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1 **Diagenetic formation of stevensite by replacement of diatom frustules in the**  
2 **sediments of the alkaline Lake Alchichica (Mexico)**

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19

20 **Abstract:**

21 Better understanding the conditions of formation of authigenic Mg-silicates and their  
22 reactivity is key to interpret the paleoenvironmental message carried by the sedimentary record  
23 and evaluate the effect of reverse weathering, a process involved in long-term climate evolution.  
24 Microbialites from most alkaline crater lakes in Mexico contain Mg-silicates except those in  
25 Lake Alchichica, where concentration of orthosilicic acid is low (< 26  $\mu\text{M}$ ). Here, we  
26 investigated the first meter of sediments in Lake Alchichica in order to check how their  
27 mineralogy compared with that of shoreline microbialites. The mineralogy and chemistry of the  
28 sediment column were determined together with the pore water chemistry, providing insights  
29 on the processes occurring during early diagenesis. Below ~3 cm in depth, diatom frustules are  
30 progressively pseudomorphized into Al-poor Mg-silicates with a composition corresponding to

31 stevensite. This diagenetic process is massive and the resulting silicate represents between 30  
32 and 53 wt.% of the sediment content at all depths. This observation questions the possibility to  
33 infer lake paleochemistry from the presence/absence of Mg-silicates in the sedimentary record.  
34 Moreover, it allowed us to refine the conditions under which Mg-silicates authigenesis occurs:  
35 the saturation of the solution should be higher or equal to the solubility of a Mg-silicate phase  
36 close to that of “amorphous sepiolite”. Although the solubility of authigenic silicates is a key  
37 parameter of reverse weathering modeling during geological times, it is still debated. In this  
38 study, we proposed a solubility constant deduced from a natural system that should be  
39 considered when modelling the formation of Mg-silicates in a natural environment. The  
40 proportion of reverse weathering associated with this solubility constant could be higher than  
41 previously predicted based on experiments and thus have a greater impact on climate stability  
42 over geological timescales.

43

44 Keywords: reverse weathering; diatoms; Mg-silicates; solubility; diagenesis

45

## 46 **A) Introduction**

47

48 Authigenic clays are reported in various sedimentary environments from marine (*e.g.* Tosca  
49 et al., 2011) to soils (*e.g.* Cuadros et al., 2016), speleothems (*e.g.* Lèveillé et al., 2002) and  
50 lacustrine environments (*e.g.* Deocampo, 2015; Wright and Barnett, 2015). They form either  
51 through direct precipitation from solution (*i.e.* neoformation) or by transformation of precursor  
52 minerals (also called neoformation by addition) (Jones, 1986; Galán and Pozo, 2011). In the  
53 sedimentary environment, transformation is mostly related to diagenesis, whereas neoformation  
54 can take place in both syngenetic (depositional) and diagenetic environments (Galan and Pozo,  
55 2011). Clays forming in evaporitic closed basins where parent rocks are enriched in

56 ferromagnesian minerals, tend to be Mg-rich minerals as reported in the American Great Salt  
57 Lake (Jones and Deocampo, 2005) and Lake Abert (Jones and Weir, 1983), in several modern  
58 East African lake basins (Jones, 1986; Deocampo, 2005, 2015) and in Mexican crater lakes  
59 (Zeyen et al., 2015). Hereafter, they are referred to as Mg-silicates (Jones, 1986). They  
60 constitute a significant part of the Mg-rich phyllosilicate family. They structurally and  
61 chemically encompass diverse phases, either rich in Al such as saponite and palygorskite or  
62 poor in Al such as sepiolite, stevensite and kerolite (Pozo and Calvo, 2018). The latter, not  
63 listed as a valid mineral species by the international mineralogical association, is considered as  
64 a stacking-disordered talc phase (Brindley et al., 1977)

65

66 Al-poor Mg-silicates have been extensively studied in modern lakes from different localities  
67 such as the Satonda crater lake (Indonesia; Arp et al., 2003; Benzerara et al., 2010), Eyasi and  
68 Ngorongoro Crater Lakes (Tanzania; Deocampo, 2005), Lake Van (eastern Anatolia; Reimer  
69 et al., 2009), Lake Clifton (western Australia; Burne et al., 2014), Atexcac, Quechulac and La  
70 Preciosa Lakes (Mexico; Zeyen et al., 2015), Dziani Dzaha Lake (Mayotte; Gérard et al., 2018,  
71 Milesi et al., 2019, 2020), Mono Lake and Great Salt Lake (USA; Souza-Egipsy et al., 2005;  
72 Pace et al., 2016), as well as peri-marine environments (*e.g.* Lèveillé et al., 2000a, 2000b;  
73 Warren, 2016). Last but not least, such authigenic Mg-silicate phases have been observed in  
74 some ancient marine and lacustrine formations showing that they can be traced back in the  
75 geological record. For example, Tosca et al. (2011) reported occurrences of talc in two early  
76 Neoproterozoic carbonate formations located in the Akademikerbreen Group (Svalbard) and in  
77 the upper Fifteenmile Group of the Ogilvie Mountains (NW Canada), which they interpreted as  
78 the product of dehydration of authigenic hydrous Mg-silicates, including kerolite. It has been  
79 shown that the identity of these phases in modern environments depends on specific  
80 geochemical factors, such as pH, dissolved CO<sub>2</sub>, salinity, alkalinity, dissolved element

81 concentration ratios, and the activities of  $\text{Mg}^{2+}$ ,  $\text{H}_4\text{SiO}_4$ ,  $\text{Al}^{3+}$ ,  $\text{Ca}^{2+}$ , and  $\text{Fe}^{3+}$  in solution (Jones  
82 and Galán, 1988; Weaver, 1989; Deocampo, 2005; Bristow and Milliken, 2011). Several  
83 experimental studies have investigated the conditions allowing their formation (*e.g.* Wollast et  
84 al., 1968; Tosca et al., 2011). They have shown that high  $\text{pH} > 8.2$  and high concentrations of  
85 Si and/or Mg are required for their precipitation (Zeyen et al., 2015; Tosca et al., 2011). In  
86 addition, high salinity favors the formation of stevensite, whereas sepiolite and kerolite are  
87 formed under lower saline conditions, with high Si/Mg ratio favoring the formation of sepiolite.  
88 Likewise, precipitation of kerolite instead of sepiolite is favored by an increase of pH and  
89 dissolved Mg concentration (Pozo and Calvo, 2018). Last, Millot et al. (1970) proposed that  
90 the Al content of Mg-silicates could be indicative of the distance to the detrital source. Millot's  
91 model shows that detrital clay inputs, mainly kaolinite, illite, chlorite and Al-rich smectite  
92 preferentially occur in the margins of saline lake basins and transition to fibrous clays  
93 (palygorskite, sepiolite) that precipitate in the center of lakes.

94

95 Overall, these minerals have received considerable attention as potential  
96 paleoenvironmental recorders of the geochemistry of the solution in which they precipitated. In  
97 addition to their value for paleoenvironmental reconstruction, better understanding the  
98 conditions of Mg-silicates formation and their reactivity is interesting for two broad questions.  
99 First, they can have a crucial role in the control of carbonate fabrics and porosity as exemplified  
100 by the study of offshore Barra Velha Formation carbonate reservoirs in Brazil (Tosca and  
101 Wright, 2018; Wright and Barnett, 2020). Second, their precipitation feeds a process called  
102 reverse weathering, which consumes alkalinity without consuming carbon and may therefore  
103 have a significant role in the evolution of climates at the geological timescale (Krissansen-  
104 Totton and Catling, 2020; Isson and Planavsky, 2018). However, there remains a few  
105 difficulties for their use as a proxy. First, there is some uncertainty about the exact value of the

106 solubility constant of the amorphous Mg-silicates, commonly considered as precursors to the  
107 formation of Mg-silicates in laboratory experiments (i.e. sepiolite, kerolite, stevensite; e.g.  
108 Tosca et al., 2011). Second, Mg-silicates may form either in the water column or in sediments  
109 during early diagenesis (Işık Ece, 1998; Bristow et al. 2009; Milesi et al. 2019, 2020). In the  
110 latter case, they do not reflect the chemistry of the waters in the depositional basin but instead  
111 that in the sediment pore waters. Moreover, several authors have suggested that some  
112 biogeochemical processes may trigger Mg-silicate formation, possibly obscuring the  
113 relationship between the identity and presence/absence of Mg-silicates and bulk chemical  
114 conditions. For instance, recent studies reported Mg-silicates formed by a local pH increase  
115 and/or nucleation on extracellular polymeric substances within microbial mats, acting as  
116 precursors of carbonate precipitation in microbialites (Bontognali et al., 2010; Burne et al.,  
117 2014; Zeyen et al., 2015; Pace et al., 2016). The association of Mg-silicates and biofilms has  
118 also been evidenced in certain speleothems (Melim and Spilde, 2018), mostly in volcanic caves  
119 (Léveillé et al., 2002; Miller et al., 2014). In some cases, Mg-silicates were shown to develop  
120 at the surface of diatom frustules (Badaut and Risacher, 1983; Bentz and Peterson, 2020). Yet,  
121 although Mg-silicates have shown a strong potential for fossilization of microbes and organic  
122 matter (Zeyen et al., 2015), the role of the microorganisms in the formation of these phases  
123 remains unclear.

124

125 Mg-silicates, characterized as poorly crystalline hydrated talc phases similar to kerolite,  
126 were reported in shallow microbialites from numerous alkaline crater lakes in Mexico (e.g.  
127 Atexcac, La Preciosa, Quechulac; Zeyen et al., 2015, 2021). By contrast, the shallow  
128 microbialites of the Mexican Lake Alchichica were essentially composed of hydromagnesite  
129 and aragonite, but did not contain kerolite, possibly because lake water had a low concentration  
130 in orthosilicic acid ( $< 26 \mu\text{M}$ ). Here, we investigated deep sediments in Lake Alchichica in

131 order to check how their mineralogy compared with that of shoreline microbialites. The  
132 mineralogy and chemistry of the first meter of the sediment column were studied together with  
133 the pore water chemistry providing insights on the processes occurring during early diagenesis.

134

## 135 **B) Geological Setting**

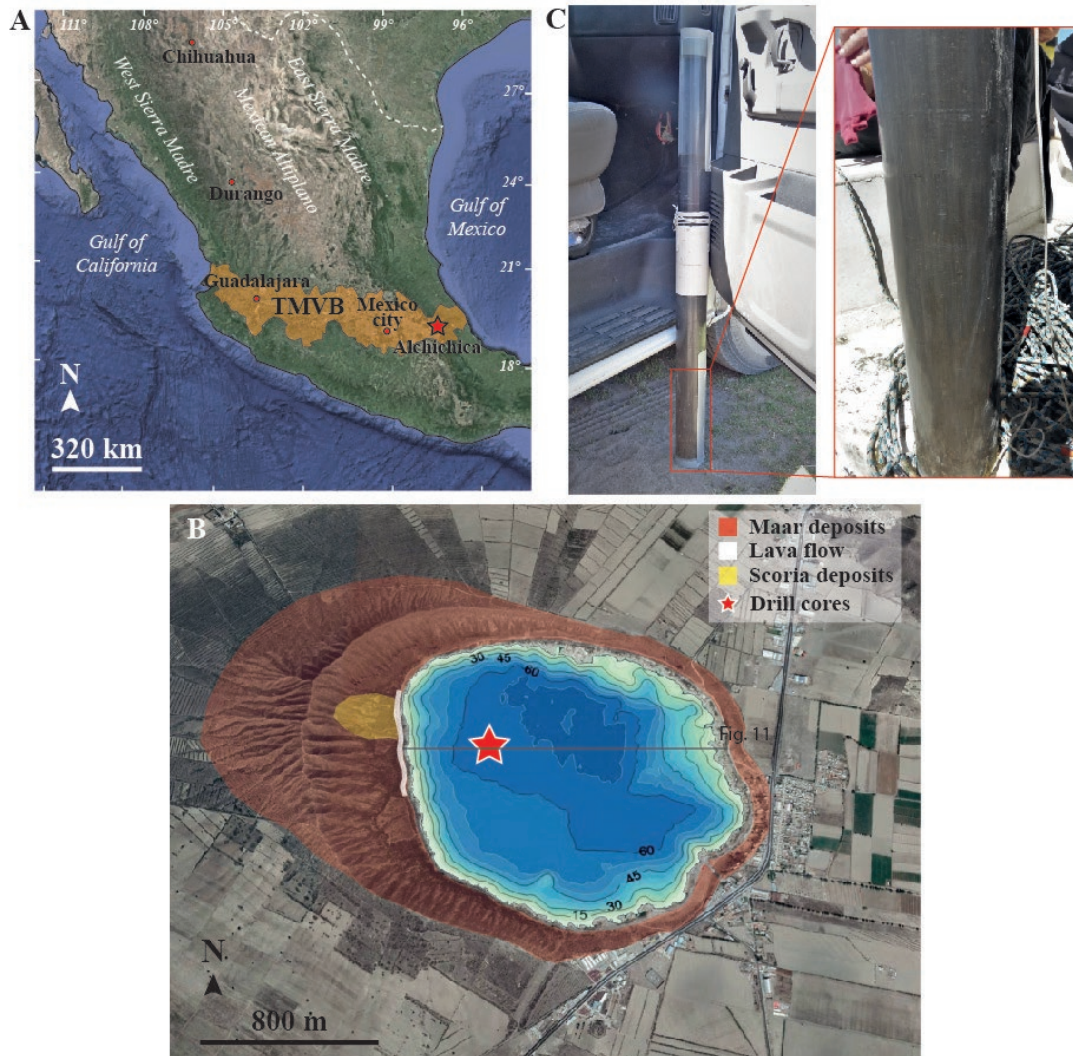
136

137 Alchichica is a crater lake located in the eastern part of the trans-Mexican volcanic belt  
138 (TMVB) in the Cuenca de Oriental region (19°24'N and 97°24'W; **Figure 1a**). The TMVB is  
139 a seismically and volcanically active zone resulting from the subduction of the Rivera and  
140 Cocos plates beneath North America at the Middle American Trench during the Neogene period  
141 (Ferrari et al., 2012). It is composed of diverse volcanic structures formed on a basement  
142 predominately composed of highly folded and faulted Cretaceous limestones and shales as well  
143 as small plutonic intrusions of granodiorite, monzonite, and syenite emplaced during the  
144 Oligocene and the Miocene (Negendank et al., 1985). These basement rocks are covered by  
145 Miocene to Pleistocene andesitic rocks, which are themselves overlain by (i) widespread  
146 pyroclastic deposits derived from both local phreatomagmatic volcanoes and large eruptions  
147 from nearby volcanoes, as well as (ii) lacustrine deposits. The crater of Alchichica is surrounded  
148 by an asymmetric ring of tephra, deposited by several Hawaiian, strombolian and  
149 phreatomagmatic eruptions dated from >114 ka to  $6.3 \pm 5.1$  ka by  $^{40}\text{Ar}/^{39}\text{Ar}$  methods (Tchamabé  
150 et al., 2020). The crater rim is mainly composed of maar deposits (i.e. pyroclastic rocks  
151 including 44 to 73 wt.% of juvenile clasts) and basaltic andesite lava flow and scoriaceous pre-  
152 maar deposits outcropping only on the west side of the crater (**Figure 1**; Tchamabé et al., 2020).

153

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Figure 1: A) Relief map of Mexico representing the location of the lake Alchichica in the trans-Mexican volcanic belt (TMVB, orange area, modified from (Ferrari et al. 2012)); B) Google Earth image of Lake Alchichica showing the surrounding volcanic deposits (from Tchamabé et al., 2020), bathymetrical data (from Alcocer & Oseguera, 2019) and the location of the drill core (red star). C) Pictures of the 85.5 cm long sediment core.

163 With a maximum depth of 65 m (Arredondo-Figueroa et al., 1983; Vilaclara et al., 1993),  
164 Alchichica is the deepest crater lake in Mexico. It is located at 2,320 m a.s.l. on a high-altitude  
165 plateau called Los Llanos de San Juan. The climate is semi-arid with an annual precipitation  
166 regime of less than 400 mm and an annual evaporation rate of about 1700 mm (Alcocer, 2021).  
167 The air temperature seasonally ranges from -5.5 to 30°C (Alcocer and Hammer, 1998). The  
168 lake is supplied by rainfall and groundwaters as suggested by Armienta et al. (2008) for all maar  
169 lakes. The water deficit between evaporation and rainfall could be partially balanced by



170 underground sources and outflows (García Martínez, 2010), yet the lake water level has  
171 decreased to a rate of about 8 cm.yr<sup>-1</sup> for several decades (Alcocer, 2021). Water in Lake  
172 Alchichica is alkaline (pH>9), subsaline (~7.9 g/L) and dominated by Na<sup>+</sup>, Cl<sup>-</sup>, HCO<sub>3</sub><sup>-</sup> and Mg<sup>2+</sup>  
173 ions. Two main factors control its chemical composition: 1) the weathering of surrounding  
174 bedrocks (basalts and Cretaceous limestones) through groundwater supply and 2) the  
175 evaporation of the lake due to the semi-arid climate (Zeyen et al., 2021). Alchichica is a  
176 monomictic lake stratified during most of the year and homogenized during winter (Arredondo-  
177 Figueroa, 2002). Two annual blooms of primary producers were observed by previous studies:  
178 a diatom bloom during the winter mixing (December-January) and a cyanobacteria bloom at  
179 the onset of the stratification period (April-May) (Alcocer et al., 2000; Vilaclara et al., 1993).  
180 Microbialites were extensively studied in Lake Alchichica and are mainly composed of  
181 hydromagnesite and aragonite in varying proportions (Kazmierczak et al., 2011; Couradeau et  
182 al., 2011, 2013; Gérard et al., 2013; Saghaï et al., 2015; Valdespino-Castillo et al., 2018; Zeyen  
183 et al., 2021). They are covered by complex microbial communities dominated by oxygenic and  
184 anoxygenic photosynthesizers (Couradeau et al., 2011; Saghaï et al., 2016; Iniesto et al., 2021).

