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1 **Temperature-dependent mechanisms of the atmospheric alteration** 2 **of a mixed-alkali lime silicate glass**

3
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11

12 **Abstract**

13 The atmospheric alteration of a mixed-alkali lime silicate glass is described as a function of time
14 and temperature. The glass hydration kinetics self-accelerates with time and is considerably
15 enhanced with the temperature. At 80°C, alkali and alkaline earth ions are retained in hydrate-
16 type environments while at 40°C, they are partially leached out forming surface carbonates, in the
17 order $\text{Ca}^{2+} > \text{Na}^{+} > \text{K}^{+}$. The temperature dependency of hydrolysis processes, the important role
18 of the hydration energy of cations and the diminished solvation properties of bound water are
19 considered to discuss experimental observations and advance in the understanding of the
20 specificities of glass atmospheric alteration.

21

22 **1. INTRODUCTION**

23 Glass is a unique material made by melting silica (from sand) with a flux: an alkali and/or
24 alkaline-earth bearing agent that will lower the melt temperature and viscosity. Because of their
25 relatively alkali-rich composition, glasses are more prone to environmental degradation than
26 other oxide-based materials such as traditional ceramics. The degradation always involves the
27 reaction of glass with water, but the physical, chemical and mass transport conditions determine
28 the extent and result of the process. In this paper, atmospheric alteration refers to the
29 degradation of glass by water in the atmosphere (*ie* when relative humidity, RH, is <100%). In the
30 field of Cultural Heritage, this condition excludes, for instance, the water condensation and rain
31 water running off experienced by stained glass windows [1,2] or the saturated environment of
32 buried archaeological objects [3]. Specifically, it is the condition of glasses conserved in
33 atmosphere in the museums, with their diverse nature (glass objects, glazes and enamels).
34 Although this condition is commonly considered as mild, it can give rise to significant damage

35 depending on the glass composition and climatic conditions (temperature, relative humidity,
36 atmospheric pollutants) [4–8] and it is an issue for curators to be able to identify stable from
37 unstable compositions and adjust the environmental parameters to protect the more alterable
38 pieces [5,9–13]. Atmospheric alteration of glass causes short-term issues also in the glass industry,
39 every time that glass is stored in confined in wet conditions ($RH > 70\%$) [14,15]. The
40 consequences of the surface defects induced by the alteration are the reduction of mechanical
41 properties of glass containers or the degradation of the quality of the thin coatings deposited on
42 float glass [16,17]. Recently, glass atmospheric alteration has become an important matter of
43 concern in the assessment of the long-term safety of geological repository of nuclear waste
44 glasses [18–20]. Indeed, it is estimated that the nuclear glass will be exposed to undersaturated
45 conditions for thousands of years [20].

46 The manifestations of the atmospheric alteration of glass are very specific, and after several
47 centuries of their description, specific and well-defined terms have emerged and been accepted in
48 both the scientific and conservation communities, the most characteristic one being “crizzling”
49 [4]. In particular, Brill [7,10] and Koob [5,11] have identified and accurately described several
50 successive stages of this deterioration that can overlap and be more or less rapid according to
51 both the glass composition and the storage conditions : (i) In an initial stage, alkali salts form on
52 the surface and give to the glass a hazy or cloudy appearance. (ii) In the second stage called
53 incipient crizzling, microcracks of the glass surface appear but they are visible under a
54 microscope only. (iii) In the third stage called full-blown crizzling, the cracking of the glass has
55 grown deeper below the surface and a uniform network of cracks becomes visible to the naked
56 eye. (iv) At the advanced stage of crizzling, flakes fall down from the deeply cracked surface.
57 Eventually, the whole glass piece may desaggregate. Note that the appearance of alkali and calcium
58 salts (carbonates or sulfates) forming a hazy veil on the glass surface, is also observed as an initial
59 stage of the atmospheric degradation of industrial soda-lime glasses [14,16,21,22]. Surface
60 cracking is even reported in the most severe conditions ($RH = 70\%$ and 80%) [22].

61 These manifestations correspond to the ambient temperature phenomena ($0 < T < 50^{\circ}\text{C}$). The
62 general mechanisms are relatively well understood [5,21,23,24]: under a wet atmosphere ($RH >$
63 $60\text{-}70\%$), alkali are leached out of the glass surface and precipitate with acid gases in the adsorbed
64 water film. The alkali-depleted and hydrated leached layer, well observed in numerous studies,
65 has different molar volume and thermal expansion coefficient than the bulk glass, causing its
66 cracking under the effect of cycling temperature and RH, after the layer thickness has reached
67 some threshold. By water diffusion through the leached layer or in the cracks, the alteration
68 progresses and the hydrated layer grows.

69 Beyond this general picture, numerous questions arise regarding the detailed mechanisms of
70 atmospheric degradation. There is no accurate understanding of the rate-controlling mechanisms,
71 nor of the effect of glass composition and structure. What is exactly the very first step, producing
72 the leached hydrated layer? How does the glass composition influence this step? What is the
73 chemical and structural nature of this alteration layer? What is the rate-controlling mechanism of
74 water diffusion through the layer and how does it depend on composition, RH and temperature?
75 In recent years, the combined effects of RH, environment and glass composition on the very first
76 stage of glass alteration have been studied using SIMS as high resolution chemical imaging and
77 depth profiling technique [23–26]. These studies have highlighted the predominant role of RH in
78 the glass surface alteration and the early mobilization of alkalis towards the surface, with a
79 tremendous effect of the surface composition and structure on the rate of this process.

80 This study is an attempt to push forward our understanding of the mechanisms of the
81 atmospheric alteration, which occur on a particular glass composition representative of a family
82 of unstable historical glasses [27]. This family is characterized by, approximately, equal weight
83 amounts of Na₂O and K₂O (~10% each), a moderate amount of CaO and MgO (~6%) and of
84 Al₂O₃ (~1%). Deciphering the mechanisms requires to measure the kinetics of the alteration
85 process and study the changes occurring from the macroscopic to the microscopic scale. In an
86 earlier paper [28] we reported the alteration study of this glass at 85 %RH and at 80°C, a high
87 temperature chosen to accelerate the process. Surprising results were found, notably we did not
88 observe the formation of a leached layer associated to alkali salts deposits on the surface. The
89 glass surface hydration was extremely rapid. In the present paper, we describe the atmospheric
90 alteration of the same glass at lower temperature (40°C to 60°C) and 85 %RH. The phenomena
91 on this temperature range were markedly different, and corresponded well to the macroscopic
92 observations reported in the literature, notably the formation of Na- and Ca-rich salts on the
93 glass surface and the crizzling [4,5,23–26]. From the time and temperature dependency of the
94 phenomena, and the precise compositional and structural analysis of the hydrated layers, we
95 propose mechanisms on both temperature ranges and discuss the possible origin of the marked
96 temperature effect.

97

98 2. METHODS

99 2.1. Sample preparation

100 Experiments were performed on one glass composition: a mixed-alkali lime silicate glass, named
101 glass A ((wt%) SiO₂ 71%, Na₂O 11%, K₂O 11%, CaO 5%, MgO 1%, Al₂O₃ 1%), representative
102 of several chemically unstable productions of the Middle Ages [27]. This glass was elaborated by

103 Saint-Gobain Recherche (France) and the composition was checked by Particle Induced X-ray
104 Emission (PIXE). Samples were prepared in the form of plates ($1 \times 1 \times 0.3 \text{ cm}^3$) polished to $1 \mu\text{m}$
105 with aqueous suspensions of CeO_2 powders. The polished plates were thoroughly cleaned before
106 the ageing tests, using a 1M sodium dodecyl sulfate (SDS) aqueous solution and ethanol. Time of
107 flight-secondary ion mass spectrometry (ToF-SIMS) analysis conducted on the pristine glass
108 plates after these preparation steps showed an outermost surface layer of about 30 nm
109 significantly depleted in Na, Ca and Mg, and slightly depleted in K (Fig. 3).

110 For bulk spectroscopy analysis, calibrated glass powders of $32\text{-}50 \mu\text{m}$, $10\text{-}20 \mu\text{m}$ and $5\text{-}10 \mu\text{m}$ size
111 fractions were also prepared by crushing and sieving the glass. After multiple ultrasonic washing
112 in acetone to remove the fines, their specific surface area σ was measured by the BET method,
113 using a Belsorp Max analyser with krypton as adsorbing gas. The results were $\sigma=1200 \pm 100$
114 cm^2/g , $3900 \pm 300 \text{ cm}^2/\text{g}$ and $7800 \pm 900 \text{ cm}^2/\text{g}$ for the powder fractions of $32\text{-}50$, $10\text{-}20$ and
115 $5\text{-}10 \mu\text{m}$ respectively.

116

117 **2.2. Ageing tests**

118 Two different devices were used in parallel to simultaneously control the relative humidity and
119 the temperature of the ageing atmosphere over various time periods: (i) a climatic chamber and
120 (ii) a hermetic box put into an oven and containing a saturated saline solution to control the RH
121 [29]. All details about these devices are given in [28]. The relative humidity was fixed at 85 RH%
122 in all experiments and the temperature was 40°C , 50°C , 60°C or 80°C [28] for various periods of
123 time from 6 h (at 80°C , [28]) to 9 months (at 40°C). With the hermetic box, the 85%RH value
124 was obtained thanks to a solution saturated in KCl. A MadgeTech sensor was inserted in the box
125 to control temperature and RH during the whole experiments. We took care to avoid any liquid
126 water run-off on the samples during the heating and cooling ramps by maintaining the RH below
127 the dew point at every temperature. All tests were performed in static mode (no cycling). In this
128 paper, they are noted V tests or VXh, VXd or VXm tests, where X is the total duration in hours
129 (h), days (d) or months (m) including the ramps. The good correspondence between the two
130 ageing methods (devices (i) or (ii)) has been checked by comparing the alteration phenomena
131 after a 2 months ageing test at 50°C , 85 RH%, for this glass composition as well as for other
132 compositions. For both ageing methods, the thickness of the alteration layer was the same (10
133 μm for glass A) and the nature and quantities of surface salts were the same (mainly Ca-bearing
134 carbonates for glass A).

135

136 **2.3. Characterization methods**

137 The glass surface and sub-surface modifications after a V test were studied at t_1 (end of the V
138 test) and over a 6 months period after t_1 , because certain aged glass surfaces continued to
139 transform under the laboratory atmosphere [28]. Sample preparation before the characterization
140 was intentionally very limited. Notably, no rinsing nor polishing were performed, because these
141 operations may be very harmful for the alteration layers integrity [28]. When alteration products
142 were scattered on the surface, we preferred keeping them and taking their presence into account
143 in the interpretation of the characterization data.

