

1 **Biogeochemical processes captured by carbon isotopes in**
2 **redox-stratified water columns: a comparative study of four**
3 **modern stratified lakes along an alkalinity gradient.**

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22 *Keywords: Carbon cycle; DIC; POC; isotopic fractionation; Precambrian analogues*
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25 **Abstract.** Redox-stratified water columns are a prevalent feature of Earth history, and ongoing environmental
26 changes tend to promote a resurgence of such settings. Studying modern redox-stratified environments has
27 improved our understanding of biogeochemical processes and element cycling in such water columns. These
28 settings are associated with peculiar carbon biogeochemical cycling owing to a layered distribution of biological
29 processes in relation to oxidant availability. Metabolisms from distinct biogeochemical layers are diverse and may
30 differently imprint the sedimentological record. Paired carbon isotope compositions of organic matter and
31 carbonates, which are commonly used to characterize these ecological dynamics, can thus vary from one stratified
32 environment to another. Changes in the organic/inorganic carbon sources and mass balance can further complicate
33 the isotopic message in stratified environments. Better understanding of these multifaceted carbon isotope signals
34 requires further evaluation of how the processes occurring in redox-stratified water columns are transferred to the
35 sediments. We therefore characterized and compared the isotopic signatures of dissolved inorganic carbon (DIC),
36 carbonate, and organic matter reservoirs, at different depths in the water column and upper sediments of four
37 stratified Mexican lakes that follow a gradient of alkalinity/salinity. Comparing these systems shows strong
38 diversity in the carbon isotope signals of the water column and sediments. Differences in inorganic carbon isotope
39 signatures arise primarily from the size of the DIC reservoir, buffering the expression of redox-dependent
40 biological processes as alkalinity increases. Combining this isotopic dataset with water-column physico-chemical
41 parameters allows us to identify oxygenic photosynthesis and aerobic respiration in the four lakes studied, while
42 anoxygenic photosynthesis is evidenced in only two of them. Sedimentary organic matter does not originate from
43 the same water-column layers in the four lakes, highlighting the ecological variability that can stem from different
44 stratified water columns, and how it is transferred or not to the sedimentary record. The least alkaline lake shows
45 higher isotopic variability, and signatures typical of methanogenesis in the sediment porewaters. This metabolism,
46 however, does not leave diagnostic isotopic signatures in the sedimentary archives (organic matter and carbonates),
47 underlining the fact that even when alkalinity does not strongly buffer the inorganic carbon reservoir, a
48 comprehensive picture of the active biogeochemical carbon cycling is not necessarily transferred to the geological
49 record.

50

51 1. INTRODUCTION

52 The carbon cycle and biogeochemical conditions prevailing at the surface of the Earth are intimately bound through
53 biological (e.g. photosynthesis) and geological processes (e.g. volcanic degassing and silicate weathering). The
54 analysis of carbon isotopes in organic matter and carbonates ($\delta^{13}\text{C}_{\text{org}}$ and $\delta^{13}\text{C}_{\text{carb}}$) in the rock record, has been used
55 to reconstruct the evolution of the biosphere and the oxygenation of the Earth's surface (e.g. Hayes et al., 1989;
56 Karhu and Holland, 1996; Schidlowski, 2001). Coupling $\delta^{13}\text{C}_{\text{org}}-\delta^{13}\text{C}_{\text{carb}}$ has frequently been used to infer the burial
57 rate of organic C, and thus the redox balance of the atmosphere and hydrosphere (e.g. Karhu and Holland, 1996;
58 Aharon, 2005; Krissansen-Totton et al., 2015; Mason et al., 2017). It has also been used to deduce the presence of
59 metabolisms like anoxygenic chemoautotrophic or methanotrophic bacteria (e.g. Hayes et al., 1999; Bekker et al.,
60 2008; Krissansen-Totton et al., 2015). Coupling $\delta^{13}\text{C}_{\text{org}}-\delta^{13}\text{C}_{\text{carb}}$ has also been used to discuss oceans stratification
61 and its effect on inorganic and organic C geochemical signatures in sediments (e.g. Logan et al., 1995; Aharon,
62 2005; Bekker et al., 2008; Ader et al., 2009). Stratification favors the expression and recording of different layers
63 of the water column, with potentially very distinct isotopic signatures. As the oceans were redox-stratified during
64 most of the Earth's history (Lyons et al., 2014; Havig et al., 2015; Satkoski et al., 2015), processes affecting the C
65 cycle were likely different from those occurring in most modern, well-oxygenated environments. This change of
66 conditions could impact the $\delta^{13}\text{C}_{\text{org}}$ signal at various scales, from changes in diversity and relative abundance of
67 microbial carbon and energy metabolism (e.g. Wang et al., 2016; Iñiguez et al., 2020; Hurley et al., 2021), to larger
68 ecological interactions (e.g. Jiao et al., 2010; Close and Henderson, 2020; Klawonn et al., 2021) and global C
69 dynamics (e.g. Ridgwell and Arndt, 2015; Ussiri and Lal, 2017).

70 Modern stratified lakes have been used as analogues of ancient redox-stratified systems to better understand the C
71 cycle in the sedimentary isotopic record (e.g. Lehmann et al., 2004; Posth et al., 2017; Fulton et al., 2018). Several
72 number of recent studies have investigated the C cycle in modern stratified water columns (e.g. Crowe et al., 2011;
73 Kuntz et al., 2015; Posth et al., 2017; Schiff et al., 2017; Havig et al., 2018; Cadeau et al., 2020; Saini et al., 2021;
74 Petrash et al., 2022), where many bio-geo-physico-chemical parameters can be directly measured, together with
75 the main C reservoirs. However, investigations of such Precambrian analogues do not necessarily include sediment
76 data, and generally focus on a single environment without integrating views from several systems.

77 In this study, we measured the concentrations and isotopic compositions of dissolved inorganic carbon (DIC) and
78 particulate organic carbon (POC) throughout the water column of four modern redox-stratified alkaline crater
79 lakes, located in the Trans-Mexican Volcanic Belt (Ferrari et al., 2012). We also measured the concentrations and
80 isotopic compositions of the sedimentary organic carbon and carbonates as well as porewater DIC from surficial
81 sediments (~ 10 cm) at the bottom of the lakes. The four lakes share similar geological and climatic contexts but
82 have distinct solution chemistries along a marked alkalinity–salinity gradient (Zeyen et al., 2021) – as well as
83 distinct planktonic communities (Iniesto et al., 2022). We therefore seek to evaluate how these environmental and
84 ecological differences are recorded in the C isotope signatures in the water columns (DIC–POC) and sedimentary
85 archives (organic matter–carbonates). The four lakes are closed lakes in endorheic basins (Alcocer, 2021; Zeyen
86 et al., 2021), which facilitates the identification of external environmental constraints (e.g. evaporation, C sources)
87 and their influence on processes occurring within the water columns. Depth profiles of the main physico-chemical
88 parameters together with trace and major elements concentrations were measured to pinpoint the dominant
89 biogeochemical processes occurring in the water columns and link them to specific C isotopes signatures.

90 First, we constrain the main DIC sources and external controls on the lakes' alkalinity. Next, we describe the
91 influence of the inter-lake alkalinity gradient on the inorganic C cycle and stratification of the lakes, and how it is
92 recorded in surficial sediments. Then, by combining POC and DIC data, we identify the sources of organic C to
93 the lakes by describing the main autotrophic reactions occurring in the water columns (e.g. oxygenic and
94 anoxygenic photosynthesis). Finally, we discuss the fate of POC, either recycled (e.g. via methanogenesis) or
95 deposited in the sediments, and how all these processes are recorded (or not) in surficial sediments.

96

97 **2. SETTING / CONTEXT**

98 **2.1. Geology**

99 The four lakes studied here are volcanic maars formed after phreatic, magmatic, and phreatomagmatic explosions,
100 related to volcanic activity in the Trans-Mexican Volcanic Belt (TMVB, Fig. 1). The TMVB originates from the
101 subduction of the Rivera and Cocos plates beneath the North America plate, resulting in a long (~1000 km) and
102 wide (90–230 km) Neogene volcanic arc spreading across central Mexico (Ferrari et al., 2012). The TMVB harbors
103 a large variety of monogenetic scoria cones and phreatomagmatic vents (maars and tuff-cones) as well as
104 stratovolcanoes, calderas, and domes (Carrasco-Núñez et al., 2007; Ferrari et al., 2012; Siebe et al., 2014). Maar
105 crater formation usually occurs when ascending magma meets water-saturated substrates, leading to successive
106 explosions and the excavation of older units (Lorenz, 1986; Carrasco-Núñez et al., 2007; Siebe et al., 2012; Chako
107 Tchamabé et al., 2020).

108 The first lake, La Alberca de los Espinos (1985 masl), is located at the margin of the Zacapu tectonic lacustrine
109 basin in the Michoacán-Guanajuato Volcanic Field (MGVF) in the central-western part of the TMVB (Fig. 1). It
110 lies on andesitic basement rocks and was dated at $\sim 25 \pm 2$ ka (Siebe et al., 2012, 2014). The other three lakes (La
111 Preciosa, Atexcac and Alchichica) are all in the same area (~ 50 km²) of the Serdan-Oriental Basin (SOB) in the
112 easternmost part of the TMVB (Fig. 1). The SOB is a closed intra-montane basin at high altitude (~ 2300 m),
113 surrounded by the Los Humeros caldera to the north and the Cofre de Perote-Citlatépel volcanic range to the east.
114 The basement is composed of folded and faulted Cretaceous limestones and shales, covered by andesitic-to-basaltic
115 lava flows (Carrasco-Núñez et al., 2007; Armienta et al., 2008; Chako Tchamabé et al., 2020). The Alchichica and
116 Atexcac craters was dated at $\sim 6\text{-}13 \pm 5\text{-}6$ ka (Chako Tchamabé et al., 2020) and 330 ± 80 ka (Carrasco-Núñez et
117 al., 2007), respectively (Table 1). The age of La Preciosa is not known.

118

119 **2.2. Climate and limnology**

120 La Alberca is a freshwater lake (0.6 psu) with a temperate to semi-humid climate (Rendon-Lopez, 2008; Sigala et
121 al., 2017). In contrast, lakes from the SOB experience a similar temperate to semi-arid climate (Armienta et al.,
122 2008; Sigala et al., 2017). The current climate of the SOB is dominated by dry conditions, reflected by higher
123 evaporation than precipitation fluxes in Lake Alchichica (~ 1686 vs. 392 mm/year; Alcocer, 2021). In La Preciosa,
124 Atexcac, and Alchichica, significant evaporation is reflected by a drop in water level, evidenced by the emersion
125 of microbialite deposits (Fig. S1; Zeyen et al., 2021). This evaporation-dominated climate strongly contributes to
126 the relatively high salinity values in these lakes (1.2–7.9 psu), ranging from sub- to hyposaline.

127 The four lakes are warm monomictic: they are stratified for about nine months of the year, mixing only when
128 thermal stratification breaks down in the cold of winter (Armienta et al., 2008). They are all closed lakes located
129 in an “endorheic” basin (Alcocer, 2021; Zeyen et al., 2021), meaning that they have no inflow, outflow, or
130 connection to other basins through surficial waters such as streams. The only water input is from precipitation and
131 groundwater inflow (quantified for Lake Alchichica; Alcocer, 2021 and references therein).

132 The four lakes are alkaline (pH ~ 9) but cover a broad range of chemical compositions (including alkalinity,
133 salinity, and Mg/Ca ratio), interpreted as reflecting different concentration stages of an initial alkaline dilute water
134 (Table 1; Zeyen et al., 2021). Variations in concentration stages may be due to differences in climate and, more
135 generally, different hydrological regimes. Microbialite deposits are found in all four lakes (Gérard et al., 2013;
136 Saghaï et al., 2016; Iniesto et al., 2021a, 2021b; Zeyen et al., 2021), and increase in abundance from lower to
137 higher alkalinity conditions (Zeyen et al., 2021).

