766 ppm in
609 ringwoodite. It would mean that the Cl content is not affected by water for such low amounts.

610 Same legend than Fig 4.

611

612 **Figure 6:**

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614 Cl and F concentrations measured in Wd and Rw, synthesized using the same experimental protocol and
615 from NaCl and NaF sources. Data from this study and from Roberge et al. (2015). Assuming a hydrous
616 transition zone, ranges of potential Cl and F concentrations in Ol, Wd, Rw are proposed.

617 Legend: white squares: hydrous wadsleyite, white triangles: hydrous ringwoodite, white circles:
618 olivines, black triangles: anhydrous ringwoodite, black square: anhydrous wadsleyite.

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621 **Figure 7**

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623 State of the art of the Cl and F budgets of the Earth, assuming a continuous supply by subduction of the
624 TZ with respect to Cl and F. Reservoirs: crust (Rudnick and Gao, 2003); upper mantle without TZ (Saal,
625 2002); TZ, Cl this study, F (Rogerge et al., 2015); lower mantle, Cl (Beyer et al., 2012), F (Kovalenko
626 et al., 2006). Degassing Fluxes are expressed in HCl ($4.3 \cdot 10^{12}$ g/yr) and HF ($0.5 \cdot 10^{12}$ g/yr), they are
627 corresponding to $4.18 \cdot 10^{12}$ g/yr Cl and $0.47 \cdot 10^{12}$ g/yr F respectively, data are from Pyle and Mather
628 (2009); subducted fluxes are from Straub and Layne (2003) and John et al. (2011).

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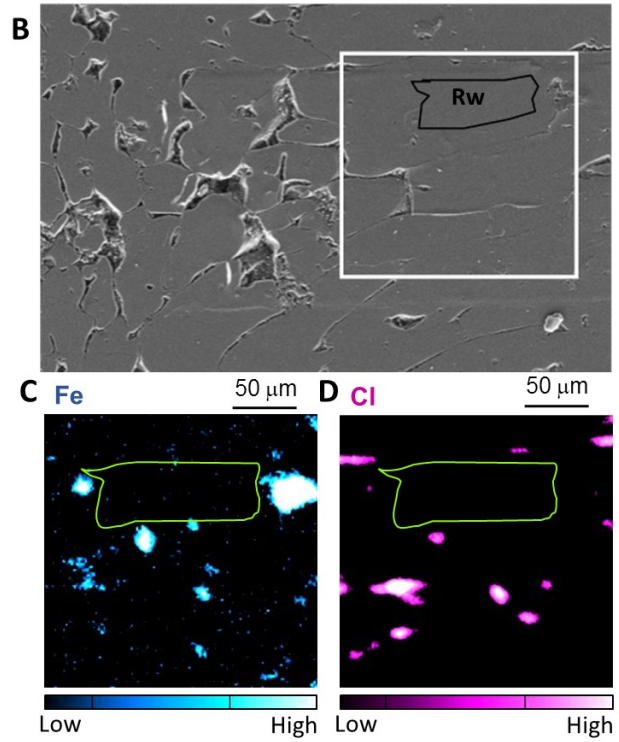
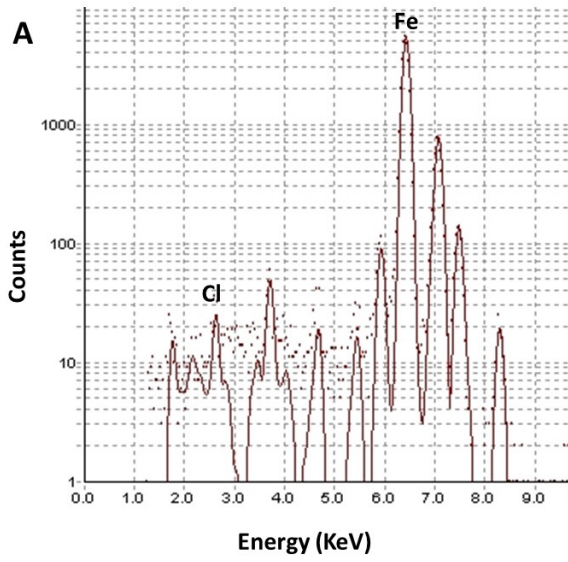
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633 **Figure 1:**