144

145 Microstructural observation of the glass surface

146 The surface states of the glass plates were characterized before the V tests (t_0) and several times
147 after the V tests (t_1 and beyond) by optical microscopy in direct or grazing light using a Keyence
148 VHX-5000 numerical microscope with a Z500 objective. This characterization was supplemented
149 by observations with a field-emission gun electronic microscope (FEG SEM) JEOL 7800F,
150 preferentially in secondary electron mode at low acceleration tension (2 to 7 kV), after making the
151 glass surface conductive with a platinum deposit about 1 nm thick. The samples were submitted
152 to a vacuum of about 1 Pa (Pt deposit) and 5.10^{-5} Pa (FEG SEM) during this characterization.

153

154 Measurement of the thickness of the alteration layers

155 Alteration layer thicknesses are noted e_a in this paper. The very small ones (some tens of nm to
156 about 200 nm) were determined from the elementary concentration profiles obtained by ToF-
157 SIMS analysis, with a ToF-SIMS V instrument (Ion-ToF company), using a O_2^+ ion beam (2 keV,
158 about 150 nA) for in-depth abrasion, a Bi^+ ion beam (25 keV, 0.3 pA) for the analysis, and an
159 unfocused electron beam of some eV for the charge neutralization. The crater is $200 \mu m \times 200$
160 μm or $300 \mu m \times 300 \mu m$ in size and at its center the analysis zone is $50 \mu m \times 50 \mu m$ or $100 \mu m$
161 $\times 100 \mu m$ respectively. The sputtering rate on aged glass plates was determined thanks to
162 mechanical profilometry of the craters (about 0.3 nm/s into the alteration layer for a current of
163 100 nA). To assess e_a , we considered the distance over which the $^{45}SiOH^+/^{28}Si^+$ ratio is halfway
164 between its near surface concentration (the value measured just after the SiOH surface spike) and
165 its baseline concentration, as a marker of the interface between the alteration layer and the
166 pristine glass, by analogy with a method described in [30] using the $^1H^+/^{28}Si^+$ ratio.

167 For the high enough altered thicknesses (> 200 nm), their evaluation by SEM observations was
168 preferred. When flakes formed on glass plates, their thickness was measured and associated to
169 the alteration layer thickness after we had checked that the flaking indeed occurred near the
170 interface between the alteration layer and the pristine glass. Otherwise, SEM observations were
171 performed on the edge, on a freshly fractured plate, preferentially in backscattered electron mode

172 to visualize the chemical contrast at the interface. These observations were performed in
173 different areas of a same aged glass plate and we report a mean thickness e_a .

174

175 Chemical analysis

176 Elementary compositions of the alteration layers were quantified by Energy Dispersive X-ray
177 spectroscopy (EDX) with the system Bruker QUANTAX 400, at the same time as SEM
178 observations. Analyses were performed either on the surface of the glass plate, outside any defect
179 or crystal, according to the operating conditions and specific quantification method described in
180 [28], or on the edge of the fresh fracture for the alteration layers thick enough. Elementary
181 composition profiles into the alteration layers, from the surface to the alteration layer-pristine
182 glass interface, were obtained by the ToF-SIMS analysis of an area free of salt deposits when
183 possible. Chemical characterization of the alteration products on the surface of glass plates were
184 performed by EDX at low acceleration tension, by X-Ray Diffraction (XRD) with an X'Pert
185 PRO PANalytical instrument using Cu-K α radiation, and by micro-Raman spectroscopy with a
186 Renishaw Invia instrument using a 532 nm solid-state laser as excitation light.

187

188 Hydration rate measurement

189 The amount of water in the alteration layer, or hydration rate, was measured on glass plates by
190 EDX according to the original method described in [28]. This local measurement corresponds to
191 the water content that has not been evaporated under vacuum, i.e. to the tightly bound water. For
192 a comparison, global weight measurements on glass powders by thermogravimetric analysis
193 (TGA) were performed on a NETZCH STA449 F3 Jupiter equipment, considering that the
194 tightly bound water disappears between about 150 and 500 °C [31].

195

196 Structural studies

197 Structural modifications of the glass at the atomic scale were studied by the mean of solid-state
198 Nuclear Magnetic Resonance (NMR) spectroscopy on pristine and bulk altered glass powder. ^1H ,
199 ^{23}Na , ^{27}Al and ^{29}Si magic-angle spinning (MAS) NMR spectra were collected on Bruker Advance I
200 and Advance II spectrometers operating at magnetic fields of 7.02 T (300WB) and 11.72 T
201 (500WB), respectively. For these experiments, about 100 mg of a sample was packed in 4 mm
202 outer diameter ZrO_2 rotor and spun from 10 kHz (7.02 T) to 12.5 kHz (11.72 T) using Bruker
203 CPMAS probes. Recycle delay applied for the complete relaxation of nuclear magnetizations was
204 about 1 s for ^1H , ^{23}Na and ^{27}Al nuclei and from 20 to 200 s for the ^{29}Si nucleus. ^1H spectra were
205 acquired using the Hahn echo pulse sequence (90- T_E -180- T_E -Acq) with a variable rotor-
206 synchronized echo delay T_E . ^1H - ^{29}Si Cross-Polarization MAS experiments (CPMAS $^1\text{H} \rightarrow ^{29}\text{Si}$)

207 were performed with contact times of about 1 ms. Chemical shifts are referenced using a
208 tetrakis(trimethylsilyl)silane sample for ^1H (0.2 ppm relative to liquid tetramethylsilane, TMS) and
209 ^{29}Si (for which the highest intensity peak is situated -9.9 ppm from that of liquid TMS), an
210 aqueous solution of NaCl for ^{23}Na (0 ppm) and an aqueous solution of AlNO_3 for ^{27}Al (0 ppm).
211 In this paper, we employ the Q^n notation to describe the structure of the silicate network at short
212 distance. Q^n represents a SiO_4 unit with n bridging oxygen atoms (NBO).

213

214 3. RESULTS

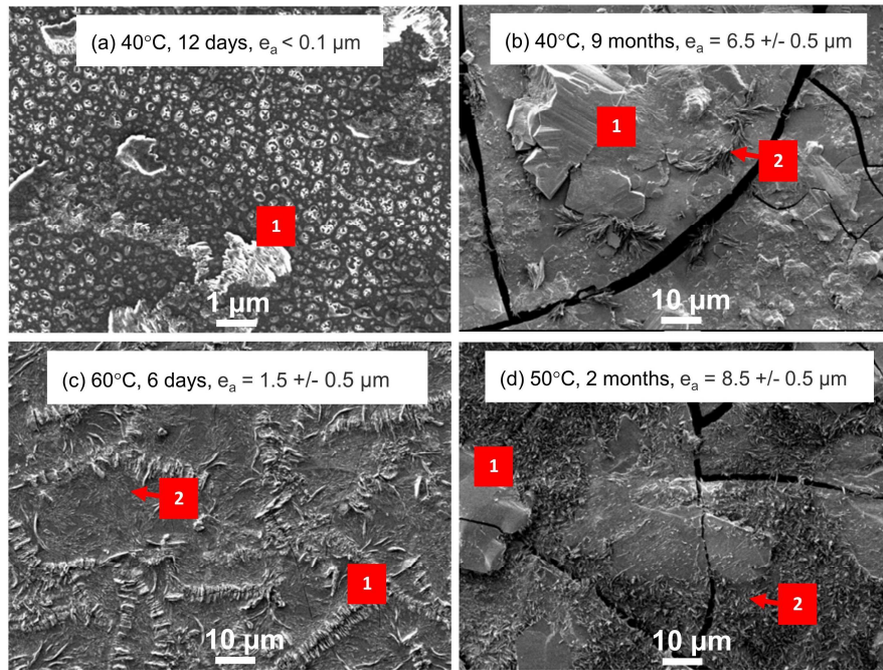
215 Contrary to the phenomena at 80°C , temperature at which no carbonate salts formed on the
216 surface of the glass plates during their ageing in the climatic chamber, in the $40\text{-}60^\circ\text{C}$ temperature
217 range the formation and growing of the carbonate salts did occur during the ageing. After a time
218 period depending on the temperature, the salts totally covered the surface. Below the crust of
219 salts, a hydrated layer partially depleted in alkali and completely depleted in Ca was observed,
220 characterized by its uniform thickness, its structural homogeneity at the SEM scale, and a flat
221 interface to the pristine glass. When the thickness of the layer reached about $1\ \mu\text{m}$, the layer
222 cracked and eventually delaminated at some locations. Except an accelerating effect of the
223 temperature, the macroscopic phenomena were similar at 40 , 50 and 60°C . The temperature of
224 40°C was chosen for an in-depth study of the kinetics and chemical and structural characteristics
225 of the alteration. After a short description of the morphological observations at 40 , 50 and 60°C ,
226 we present the results on the kinetics (as given by the alteration layer thickness as a function of
227 time), and the detailed chemical and structural study of the alteration layer that forms at 40°C , 85
228 RH%. The results obtained at 80°C , 85 RH% are given for comparison. Note that we name
229 “alteration layer” the entire layer composed of the hydrated layer and the crust of salts on its
230 surface.

231

232 3.1. Microstructure of the alteration layers as a function of time and temperature

233 The SEM images of a selection of altered surfaces are shown in Fig. 1, and the edge of fresh
234 fractures, or of delaminated flakes are shown in Fig. 2. Optical microscopy images, more
235 informative about the optical macroscopic appearance of the samples, as well as complementary
236 SEM images are given in the Data in Brief associated to this paper. The sample aged at 40°C for
237 12 days provides the earliest stage of alteration in our series of experiments. Two kinds of salts
238 have appeared at this early stage: micrometric salts with sharp angles and faces and
239 submicrometric salts with no specific shape (Fig. 1a). From SEM-EDX and microRaman
240 analysis, the big microscopic salts are assigned to calcite and labelled with number 1 on the

241 images. The glass surface is still visible below the salts, which appear with a shallow white
242 boundary due to electronic charge effects. At this stage, the alteration layer thickness is less than
243 100 nm as described later. The coverage of the surface by salts is almost complete after 3 months
244 of ageing. At this ageing duration and beyond, scattered Na enriched carbonates with a
245 characteristic needle-like shape sometimes referred to as “crow’s foot”[14] are observed and
246 identified by EDX (Na and C enrichment). These salts are labelled with number 2 in Fig. 1b, c
247 and d. Immediately next to the calcite crystals, other small and shapeless precipitates enriched in
248 Ca, C and O are also observed (Data in Brief Fig. 1d, label 3) and assigned to vaterite. After 9
249 months of ageing at 40°C, the longest experimental period of the study, the salts form a
250 continuous crust on the top of the glass hydrated layer, which is about 6.5 µm thick (Fig. 2b).
251 The calcite is always recognizable as distinctive crystal stacks with large faces (Fig. 1b, label 1), as
252 are the needle-like Na enriched carbonates (Fig. 1b, label 2). Due to the insufficient amount of
253 salts, only the calcite phase was identified by XRD. The microRaman spectrum of a glass powder
254 (6-10 µm) altered for 6 months at 40°C, bears the distinctive peaks of calcite (283, 715 and 1087
255 cm⁻¹) and an additional peak at 1071 cm⁻¹ that could indicate the presence of pirssonite, a mixed
256 soda-lime carbonate of composition Na₂Ca(CO₃)₂·2H₂O (RRUFF database, reference R040146).
257 At 40°C, the cracking of the hydrated layer starts after 3 months of ageing, when the hydrated
258 layer is 1±0.2 µm thick. The cracks cut across the salts (see for instance Fig. 1d, the calcite crystal
259 on the left). It means that they have occurred after the formation of the salts, possibly during the
260 ramp to ambient temperature and RH, because of stresses induced by differential dilatation and
261 loss of water. The delamination, which opens an air gap below the flakes, is well put in evidence
262 by the observation of circular interference patterns by optical microscopy (Newton rings) (Data
263 in Brief Fig. 1b). It begins after 6 months of ageing (hydrated layer thickness is 2.5±0.2 µm).
264 Flakes are about 100 µm in length, a size that seems independent of the temperature and time
265 (compare Fig. 1b and d).