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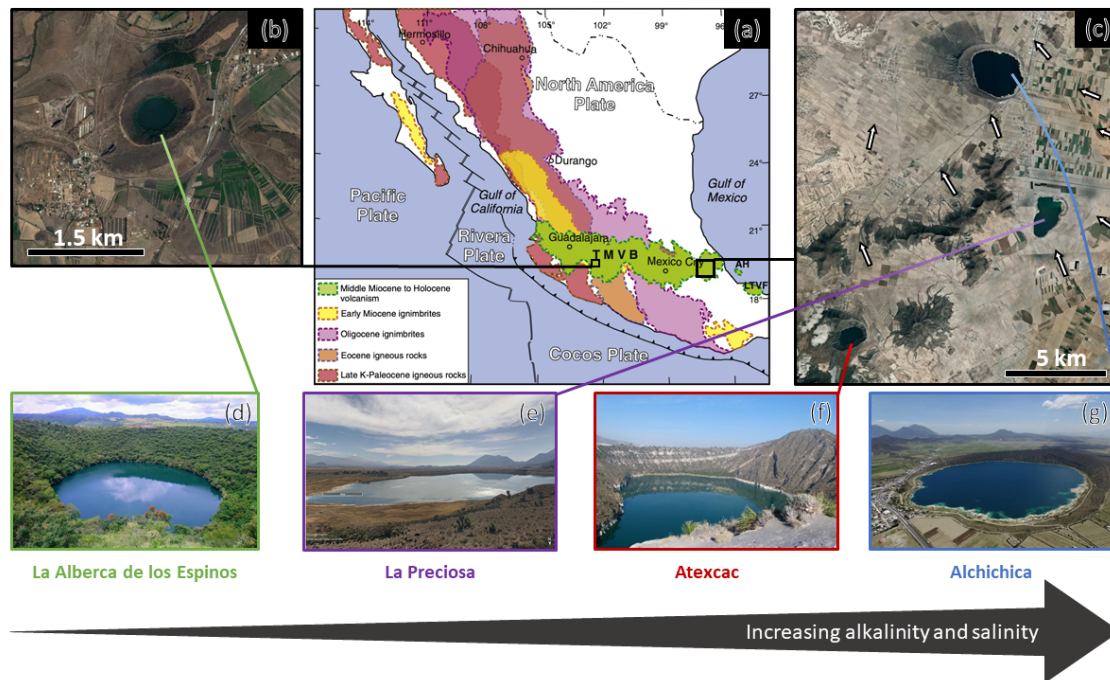
139 3. METHOD

140 3.1. Sample Collection

141 The sediment core from Lake La Preciosa was collected in May 2016. All other samples were collected in May
142 2019. The depth profiles of several physico-chemical parameters were measured in the water columns of the four
143 lakes using an YSI Exo 2 multi-parameter probe: temperature, pH, ORP (oxidation reduction potential),
144 conductivity, O₂, chlorophyll a, phycocyanin, and turbidity. Precisions for these measurements were 0.01 °C, 0.1
145 pH unit, 20 mV, 0.001 mS/cm, 0.1 mg/L, 0.01 µg/L, 0.01 µg/L and 2% FTU unit, respectively. The ORP signal
146 was not calibrated before each profile and is thus used to discuss relative variations over a depth profile.
147 Measurements of the aforementioned parameters served to pinpoint depths of interest for further chemical and
148 isotopic analyses, notably around the redoxcline of the lakes. Water samples were collected with a Niskin bottle.
149 Particulate matter was collected on pre-combusted (2 h at 490°C) and weighted glass fiber filters (Whatman GF/F,
150 0.7 µm) and analyzed for particulate organic carbon (POC), major and trace elements. Between 1.5 and 5 L of lake
151 water was filtered before the GF/F filters became clogged. The processed solution was filtered again at 0.22 µm
152 with Filtropur S filters (pre-rinsed with lake water filtered at 0.7 µm) for analyses of dissolved inorganic carbon
153 (DIC), and major, minor, and trace ions.

154 Sediment cores were collected using a 90 mm Uwitec corer close to the deepest point of each lake’s water column
155 (Table 1), where anoxic conditions prevail almost all year long. Cores measured between 20 and 85 cm in length.
156 Slices of about 2-3 cm were cut under anoxic conditions, using a glove bag filled with N₂ (anoxia was monitored
157 using a WTW3630 equipped with a FDO O₂ optode). Interstitial porewater was drained out of the core slices using
158 Rhizons in the glove bag. Sediments were transported back to the laboratory within aluminized foils (Protpack,
159 UK). Sediments were then fully dried in a laboratory anoxic N₂-filled glove box.

160



161 Figure 1. Geographical location and photographs of the four crater lakes. (a) Geological map from Ferrari et al.
 162 (2012) with black squares showing the location of the four studied lakes within the Trans-Mexican Volcanic Belt
 163 (TMVB). (b, c) Close up © Google Earth views of La Alberca de los Espinos and the Serdan-Oriental Basin
 164 (SOB). The white arrows represent the approximate groundwater flow path (based on Silva-Aguilera, 2019). (d-
 165 g) Photographs of the four lakes (d from © Google Image [‘enamoredemexicowebiste’], e from © Google Earth
 166 street view, and g from © ‘Agencia Es Imagen’).

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Lake	General location	Sampling location	Elevation (masl)
Alchichica	Serdan Oriental Basin, eastern TMVB	19°24'51,5" N; 97°24'09,9" W	2320
Atexcac	Serdan Oriental Basin, eastern TMVB	19°20'2.2" N; 97°26'59.3" W	2360
La Preciosa	Serdan Oriental Basin, eastern TMVB	19°22'18.1" N; 97°23'14.4" W	2330
La Alberca de los Espinos	Zacapu Basin, MGVF, central TMVB	19°54'23.9" N; 101°46'07.8" W	1985

168

Lake	Lake Basement	Age	Max Depth (m)	Alkalinity (mmoles/L)	Salinity (psu)	pH
Alchichica	limestone, basalts	6-13 ± 5-6 ka	63	~35	7.9	9.22
Atexcac	limestone, andesites, basalts	330 ± 80 ka	39	~26	7.4	8.85
La Preciosa	limestone, basalts	Pleistocene	46	~13.5	1.15	9.01
La Alberca de los Espinos	andesite xenoliths	25 ± 2 ka	30	~7	0.6	9.14

169

170 Table 1. General information about the lakes studied. Abbreviations: TMVB: Trans-Mexican Volcanic Belt;
 171 MGVF: Michoacán-Guanajuato Volcanic Field; masl.: meters above sea level. NB: Sampling took place in
 172 May 2019, except for La Preciosa sediments, sampled in May 2016.

173 **3.2. Dissolved inorganic carbon (DIC) concentration and isotope measurements**

174 Twelve mL of the 0.7- μm filtered lake water was filtered at 0.22- μm directly into hermetic Exetainer® tubes to
175 avoid exchange between DIC and atmospheric CO_2 . The DIC concentrations and isotopic compositions were
176 measured at the Institut de Physique du Globe de Paris (IPGP, France), using an Analytical Precision 2003 GC-
177 IRMS, running under He-continuous flow, following the protocol described by Assayag et al. (2006). A given
178 volume of the solution was extracted from the Exetainer® tube with a syringe, while the same volume of helium
179 was introduced to maintain stable pressure and atmospheric- CO_2 -free conditions within the sample tubes. The
180 collected sample was inserted into another Exetainer® tube, pre-filled with a few drops of 100% phosphoric acid
181 (H_3PO_4) and pre-flushed with He gas. Under acidic conditions, DIC quantitatively converts to gaseous and aqueous
182 CO_2 , which equilibrates overnight within the He-filled head space of the tube. Quantification and isotopic analyses
183 of released gaseous CO_2 were then carried out by GC-IRMS using internal standards of known composition that
184 were prepared and analyzed via the same protocol. Each measurement represented an average of four injections in
185 the mass spectrometer. Chemical preparation and IRMS analysis were duplicated for all the samples. The $\delta^{13}\text{C}_{\text{DIC}}$
186 reproducibility calculated for the 65 samples was better than ± 0.2 ‰, including internal and external
187 reproducibility. Standard deviation for [DIC] was 0.6 ± 0.9 mmol/L on average.

188 Specific DIC speciation, i.e., $\text{CO}_{2(\text{aq})}$, HCO_3^- and CO_3^{2-} activities, was computed using Phreeqc with the full
189 dissolved chemical composition of each sample as an input. It should be noted that these results are calculated
190 from theoretical chemical equilibria and do not necessarily take into account local kinetic effects, which, for
191 example, could lead to local exhaustion of $\text{CO}_{2(\text{aq})}$ where intense photosynthesis occurs.

192

193 **3.3. Particulate organic carbon and nitrogen (POC / PON)**

194 Particulate organic matter from the lake water columns was collected on GF/F filters, dried at room temperature
195 and ground in a ball mill before and after decarbonation. Decarbonation was performed with 12N HCl vapors in a
196 desiccator for 48 h. Aliquots of dry decarbonated samples (25 - 70 mg) were weighed in tin capsules. The POC
197 and PON contents and $\delta^{13}\text{C}_{\text{POC}}$ were determined at the Laboratoire Biogéosciences (Dijon, France) using a Vario
198 MICRO cube elemental analyzer (Elementar, Hanau, Germany) coupled in continuous flow mode with an
199 IsoPrime IRMS (Isoprime, Manchester, UK). The USGS 40 and IAEA 600 certified materials used for calibration
200 showed reproducibility better than 0.15 ‰ for $\delta^{13}\text{C}$. External reproducibility based on triplicate analyses of
201 samples (n=23) was 0.1 ‰ on average for $\delta^{13}\text{C}_{\text{POC}}$ (1SD). External reproducibility for POC and PON
202 concentrations was 0.001 and 0.005 mmol/L on average, respectively (i.e. 3 and 7 % of measured concentrations).

203

204 **3.4. Geochemical characterizations of the sediments**

205 Sedimentary organic carbon (SOC), sedimentary organic nitrogen (SON), and their isotopic compositions were
206 measured on carbonate-free residues of the first 12 cm of the sediment cores, produced after overnight 1N HCl
207 digestion. Plant debris (mainly found in La Alberca and Atexcac) was identified upon initial sediment grinding in
208 an agate mortar and analyzed separately. Aliquots of dried decarbonated samples (~ 4-70 mg) were weighed in tin
209 capsules. The SOC and SON contents and $\delta^{13}\text{C}$ were determined at the Laboratoire Biogéosciences (Dijon) using

210 a Vario MICRO cube elemental analyzer (Elementar GmbH, Hanau, Germany) coupled in continuous flow mode
211 with an IsoPrime IRMS (Isoprime, Manchester, UK). The USGS 40 and IAEA 600 certified materials used for
212 calibration had a reproducibility better than 0.2 ‰ for $\delta^{13}\text{C}_{\text{SOC}}$. Sample analyses (n=67) were at least duplicated
213 and showed an average external reproducibility of 0.1 ‰ for $\delta^{13}\text{C}$ (1SD). External reproducibility for SOC and
214 SON contents was 0.1 and 0.03 wt. %, respectively.

215 Carbon isotope compositions of sedimentary carbonates were analyzed at the Laboratoire Biogéosciences (Dijon)
216 using a ThermoScientific™ Delta V Plus™ IRMS coupled with a Kiel VI carbonate preparation device. External
217 reproducibility was assessed by multiple measurements of NBS19 standard and was better than ± 0.1 ‰ (2σ).
218 Total carbonate concentration was determined by mass balance after decarbonation for SOC analysis.

219 Mineralogical assemblages of sediments were determined on bulk powders by X-Ray diffraction (XRD) at the
220 Laboratoire Biogéosciences (Dijon). Samples were ground in an agate mortar. Diffractograms were obtained with
221 a Bruker D8 Endeavor diffractometer with $\text{CuK}\alpha$ radiation and LynxEye XE-T detector, under 40 kV and 25 mA
222 intensity. Mineral identification was based on COD (“Crystallography Open Database”) and BGMN databases.
223 Mineral abundances were estimated by Rietveld refinement analysis implemented in the Profex software.

224 Solid sulfide concentrations were determined on dry bulk sediments from La Alberca Lake after a wet chemical
225 extraction using a boiling acidic Cr(II)-solution as detailed in Gröger et al. (2009).

226

227 **3.5. Major and trace elements concentrations**

228 Dissolved and particulate matter elemental compositions were measured at the Pôle Spectrométrie Océan
229 (Plouzané, France) by inductively coupled plasma–atomic emission spectroscopy (ICP-AES, Horiba Jobin) for
230 major elements and by high-resolution ICP–mass spectrometry using an Element XR (HR-ICP-MS, Thermo Fisher
231 Scientific) for trace elements. Major element measurement reproducibility based on internal multi-elemental
232 solution was better than 5%. Trace elements were analyzed by a standard-sample bracketing method and calibrated
233 with a multi-elemental solution. Analytical precision for trace elements was generally better than 5%. Dissolved
234 sulfate concentrations were analyzed by ion chromatography at the IPGP (Paris, France) with uncertainty lower
235 than 5%.

