

1 **Use of water isotopes and chemistry to infer the type and degree of**
2 **exchange between groundwater and lakes in an esker complex of**
3 **northeastern Ontario, Canada**

4
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12
13 **Abstract**

14
15 While interactions between groundwater and lake-water influence water chemistry, water balance,
16 aquatic organisms, biochemical cycles and contamination levels, they remain a poorly-studied component
17 of lake hydrology. Identifying the controls of groundwater-lake water interactions at the landscape level
18 and classifying lakes into different categories based on their degree of interaction with the groundwater can
19 provide insights on a lakes' sensitivity and vulnerability to environmental stressors, and provide baseline
20 conditions for comparison to future changes that are important for water management and conservation. To
21 this end, water chemistry and water isotopic composition were investigated in a set of 50 boreal lakes
22 located at different elevations in an esker system near Timmins, Ontario. Analyses focused on stable
23 isotopic ratios of hydrogen and oxygen, as well as specific conductance as indicators of the position of a
24 lake with respect to the influence of groundwater. Both isotopic composition and specific conductance
25 distinguished higher-elevation groundwater-recharge lakes from lower-elevation groundwater-discharge
26 lakes. Groundwater-recharge lakes, characterized by enriched isotopic values and low values of specific
27 conductance, were located above the hydraulic midline elevation of the study lakes. In contrast,
28 groundwater-discharge lakes were isotopically depleted and had higher values of specific conductance, and
29 occurred below the hydraulic midline of the study lakes. An intermediate group of lakes was also defined
30 (termed seepage lakes) and had intermediate isotopic and water chemistry characteristics compared to
31 recharge and discharge lakes. Differences in water geochemistry between field campaigns revealed that
32 upland groundwater-recharge lakes showed evidence of evaporative drawdown, indicating sensitivity to
33 short-term changes in climate, whereas the lower-elevation groundwater-discharge lakes showed little

34 variation between seasonal samples, and consequently would likely be affected only by hydroclimatological
35 changes of greater duration and **magnitude**.

36

37 **Keywords**

38

39 Water balance, landscape limnology, groundwater, hydrochemistry, lake-water chemistry, stable isotopes

40

41 **1 Introduction**

42

43 Nearly all surface waters, including lakes, interact with groundwater to some extent (Winter *et al.*,
44 1998). The degree to which lakes interact with groundwater strongly influences lake-water chemistry and
45 water balance as well as aquatic biota and biochemical cycles (Rosenberry *et al.*, 2015; Lewandowski *et al.*,
46 2015). These characteristics will influence the **sensitivity of** lakes to environmental stressors such as
47 climate change, and other anthropogenic disturbances. Future climate change will likely affect lakes
48 differently depending on their connectivity to groundwater, further increasing our need to better understand
49 the relations between lake hydrology and lake-water geochemistry. Further, understanding the climatic
50 controls on water **balance is** essential for informed ecosystem management and conservation practices
51 (Winter *et al.*, 1998).

52

53 Groundwater-lake water interactions are an often poorly-studied component of lake hydrology.
54 However, recent advances in isotopic techniques and modelling approaches have enabled researchers the
55 opportunity to better understand hydrological processes in lakes at a local to regional scale (Fleckenstein *et al.*,
56 2010; Rosenberry *et al.*, 2015). This has included studies using numerical simulations (e.g. Winter *et al.*,
57 1998), as well as empirical studies at a range of different spatial and temporal scales. Studies at the
58 regional scale (e.g. Gorham *et al.*, 1983; Webster *et al.*, 2000) emphasize differences in climate and surface
59 geology as being the important drivers of changes in lake-water chemistry. By contrast, empirical studies
60 performed at a local to regional scale (e.g. Webster *et al.*, 1996; Quinlan *et al.*, 2003; Martin and Soranno,
61 2006; Ala-aho *et al.*, 2013) indicate the importance of landscape position and also lake morphology as being
62 important in understanding lake-groundwater interactions. At the scale of an individual lake, a number of
63 studies have shown that groundwater-lake water interactions can vary temporally according to changes in
64 seasonality and longer-term changes in hydroclimatic conditions (e.g. Kenoyer and Anderson, 1989;
65 LaBaugh *et al.*, 1997; Sebestyen and Schneider, 2001; Schuster *et al.*, 2003; Arnoux *et al.*, 2017b).

66

67 The vast majority of past empirical studies have relied on studying aspects of water chemistry (e.g.
68 dissolved ions), as an indicator of hydrological connectivity between lakes, **because** such approaches can
69 **reveal** signals of important processes such as mineral weathering and dissolution (Bertrand *et al.*, 2014).
70 Chemical tracers are often referred to as non-conservative tracers because their composition changes as
71 they react with catchment materials (Kendall and McDonnell, 1998). By contrast, other studies relied on
72 the isotopic composition of water as a hydrological tracer (e.g. Turner *et al.*, 2010; Isokangas *et al.*, 2015).
73 The latter are labelled to as conservative tracers because they are relatively conservative in reactions with
74 catchment materials and retain their distinctive values until they mix with other water sources or they
75 evaporate (Kendall and McDonnell, 1998). In addition to being a good indicator of source water, stable
76 isotopes of water constitute an efficient and cost-effective means to quantify lake evaporation and water
77 balance status of water bodies using the Craig-Gordon model (Craig and Gordon, 1965), given that the
78 isotopic composition of lake-water and precipitation as well as air temperature and relative humidity are
79 known, or can be estimated (Gibson and Edwards, 2002). The Craig-Gordon model is well established and
80 has been used extensively to investigate the spatial and temporal variability in lake-water balance in remote
81 boreal regions of Canada (e.g. Wolfe *et al.*, 2007; Bouchard *et al.*, 2013; Tondu *et al.*, 2013; Turner *et al.*,
82 2014a; Turner *et al.*, 2014b). However, precipitation and groundwater often display similar isotopic
83 signatures as they both tend to retain their original isotopic composition because they undergo little to no
84 evaporation (Gibson and Edwards, 2002; Gibson *et al.*, 2008; Yi *et al.*, 2008). This makes the distinction
85 of the relative influence of groundwater and precipitation in lake-water balance challenging. Given this, the
86 combination of chemical and isotopic approaches has the ability to produce more reliable interpretations,
87 especially if the two approaches converge on a mutually reinforcing interpretation. Chemical and isotopic
88 tracers have been widely used together to investigate the connectivity between groundwater and lake water
89 within a single lake (e.g. Labaugh *et al.*, 1997; Schuster *et al.*, 2003; Rautio and Korkka-Niemi, 2011) or
90 for a cluster of a few selected lakes (e.g. Arnoux *et al.*, 2017a; Arnoux *et al.*, 2017b; Gurrieri and Furniss,
91 2004; Katz *et al.*, 1997; Turner and Townley, 2006). Nonetheless, studies that have combined chemical and
92 isotopic approaches to investigate the connectivity between groundwater and lake water **at the landscape**
93 **level and for a large number of lakes in a region** are lacking.

94

95 The main objective of this study is to examine the importance of landscape position on groundwater
96 connectivity by scrutinising both water chemistry and isotopic composition of water in a boreal esker
97 complex in northeastern Ontario. **The use of water tracers was preferred to direct measurements because**
98 **tracers (i) have proven to be good indicators of groundwater-lake water interactions and (ii) constitute a**
99 **time and cost-effective approach that can be applied at a greater spatial scale.** Investigating such interactions
100 in the context of esker hydrology is particularly relevant as eskers consist of porous and permeable materials

101 that facilitate groundwater flows, are widespread in boreal regions (Ala-aho et al., 2015) and constitute one
102 of the most common type of aquifers in boreal regions of Canada (Cloutier et al., 2007). Results from this
103 study will be used to develop a lake typology (i.e. a classification or generalisation of lakes into different
104 categories) of hydrological connectivity (Newton and Driscoll, 1990; Bertrand et al., 2014). The resultant
105 typology will provide insights on lakes' sensitivity to environmental stressors over time such as climate
106 change, acidification or pollution by accounting for variation among lake types through continuous
107 monitoring. This classification will also improve and simplify water management and conservation goals
108 as each lake type requires the same management strategies in a region where cottage development,
109 recreational fishing, mining activities and aggregate extraction is prevalent (Cochrane, 2006). Additionally,
110 the typology will provide an important baseline for comparison to future hydrological regimes that may
111 altered them and be used for site-selection and interpretations of past hydrological changes from
112 stratigraphic analysis of isotopic and geochemical indicators in sediment cores from lakes. Finally, the
113 sensitivity of the proposed typology will be assessed by investigating if short-term variations of lake-water
114 characteristics are more readily detected in higher-elevation groundwater-recharge lakes (i.e. lakes that
115 receive the majority of their water from precipitation and feed the groundwater system) in comparison to
116 lower-elevation groundwater-discharge lakes (i.e. lakes that receive the majority of their water from
117 groundwater).