185

## 186 **C) Material and Methods**

### 187 **a. Sample collection**

188

189 Lake water was sampled with a Niskin bottle in May 2019 in the center of the lake at 0,  
190 5, 10, 20, 30, 40, 50, 55 and 60 m depth. Temperature, pH and salinity were measured *in situ*  
191 with a YSI Exo 2 multi-parameter probe. Hundreds of milliliters of solution were systematically  
192 filtered using 0.22 µm Filtropur S filters. For major cation analysis, 30 mL of solution were  
193 acidified with nitric acid (2%). One hundred and thirty milliliters were used for measurements  
194 of anion and orthosilicic acid (H<sub>4</sub>SiO<sub>4</sub>) concentrations, without pre-acidification. For dissolved

195 inorganic carbon (DIC) analysis, 12 mL were placed in airtight Labco Exetainer® tubes in order  
196 to avoid exchange of DIC with atmospheric CO<sub>2</sub>.

197

198 The sediment core was collected from the bottom of the lake at 61 m in depth during  
199 the same field campaign in May 2019, using a gravitational Uwitec corer with a diameter of 90  
200 mm. The total length of the recovered core was 85.5 cm (**Figure 1c**). The sediment core was  
201 then cut in two in order to allow its processing in a glove-bag. The two cores were transferred  
202 into a glove-bag and placed under anoxic conditions (N<sub>2</sub> atmosphere) immediately after  
203 collection. They were then processed and split into cm-scale fractions along the core's vertical  
204 axis. The porewater was separated from the solid phases once extracted from the core barrel  
205 using Rhizon samplers connected via tubing to syringes (filtration to 0.2 µm). During all the  
206 slicing process, oxygen levels were monitored with a Multi 3630 IDS WTW oxygen meter  
207 (equipped with a FDO 925-3 O<sub>2</sub> optode) and were always below the detection limit of 0.1 mg/L.  
208 The pH of the porewaters were measured (with a Sentix 940 IDS WTW electrode) immediately  
209 after water collection within the glove-bag. Sediments were packaged under nitrogen and  
210 transported in heat sealed airtight bags (PP016 H Protective Packaging Ltd) with O<sub>2</sub> trap  
211 (Anaerocult® A), then fully dried in an anoxic N<sub>2</sub>-filled glove box in the lab.

212

### 213 **b. Chemical analyses of solutions**

214

215 Chemical analyses for major elements (Al, B, Ca, K, Li, Mg, Na, Mn, Si, S, Cl) are  
216 summarized in **Table 1**. They were carried out at the Laboratoire Géoscience Océan (UBO,  
217 Brest, France) using inductively coupled plasma-atomic emission spectroscopy (ICP-AES).  
218 Anion concentrations (F<sup>-</sup>, Cl<sup>-</sup>, Br<sup>-</sup>, and SO<sub>4</sub><sup>2-</sup>) were measured by ion chromatography and  
219 orthosilicic acid (H<sub>4</sub>SiO<sub>4</sub>) concentrations were determined by continuous flow colorimetric

220 analyses at the Institut de physique du globe de Paris (IPGP, Paris, France). The uncertainty on  
221 the concentration measurements of cations, anions and orthosilicic acid was lower than  $\pm 5\%$ .

222 Dissolved inorganic carbon (DIC) concentrations were determined at the IPGP, using  
223 an “Analytical Precision 2003” Gas Chromatograph – Isotope Ratio Mass Spectrometer  
224 (running under He-continuous flow) following the analytical protocol detailed in Assayag et al.  
225 (2006). An aliquot of the water sample was injected into a Labco Exetainer<sup>®</sup> tube pre-loaded  
226 with H<sub>3</sub>PO<sub>4</sub> and He. After the dissolved CO<sub>2</sub> has equilibrated with the head-space gas (for one  
227 night), gases (i.e. He and CO<sub>2</sub>) were sampled and quantified with the GC-IRMS by comparison  
228 with internal standards of known concentrations. Each measurement represents an average of  
229 four injections in the mass spectrometer and all samples were duplicated (including chemical  
230 preparation and analysis). Standard deviation for [DIC] is 1 mM (n=33), on average.

231 Activities of cations, anions and orthosilicic acid as well as saturation indices of the  
232 solutions were calculated using the software PHREEQC 3 (Parkhurst and Appelo, 2013) and  
233 the thermodynamic database “thermoddem.dat” (Blanc et al., 2012).

234

### 235 **c. Bulk analyses of sediments**

236

237 A quantity of about 10 g for each sediment sample was finely ground and homogenized  
238 in an agate mortar and aliquots of the same powdered sample were used for the different bulk  
239 analyses.

240

#### 241 *X-ray powder diffraction*

242 For X-ray diffraction (XRD) analyses, each core sample powder was placed onto 3 cm  
243 diameter sample holders and leveled to form a flat surface for analysis. XRD analyses were  
244 performed using a Panalytical Xpert Pro transmission diffractometer with a cobalt anode

245 operated at 45 kV and 40 mA and a slit of 0.5° at 240 mm radius distance. The 2θ scan was  
246 performed in the continuous mode from 4° to 110° (2θ) with a step of 0.024°.

247 Each diffractogram was corrected for small relative displacements of sample height  
248 using the alignment with major aragonite diffraction peaks (2θ-Co angles 30.52° and 53.77°),  
249 dominant in all samples. The background signal was fit so that the contribution of X-ray low  
250 crystalline phases could be removed from peak analysis. Automatic background determination  
251 did not produce adequate results given the presence of magnesium silicate and opaline silica in  
252 the sample generating broad diffraction signals, but these were used to assess the relative  
253 presence of the two low crystalline materials (**Figure S1**). The background signal was manually  
254 fit with an interpolated spline for each diffractogram. Peak identification was performed using  
255 pattern search on the X-ray diffraction database of reference spectra “Crystallography Open  
256 Database (COD)” (from <http://www.crystallography.net/>).

257 Structural data of identified minerals were obtained from the crystallography open  
258 database (COD). The structure and abundance of crystalline minerals were estimated using a  
259 Rietveld refinement analysis implemented in the Highscore Plus software (**Figure S2**). XRD  
260 identifications were cross-checked back and forth with results from energy dispersive X-ray  
261 spectrometry (EDXS) elemental composition data obtained by scanning electron microscopy  
262 (SEM) to more confidently identify the mineralogy of (core) samples.

263

#### 264 ***Fourier transform infrared spectroscopy analyses***

265 For attenuated total reflectance (ATR) - Fourier transform infrared (FTIR) spectroscopy  
266 analyses, ~1 mg of sediment powder was pressed against a single-reflection diamond prism.  
267 ATR spectra were recorded between 400 and 4000 cm<sup>-1</sup> with a resolution of 1 cm<sup>-1</sup>, using a  
268 Nicolet 6700 FTIR spectrometer equipped with a deuterated triglycine sulfate (DTGS) detector.  
269 Spectra were corrected from the atmospheric signal.

270

271 ***Bulk chemical analyses of sediments***

272 Concentrations of major elements were measured on 30 sediment samples using an ICP-  
273 AES ICap 6500 (Thermo Fischer) after alkali fusion of rock samples with LiBO<sub>2</sub> followed by  
274 dissolution in HNO<sub>3</sub> at the Service d'Analyse des Roches et Minéraux (SARM, Centre de  
275 Recherches Pétrographiques et Géochimiques, Nancy, France). About two grams of ground  
276 powder were used for these analyses. The uncertainties on the major element measurements,  
277 between 1 and 25% depending on their concentrations, were assessed from relative standard  
278 deviations calculated on a minimum of 30 measurements of reference geochemical standards.

279

280 ***Microscopy analyses of sediments***

281 Around 5 mm<sup>3</sup> of powder (<100 µm) from 8 samples (at 0.5, 2, 4, 8.5, 37.3, 41.5, 44.5 and  
282 84.8 cm) collected all along the core length were progressively impregnated in hard grade LR-  
283 white resin (Polysciences, Inc.) by incubating them at 6°C for 18 h in (1:2) then (2:1) mixture  
284 of LR-white/ethanol and finally in pure LR-white resin. After 1 h at room temperature, the resin  
285 was polymerized by incubation at 40°C for 1 h and at 60°C for 24 h. Once polymerization was  
286 completed, inclusions were polished with diamond polishing paper with a grain size down to a  
287 ¼ of a micrometer and rinsed in deionized water with ultrasounds for 10 minutes. Scanning  
288 electron microscopy (SEM) analyses were performed using a Zeiss Ultra 55 field emission gun  
289 (FEG) SEM. Backscattered electron (BSE) images were acquired using an angle selective  
290 backscattered (AsB) detector at an accelerating voltage of 15 kV, a working distance of ~7.5  
291 mm and a 60 mm aperture at high current. The elemental composition of mineral phases was  
292 determined by energy dispersive X-ray spectrometry (EDXS) using an EDS QUANTAX  
293 detector after copper calibration. Semi-quantification of the spectra was achieved using the  
294 ESPRIT software package (Bruker) and the phi-rho-z method.

295 An electron-transparent foil (~100 nm in thickness) was cut by focused ion beam (FIB)  
296 milling from one of the embedded samples, to further analyze by transmission electron  
297 microscopy (TEM) a diatom replaced by Mg-silicates (at 16-19 cm in depth). This was  
298 performed at IMPMC using a Zeiss Crossbeam Neon40. We used the lift-out procedure as  
299 described by Benzerara et al. (2005). A FIB-assisted Pt deposit was first made. A 30 kV Ga+  
300 beam operated at ~5 nA was then used for initial milling, resulting in rough excavation from  
301 both sides of the thin foil. An *in situ* micromanipulator was attached to the foil by FIB-assisted  
302 platinum deposition prior to separation of the foil (at ~100 pA). The thin foil was transferred to  
303 a TEM grid and welded to it. Further thinning of the foil was performed with the beam operated  
304 at a low, ~100 pA current. A last cleaning step was performed at low acceleration tension (~3  
305 kV).

306 The electron-transparent foil was then analyzed by TEM using a JEOL 2100F operating at  
307 200 kV and equipped with a field emission gun (FEG-TEM). Scanning transmission electron  
308 microscopy (STEM) observations were performed in the high-angle annular dark field mode  
309 (HAADF) and a probe size of 1 nm. EDXS mapping was performed using the STEM mode.

310

## 311 **D) Results**

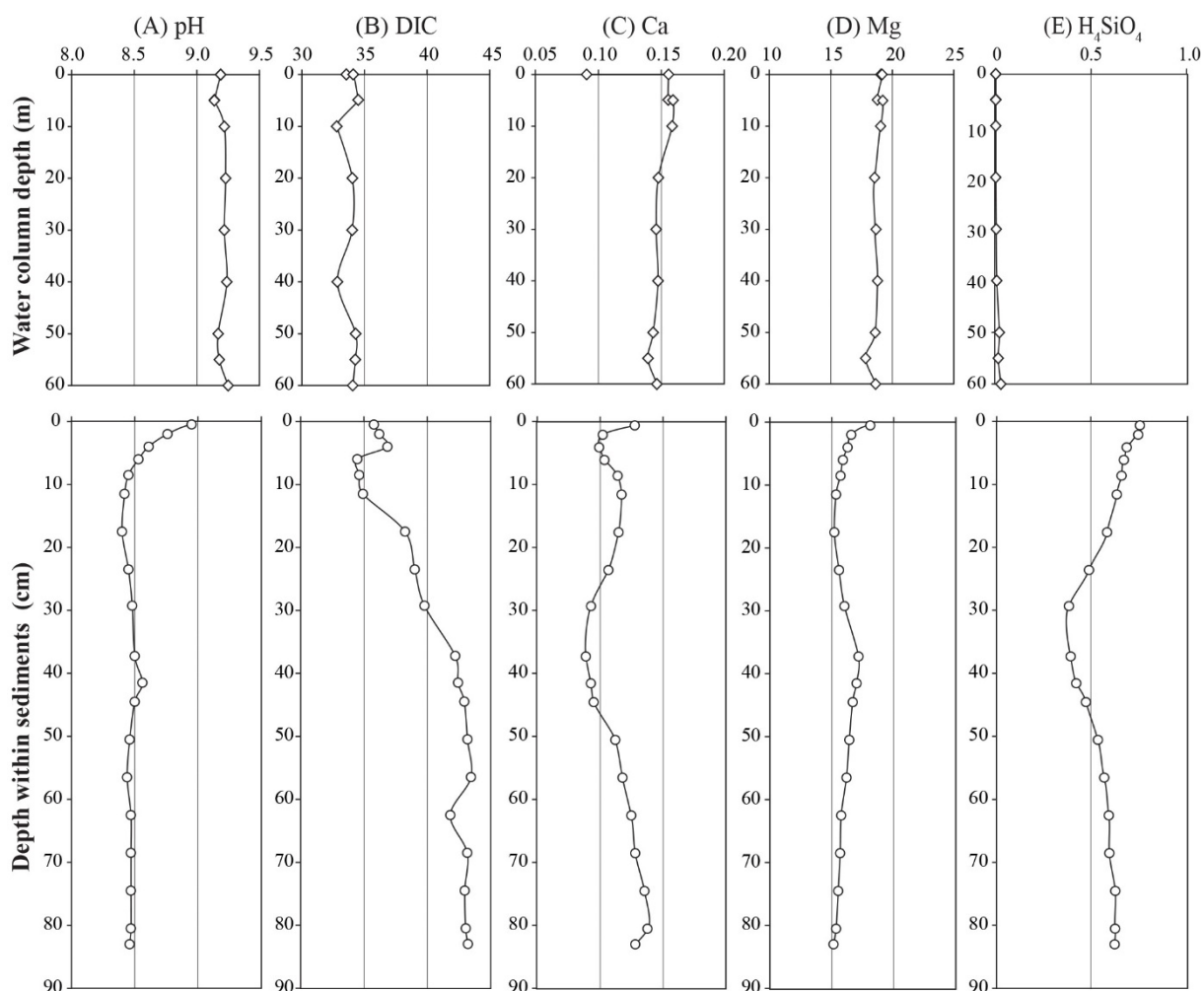
312

### 313 *Solution chemistry of the water column and the sediment pore water*

314

315 Solutions at all depths in the water column of Lake Alchichica are dominated by sodium,  
316 chloride and magnesium (**Figure 2; Table 1**). In the water column, the temperature decreases  
317 from 19.5 to 15.2°C with depth and the chemical composition remains stable, with a pH of 9.2  
318 and concentrations of DIC at 33.9 mM, Mg at 18.8 mM and Ca at 0.1 mM (**Figure 2; Table 1**).  
319 The concentration of orthosilicic acid in the water column is below the detection limit (0.1 μM)

320 from the surface down to 30 m in depth, then increases to 26  $\mu\text{M}$  at 60 m (maximum depth of  
 321 the lake). In the sediments, pore waters are also dominated by sodium, chloride and magnesium  
 322 (**Figure 2; Table 1**). The  $[\text{Mg}]$  decreases in the first 20 centimeters from 18.1 to 15.2 mM and  
 323 then remains constant with a peak at 17.1 mM at 34.5-40 cm depth. The pH also decreases down  
 324 to 8.47 in the first ten centimeters of the sediments and then remains constant. The concentration  
 325 of orthosilicic acid in porewater is significantly higher than in the lake water. It decreases from  
 326 760  $\mu\text{M}$  at 0.5 cm down to 380  $\mu\text{M}$  at 28-40 cm and raises again to 620  $\mu\text{M}$  at 83 cm deep. The  
 327 DIC concentration globally increases from the top to the bottom of the core, from 35.8 mM to  
 328 43.2 mM. The  $[\text{Ca}]$  varies non-monotonously between 0.09 and 0.14 mM (**Figure 2; Table 1**).



329  
 330 **Figure 2: Depth profiles of (A) pH, (B) dissolved inorganic carbon (DIC), (C) Ca, (D) Mg and**  
 331 **(E)  $\text{H}_4\text{SiO}_4$  concentrations in mM along the water column (top part, diamonds) versus in**  
 332 **porewater in the first 83 cm of sediments (bottom part, circles).**  
 333

334 **Table 1.** pH, DIC and major element concentrations measured in lake and pore waters. Relative  
 335 standard deviation is  $\leq 5\%$  for DIC and element concentrations and  $\pm 0.02$  units for pH values.  
 336 Charge balance (Bal.) in % is calculated as  $100 * (\Sigma \text{cations} - \Sigma \text{anions}) / ((\Sigma \text{cations} + \Sigma \text{anions}) / 2)$ .