236

237 **4. RESULTS**

238

239 **4.1. Lake La Alberca de los Espinos**

240 Stratification of the water column was well defined in La Alberca de los Espinos (Fig. 2). Temperature was higher
241 than in the other lakes (decreasing from ~ 23 °C at the surface to 16.5 °C at depth). Dissolved O_2 was oversaturated
242 at the lake surface (118 %, *i.e.*, 7.9 mg/L), rapidly decreasing to 0 between ~ 5 and 12 m, while the oxidation
243 reduction potential (ORP) only decreased below 17 m depth. The offset between O_2 exhaustion and ORP decrease
244 can be explained by the presence of other oxidant species and/or extended chlorophyll a peaks (supplementary
245 text 1). Conductivity decreased from 1.20 to 1.17 mS/cm at 16 m before increasing to 1.27 mS/cm at 26 m (salinity

246 between 0.58 and 0.64 psu). Chlorophyll a (Chl a) averaged 3.1 $\mu\text{g/L}$, and showed a profile with at least three
247 distinctive peaks, (i) between 6 and 9.5 m, (ii) around 12.5 m and (iii) between 16 and 19 m, all reaching $\sim 4 \mu\text{g/L}$.
248 The turbidity profile showed a pronounced increase from 16 to 19 m. The pH profile showed important variation
249 from 9.15 at the lake surface to 8.75 between 6.5 and 10 m, further decreasing to 7.5 between 16 and 26 m. Based
250 on the temperature profiles, epi-, meta- and hypolimnion layers of Lake La Alberca de los Espinos in May 2019
251 broadly extended from 0-5, 5-12 and 12-30 m, respectively (Fig. 2). The conductivity and pH profiles, however,
252 show that different conditions prevail at the top and bottom of the hypolimnion.

253 Dissolved inorganic carbon (DIC) concentration progressively increased from 6.8 mM at 5 m to 8.7 mM at 26 m.
254 The pCO_2 calculated for surface waters was near equilibrium with atmospheric $\text{pCO}_{2\text{atm}}$, but strongly increased
255 with depth, up to ~ 40 times the $\text{pCO}_{2\text{atm}}$ (Table S2). The $\delta^{13}\text{C}_{\text{DIC}}$ first decreased from about -2.5‰ to -4.1‰
256 between 5 and 10 m, before increasing again up to -2‰ at 25 m. Particulate organic carbon (POC) concentrations
257 reached minimum values of 0.02 mM at 10 m but rose to maximum values in the hypolimnion (0.06 mM). The
258 C:N molar ratio of particulate organic matter (POM) progressively decreased from 8.5 at the surface to less than
259 6.5 in the hypolimnion. The $\delta^{13}\text{C}_{\text{POC}}$ had minimum values at 10 and 17 m (-28.3 and -29‰ , respectively). Above
260 and below these depths, $\delta^{13}\text{C}_{\text{POC}}$ averaged $-26.4 \pm 0.5 \text{‰}$.

261 Dissolved sulfates as measured by chromatography were only detectable at 5 m with a low concentration of 12 μM ,
262 while total dissolved S measured by ICP-AES, showed values in the hypolimnion higher than in the upper layers
263 (~ 10.3 vs. $7.4 \mu\text{M}$, Table S4). Dissolved Mn concentrations decreased from 1.5 to 0.5 μM between 5 and 10 m,
264 then increased to 2 μM at 25 m. Aqueous Fe was only detectable at 25 m with a concentration of 0.23 μM
265 (Table S4). In parallel, particulate S concentrations increased with depth, with a marked increase from 0.1 to
266 0.6 μM between 20 and 25 m. Increase in particulate S was correlated with a 25-fold increase in particulate Fe
267 (from 0.2 to 5.97 μM). Particulate Mn showed a peak between 17 and 20 m around 1 μM , contrasting with values
268 lower than 0.15 μM in the rest of the water column (Fig. 2, Table S5).

269 In the first centimeters of sediments, DIC concentration in the porewater varied between ~ 11 and 12 mM and
270 $\delta^{13}\text{C}_{\text{DIC}}$ varied between $+8$ and $+10 \text{‰}$ (Figs. 3, 4). Surficial sedimentary carbonates corresponded to calcite and
271 had a $\delta^{13}\text{C}$ around -1.5‰ . Sedimentary organic matter had a $\delta^{13}\text{C}_{\text{SOC}}$ increasing from ~ -29.4 to -25.5‰ and a
272 C:N molar ratio varying between 11.6 and 14.3 (Figs. 3, 4; Table S3).

273

274 4.2. Lake La Preciosa

275 Lake La Preciosa was also stratified at the time of sample collection (Fig. 2). Temperature decreased from $\sim 20 \text{°C}$
276 at the surface to 16°C at 15 m depth. Conductivity showed the same trend with values between 2.24 and
277 2.22 mS/cm (salinity around 1.15 psu). Dissolved O_2 was oversaturated at the lake surface (120 %, i.e., 8.4 mg/L),
278 rapidly decreasing to 0 between ~ 8 and 14 m, while the ORP decreased right below 16 m. Chl a concentration
279 averaged 3 $\mu\text{g/L}$ and recorded the highest peak compared to the other lakes (about 9 $\mu\text{g/L}$ at 10 m) before
280 decreasing to 0.7 $\mu\text{g/L}$ below 15 m. Turbidity showed a large peak between 16 and 19 m. The pH showed a small
281 decrease from 9 to 8.8 between the surface and 15 m depth. Based on the temperature profiles, epi-, meta- and
282 hypolimnion layers of La Preciosa in May 2019 broadly extended from 0-6, 6-15 and 15-46 m, respectively
283 (Fig. 2).

284 The DIC concentration was constant throughout the water column at 13.3 mM, with an exception at 12.5 m, where
 285 it decreased to 11.5 mM (Fig. 3, Table S1). Calculated pCO₂ at the surface represented about two times the
 286 atmospheric pCO_{2atm} (Table S2). The δ¹³C_{DIC} decreased from about 0.5 ‰ to -0.36 ‰ between the surface and the
 287 hypolimnion. The POC concentration decreased from ~0.06 mM in the epi-/metalimnion to 0.02 mM in the
 288 hypolimnion. Similarly, (C:N)_{POM} decreased from ~11.2 in the epi-/metalimnion to 7.6 in the hypolimnion. The
 289 δ¹³C_{POC} increased downward from ~ -27 to -25 ‰ to with a peak to -23.5 ‰ at 15 m.

290 In the first 10 cm of sediments, δ¹³C_{SOC} values increased downwards from ~ -25.5 to -23.2 ‰ and C:N molar ratio
 291 from 9.8 to 11 (Figs. 3, 4; Table S3). Carbonates corresponded to aragonite and calcite and had a bulk C isotope
 292 composition averaging 2.6 ‰ (Table S3). Porewaters from the 2016 La Preciosa core were not retrieved.

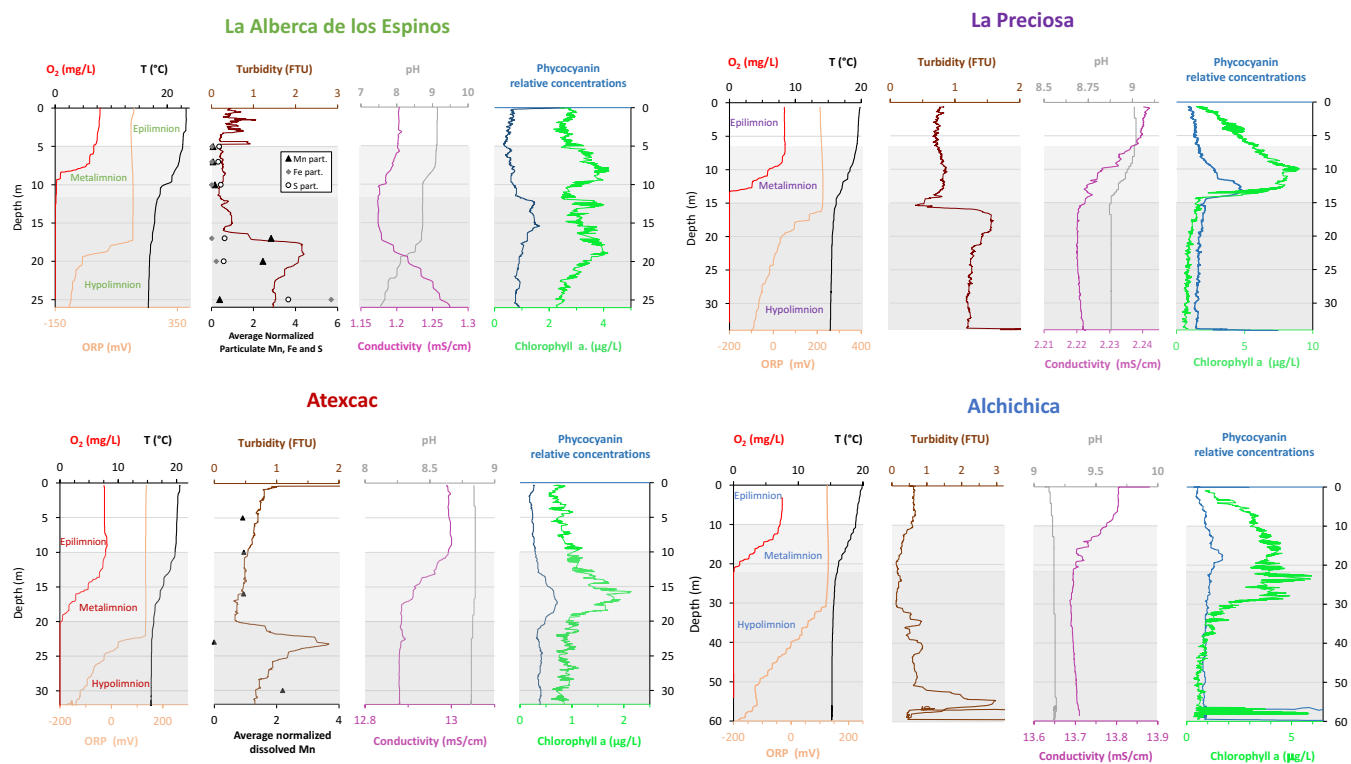


Figure 2. Physico-chemical parameters depth profiles in La Alberca de los Espinos , La Preciosa, Atexcac and Alchichica in May 2019 including: dissolved oxygen concentration (mg/L), water temperature (°C), oxidation-reduction potential (ORP, mV), turbidity (Formazin Turbidity Unit), pH, conductivity (mS/cm), phycocyanin and chlorophyll a pigments (µg/L). Absolute values for phycocyanin concentrations were not determined; only relative variations are represented (with increasing concentrations to the right). Discrete concentration values of dissolved Mn in Atexcac and particulate Mn, Fe and S in La Alberca, normalized by their respective average are represented. Epi-, meta- and hypo-limnion layers are depicted for each lake according to temperature profiles. The three layers closely corresponded to oxygen-rich, oxygen-poor and intermediate zones (except in La Preciosa where the oxycline was slightly thinner than the thermocline layer, ~5 vs. 8 m).

293

294 **4.3. Lake Atexcac**

295 Stratification of the Lake Atexcac water column was also very well defined (Fig. 2). Temperature decreased from
 296 ~ 20.6 °C at the surface to reach 16 °C below 20 m. Conductivity showed the same trend with values between 13
 297 and 12.8 mS/cm near the surface (salinity around 7.4 psu). Dissolved O₂ was slightly oversaturated at the lake

298 surface (115 % or 7.6 mg/L), rapidly decreasing to 0 mg/L between ~ 10 and 20 m, while ORP signal decreased
299 below a depth of 22 m. Chl a averaged 1 µg/L and showed a narrow peak centered at around 16 m, reaching
300 ~2 µg/L. Turbidity showed a pronounced increase below 20 m, peaking at 23.3 m and returning to surface values
301 at 26 m. The pH remained around 8.85 throughout the water column. Based on the temperature profiles, the epi-,
302 meta- and hypolimnion of Atexcac in May 2019 broadly extended from 0-10, 10-20 and 20-39 m, respectively
303 (Fig. 2).