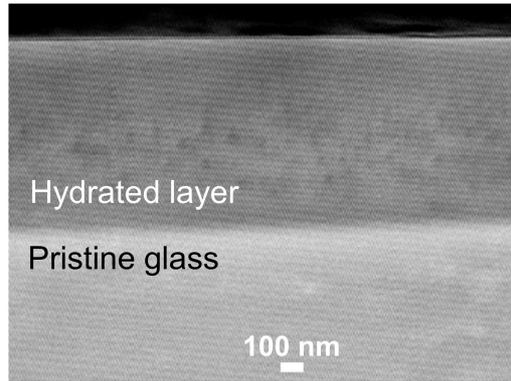
266

267 **Fig. 1** Surface states of glass A plates immediately after an ageing (t_1 time) at 40 °C (a,b), 60 °C
 268 (c) or 50 °C (d), 85 RH%. Secondary electrons SEM images. e_a : alteration layer thickness. Label 1
 269 depicts calcite crystals and label 2 sodium-bearing carbonate crystals.

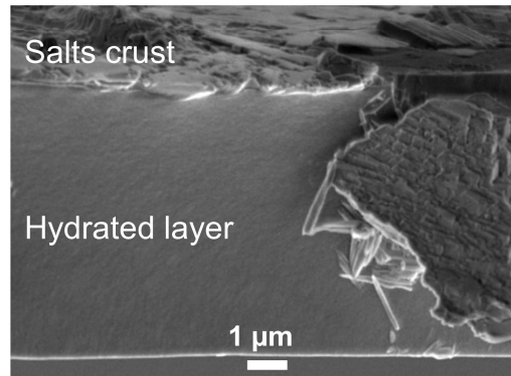
270

271 At 50°C and 60°C, the morphology is comparable but the alteration is more rapid. After 2
 272 months of ageing at 50°C, the alteration layer is composed of an irregular crust of carbonates of
 273 thickness up to about 2 μm, and underneath, of a hydrated layer about 8.5 μm thick (Fig. 1d and
 274 Fig. 2c). At 60°C and 6 days of ageing, the carbonate salts almost cover the surface, the hydrated
 275 layer underneath is already 1.5 μm thick and crossed by cracks (Fig. 1c). Vaterite crystals (CaCO_3)
 276 were identified (DRX, Raman) together with calcite crystals (CaCO_3). The morphology of the
 277 calcite crystals is different than at lower temperature: small calcite crystal plates perpendicular to
 278 the surface are stacked as chains of several 10 μm length. This microstructure characterized by
 279 more numerous and smaller crystals is likely related to a higher nucleating rate of calcite at this
 280 temperature. The visual appearance of this sample was completely opaque and white.

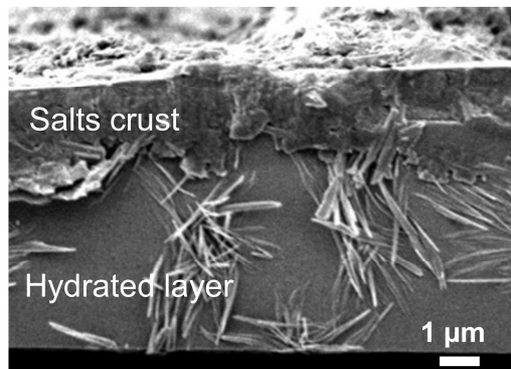
281



(a) 40°C, 3 months, $e_a = 1 \pm 0.2 \mu\text{m}$



(b) 40°C, 9 months, $e_a = 6.5 \pm 0.5 \mu\text{m}$



(c) 50°C, 2 months, $e_a = 8.5 \pm 0.5 \mu\text{m}$

282

283 **Fig. 2** SEM images of the edge of glass A plates after an ageing at 40 °C (a,b) or 50°C (c), 85
 284 RH%. Secondary electron mode (b, c) or backscattered electron mode (a). e_a : alteration layer
 285 thickness.

286

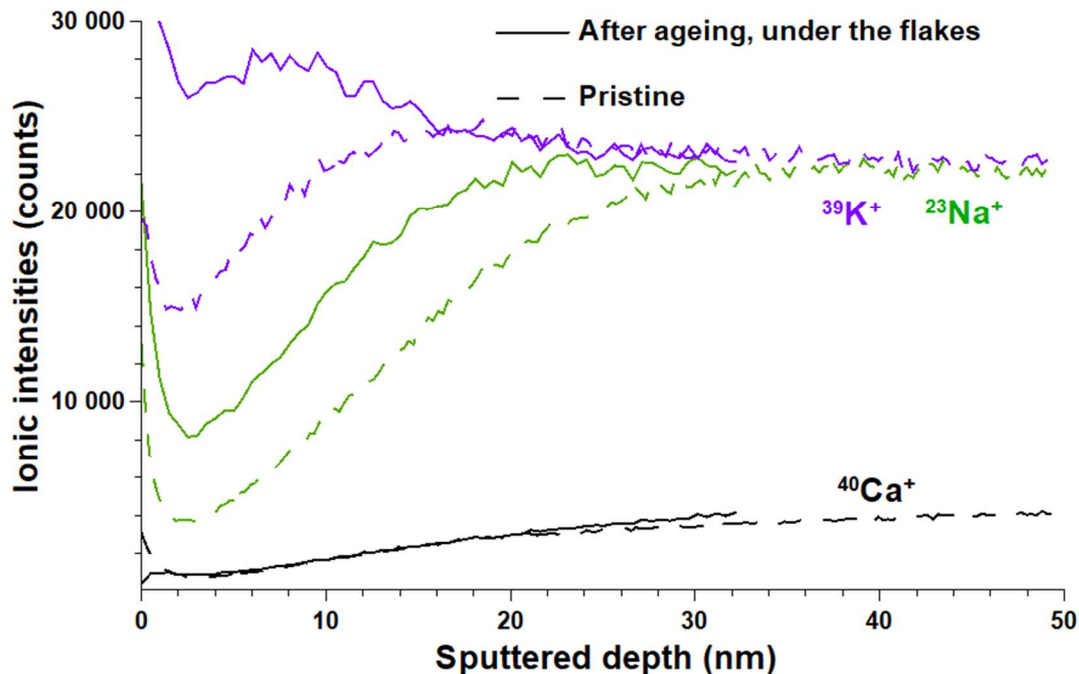
287 3.2. Growing kinetics of the alteration layer

288 As described above, the atmospheric alteration in the 40-60°C temperature range manifests by
 289 the concomitant formation of a hydrated layer and of carbonate salts on the top of it. At 80°C,
 290 only the formation of the hydration layer is observed during the ageing. Carbonate salts form

291 after the end of the ageing test, when samples have cooled down to ambient temperature and RH
292 [28]. Interestingly, calcite is never observed on these 80°C aged samples, but only Na carbonates,
293 among which trona only could be identified (a hydroxycarbonate with composition
294 $\text{Na}_3(\text{HCO}_3)(\text{CO}_3)\cdot 2\text{H}_2\text{O}$) [28]. Whatever the ageing duration and temperature (40, 50, 60 and
295 80°C), the hydrated layer has always the same microscopic characteristics: it has a uniform
296 thickness, delimited by a flat and sharp interface with the pristine glass at the SEM scale (about
297 0.1 μm , Fig. 2a). The dark contrast with the pristine glass is due to partial loss of alkali and lime
298 and hydration (see later), but also probably to a reduced density. The granularity observed at very
299 small scale on the SEM images of the edge is of topological nature (by comparing secondary and
300 back-scattered electrons images). It probably reflects structural inhomogeneity at the nanometer
301 scale. At last, the flatness of the alteration layer-pristine glass interface again demonstrates that
302 the cracking probably occurred during the ramp of temperature and RH at the end of the ageing,
303 otherwise irregularities of the interface around the cracks would be expected.

304 After the layer has reached one micrometer thickness (1 μm), it cracks and eventually forms
305 delaminated flakes. The concentration profile of the glass beneath the detached flakes was
306 measured by ToF-SIMS for the sample altered at 50°C for 2 months. It is compared with the
307 profile of the pristine, polished glass plates in Fig. 3 (only ionic intensities of $^{39}\text{K}^+$, $^{23}\text{Na}^+$ and
308 $^{40}\text{Ca}^+$ are depicted). The surface of the pristine glass is depleted in Na and K over about 30 nm,
309 due to the polishing by CeO_2 powder in water. The surface of the glass below the detached flakes
310 is less depleted in Na in comparison, and not depleted in K. On the contrary, there is a slight K-
311 enrichment over the first 20 nm below the surface. It means that the delamination occurs within
312 20 nm of the hydration front. A local K-enrichment around the hydration front has been
313 reported in another study of glass alteration in silica-saturated solution [32]. From this result, it
314 can be considered that the thickness of the flakes corresponds to the thickness of the hydrated
315 layer. The conclusion was the same at 80°C, because the EDX analysis of the glass below the
316 flakes was identical to that of the pristine glass (Table 4 in [28]).

317



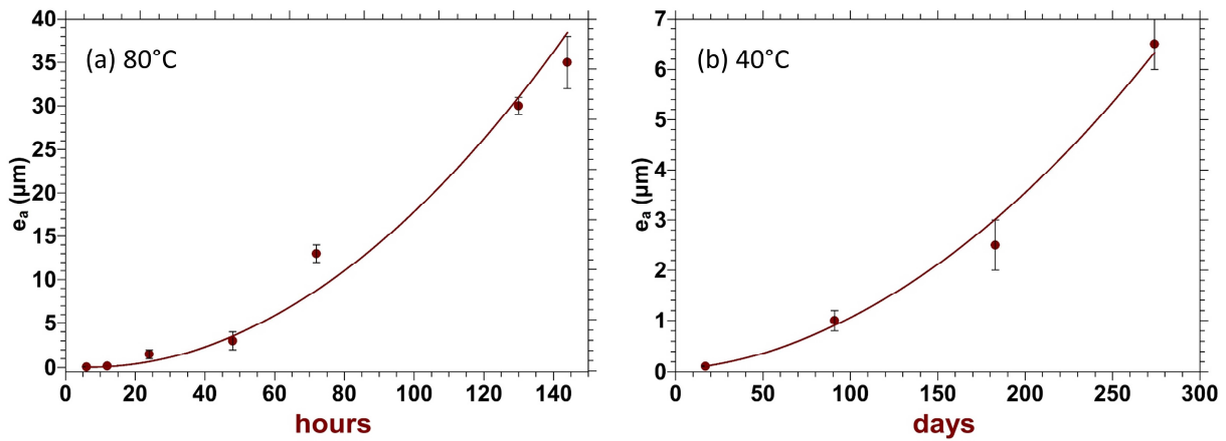
318
 319 **Fig. 3** Ionic intensities in $^{39}\text{K}^+$, $^{23}\text{Na}^+$ and $^{40}\text{Ca}^{2+}$ measured by TOF-SIMS on a pristine glass plate
 320 after polishing and washing (dashed curve) and on the glass surface below the flakes of a glass
 321 plate aged at 50°C and 85 %RH for 2 months (continuous curve).