	Depth	pH	T °C	DIC mM	Cl mM	Na mM	Mg mM	SO <sub>4</sub> <sup>2-</sup> mM	K mM	B mM	Li mM	Ca mM	Mn mM	H <sub>4</sub> SiO <sub>4</sub> mM	Al μM	Bal. %
Lake water																
AL19_0m-ext	0 m	9.19	19.6	34.1	116	108	19.1	12.8	4.96	3.79	0.35	0.16	<DL	<DL	<DL	-14.2
AL19_4.9m	4.9	9.14	19.2	34.5	107	106	18.8	11.8	4.91	3.79	0.35	0.16	<DL	<DL	0.96	-9.5
AL19_5m	5	9.14	19.2		107	108	19.3	11.9	5.01	3.84	0.35	0.16	<DL	<DL	<DL	-8.0
AL19_10m	10	9.22	19.2	32.8	106	107	19.1	11.8	5.00	3.83	0.35	0.16	<DL	<DL	<DL	-7.3
AL19_20m	20	9.23	16.7	34.0	106	105	18.6	11.8	4.90	3.75	0.34	0.15	<DL	<DL	<DL	-10.0
AL19_30m	30	9.22	15.5	34.0	106	105	18.7	11.7	4.90	3.75	0.34	0.15	<DL	0.002	<DL	-9.7
AL19_40m	40	9.24	15.3	32.8	106	106	18.8	11.8	4.87	3.74	0.35	0.15	<DL	0.005	<DL	-9.0
AL19_50m	50	9.17	15.2	34.3	108	104	18.7	12.0	4.82	3.75	0.34	0.14	<DL	0.018	<DL	-12.6
AL19_55m	55	9.18	15.2	34.3	109	100	17.9	12.0	4.62	3.58	0.32	0.14	<DL	0.013	<DL	-16.9
AL19_60m	60	9.25	15.2	34.1	112	104	18.7	12.2	4.84	3.75	0.34	0.15	<DL	0.026	<DL	-14.3
Sediment porewater																
AL19_C2a_0.5cm	0-1 cm	8.95	N.D.	35.8	97.4	109	18.1	9.92	5.05	3.85	0.35	0.13	0.02	0.76	2.01	2.4
AL19_C2a_2cm	1-3	8.76	N.D.	36.2	99.8	105	16.6	9.50	5.03	3.71	0.34	0.10	0.012	0.75	<DL	8
AL19_C2a_4cm	3-5	8.61	N.D.	36.8	99.0	106	16.3	9.21	5.03	3.74	0.34	0.10	0.012	0.69	<DL	5.9
AL19_C2a_6cm	5-7	8.53	N.D.	34.5	99.6	105	15.9	9.09	5.06	3.68	0.33	0.10	0.013	0.67	0.08	7.7
AL19_C2a_8.5cm	7-10	8.45	N.D.	34.6	100.2	104	15.7	8.70	5.06	3.72	0.33	0.11	0.022	0.66	1.16	8.1
AL19_C2a_11.5cm	10-13	8.42	N.D.	34.9	98.9	105	15.4	8.26	5.08	3.77	0.33	0.12	0.028	0.64	0.45	8.8
AL19_C2a_17.5cm	16-19	8.4	N.D.	38.2	98.2	103	15.2	8.06	5.02	3.71	0.32	0.11	0.036	0.59	0.47	10.2
AL19_C2a_23.5cm	22-25	8.45	N.D.	39.0	96.2	103	15.6	7.40	5.02	3.74	0.31	0.11	0.011	0.49	<DL	8.1
AL19_C2a_29.3cm	28-30.5	8.48	N.D.	39.8	94.6	100	16.0	7.00	4.96	3.68	0.30	0.09	0.011	0.38	<DL	9.1
AL19_C2a_37.3cm	34.5-40	8.5	N.D.	42.2	93.2	102	17.1	6.16	5.06	3.80	0.30	0.09	0.011	0.39	<DL	4.4
AL19_C2a_41.5cm	40-43	8.56	N.D.	42.4	94.4	100	17.0	5.56	4.93	3.62	0.29	0.09	0.011	0.42	<DL	6.1
AL19_C2b_44.5cm	43-46	8.5	N.D.	42.9	96.6	99.4	16.7	5.62	4.89	3.59	0.29	0.09	0.012	0.47	0.44	8.9
AL19_C2b_50.5cm	49-52	8.46	N.D.	43.2	97.2	99.4	16.4	5.47	4.87	3.57	0.28	0.11	0.012	0.54	0.34	9.6
AL19_C2b_56.5cm	55-58	8.44	N.D.	43.4	95.3	97.8	16.2	5.13	4.81	3.56	0.28	0.12	0.013	0.57	<DL	8.5
AL19_C2b_62.5cm	61-64	8.47	N.D.	41.8	95.4	97.0	15.8	4.84	4.79	3.52	0.27	0.12	0.013	0.59	0.27	10.2
AL19_C2b_68.5cm	67-70	8.47	N.D.	43.1	94.6	96.3	15.7	4.90	4.71	3.48	0.27	0.13	0.002	0.60	0.50	10.3
AL19_C2b_74.5cm	73-76	8.47	N.D.	43.0	96.7	96.2	15.5	4.49	4.74	3.50	0.26	0.14	0.002	0.63	0.71	11.4
AL19_C2b_80.5cm	79-82	8.47	N.D.	43.0	94.2	95.6	15.3	4.20	4.68	3.52	0.26	0.14	0.002	0.63	0.31	10.3
AL19_C2b_83cm	82-84	8.46	N.D.	43.2	93.8	95.9	15.1	4.09	4.65	3.48	0.26	0.13	0.002	0.62	0.03	2.1

337

338

339 Saturations of the solutions with respect to authigenic silicate and silica phases are best

340 visualized in a diagram, where the logarithm of the  $a(\text{Mg}^{2+})/a(\text{H}^+)^2$  activity ratio is plotted

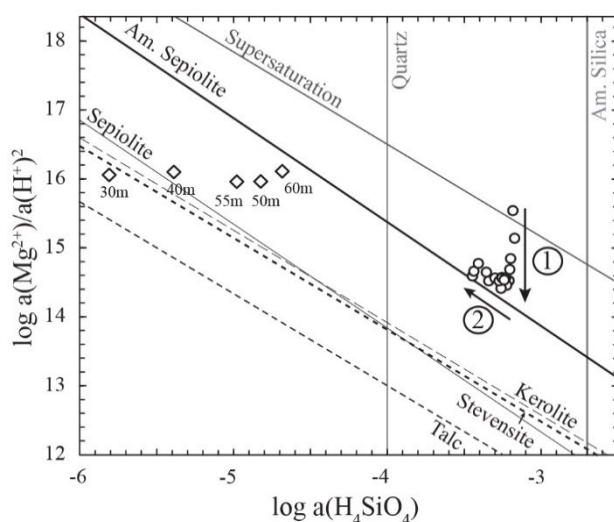
341 against the logarithm of the orthosilicic acid activity ( $a(\text{H}_4\text{SiO}_4)$ ) (**Figure 3**). As the orthosilicic

342 acid concentration increases with depth in the water column, the lake water reaches saturation

343 with respect to several low-temperature Mg-silicates such as sepiolite, kerolite, talc and



344 stevensite below 30 m in depth. However, all lake waters down to 60 m remain undersaturated  
 345 with respect to “amorphous sepiolite”, as defined by Wollast et al. (1968). By contrast, all  
 346 sediment pore solutions are saturated/supersaturated with respect to “amorphous sepiolite”.  
 347 Two trends can be detected: (1) from 0 to 11.5 cm in depth, pore waters align along a vertical  
 348 trend, i.e. at a constant  $a(\text{H}_4\text{SiO}_4)$ , located between the solubility lines of quartz and amorphous  
 349 silica (arrow 1 in **Figure 3**); (2) from 11.5 to 84.8 cm, pore waters follow a trend parallel to the  
 350 “amorphous sepiolite” solubility line (arrow 2 in **Figure 3**). Only one sample of pore water  
 351 from the top of the sediments (0-1 cm) lies above the critical supersaturation line for  
 352 homogeneous nucleation of Mg-silicates as defined by Tosca et al. (2011) and Tosca and  
 353 Masterson (2014).



354  
 355 **Figure 3: Solubility diagram in the  $\log[a(\text{Mg}^{2+})/a(\text{H}^+)^2] - \log[a(\text{H}_4\text{SiO}_4)]$  space determined**  
 356 **at 25°C.** Lake (diamond) and pore waters (dots) of Alchichica are plotted against the solubility  
 357 lines of talc (Jones and Galan, 1988), stevensite (Chahi et al., 1997), kerolite and sepiolite  
 358 (Stoessel, 1988), amorphous sepiolite (Wollast et al., 1968), quartz and amorphous silica  
 359 (Truesdell and Jones, 1974). The “supersaturation” line results from experiments conducted by  
 360 Tosca et al. (2011) and Tosca and Masterson (2014) and relates to homogeneous nucleation of  
 361 Mg-silicate phases from solution. The arrows 1 and 2 show the main trends followed by  
 362 Alchichica sediment pore waters respectively between 0 and 11.5 cm and between 11.5 and  
 363 84.8 cm. Above 30 m in the lake waters, silica is below the detection limit and hence data are  
 364 not represented in this diagram.

365  
 366 Saturations of solutions with respect to several carbonate phases are visualized in other  
 367 diagrams, where the logarithms of  $a(\text{Ca}^{2+})$  or  $a(\text{Mg}^{2+})$  are plotted against the log of  $a(\text{CO}_3^{2-})$

368 (Figure S3). These diagrams clearly show that lake and pore waters are supersaturated with  
369 respect to calcite and aragonite at all depths but only lake waters reach saturation with  
370 monohydrocalcite (MHC; Figure S3a). The water column of Lake Alchichica is oversaturated  
371 at all depths with amorphous Mg carbonate (AMC) as defined by Fukushi and Matsumiya  
372 (2018) (Figure S3b) and therefore with hydromagnesite as well. All pore waters but those  
373 between 8.5 and 23.5 cm, are saturated or supersaturated with hydromagnesite.

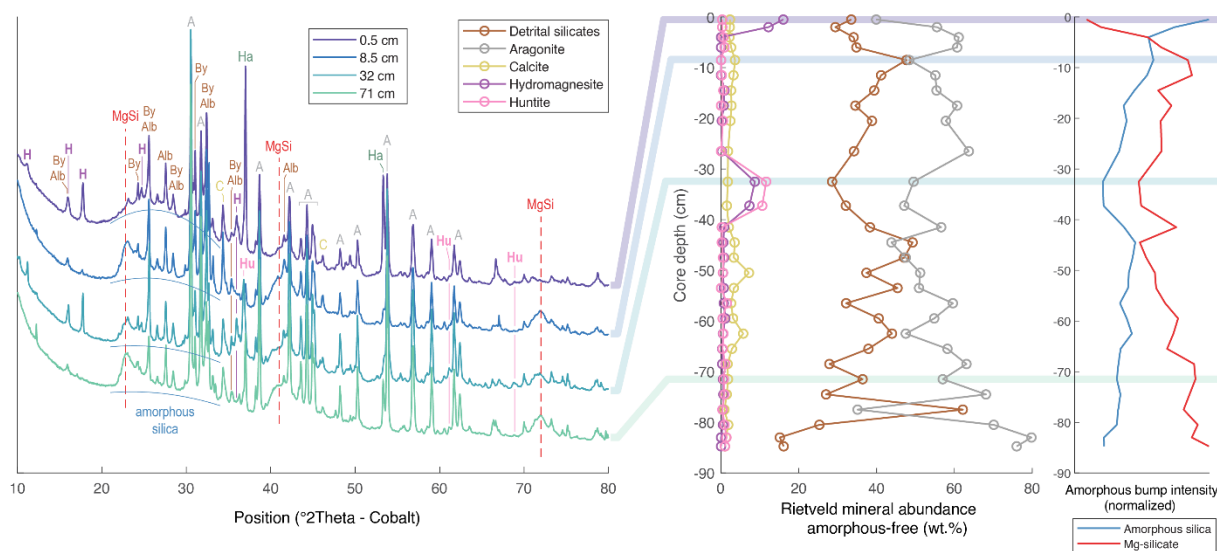
374

### 375 *Bulk mineralogy*

376

377 X-ray diffraction analyses highlight that aragonite is the dominant mineral species in  
378 the sediments with an abundance varying between 40 and 80 wt.% (amorphous-free) as  
379 estimated by Rietveld (Figures 4 and S4, Table S1). Detrital silicate minerals (albite,  
380 bytownite, quartz and pigeonite) are also detected in all diffractograms, their total abundance  
381 varying from 15 to 60 wt.% (amorphous-free). The relative abundance of aragonite and detrital  
382 species are anticorrelated to some degree along the core, but other mineral phases exhibit key  
383 variations with depth. In particular, substantial amount of hydromagnesite is identified in the  
384 first two samples near the sediment surface (up to 15 wt.% in the top 3 cm of the core), but is  
385 absent at depth (< 1 wt.%) except within a specific interval near 35 cm in depth where  
386 hydromagnesite is clearly detected (~ 8 wt.%) along with huntite (~ 11 wt.%), a trigonal  
387 carbonate phase of formula  $Mg_3Ca(CO_3)_4$ . Poorly crystalline magnesium silicates and opaline  
388 silica can also be detected by XRD (Figure S1). The estimation of the intensity of the broad x-  
389 ray scattering signal related to silica reveals that it is most intense near surface but significantly  
390 decreases within the sediment below, with the lowest signal near 35 cm. By contrast, the  
391 magnesium silicate, identified as a kerolite or a stevensite based on broad diffraction peaks at  
392  $22.7^\circ$ ,  $41.2^\circ$  and  $71.8^\circ$   $2\theta$  angles (see Zeyen et al., 2015 for more details about XRD

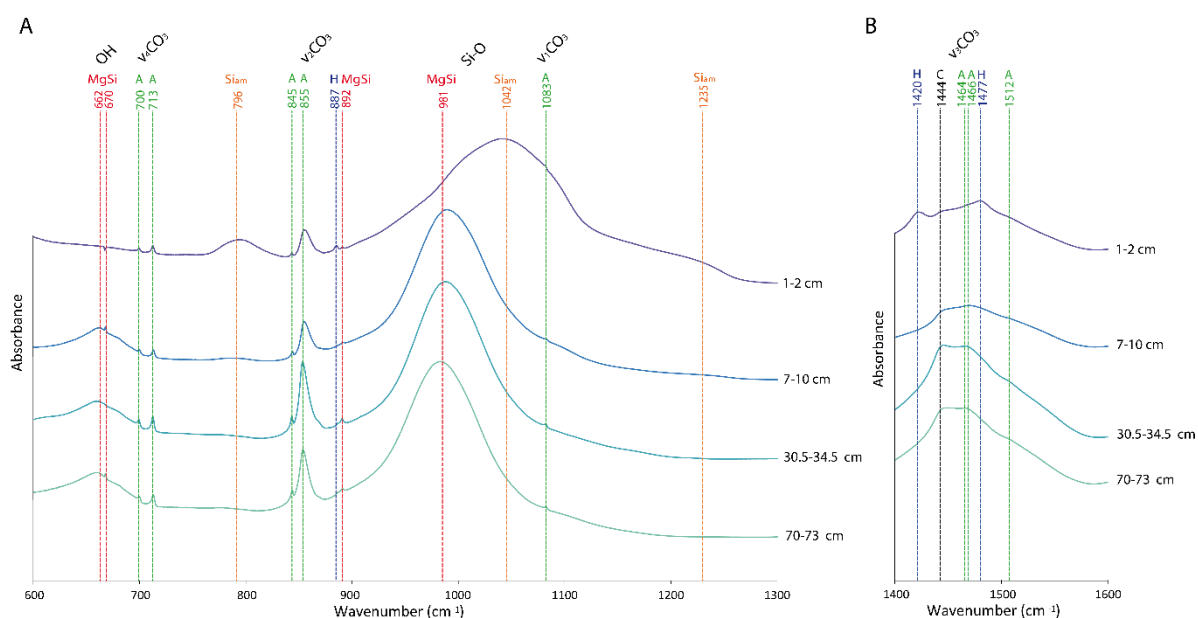
393 identification), is not detected in the first two samples at the surface but clearly detected deeper  
 394 on, with a low relative intensity near 35 cm. Other minor minerals are detected by XRD in all  
 395 samples, such as calcite (average abundance of 2.6 wt.% as suggested by Rietveld), and halite,  
 396 which likely formed when the samples containing Cl-rich pore waters were dried.



397  
 398 **Figure 4: X-ray diffraction analyses of Alchichica sediment core samples. (Left panel)**  
 399 Selected diffraction patterns (at 0.5 cm, 8.5 cm, 32.5 cm and 71.5 cm depth) highlighting the  
 400 key crystalline mineral assemblages used for Rietveld refinement analysis (A: aragonite, Alb:  
 401 albite, By: bytownite, H: hydromagnesite, Hu: huntite, Ha: halite, C: calcite, see **Figure S4** for  
 402 all patterns), along with the broad diffraction signals of poorly crystalline magnesium silicate  
 403 (MgSi, kerolite or stevensite) and amorphous silica. (**Middle panel**) Estimated crystalline  
 404 mineral relative abundances and (**Right panel**) relative intensity of XRD signals affiliated to  
 405 poorly crystalline phases (see also **Figure S1**) in the core with depth.  
 406

407 FTIR analyses of the sediment samples show an evolution of their mineralogical  
 408 composition with depth in perfect agreement with XRD analyses (**Figures 5, S4 and S5**). The  
 409 aragonite signal is present in all samples at all depths (**Figure S5**), as attested by bands at 700,  
 410 713, 845, 855, 1083, 1464 and 1512  $\text{cm}^{-1}$  (Zeyen et al., 2015). In surface sediment spectra  
 411 (down to 10 cm), bands at 796, 1059 and 1230  $\text{cm}^{-1}$ , characteristic of amorphous silica (Nied et  
 412 al., 2015; d’Espinose de la Caillerie et al., 1995) are detected as well as bands at 887, 1420 et  
 413 1477  $\text{cm}^{-1}$ , characteristic of hydromagnesite (Farmer, 1974). These bands disappear in deeper  
 414 samples except for the 37.3 cm depth. By contrast, Mg-silicates (e.g., kerolite or stevensite) are

415 absent in surface samples but detected in FTIR spectra of all sediments deeper than 10 cm as  
 416 shown by the presence of OH absorption bands at 662 and 670  $\text{cm}^{-1}$  (Tosca & Masterson, 2014),  
 417 Si-OH stretching band at 892  $\text{cm}^{-1}$  and Si-O stretching band at 981  $\text{cm}^{-1}$  (Zeyen et al., 2015).  
 418 Only the sediment samples at 34.5-40 cm in depth appeared anomalous in that trend: in  
 419 particular it notably shows the reappearance of hydromagnesite as attested by the bands at 887  
 420 and 1420  $\text{cm}^{-1}$ , but also the presence of huntite as attested by the bands at 870, 878, 891, 1442,  
 421 1511 and 1551  $\text{cm}^{-1}$  (Kangal et al., 2005, Hollingbery and Hull, 2012).