304 The DIC concentration was around 26 mM throughout the water column, except at 23 m where it decreased to
305 24.2 mM (Fig. 3, Table S1). Calculated pCO₂ was about five times higher than the atmospheric pCO_{2atm} (Table S2).
306 The δ¹³C_{DIC} was stable around 0.4 ‰ in the epi-/metalimnion, but increased to 0.9 ‰ at 23 m and reached 0.2 ‰
307 minimum values at the bottom of the lake. The POC concentration was ~ 0.05 mM in the epi-/metalimnion,
308 decreasing to 0.02 mM in the hypolimnion. The C:N molar ratio of POM showed the same depth profile, decreasing
309 from ~9.6 in the epi-/metalimnion to 6.6 in the hypolimnion (Fig. 3). The δ¹³C_{POC} showed minimum values in the
310 epi-/metalimnion (-29.3 ‰ at 16 m) and increased to -26.5 ‰ in the hypolimnion.

311 Dissolved sulfate concentration was relatively stable at ~ 2.51 mM throughout the water column but increased to
312 2.64 mM at 23 m. Dissolved Mn concentration was constant at 1 µM down to 16 m before dropping to 0 at 23 m,
313 and increasing again to 2.35 µM at 30 m (Fig. 2; Table S4). Similar depth profiles were found for other heavy
314 elements as well, including Cu, Sr, Ba or Pb among others.

315 In the first 12 cm of sediments, DIC concentration in the porewater varied between ~ 21 and 26 mM, and δ¹³C_{DIC}
316 was around 0 ‰. Carbonates corresponded to aragonite and calcite and had a bulk C isotope composition between
317 2.1 and 2.6 ‰ (Table S3). Sedimentary organic matter had a δ¹³C_{SOC} averaging -26.8 ± 0.1 ‰ and a C:N molar
318 ratio increasing from 8 to 10 (Figs. 3, 4; Table S3).

319

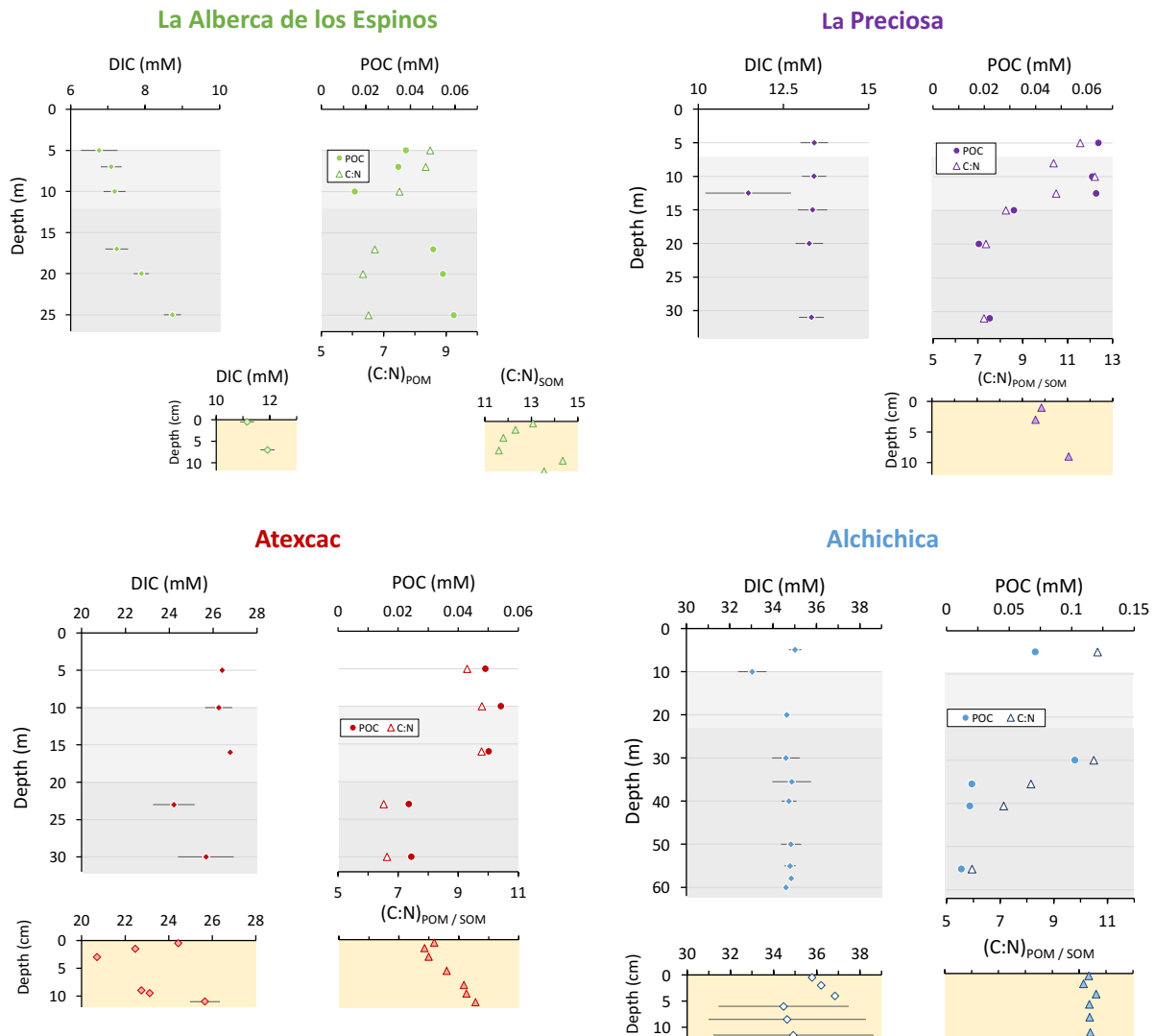


Figure 3. Concentrations in mmol/L (mM) of DIC, DOC, POC and sum of all three reservoirs, C:N molar ratios of POM as a function of depth in the water columns, as well as DIC concentrations in the surficial sediment porewaters and C:N molar ratios of sedimentary OM. Porewaters from La Preciosa's 2016 core were not retrieved.

320

321 4.4. Lake Alchichica

322 The water column of Lake Alchichica showed a pronounced stratification compared to previous years at the same
 323 period (Fig. 2, Fig. S2; Lugo et al., 2000; Adame et al., 2008; Macek et al., 2020). Temperature decreased from
 324 ~ 20 °C at the surface to 15.5 °C at depths below 30 m. Conductivity showed the same trend with values between
 325 around 13.8 mS/cm (salinity decreasing from 7.9 to 7.8 psu). Dissolved O₂ was slightly oversaturated at the lake
 326 surface (112 % or 7.5 mg/L), rapidly decreasing to 0 mg/L between ~ 10 and 20 m. The ORP followed a similar
 327 trend but decreasing below 30 m only. The offset between O₂ exhaustion and decrease of the ORP can be explained
 328 by the presence of other oxidant species and/or extended Chl a peaks (supplementary text 1). Chl a averaged
 329 2 µg/L, with a broad peak extending from ~ 7 to 29 m (averaging 4 µg/L and showing a narrow 6 µg/L maximum
 330 values at 23 m). Then, it decreased to minimum values of ~ 0.5 µg/L in the lower water column. The pH remained

331 constant at ~ 9.2 over the whole water column. Based on the temperature profiles, the epi-, meta- and hypolimnion
 332 layers of Lake Alchichica in May 2019 extended from 0-10, 10-20 and 20-63 m, respectively (Fig. 2).

333 The DIC concentration was around 34.8 mM throughout the water column, except at 10 m where it decreased to
 334 33 mM (Fig. 3; Table S1). Calculated $p\text{CO}_2$ was about three times higher than the atmospheric $p\text{CO}_{2\text{atm}}$ (Table S2).
 335 The $\delta^{13}\text{C}_{\text{DIC}}$ decreased from 2 to ~ 1.5 ‰ between 5 and 60 m depth (Fig. 4; Table S1). The POC concentration
 336 was ~ 0.09 mM in the epi-/metalimnion, decreasing to 0.02 mM in the hypolimnion. The $\delta^{13}\text{C}_{\text{POC}}$ increased from
 337 -26.5 ‰ in the top 30 m to -24.1 ‰ at 55 m. The C:N molar ratio of POM showed a similar profile with values
 338 around 10.5 down to 30 m, progressively decreasing towards 5.9 at 55 m (Fig. 3; Table S1).

339 In the first 12 cm of sediments, porewater DIC had a concentration of ~ 35.5 mM and $\delta^{13}\text{C}_{\text{DIC}}$ decreased from 0.4
 340 to -0.5 ‰. Solid carbonates were contained within several phases (aragonite, hydromagnesite, huntite and calcite)
 341 and had a bulk C isotope composition around 4.6 ‰ (Table S3). Sedimentary organic matter had a $\delta^{13}\text{C}_{\text{SOC}}$
 342 increasing from -25.7 to -24.5 ‰ and a constant C:N molar ratio slightly higher than 10 (Figs. 3, 4; Table S3).

343

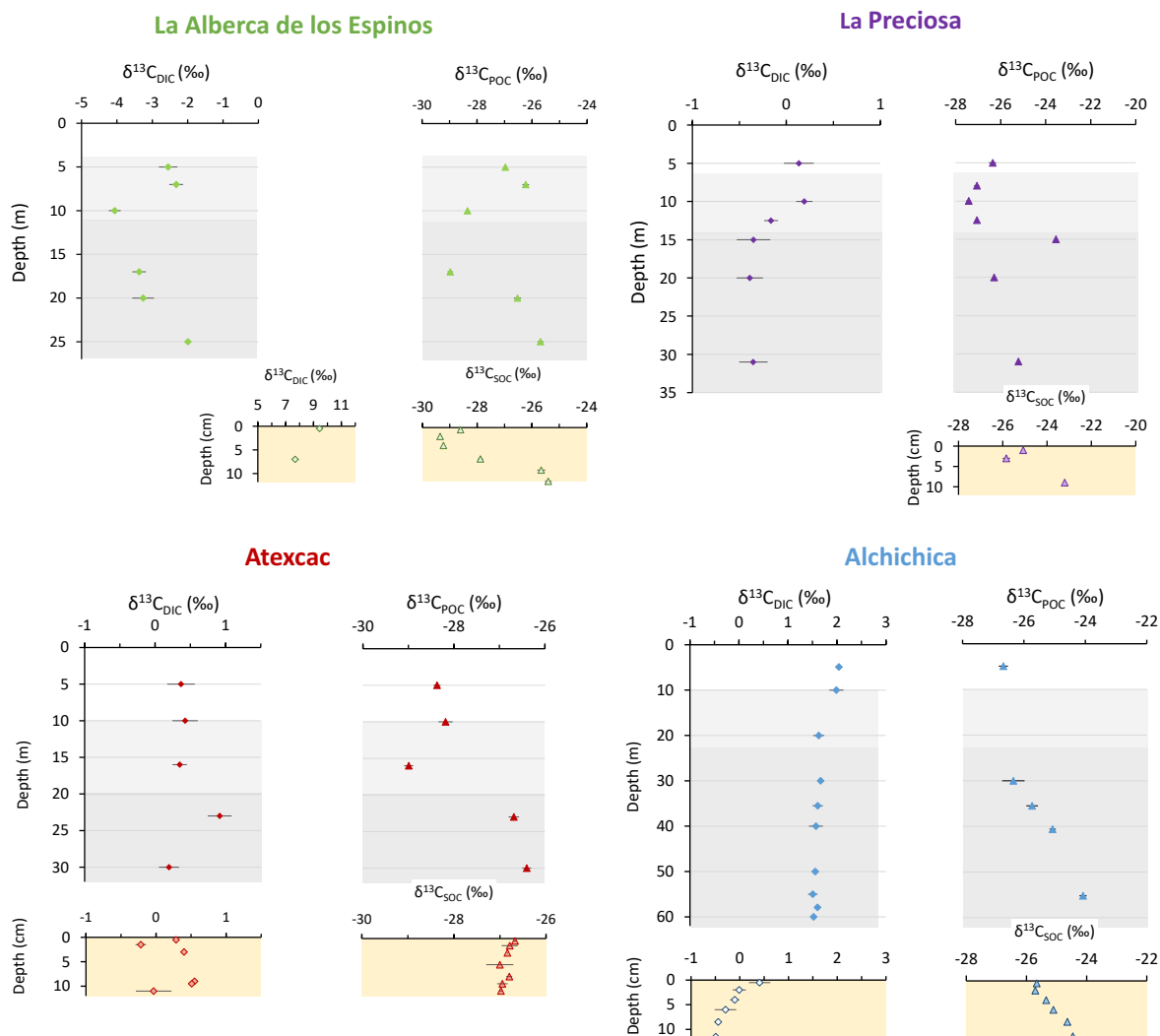


Figure 4. Isotopic compositions of DIC and POC reservoirs as a function of depth in the water columns, as well as isotopic compositions of the porewater-DIC and total organic carbon from the surficial sediments.