322
 323 The hydrated layer thickness was then measured by ToF-SIMS profiling at short ageing time (6h,
 324 12h at 80°C and 17 days at 40°C), and by SEM at longer ageing time. Values are given in Table 1
 325 and plotted as a function of time in Fig. 4.

326 In the timescale of the experiments, the growing of the hydration layer tends to accelerate at
 327 80°C and at 40°C. Surprisingly, the hydration kinetics is neither at the square-root of time nor
 328 linear, indicating that it is neither simply diffusion controlled nor surface-controlled. Several
 329 mechanisms interplay to explain this acceleration that will be discussed later. A complex time
 330 dependency has also been found at room temperature for soda-lime silicate glasses, although the
 331 very short alteration depths hinders accurate measurement of the kinetics [26].

332 Moreover, we emphasize that the time and e_a scales are very different in Fig. 4a (80 °C) and Fig.
 333 4b (40 °C). By neglecting the acceleration and assuming a linear rate, the alteration layer thickness
 334 increases by approximately 6 $\mu\text{m}/\text{day}$ at 80 °C and by about 24 nm/day at 40 °C, which would
 335 correspond to a very high apparent activation energy of about 130 kJ/mol. Such a value far
 336 exceeds the activation energy reported for silicate network hydrolysis (about 80-85 kJ.mol⁻¹ for
 337 silicate glasses [33,34]). This result suggests that i) either the rate-controlling mechanisms are
 338 different at 80 °C and at 40 °C, or ii) the rate-controlling mechanisms are the same but their

339 kinetics parameters (activation energy, prefactor) have changed with the temperature.



340
 341 **Fig. 4** Alteration layer thicknesses (e_a) of glass A as a function of the ageing duration (numerical
 342 values are reported in Table 1). The lines are a guide for the eyes. (a) 80 °C, 85 RH% (climatic
 343 oven). (b) 40 °C, 85 RH% (V17d: climatic oven, other ageing durations: hermetic box containing
 344 a saline solution of KCl in H₂O).

345

80 °C							
	6 h	12 h	24 h	48 h	72 h	130 h	144 h
e_a (μm)	0.055 ± 0.01 *	0.07 ± 0.01 *	1.5 ± 0.5	3 ± 1	13 ± 1	30 ± 3	35 ± 3
40 °C							
	17 d	3 m	6 m	9 m			
e_a (μm)	0.11 ± 0.01 *	1 ± 0.2	2.5 ± 0.2	6.5 ± 0.5			

346 **Table 1.** Alteration layer thicknesses (e_a) of glass A developed with various ageing durations at 80
 347 °C (climatic oven) or at 40 °C (V17d: climatic oven, other ageing durations: hermetic box
 348 containing a saline solution of KCl in H₂O), 85 %RH, as determined by SEM observations or (*)
 349 from ToF-SIMS profiles.

350

351 3.3. Composition of the alteration layer

352 The composition of the alteration layer has been measured by EDX with an acceleration tension
 353 of 7 kV, on the surface, in an area free of carbonate salts, for the samples aged for 3 months or 6
 354 months at 40°C. At this tension, the analyzed depth is equal to, or less than, 0.6 μm for all the
 355 elements of the glass (calculation with the STRATAGEM software). Therefore, the alteration
 356 layer only contributes to the chemical analysis. The glass plate aged for 9 months at 40°C has
 357 developed an alteration layer of 6.5 μm in thickness, which has started to delaminate. It was thus
 358 possible to measure the composition on the edge, by scanning a 1 μm² area at two distinct

359 locations: near the surface (location 1) and near the interface with the pristine glass (location 2).
 360 For the different analyses, the mass elemental fractions totalized 92.5% to 95%. The difference is
 361 attributed to the contribution of protons and to porosity (because the contribution of protons is
 362 of the order of 1 wt% according to the estimated H₂O content, see below). The elemental
 363 fractions reported in Table 2 are normalized to 100% and compared with the composition of the
 364 pristine glass A. The elemental ratio Na/Si, K/Si and Ca/Si and the O weight fraction calculated
 365 from the cationic composition are also given in Table 2. The difference between the measured O
 366 and the calculated O is attributed to water, it allows the calculation of a hydration rate given in
 367 the last column. Note that because we used the pristine glass A as a standard for these analyses,
 368 the error is minimized and estimated to ~1% for the major elements.
 369

	O	Na	Mg	Al	Si	K	Ca	calculated O	Na/Si	K/Si	Ca/Si	H ₂ O* wt%
A pristine	59.5	7.58	0.42	0.33	25.2	4.98	1.90	59.5	0.30	0.20	0.075	-
A V3m surface	63.5	5.29	0.58	0.31	25.2	4.95	0.13	56.7	0.21	0.20	0.005	6
A V6m surface	64.4	5.87	0.63	0.29	23.6	5.06	0.11	58.3	0.25	0.21	0.004	9
A V9m edge, 1	64.0	4.72	0.37	0.34	25.6	4.95	0.00	57.0	0.18	0.19	0.00	6.1
A V9m edge, 2	63.73	4.71	0.27	0.42	25.76	5.12	0.00	57.32	0.18	0.20	0.00	5.6

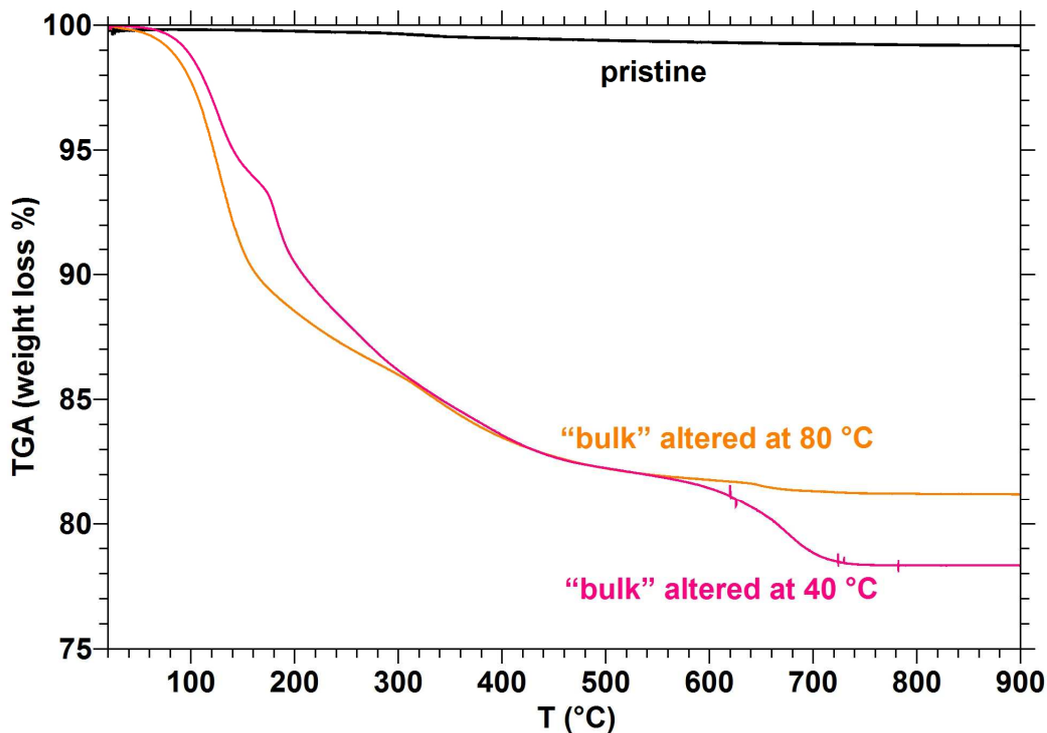
370 **Table 2.** EDX average composition (atom% element with 100% normalization) of the alteration
 371 layers of glass A analyzed at the surface or the edge of the glass plates (7 kV acceleration tension)
 372 after a V test at 40 °C, 85 RH%, for various ageing duration (hermetic box containing a saline
 373 solution of KCl in H₂O). Pristine glass A is used as standard and matrix effects are considered by
 374 using the data processing software STRATAGem. * Average hydration rate in weight %
 375 estimated from the comparison between the oxygen atomic fraction obtained by EDX and the
 376 one calculated by stoichiometry (calculated O column). Concentration ratios R/Si (R=Na, K, Ca)
 377 are calculated from EDX results.
 378

379 These chemical analyses indicate that the hydrated layer is partially depleted in Na (about 40% of
 380 the Na is lost after 9 months), and totally depleted in Ca. It is not depleted in K. Concerning Mg,
 381 it is still present in the layer but as a minor element, its analysis is not accurate enough to

382 comment the variation. Its presence could not be put in evidence in the carbonate precipitates on
383 the surface.

384 The hydration rate of the layer is 6 to 9 wt%. It can be compared to the hydration rate estimated
385 by thermogravimetric analysis of bulk altered glass powder. According to the thickness of the
386 alteration layer after 6 months of ageing (2.5 μm), the 5-10 μm powder fraction aged for 6
387 months should be altered in the bulk and this has been verified by ^{27}Al and ^{23}Na MAS NMR.
388 Three thermograms are shown in Fig. 5: the 5-10 μm powder aged for 6 months at 40°C, the 32-
389 50 μm powder aged for 72h at 80°C [28] and a pristine glass A powder. The mass loss below 150-
390 200°C is attributed to weakly bound molecular water, it is around 10 wt% at both 80°C and 40°C
391 (the kink on the 40°C powder thermogram is possibly due to the dehydration of
392 hydroxycarbonates). We observed that the mass loss in this temperature range depends on the
393 conservation conditions of the powders (inside or outside a dessicator for instance), indicating an
394 equilibrium of this weakly bound water with the atmosphere (RH). This water is lost when the
395 samples are put under vacuum in the SEM chamber, so that the mass loss that should be
396 compared with the SEM hydration rate is that measured beyond 150-200°C. On this temperature
397 range, there is a mass loss event between 600 and 700°C in the 40°C altered glass powder, which
398 corresponds to the decomposition of carbonates (CO_2 departure). This event is minimized in the
399 80°C altered powder because no carbonates formed at this temperature as described before (a
400 few carbonates appeared after the ageing). The hydration rate evaluated by the mass loss in the
401 150-500°C range, taking into account the contribution of carbonates to the total mass of the
402 40°C altered powder, is about 8 wt%, in very good agreement with the EDX hydration rate
403 (Table 2). The hydration rate of the 80°C altered powder is similar. The mass of CO_2 in the
404 carbonates is 4.4 wt% of the mass of the 40°C altered powder. It is consistent with the calculated
405 mass of CO_2 in calcite by assuming that all the Ca in this altered glass powder has formed calcite
406 (3.9 wt%), the difference being assigned to the presence of Na-bearing carbonates. This
407 consistency between glass plates data and glass powder data is important to emphasize because it
408 gives robustness to our structural study based on the analysis of altered glass powders.