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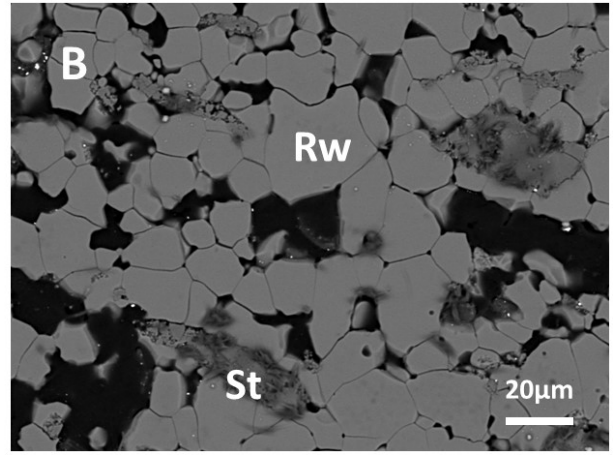
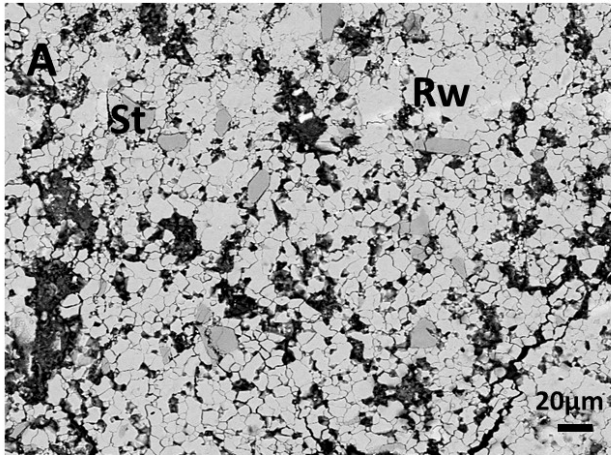
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639 **Figure 2:**