344 5. DISCUSSION

345 346 5.1. Inorganic Carbon: origins and implications of the alkalinity/DIC gradient

347 348 5.1.1 Sources of DIC and origin of the inter-lake alkalinity gradient

349 Salinity and DIC concentration gradually increase from La Alberca de los Espinos (0.6 psu, 7 mM) to Alchichica
350 (7.9 psu, 35 mM), while La Preciosa (1.15 psu, 13 mM) and Atexcac (7.44 psu, 26 mM) have intermediate values
351 (Table 1 and S1). This trend matches the alkalinity gradient (with values of ~ 8, 15, 32 and 47 meq/L, Fig. S3a)
352 previously described for these lakes (Zeyen et al., 2021), consistent with the fact that alkalinity is mainly composed
353 of HCO_3^- and CO_3^{2-} ions in most natural waters. This alkalinity gradient may result from different concentration
354 stages of an initial dilute alkaline water (Zeyen et al., 2021), ultimately controlled by differences in hydrological
355 regime between the four lakes. In the SOB, the weathering of basaltic/andesitic bedrock (Armienta et al., 2008;
356 Carrasco-Núñez et al., 2007; Lelli et al., 2021) and Cretaceous limestone (with $\delta^{13}\text{C} \approx 0 \pm 1 \text{‰}$; Gonzales-Partida
357 et al., 1993; Armstrong-Altrin et al., 2011) favors the inflow of more alkaline and DIC-concentrated groundwater
358 than in La Alberca, which lies on an essentially basaltic basement (Rendon-Lopez, 2008; Siebe et al., 2014; Zeyen
359 et al., 2021). The SOB is currently experiencing higher rates of evaporation than precipitation (Alcocer, 2021),
360 which may play an important role in concentrating solutes and decreasing the water level in La Preciosa, Atexcac,
361 and Alchichica (Anderson and Stedmon, 2007; Zeyen et al., 2021). Substantial “sub-fossil” microbialite deposits
362 emerge well above the current water level in lakes Atexcac and Alchichica, confirming this fall in water level
363 (~15 m for Atexcac, and ~5 m for Alchichica). Scattered patches of microbialites emerge at La Preciosa
364 (suggesting a water level decrease of ~6 m). By contrast, emerged microbialites are virtually absent in Lake La
365 Alberca de los Espinos (Fig. S1).

366 Additional local parameters such as variable groundwater paths and fluxes (Furian et al., 2013; Mercedes-Martín
367 et al., 2019; Milesi et al., 2020; Zeyen et al., 2021) most likely play a role in explaining some of the variation in
368 DIC concentration between lakes. La Preciosa’s water composition significantly differs from that of Atexcac, and
369 Alchichica, despite a similar geological context and climate (all are located within 50 km², Fig. 1). Groundwater
370 in the SOB area becomes more saline as it flows towards the center of the basin and through the crater lakes (Silva
371 Aguilera, 2019; Alcocer, 2021). Since groundwater flows through La Preciosa first, its ionic strength (including
372 DIC concentration) increases as it enters Alchichica (Silva Aguilera, 2019; Alcocer, 2021; Lelli et al., 2021).
373 Different regimes of volcanic CO₂ degassing into these crater lakes may also contribute to variation in the C mass
374 balance and $\delta^{13}\text{C}_{\text{DIC}}$ values between the four lakes. Near the lakes from the SOB area, geothermal fluids derived
375 from meteoric waters have been shown to interact with deep volcanic fluids as well as the calcareous basement
376 rocks (Peiffer et al., 2018; Lelli et al., 2021). In the water column of La Alberca, $\delta^{13}\text{C}_{\text{total}}$ averages -4.8 ‰ (Havas
377 et al., submitted). This isotopic composition is very similar to signatures of mantle-CO₂ (Javoy et al., 1986; Mason
378 et al., 2017), which could buffer the overall C isotope composition of this lake. La Alberca is located on top of a
379 likely active normal fault (Siebe et al., 2012), favoring the ascent of volcanic gases.

380 Differences in the remineralization rate of organic carbon (OC) could also contribute to the heterogeneous DIC
381 content among the lakes. However, assuming that all OC from the lakes ultimately remineralized into DIC, it
382 would still represent only a small proportion of the total carbon (9 % for La Alberca, ~5 % for La Preciosa and
383 Alchichica, and 16 % for Atexcac, Havas et al., submitted). From an isotopic mass balance perspective, Lake La

384 Alberca exhibits more negative $\delta^{13}\text{C}_{\text{DIC}}$ (and $\delta^{13}\text{C}_{\text{carb}}$), slightly closer to OC signatures, whereas the $\delta^{13}\text{C}_{\text{DIC}}$ of the
385 three SOB lakes lie very far from OC isotopic signatures (Fig. 4). Dense vegetation surrounds La Alberca (Fig. S1),
386 making it the only lake in this study where OC respiration could be a significant source of inorganic C to the water
387 column (potentially influencing the P_{CO_2} , [DIC] and pH profiles described above).

388 In summary, a combination of very local and external environmental factors generates the contrasting water
389 chemistries of the lakes, notably a gradient in their alkalinity/[DIC]. This chemical variability stems from the exact
390 nature of the basement rocks, the distinct groundwater flow paths feeding the lakes, differences in evaporation
391 rates, and potentially different volcanic- CO_2 degassing regimes.

392

393 **5.1.2 Influence of alkalinity on physico-chemical stratification in the four lakes**

394 Stratified water columns can sustain strong physico-chemical gradients, where a wide range of biogeochemical
395 reactions impacting the C cycle can take place (e.g. Jézéquel et al., 2016). In the four lakes studied here, the
396 evolution of pH with depth exemplifies the interplay between the alkalinity gradient, the physico-chemical
397 stratification of the lakes, and their respective C cycle. The pH shows a stratified profile in La Alberca and La
398 Preciosa, but remains constant in Atexcac and Alchichica. The decline in pH at the oxycline of La Preciosa is
399 associated with the decrease in POC and chlorophyll a concentrations and $\delta^{13}\text{C}_{\text{DIC}}$ values, reflecting the impact of
400 oxygen respiration (i.e. carbon remineralization) at this depth (Figs. 2-4). In La Alberca, the surface waters are
401 markedly more alkaline than the bottom waters, with a two-step decrease in pH occurring at 8 m and 17 m (with a
402 total drop of 1.5 pH unit). As in La Preciosa, this pH decrease likely results from high OM respiration, although
403 input of volcanic acidic gases (e.g. dissolved CO_2 with $\delta^{13}\text{C} \sim -5\text{‰}$) might also contribute to the pH decrease in
404 the bottom waters, reflected by negative $\delta^{13}\text{C}_{\text{DIC}}$ signatures and an increase of [DIC] and conductivity in the
405 hypolimnion (Figs. 2 and 4). By contrast, while the same evidence for oxygen respiration ([POC], chlorophyll a)
406 can be detected in the other two lakes, it does not impact their pH profile in a similar way (Fig. 2). This result
407 suggests that the acidity generated by OM respiration (and possibly volcanic- CO_2 degassing) is buffered by the
408 much higher alkalinity measured in these two lakes.

409 External forcings such as lake hydrology and fluid sources thus impact the alkalinity buffering capacity of these
410 lakes and influence the vertical pH profile of the water columns, which is particularly important considering the
411 critical interplay between pH and biogeochemical reactions affecting the C cycle (e.g. Soetaert et al., 2007).

412

413 **5.1.3 Sinks of DIC along the alkalinity gradient**

414 Interplay between pH and sources of alkalinity/DIC in the lakes also has a strong impact on their C storage capacity
415 as it can result in different fluxes of the C sinks (inorganic and organic C precipitation / sedimentation, CO_2
416 degassing).

417 Alkaline pH can store large quantities of DIC because it favors the presence of HCO_3^- and CO_3^{2-} species over
418 H_2CO_3^* (the intermediate species between gaseous $\text{CO}_{2(\text{g})}$ and bi-/carbonate ions, defined here as the sum of H_2CO_3
419 and $\text{CO}_{2(\text{aq})}$). Carbonate and bicarbonate ions represent over 99% of total DIC in the four lakes (Table S2). In La

420 Alberca de los Espinos, the lake with the lowest DIC, the surface water pCO₂ is slightly lower than atmospheric
421 pCO_{2atm} (Table S2). By contrast, large amounts of CO₂ degas at the surface of the SOB lakes, as indicated by their
422 elevated surface water pCO₂, from 2 to 5 times higher than atmospheric pCO_{2atm} (Table S2). These different CO₂
423 degassing potentials are consistent with the notion that higher DIC concentrations favor CO₂ degassing through
424 higher pCO₂ (e.g. Duarte et al., 2008). Although La Alberca and Alchichica (the two endmembers of the alkalinity
425 gradient) have the same surface water pH, CO₂ degassing is three times higher at Alchichica, for a given value of
426 gas transfer velocity.

427 Another important C sink for these lakes is the precipitation of carbonate minerals, found in the microbialites and
428 lake sediments. Lake alkalinity and resulting mineral saturation index greatly influence the amount of C
429 precipitated from the lake waters. Although the four lakes are supersaturated with aragonite, calcite and the
430 precursor phase monohydrocalcite, they present highly contrasted amounts of carbonate deposits (Zeyen et al.,
431 2021). The occurrence of microbialites increases along the alkalinity gradient, with limited presence at La Alberca,
432 and more massive deposits at Atexcac and Alchichica (Zeyen et al., 2021; Fig. S1). Similarly, surficial sediments
433 contain only 16 wt. % for La Alberca, but from 40 to 62 wt. % carbonates for the SOB lakes (Table S3). Thus, the
434 SOB lakes seem to bury more C than La Alberca de los Espinos. Nonetheless, the data from May 2019 indicate
435 that La Alberca was the only one of the four lakes with a pCO₂ slightly lower than atmospheric pCO_{2atm}, thus
436 representing a net sink of C. Classifying the three other lakes as net C sources or sinks – notably in order to see
437 the influence of their respective position in the alkalinity gradient – will require a more detailed description of C
438 in- and out-fluxes since they all store and emit significant amounts of C (as organic and inorganic C deposits and
439 via CO₂ degassing, respectively). However, such a C budget is out of the scope of the present study.

440

441 **5.1.4 Isotopic signatures of inorganic C in the four lakes ($\delta^{13}\text{C}_{\text{DIC}}$ and $\delta^{13}\text{C}_{\text{Carbonates}}$)**

442 The DIC isotopic composition of the lakes (between ~ -3 and +2 ‰ on average; Table S1) is consistent with the
443 DIC sources described above. The lower $\delta^{13}\text{C}_{\text{DIC}}$ in La Alberca is consistent with influence of remineralized OC
444 and/or volcanic CO₂. The $\delta^{13}\text{C}_{\text{DIC}}$ in the SOB lakes suggests groundwater $\delta^{13}\text{C}_{\text{DIC}}$ values resulting from the
445 dissolution of the Cretaceous limestone basement.