409



410 **Fig. 5** Thermogravimetric analysis of pristine glass A powder and “bulk” altered glass A powders
 411 after an ageing at 80 °C (32-50 μm , V72h, climatic oven) or at 40 °C (5-10 μm , V6m, hermetic
 412 box containing a saline solution of KCl in H_2O), 85 %RH. Bulk alteration was checked by ^{27}Al
 413 MAS NMR.

414
 415 By comparing the EDX analysis at the top of the hydrated layer (Table 2, 9 months, location 1)
 416 or at the bottom (Table 2, 9 months, location 2), we notice that the hydrated layer is chemically
 417 homogeneous. The absence of chemical gradients points out the fact that all the water-glass
 418 reactivity occurs at the interface (within 0.1 μm that is our SEM-EDX spatial resolution), leaving
 419 behind a hydrated layer that is apparently at equilibrium with the atmosphere and the carbonate
 420 salts on the surface. Nevertheless, this equilibrium is only apparent because the Na content of the
 421 hydrated layer slowly decreases as a function of the ageing time (compare 3 months and 9
 422 months in Table 2). Another remarkable feature is the complete absence of K depletion,
 423 consistent with the absence of K in the carbonate precipitates. There is a striking contrast
 424 between the behavior of K (not at all depleted) and Ca (totally depleted).

425 An attempt to examine the interface between the hydrated layer and the pristine glass has been
 426 carried out by TOF-SIMS profile analysis of glass plates aged at 40°C for 17 days and for 3
 427 months. Because of the presence of carbonates on the surface, this attempt was not successful in

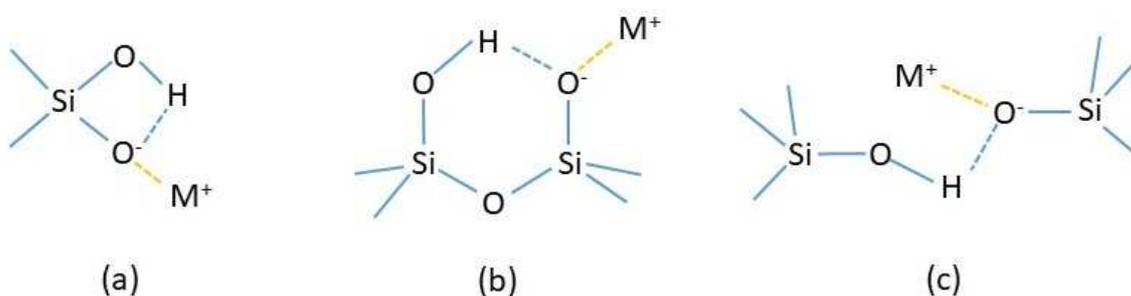
428 accurately profiling the interface, but it confirmed the extent of the depletion of the various
429 cationic elements. The description of this result is placed in the Data in Brief associated to this
430 paper.

431

432 3.4. Atomic structure of the alteration layer

433 Glass powders of 5-10 μm , 10-20 μm and 32-50 μm have been aged for 6 months at 40°C, or 2
434 months at 50°C, or 12 days at 60°C (85 %RH) respectively, in order to produce alteration in the
435 bulk. The spectra of the bulk altered glass powder (32-50 μm) aged at 80°C published in [28] are
436 reported also for comparison. The ^1H MAS NMR spectra at the different temperatures are
437 compared in Fig. 6a while the Hahn echo (HE) acquisitions of the 40°C altered powder are
438 shown as a function of varying echo time T_E in Fig. 6b (peaks of the species experiencing the
439 strongest homonuclear ^1H - ^1H dipolar interactions – thus in a dense proton network – decrease
440 faster; thus using a variable echo delay helps identify different environments). These series of HE
441 spectra put in evidence at least 5 hydrated species in the aged glass: a broad signal at about +5
442 ppm, two high frequency signals at +10 ppm and +14 ppm and two weak, narrow signals at
443 about 0-+3 ppm. The broad signal centered at +5 ppm dominates in the spinning sidebands
444 manifold, and its relative intensity decreases rapidly with increasing T_E . Following the literature
445 [35,36], this behavior allows to assign it to molecular water, because the strong homonuclear
446 dipolar coupling in H_2O associated to its high concentration induce shorter spin-spin relaxation
447 time (T_2) and irreversible loss of this signal in the central band of the HE spectra at high echo
448 time, with respect to the other hydrated species with weaker homonuclear dipolar coupling.
449 Moreover, the relative intensity of this signal decreases faster in the spinning sidebands than in
450 the central band (this is particularly visible in the $T_E=120 \mu\text{s}$ spectrum in Fig. 6b). It means that
451 molecular water does not account for all the intensity at about 5 ppm. Note that because of these
452 irreversible losses that depend on the hydrated species (different T_2 values), it is not really
453 possible to quantify the proportion of these species by simulating the central band intensity.
454 Besides, the widths of these spectra have two main origins: i) the distribution of isotropic
455 chemical shifts (inhomogeneous width) and ii) the residual homogeneous width due to the
456 incomplete averaging (i.e., cancellation) of the dipolar couplings by the MAS. It is highly probable
457 that in our glasses, using a MAS spinning frequency of 10 kHz, the inhomogeneous width
458 dominates, as it has been demonstrated in hydrous aluminosilicate glasses [37].
459 By increasing the echo time, the water signal is filtered and the other contributions with longer T_2
460 appear. The two peaks at +10 ppm and +14 ppm most likely belong to the same broad and
461 asymmetrical distribution of intensity between +14 and +3 ppm, with most of the intensity on

462 the intermediate to high frequency side. This broad component corresponds to OH groups with
 463 a distribution of hydrogen bonding strength. Indeed, a correlation between the chemical shift and
 464 the O-H...O distance, which measures the strength of the hydrogen bonding, has been reported
 465 for long in the literature and supported by theoretical calculations [35,37]. The O-H...O
 466 distances that give account of the +10 and +14 ppm chemical shifts are 2.65 Å and 2.50 Å
 467 respectively. Such high ^1H isotropic chemical shifts revealing very strong hydrogen bonding of
 468 the OH groups are commonly observed in alkali silicate crystal hydrates [35,38,39] and in
 469 hydrous alkali silicate glasses [40–42]. They are not observed in hydrous silica [43], nor in glasses
 470 altered by immersion in liquid water, which are depleted in alkalis and of which ^1H isotropic
 471 chemical shifts do not exceed 8 ppm [44]. Therefore, they are related to the presence of alkalis
 472 and non-bridging oxygens (NBO) and indicate hydrogen bonding between a Si-OH group and a
 473 Si-O(M^+) group possibly connected to the same Si [38,45] with M^+ being an alkali, as in the
 474 following picture where (a) is an intratetrahedral H-bonding in a hydrated Q^2 species, (b) is a H-
 475 bonding between neighboring tetrahedra and (c) is a H-bonding between adjacent, but not
 476 neighboring tetrahedra:



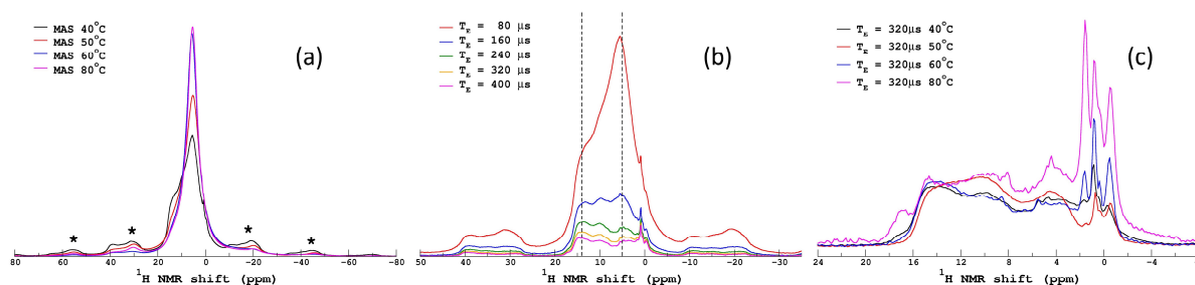
477
 478 In the cases (b) and (c), the two tetrahedra are noted $\text{Q}^3\text{-H}$ and $\text{Q}^3\text{-M,H}$ to refer to the H-bond
 479 of this species. The O(H)-O- distance is expected to be about 2.64 Å in the intratetrahedral H-
 480 bonding (a), between 2.55 Å and 2.75 Å in H-bonding between neighboring tetrahedra
 481 depending on the nature of the alkali, in the order $\text{K} < \text{Na} < \text{Li}$ (b) [40], and distributed in (c). Note
 482 that in these structural arrangements, the protons are almost structurally equivalent to modifier
 483 cations. Such a proton species exists in KHSi_2O_5 crystal structure (with $\delta_{iso} = 15.6$ ppm) and it
 484 has been proposed that similar NaHSi_2O_5 units exist in sodium silicate crystal hydrates like
 485 sodium disilicate hydrates (^1H $\delta_{iso} = 15.3$ and 18.7 ppm, [38]) and hydrates with high silica
 486 content (octosilicate, ^1H $\delta_{iso} = 16$ ppm, magadiite, ^1H $\delta_{iso} = 14.9$ ppm and kenyaite, ^1H $\delta_{iso} =$
 487 15 ppm, [40]) with the difference that Na^+ ions are also surrounded by water molecules. In these
 488 hydrates, by examining the ^1H NMR line widths and the ^2H (D) quadrupolar parameters of OD
 489 groups in deuterated samples, the protons of the H-bonded OH may be considered as rigid with
 490 respect to the NMR time scale, and they do not exchange with protons of water molecules

491 [39,45].

492 At last, narrow, weak signals at low chemical shift (0-+3 ppm) are assigned to isolated OH
493 groups, because of their weak H-H dipolar coupling and low chemical shift. These species may
494 be isolated Si-OH or M-OH groups, possibly in crystalline environments [35,40].