118

119 **2 Study area**

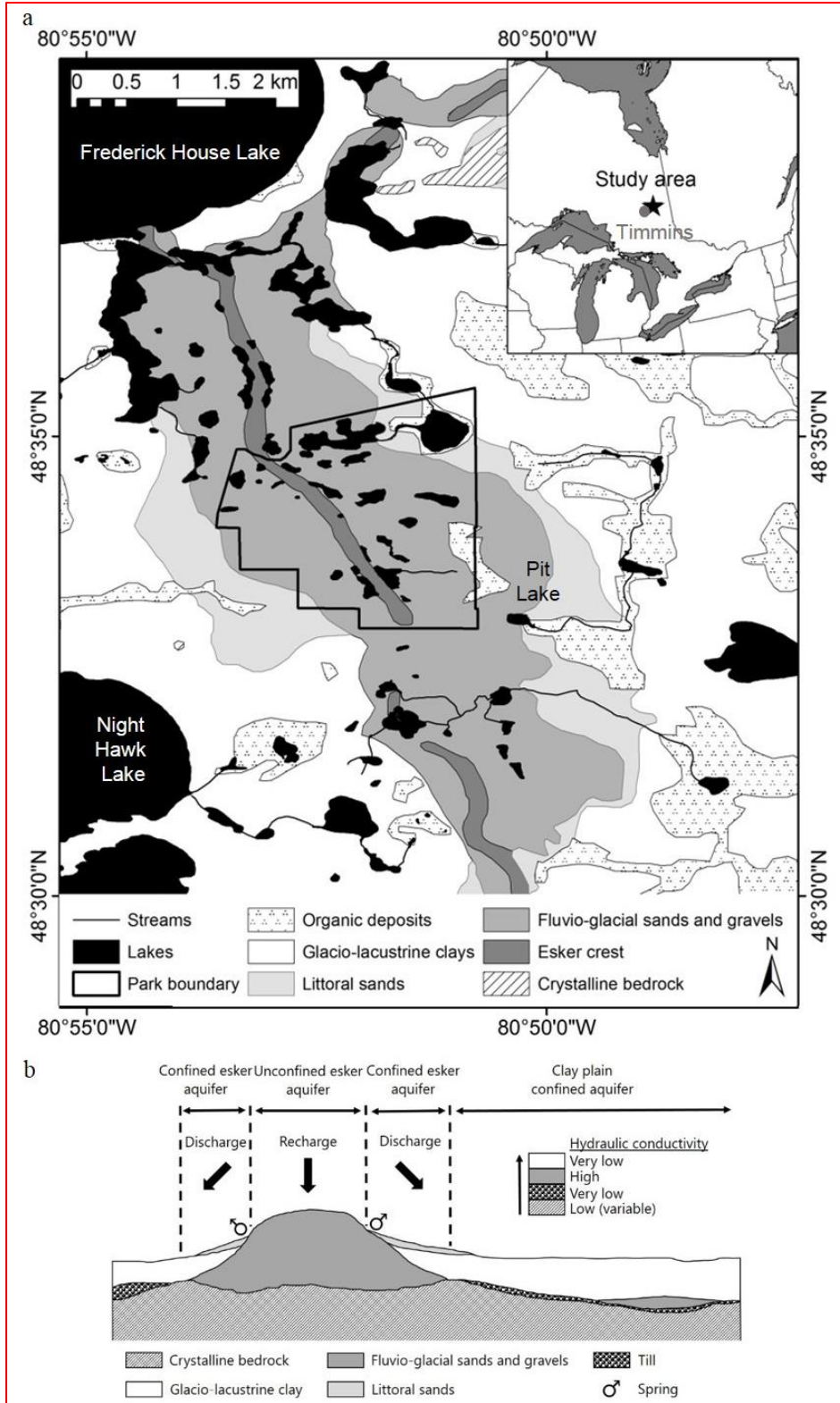
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121 The study area is a portion of the Kettle Lake Esker between the southern shore of Frederick House
122 Lake, and the northern shore of Night Hawk Lake, a region located approximately 35-km east of Timmins,
123 Ontario, Canada (Fig. 1a). The Timmins region is characterized by a humid continental climate (dfb in the
124 Köppen climate classification) with a mean annual temperature of 1.8°C and average precipitation of 835
125 mm (Environment Canada, 2015). This region has long cold winters and lakes are covered with ice from
126 early November until early April. Summers are usually wet and mean air temperatures are 17.5°C in July.
127 Many of the study lakes and streams are located within Kettle Lakes Provincial Park, as well as in lower
128 elevation regions that have been moderately influenced by human activities (Cochrane, 2006). The study
129 region is covered with well-drained orthic humo-ferric podzols, while surrounding clay plains are for the
130 most part covered with moderately to poorly drained orthic gray luvisols and gleyed gray luvisols
131 (OMNDM, 2006). Jack pine, poplar, black spruce, white birch, trembling aspen and balsam fir dominate
132 the well-drained areas, with a dominance of spruce in poorly drained regions.

133

134 The regional landscape is dominated by landforms and deposits of the Laurentide Ice Sheet during
135 the last glacial maximum and deglaciation of the region was essentially complete approximately 10,000
136 years ago (Dyke, 2004). The ablation of the ice sheet was particularly dynamic and led to the formation of
137 relatively large eskers composed of long sinuous ridges of coarse grained glaciofluvial sediments in
138 deposits oriented in a north-south direction and mantling the crystalline bedrock (Cloutier et al., 2007). The
139 retreat of the ice sheet was accompanied by ponding of glacial meltwaters that led to the development of
140 glacial Lake Ojibway that submerged most of the region (Roy et al., 2011) and the widespread deposition
141 of glaciolacustrine clay (Nadeau, 2011), followed by the drainage of Lake Ojibway into Hudson Bay ~8,200
142 years ago (Roy et al., 2011). As Lake Ojibway levels dropped, wave action eroded the surface of the esker
143 and redistributed some sand materials on the flanks on the esker, forming lateral littoral sand units that
144 drape the glaciolacustrine clays (Cloutier et al., 2007) (**Fig. 1b**). The numerous kettle lakes on the esker
145 formed as glacial ice was trapped in the outwash materials melted (Nadeau, 2011).

146
147 The esker stratigraphy ensures that its groundwater system is highly localized because the esker
148 generally have a high hydraulic conductivity due to their coarse texture. The esker is surrounded by
149 underlying bedrock and the adjacent fine-grained glaciolacustrine deposits, both characterized by very low
150 hydrological conductivity (Stauffer and Wittchen, 1992). As a consequence, the esker can be conceptually
151 partitioned into zones of unconfined aquifers in its center where coarse material is present at the surface
152 and zones of confined aquifers at its edges when fine-grained sediment mantles the core of the esker (**Fig.**
153 **1b**) (Cloutier et al., 2007). Thus, the recharge of the esker will occur through infiltration of precipitation in
154 the unconfined aquifer and discharge will take place on the esker flanks at the contact of the clay, where
155 most groundwater springs emerge (Cloutier et al., 2007). Confined aquifers found on the surrounding clay
156 plain are often covered by peatlands and shallow lakes fed by groundwater springs on the edges of the esker
157 or by streams that drain the esker (Rossi et al., 2012).



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Fig. 1: Location of the study area in north east Ontario near Timmins, Ontario (Based on: Richard and McClenaghan, 2000) (a) and conceptual geological transverse section of an esker complex at latitude 48°35'0"N (b) (Modified from: Veillette et al., 2004)

163 **3 Methods**

164

165 *3.1 Geomatic and lake morphometric data*

166

167 Lakes and other geographic features were digitized from Google Earth using the imagery dating from
168 7/26/2005. Maximum lake depths Z_{max} were obtained from the Ontario Ministry of Natural Resources and
169 Forestry (OMNRF, 2011). Digital Elevation Models (DEM) from the study area were obtained from Natural
170 Resources Canada with a spatial resolution of 20 m (NRC, 2013). Lake elevation values were calculated as
171 the mean elevation of the lake in ArcGIS 10.3 from the available DEM, and lake watershed slopes were
172 calculated from the DEM in ArcGIS in a 100 m buffer zone surrounding each lake. This approach was used
173 because of the coarse spatial resolution of the DEM and the close proximity of the lakes made clear
174 individual lake watershed delineation impossible. Because all lakes in the study area are kettle lakes, which
175 are characterized by steep shore slopes, initial buffer zones of different widths were produced. The buffer
176 width of 100 m was chosen as this distance showed the best correlation with water tracers.

177

178 *3.2 Field measurements and water sample collection*

179

180 Water samples were collected during three field campaigns (7-14 June 2013, 16-23 June 2014, and
181 12-19 August 2014). 50 lakes were sampled (29, 28 and 50 lakes during the June 2013, June 2014 and
182 August 2014 campaign respectively), as well as a number of streams (lake outlets and lake inlets) and
183 groundwater springs. The lakes were chosen to achieve a wide spatial coverage over this region
184 encompassing a gradient of lake elevation as well as lake types. A few lakes located in the area of interest
185 could not be sampled due to poor accessibility. Groundwater springs and streams flowing into the study
186 lakes were also sampled to characterize the local groundwater-surface water interactions.

187

188 Temperature and specific conductance of the water (corrected to 25°C) at each site were measured
189 with an YSI Salinity Conductivity Temperature meter (accuracy $\pm 0.1 \mu\text{S}/\text{cm}$ and $\pm 0.1^\circ\text{C}$). The pH was
190 measured with an UP-5 Denver Instrument pH meter calibrated before usage (accuracy ± 0.1). Water
191 samples were collected using 1 L Nalgene bottles that were tripled-rinse with distilled water prior to use
192 and again with sample water while sampling (Louiseize et al., 2014). To prevent cross-contamination, the
193 tripled-rinsing of bottles with sampling water was carried away from the final sampling point in the lake or
194 downstream of stream/spring sampling points. Bottles were completely filled in order to avoid headspace
195 air and subsequently filtered within 3 hours of sampling.

196 Samples collected for water stable isotope, dissolved ion analysis and dissolved inorganic nitrogen
197 were vacuum filtered with 0.45 μm Millipore Isopore filters. Vacuum filtering apparatus were triple-rinse
198 with distilled water and the lake water before and after use for each sample. The filtrates were then poured
199 and placed into pre-cleaned 20ml plastic scintillation vials. Three replicates were collected for each sample.
200 All vials were previously rinsed with filtered sample water that was discarded. Vials were completely filled
201 to avoid any headspace, sealed with Parafilm™ to avoid evaporation, and kept in a cool and dark fridge
202 prior to analysis.

203
204 Samples for total dissolved nitrogen (TDN), dissolved organic carbon (DOC) and dissolved inorganic
205 carbon (DIC) analysis were filtered on-site using a glass filtration and Whatman GF/F glass fiber filters to
206 avoid any *ex situ* carbon addition. Filters were pre-ashed at 550°C for two hours and wrapped with pre-
207 ashed aluminum foil prior to utilization to eliminate any residual organic matter (Lamoureux and
208 Lafrenière, 2014). The glass filtration apparatus was triple-rinsed with distilled water and lake water before
209 and after use for each sample and was wrapped with new clean pre-ashed aluminum foil overnight. Filtrates
210 were then poured and stored into pre-cleaned 45-ml amber EPA vials with Teflon-lined septa with no
211 headspace (Louiseize et al., 2014). Two replicates were collected for each sample. Vials were labelled and
212 were kept cool and in the dark in a fridge prior to analysis.

213 214 3.3 Laboratory analysis

215
216 The stable isotope ratios of water ($\delta^{18}\text{O}$ and $\delta^2\text{H}$) were measured using a Los Gatos Research Liquid–
217 Water Isotope Analyzer (LGR) in the FaBRECC laboratory at Queen’s University, which vaporizes injected
218 sample and measures its absorbance relative to Vienna Standard Mean Ocean (‰ V-SMOW). All runs
219 contain 6 replicate analyses and 3 standards produced by Los Gatos bracketing every 3 samples (e.g.
220 LGR1A, $\delta^{18}\text{O}=-19.50\text{‰}$ and $\delta^2\text{H}=-154.3\text{‰}$; LGR2A, $\delta^{18}\text{O}=-16.14\text{‰}$ and $\delta^2\text{H}=-123.6\text{‰}$; LGR3A, $\delta^{18}\text{O}=-$
221 13.10‰ and $\delta^2\text{H}=-96.4\text{‰}$; LGR4A, $\delta^{18}\text{O}=-7.69\text{‰}$ and $\delta^2\text{H}=-51.0\text{‰}$; LGR5A, $\delta^{18}\text{O}=-2.80\text{‰}$ and $\delta^2\text{H}=-$
222 9.5‰). Sample reproducibility (1σ) was based on repeated measurements of samples and standards fixed
223 at 0.25‰ for $\delta^{18}\text{O}$ and at 1.5‰ for $\delta^2\text{H}$.

224
225 Concentrations of inorganic ions (Ca, Mg, K, Na, Cl, SO_4) were measured by liquid ion
226 chromatography with a Dionex ICS-3000. Detection limits were as follows: 0.5 ppm for Ca, 0.01 ppm for
227 Mg, 0.2 ppm for K, 0.3 ppm for Na, 0.05 ppm for Cl and 0.1 ppm for SO_4). Concentrations of dissolved
228 inorganic species (NO_2 - NO_3 and NH_4) were measured by colorimetry using an Astoria Pacific FASpac II
229 Flow Analyser (detection limits of 0.01 ppm). Concentrations of DOC, DIC and TDN were measured by

230 high-temperature combustion and nondispersive infrared sensor and chemiluminescent detection using a
 231 Shimadzu TOC-VPCH/TNM equipped with a high-sensitivity catalyst system (detection limits of 0.08 ppm
 232 for DOC and 0.015 ppm for TN) (Louiseize et al., 2014). Total dissolved carbon (TDC) was calculated as
 233 the sum of DOC and DIC.