446 By controlling DIC speciation (H₂CO₃/CO_{2(aq)}, HCO₃⁻, CO₃²⁻), pH also strongly influences $\delta^{13}\text{C}_{\text{DIC}}$. Indeed, there
447 is a temperature-dependent fractionation of up to 10 ‰ between the different DIC species (Emrich et al., 1970;
448 Mook et al., 1974; Bade et al., 2004; Table S6). The Mexican lakes present $\delta^{13}\text{C}_{\text{DIC}}$ values that are common for
449 lakes with a pH around 9 (Bade et al., 2004), where DIC is dominated by HCO₃⁻. However, the pH values of the
450 four lakes studied here are too similar to explain the significant difference between their $\delta^{13}\text{C}_{\text{DIC}}$ (Fig. 4; $p=4.2 \times 10^{-3}$
451 for La Preciosa and Atexcac, which have the closest $\delta^{13}\text{C}_{\text{DIC}}$). Part of the variability of $\delta^{13}\text{C}_{\text{DIC}}$ among the lakes
452 may result from their distinct evaporation stages, as the mean $\delta^{13}\text{C}_{\text{DIC}}$ values of the lakes broadly correlate with
453 their salinity/alkalinity (Fig. S3b). Evaporation generally increases the $\delta^{13}\text{C}_{\text{DIC}}$ of residual waters by increasing
454 lake pCO₂ and primary productivity, which bolsters CO₂ degassing and organic C burial, both having low $\delta^{13}\text{C}$
455 compared to DIC (e.g. Li and Ku, 1997; Talbot, 1990). Accordingly, the pCO₂ of La Alberca is lower than that of
456 the other lakes (Table S2). The $\delta^{13}\text{C}_{\text{DIC}}$ in lakes with lower DIC concentrations is expected to be more easily
457 influenced by exchanges with other carbon reservoirs, such as organic carbon (through photosynthesis/respiration),

458 or other DIC sources (e.g., depleted volcanic CO₂ or groundwater DIC), compared with buffered, high DIC lakes
 459 (Li and Ku, 1997). As a result, the low DIC/alkalinity concentration in La Alberca features the lowest δ¹³C_{DIC} of
 460 the four lakes, likely reflecting organic and/or volcanic C influence and thus higher responsiveness to
 461 biogeochemical processes of the inorganic C reservoir. By contrast, the three SOB lakes exhibit δ¹³C_{DIC} with less
 462 internal variability, with a maximum amplitude of 0.7 ‰ within a single water column.

463 Surficial sedimentary carbonates are in isotopic equilibrium with the δ¹³C_{DIC} of the water columns, within the
 464 uncertainty of δ¹³C_{DIC} measurement, and more specifically with the δ¹³C_{DIC} values at the oxycline/thermocline of
 465 the lakes (Tables S6 and S7). The δ¹³C_{DIC} at equilibrium with carbonates is estimated by correcting the carbonates
 466 C isotope composition (δ¹³C_{carb}) by the fractionation value between DIC and the different carbonate mineralogies
 467 (supplementary text S2). Therefore, the δ¹³C_{carb} also follows and reflects the alkalinity gradient, with the lowest
 468 δ¹³C_{carb} found in the surficial sediments of La Alberca (~ -1.5 ‰), intermediate values in La Preciosa and Atecac
 469 (~2.5 ‰), and the highest values in Alchichica (~ +4.6 ‰) (Table S3).

470

471 In summary, although all four lakes present the same general structure and environmental conditions (i.e. tropical
 472 alkaline stratified crater lakes), external and local factors (e.g. hydrology, fluid sources, and stratification
 473 characteristics) result in contrasting water chemistry compositions, which have a critical impact on the physico-
 474 chemical depth profiles of each lake and their biogeochemical carbon cycle functioning. These external factors
 475 represent a first-order control on the size, isotopic composition, and responsiveness to biogeochemical processes
 476 of the inorganic C reservoir. Lakes with the highest alkalinity/DIC content will poorly record internal biological
 477 processes. Interestingly, C storage in mineral carbonates seems to be significant in watersheds where carbonate
 478 deposits pre-exist in the geological substratum (here, the Cretaceous limestone basement), providing more alkaline
 479 and C-rich sources.

480

Symbols	Mathematical Expression	Signification
δ ¹³ C _X	$\left(\frac{\left(\frac{^{13}\text{C}}{^{12}\text{C}} \right)_X}{\left(\frac{^{13}\text{C}}{^{12}\text{C}} \right)_{VPDB}} - 1 \right) * 1000$	Relative difference in ¹³ C: ¹² C isotopic ratio between a sample of a given C reservoir and the international standard "Vienna Pee Dee Bee", expressed in permil (‰). δ ¹³ C _{total} represents the weighted average of δ ¹³ C for all DIC and POC.
Δ ¹³ C _{X-Y}	$= \delta^{13}\text{C}_X - \delta^{13}\text{C}_Y \approx 1000 \ln \alpha_{X-Y}$	Apparent isotopic fractionation between two reservoirs 'X' and 'Y'. Difference between their measured C isotope compositions approximating the fractionation α in ‰.
ε _{X-CO2}	$= (\alpha_{X-\text{CO}_2} - 1)1000 \approx \delta^{13}\text{C}_X - \delta^{13}\text{C}_{\text{CO}_2}$	Calculated isotopic fractionation between a reservoir 'X' and CO _{2(aq)} . α _{X-CO2} is calculated as (δ ¹³ C _X +1000)/(δ ¹³ C _{CO2} +1000) where δ ¹³ C _X is measured and δ ¹³ C _{CO2} is computed based on DIC isotopic composition and speciation (see supplementary text S3).

481

482 Table 2
483 Index for mathematical notations used in the text including C isotopic composition of a reservoir X ($\delta^{13}\text{C}_X$),
484 isotopic discrimination between the two carbon reservoirs X and Y ($\Delta^{13}\text{C}_{X-Y}$). In the main text, we report organic
485 C isotope discrimination *versus* both bulk DIC ($\Delta^{13}\text{C}_{\text{POC-DIC}}$) – in a way to facilitate studies intercomparison and
486 because it is the commonly reported raw measured data (Fry, 1996) – and calculated $\text{CO}_{2(\text{aq})}$ ($\epsilon_{\text{POC-CO}_2}$) in order to
487 discuss the intrinsic isotopic fractionations associated with the lakes metabolic diversity. All C isotope values and
488 fractionations are reported relative to the international standard VPDB (Vienna Pee Dee Belemnite).

489

490

491 **5.2. Particulate organic carbon: from water column primary production to respiration recycling and** 492 **sedimentary organic matter**

493

494 **5.2.1. Particulate organic C sources**

495 *Primary productivity by oxygenic photosynthesis in the upper water column*

496 The C:N ratios of water-column POM and bulk organic matter in the sediments of the four lakes ranged from 6 to
497 13 (Fig. 3), close to the phytoplankton Redfield but much lower than land plant ratios. Yet abundant vegetation
498 covers the crater walls of La Alberca and, to a lesser extent, of Atexcac, and some plant debris was found in the
499 sediment cores of these two lakes. Its analysis resulted in high C:N ratios (between 24 and 68), typical of plant
500 tissues and significantly higher than those of the bulk sediment and water column organic matter. Thus, the
501 allochthonous organic carbon in these two lakes does not significantly contribute to their bulk organic signal. All
502 four crater lakes are endorheic basins, with no surface water inflow or outflow, supporting the predominantly
503 autochthonous origin of organic carbon sources (Alcocer et al., 2014b), from planktonic autotrophic C fixation.

504 The importance of planktonic autotrophic C fixation as a major source of POC in the four lakes is further supported
505 by the assessment of the isotopic discrimination between DIC and organic biomass, expressed as $\Delta^{13}\text{C}_{\text{POC-DIC}}$ and
506 $\epsilon_{\text{POC-CO}_2}$ (Table 2). The $\Delta^{13}\text{C}_{\text{POC-DIC}}$ varies between ~ -29 and -23 ‰ (corresponding to $\epsilon_{\text{POC-CO}_2}$ between ~ -19 and
507 -13 ‰) throughout the four water columns, within the typical range of planktonic oxygenic phototrophs (Pardue
508 et al., 1976; Sirevag et al., 1977; Thomas et al., 2019). Yet these values exhibit variability – both within a single
509 water column (up to 4.5 ‰) and among the four lakes (up to 6 ‰, Figs. 4 and 5). The $\Delta^{13}\text{C}_{\text{POC-DIC}}$ variability may
510 reflect several abiotic and biotic factors.

511 Notably, lower DIC availability in La Alberca and La Preciosa probably makes the carboxylation step less limiting
512 during photosynthesis (e.g. O’Leary, 1988; Descolas-Gros and Fontungne, 1990; Fry, 1996), decreasing $|\epsilon_{\text{POC-CO}_2}|$
513 in these lakes (between 14.5 and 17.7 ‰ at the peak of Chl. a) compared with Atexcac and Alchichica (Fig. 5a;
514 between 17.5 and 19.2 ‰). Lower $\text{CO}_{2(\text{aq})}$ availability and/or higher reaction rates result in transport-limited rather
515 than carboxylation-limited fixation, with smaller C isotope fractionation between POC and DIC (Pardue et al.,
516 1976; Zohary et al., 1994; Fry, 1996; Close and Henderson, 2020). The isotopic fractionation associated with
517 diffusion is much smaller than with carboxylation, and a higher proportion of the DIC entering the cells is
518 converted into organic biomass (e.g. Fogel and Cifuentes, 1993). We consistently notice a correlation among the

519 lakes between $a(\text{CO}_2)_{\text{aq}}$ (or [DIC]) and $|\epsilon_{\text{POC-CO}_2}|$ at depths where oxygenic photosynthetic peaks (Fig. 6).
 520 Furthermore, La Alberca and La Preciosa are considered less oligotrophic than the two other lakes (Lugo et al.,
 521 1993; Vilaclara et al., 1993; Havas et al., submitted), with higher chlorophyll a contents and thus smaller $|\epsilon_{\text{POC-CO}_2}|$
 522 (Fig. 5). Higher water temperatures in La Alberca de los Espinos (by ~ 3 °C) could also partly contribute to a
 523 smaller $|\epsilon_{\text{POC-CO}_2}|$ in this lake (Sackett et al., 1965; Pardue et al., 1976; Descolas-Gros and Fontungne, 1990).

524 Unlike $\delta^{13}\text{C}_{\text{DIC}}$, organic carbon isotope signatures do not evolve linearly with the alkalinity/salinity gradient,
 525 suggesting other lake- and microbial-specific controls on these signatures. These controls include: diffusive or
 526 active uptake mechanisms, specific carbon fixation pathways, the fraction of intracellular inorganic carbon
 527 released out of the cells, cell size and geometry (Werne and Hollander, 2004 and references therein) and
 528 remineralization efficiency. Moreover, an increasing amount of isotopic data has evidenced a significant variability
 529 of the isotopic fractionation achieved by different purified RuBisCO enzymes ($\epsilon_{\text{RuBisCO}}$, Iñiguez et al., 2020), and
 530 even by a single RuBisCO form (Thomas et al., 2019). Thus, caution should be paid to the interpretation of the
 531 origin of small isotopic variations of the biomass in distinct environmental contexts because RuBisCO alone can
 532 be an important source of this variability (Thomas et al., 2019).

533

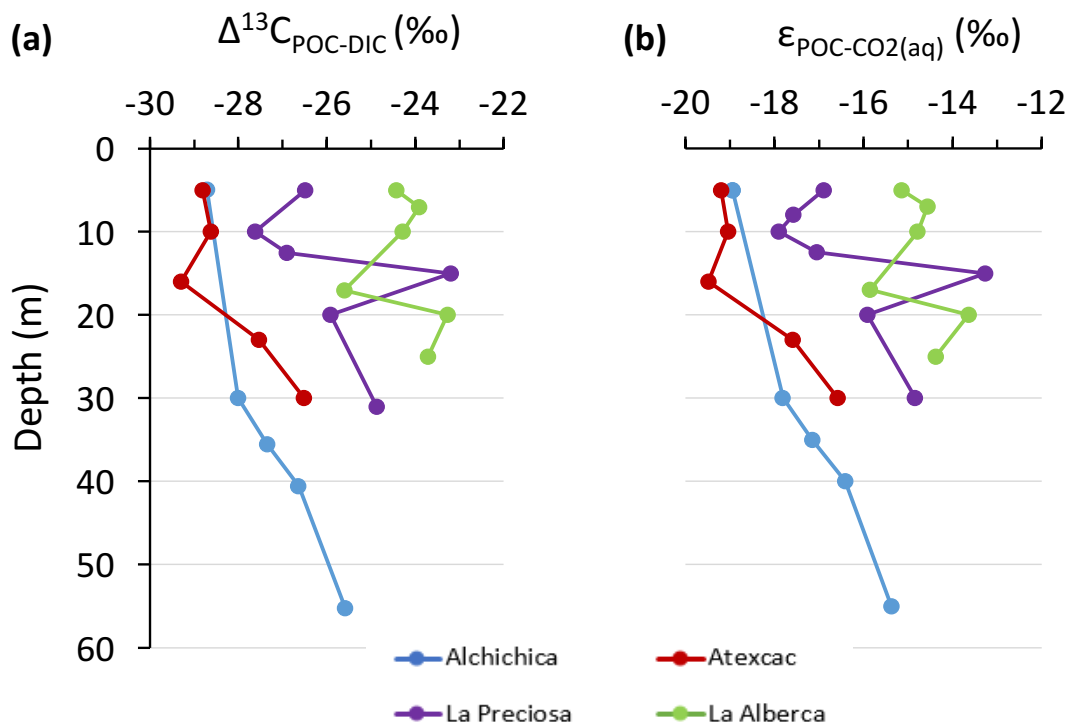


Figure 5. Isotopic fractionations between POC and DIC in the water columns of the four lakes, expressed as a) $\Delta^{13}\text{C}_{\text{x-y}}$ and b) $\epsilon_{\text{POC-CO}_2}$. Refer to Table 2 for more detail about the Δ and ϵ notations.