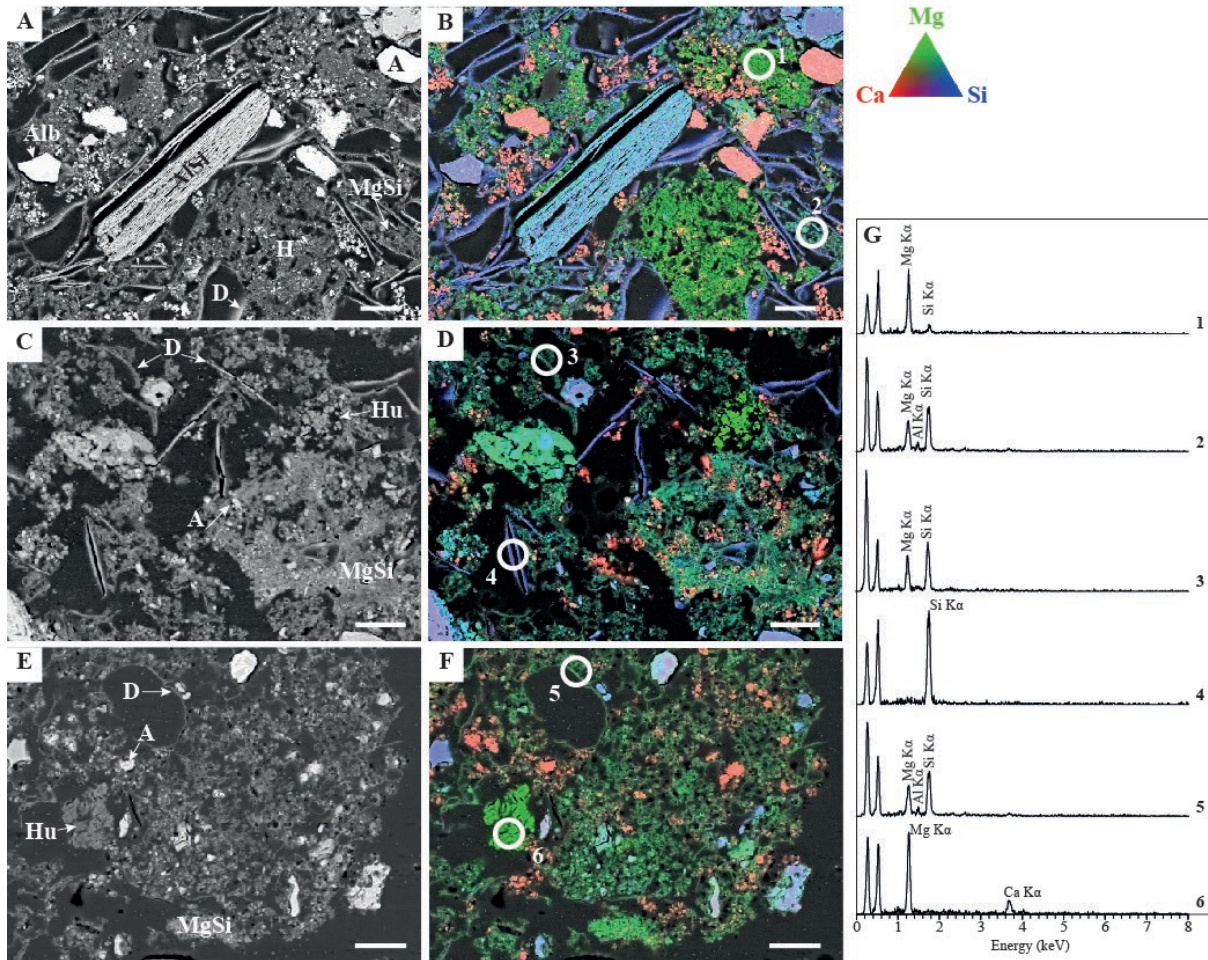


422  
 423 **Figure 5: Selected FTIR spectra of Alchichica sediment samples (at increasing depth from**  
 424 **top to bottom, same depths as those shown in Fig. 4).** (A) Spectra in the 600 to 1300  $\text{cm}^{-1}$   
 425 wavenumber range. (B) Spectra of the same samples from 1400 to 1600  $\text{cm}^{-1}$ . The dashed lines  
 426 show characteristic bands of silicates [red for authigenic magnesium silicates (MgSi), orange  
 427 for amorphous silicates (Si<sub>am</sub>)] and carbonates [green for aragonite (A), black for calcite (C),  
 428 blue for hydromagnesite (H)] in the samples.  
 429

430  
 431 Observations by scanning electron microscopy and EDXS analyses of 8 samples also  
 432 confirm the evolution of the mineralogical composition of the sediments with depth in total  
 433 agreement with FTIR and XRD data. SEM-EDXS analyses show that the sediments contain  
 434 abundant fine-grained Ca-carbonates chemically consistent with aragonite at all depths. By

435 contrast, Mg-rich non-silicate minerals (containing Mg, O and C only) are detected only in the  
436 first ten centimeters of sediments and at 34.5-40 cm and likely correspond to hydromagnesite  
437 as supported by XRD and FTIR analyses (**Figure 6**). A Ca- and Mg-rich mineral interpreted as  
438 huntite based on a Mg/Ca ratio of about 3 is locally observed in samples C2a-05 (7-10 cm),  
439 C2a-10 (34.5-40 cm), C2a-11 (40-43 cm) and C2b-12 (43-46 cm). Huntite grains are relatively  
440 small, measuring less than 20  $\mu\text{m}$  in size. Regarding Si-rich mineral phases, various grains are  
441 best interpreted as detrital minerals. The stoichiometry of these grains together with XRD  
442 analyses allowed to identify them as alkaline feldspars and quartz (**Table S2**). Other detrital  
443 minerals rich in Mg, Al, Si, and K show elemental Si/Al and Mg/Al ratios consistent with  
444 olivine and pyroxenes (**Table S2**). Last, SEM images show that diatom frustules are very  
445 abundant in the sediments at all depths, most of them are broken (and potentially degraded  
446 during sample preparation) but show a nicely preserved fine ultrastructure with e.g., visible  
447 nanopores (Figures 6-8). However, notable chemical differences are observed depending on  
448 depth in the sediments. In the first ten centimeters, the frustules are composed of silica.  
449 Morphologically well-preserved frustules become less abundant below 10 cm depth and the  
450 silica composing them gradually transforms to a poorly crystalline Mg-silicate phase, as shown  
451 by SEM- and TEM-EDXS as well as SAED (**Figures 6, 7 and 8**). Despite this  
452 chemical/mineralogical transformation, the ultrastructure sometimes remains finely preserved,  
453 showing that this mineralogical transformation is pseudomorphic. The average Mg/Si ratio of  
454 this phase as analyzed by EDXS is  $0.74 \pm 0.18$  (n=12), which is consistent with kerolite  
455  $[\text{Mg}_3\text{Si}_4\text{O}_{10}(\text{OH})_2 \cdot \text{H}_2\text{O}]$  and stevensite  $[(\text{Ca},\text{Na})_x\text{Mg}_{3-x}(\text{Si}_4\text{O}_{10})(\text{OH})_2 \cdot n\text{H}_2\text{O}]$ . The stoichiometry  
456 estimated more finely as  $\text{Na}_{0.12}\text{K}_{0.009}\text{Ca}_{0.31}\text{Fe}_{0.2}\text{Al}_{0.15}\text{Mg}_{2.65}\text{Si}_4\text{O}_{10}(\text{OH})_2 \cdot n\text{H}_2\text{O}$  is more  
457 consistent with Al-poor stevensite (Al =  $0.19 \pm 0.26$  at.%, n=12). Some grains of this phase are  
458 also observed away of the diatoms and not just replacing them. In few cases, Al-containing Mg-  
459 silicates were detected but mostly in the vicinity of detrital grains in Lake Alchichica sediments

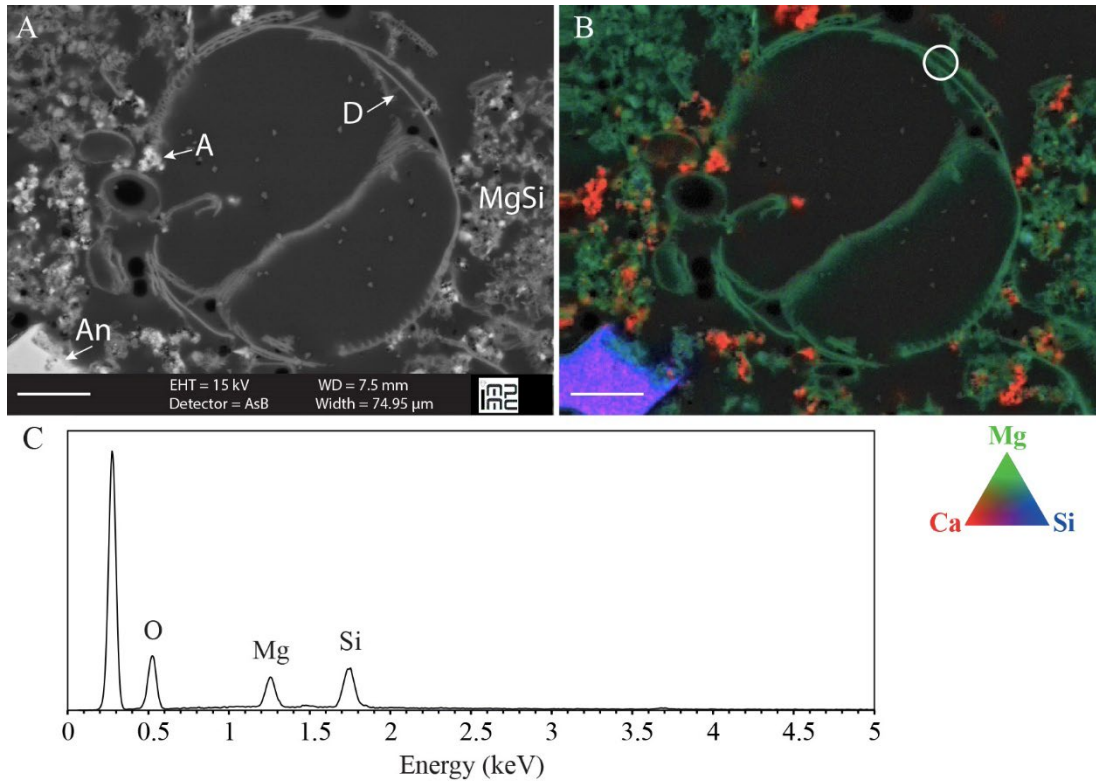
460 (Al = 2.41±0.85 at.%, n=20; **Figure S6**). In the literature, up to 1 wt.% of Al is commonly  
 461 reported in sepiolite, kerolite and stevensite (e.g. Stoessell, 1988; Capet, 1990), sometimes  
 462 reaching up to 5 wt.% (e.g. Rhouta et al. 2008 and references therein).



463  
 464 **Figure 6: Evolution of the sediment mineralogy with depth.** Backscattered electron (BSE)  
 465 images (left) and corresponding EDXS maps (right) of Mg (green), Ca (red) and Si (blue) at 1-  
 466 3 cm (A, B), 7-10 cm (C, D) and 34.5-40 cm depth in the sediment core (E, F). (G) EDXS  
 467 spectra of areas circled in white. On the chemical composition maps, aragonite appears in red,  
 468 silica in blue, huntite and hydromagnesite in light green and Mg-silicates in dark green.  
 469 Hydromagnesite is mainly observed at 1-3 cm (spectra 1), whereas huntite is only detected at  
 470 7-10 cm and 34.5-40 cm (spectra 6). Mg-silicates with very low amount of Al (0.19±0.26 at.%  
 471 in average, n=12) are detected in all the samples (spectra 2). Diatoms are progressively  
 472 transformed into Mg-silicate. At 1-3 cm depth, they are mostly composed of silica, whereas at  
 473 7-10 cm depth, some are transformed into Mg-silicates (spectra 3) while others are not (spectra  
 474 4). Then, at 34.5-40 cm depth, all diatoms are replaced by Mg-silicates (spectra 5). Scale bars  
 475 correspond to 20 μm. H: hydromagnesite; MgSi: Mg-silicate; D: diatom; A: aragonite; Hu:  
 476 huntite; AlSi: detrital Al-silicate (containing Al, Si, Fe, K, Na, Mg).

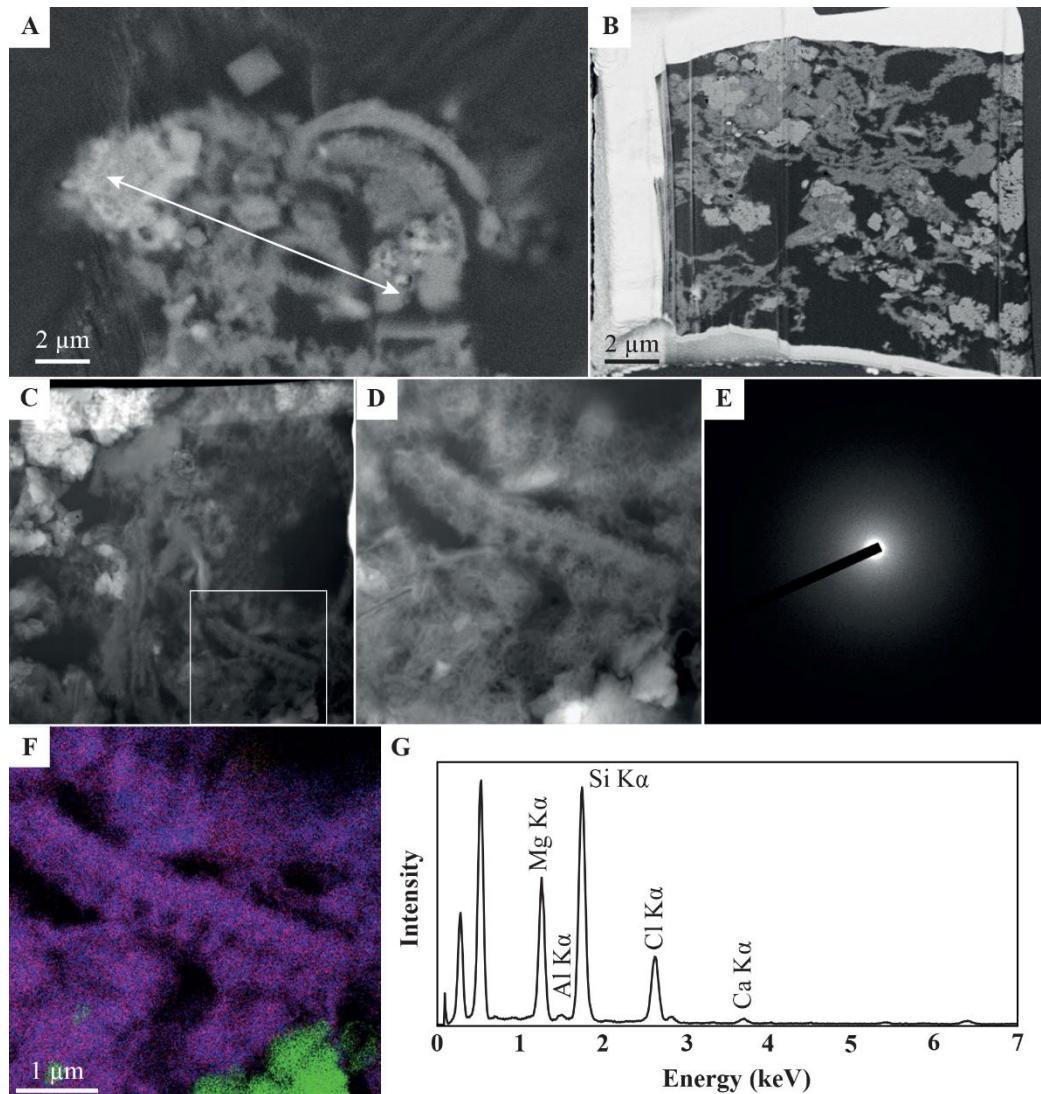
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478



479

480 **Figure 7: Diatom composed of Mg-silicates at a 41.5 cm depth in the sediment core.** (A)  
 481 Backscattered electron (BSE) image, (B) corresponding EDXS map of Mg (green), Ca (red)  
 482 and Si (blue) and (C) EDXS spectra of the zone highlighted by a white circle. On the chemical  
 483 composition map, aragonite appears in red, Mg-silicates in dark green and anorthite in pink-  
 484 blue (at bottom left corner). Scale bars correspond to 10  $\mu$ m. An: anorthite, A: aragonite, D:  
 485 diatom, MgSi: Mg-silicate.  
 486



487

488 **Figure 8: Scanning transmission electron microscopy (STEM) analyses of a diatom**  
 489 **pseudomorphosed into Mg-silicates.** (A) Scanning electron microscopy image showing  
 490 sediment particles collected at 16-19 cm and embedded in resin. The white line indicates where  
 491 the FIB section was excavated. (B) Image of the FIB foil. Aragonite appears in bright and Mg-  
 492 silicates in grey. (C) STEM image in high-angle annular dark field mode (HAADF) showing  
 493 sediment particles. (D) Zoom showing a diatom fragment with a fibrous texture. (E) Electron  
 494 diffraction pattern measured on the diatom fragment, indicating that it is amorphous. (F)  
 495 Overlay of magnesium (red), silicon (blue) and calcium (green) elemental maps as measured  
 496 by energy dispersive x-ray spectrometry (EDXS). (G) EDXS spectrum obtained on the diatom  
 497 fragment showing major Mg and Si peaks. Cl is attributed to resin. A small Al peak can be  
 498 observed.