495 The comparison of the MAS spectra in Fig. 6a indicates that the relative proportion of molecular
496 water and OH groups vary with the temperature of ageing in a regular way, with more molecular
497 water at high temperature. The total amount of water (H_2O+OH) has not been measured in
498 these powders at the time of the NMR experiment, so that it is difficult to draw any conclusion
499 from this tendency. As we have noticed with the TGA measurements, there is an equilibrium
500 between the weakly bound molecular water in the layer and the atmospheric humidity. All the
501 spectra have been measured in the same atmosphere. Either the alteration layer produced at high
502 temperature (60°C, 80°C) takes more molecular water, or the relative proportion of the species
503 differ: the answer needs a careful study with control of the total H_2O content.

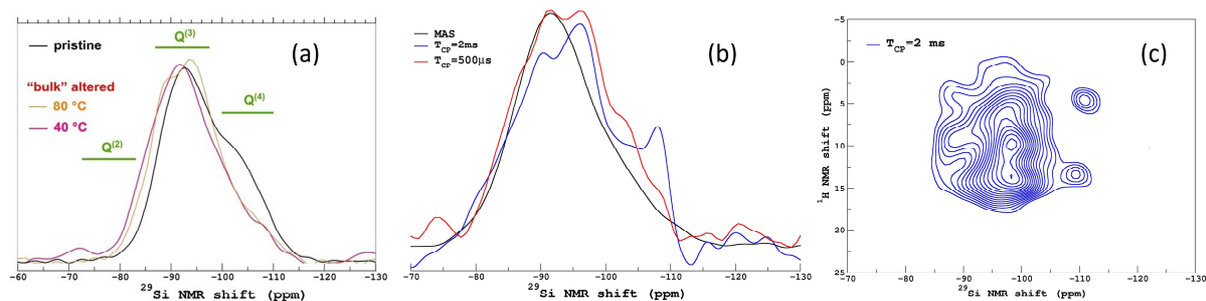
504 The HE acquisitions ($T_E = 320 \mu s$) of the glass powders aged at the different temperatures are
505 given in Fig. 6c, to compare the spectra after filtering the water (and Si-OH species close to this
506 water having the shortest T_2). These spectra are very similar regarding the number, positions and
507 relative intensities of the peaks. This points out a structural similarity of the hydrated layer
508 produced in the 40°C-80°C temperature range. The quantitative analysis of the Si-OH and water
509 speciation would require a deeper experimental study.



510 **Fig. 6** (a) 1H MAS NMR spectra of bulk altered glass A powders after an ageing at 40°C, 50°C,
511 60°C, 80°C and 85 %RH. The spectra are normalized to the same area including the spinning
512 side bands. *: spinning sidebands. (b) Hahn echo acquisitions with variable echo delays T_E of the
513 bulk altered powder produced at 40°C, 85 RH%. (c) Hahn echo acquisitions ($T_E = 320 \mu s$) of the
514 bulk altered glass A powders produced at the different temperatures. The spectra are normalized
515 in height to the peak intensity centered around +14 ppm.

516
517
518 The ^{29}Si MAS NMR spectra of the pristine and altered glass A at 40°C and 80°C are depicted in
519 Fig. 7a. Intermediate temperatures give similar spectra and are therefore not shown. The

520 simulation of the pristine glass spectrum gives 35% of Q^4 species at -103 ppm and 65% of Q^3
521 species at -92 ppm, in excellent agreement with the glass A composition assuming homogeneous
522 distribution of the NBOs in the silicate network, and with the chemical shifts in the literature
523 [46]. After ageing in humid atmosphere at both temperatures, the Q^4 band decreases very
524 significantly while the Q^3 band maintains its intensity and is enlarged on the high frequency-side.
525 The apparition of a few Q^2 units might explain part of this enlargement, but the main part is
526 probably owed to the widening of the Q^3 chemical shift distribution because of a greater variety
527 of Q^3 species (Q^3 -H and Q^3 -Na and all the intermediate Q^3 -H,Na species with varying H-
528 bonding strength). We notice that there is a double-peak in the Q^3 component of the glass
529 powder aged at 80°C, which is not visible in the powder aged at 40°C. This double-peak appears
530 in the $^1\text{H} \rightarrow ^{29}\text{Si}$ CP MAS spectrum of this latter powder, with a long contact time (2 ms, Fig.
531 7b). Indeed, although no efficient $^1\text{H} \rightarrow ^{29}\text{Si}$ cross-polarization could be achieved on the 80°C
532 altered powder (possibly because of a high amount of short T_2 H species), it was possible to
533 acquire $^1\text{H} \rightarrow ^{29}\text{Si}$ CP MAS and HETCOR spectra on the 40°C altered powder, shown in Fig. 7b
534 and 7c. At short contact time (0.5 ms), all the hydrated species are in the vicinity of a wide
535 distribution of Q^3 (and possibly Q^2) units. However, at long contact time (2 ms), two additional
536 Q^n species show up at a longer distance to protons: a Q^4 signal at about -109 ppm, remarkably
537 narrow, and a Q^3 signal at about -96 ppm analogous to the low-frequency peak in the 80°C
538 altered powder. Thus, we propose to assign the Q^3 low-frequency peak (-96 ppm) to Q^3 -Na
539 species, or to Q^3 -Na,H species undergoing weak H-bonding (long O---H distance), and the high-
540 frequency peak (-88 ppm) to Q^3 -H species, or to Q^3 -Na,H species undergoing strong H-bonding
541 (short O---H distance). This assignment is consistent with the fact that Q^3 -Na species are
542 expected to be more abundant in the 80°C altered glass, because it has retained all the alkali [28]
543 with respect to the 40°C altered glass that has lost up to 40% of the alkalis. On the other hand,
544 the narrow Q^4 signal is not expected in a glass. Its ^{29}Si chemical shift is the same as in cristobalite
545 [47]. This suggests the possible presence of locally ordered re-polymerized areas in the hydrated
546 glass at 40°C, in agreement with hints of a re-polymerized component in the corresponding
547 Raman spectra (see below).
548



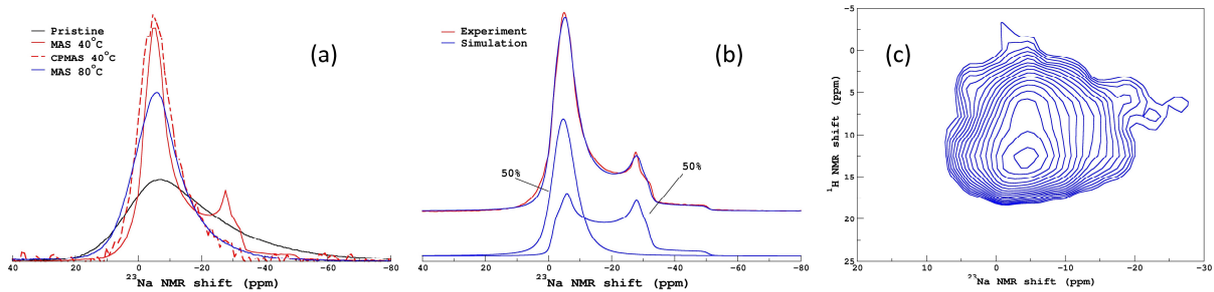
549
 550 **Fig. 7** (a) ^{29}Si MAS NMR spectra of bulk altered glass A powders after an ageing at 40°C or 80
 551 °C, 85 %RH. The ^{29}Si NMR MAS spectrum of pristine glass A powder is reported for reference.
 552 (b) CPMAS ^1H - ^{29}Si acquisition for two contact times (T_{CP}) of the bulk altered glass A powder
 553 aged at 40°C, 85 RH% and (c) HETCOR ^1H - ^{29}Si acquisition of the same powder for $T_{\text{CP}}=2$ ms.

554
 555 This NMR study has been completed by ^{23}Na MAS NMR and $^1\text{H} \rightarrow ^{23}\text{Na}$ CP experiments on the
 556 altered powders, reported in Fig. 8. The ^{23}Na MAS NMR spectrum of the pristine glass A shows
 557 a broad asymmetrical profile which is typical of Na in alkali silicate glasses [48,49]. The
 558 broadening originates from the distribution of both the isotropic chemical shift and the second-
 559 order quadrupolar interaction. These spectra have been fitted following the procedure described
 560 in [50]. The fit of the 40°C altered powder is shown in Fig. 8b and the fit parameters are reported
 561 in Table 3.

562 Comparing the pristine glass A and the glass powder aged at 80°C, the ^{23}Na NMR parameters
 563 indicate a very slight increase of the Na-O mean distance, and an environment that has become
 564 more centrosymmetrical with the alteration (see discussion in [28]). The ^{23}Na NMR spectrum of
 565 the glass powder aged at 40°C bears two signals, the signal of Na^+ in the hydrated glass, having a
 566 $^{23}\text{Na} \delta_{\text{iso}} \sim -2.7$ (2.3) ppm and $C_Q \sim 1.1$ (0.5) MHz (standard deviation under brackets) and the
 567 sharp signal of Na^+ in a crystalline phase that may possibly be assigned to Na, Ca mixed hydrated
 568 carbonates such as gaylussite ($\text{Na}_2\text{Ca}(\text{CO}_3)_2 \cdot 5\text{H}_2\text{O}$) that has been identified on the Raman
 569 spectrum (Fig. 9) although no ^{23}Na NMR reference spectrum could be found to confirm. The ^1H
 570 $\rightarrow ^{23}\text{Na}$ CPMAS NMR spectrum does not exactly overlap the MAS spectrum (Fig. 8a). This
 571 indicates a complex distribution of environments for Na in this powder aged at 40°C, with, in
 572 average, a more centrosymmetric character (lower mean C_Q) and larger Na-O distances (lower
 573 mean δ_{iso}) than in the powder aged at 80°C. The $^1\text{H} \rightarrow ^{23}\text{Na}$ correlated component and the 80°C
 574 altered glass have very similar ^{23}Na parameters although (Table 3).

575 In the $^1\text{H} \rightarrow ^{23}\text{Na}$ HETCOR spectrum in Fig. 8c, the sharp signal is correlated with protons at
 576 low-frequency only, while the signal of Na^+ in the hydrated glass is correlated with all the proton
 577 frequencies, in particular that around +14 ppm consistently with their attribution to OH groups

578 forming hydrogen bonds with NBOs. Thus, Na⁺ ions are in close vicinity of all the hydrated
 579 species (OH and molecular H₂O) in the glass hydrated at 40°C as well as in the glass hydrated at
 580 80°C [28]. However, at 40°C, part of the Na⁺ ions have migrated and formed carbonates.