234

235 3.4 Water balance calculations

236

237 Monthly precipitation isotopic data are available from February, 1997 to November, 2010.
 238 Precipitation isotopic data were collected at Bonner Lake, about 125-km NW of the study area by the
 239 Canadian Network for Isotopes in Precipitation (CNIP) (Birks et al., 2010). The general water (Eq. 1) and
 240 isotope balance (Eq. 2) of a well-mixed lake may be written respectively as follow (Darling et al., 2005):

241

$$\frac{dV}{dt} = I - Q - E \quad \text{Eq. 1}$$

242

$$\frac{Vd\delta_L + \delta_L dV}{dt} = I\delta_I - Q\delta_Q - E\delta_E \quad \text{Eq. 2}$$

243

244 where V is the volume of the lake, t is time, dV is the change of volume over time dt , I is instantaneous
 245 inflow where $I = I_F + I_G + P$ (I_F being surface inflow, I_G groundwater inflow and P precipitation on the
 246 lake surface); Q is instantaneous outflow where $Q = Q_R + Q_G$ (Q_R is surface outflow and Q_G is groundwater
 247 outflow), E is evaporation; and δ_L , δ_I , δ_Q and δ_E are the isotopic compositions of the lake, inflow, outflow
 248 and evaporative flux respectively. Assuming (i) that the lake maintains a near-constant volume on the long-
 249 term (*i.e.*, $dV = 0$ and $dt \rightarrow \infty$) (Darling et al., 2005) and (ii) that physical outflow does not cause isotopic
 250 fractionation (*i.e.*, $\delta_Q = \delta_L$) (Gibson and Edwards, 2002; Yi et al., 2008), Eq. 1 and 2 can be simplified and
 251 rewritten as follows:

252

$$I = Q + E \quad \text{Eq. 3}$$

253

$$I\delta_I = Q\delta_Q + E\delta_E \quad \text{Eq. 4}$$

254

255 E can be related to I assuming that the lakes are in isotopic steady state (*i.e.* undergoing evaporation while
 256 maintaining constant volume). This assumption seems well justified as most of the lakes have had sufficient
 257 time in the past to reach their isotopic steady-state which is reflective of the local climate and their mean

258 hydrological status, and can be defined by its water balance, which corresponds to the ratio of the total
 259 inflow to the evaporation rate (Isokangas et al., 2015). The evaporation-to-inflow ratio of the lake E_L/I_L can
 260 be calculated by combining Eq. 3 and 4 (Gibson and Edwards, 2002; Yi et al., 2008):

$$\frac{E_L}{I_L} = \frac{\delta_I - \delta_L}{\delta_E - \delta_L} \quad \text{Eq. 5}$$

262 where δ_I was computed as the intersection of the Local Meteoric Water Line (LMWL) with the Local
 263 Evaporation Line (LEL) (Gibson et al., 1993; Yi et al., 2008); δ_L is the isotopic composition of the lake
 264 water sample and δ_E was estimated using the Craig-Gordon model (Craig and Gordon, 1965) formulated
 265 by Gonfiantini (1986) as follows:

$$\delta_E = \frac{(\delta_L - \varepsilon^*)/\alpha^* - h\delta_A - \varepsilon_k}{1 - h + \varepsilon_k} \quad \text{Eq. 6}$$

267 where δ_L is the isotopic composition of lake water, ε^* is the equilibrium isotopic separation term, α^* is the
 268 liquid–vapour equilibrium fractionation factor, h is the relative humidity, δ_A is the isotopic composition of
 269 the local atmospheric moisture, and ε_k is the kinetic separation term between the liquid and vapour phases.
 270 The ε^* and α^* parameters which are temperature dependent can be calculated using empirical equations for
 271 $\delta^{18}\text{O}$ as follows (Horita and Wesolowski, 1994):

$$\varepsilon^* = -7.685 + 6.7123 \left(\frac{10^3}{T} \right) - 1.6664 \left(\frac{10^6}{T^2} \right) + 0.35041 \left(\frac{10^9}{T^3} \right) \quad \text{Eq. 7}$$

$$\alpha^* = \exp \left(-\frac{7.685}{10^3} + \frac{6.7123}{T} - \frac{1666.4}{T^2} + \frac{350410}{T^3} \right) \quad \text{Eq. 8}$$

275 where T is the air temperature in Kelvins. ε_k (Eq. 9) is expressed for $\delta^{18}\text{O}$ by (Gonfiantini, 1986):

$$\varepsilon_k = (0.0142 (1 - h))1000 \quad \text{Eq. 9}$$

278 The equation for δ_E was modified according to Gibson and Edwards (2002) to directly utilize isotopic data
 279 in per mil rather than as a decimal fraction and expressed as follows:

281

$$\delta_E = \frac{\alpha^* \delta_L - h \delta_A - \varepsilon}{1 - h + 10^{-3} \varepsilon_k} \quad \text{Eq. 10}$$

282

283 where ε is the total isotopic separation factor that includes both ε^* and ε_k expressed as:

284

$$\varepsilon = \varepsilon^* + \varepsilon_k \quad \text{Eq. 11}$$

285

286 δ_A was originally estimated with the original model that assumes isotopic equilibrium between atmospheric
287 moisture and precipitation as follows (Gibson, 2002):

288

$$\delta_A = \frac{\delta_p - \varepsilon^*}{1 + 10^{-3} \varepsilon^*} \quad \text{Eq. 12}$$

289

290 where δ_p was computed as the average isotopic composition of annual precipitation from February 1997 to
291 November 2010 (data collected by CNIP). The same procedure was used to calculate the evaporation-to-
292 inflow ratio of streams and groundwater springs.

293

294 However, the hypothesis of steady-state may not be valid for some small lakes that undergo
295 significant changes in lake levels as evaporation progresses. For those lakes in non-steady state (here
296 defined as non-alkaline high-elevation lakes), the evaporative loss fraction of the lake volume (f) was
297 calculated using the original equation of Gonfiantini (1986) rearranged as follows (Skrypek et al., 2015):

298

$$f = 1 - \left[\frac{\delta_L - \delta^*}{\delta_p - \delta^*} \right]^{\frac{1}{m}} \quad \text{Eq. 13}$$

299 where δ_p is the initial value of water in the lake that undergoes evaporation, δ_L the final value of water in
300 the lake that undergoes evaporation, δ^* is the limiting isotope enrichment factor defined as follows (Skrypek
301 et al., 2015):

302

$$\delta^* = \frac{h \delta_A + \varepsilon}{h - \frac{\varepsilon}{1000}} \quad \text{Eq. 14}$$

303 and m is the enrichment slope defined as follows (Skrypek et al., 2015):

304

$$m = \frac{h - \frac{\varepsilon}{1000}}{h + \frac{\varepsilon_k}{1000}} \quad \text{Eq. 14}$$

305

306 3.5 Statistical analysis

307

308 Linear regressions were used to assess the degree of co-variability between quantitative variables
309 while logistic regressions were utilized to assess the relations between binary variables and quantitative
310 variables at the 0.05 level. Breakpoint analysis or segmented regression was used to detect any change of
311 trends in water tracers along an elevation gradient and to produce subsequent higher-level groupings of
312 lakes. Breakpoints that were significant at the 0.05 level were averaged to obtain the elevation of the
313 “breakpoint line”. A non-metric multidimensional scaling (NMDS) was run to assess the differences among
314 lake types in a 2-dimensional ordination space using non-scaled values of electrical conductance, Ca, $\delta^{18}\text{O}$
315 and $\delta^2\text{H}$ as input variables, Euclidean distance as dissimilarity measure, and without rotation. A Wilcoxon
316 signed-rank test was subsequently applied as a post-hoc analysis for all lake-water variables that were above
317 detection limits to determine if differences among the different types of lakes were statistically significant
318 at the 0.05 level as most of the Shapiro-Wilk test for normality revealed that most variables were not
319 normally distributed. An analysis of similarity (ANOSIM) was also carried out as a complement to
320 determine if within group similarity was significantly greater than in-between group similarity at the 0.05
321 level. All statistical analyses were performed in R 3.4 on the data from the August 2014 campaign as it was
322 the one with the most samples.

323

324 **4 Results**

325

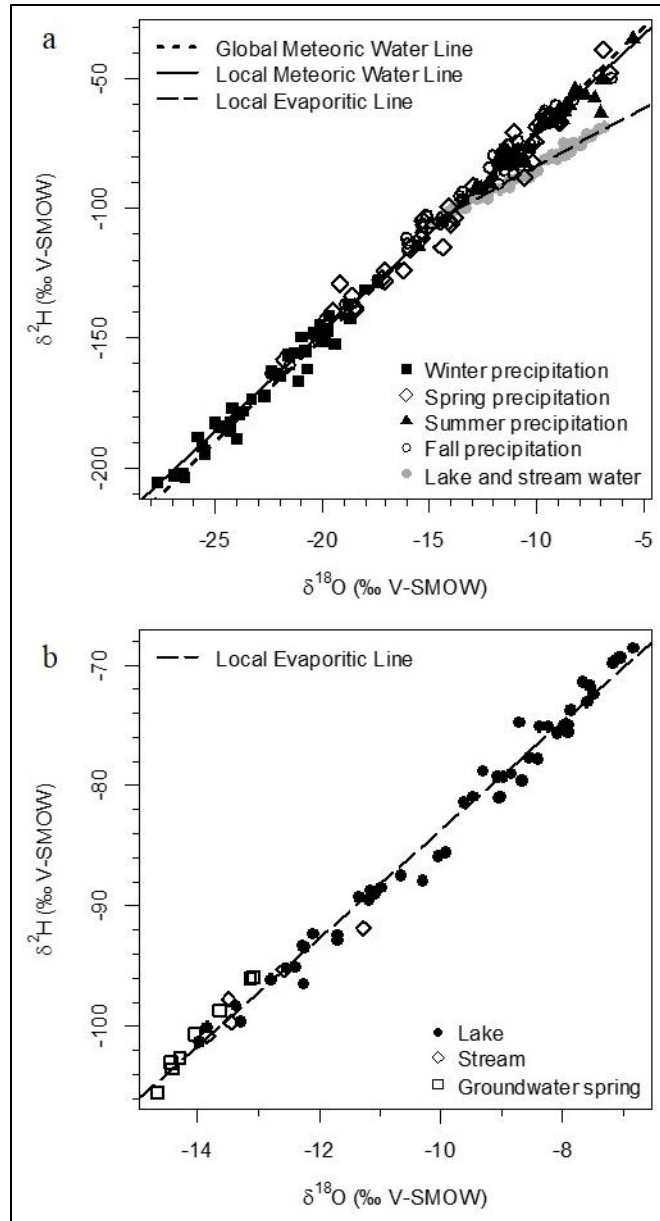
326 4.1 Temperature and water stable isotopes

327

328 During the August 2014 field campaign, all types of bodies of water differed from one another by
329 their temperature: springs are characterized by low temperatures (6 -12°C) whereas streams have slightly
330 higher temperature (12 -16°C) and lakes ranged from (16 -18°C). It should be noted however that spring
331 temperature is not an accurate indication of groundwater temperature due to alterations induced by the
332 velocity of the discharging spring and the resulting warming of the discharging water as it reaches the
333 surface.