499

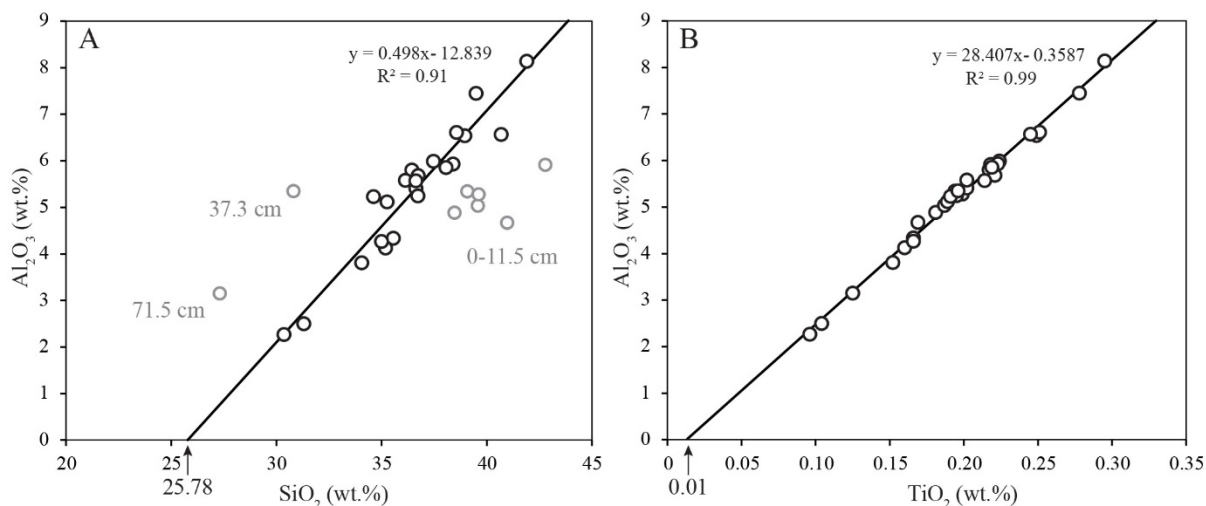
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501 *Bulk chemistry of the sediment solid phases*

502



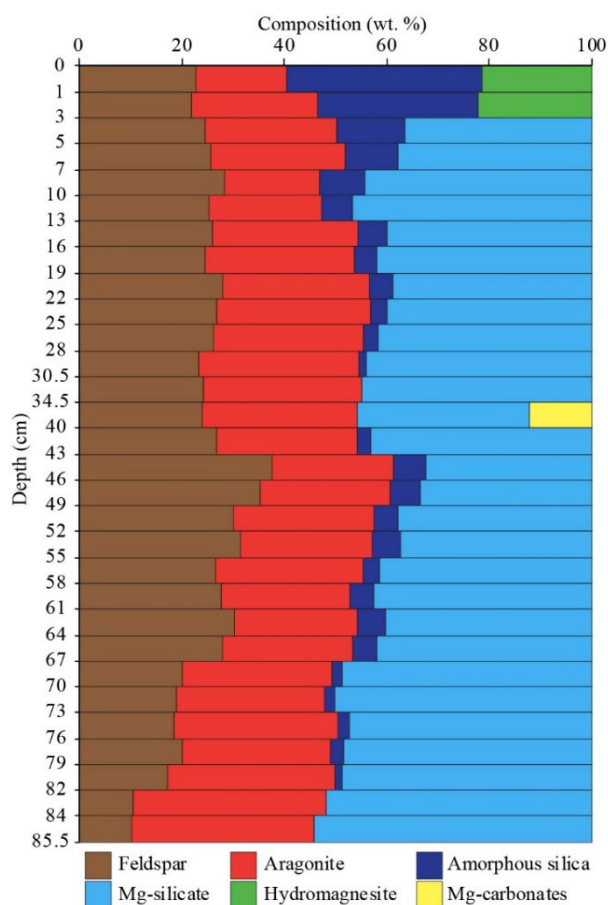
503 The bulk chemical composition of the sediments mostly consists of silicon, calcium,  
 504 magnesium and aluminum (**Table 2**). Titanium content shows a strong positive correlation with  
 505 aluminum with an y-intercept of about 0 in all sediments interpreted as a result of the shared  
 506 detrital origin of these two elements (**Figure 9b**). SiO<sub>2</sub> and Al<sub>2</sub>O<sub>3</sub> also correlate positively for  
 507 most of the samples suggesting that Al is mostly found in silicates (**Figure 9a**). However, since  
 508 [SiO<sub>2</sub>] = 25.78 wt.% when [Al<sub>2</sub>O<sub>3</sub>] = 0, phases other than detritic Al-containing silicates  
 509 contribute massively to the total concentration of silica in the sediments. Based on all  
 510 mineralogical analyses shown above, we suggest that those phases are the amorphous silica of  
 511 diatoms and/or authigenic Mg-silicates, depending on the depth in the core. Moreover, **Figure**  
 512 **9** suggests that the top sediments (from 0 to 11.5 cm) are enriched in non-detrital silica possibly  
 513 due to the accumulation of diatoms, whereas samples at 37.3 and 71.5 cm are depleted in non-  
 514 detrital silica, in agreement with FTIR and XRD data.



515  
 516 **Figure 9:** Correlation between bulk concentrations of (A) Al<sub>2</sub>O<sub>3</sub> and SiO<sub>2</sub> and (B) Al<sub>2</sub>O<sub>3</sub> and  
 517 TiO<sub>2</sub> in the sediments of Alchichica. Symbolized by grey dots, the top sediments (from 0 to  
 518 11.5 cm) are enriched in non-detrital silica possibly due to the accumulation of diatoms,  
 519 whereas samples at 37.3 and 71.5 cm are depleted in non-detrital silica, in agreement with FTIR  
 520 and XRD data.

521  
 522  
 523

524 The semi-quantitative evolution of the mineral composition with depth was assessed  
 525 based on bulk chemical analyses of the sediments and relying on XRD and FTIR analyses  
 526 (Figure 10, Table S3). The feldspars (albite and bytownite) and aragonite contents are  
 527 relatively constant along the core, between 40 and 60 wt.%. The uppermost sediments (0–3 cm)  
 528 also show more than 30 wt.% of amorphous silica and 20 wt.% of hydromagnesite. Then,  
 529 amorphous silica decreases with depth down to 3 wt.% in average, while Mg-silicates  
 530 accumulate to reach up to 40 wt.% on average. As hydromagnesite is not detected either by  
 531 XRD (Figure S4), FTIR (Figure S5) or SEM (Figure 6) below 3 cm in depth, it is no longer  
 532 considered in the calculations. Between 30 and 40 cm in depth, sediments are devoid of silica.  
 533 Concurrently, between 35 and 40 cm, they show a lower Mg-silicates content (34 wt.%) and a  
 534 relatively high amount of Mg carbonates (10 wt.% of huntite and hydromagnesite).



535  
 536 **Figure 10:** Estimated mineral content of the sediment (wt. %) as function of depth based on  
 537 bulk chemical analyses. Minerals stoichiometry used for the quantification of mineral phases is  
 538 given in Table S3.

539

540 **E) Discussion**

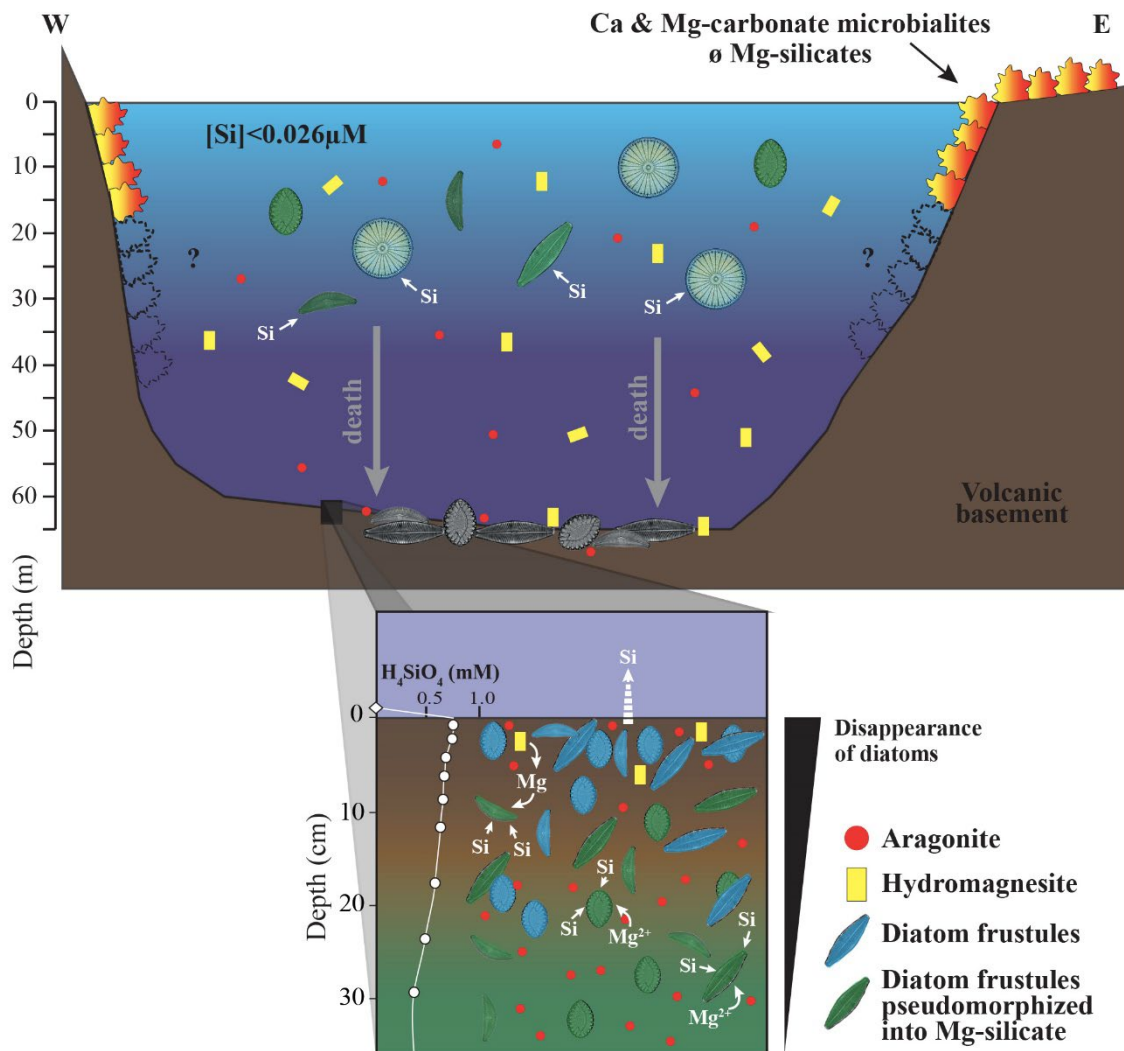
541

542 **a) Massive formation of Mg-silicates in the sediments of Lake Alchichica**

543

544 The uppermost sediments (0-3 cm) are dominated by hydromagnesite, aragonite, diatom  
545 frustules which settled down the water column, detrital alkaline feldspars and quartz inherited  
546 from the erosion of the surrounding volcanic rocks (**Figures 10 and 11**). The presence of  
547 aragonite and hydromagnesite in the first centimeters of sediment suggests that they precipitated  
548 in the water column or at the sediment–water interface, where solutions are oversaturated with  
549 all calcium and magnesium carbonate phases, including their potential precursors, i.e.,  
550 amorphous phases (Zeyen et al., 2021). This is also consistent with the fact that these phases  
551 are found in shallow microbialites populating Lake Alchichica (Zeyen et al., 2021). However,  
552 below ~3 cm in depth, hydromagnesite rapidly disappears, and diatom frustules are  
553 progressively replaced by Al-poor Mg-silicates with a composition corresponding to stevensite,  
554 as evidenced by SEM-EDXS and STEM (**Figures 6-8**). These Mg-silicates then rapidly become  
555 a dominant phase detected by XRD, FTIR and SEM-EDXS (**Figures 4-6**), and represent  
556 between 30 and 53 wt.% of the sediment content at all depths below ~3 cm as calculated from  
557 the bulk chemistry and relying on FTIR and XRD results (**Figure 10**). Thereafter, we explain  
558 how this massive amount of Mg-silicates results from a diagenetic process occurring within the  
559 sediments and not past variations of the nature of the mineral phases settling down in the water  
560 column.

561



562

563 **Figure 11: Summarizing sketch of the mineralogical evolution of the first centimeters of**  
 564 **sediments in Lake Alchichica compared to that of the shallow microbialites populating**  
 565 **lake waters.** In Lake Alchichica, microbialites form at low concentrations of  $H_4SiO_4$  in the lake  
 566 because the living diatoms are an active  $SiO_2$  sink. They are mainly composed of aragonite (in  
 567 red) and hydromagnesite (in yellow). In contrast, in the sediments, hydromagnesite rapidly  
 568 disappears and diatom frustules (in blue) are progressively pseudomorphized by Al-poor Mg-  
 569 silicates, which become a dominant phase (in green). Therefore, the authigenesis of Mg-  
 570 silicates in the sediments is mainly controlled by the spontaneous dissolution of diatom frustules  
 571 which reduces the diffusion rate of  $H_4SiO_4$  from the sediments to the water column.  
 572

573 Large amounts of authigenic Mg-silicates in modern environments are rarely described in  
 574 the literature. Badaut and Risacher (1983) and more recently Bentz and Peterson (2020)  
 575 reported authigenesis of Mg-silicates from diatom frustules (from 25 to 100% of the total  
 576 diatom content) in the sediments of Bolivian salars. High amounts of Mg-silicates ( $\sim 296 \mu\text{mol}$   
 577  $Si \text{ g}^{-1}$  sediments) were also described in the Amazon delta sediments (Michalopoulos and Aller,

1995, 2004) and in Lake Dziani Dzaha (Mayotte), including Al-poor Mg-silicate associated with shallow microbialites and up to 20 wt.% of Mg-saponite in the sediments (Gérard et al., 2018; Milesi et al., 2019, 2020). Interestingly, the high proportion of Mg-silicates in Lake Alchichica sediments contrasts with the absence of authigenic Mg-silicates in Alchichica shallow microbialites (Zeyen et al., 2019). Zeyen et al. (2021) systematically assessed the presence/absence of authigenic Mg-silicates in microbialites from 10 Mexican alkaline crater lakes. They showed that microbialites from most lakes contained Mg-silicates except those in Lake Alchichica. The authors suggested that this difference may result from the very low orthosilicic acid concentration in Lake Alchichica ( $64 \pm 94 \mu\text{M}$ ) compared with other lakes (e.g., 1.10 and 0.58 mM in Lakes Atexcac and La Preciosa, respectively; Zeyen et al., 2021), and therefore that water was not saturated enough to allow spontaneous precipitation. They also speculated about the potential causes for a lower acid orthosilicic acid concentration in the water column of Lake Alchichica, mentioning variations in the nature of the weathered protolith and the active extraction of orthosilicic acid by diatoms, similar to what has been shown in oceans (Conley et al., 2017). As explained thereafter, the formation of Mg-silicates in Lake Alchichica sediments results from a diagenetic process which locally increases the orthosilicic acid concentrations highly enough to allow the precipitation of these phases. The source of the orthosilicic acid is discussed in the next section.

596

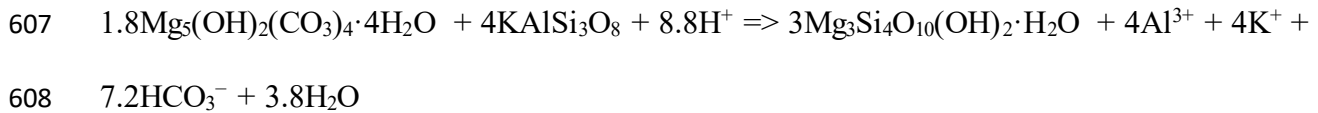
#### 597 **b) Geochemical processes involved in the formation of Mg-silicates**

598

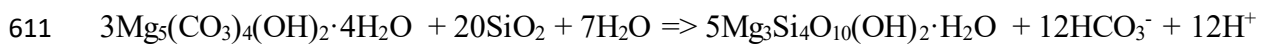
599 Here, we discuss the processes that may explain the evolution at depth of the mineralogical  
600 composition of Lake Alchichica sediments, i.e., the correlated disappearance of Mg-carbonates  
601 and amorphous silica and appearance of Mg-silicates. In the sediments of Lake Alchichica, the

602 formation of Mg-silicates may result from three different mechanisms providing alternate Mg  
603 and/or Si sources:

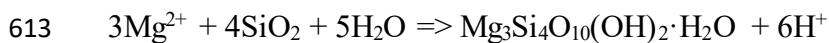
604 (1) a reaction between detrital silicates as the main source of Si and hydromagnesite as the  
605 source of Mg, along the same mechanism as that mentioned by Milesi et al. (2018) for the  
606 sediments of Lake Dziani Dzaha (Mayotte), and following the equation:



609 (2) a reaction between diatom frustules as the main source of Si and hydromagnesite as the  
610 source of Mg:



612 (3) the same reaction as in (2) with aqueous  $\text{Mg}^{2+}$  from porewater instead of hydromagnesite:



614

615 Detrital silicates, which represent 12 to 39 wt.% of the sediment content, could be a  
616 source of silica for the formation of Mg-silicates similarly as in Lake Dziani Dzaha. In Dziani  
617 Dzaha, the diagenetic Mg-silicates contain significant amounts of Al (3.65 wt.% in average;  
618 Milesi et al., 2018). Accordingly, we observed some Al-containing Mg-silicates. However, they  
619 were rare and mostly spatially associated with altered detrital silicate phases (**Figures 6 and**  
620 **S6**). By contrast, diatom frustules appear as a more likely source of  $\text{SiO}_2$  involved in the  
621 diagenetic formation of Mg-silicates in Lake Alchichica sediments. First, they are abundant  
622 enough (around 35 wt.% of the uppermost sediments) to provide all Si necessary for the  
623 formation of the Mg-silicates detected in the sediments. Second, there is very good correlation  
624 between the disappearance of the FTIR bands of amorphous silica and the concomitant  
625 appearance of silicates bands below 10 cm (**Figure 5**). Last, we observed by SEM and TEM a  
626 massive pseudomorphic replacement of the amorphous silica composing diatoms by Mg-

627 silicates, providing direct images of this diagenetic process (**Figures 7-8**). It can be noted that  
628 some Mg-silicate grains do not form as pseudomorphic replacements of diatoms but likely  
629 precipitate in the porewater containing high orthosilicic acid concentrations dissolved from  
630 diatoms. Authigenesis of Mg-silicates on diatom frustules was previously observed in the  
631 sediments of Bolivian saline lakes (Badaut, 1979; Hoffert, 1980; Badaut and Risacher, 1983;  
632 Bentz and Peterson, 2020) and marine settings (Michalopoulos and Aller, 2004). The  
633 pseudomorphic replacement of the amorphous silica of diatoms by Mg-silicates was also  
634 occasionally observed in some microbialites (e.g., Zeyen et al. 2021) but this did not produce  
635 the massive amounts of Mg-silicates as observed in Lake Alchichica sediments.