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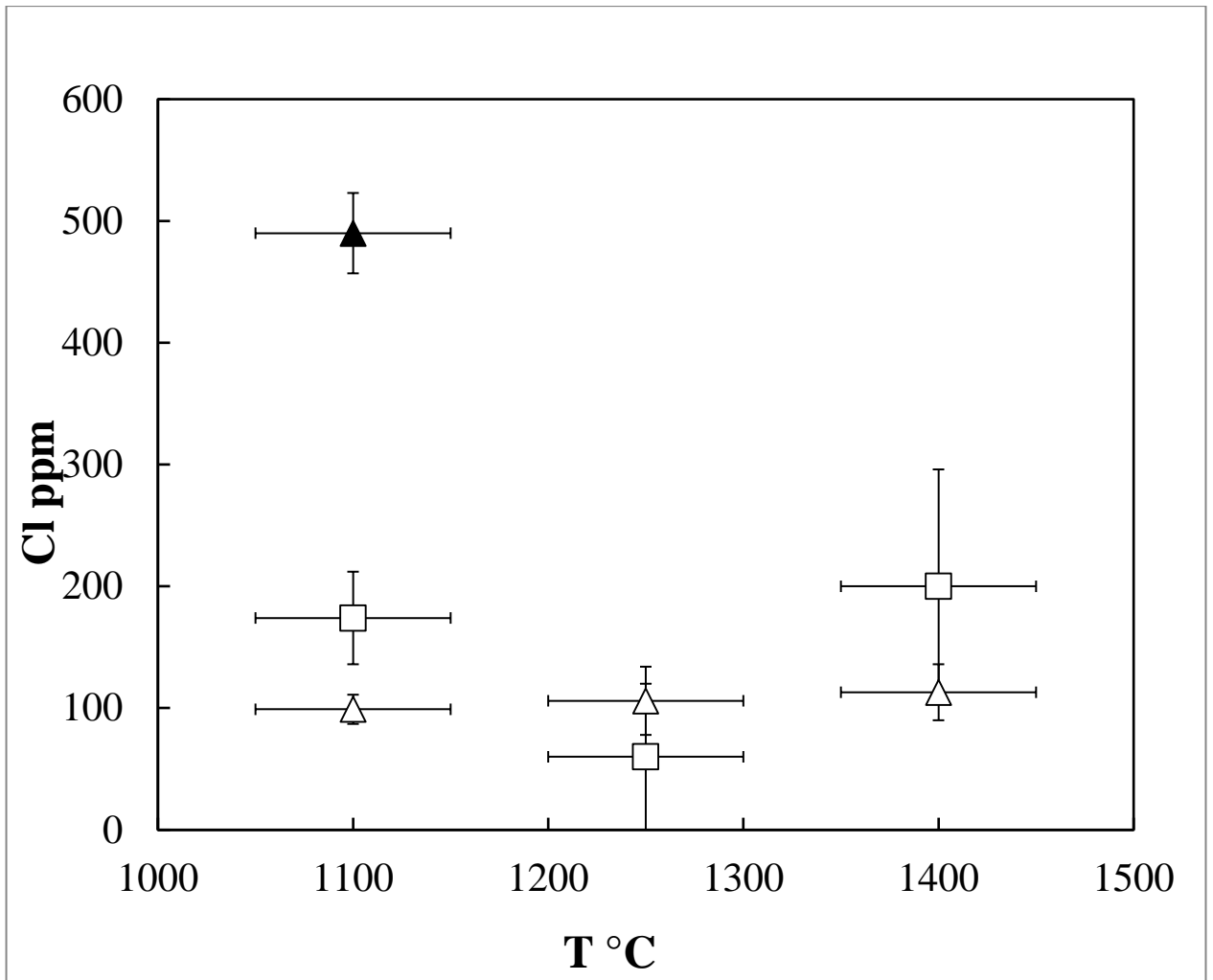
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654 **Figure 3:**
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660 **Figure 4:**