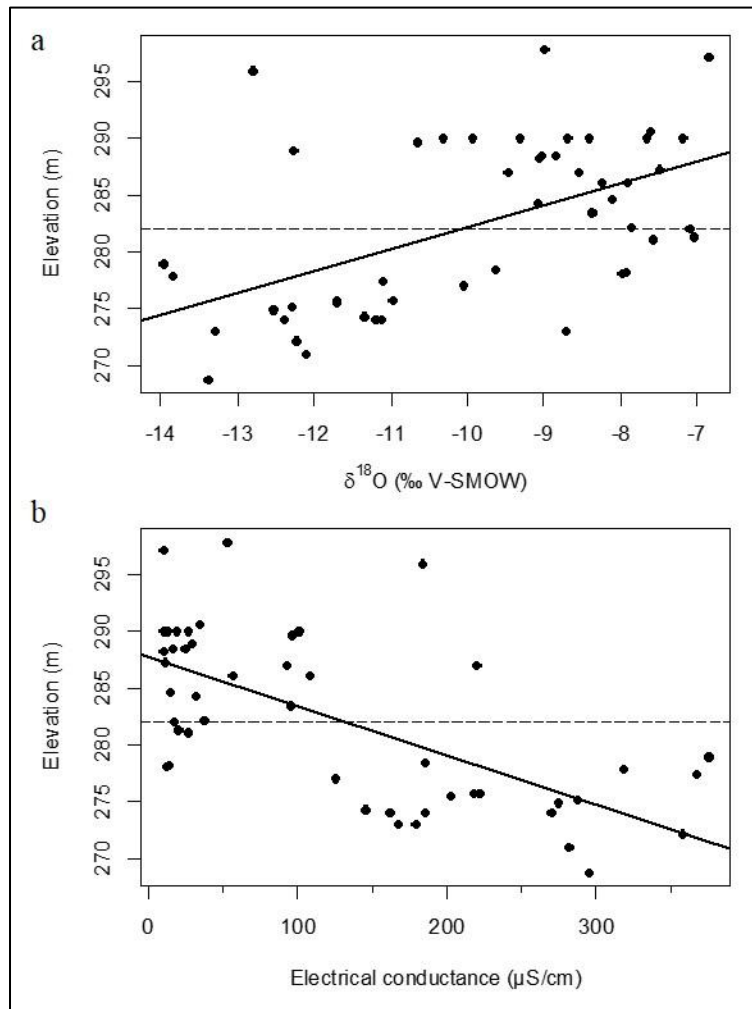
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335 Monthly precipitation isotopic data from CNIP from February, 1997 to November, 2010 from Bonner
336 Lake, about 125-km NW of the study area, show progressive enrichment in values between winter, spring,
337 fall and summer on the global and local meteoric water lines, which are similar (**Fig. 2a**). The water samples
338 (*i.e.* groundwater springs, streams and lakes) displayed a wide range of isotope values (-14.7‰ to -6.8‰
339 for $\delta^{18}\text{O}$ and -105.5‰ to -68.6‰ for $\delta^2\text{H}$), which fell on a Local Evaporation Line ($r = 0.99$, $p < 0.001$)
340 (**Fig. 2b**). Groundwater springs have isotopic values similar to mean annual precipitation (-14.7‰ to -
341 13.1‰ for $\delta^{18}\text{O}$ and 105.5‰ to -96.0‰ for $\delta^2\text{H}$) while being more depleted as summer precipitation is more
342 enriched in heavy isotopes than groundwater (**Fig 2a**). Streams have comparable isotopic composition to
343 groundwater springs (-13.8‰ to -11.3‰ for $\delta^{18}\text{O}$, and -100.8‰ to -91.9‰ for $\delta^2\text{H}$). By contrast, lakes are
344 characterized by large variations in water isotopic composition, ranging from values comparable to
345 groundwater springs (*ca.* -14‰ for $\delta^{18}\text{O}$ and -100‰ for $\delta^2\text{H}$) to more enriched values (*ca.* -6‰ for $\delta^{18}\text{O}$
346 and -70‰ for $\delta^2\text{H}$).
347



348 **Fig. 2:** Isotopic composition of precipitation in relation to the Global Meteoric Water and Local Meteoric
 349 Water lines ($r = 0.99$, $n = 166$, $p < 0.001$) (a), and isotopic composition of collected water samples in
 350 relation to the Local **Evaporation** Line ($r = 0.99$, $n = 68$, $p < 0.001$) (b).
 351
 352

353 A significant correlation exists between $\delta^{18}\text{O}$ and elevation ($r = 0.53$, $n = 50$, $p < 0.001$) (**Fig. 3a**),
 354 suggesting that elevation is **an important** variable explaining **the** isotopic composition of **water** in the **study**
 355 lakes. Lakes sampled at lower elevations are more depleted while lakes sampled at higher elevations are
 356 enriched in ^{18}O and ^2H . **Another significant correlation occurs between $\delta^{18}\text{O}$ and the steepness of the slopes**
 357 **surrounding the lake** ($r = -0.33$, $n = 48$, $p = 0.02$), which suggests that morphometric factors may also
 358 **influence lake-water balance.**



360
 361 **Fig. 3:** Relation between $\delta^{18}\text{O}$ in water samples and elevation ($r = 0.53$, $n = 50$, $p < 0.001$) (a), and
 362 specific **conductance** and elevation ($r = -0.67$, $n = 50$, $p < 0.001$) (b). The horizontal dashed line indicates
 363 the position of a breakpoint analysis which occurs at an elevation of ~ 282 m asl, in both cases (**Tab. 1**)
 364

365 Evaporation-to-inflow ratio calculations (**E/I**) and **evaporative loss fractions of the lake volume (f)**
 366 show that groundwater springs and streams have E/I ratios close to 0 due to their short residence times
 367 while lakes have E/I or f values ranging from values similar to groundwater springs and streams to **near E/I**
 368 **or f ~ 1** owing to their longer residence times that expose them to evaporation (**Fig. 4a**).
 369

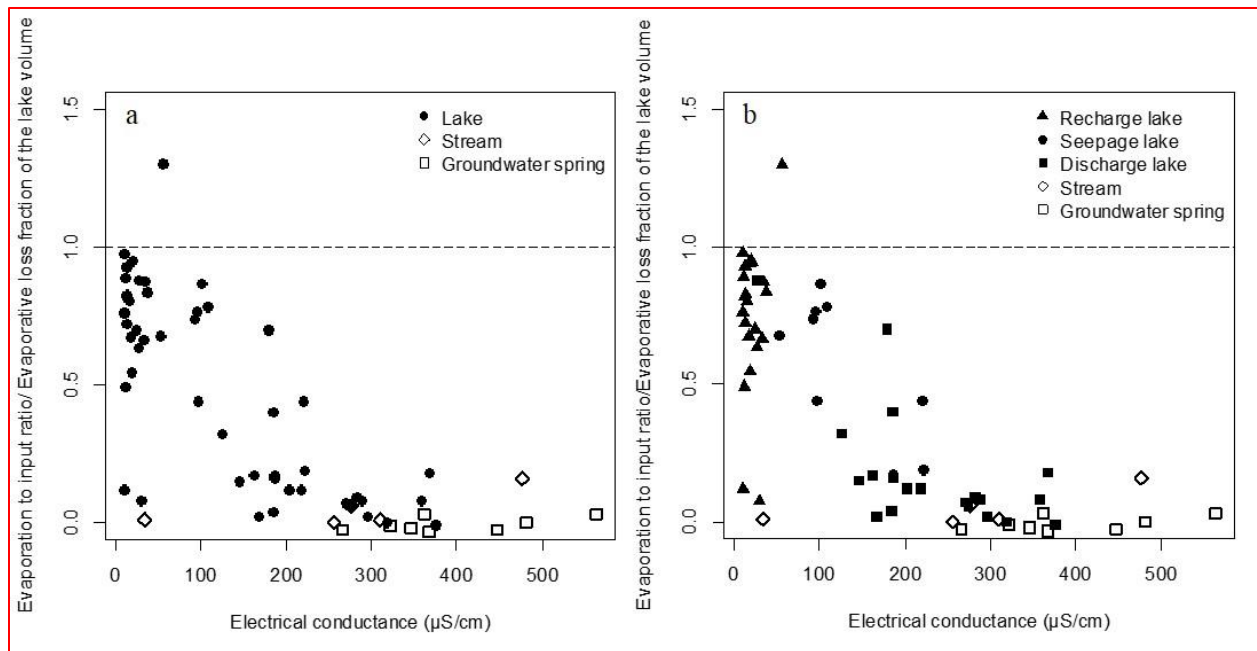


Fig. 4: Relation between electrical conductance and calculated evaporation-to-inflow ratios. Lakes in panel b are coded according to a classification scheme developed in this paper

4.2 *Solutes and dissolved organic matter*

As is the case with stable isotope values, water bodies reveal a wide range for non-conservative ions, and the overall sum of ions indicated by lake-water specific conductance, all of which are significantly correlated (see correlation matrix **Tab. A1**). Groundwater springs have the highest specific conductance (300-550 $\mu\text{S}/\text{cm}$) while streams have values *ca.* 300 $\mu\text{S}/\text{cm}$. Lakes show a wide range of solute content and values of specific conductance, from values similar to groundwater springs (*ca.* 400 $\mu\text{S}/\text{cm}$) to very low values similar to precipitation (as low as 10 $\mu\text{S}/\text{cm}$) (**Fig. 3b**). A significant correlation exists between the specific conductance and elevation ($r = -0.67$, $n = 50$, $p < 0.001$), suggesting that elevation is an important variable explaining specific conductance in lakes. There was also a significant relationship between the specific conductance and the ratio of perimeter to surface area of the lakes ($r = 0.48$, $n = 48$, $p < 0.001$), which also suggests that lake morphology may also be important.

Only 36 lakes were sampled for dissolved organic carbon (DOC) and nitrogen (TN). Unlike non-conservative ions and conservative isotopic tracers, no significant correlations were found between dissolved organic elements or TN and elevation ($r = -0.04$, $n = 36$, $p = 0.84$ for DOC; $r = 0.05$, $n = 36$, $p = 0.77$ for TN). However, significant, or marginally significant correlations were observed between DOC and perimeter to area ratio ($r = 0.54$, $n = 33$, $p < 0.001$) with more elongated lakes having higher concentrations of DOC, and between DOC and mean lake depth, with deeper lakes having lower concentrations of DOC

393 (r = -0.58, n = 25, p = 0.11). A similar pattern was observed between TN and perimeter to area ratio (r =
 394 0.60, n = 33, p = 0.002) and between TN and mean lake depth (r = -0.71, n = 25, p = 0.02).

395

396 4.3 Correlations between water tracers

397

398 There is a strong and significant correlation between lake-water isotopic values and specific
 399 conductance (r = 0.80, n = 50, p < 0.001). However, the slopes of the linear regressions for the water $\delta^{18}\text{O}$
 400 (Fig. 3a) and specific conductance (Fig. 3b) do not match the data points perfectly as there seem to be a
 401 distinct transition between similar values found in higher- and lower-elevation lakes, which was further
 402 examined using a breakpoint analysis of the lake water properties (the later undertaken to detect any step-
 403 wise changes in trends). Nine available environmental variables had a statistically significant breakpoint
 404 (i.e. an era of the line where the relationship between the variables changes trends) when regressed over
 405 elevation and significant breakpoints were within a narrow range of elevation with a mean of 282.4 m asl
 406 (Tab. 1).