636

637 In the case of a replacement of diatoms by Mg-silicates, two sources of Mg can  
638 contribute to this process: hydromagnesite and/or dissolved  $Mg^{2+}$ . In Lake Alchichica  
639 sediments, hydromagnesite disappears below the first 3 cm of sediments, possibly entirely  
640 consumed by reaction (2). Using chemical and mineralogical data, we calculated that the 20  
641 wt.% of hydromagnesite present in the uppermost sediments (at 0-3 cm depth) represent 0.17  
642 mmol of Mg /g sediment. However, 42 wt.% of Mg-silicates are formed along the core, which  
643 represent 0.24 mmol of Mg /g sediment on average (at 41-85 cm; n=16). Therefore, some  
644 dissolved  $Mg^{2+}$  may also contribute to the formation of Mg-silicates by reaction (3). In addition,  
645 between 30 and 40 cm depth, hydromagnesite is not totally consumed. Therefore, we suggest  
646 that a lower input of diatoms to the sediments from this depth, at the time they settled down,  
647 limited the reaction of Mg-silicates formation, preserving some hydromagnesite. Overall, this  
648 suggests that the amount of amorphous silica provided by the diatom frustules (and therefore  
649 the ecological dynamics of diatom populations in the lake) is one major control of the extent of  
650 Mg-silicates authigenesis in Lake Alchichica sediments.

651

652           **c) Geochemical conditions necessary for the authigenesis of Mg-silicates in**  
653           **sediments**

654  
655           While the replacement of amorphous silica by Mg-silicates is massive in Lake Alchichica  
656 sediments and possibly other lakes, it does not obviously occur in all lakes. Here we discuss the  
657 conditions necessary for this process to take place. In this purpose, the chemical composition  
658 of the porewater is helpful. Pore waters in the first 10 centimeters of the sediments had a  
659 constant orthosilicic acid activity (**Figure 3**), around  $661 \pm 24$  on average, suggesting a potential  
660 control by the solubility of a silica phase that would have a  $\log(K_s)$  of -3.18  
661 ( $\log(K_s) = \log(a[\text{H}_4\text{SiO}_4])$ ; Figure S7). This value is consistent with what is known for the  
662 solubility of diatom silica. Indeed, the solubility constant of diatom frustules is known to be  
663 variable and a function of the physicochemical properties of the solid material (e.g. degree of  
664 condensation and order of the silica framework, water and impurity contents, specific surface  
665 area; Willey, 1980; Hurd and Birdwhistell, 1983; Beck et al., 2002; Dixit and Van Cappellen,  
666 2002; Gehlen and Van Raaphorst, 2002; Loucaides et al., 2012a,) and of the surrounding  
667 solution (e.g. temperature, salinity, pressure; Loucaides et al., 2012b). As a result, it likely  
668 varies substantially between different diatom species. Overall, it has been shown that  
669 diatomaceous silica solubilities exhibit a large variability from 200 to 1600  $\mu\text{M}$  in seawater  
670 ( $\log(K_s)$  between -3.70 and -2.79; Loucaides et al., 2012a, 2012b and references therein).  
671 Badaut and Risacher (1983) observed similar solubilities in continental saline lakes around -3.  
672 Therefore, the  $\log(K_s)$  of -3.18 tentatively inferred from the chemical measurements in the  
673 porewater of Lake Alchichica is consistent with an equilibrium of porewaters with the  
674 amorphous silica phase composing the abundant diatoms of the sediments.

675



676 Below 11.5 cm in depth in the sediments, the activity of orthosilicic acid is no longer  
677 constant, which is consistent with the observed disappearance of silica in the solid by, e.g.,  
678 FTIR. However, the  $\log[a(\text{Mg}^{2+})/a(\text{H}^+)^2] / \log[a(\text{H}_4\text{SiO}_4)]$  becomes constant (see trend 2 in  
679 **Figure 3**), i.e., in a  $\log[a(\text{Mg}^{2+})/a(\text{H}^+)^2]$  vs  $\log[a(\text{H}_4\text{SiO}_4)]$  graph, porewaters become aligned  
680 along a trend parallel to the solubility line of the “amorphous sepiolite” defined by Wollast et  
681 al. (1968) at 25°C (with a  $\log(K_s)$  of 18.78). The  $\log(K_s)$  inferred for the Mg-silicate phase  
682 observed in Alchichica sediments at 15°C is 19.26 (Figure S7). This strongly suggests that  
683 below 11.5 cm depth, a Mg-silicate phase with this solubility controls the activities of protons,  
684  $\text{Mg}^{2+}$  and orthosilicic acids in the porewater. This is again in agreement with the fact that such  
685 a phase becomes a dominant Si and Mg-bearing phase in the sediments at these depths. This  
686 also answers the question of the geochemical conditions necessary for the formation of Mg-  
687 silicates in sediments: the ionic product of the solution should be higher or equal to the solubility  
688 of a Mg-silicate phase close to that of “amorphous sepiolite”.

689  
690 Tosca et al. (2011) suggested that a much higher saturation of the solution was needed so  
691 that precipitation could spontaneously occur. According to them, the different solubility  
692 constants obtained by their study vs. that by Wollast et al. (1968) were related to the source of  
693 Si used in their experiments (i.e., respectively sodium metasilicate nonahydrate *versus*  
694 tetraethyl orthosilicate). Here, the observations point to a phase less soluble than that formed in  
695 Tosca et al. (2011)'s experiments. However, the critical supersaturation line defined by these  
696 authors refers to the conditions allowing homogeneous nucleation of Mg-silicates. In  
697 Alchichica sediments, the replacement of the diatom frustules by Mg-silicates likely involves  
698 heterogeneous nucleation, which is known to occur at lower saturations than homogenous  
699 nucleation. Therefore, this may explain why the solubility of the Mg-silicate phase formed in  
700 the sediments of Lake Alchichica is lower than the one determined by Tosca et al. (2011) and

701 closer to that of the “amorphous sepiolite” defined by Wollast et al. (1968). In any case, we  
702 suggest that the solubility constant determined here should be considered when modelling the  
703 formation of Mg-silicates in a natural environment.

704

705 Now that the geochemical conditions necessary for the formation of Mg-silicates are better  
706 assessed, we can understand some other observations. First, despite oversaturation relative to  
707 the reported solubilities of Mg-silicate phases such as kerolite, stevensite, talc and sepiolite  
708 below 30 m depth, the whole water column of Lake Alchichica remains undersaturated with  
709 “amorphous sepiolite”. This explains why no Mg-silicate was observed in the mineral  
710 assemblage of the uppermost sediments, suggesting that they do not precipitate from the lake  
711 water. Second, porewaters in Lake Dziani Dzaha sediments were undersaturated with  
712 “amorphous sepiolite” at almost all depths, explaining why Al-poor Mg-silicates could not form  
713 there and therefore were not observed (**Figure S7**). One reason why the saturation in Lake  
714 Dziani sediments might be lower than in Lake Alchichica and Bolivian saline lakes sediments  
715 (Badaut and Risacher, 1983) may relate to the low abundance of diatoms in the water column  
716 of this lake (Gérard et al., 2018). However, Mg-silicates detected in the sediments of Lake  
717 Dziani Dzaha were rich in Al (Milesi et al., 2019) in contrast to those observed in Alchichica  
718 sediments. Al-rich Mg-silicates have a lower solubility (Milesi et al., 2020), explaining why  
719 those phases may form in Lake Dziani Dzaha sediments.

720 In Lake Dziani Dzaha, the formation of Mg-silicates from detrital silicates by reaction (1)  
721 consumes protons and can therefore be driven by a pH decrease. Milesi et al. (2019, 2020)  
722 proposed that oxidation of organic matter and input of volcanic CO<sub>2</sub> tend to decrease the pH of  
723 porewaters, which promotes destabilization of hydromagnesite and ultimately allows Mg  
724 supply for the formation of saponite. In Lake Alchichica sediments, the decrease of pH from  
725 9.25 to 8.46 and the DIC increase from 34.1 to 43.2 mM may as well reflect oxidation of organic

726 matter and/or input of volcanic CO<sub>2</sub>. However, reactions (2) and (3) based on diatom  
727 transformation produce protons instead of consuming them. Therefore, these reactions are  
728 favored by a pH increase, provided that concentrations of Mg and Si are high enough to reach  
729 the supersaturation with respect to Mg-silicates precursors, i.e. “amorphous sepiolite”. We thus  
730 suggest that in Alchichica sediments, the spontaneous dissolution of the diatom frustules and  
731 not organic matter mineralization drives Mg-silicates authigenesis.

732

#### 733 **d) Some implications of massive authigenesis of Mg-silicates**

734

735 We can note three possible geochemical implications associated with the massive  
736 authigenesis of Mg-silicates in Lake Alchichica sediments.

737 First, the precipitation of Mg-silicates in Alchichica sediments imposes lower dissolved  
738 H<sub>4</sub>SiO<sub>4</sub> concentrations in porewaters than if they were in equilibrium with the silica of diatoms  
739 (such as in the top 10 cm of sediment, **Figure 3**). Therefore, we suggest that H<sub>4</sub>SiO<sub>4</sub>  
740 concentration gradient between the sediments and the water column is also lower, decreasing  
741 the diffusion rate of this species from the sediments to the water column. Accordingly, although  
742 the H<sub>4</sub>SiO<sub>4</sub> concentration increases with lake water depth from below the detection limit to  
743 0.026 mM at 60 m depth and 0.084 mM at the sediment interface, it is much lower than the pore  
744 water concentration of 0.76 mM (**Table 1**). This low Si release rate from the sediments, in  
745 addition to a possibly higher abundance of diatoms in the water column, could ultimately favor  
746 the low concentrations of H<sub>4</sub>SiO<sub>4</sub> observed in Lake Alchichica compared with other lakes  
747 around (namely Lakes Atexcac and La Preciosa).

748 Second, the diagenetic formation of Mg-silicates calls for care when interpreting the  
749 paleoenvironmental message carried by the sedimentary record. The presence/absence of Mg-  
750 silicates has been suggested to provide information about the paleochemistry of lakes. However,

751 in Lake Alchichica, we observed on the one hand shallow microbialitic deposits without Mg-  
752 silicate and on the other hand deep sediments dominated by Mg-silicates. A basin-wide  
753 assessment should therefore be considered whenever possible. In the modern system of Lake  
754 Alchichica, it is clear that stromatolites form at low concentrations of  $\text{H}_4\text{SiO}_4$  in the lake  
755 because the living diatoms are an active  $\text{SiO}_2$  sink, whereas in the sediments, the dead diatoms  
756 are a source of Si that feeds the Mg-silicates precipitation. How likely can we find signatures  
757 of this process in ancient sediments? Although fossilization of microorganisms by silicates was  
758 occasionally observed in modern microbialites (Zeyen et al., 2015), the preservation of diatom  
759 morphologies/structures in sediments is not systematic (Michalopoulos and Aller, 2004). In  
760 Alchichica, they rather gradually disappear through the massive authigenesis of Mg-silicate  
761 around diatoms (**Figure 6**). Therefore, the presence/absence of diatoms is not a robust criterion  
762 to identify this diagenetic process in the past.

763 Last, better understanding silicate authigenesis in a variety of environments is key to  
764 evaluate the effect of reverse weathering and especially its involvement in the long-term cycles  
765 of C and Si (Mackenzie and Kump, 1995). Reverse weathering refers to the reactions that  
766 produce authigenic silicates, which consumes soluble cations generated by silicate weathering,  
767 but unlike carbonates do not consume carbon (e.g., Isson and Planavsky, 2018; Krissansen-  
768 Totton and Catling, 2020). Therefore, it affects not only Si and C cycles but also the climate by  
769 retaining  $\text{CO}_2$  within the atmosphere-ocean system. While robust evidence for clay authigenesis  
770 in modern environments has emerged (Badaut and Risacher, 1968; Michalopoulos and Aller,  
771 1995; Wallmann et al., 2008; Ehler et al., 2016; Bentz and Peterson, 2020), both laboratory  
772 studies (Michalopoulos and Aller, 1995) and analyses of natural marine sediments (Wallmann  
773 et al., 2008; Ehler et al., 2016) suggest that it is almost exclusively restricted to environments  
774 characterized by extremely high rates of biogenic opal remineralization (Michalopoulos and  
775 Aller, 2004; Ehler et al., 2016). This type of environments is supposedly rare. Consequently,

776 reverse weathering is generally estimated to constitute only a relatively minor (<10%)  
777 component of total dissolved silica export in the modern oceans (e.g. Wallmann et al., 2008).  
778 Here, we show a continental system in which Mg-silicate authigenesis by diatom  
779 remineralization is major. This allowed us to refine the conditions under which authigenesis  
780 occurs. Furthermore, models have tried to quantify reverse weathering over geological times  
781 (Isson and Planavsky, 2018; Krissansen-Totton and Catling, 2020). In the Precambrian, before  
782 the advent of significant biogenic silica precipitation, ocean silica concentrations were likely  
783 elevated (Siever, 1992; Conley et al., 2017), deriving from continental and seafloor weathering  
784 (Isson and Planavsky, 2018). Isson and Planavsky (2018) argued that these elevated silica levels  
785 may have increased the reverse weathering flux, maintained high levels of CO<sub>2</sub>, and warmed  
786 the Precambrian climate. Additionally, these authors proposed that reverse weathering is  
787 strongly pH dependent, and thus may have buffered ocean pH and stabilized the atmosphere-  
788 ocean pCO<sub>2</sub> during the Precambrian. A key parameter in these models is the solubility of  
789 authigenic silicates, which has been determined by laboratory experiments and is still debated  
790 (Wollast et al., 1968; Tosca et al., 2011). Here, we provide a solubility constant deduced from  
791 a natural system, lower than the one proposed by Tosca et al. (2011) and Tosca and Masterson  
792 (2014). Therefore, despite there is no observational evidence to date suggesting a greater  
793 importance of reverse weathering in the past, its proportion associated with this solubility  
794 constant could be higher than predicted by Tosca et al. (2011) and thus have a greater impact  
795 on pH buffering potential of the oceans and climate regulation. We also showed that Mg-silicate  
796 authigenesis in Alchichica is strongly driven by diatoms, which appeared late in geological  
797 history, likely during the Cretaceous (Girard et al., 2020). Earlier organisms such as sponges,  
798 radiolarians and some bacteria (Li et al., 2022) may have produced silica-rich sediments by  
799 biomineralization as well in the past, which may have fed this reverse weathering process in  
800 the same way.

801

## 802 F) Conclusion

803

804 Alchichica lacustrine sediments exhibit a particular mineralogical evolution in their first  
805 centimeters, which can be explained by early diagenetic reactions (**Figure 11**). Below ~3 cm in  
806 depth, hydromagnesite rapidly disappears and diatom frustules are progressively  
807 pseudomorphized by an Al-poor Mg-silicate phase with a composition corresponding to  
808 stevensite. Interestingly, the replacement of amorphous silica by Mg-silicates is massive in  
809 Lake Alchichica sediments and contrasts with the absence of authigenic Mg-silicates in  
810 Alchichica shallow microbialites, mainly composed of aragonite and hydromagnesite (**Figure**  
811 **11**). This observation raises questions about the paleoenvironmental message carried by the  
812 sedimentary record, since the presence/absence of Mg-silicates is considered to provide  
813 information about the paleochemistry of lakes. However, in Lake Alchichica, microbialites  
814 form at low concentrations of  $\text{H}_4\text{SiO}_4$  in the lake because the living diatoms are an active  $\text{SiO}_2$   
815 sink, whereas in the sediments, the spontaneous dissolution of the diatom frustules drives the  
816 Mg-silicates precipitation (**Figure 11**). Therefore, an appraisal at the basin scale should be  
817 considered whenever possible. Moreover, this study confirms that the preservation of diatom  
818 morphologies/structures in sediments is not systematic, and not a robust criterion to identify  
819 this process in the past.

820 In Lake Alchichica sediments, Mg-silicate authigenesis by diatom pseudomorphosis occurs  
821 in porewaters saturated with “amorphous sepiolite” as defined by Wollast et al. (1968) (**Figure**  
822 **3**). The solubility constant deduced from this study ( $\log(K_s) = 19.26$  at  $15^\circ\text{C}$ ) is thus lower than  
823 the one proposed by Tosca et al. (2011) and Tosca and Masterson (2014) ( $\log(K_s) = 33.82$  at  
824  $25^\circ\text{C}$ ). We propose that it should be considered when modelling the formation of Mg-silicates

825 in a natural environment. The reassessment of this higher solubility constant suggests that  
826 reverse weathering may have a greater impact on climate stabilization than previously thought.

827

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838 France). The authors declare no conflict of interest.

839

## 840 **Data Availability Statement**

841 The data that supports the findings of this study are included in this article and available in  
842 supplementary material.