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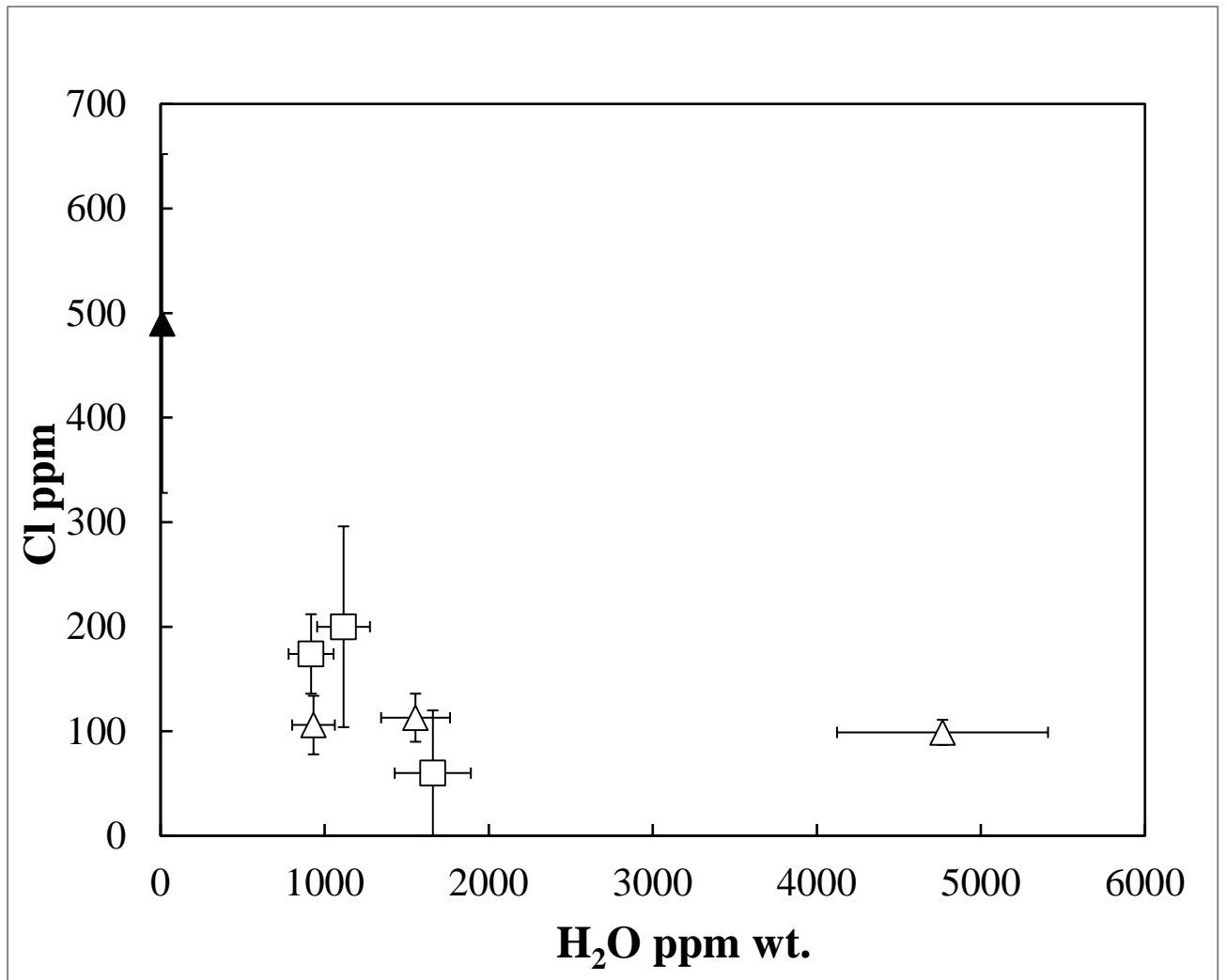