407

408 **Tab. 1:** Results showing significant breakpoints in nine water chemistry variables and lake elevation
 409 (lower and upper elevation ranges represent the standard deviation)

Environmental Variable	Mean Elevation	Lower Elevation	Upper Elevation
$\delta^{18}\text{O}$	282.2	280.2	284.2
$\delta^2\text{H}$	282.0	279.9	284.1
<i>d</i> *	282.2	280.3	284.1
EC	284.0	281.4	286.6
Ca^{2+}	284.2	281.9	286.5
Mg^{2+}	282.0	280.3	283.7
K^+	281.8	278.2	285.4
DIC	281.5	279.6	283.4
TC	281.8	280.2	283.4
Breakpoint line	282.4		

410

**d* corresponds to deuterium excess

411

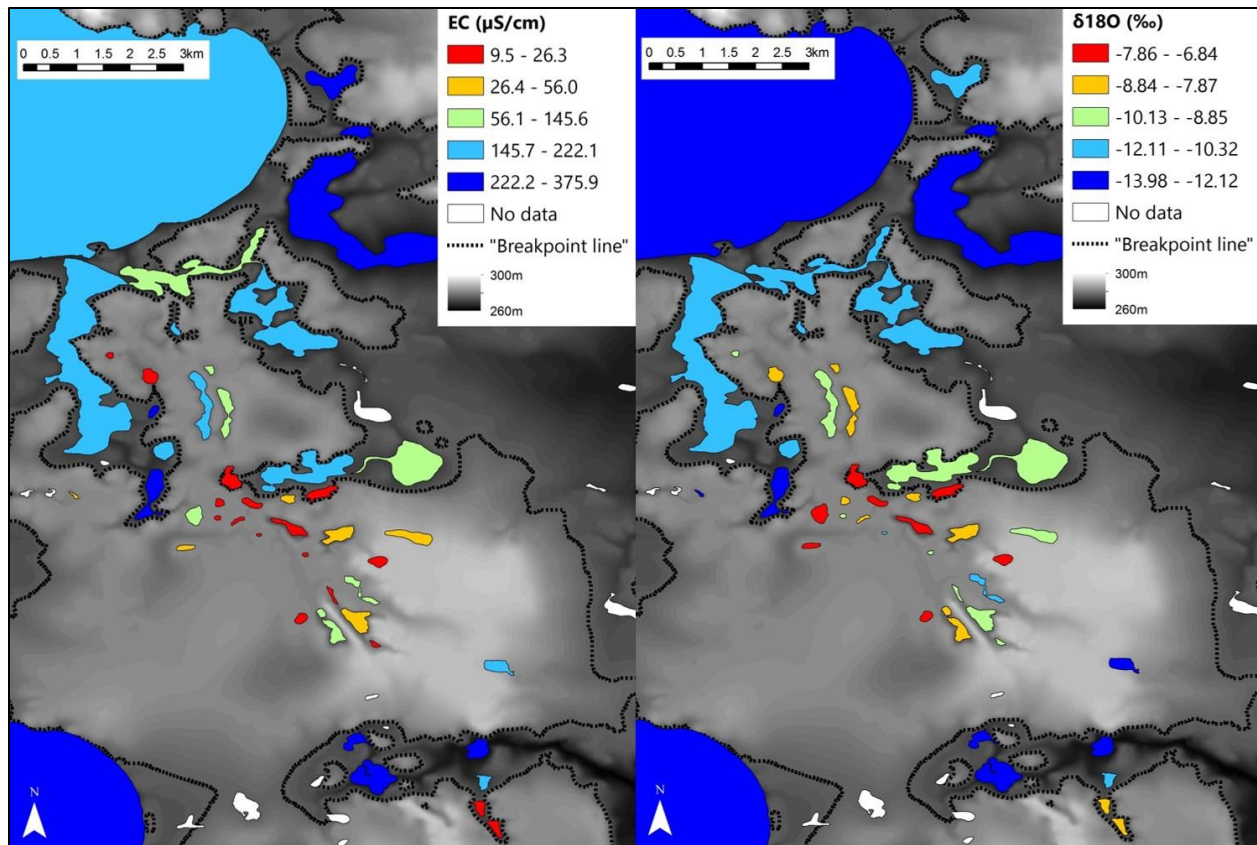
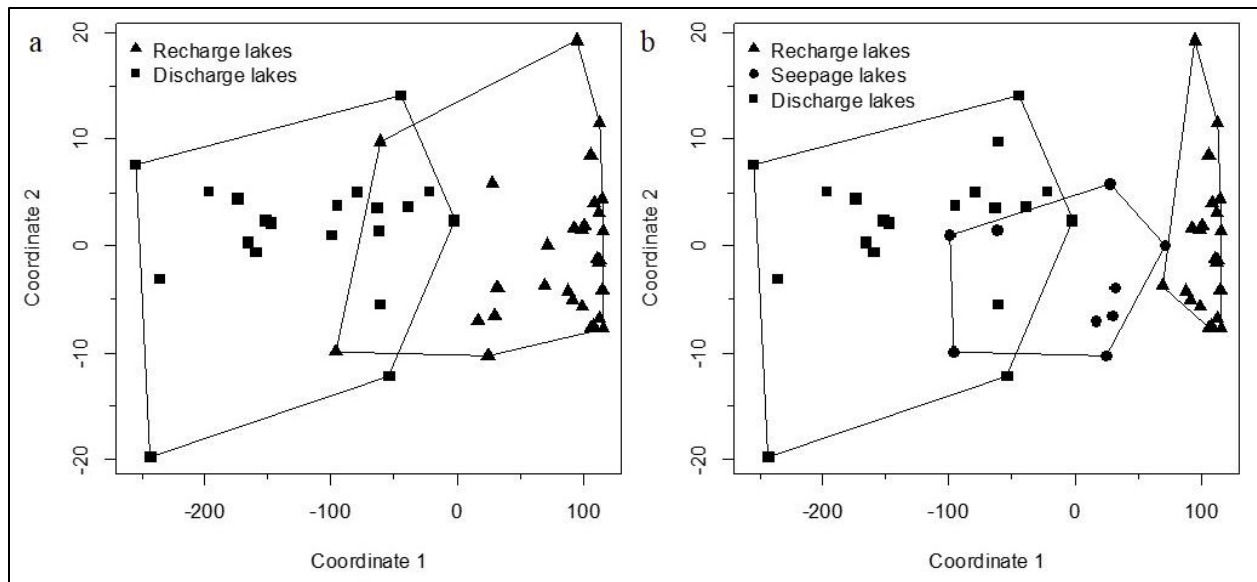


Fig. 5: Spatial depiction between elevation and lake-water specific conductance ($\mu\text{S}/\text{cm}$) (a), and lake-water $\delta^{18}\text{O}$ (‰) (b). The elevation of the breakpoint line (284 m) is shown as a dashed line

To further assess differentiation of the lakes in terms of conservative and non-conservative water tracers, a non-metric multidimensional scaling (NMDS) based on these factors was undertaken, with lakes above an elevation of 282 m coded as groundwater-recharge lakes, and lakes below 282 m coded as groundwater-discharge lakes, indicating good separation of the groups based on water geochemistry (**Fig. 6a**).

The existence of two distinctive types of lakes was used to develop a lake typology to explain changes in water biogeochemistry across the studied lakes. In order to better understand how water tracers vary in those two zones, individual **Wilcoxon signed-rank test** were undertaken for all lake-water variables that were above detection limits (**Tab. A2**). All anion and cations (and correlated variables including specific conductance), with the exception of **K**, individually were significantly different above and below an elevation of 282 m asl. This was also the case of the isotopic variables. Variables that were not significantly different included: **K and TN**.



430
 431 **Fig. 6:** Non-metric multidimensional scaling (NMSD) applied to conservative and non-conservative water
 432 tracers. Lakes that occurred at an elevation of greater than 282 m asl are labelled as 'recharge' lakes,
 433 whereas lakes as located at an elevation of less than 282 m are labelled as 'discharge' lakes (a). A third
 434 group of lakes (labelled 'seepage') is further discerned based on chemical and isotopic characteristics (b).
 435 **The model converged after 35 iterations.** Stress value for the NMSD is low (stress = 0.023)
 436

437 4.4 Short-term evolution of water tracers

438
 439 Short-term water balance variability was observed towards the end of the growing season **in 2014**
 440 **(Fig. A1b,d in the appendix)** and **between the sampling campaign of 2013 and 2014 (Fig. A2a,c)**. Lakes
 441 located at an elevation greater than 282 m asl underwent marked changes in water balance during the three
 442 sampling campaigns whereas lakes located below 282 m asl underwent little or no change, especially the
 443 lower lakes **(Fig. A2a,b)**. Similarly, the water-chemistry gradient between upland solute-poor lakes and
 444 lowland solute-rich lakes changed seasonally and between years. Lakes above 282 m asl did not respond
 445 chemically to short-term hydroclimatic change while lakes below 282 m asl displayed significant solute
 446 changes **(Fig. A2c,d)**.

447
 448 **5 Discussion**

449
 450 5.1 Interpretation of water tracers

451
 452 The wide range of lake isotopic values, **E/I and f** ratios suggest that lakes on the esker are
 453 heterogeneous in terms of water balance and hydrological characteristics. Groundwater springs have the

454 most depleted isotopic values because their water-residence times are short and they only undergo limited
455 evaporation (Gazis and Feng, 2004). Streams display isotopic values similar to groundwater springs **because**
456 they originate from groundwater and experience relatively low evaporation rates due to more continuous
457 water flow. The wide range of isotopic values for lakes can be explained by their position in the landscape,
458 particularly relative to their location in the esker aquifer system for which elevation appears to be a good
459 proxy. This suggests that lowland lakes are primarily fed by groundwater inflow while upland lakes receive
460 a much lower contribution from the local groundwater in their respective water budgets. Nonetheless, it is
461 challenging to distinguish the influence of groundwater from precipitation in lake-water balance as they
462 both have similar isotopic signatures (**Fig. 2b**) (Gibson and Edwards, 2002; Gibson *et al.*, 2008; Yi *et al.*,
463 2008). Yet, given that: i) our study was carried out on a small spatial scale (*i.e.* a rectangular zone of ~12
464 km by ~6 km); ii) the close proximity of the lakes; iii) the terrain homogeneity (with boreal forest as the
465 dominant land cover); iv) the limited topography; and v) the good correlation between water isotopes and
466 specific conductance ($r = 0.63$, $n = 50$, $p < 0.001$), it is unlikely that there are significant differences in
467 terms of precipitation patterns within our study area. Therefore, it is a reasonable assumption that
468 groundwater connectivity is the main control on lake water balance. Lake isotopic values can also be
469 influenced by the isotopic composition of surface inflowing waters. But those are **mainly** groundwater
470 springs (which are made up of groundwater), their volume **is small** in comparison to the lake volume and,
471 some of those streams are intermittent in the sense that they were not flowing during each of the three field
472 campaigns. Thus, it can reasonably be assumed that this influence is limited.