843

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1 **Supplementary Information**

2 **Diagenetic formation of stevensite by replacement of diatom frustules in the**  
3 **sediments of the alkaline Lake Alchichica (Mexico)**

4  
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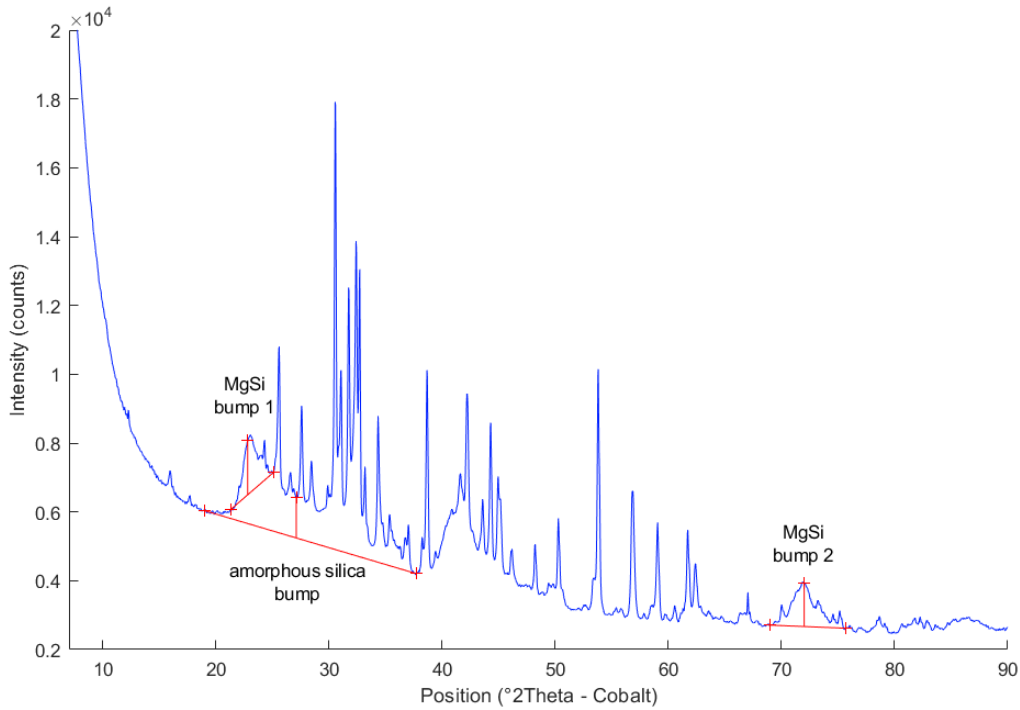
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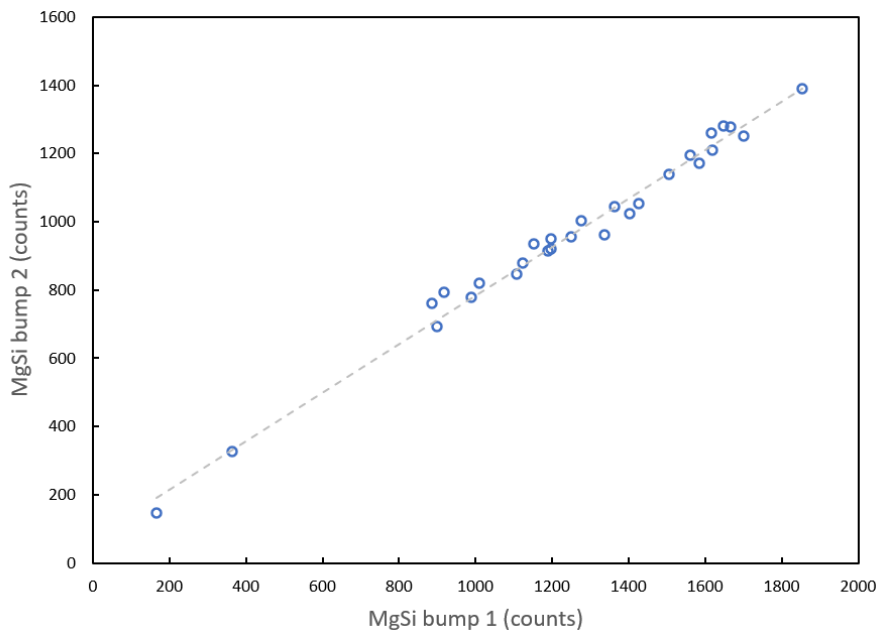
17 *<sup>f</sup>Ecologie, Systématique et Evolution, CNRS, Université Paris-Saclay, AgroParisTech, Orsay,*  
18 *France*

19 *<sup>g</sup>Departamento de Ecología y Recursos Naturales, Universidad Nacional Autónoma de México,*  
20 *DF México, Mexico*

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22 \*Corresponding author: [karim.benzerara@upmc.fr](mailto:karim.benzerara@upmc.fr)  
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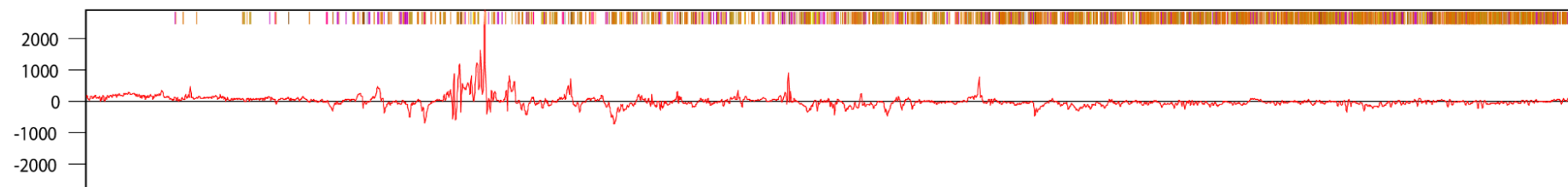
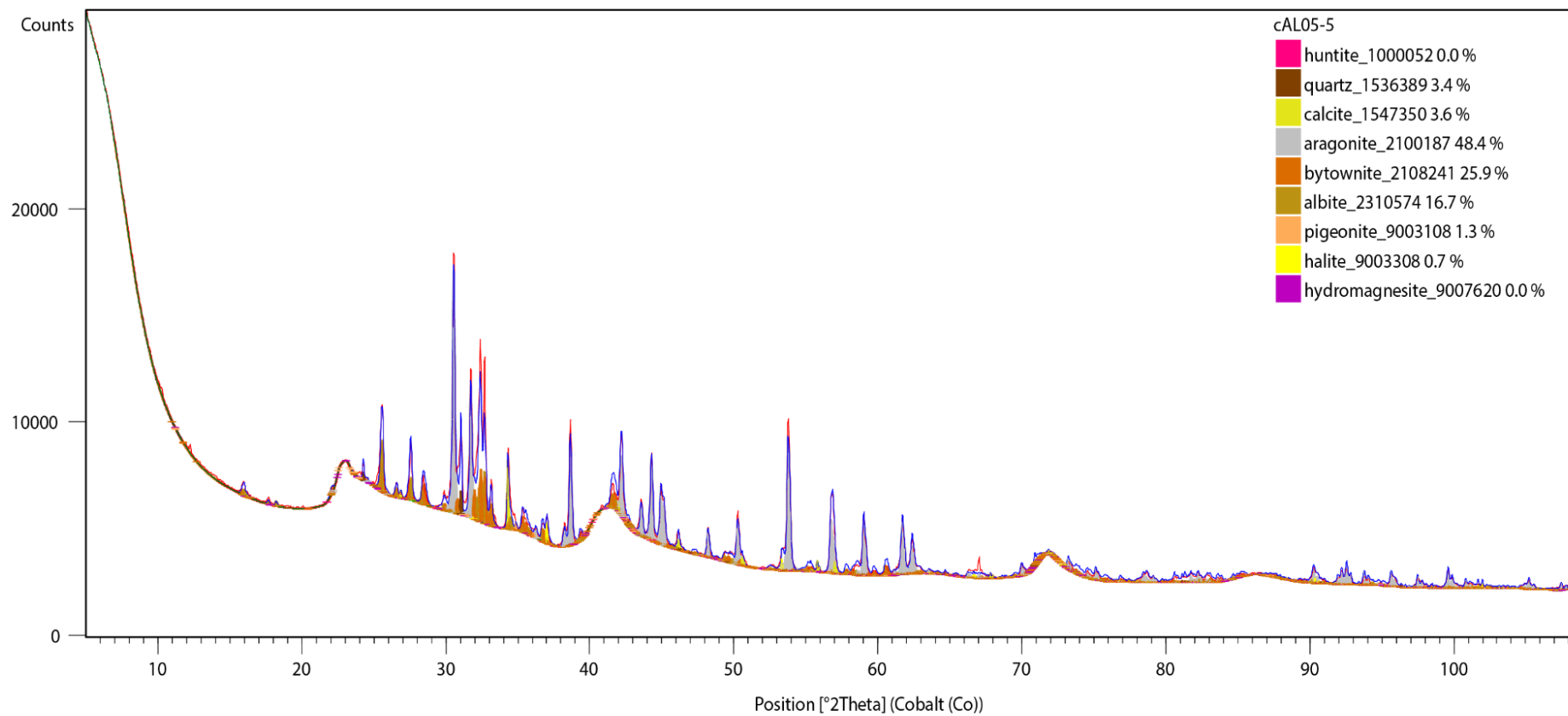


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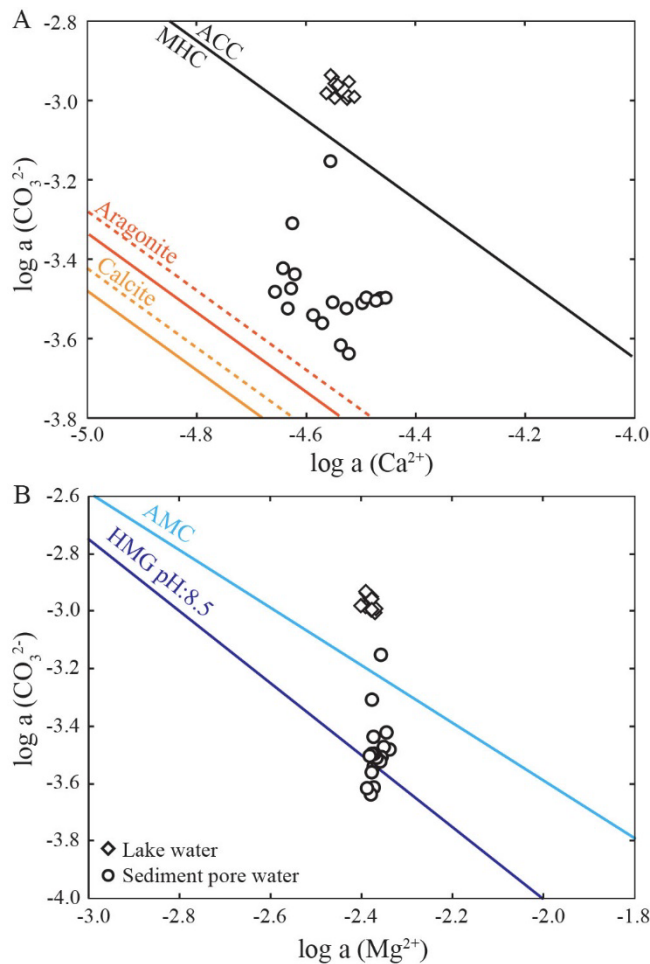
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26 **Figure S1:** Poorly crystalline phases produce broad diffraction signal on the diffractograms,  
 27 which intensity could be estimated. (Top) Example diffractogram (sample AL19\_C2a\_05) in  
 28 blue with metric used to estimate “bumps” signal intensity (in red). For each signal, a linear  
 29 baseline corrects for background within the diffraction windows. Then, the intensity is  
 30 estimated by the signal height near the center of the window, at an angle where no interference  
 31 from major diffraction peaks is found across all samples. (Bottom) Intensity of Mg-silicate  
 32 bump 1 and 2 correlation for all samples highlighting the signal produced from the same low  
 33 crystalline phase.

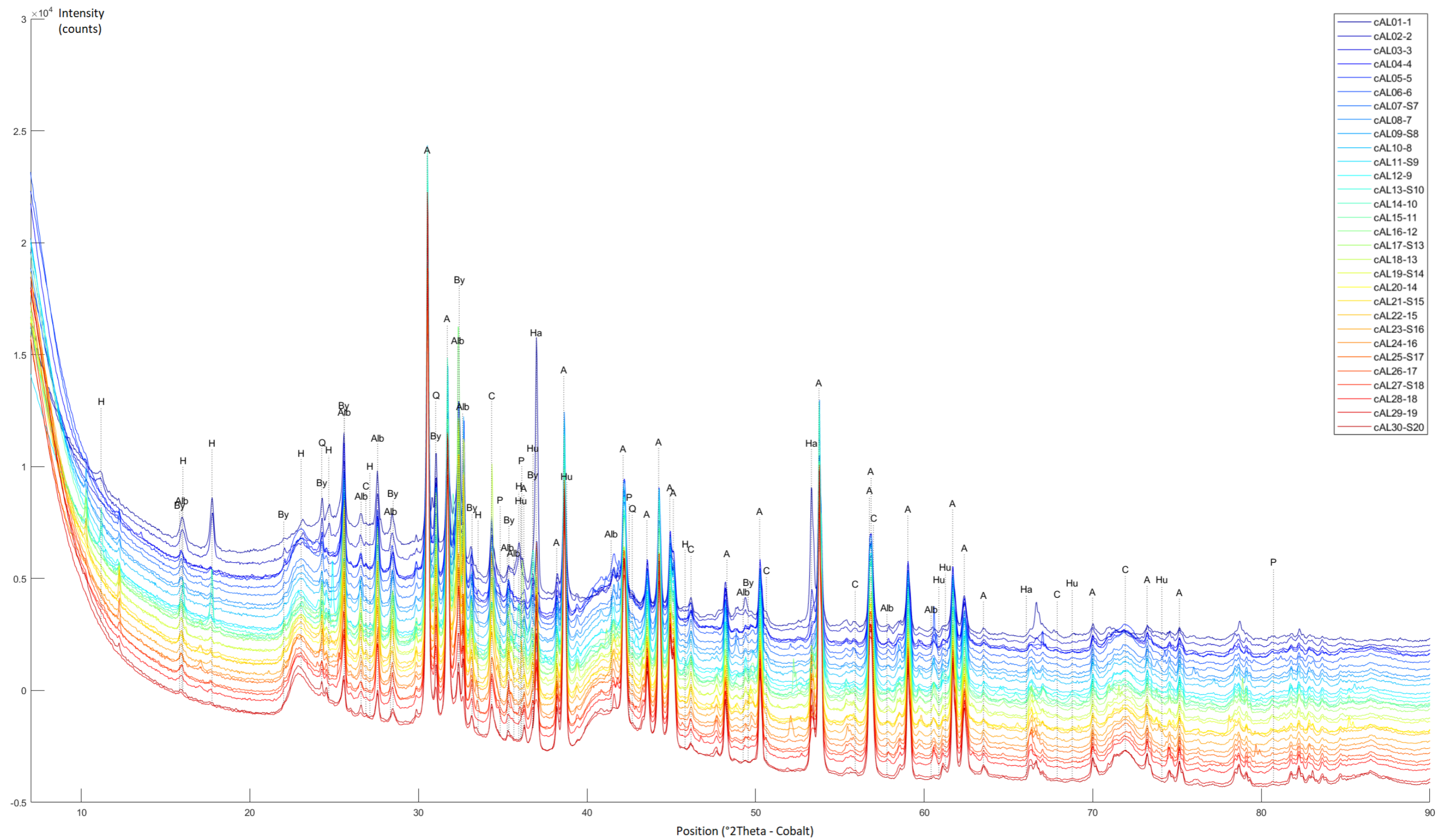


35 **Figure S2:** Example Rietveld fit result (sample AL19\_C2a\_05) showing the actual data in red and the modeled diffractogram in blue. Each Rietveld  
36 patterns of the minerals is shown as a solid profile fill. The background signal was fit so that the contribution of X-ray low crystalline phases could  
37 be removed from peak analysis. Bottom panel shows the residual difference with fit model.