581
 582 **Fig. 8** (a) ²³Na MAS NMR spectra of pristine and bulk altered glass A powder after an ageing at
 583 80 °C, 85 %RH (32-50 μm, V72h, climatic oven) or at 40 °C, 85 %RH (5-10 μm, V6m, hermetic
 584 box containing a saline solution of KCl in H₂O). Spectra are normalized to the same area. (b) Fit
 585 and decomposition of the spectrum of the 40°C altered powder. (c) HETCOR ¹H → ²³Na
 586 acquisition with a contact time of 0.8 ms of the 40°C altered powder.
 587

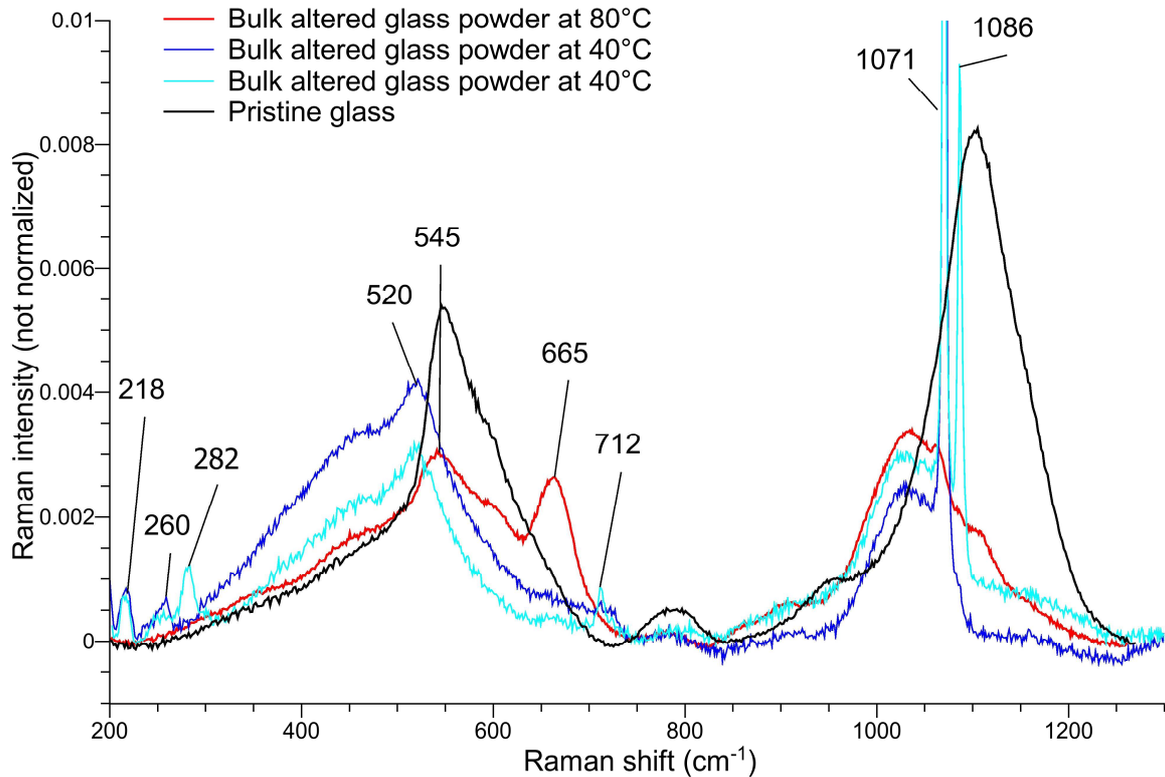
	²³ Na δ _{iso}	C _Q (MHz)	η _Q	Assignment and proportion
Pristine glass A	+3 (9)	4.3 (1.4)	0.6	
Bulk altered glass 80°C	-1 (4)	1.6 (0.7)	0.6	
Bulk altered glass 40°C MAS	-2.7 (2.3)	1.1 (0.5)	0.6	Large component 50%
	4.8	3.9	0.15	Crystallized component 50%
Bulk altered glass 40°C CP MAS ¹ H → ²³ Na	-1 (3)	1.8 (0.4)	0.6	

588 **Table 3.** ²³Na NMR fit parameters of the bulk altered glass A powder aged at 40°C. The
 589 parameters of the pristine glass and 80°C altered powder are also given for comparison.

590
 591 The Raman spectra of the pristine and bulk altered glass powders are compared in Fig. 9. The
 592 Raman spectrum of the pristine glass A is typical of alkali silicate glasses with O/Si ratio between
 593 2.25 (tetrasilicate composition) and 2.5 (disilicate composition), of which silicate network is
 594 mainly composed of Q⁴ and Q³ units. In the high-frequency range, the band at 1100 cm⁻¹ is
 595 assigned to the stretching modes of Q³ units while the small band at 950 cm⁻¹ is related to Q²
 596 units [51,52] likely bound to Ca²⁺ and Mg²⁺ cations [53]. In the intermediate frequency range
 597 (300-700 cm⁻¹), the Raman intensity stems from the Si-O-Si bending modes of the silicate

598 network, and directly reflects the Si-O-Si angle distribution. In alkali silicate glasses, this
599 distribution is bimodal, with high and widely distributed Si-O-Si angles in polymerized silica-type
600 regions, and weaker, less distributed Si-O-Si angles in depolymerized sodo-silicate type regions.
601 The first population is responsible of the broad low-intensity signal around 460 cm^{-1} , while the
602 second population gives rise to the peak at 545 cm^{-1} [54]. The Raman spectra of the bulk altered
603 glass powders are severely modified. The Q^n stretching band downshifts to 1030 cm^{-1} for the two
604 ageing temperatures 40°C and 80°C . The interpretation of the Raman intensity in the $1020\text{-}1040$
605 cm^{-1} range is controversial. Zotov and Keppler assign it to Q^3 species distinct from those causing
606 the 1100 cm^{-1} band [55]. Because intensity in this region is present in alkali tectosilicate glasses
607 [56] and following theoretical results [57], Le Losq et al. ascribe it to asymmetrical stretching of
608 polymerized units [58]. Considering the NMR results described above on these altered powders,
609 we prefer the interpretation of Zotov and Kepler and propose to assign this band to Q^3 units
610 with strong hydrogen bonding of their NBO (these units will be noted $Q^3(\text{Na},\text{H})$ in this paper). It
611 is expected that this hydrogen bonding induces a low frequency shift of the Q^3 stretching band,
612 by weakening the Si-NBO bond. Beside, the well-defined Q^2 peak at 950 cm^{-1} interestingly
613 disappears, and a weak intensity signal around 910 cm^{-1} appears that has been clearly identified as
614 the stretching mode of Si-OH bonds [55,57]. The decrease of the 800 cm^{-1} band, ascribed to the
615 “cage-like” motion of Si in the Q^4 tetrahedron has been observed by Zotov and related to the
616 depolymerization of the network. The intermediate frequency range shows more variations
617 according to the ageing temperature. At 80°C , the Si-O-Si bending band at 545 cm^{-1} is still
618 present with a decreased intensity, and a new narrow band at 665 cm^{-1} has appeared. The
619 spectrum of the 80°C altered powder is very close to the spectra of Cultural Heritage glass replica
620 aged at 60°C in humid atmosphere with 100% RH published by Robinet et al. [59]. The narrow
621 band at $665\text{-}670\text{ cm}^{-1}$ has been proposed as a characteristic feature of these alteration conditions.
622 It is assigned to Si-O-Si symmetrical bending in depolymerized and isolated components of the
623 network, such as Q^2 chains in pyroxenes (of which Si-O-Si bending frequencies are in the range
624 $650\text{-}700\text{ cm}^{-1}$) [60]. Actually, it strongly evokes the intense and characteristic band at $668\text{-}672\text{ cm}^{-1}$
625 in calcium silicate hydrate (C-S-H), which consists in Q^2 chains and Q^1 dimers [61]. This
626 assignment is consistent with the retention of Ca^{2+} ions in the hydrated layer at 80°C . In the
627 spectra of the powder aged at 40°C , this peak is absent, and the intensity of the broad band with
628 maximum at 460 cm^{-1} has markedly increased, which can be related to partial re-polymerization
629 of the network. Moreover, the Si-O-Si bending peak originating from the depolymerized
630 component of the network has shifted to lower frequency, from 545 cm^{-1} (characteristic of the
631 NS3 composition [54]) to 520 cm^{-1} (between NS4 and NS7 composition). This downshift

632 indicates a lower content of alkalis in the altered glass, consistently with the chemical analysis of
 633 the alteration layer that shows a loss of up to 40% of the Na^+ ions. On the other hand, the sharp
 634 peaks in the spectra of the 40°C altered powder correspond to carbonates, among them calcite
 635 (peaks at 282, 712 and 1086 cm^{-1}), gaylussite ($\text{Na}_2\text{Ca}(\text{CO}_3)_2 \cdot 5\text{H}_2\text{O}$, peaks at 260 and 1071 cm^{-1}),
 636 the peak at 218 cm^{-1} remaining unidentified.



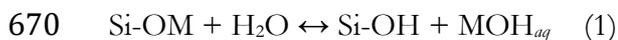
637
 638 **Fig. 9** Raman spectra of the pristine glass A (black curve) and of bulk altered glass A powder
 639 after an ageing at 80 °C, 85 %RH (32-50 μm , V72h, climatic oven) (red curve) or at 40 °C, 85
 640 %RH (5-10 μm , V6m, hermetic box containing a saline solution of KCl in H_2O). Because of the
 641 presence of carbonates in the powder aged at 40°C, there is a variability in the Raman spectra, so
 642 that two spectra are shown for this powder (blue and cyan curves).

643
 644 **4. DISCUSSION**

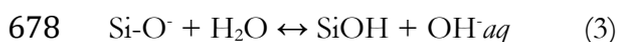
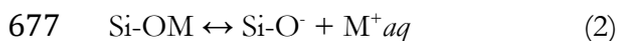
645 The alteration layer produced in glass A in atmospheric conditions (85 %RH) is composed of a
 646 hydrated layer with no depletion of any chemical element at 80°C, and of a hydrated layer
 647 depleted in Ca (totally) and Na (partially) with carbonate salts on the surface at 40°C. The
 648 concentration of tightly bound water is about 5 to 10 wt% with no clear indication of a different
 649 water content at 80°C and at 40°C. This water is present as molecular H_2O and as Si-OH species
 650 with a wide distribution of H-bonding strengths, the strongest ones connecting H to NBO's like
 651 in alkali silicate hydrates. From ^{29}Si NMR and Raman spectra, the silicate network has

652 depolymerized with the ageing and bears primarily Q³-H (Si-OH) and Q³-M,H (Si-OM---H with
653 M = Na or K) units, although at 40°C a part of the network has also re-polymerized, possibly
654 after the departure of some Na⁺ and Ca²⁺ ions. We emphasize that the structure of the hydrated
655 layer produced in these atmospheric conditions markedly differs from the structure of the
656 alteration layer, often called « alteration gel », produced in immersion conditions. In the latter
657 conditions, the gel is completely depleted in alkalis and the silicate network has re-polymerized,
658 following either *in situ* condensation reactions or silica precipitation from the solution [62]. In
659 atmospheric conditions, the hydrated glass still contains a significant amount of alkalis and
660 NBO's, and is closer to an alkali silicate hydrate than to an alteration gel. As in hydrates, the
661 dense network of hydrogen bonds probably contributes to the stability of the layer [63].