473
474 Similar patterns are observed with the major ions in water. Groundwater springs have the highest
475 solute **concentrations** likely due to chemical processes associated with mineral surface exchanges and
476 weathering (Ala-aho *et al.*, 2013). Lakes, however, displayed a wide range of water chemistries, and range
477 from high solutes characteristic of groundwater to values close to zero, typical of precipitation (**Fig. 2**),
478 suggesting that the heterogeneity of lakes on the esker are a result to the degree to which they interact with
479 groundwater. As it is the case for isotopic values, chemical composition of a given lake depends on
480 elevation. Lakes sampled at lower elevation are higher in solutes; this indicates that lowland lakes reflect
481 interaction with intermediate or regional groundwater flows subject to more mineral weathering and
482 dissolution (Tóth, 1963). Upland lakes on the other hand reflect interaction with local groundwater flow
483 paths with correspondingly reduced mineral weathering and dissolution (Tóth, 1963). There are only minor
484 differences in terms of the relative solute composition among the samples suggesting that the esker
485 subsurface material is geochemically relatively homogeneous and reflects the carbonate-rich nature of the
486 glaciofluvial outwash that makes up the esker (Cummings *et al.*, 2011). The carbonate-rich sediment

487 originates from Paleozoic carbonates of the Hudson sedimentary Platform *ca.* 150 km to the north and is
488 localized to glacial surficial sediments (Roy et *al.*, 2011).

489
490 Other characteristics of the lake water (temperature and dissolved organic matter) do not reflect the
491 changes seen in the stable isotopes and ionic concentration. Water temperatures of groundwater springs are
492 coolest because the temperature of groundwater is typically close to the mean annual temperature of the
493 region, while lake-water temperature varies strongly with season. There is little difference in temperature
494 between lakes and the latter are mainly the result of lake morphology than their connection with cool
495 groundwater as lakes of smaller volumes have a lower thermal inertia than larger ones. Differences in
496 dissolved organic content between the lakes is also a result of lake morphology as smaller lakes tend to
497 have higher concentrations as they mix more easily and receive greater inputs due to their high catchment
498 size-lake volume ratios (Knoll et *al.*, 2015). However, upland lakes tend to have slightly higher amounts
499 likely due to their higher water-residence times.

500

501 5.2 Lake hydrological classification

502

503 The correlation between lake-water $\delta^{18}\text{O}$ and specific conductance ($r = 0.80$, $n = 50$, $p < 0.001$) shows
504 there is a clear relation between conservative water tracers (which are indicative of water source and
505 evaporation) and non-conservative water tracers (which are indicative of water flow paths), signifying that
506 lowland lakes will receive a significant portion of their water as groundwater flows and will geochemically
507 reflect this origin, while upland lakes will receive most of their water through precipitation and will be
508 geochemically dilute.

509

510 Consequently, the observed breakpoint elevation appears to correspond to an effective hydraulic
511 separation between groundwater recharge and discharge areas (Winter et *al.*, 1998). This contrast between
512 the contribution of groundwater flow is evident in the distribution of lake-water composition (**Fig. 5**, **Fig.**
513 **A1**). Lakes characterized by groundwater discharge are spatially distinct from higher elevation lakes in the
514 groundwater recharge zone. Lake position has been used as a classification criterion in several studies (e.g.
515 Winter, 1997; Born et *al.*, 1979). Thus, upland lakes in the recharge zone are known as groundwater-
516 recharge lakes or recharge lakes and, conversely, lowland lakes in the discharge zone, also called underflow
517 zone, will be referred to as groundwater-discharge lakes or discharge lakes (**Fig. 7**) (Winter et *al.*, 1998).
518 Because discharge lakes receive a substantial amount of water from groundwater, they are considered to be
519 groundwater-fed or minerotrophic whereas recharge lakes which receive the majority of their water from
520 precipitation and feed the aquifer are said to be precipitation-fed or ombrotrophic (Webster et *al.*, 1996).

521 NMDS (**Fig. 6a**) and **Wilcoxon signed-rank test** (**Tab. A2**) analysis showed that all conservative and non-
522 conservative tracers are statistically different between the discharge zone and the recharge zone, except for
523 TN and K. An analysis of similarity (ANOSIM) between of the recharge and discharge lakes shows that
524 within group similarity is significantly greater than between group similarity, as illustrated by a large and
525 significant r value ($r = 0.77$, $n = 50$, significance = 0.001 on 1000 permutations).

526
527 Nevertheless, a few solute-rich alkaline lakes are located in the groundwater recharge zone which is
528 supposed to be depleted in solutes (**Fig. 5a**). Furthermore, the two primary groups in ordinal space display
529 a small overlap (**Fig. 6a**), suggesting the existence a third category of hybrid lakes referred to as seepage
530 or flow-through lakes (Winter, 1976; Webster *et al.*, 1996; Winter *et al.*, 2003). In those lakes, water comes
531 in as groundwater in-seepage and is returned to the groundwater system as out-seepage (Anderson and
532 Munter, 1981). Seepage lakes can be found both in the recharge zone and the discharge zone and they **can**
533 **contribute** a recharge or a discharge function (Anderson and Munter, 1981). In the groundwater recharge
534 zone, seepage lakes differ from recharge lakes by their pH, which is more alkaline while in the groundwater
535 discharge zone, seepage lakes differ from discharge lakes by the absence of an outlet, meaning they receive
536 less input than discharge lakes (**Fig. 8**). Lake-water chemistry and the presence of outlets have been used
537 in several studies as a classification criterion (e.g. Newton and Driscoll, 1990; Winter, 1977). NMDS (**Fig.**
538 **6b**) and **Wilcoxon signed-rank test** (**Tab. A3**) analysis showed that all conservative and non-conservative
539 tracers are statistically different between the three types of lakes, except for **SO₄**.

540
541 As noted earlier, lakes can first be classified according to their location within the groundwater
542 system, particularly above and below the hydraulic midline, in this case, at or near the 282 m elevation in
543 this study that indicated a breakpoint in the many isotopic and limnological variables (282.4 m) associated
544 with differences in hydrological inputs. In this study, elevation appears to be a proxy of the boundary
545 between the groundwater recharge and discharge zones on the esker. Seepage lakes, however, can be found
546 on each side of the hydraulic midline, thus generating additional classes of lakes (see conceptual diagram,
547 **Fig. 7**) distinguished from on the presence or absence of a lake outlet (**Fig. A1**) (Stauffer and Wittchen,
548 1992). Lakes with outlets were classified as “groundwater discharge lakes” and lakes without outlets as
549 “discharge seepage lakes”.

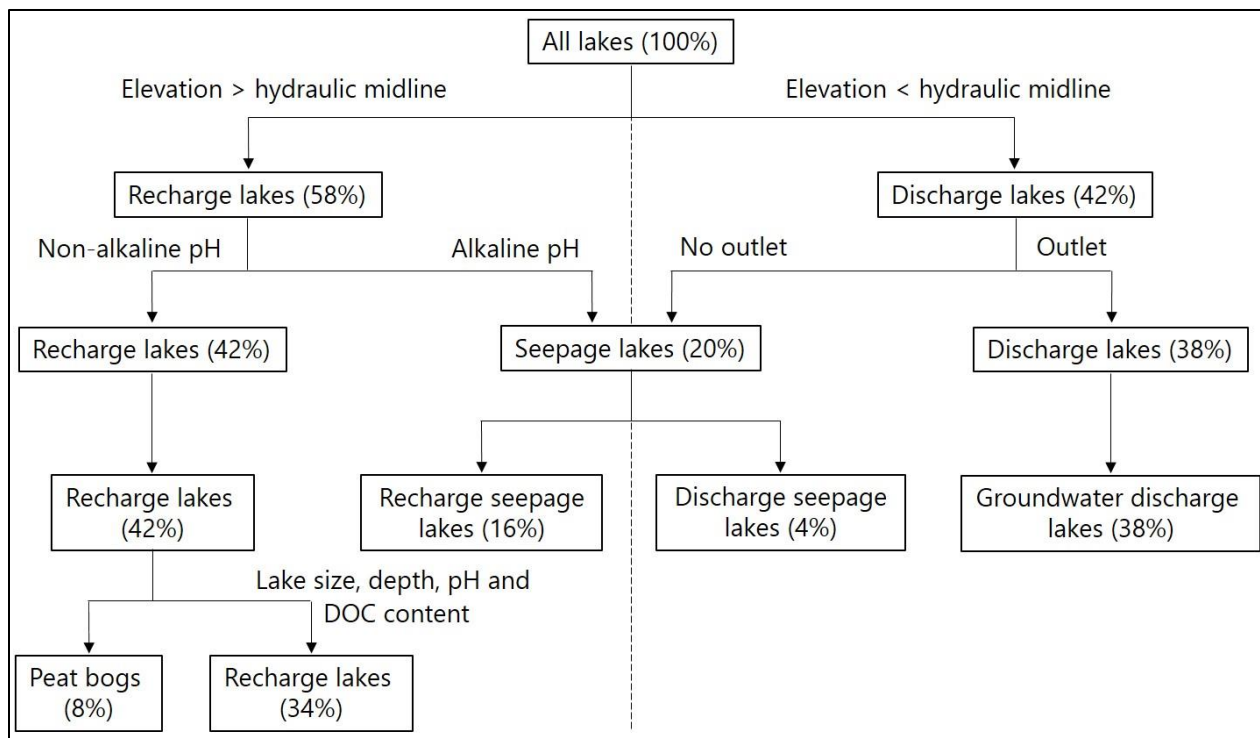
550
551 Lakes with outlets are typically found at the contact between the impermeable glaciolacustrine clay
552 plain and the permeable glaciofluvial esker sand contact where groundwater springs emerge (**Fig. 1b**).
553 Furthermore, in the study area, there is a significant relation between elevation and lake watershed
554 maximum slope ($r = 0.75$, $n = 50$, $p < 0.001$). Lithology is also a variable used as a classification criterion

555 in several studies (Winter, 1997; Martin *et al.*, 2011). Lakes located at the edge of the esker tend to have
556 steeper watershed slopes and have low elevations, which increases the likelihood of lakes to be in contact
557 with deeper groundwater flows (Winter, 1976). These lakes receive a substantial inflow of groundwater
558 due to their geological setting and their water isotopic and chemical composition is similar to the one of
559 groundwater springs and outlets (Ala-aho *et al.*, 2013).

560

561 Only two lakes in the groundwater discharge zone are without outlets and are classified as discharge
562 seepage lakes. The following hypothesis could potentially explain the absence of outlet: the ability of the
563 lakes to lose water to groundwater; or a reduced input from groundwater or surface inflow. Yet, some lakes
564 with outlets, like Pit Lake, can be found at elevations corresponding to the groundwater recharge zone (**Fig.**
565 **5**). The presence of Pit Lake at a distinctly high elevation (296 m) is nonetheless explained by the location
566 of this lake at the clay-sand interface (**Fig. 1b**). This suggests that elevation is just a putative variable or, in
567 other words, a proxy of the actual hydraulic midline delineated by the clay-sand interface. This is confirmed
568 numerically by applying a logistic regression to the presence or absence of a lake outlet and lake elevation,
569 which provided a poor relation (Mc-Fadden $r = 0.40$, $n = 50$, $p < 0.01$). On the other hand, the logistic
570 regression of the presence or absence of a lake outlet to the closest distance to the sand-clay contact provided
571 a better relation (Mc-Fadden $r = 0.69$, $n = 50$, $p < 0.001$), although it is challenging to find the real clay-
572 sand interface as the lateral sands can mask the real exact location (**Fig. 1b**) and surficial geological maps
573 have errors in tens of meters in comparison to field observations.