535 Anoxygenic autotrophs commonly thrive in anoxic
 536 bottom waters of stratified water bodies (e.g.
 537 Pimenov et al., 2008; Zyakun et al., 2009; Posth et
 538 al., 2017; Fulton et al., 2018; Havig et al., 2018).
 539 They have been identified at different depths in the
 540 four Mexican lakes (Macek et al., 2020; Iniesto et al.,
 541 2022). In our samples collected during the
 542 stratification period, anoxygenic autotrophs appear to
 543 have a distinct impact on the C cycle of La Alberca
 544 and Atexcac only. Lake Atexcac records a
 545 concomitant decrease in [DIC] and increase in
 546 $\delta^{13}\text{C}_{\text{DIC}}$ in the anoxic hypolimnion at 23 m, below the
 547 peak of Chl a, suggesting autotrophic C fixation by
 548 chemoautotrophy or anoxygenic photosynthesis. The
 549 calculated $\epsilon_{\text{POC-CO}_2}$ at 23 m (-17.3 ‰) is consistent
 550 with C isotope fractionation by purple- and green-
 551 sulfur-anoxygenic bacteria (PSB and GSB), while
 552 $\epsilon_{\text{POC-CO}_2}$ in La Alberca's hypolimnion (~ -15 ‰) is
 553 closer to GSB canonical signatures (Posth et al., 2017
 554 and references therein) (Fig. 5b). In La Alberca,
 555 anoxygenic primary productivity is suggested by
 556 increasing POC concentrations below the oxycline,
 557 showing a distinct isotopic signature (Figs. 4 and 5).
 558 We also observe a Chl a peak in the anoxic
 559 hypolimnion of this lake (Fig. 2), which likely
 560 represents a bias of the probe towards some
 561 bacteriochlorophyll pigments typical of GSB (see
 562 supplementary text S4). In Atexcac, C fixation by
 563 anoxygenic autotrophs at 23 m causes a shift in the DIC
 564 reservoir, while oxygenic photosynthesis at 16 m does
 565 not, suggesting that anaerobic autotrophs are the main
 566 autotrophic metabolisms in this lake (in terms of DIC
 567 uptake). In La Alberca, the increase in [POC] to
 568 maximum values below the oxycline also supports the
 569 predominance of anoxygenic *versus* oxygenic
 570 autotrophy (Fig. 3), similarly to other stratified water
 571 bodies exhibiting primary production clearly dominated
 572 by anoxygenic metabolisms (Fulton et al., 2018).

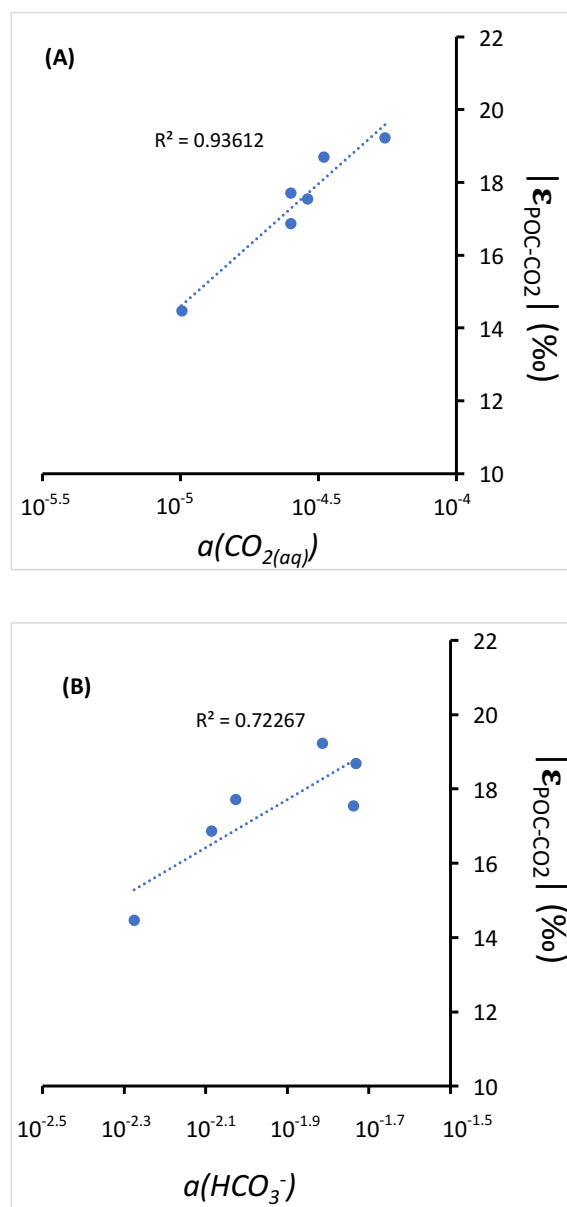


Figure 6.

Cross plots of DIC species activities *versus* absolute values of calculated C isotopic fractionations between POC and CO_2 at depths of peak oxygenic photosynthesis where data was available (5 and 30 m for Alchichica, 16 m for Atexcac, 10 and 12.5 m for La Preciosa and 7 m for La Alberca). (A) Dissolved $\text{CO}_{2(\text{aq})}$ activity and (B) bicarbonate activity as functions of $|\epsilon_{\text{POC-CO}_2}|$ in ‰ plus linear correlation trends and corresponding R^2 .

573 Lastly, at 23 m in Atexcac and 17 m in La Alberca, we find a striking turbidity peak precisely where the redox
574 potential and the concentration of dissolved Mn drops (Fig. 2). In Atexcac, the concentration in dissolved metals
575 such as Cu, Pb, or Co also drops at 23 m (Fig. S4). In La Alberca, a peak of particulate Mn concentration is detected
576 at 15 m (Fig. 2; data unavailable for Atexcac). This peak is most likely explained by the precipitation of Mn mineral
577 particles, where reduced bottom waters meet oxidative conditions prevailing in the upper waters. These oxidized
578 Mn phases can be used as electron acceptors during chemoautotrophy (Havig et al., 2015; Knossow et al., 2015;
579 Henkel et al., 2019; van Vliet et al., 2021). Even at a low particle density, such phases can catalyze abiotic oxidation
580 of sulfide to sulfur compounds, which in turn can be used and further oxidized to sulfate by phototrophic or
581 chemoautotrophic sulfur-oxidizing bacteria (van Vliet et al., 2021). Autotrophic sulfur oxidation is also consistent
582 with the small increase in $[\text{SO}_4^{2-}]$ observed at 23 m in Atexcac (Table S4).

583 In summary, combined POC and DIC data allowed us to recognize the most representative autotrophic
584 metabolisms in the Mexican lakes. The upper water columns are all dominated by oxygenic photosynthesis. Lower
585 in the water columns, anoxygenic photosynthesis and/or chemoautotrophy were found to have a noticeable impact
586 on POC and DIC reservoirs in La Alberca and Atexcac only. Their activity was associated with metal elements
587 cycling. More specifically in La Alberca, the anoxygenic phototrophs correspond to GSB.

588

589

590 **5.2.2. Sinks of particulate organic carbon: respiration and sedimentation**

591 *Aerobic respiration at the oxycline*

592 At the oxycline of stratified water bodies, aerobic respiration of OM by heterotrophic organisms favors the
593 transition from oxygenated upper layers to anoxic bottom waters. In the water column of the four lakes, $\Delta^{13}\text{C}_{\text{POC-DIC}}$
594 (and $\epsilon_{\text{POC-CO}_2}$) show increasing values in the hypolimnion, and especially below the chlorophyll a peaks (Figs. 2
595 and 5). The $\Delta^{13}\text{C}_{\text{POC-DIC}}$ trend correlates with increasing $\delta^{13}\text{C}_{\text{POC}}$, decreasing $(\text{C:N})_{\text{POM}}$ ratios as well as decreasing
596 POC concentrations except in La Alberca (Figs. 3 and 4). Decreasing POC concentrations near the oxycline and
597 redoxcline are consistent with the fact that part of the upper primary production is degraded deeper in the water
598 columns and/or that there is less primary production in the anoxic bottom waters. Increased $\delta^{13}\text{C}_{\text{POC}}$ in the
599 hypolimnion of the lakes is consistent with heterotrophic activity and points out that POC at these depths could
600 mainly record secondary production rather than being a residue of sinking degraded OM formed by primary
601 production. Heterotrophic bacteria preferentially grow on available ^{13}C -enriched amino acids and sugars, thus
602 becoming more enriched than their C source (Williams and Gordon, 1970; Hayes et al., 1989; Zohary et al., 1994;
603 Briones et al., 1998; Lehmann et al., 2002; Jiao et al., 2010; Close and Henderson, 2020). The decrease in C:N
604 ratios in the POM also reinforces this conclusion since secondary heterotrophic bacteria biomass generally have
605 C:N between 4 and 5 (Lehmann et al., 2002), whereas residual degraded OM from primary producers would carry
606 higher C:N signatures (van Mooy et al., 2002; Buchan et al., 2014). These latter signatures are not recorded by
607 POM in the lower water columns of the lakes (Fig. 3).

608 The $\delta^{13}\text{C}_{\text{DIC}}$ signatures in La Preciosa and Alchichica are consistent with the mineralization of OM as they exhibit
609 lower values below the oxycline than in surficial waters (Figs. 2 and 4). Similarly to what is observed in several
610 other water bodies and notably stratified water columns such as the Black Sea (e.g. Fry et al., 1991), surface

611 photosynthesis increases $\delta^{13}\text{C}_{\text{DIC}}$ by fixing light DIC, while respiration transfers light OC back to the DIC pool at
612 depth. Such a decrease in $\delta^{13}\text{C}_{\text{DIC}}$ can also be seen in the oxycline of Lake La Alberca between 7 and 10 m.

613
614 *Influence of methanogenesis in Lake La Alberca de los Espinos*

615 La Alberca shows the least saline/alkaline water column and most peculiar geochemical depth profiles among the
616 four lakes. Notably, its [DIC] and $\delta^{13}\text{C}_{\text{DIC}}$ (the lowest of the studied lakes) increase from the lower metalimnion to
617 the hypolimnion, and further into the first cm of sediment porewaters, with $\delta^{13}\text{C}_{\text{DIC}}$ reaching almost 10 ‰ (Figs. 3;
618 4). The calculated CO_2 partial pressure (P_{CO_2}) increases downward from slightly less than $1 \times P_{\text{CO}_2\text{-atm}}$ near the lake
619 surface up to almost 40x at the bottom of the lake (Table S2).

620 While the increase of [POC] at depth may contribute to the observed $\delta^{13}\text{C}_{\text{DIC}}$ increase, by mass balance, it should
621 also lower the [DIC] instead of increasing it. Similarly, the sinking of POC at depth followed by its
622 remineralization into DIC cannot explain the $\delta^{13}\text{C}_{\text{DIC}}$ trend since it would lower the $\delta^{13}\text{C}_{\text{DIC}}$ in the hypolimnion
623 (Fig. 4). Overall, these observations require that a significant source of inorganic ^{13}C -rich carbon fuels the bottom
624 waters of La Alberca de los Espinos. The source of heavy carbon most likely results from methanogenesis, which
625 consumes organic carbon in the sediments and produces ^{13}C -depleted methane and ^{13}C -rich carbon dioxide
626 diffusing upward in the water column (i.e. acetoclastic methanogenesis, dominant in lacustrine contexts, Whiticar
627 et al., 1986). Methanogenesis, as an “alternative” OM remineralization pathway would be favored in La Alberca,
628 because it is relatively rich in OM (notably with high [DOC], Havas et al., submitted), and depleted in SO_4^{2-}
629 (Wittkop et al., 2014; Birgel et al., 2015; Cadeau et al., 2020) compared with the three other Mexican lakes. Based
630 on the $\delta^{13}\text{C}_{\text{SOC}}$ and porewater $\delta^{13}\text{C}_{\text{DIC}}$, we can tentatively calculate the methane isotopic signature in La Alberca
631 (see supplementary text S5). The resulting $\delta^{13}\text{C}_{\text{CH}_4}$ in the first 10 cm of sediments is between -59 and -57 ‰, which
632 is consistent with the range of isotopic composition of methane after biogenic methanogenesis (Whiticar et al.,
633 1986).