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674 **Figure 5:**

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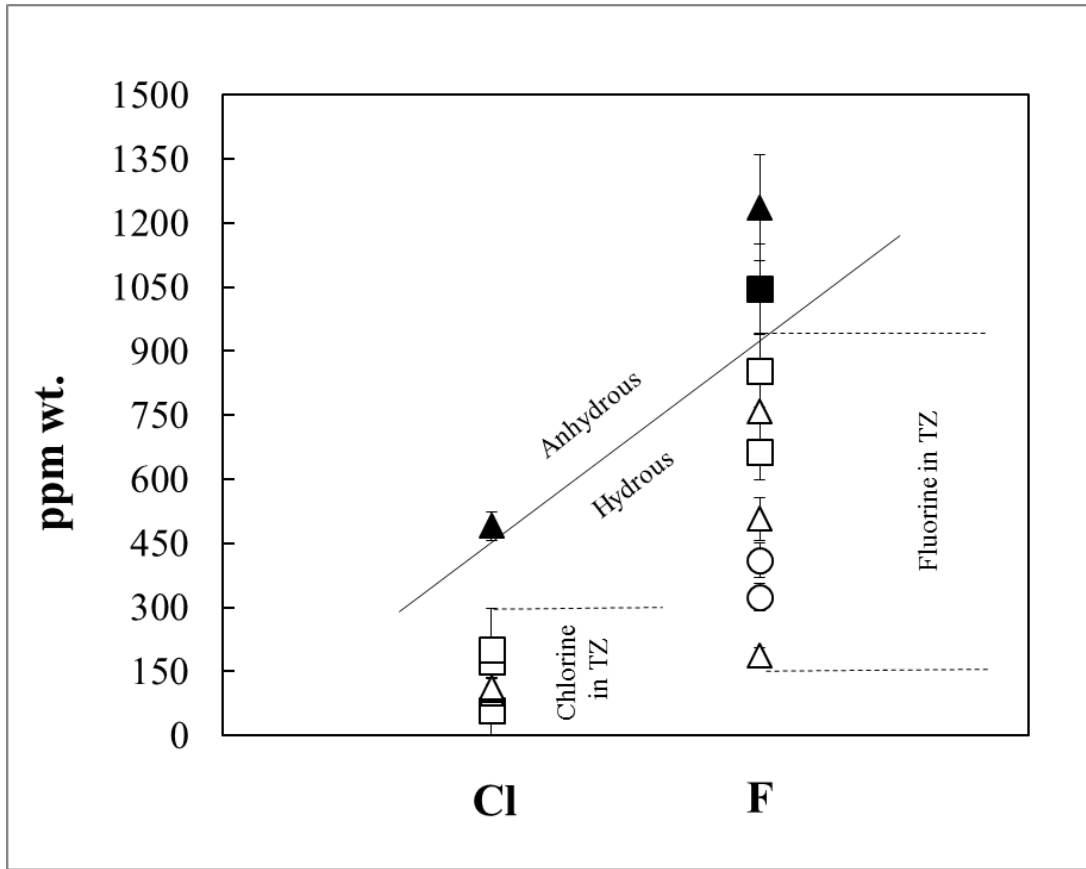
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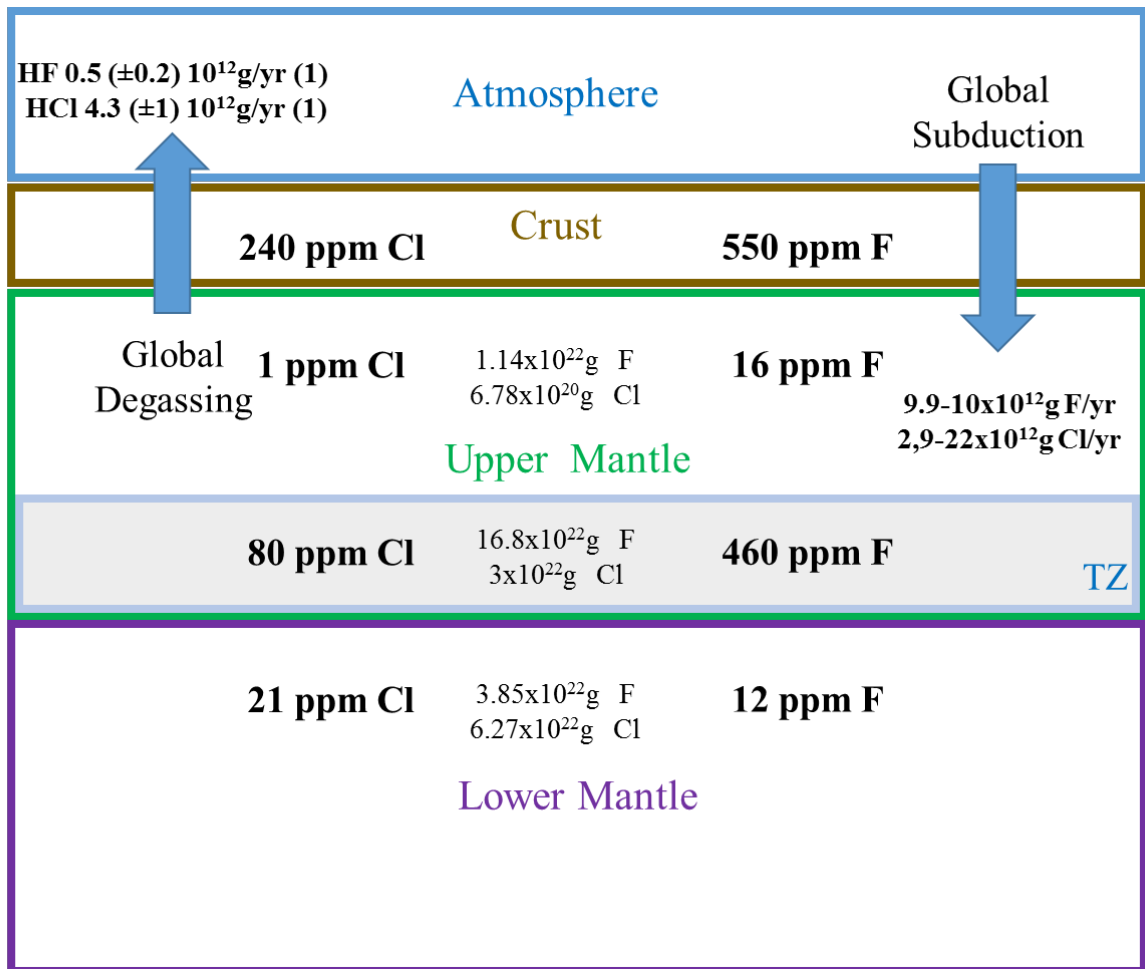
678 **Figure 6:**