574



575
 576 **Fig. 7:** Illustration of a lake typology for the study lakes. Lake elevation was used as the primary factor in
 577 separating lakes, with elevation being an effective surrogate for dividing lakes based on contribution of
 578 groundwater flow to a lake. The lakes were then separated based on presence of an outlet, as well as lake
 579 water pH and other morphological characteristics of the lakes
 580

581 Some lakes in the recharge zone, called “recharge seepage lakes”, have an alkaline or a circumneutral
 582 pH and higher solute content, signifying that they interact to some extent with groundwater. This could
 583 occur for a number of reasons related to the existence of significant groundwater input including: i) slightly
 584 steeper lake watershed slopes; ii) geographical proximity to the clay-sand interface; and iii) lake relative
 585 deepness allowing interactions with deeper groundwater flows, or a combination of those.

586
 587 Recharge lakes can be further subdivided into two types of recharge lakes: “classical” recharge lakes
 588 and peat bogs (**Fig. 7**), adding a fifth type of lake in the typology (Newton and Driscoll, 1990). Sampled
 589 peat bogs are very small and shallow lakes rimmed by floating mats of vegetation (typically less than 1 ha
 590 and 1 to 2 m deep), acidic, characterized by a very low amount of solutes (maximum 30 $\mu\text{S}/\text{cm}$), a relatively
 591 high amount of dissolved organic carbon (above 10 ppm) and by water isotopic composition controlled by
 592 short-term hydroclimatic conditions (enriched in heavy water isotopes during drier periods and similar to
 593 the isotopic composition of precipitation during wet periods). This could result from relative hydrological
 594 isolation from the groundwater system due to the thick layer of peat at their bottom formed by the successive
 595 accumulation of sedge and sphagnum characterized by a low hydraulic conductivity (Newton and Driscoll,

596 1990). As a consequence, direct precipitation would be the predominant source of water, making these lakes
597 sensitive to hydroclimatic variability.

598
599 A sixth type of lake can be added to the proposed lake typology: ephemeral kettle ponds, commonly
600 called dry kettles, even though they are not lakes *per se*. They can be identified on geological maps, aerial
601 photography and satellite imagery. Dry kettles consist of small kettle holes adjacent to the esker crest that
602 are usually but not always too high in elevation and shallow to be connected to the water table and will be
603 dry during the summer months and most of the year, creating patches of bare land or wetlands in the forested
604 landscape.

605
606 Based on this typology, 42% of the sampled lakes were recharge lakes, 20% seepage lakes and 38%
607 discharge lakes. Other studies report slightly different proportions (e.g. Anderson and Munter, 1981) as
608 might be expected based on the geographical location, the size of the study area, and other factors relative
609 to the groundwater system. The subdivision of lakes into recharge and discharge lakes is consistent with
610 Ala-aho et al. (2013) who worked in similar settings and established their classification based on water
611 solutes. The three-fold typology compares well with Turner et al. (2010) who also established a three
612 category lake typology although the terminology of the three lake types differ and E/I and f values were
613 used as the primary classification criteria. Hence, the threefold typology of **Fig. 7** is particularly relevant
614 for water resources management in esker complexes as it uses readily available variables (*i.e.* elevation,
615 specific conductance/pH and presence/absence of an outlet).

616
617 Discharge (ground-water discharge) lakes have the highest pH, and amounts of solutes, while the
618 opposite is observed for recharge lakes. Seepage lakes have intermediate values compared to the other two
619 lake types. The same patterns are observed for isotopic values: recharge lakes tend to be enriched in heavier
620 water isotopes, discharge lakes tend to be depleted in heavier isotopes, and seepage lakes characterized by
621 intermediate values. This reinforces the interpretation that seepage lakes are a hybrid type between recharge
622 and discharge lakes. Seepage lakes thus tend to be closer to recharge lakes on an isotopic basis, but more
623 similar to discharge lakes on a chemical basis. This is explained by the fact that most sampled seepage lakes
624 are located in the recharge zone and those receive some intermediate groundwater flows.

625
626 Recharge lakes contain higher amounts of dissolved organic matter likely due to substantial water
627 residence times, whereas seepage lakes and discharge lakes can have sustained inflow of groundwater that
628 decreases water residence times. The concentration of DOC in discharge lakes is relatively higher than
629 expected as groundwater usually contain limited organic matter. This is likely an artefact of lake

630 morphology and anthropogenic activities. Indeed, values of DOC for discharge lakes display a high
631 standard deviation due to a subgroup of smaller lakes that are relatively deep and without cottages,
632 combined with another group of larger lakes that are relatively shallow with the presence of cottages.
633 Relatively shallow lakes are known to have significantly higher amounts of DOC than deeper ones as a
634 result of smaller volumes relative to inputs, and potential reworking of bottom organic matter and nutrients
635 due to the shallow water columns. DOC values for discharge lakes also contain two sizeable outliers that
636 skew the average and median: Nighthawk Lake and Frederick House Lake, both of which are two regional
637 high-order and large lakes (10,701 ha and 3,888 ha respectively) that are relatively shallow (maximum
638 depth of 4.6 m and 12.0 m respectively) and are heavily used for recreational purposes.

639

640 5.3 Lake morphometry and water geochemistry

641

642 There are additional morphometric factors that influence lake water isotopic composition in these
643 settings. Evaporation over a water body is enhanced when there is a strong gradient of temperature and
644 water vapour pressure between the lake water and the adjacent air. Assuming that there is no large
645 microclimate differences within the esker complex, isotopic values are influenced by: i) the lake fetch ($r =$
646 -0.26 , $n = 48$, $p = 0.07$) as wind is more efficient at removing moisture over a long distance (Granger and
647 Hedstrom, 2011); ii) relative depth ($r = -0.32$, $n = 32$, $p = 0.07$) as lakes that have a large surface compared
648 to their depth are proportionally more exposed to the atmosphere and thus more susceptible to evaporation;
649 and iii) the steepness of the slopes surrounding the lake ($r = -0.33$, $n = 48$, $p = 0.02$). Only the latter variable
650 has a significant relation with the isotopic composition of the lake. Steep slopes tend to reduce evaporation
651 rates by blocking air flows over the lake, thus reducing wind speed, water-air temperature and water vapour
652 pressure contrasts, and to increase the likelihood of lakes to be in contact with deeper groundwater flows
653 (Winter, 1976).

654

655 Lake-water chemistry can also be influenced by lake morphometry. Lake depth has been known to
656 play a role (Winter, 1976). It seems that *a priori* there is no relation between lake maximum depth and
657 specific conductance ($r = 0.09$, $n = 39$, $p = 0.59$). However, to determine whether the relation between
658 specific conductance and lake maximum depth varied with lake landscape position, two separate
659 regressions analyses were carried out for recharge ($r = 0.01$, $n = 16$, $p = 0.66$) and discharge lakes ($r = 0.30$,
660 $n = 23$, $p = 0.05$). The two regressions show that maximum depth can act as a control on specific
661 conductance only in lakes in the discharge zone: the deeper the lake, the deeper solutes-rich groundwater
662 flows seep into the lake. It is also expected that small lakes with a large ratio of perimeter to surface area (r

663 = 0.48, $n = 48$, $p < 0.001$) will have a higher amounts of solutes as the majority of groundwater seepage
664 into the lake is typically localised near the shoreline (Rosenberry *et al.*, 2015).

665
666 Water-quality tracers (DOC and TN) were not correlated to elevation, suggesting there is no apparent
667 relationship between groundwater inflow and water quality. The correlation between water-quality tracers
668 and the ratio of lake perimeter to surface area may suggest a relation between lake morphometry and water
669 quality. Elongated lakes have proportionally a greater interface with terrestrial ecosystem, which favours
670 the inflow of nutrients in lakes while mean depth is seen as an indicator of water residence times and rates
671 of mixing (Knoll *et al.*, 2015; Mulholland, 2003).

672
673 Given this, it can be argued that lake morphometry can play a significant role on lake-water
674 biogeochemistry and introduce some complexity in the lake typology. As a fact, some lake typologies
675 incorporate morphometric features as defining characteristics (e.g. Knoll *et al.*, 2015; Martin *et al.*, 2011;
676 Winter, 1977). Even though water tracers show a stronger correlation with elevation than with
677 morphometric features, it cannot be assessed with certainty that elevation accounts for much more variance
678 in isotopic composition and specific conductance as certain variable that influence lake isotopic
679 composition (e.g. lake network number which measures connections to other lakes, specific yield, *etc.*) and
680 lake-water chemistry (e.g. lake to catchment area ratio, soil type, specific yield, *etc.*) are not available for
681 comparison. In other studies, land use has been identified as a key variable explaining lake water balance
682 (as canopy cover reduce evaporative losses) and water chemistry (as substrate and water pathways influence
683 water chemistry) (e.g. Turner *et al.*, 2010; Turner *et al.*, 2014a and Turner *et al.*, 2014b). However, given
684 the predominance of the boreal forest in our study area, the impact of land use on water geochemistry has
685 been discarded as an explanatory variable. All those variables can trump the influence elevation has on
686 water balance and water chemistry between lake basins.