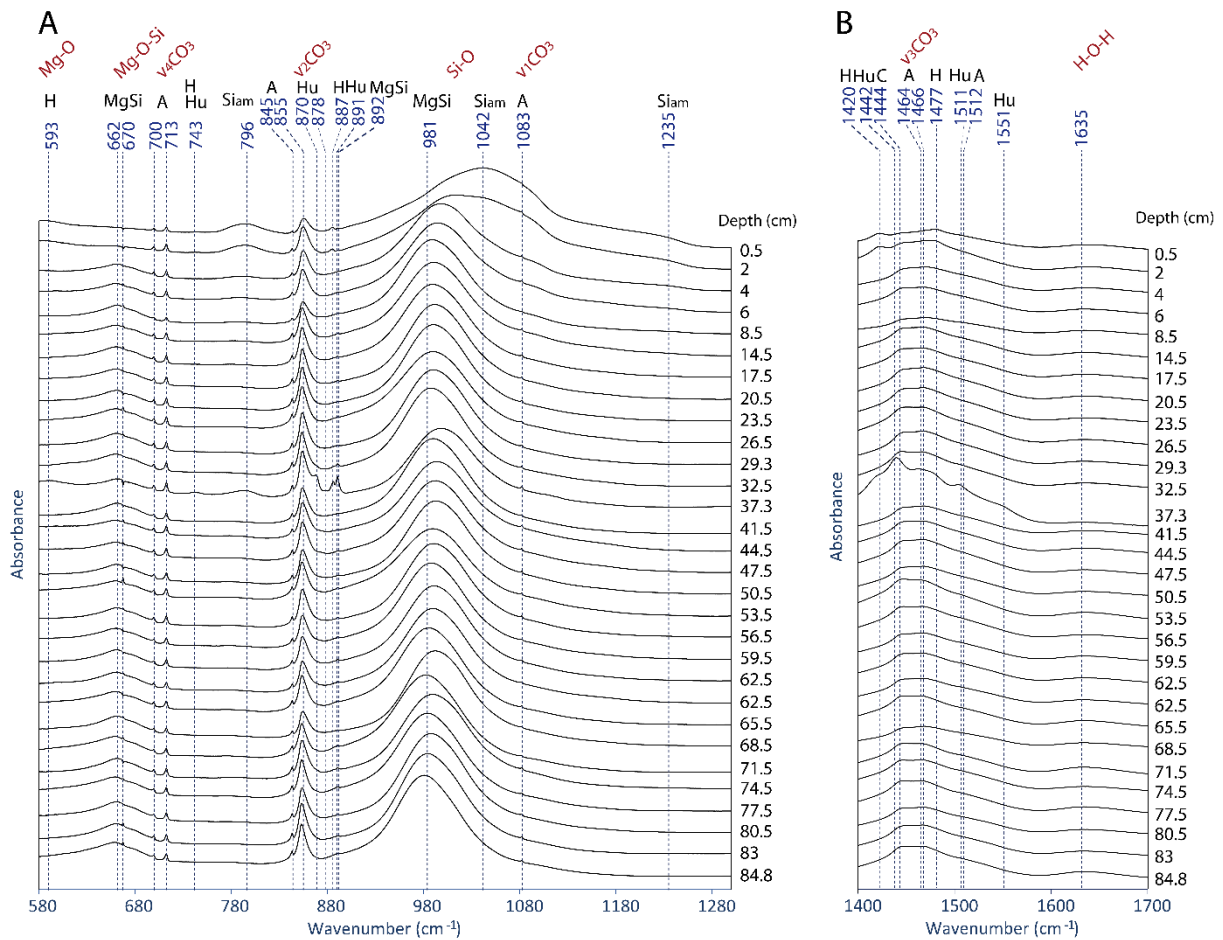


38  
 39 **Figure S3: Solubility diagram in the a)  $\log[a(\text{CO}_3^{2-})] - \log[a(\text{Ca}^{2+})]$  and b)  $\log[a(\text{CO}_3^{2-})] -$**   
 40  **$\log[a(\text{Mg}^{2+})]$  spaces.** Diamonds correspond to aqueous solutions from the water column  
 41 whereas circles correspond to porewaters. The solubilities of calcite and aragonite were  
 42 calculated at 15°C (dashed lines) and 25°C (solid lines) using SUPCRTBL software (Zimmer  
 43 et al., 2016). The solubilities of monohydrocalcite (MHC) and amorphous Mg carbonate  
 44 (AMC) were defined by Fukushi and Matsumiya (2018) and the solubility of amorphous Ca  
 45 carbonate (ACC) was measured by Kellermeier et al. (2014). The solubility line of  
 46 hydromagnesite (HMG) was determined at 15°C using the solubility coefficient defined by  
 47 Robie and Hemingway (1973) and with an average pore water pH of 8.5.

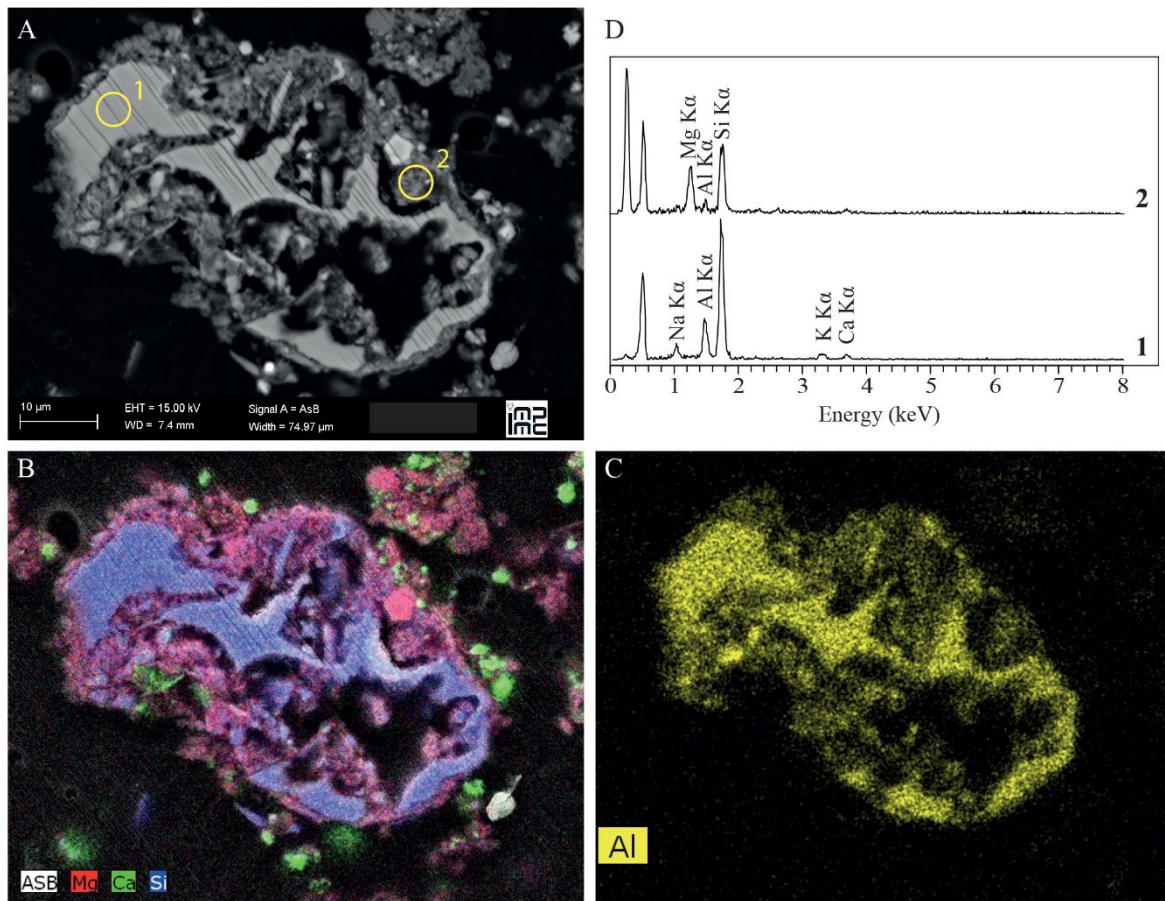


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**Figure S4: All diffractograms collected on samples from shallowest to deepest.** Vertical annotations highlight the location of diffraction peaks from mineral species used in Rietveld fit (A: aragonite, Alb: albite, By: bytownite, H: hydromagnesite, Hu: huntite, Ha: halite, C: calcite, Q: Quartz, P: Pigeonite)

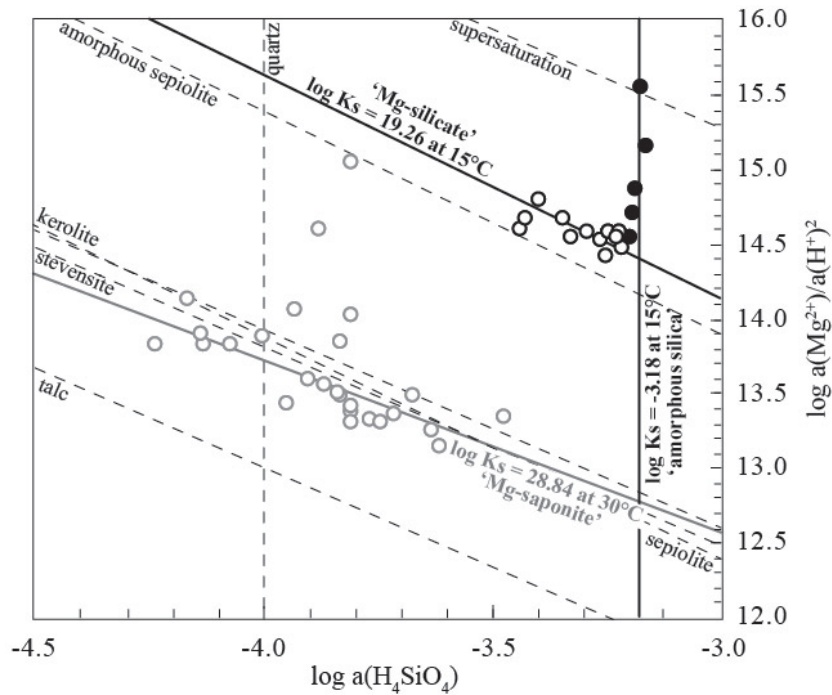


51  
 52 **Figure S5: FTIR spectra of the Alchichica sediment cores from the shallowest to the**  
 53 **deepest (from top to bottom).** (A) Spectra in the 580 to 1280 cm<sup>-1</sup> wavenumber range of  
 54 Alchichica sediment cores. The dashed lines show characteristic bands of silicates [authigenic  
 55 magnesium silicates (MgSi), amorphous silica (Si<sub>am</sub>)] and carbonates [aragonite (A), calcite  
 56 (C), hydromagnesite (H), huntite (Hu)] in the samples. (B) Spectra of the same samples in the  
 57 1400 to 1700 cm<sup>-1</sup> wavenumber range.  
 58



59

60 **Figure S6:** Mg-silicates formed around a detrital feldspar grain. (A) Backscattered electron  
 61 (BSE) image, (B) corresponding EDXS maps of Mg (red), Ca (green) and Si (blue) and (C) Al  
 62 (yellow). (D) EDXS spectra of the zones highlighted by yellow circles. On the chemical  
 63 composition map B, aragonite appears in green, Mg-silicates in dark red and Na-Ca-K feldspar  
 64 in blue. The Mg-silicates found in the vicinity of detrital grains contain  $2.41 \pm 0.85$  at.% of Al in  
 65 average ( $n=20$ ).  
 66



67

68 **Figure S7: Solubility diagram in the  $\log[a(\text{Mg}^{2+})/a(\text{H}^+)^2] - \log[a(\text{H}_4\text{SiO}_4)]$  space**  
 69 **determined at 25°C.** Pore waters of Lakes Alchichica (in black) and Dziani Dzaha (Mayotte,  
 70 in grey) are plotted against the equilibrium solubility lines of talc (Jones and Galan 1988),  
 71 stevensite (Chahi et al. 1997), kerolite and sepiolite (Stoessell 1988), “amorphous sepiolite”  
 72 (Wollast et al. 1968), quartz and amorphous silica (Truesdell and Jones 1974) and the  
 73 “supersaturation” line from Tosca et al. (2011) and Tosca and Masterson (2014). The solid  
 74 lines represent the solubility of Mg-saponite (in grey), amorphous silica and poorly crystalline  
 75 Mg-silicate (in black) deduced from the analysis of sediment porewaters respectively from  
 76 Dziani (Milesi et al. 2020) and Alchichica between 0 and 13 cm and between 16 and 84 cm  
 77 (this study).

78

79 **Table S1.** Sediment composition in wt.% estimated by Rietveld (amorphous free).

	Albite	Aragonite	Augite	Bytownite	Calcite	Halite	Huntite	Hmg	Illite	Pigeonite	Plagioclase	Quartz	Total
AL19_C2a_01	12.0	39.9	0.4	17.0	2.3	8.6	0.3	16.0	0.0	1.0	0.0	3.1	100.6
AL19_C2a_02	13.0	55.5	0.0	13.0	2.3	0.0	0.3	12.2	0.0	0.7	0.0	2.7	99.6
AL19_C2a_03	15.0	61.1	0.0	15.0	2.2	2.5	0.6	0.0	0.0	0.6	0.0	3.5	100.4
AL19_C2a_04	12.0	60.7	0.0	19.0	2.6	0.1	0.9	0.0	0.0	0.8	0.0	3.0	99.1
AL19_C2a_05	17.0	48.4	0.0	26.0	3.6	0.7	0.0	0.0	0.0	1.3	0.0	3.4	100.3
AL19_C2a_06	15.0	55.1	0.0	22.0	3.2	0.2	0.2	0.0	0.0	1.3	0.0	2.8	99.8
AL19_C2a_S7	17.0	55.4	0.0	18.0	2.6	0.2	0.6	0.9	0.2	1.5	0.0	2.9	99.3
AL19_C2a_07	12.0	60.7	0.0	19.0	2.7	0.1	0.0	0.7	0.8	1.4	0.0	2.1	99.5
AL19_C2a_S8	14.0	57.8	0.1	20.0	2.4	0.1	0.5	0.2	0.3	1.3	0.0	3.4	100.1
AL19_C2a_S9	12.0	63.7	0.3	17.0	1.8	0.0	0.0	0.2	0.0	2.1	0.1	2.7	99.9
AL19_C2a_S10	12.0	49.5	0.3	13.0	1.7	0.1	11.6	8.7	0.0	1.3	0.0	2.0	100.2
AL19_C2a_10	15.0	47.1	0.0	14.0	1.5	0.2	10.6	7.3	0.2	1.0	0.0	2.1	99.0
AL19_C2a_11	18.0	56.6	0.1	16.0	2.0	1.4	0.2	0.8	0.8	1.4	0.2	2.6	100.1
AL19_C2b_12	17.0	43.8	0.1	27.0	3.5	2.0	0.2	0.6	0.1	1.6	0.2	3.4	99.4
AL19_C2b_S13	16.0	47.5	0.3	23.0	3.3	0.5	0.5	0.8	0.7	3.6	0.2	4.0	100.4
AL19_C2b_13	15.0	51.2	0.0	18.0	7.2	2.8	0.5	0.5	0.2	1.2	0.1	3.1	99.8
AL19_C2b_S14	15.0	50.9	0.0	26.0	3.3	0.1	0.0	0.7	0.0	1.3	0.1	3.0	100.4
AL19_C2b_14	9.0	59.6	0.4	19.0	2.6	2.7	1.5	0.7	0.3	1.1	0.1	2.5	99.6
AL19_C2b_S15	10.0	54.8	0.1	26.0	3.1	0.2	0.4	1.1	0.0	1.4	0.0	3.0	100.1
AL19_C2b_15	17.0	47.5	0.0	22.0	5.7	1.9	0.6	0.5	0.0	2.3	0.0	2.7	100.2
AL19_C2b_S16	17.0	58.2	0.0	17.0	2.8	0.2	0.4	0.3	0.1	1.1	0.0	2.8	99.9
AL19_C2b_16	13.0	63.1	0.0	12.0	1.7	4.3	1.1	0.3	0.8	0.7	0.0	2.1	99.2
AL19_C2b_S17	14.0	57.0	0.3	19.0	1.8	2.7	0.7	0.5	0.5	1.0	0.0	2.1	99.6
AL19_C2b_17	9.0	68.1	0.0	15.0	1.5	1.3	0.9	0.6	0.4	0.5	0.0	2.5	99.8
AL19_C2b_S18	8.3	35.1	0.0	52.0	0.9	0.4	0.3	0.4	0.4	0.5	0.0	1.4	99.7
AL19_C2b_18	12.0	70.1	0.0	10.0	1.9	1.8	0.9	0.5	0.4	1.3	0.0	2.0	100.8
AL19_C2b_19	6.0	79.8	0.0	8.0	1.3	3.0	1.4	0.0	0.0	0.2	0.0	0.9	100.7
AL19_C2b_S20	11.0	76.0	0.1	2.0	1.0	4.0	1.0	0.0	1.0	1.0	0.0	2.0	99.1

81 **Table S2.** Stoichiometry of the mineral phases estimated by EDXS.

	Si/O	Mg/O	Ca/O	Fe/O	Al/O	Na/O	Formula
Bytownite (n=5)	0.37		0.07		0.22	0.07	(Ca <sub>0.5</sub> Na <sub>0.5</sub> )(Al(Al <sub>0.5</sub> Si <sub>0.5</sub> ))Si <sub>2</sub> O <sub>8</sub>
Albite (n=5)	0.38				0.15	0.13	NaAlSi <sub>3</sub> O <sub>8</sub>
Pyroxene (n=7)	0.29	0.13	0.13	0.03			(Ca <sub>0.45</sub> Mg <sub>0.45</sub> Fe <sub>0.1</sub> )Si <sub>2</sub> O <sub>6</sub>
Olivine (n=12)	0.19	0.31	0.02	0.08			(Fe <sub>0.3</sub> Mg <sub>0.7</sub> ) <sub>2</sub> SiO <sub>4</sub>
Quartz (n=5)	0.50						SiO <sub>2</sub>

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84 **Table S3.** Method for quantification of mineral phases.

Mineral	Chemical Formula	Calculation***
Feldspars*	Ca <sub>0.25</sub> Na <sub>0.25</sub> Al <sub>1.25</sub> Si <sub>2.75</sub> O <sub>8</sub>	Felds = [Al] / 1.25
Aragonite	CaCO <sub>3</sub>	A = [Ca] – (Felds x 0.25) – (Mg <sub>Carb</sub> x 0.6)
Mg-silicates	Mg <sub>3</sub> Si <sub>4</sub> O <sub>10</sub> (OH) <sub>2</sub>	MgSi = [Mg] / 3 or ([Si] – Felds x 2.75) / 4**
Hydromagnesite	Mg <sub>5</sub> (CO <sub>3</sub> ) <sub>4</sub> (OH) <sub>2</sub> 4H <sub>2</sub> O	H = [Mg] / 5
Amorphous silica	SiO <sub>2</sub>	Si <sub>am</sub> = [Si] – (Felds x 2.75) – (MgSi x 4)
Mg-carbonates**	Ca <sub>0.6</sub> Mg <sub>3.8</sub> (CO <sub>3</sub> ) <sub>4</sub> (OH) <sub>0.8</sub> (H <sub>2</sub> O) <sub>1.6</sub>	Mg <sub>Carb</sub> = ([Mg] – MgSi x 3) / 3.8

\* Mix 50/50 Albite-Bytownite based on XRD results (at all depths)

\*\* Mix 40/60 Hydromagnesite-Huntite based on XRD results (at 30.5-40 cm depth)

\*\*\* [X] are the concentrations in mol.% measured by ICP-AES

85

## 86 **References**

87

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