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 706 **Figure 7**
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Table 1: Starting materials, experimental conditions and sample descriptions.

Sample	P (GPa)	T (°C)	Time (min)	Starting material	Phases
<i>References</i>					
83(*)	15	1400	20	Pure SC	Wadsleyite (>400 μm)
H3698(*)	22	1400	20	Pure SC	Ringwoodite (>400 μm)
<i>Anhydrous</i>					
87_Cl(*)	14	1350	420	SCP + 5wt.% NaCl + 2 wt.% Mg(OH) ₂	Wadsleyite (20 μm) Enstatite NaCl-glass
S5551(**)	20	1100	240	SCP + 5wt.% NaCl + 2 wt.% Mg(OH) ₂	Ringwoodite(40 μm) Stishovite (10 μm) NaCl-glass
<i>Hydrous</i>					
40_Cl(*)	14	1100	240	SP + 5wt.% NaCl + 2 wt.% Mg(OH) ₂	Wadsleyite (40 μm) Enstatite NaCl-glass
86(*)	15	1250	540	SCP + 5wt.% NaCl + 2 wt.% Mg(OH) ₂	Wadsleyite (30 μm) Enstatite (10 μm) NaCl-glass
88_Cl(*)	14	1400	360	SCP + 5wt.% NaCl + 2 wt.% Mg(OH) ₂	Wadsleyite (70 μm) Enstatite (10 μm) NaCl-glass
S5553(***)	20	1100	240	SCP + 5wt.% NaCl + 2 wt.% Mg(OH) ₂	Ringwoodite (>200 μm) stishovite(15 μm) NaCl-glass
H3694(*)	22	1250	240	SCP + 5wt.% NaCl + 2 wt.% Mg(OH) ₂	Ringwoodite (100 μm) Stishovite (20 μm) NaCl-glass
H3697(*)	22	1400	240	SCP + 5wt.% NaCl + 2 wt.% Mg(OH) ₂	Ringwoodite (> 300μm) Stishovite (20 μm) NaCl-glass

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SC San Carlos olivine, SCP San Carlos olivine powder + SiO₂; SP synthetic powder of SiO₂ + MgO + FeO; (*) Au-Pd capsule, (**) Re capsule, (***) Pt capsule, NaCl pure salt, Mg(OH)₂ brucite, Larger grain sizes are given in brackets when measured.

724

725 **Table 2:** Representative analyses of wadsleyites and ringwoodites. Major elements are from EPMA, Cl from PIXE, H₂O from ERDA.