662 In the hydrated layer, Na⁺ ions are surrounded by NBO's, OH groups and molecular water as
663 inferred from the ¹H → ²³Na CP NMR spectra. They are thus totally or partially solvated
664 explaining that they are finally detached from the NBO's and leached out of the glass. There is
665 no structural information on the environment of K⁺ and Ca²⁺, but because leaching of K⁺ is
666 never observed in these experiments, we may propose that K⁺ ions are still primarily bound to
667 NBO's in the hydrated glass. On the contrary, at 40°C, Ca²⁺ ions are primarily solvated because
668 they are totally leached out, forming carbonates on the glass surface. These observations can be
669 summarized by considering the following equilibrium in the hydrated glass:



671 Where M are alkali (Na⁺ or K⁺) or half a Ca²⁺ ion and *aq* refers to the hydrated state of MOH.
672 This well-known equilibrium has been referred to as « ion exchange » [64] but it is more exactly
673 an acid-base reaction of water with the NBO sites, implying diffusion of molecular water and its
674 immobilization by the formation of silanol groups [65,66]. It can also be viewed as a
675 complexation equilibrium of the M cations with the silicate polyanionic network, associated to
676 the acid-base reaction ((1) = (2)+(3)) :



679 Reaction (2) points out the fact that there is a balance between the solvation (or hydration) of the
680 M⁺ cations and their inner sphere complexation on the silicate NBO sites. A similar balance
681 governs the selectivity of clays for alkali ions and results in the order Cs⁺ > K⁺ > Na⁺ > Li⁺,
682 because the energy of hydration that goes in the reverse order overrides the energy of
683 complexation in the interlayer (Teppen and Miller, 2006; Rotenberg et al., 2009). Similar ionic
684 selectivity has been found in the alteration gels of the International Simple Glass (ISG) formed in
685 Si-saturated solutions with increasing alkali salts concentrations : the K⁺ or Cs⁺ ions provided by

686 the solution almost totally substitute the Ca^{2+} ions in the gels, while Na^+ and Li^+ ions do not [69].
687 The authors interpreted these results by analogy with the selectivity in clays, suggesting that it is
688 mainly determined by the affinity of cations for hydration, rather than for pore surface
689 complexation: K^+ and Cs^+ are “hydrophobic” cations that prefer complexation. We propose to
690 use the same analogy to explain our observation that cationic species are detached of the NBO’s
691 in the order $\text{Na}^+ > \text{K}^+$. Na^+ ions are more stabilized by solvation than K^+ and are thus more
692 quantitatively converted to solvated species, able to migrate under a chemical potential gradient.
693 Moreover, in alkali disilicate hydrates, K^+ are not coordinated to molecular H_2O and form
694 “ KHSi_2O_5 ” units, while Na^+ are coordinated to NBO *and* H_2O [39]. It is then possible that Si-
695 OK units are particularly stabilized in the H-bonding network, adding another energy term
696 favouring the left side of reaction (2). Concerning Ca^{2+} ions, their observed behaviour is striking
697 as it is very contrasted with the temperature, revealing both the specificity of atmospheric
698 alteration phenomena and the temperature-induced change in their mechanisms. At 80°C , the
699 Ca^{2+} ions remain in the alteration layer and locally form C-S-H-type structures evidenced by the
700 665 cm^{-1} sharp band in the Raman spectra. At 40°C on the contrary, Ca^{2+} ions are totally leached
701 out of the alteration layer, forming calcium carbonates on the surface. This behaviour is in
702 contrast to the one expected considering the calcite solubility, which decreases when temperature
703 rises. The hydration energy of Ca^{2+} is three times that of Na^+ so that Ca^{2+} are markedly stabilized
704 by hydration [70]. Following the preceding discussion, this probably explains that Ca^{2+} ions are
705 more quantitatively extracted out of the hydrated layer than Na^+ at 40°C . Because the enthalpies
706 of hydration and complexation are likely both modest, the dominance of the hydration term is
707 expected to be similar at 40°C and at 80°C . So, why aren’t Ca^{2+} ions (and Na^+ ions) also
708 extracted at 80°C ? To answer this question, we consider the specificity of atmospheric alteration
709 that lies in the scarcity of water with respect to the silicate solid phase. In the first stage of the
710 alteration, it may be assumed that all the water molecules are bound to the surface or within the
711 hydrated layer, consequently losing part of their solvating properties. Such an effect has been
712 observed in several molecular dynamics simulations of water confined in silica nanopores [71–
713 73]. Then, the complete solvation of the M^+ ion and its detachment from the NBO may be
714 considerably slowed down. If water molecules are engaged in another reaction as the Si-O-Si
715 bond hydrolysis, as proposed below, then the M^+ ions may never be completely solvated and
716 extracted.

717 Next to the acid-base reaction on the NBO sites (reaction (1)), there is another reaction called
718 hydrolysis of the Si-O-Si bonds:



720 This reaction *only* is responsible for the silicate network depolymerisation when shifted to the
721 right. It is well-known that this reaction can be catalysed either by protonic assistance (acid pH)
722 or by nucleophilic attack by OH⁻ ions (basic pH). The activation energy of this reaction is high
723 (86 kJ.mol⁻¹ according to [34]) and certainly higher than the activation energy for hydration.
724 Therefore, this reaction is considerably enhanced with temperature, going much faster to the
725 right at 80°C with respect to 40°C. This provides the basis of our hypothesis to explain the
726 different phenomena observed for atmospheric alteration of glass A at 80°C and at 40°C. Indeed,
727 in addition to the different behaviour of Na⁺ and Ca²⁺ cations described above, there is a
728 considerable temperature-induced increase of the kinetics of the glass hydration, as measured by
729 the growth of the hydrated, depolymerized layer. We propose that it is due to the increase of the
730 rate of the hydrolysis reaction (4), relatively to the rate of reaction (1). Water molecules are
731 primarily engaged in network hydrolysis rather than in acid-base dissociation onto NBO sites at
732 80°C. On the contrary at 40°C, the rates of the two reactions converge and water molecules
733 have the time to solvate M⁺ and OH⁻ ions that then migrate towards the surface to form
734 carbonates. Note that Si-O-Si hydrolysis, instead of acid-base dissociation has been inferred from
735 the study of the hydration process of sodium disilicate, forming Q² [(SiO)₂Si-(OH)O⁻Na⁺] rather
736 than Q³ [(SiO)₃Si-OH] groups. The authors suggested a possible stabilization of these Q² groups
737 by the strong H-bonding [63].

738 Another striking feature of the atmospheric alteration of glass A is the acceleration of the
739 kinetics, observed both at 80°C and at 40°C. The simple “diffusion control” or “surface control”
740 models cannot account for this complex kinetics. Given our understanding of the alteration
741 phenomena described above, the network hydrolysis (reaction (4)) is the rate-controlling step at
742 least at 80°C. This reaction is accelerating with the ageing time and hydrated layer thickness (in
743 the timescale of our experiments). Wang et al. have highlighted the existence of self-accelerating
744 mechanisms in the aqueous alteration of silicate glasses, due to the catalytic effect of OH⁻ ions
745 that are released by the acid-base reaction step [74,75]. We propose that this effect also operates
746 in atmospheric conditions. After a time period depending on the availability and solvation
747 properties of water in the hydrated layer, M⁺ and OH⁻ ions end up being released and catalyse the
748 network hydrolysis, accelerating the glass hydration rate. It is probable that OH⁻ ions stay trapped
749 in the H bonding network, as well as their precursor form NBO+H₂O, and that both species
750 contribute to the acceleration of the Si-O-Si bond hydrolysis.

751 At last, we have attempted to rationalize our observations of the atmospheric alteration
752 phenomenology of glass A using a molecular-scale approach based on the relative rates of
753 reactions (1) and (4). This also corresponds to the basis of the incongruent dissolution model for

754 glasses [76]. Two considerations have been added: the important role of the hydration of cations,
755 and the bound or *ice-like* character of water molecules due to their confinement and scarcity.
756 These considerations are very recent in the glass alteration community. On the other hand, an
757 alternative model for glass dissolution has been proposed on the basis of the very sharp interface
758 between the gel and the pristine glass. It consists in the coupling of surface congruent dissolution
759 and precipitation at the interface (CIDP for Coupled Interface Dissolution Precipitation) [32].
760 We also observe many features that evoke CIDP, as the sharpness of the interface evidenced by
761 the flaking phenomenon and the chemical homogeneity of the hydrated layer. However, we think
762 that the molecular aspects that we have discussed, because they occur at the molecular scale near
763 the alteration front, are relevant to both models. It is possible that the properties of confined
764 water specific to atmospheric conditions preclude any meaningful distinction between the two
765 descriptions of glass alteration.

766

767 5. CONCLUSION

768 In this study, the atmospheric alteration (RH = 85 %) of a mixed alkali lime silicate glass
769 representative of a family of unstable compositions of the Cultural Heritage has been described
770 in detail as a function of temperature and time. It is the first time that compositional information
771 on the hydrated layer is accompanied by structural data (multinuclear NMR and Raman
772 spectroscopies) obtained on bulk altered glass powders. First, this study provides a clear evidence
773 that in atmospheric conditions, the hydrolysis of the silicate network (Si-O-Si bonds) is mainly
774 responsible for the glass hydration and controls its kinetics. The leaching of alkali and alkaline-
775 earth cations is limited, probably because of the scarcity and tightly H-bonded character of water,
776 which diminish its solvation property. At 40°C, the cations are leached in the order $\text{Ca}^{2+} > \text{Na}^+$
777 $\gg \text{K}^+$, possibly revealing the role of the hydration energy in the leaching process. At 80°C, no
778 leaching is even observed and the hydration by network hydrolysis occurs very rapidly. The
779 thermally activated character of hydrolysis induces an acceleration of this process with respect to
780 other processes (acid-base dissociation of water on NBO sites, cation solvation, cation migration
781 towards the surface). In addition, we suggest that the time acceleration of the hydration kinetics
782 at both temperatures points out another effect: the self-accelerating property of the system
783 towards hydrolysis, due to the catalytic effect of the OH^- ions trapped in the H-bonding network,
784 or of their precursor H_2O species that are H-bonded to NBOs. This effect may contribute to the
785 very strong temperature dependency of the hydration kinetics, because at 80°C more NBOs and
786 alkalis are retained in the layer. As a consequence of the partial alkali retention and network
787 depolymerization, the hydrated layer structure is closer to an unstable alkali silicate hydrate than

788 to an alteration gel (as produced in immersion conditions), at least in the first stages of the
789 alteration. This knowledge is important in the research for efficient surface chemical treatments
790 aimed at preventing glass objects from atmospheric degradation in museum collections.

791

792 **DATA AVAILABILITY**

793 The raw data required to reproduce these findings are available to download from Mendeley
794 Dataset <http://dx.doi.org/10.17632/mnjkhzr6sx.3>.

795

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1004

1005 **SUPPLEMENTARY MATERIALS**

1006 See Data in Brief zip file.