634 Upward diffusing methane may be either (i) partly lost from the lake’s surface (i.e. escaping the system) by
635 degassing or (ii) totally retained in the water column by complete oxidation (either abiotically by oxygenated
636 surface waters or biologically by methanotrophic organisms). The oxidation of CH_4 in the water column should
637 lead to the formation of ^{13}C -depleted carbon dioxide that would mix back with the lake DIC (and notably with
638 heavy methanogenic CO_2 produced at depth) and/or ^{13}C -depleted biomass (as POC or SOC) if it occurs through
639 methanotrophy. Thus, the net effect of combined methanogenesis and methane oxidation is expected to (i) generate
640 a $\delta^{13}\text{C}_{\text{DIC}}$ gradient from high to low values between the sediment porewaters and the oxycline as proposed
641 elsewhere (Assayag et al., 2008; Wittkop et al., 2014) and (ii) progressively lower sedimentary $\delta^{13}\text{C}_{\text{SOC}}$ in the case
642 of methanotrophy. Abiotic oxidation of methane by dioxygen is consistent with the observation that $\delta^{13}\text{C}_{\text{DIC}}$
643 decreases from porewaters ($\sim +10$ ‰) to the oxycline (-4 ‰), reaching minimum values where dissolved- O_2 starts
644 to appear (Fig. 2). Microbial anaerobic oxidation of methane (AOM) could occur at the 17 m depth through Mn-
645 oxide reduction (Cai et al., 2021; Cheng et al., 2021) and possibly bacterial sulfate-reduction closer to the water-
646 sediment interface, as inferred for the surficial sediments of meromictic Lake Cadagno (Posth et al., 2017). Indeed,
647 we observe a net increase of particulate Fe and S concentrations at a depth of 25 m and a peak of solid sulfide
648 minerals in the surficial sediments (Fig. S5). However, $\delta^{13}\text{C}_{\text{SOC}}$ and $\delta^{13}\text{C}_{\text{POC}}$ are far from calculated $\delta^{13}\text{C}_{\text{CH}_4}$,

649 suggesting that AOM is not a major process in the bottom lake waters and surface sediments (Lehmann et al.,
650 2004) and thus that methanotrophy is not the main CH₄ oxidation pathway in Lake La Alberca.

651 Alternatively, if some portion of the methane escaped oxidation and degassed out of the lake, $\delta^{13}\text{C}_{\text{DIC}}$ would likely
652 be driven to extreme positive values with time (Gu et al., 2004; Hassan, 2014; Birgel et al., 2015; Cadeau et al.,
653 2020). Methane escape is not consistent with the average $\delta^{13}\text{C}_{\text{DIC}}$ in La Alberca (~ -3 ‰; Fig. 4), unless an
654 additional counterbalancing source of DIC to this lake exists. This source of DIC could be volcanic CO₂-degassing
655 (see section 5.1.1). Such a contribution may maintain the lake's average $\delta^{13}\text{C}_{\text{total}}$ close to a mantle isotopic signature
656 and notably away from extreme positive values if CH₄-escape dominated. It is also possible that volcanic CO₂
657 degassing is coupled to methanogenesis by CO₂ reduction in addition to the acetoclastic type described above.

658 Although volcanic CO₂ could be an important source in the C mass balance of Lake La Alberca, we note that it
659 cannot explain the very positive $\delta^{13}\text{C}_{\text{DIC}}$ in the sediment porewaters alone, thus bolstering the identification of
660 methanogenesis. Importantly, this methane cycle is cryptic to the sediment record, as it is evidenced in the
661 dissolved inorganic C phase, but not in the sedimentary organic matter or carbonates. This is a consequence of the
662 lake's stratified nature, where the location of carbonate precipitation and methane production is decoupled.

663

664 *Transfer of OM from the water column to the surficial sediments*

665 The OC content in the first 12 cm of the sediment cores from the four lakes ranges from 1 to 13 wt. % (Table S3).
666 These concentrations are relatively elevated considering the predominantly autochthonous nature of OC and the
667 oligotrophic conditions in these lakes (Alcocer et al., 2014; Havas et al., submitted). In Lake Alchichica, the recent
668 OC burial flux in the sediment was estimated to represent between 15 and 26 g.yr⁻¹.m⁻² (Alcocer et al., 2014).
669 These values are within the range observed for small lakes around the world (Mulholland and Elwood, 1982; Dean
670 and Gorham, 1998; Mendonça et al., 2017), though most of them receive allochthonous OM inputs. Different
671 factors can favor the preservation of OM including lower respiration and oxidation rates due to anoxic bottom
672 waters and scarce benthic biota and/or high sedimentation rates (Alcocer et al., 2014). Anaerobic respiration clearly
673 occurs in the four lakes to some extent, as detailed for La Alberca, and as seen in the surficial sediment data of the
674 other lakes as well (decreasing $\delta^{13}\text{C}_{\text{DIC}}$ in Alchichica, increasing C:N ratio in Atexcac and La Preciosa; Table S3).
675 Nonetheless, the anoxic conditions prevailing in the hypolimnion most of the year are significantly more favorable
676 to OM preservation than oxic conditions (Sobek et al., 2009; Kuntz et al., 2015). While the yearly mixing oxidizes
677 most of the water column during the winter, it also generates a bloom of diatoms which fosters OM production
678 (through shuttling up of bio essential nutrient such as N and Si) and development of anoxia (e.g. Adame et al.,
679 2008). In Alchichica, the large size of some of the phytoplankton was also suggested to favor OM preservation
680 (Adame et al., 2008; Ardiles et al., 2011). Because bacterial sulfate reduction (BSR) is a major remineralization
681 pathway in SO₄-rich environments (e.g. Jørgensen, 1982), the low sulfate content in La Alberca probably favors
682 the preservation of high TOC in the sediments. Even though, appreciable BSR rates may occur in this lake (see
683 discussion above and Fig. S5), similarly to other sulfate-poor environments due to rapid S-cycling (e.g. Vuillemin
684 et al., 2016; Friese et al., 2021). Again, a complete mass-balance of these lakes C fluxes will be required to estimate
685 their net C emission or sequestration behavior.

686 Although the nature and geochemical signatures of the OM that deposits in the sediments may vary throughout the
687 year, it is interesting to infer from what part(s) of the water column surficial sedimentary OM comes during the
688 stratified seasons. In the three lakes from the SOB, $\delta^{13}\text{C}_{\text{SOC}}$ and $(\text{C:N})_{\text{SOM}}$ signatures of the surficial sedimentary
689 OM lie somewhere between POM signatures from the upper water column and from the hypolimnion (Figs. 3, 4).
690 More precisely, in Alchichica, the most surficial $\delta^{13}\text{C}_{\text{SOC}}$ and $(\text{C:N})_{\text{SOM}}$ signatures (-25.7 ‰ and 10.4, respectively)
691 are much closer to values recorded in the upper water column (~ -26.5 ‰ and 10.5, respectively), implying that
692 the upper oxygenic photosynthesis production is primarily recorded. It is consistent with previous studies
693 suggesting that most of the phytoplankton biomass being exported is composed of diatoms (Ardiles et al., 2011).
694 In Lake Atexcac, however, $\delta^{13}\text{C}_{\text{SOC}}$ and $(\text{C:N})_{\text{SOM}}$ signatures (~ -26.8 ‰ and 8, respectively) are closer to values
695 recorded in the hypolimnion (~ -26.5 ‰ and 6.5, respectively) suggesting that SOM records mostly the anaerobic
696 primary production.

697 In La Alberca, surficial $\delta^{13}\text{C}_{\text{SOC}}$ is markedly more negative (by ~ 2 to 3 ‰) than the deepest and shallowest water
698 column values (Fig. 4), but close to what is recorded at the redoxcline depth of 17 m. However, the $(\text{C:N})_{\text{SOM}}$
699 values are much higher than what is measured anywhere in the water column, which is consistent with OM
700 remineralization by sulfate-reduction and methanogenesis in the sediments of this lake. Therefore, OM
701 biogeochemical signatures in the surficial sediments of La Alberca could be strongly influenced by early diagenesis
702 occurring at the water-sediment interface – despite favorable conditions for OM preservation.

703 In summary, the OM depositing at the bottom of these stratified lakes does not always record geochemical
704 signatures from the same layers of the water columns and can be modified by very early diagenesis. It does not
705 necessarily record the signatures of primary production by oxygenic photosynthesis from the upper column. For
706 example, in Lake Atexcac, sedimentary OM records primary production by anoxygenic photosynthesis, even
707 though POC concentration is highest in the upper water column. This result highlights the diversity of geochemical
708 signatures that can stem from continental environments despite their geographical, geological, and climatic
709 proximity. A deeper understanding of the OM transfer process from water column to sediment will require more
710 detailed analyses and comparison of the different OM pigments and molecules and could have strong implications
711 for the interpretation of the fossil record in deep anoxic time.

712

713 6. CONCLUSIONS AND SUMMARY

714 The carbon cycles of four stratified alkaline crater lakes were described and compared based on the concentration
715 and isotopic compositions of DIC and POC in the water columns and surficial (~10 cm) sedimentary carbonates
716 and organic carbon. Overall our study shows the wide diversity of geochemical signatures found in continental
717 stratified environments despite similar geological and climatic contexts. We identify different regimes of C cycling
718 in the four lakes due to different biogeochemical reactions related to slight environmental and ecological
719 variations. In more detail, we show that:

720 - External abiotic factors, such as the hydrological regime and the inorganic C sources in the lakes, control
721 their alkalinity and thus, the buffering capacity of their waters. In turn, these differences in buffering
722 capacity constrain variations in pH along the stratified water columns as well as the inorganic C isotope
723 signatures recorded in the water columns and sediments of the lakes. The $\delta^{13}\text{C}_{\text{carb}}$ reflects the abiotic

724 factors generating the alkalinity gradient, but it is poorly representative of biological processes in lakes
725 with high alkalinity. The external environmental factors further impact the C mass balance of the lakes
726 with probable consequences on their net C-emitting or -sequestering status.

- 727 - Based on POC and DIC concentrations and isotopic compositions, combined with physico-chemical
728 parameters, we are able to identify the activity of oxygenic photosynthesis and aerobic respiration in the
729 four lakes studied. Anoxygenic photosynthesis and/or chemoautotrophy are also evidenced in two of the
730 lakes, but their POC and DIC signatures can be equivocal.
- 731 - Methanogenesis is evidenced in the surficial sediments of the OM-rich Lake La Alberca de los Espinos
732 and influences the geochemical signatures lower in the water column. However, it is recorded only in
733 analyses of porewater dissolved species, but not imprinted in the sedimentary archives (OM and
734 carbonates).
- 735 - The SOM geochemical signatures of these stratified lakes do not all record the same “biogeochemical
736 layers” of the water column (e.g. anaerobic vs. aerobic metabolisms), and, in some cases, can be greatly
737 modified by early diagenesis.

738

739 **Author Contributions**

740 RH and CT designed the study in a project directed by PLG, KB and CT. CT, MI, DJ, DM, RT, PLG and KB
741 collected the samples on the field. RH carried out the measurements for C data; DJ the physico-chemical parameter
742 probe measurements and EM provided data for trace and major elements. RH and CT analyzed the data. RH wrote
743 the manuscript with important contributions of all co-authors.

744

745 **Competing Interests**

746 The authors declare that they have no conflict of interest.

747

748 **Disclaimer**

749

750 **Acknowledgements**

751 This work was supported by Agence Nationale de la Recherche (France; ANR Microbialites, grant number ANR-
752 18-CE02-0013-02). The authors thank Anne-Lise Santoni, Elodie Cognard, Théophile Cocquerez and the GISMO
753 platform (Biogéosciences, Université Bourgogne Franche-Comté, UMR CNRS 6282, France). We thank Céline
754 Liorzou and Bleuenn Guéguen for the analyses at the Pôle Spectrométrie Océan (Laboratoire Géo-Océan, Brest,
755 France) and Laure Cordier for ion chromatography analyses at IPGP (France). We thank Nelly Assayag and Pierre
756 Cadeau for their help on the AP 2003 at IPGP.

757

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