726 Wd= wadsleyite, Rw =ringwoodite, Nb =number of EPMA analysis, Mg#=Mg/(Fe+Mg) ratio atomic, (Fe+Mg)/SiO₂ atomic ratio. Error in brackets is the standard deviation
 727 for EPMA analysis and is in (%) for PIXE and ERDA.

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Sample	P ± 1GPa	T ±50°C	Time min	Phase	Oxydes (%wt)									Nb	Mg#	(Mg+Fe)/Si	(ppm)	
					SiO ₂	FeO	MnO	Al ₂ O ₃	Na ₂ O	MgO	CaO	TiO ₂	Total				H ₂ O	Cl
<i>Reference</i>																		
83	15	1400	20	Wd	41.55 (0.09)	9.21 (0.10)	0.12 (0.04)	0.11 (0.02)	0.02 (0.02)	48.48 (0.29)	0.02 (0.03)	0.01 (0.01)	99.63 (0.38)	9	0.90	1.92		
H3698	22	1400	20	Rw	41.50 (0.31)	9.70 (0.14)	0.12 (0.05)	0.04 (0.04)	0.01 (0.01)	48.79 (0.27)	0.07 (0.02)	0.01 (0.02)	100.39 (0.38)	14	0.90	1.95		
<i>Anhydrous</i>																		
87_Cl	14	1350	420	Wd	41.38 (0.39)	8.56 (0.16)	0.08 (0.03)	0.07 (0.02)	0.16 (0.03)	48.87 (0.42)	0.03 (0.03)	0.03 (0.02)	99.18 (0.21)	7	0.91	1.93		ND
S5551	20	1100	240	Rw	41.40 (0.86)	9.00 (0.33)	0.14 (0.04)	0.05 (0.03)	0.06 (0.04)	47.84 (0.93)	0.04 (0.01)	0.01 (0.01)	98.54 (0.29)	4	0.90	1.91		490 (33)
<i>Hydrous</i>																		
40_Cl	14	1100	240	Wd	41.13 (0.34)	8.84 (0.20)	0.02 (0.02)	0.24 (0.04)	0.22 (0.05)	49.09 (0.52)	0.02 (0.01)	0.02 (0.03)	99.58 (0.75)	6	0.91	1.96	917 (15)	174 (22)
86	15	1250	540	Wd	41.49 (0.46)	8.57 (0.13)	0.09 (0.03)	0.08 (0.01)	0.19 (0.03)	48.88 (0.34)	0.00 (0.00)	0.02 (0.02)	99.32 (0.47)	6	0.91	1.93	1659 (14)	60 (60)
88_Cl	14	1400	360	Wd	41.55 (0.20)	8.04 (0.20)	0.05 (0.03)	0.09 (0.02)	0.19 (0.02)	49.65 (0.24)	0.01 (0.01)	0.02 (0.02)	99.6 (0.35)	6	0.92	1.94	1115 (14)	200 (48)
S5553	20	1100	240	Rw	41.63 (0.22)	7.63 (0.13)	0.07 (0.03)	0.03 (0.02)	0.04 (0.01)	49.42 (0.04)	0.00 (0.00)	0.02 (0.02)	98.84 (0.42)	3	0.92	1.92	4766 (13)	99 (12)
H3694	22	1250	240	Rw	40.49 (0.55)	7.67 (0.07)	0.04 (0.04)	0.07 (0.00)	0.09 (0.04)	49.97 (0.42)	0.00 (0.01)	0.03 (0.02)	98.36 (0.30)	3	0.92	2.00	932 (14)	106 (26)
H3697	22	1400	240	Rw	41.37 (0.37)	8.18 (0.34)	0.07 (0.02)	0.06 (0.02)	0.10 (0.05)	49.57 (0.35)	0.01 (0.01)	0.02 (0.02)	99.38 (0.29)	7	0.92	1.95	1554 (13)	113 (20)

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