687

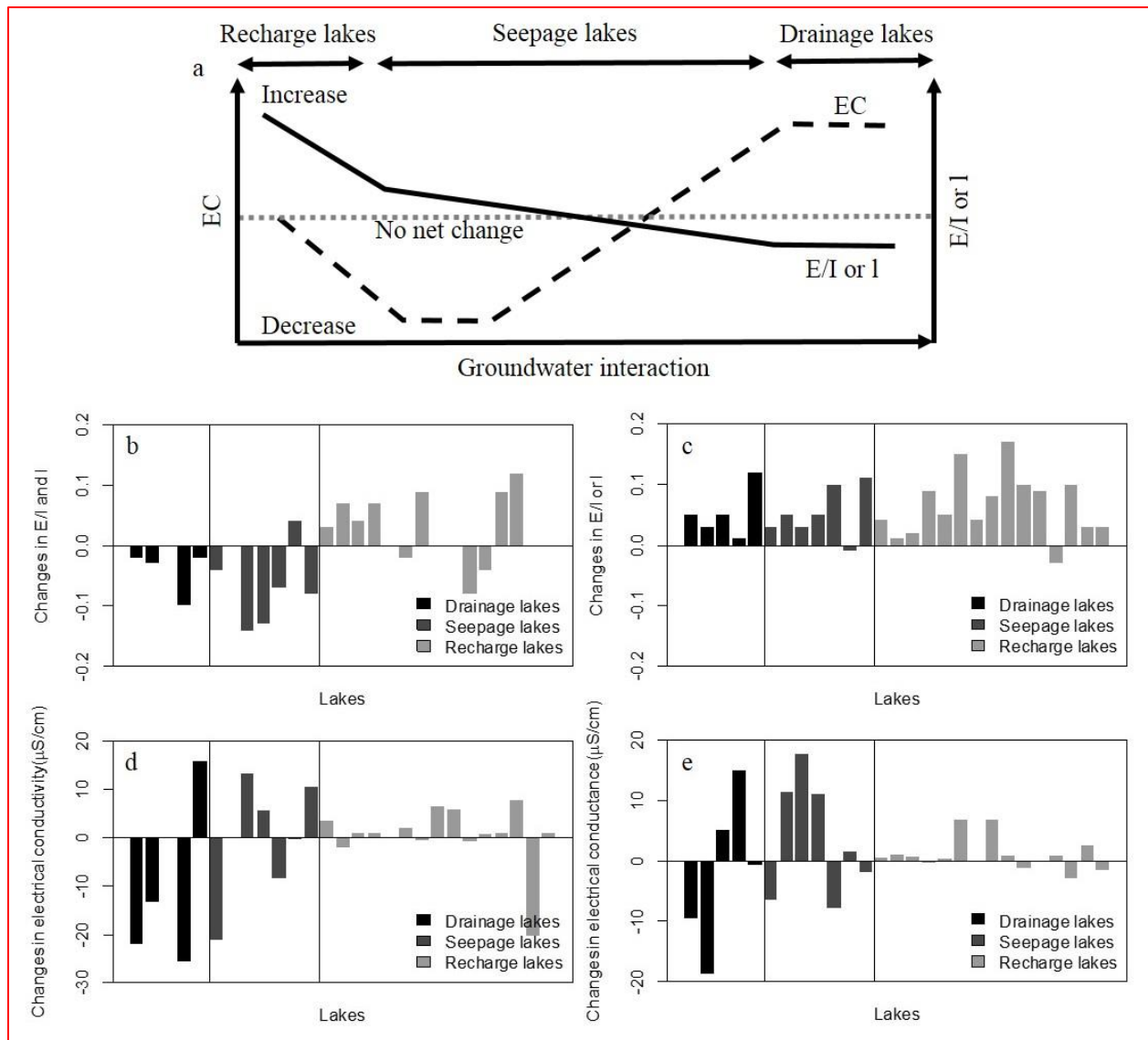
688 5.4 Implications for water balance, hydrochemistry and response to external stressors

689

690 Results from the study show that the water balance of individual lakes is highly dependent on the
691 nature of their interactions with groundwater that is in turn determined by landscape position at highly
692 localized scale as shown in the conceptual model in Fig 8a. Discharge lakes undergo minimal to negative
693 changes in water balance as the continuous flow of isotopically light groundwater masks evaporative
694 enrichment whereas recharge lakes are highly sensitive to evaporation as they rely on precipitation as their
695 primary source of water. Seepage lakes constitute a hybrid between recharge and discharge and their water
696 balance is between the other two lake types (Fig. 4b). These patterns of change were observed when

697 comparing changes in E/I and f ratios between sampling campaigns: E/I and f ratios experienced negative
698 noticeable changes in discharge lakes and small changes in seepage lakes while positive changes in E/I and
699 f were observed in recharge lakes (Fig. 8b and Fig. 8c). Consequently, upland recharge lakes will be more
700 prone to evaporative drawdown and therefore more sensitive to short-term climate change and droughts,
701 while discharge lakes will be buffered by groundwater inflow and affected by hydroclimatological changes
702 of greater duration and persistence that alter water table position (Fig. 8a). Seepage lakes will presumably
703 be sensitive to drought but not to the same extent as recharge lakes as they have stronger interactions with
704 groundwater (Fig. 8a). This suggests that the esker hydrological system can respond to a large-scale
705 hydroclimatic forcing (e.g. prolonged drought) in a manner that affects individual lakes differently. The
706 degree of interaction with groundwater by an individual lake will also dictate the response to strong
707 hydroclimatic forcings and introduce time lags (Webster et al., 2000).

708



709
 710 **Fig. 8:** Conceptual model of the relationship between the direction and magnitude of lake E/I or f and EC
 711 changes during drier conditions to lake type defined by the degree to which lakes interact with
 712 groundwater (modified from Webster *et al.* (1996)) (a) and observed changes in E/I or f between June
 713 2013 and June 2014 (E/I 2014 – E/I 2013) (b) and changes in E/I between August 2014 and June 2014
 714 (E/I or f Aug – E/I or f Jun) (c), and changes in specific conductance between June 2013 and June 2014
 715 (EC 2014 – EC 2013) (d) and changes in electrical conductance between August 2014 and June 2014 (EC
 716 Aug – EC Jun) (e) by lake type

717
 718 Similarly, the water-chemistry gradient between upland solute-poor lakes and lowland solute-rich
 719 lakes could make recharge lakes more vulnerable to acidification than seepage and discharge lakes. The
 720 effects of changing hydroclimatic conditions on lake-water chemistry across the landscape observed during
 721 successive sampling in June 2013, June 2014 and August 2014 resulted in heterogeneous pattern depending
 722 on the lake type, consistent with the findings of Webster *et al.* (1996). Recharge lakes did not respond

723 chemically to short-term hydroclimatic change as they feed the groundwater system by lake out-seepage
724 thus producing no net change in solutes, while seepage and discharge lakes displayed significant solute
725 changes (**Fig. 8b and Fig. 8c**). Based on the limited sampling frequency, it is difficult to draw conclusions
726 on which of the seepage or discharge lake types undergo the most chemical variation. **A greater temporal**
727 **resolution of sampling would provide more insights on short-term changes in hydrochemistry in relation to**
728 **short-term hydroclimatic fluctuations and reinforce positively our interpretations.** However, Webster *et al.*
729 (1996) suggest that discharge lakes and seepage lakes respond chemically to evaporative drawdown in
730 opposite ways, with seepage lakes showing a decline in solutes during droughts as inputs from groundwater
731 diminish due to the lowering of the water table (**Fig. 8a**). By contrast, discharge lakes have been noted as
732 susceptible to evaporative enrichment of solutes and increased relative contribution of solute-rich
733 groundwater during drought periods (Kratz *et al.*, 1997). For these reasons, Webster *et al.* (1996) suggested
734 that climate change could amplify anthropogenic impacts and make lakes more vulnerable to other stressors,
735 such as lake acidification.

736
737 Due to their varied hydrological characteristics, the lake types identified in this study will have a
738 different susceptibility to direct anthropogenic impacts. Because recharge lakes have lower groundwater
739 inflow, they are characterized by relatively long water-residence times, making them highly vulnerable to
740 inputs **and pollutants** in comparison to seepage lakes and discharge lakes that tend to have a greater
741 watershed, which results in increased flushing. **Given this situation, cottage development should be limited**
742 **in the groundwater recharge zone and/or stricter regulations should apply on cottage septic tanks in this**
743 **zone to prevent the downstream contamination and subsequent degradation of water quality in the**
744 **groundwater discharge zone delimited in Fig 5.** However, it can be argued that differences in material
745 permeability between the esker and the clay plain can produce the opposite effect **as clays and organic**
746 **deposits on the esker flanks act as an aquitard that locally confine the aquifer (Fig. 1b) (Rossi et al., 2012).**
747 Sand extraction and mining activities in the groundwater discharge zone could potentially influence the
748 water levels of upland lakes in the recharge zone (Klove *et al.*, 2011). **Indeed, material excavation and**
749 **tunnel construction can cause the desiccation of groundwater-dependent systems by reversing flow patterns**
750 **in the recharge area, and increase the discharge from the esker after ditching on the esker flanks and drain**
751 **groundwater-dependent systems in the recharge area (Klove et al., 2011; Rossi et al., 2012).** As a result,
752 **any excavation project in those settings should not be conducted without an incidence study that investigates**
753 **and takes into account local groundwater flow patterns and hydrological connections between the recharge**
754 **and discharge areas.** This potential was acknowledged by an impact study undertaken for an aggregate pit
755 project in the study area, and one recommendation was the construction of an engineered frozen earth

756 barrier to prevent ground water flow into the proposed pit in order to minimize the effects on the water table
757 and surrounding lake levels (Cochrane, 2006).

758
759 Finally, when considering groundwater-fed lakes for the purposes of paleohydrological
760 reconstruction (e.g. Laird et al., 2012), it is critical to explore their modern hydrology to be able to **have a**
761 **baseline for comparison to future hydrological regimes that may alter them and subsequently** correctly
762 interpret the probable causes of isotopic, chemical, and biological change and variability recorded in the
763 sediment through time along with their potential hydroclimatic drivers. Because the degree to which lakes
764 interact with groundwater produces differences in hydrologic response to **the** same hydroclimatic forcing,
765 the interpretation of paleolimnological records can be complex (Fritz, 2000) and multiple site selection
766 **seems necessary** as a discharge lake may show long-term stability while a recharge lake may display
767 significant short-term and long-term variability (Bennett et al., 2007).

768

769 **6 Conclusion**

770

771 Lakes located in an esker complex in northeast Ontario showed strong systematic and localized
772 differences in terms of water balance and hydrochemistry, similar to other esker complexes in other settings.
773 Results from this study indicated that elevation is a critical factor explaining water chemistry and water
774 balance across the landscape. As eskers are structurally complex and often characterized by high hydraulic
775 conductivity, groundwater interactions are an important component of lake water hydrology. Low-elevation
776 lakes are likely interconnected with solute-rich intermediate and regional groundwater flowpaths while
777 upland lakes are only interacting with local solute-poor groundwater flowpaths. This threshold in water
778 chemistry is also accompanied by strong contrasts in lake-water balance. Upland lakes tend to be
779 isotopically-enriched and more sensitive to evaporation, while lowland lakes are more depleted and subject
780 to groundwater inflows. Limnological variables including lake depth, DOC and forms of nitrogen were
781 **weakly** related to elevation, and likely influenced by lake morphometry and watershed activities. Thus,
782 these results **are in agreement with** other studies that indicate that at the local scale, landscape position is
783 the main control on lake water chemistry and balance **while lake morphometric characteristics explain**
784 **additional variance.**

785

786 The physical and chemical characteristics of lake water allowed the development of a lake typology
787 that is made up of three main types of lakes: i) higher-elevation groundwater-recharge lakes, essentially fed
788 by precipitation, **are** characterized by higher evaporations rates and lower amounts of solutes; ii) seepage
789 lakes, that both gain and lose water to the groundwater, **are** characterized by intermediate rates of

790 evaporation and amounts of solutes; and iii) lower-elevation groundwater-discharge lakes, are continuously
791 fed by groundwater inflow and characterized by almost no evaporation rates and higher amounts of solutes.
792 The obtained typology provides insights about lake vulnerability to environmental stressors, particularly
793 short- and long-term hydroclimatic change. Recharge lakes will be more prone to evaporative drawdown
794 and therefore more sensitive to short-term droughts, while discharge lakes will be buffered by groundwater
795 inflow and affected by hydroclimatological changes of greater persistence. Similarly, recharge lakes will
796 likely be more subject to other anthropogenic impacts, in comparison to discharge lakes.

797

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799

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810

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1084 **Author contributions**

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1086 Maxime Boreux collected data in the field and processed samples in the lab, conducted data analysis and
1087 interpretation (with input from all authors), generated figures and tables. Scott Lamoureux provided
1088 guidance, funding, reviewed and edited the manuscript in his function as supervisor, as did Brian Cumming.
1089 The manuscript was written by Maxime Boreux with input from all authors.

1090 **Data availability**

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1092 The data can be made available by contacting the corresponding author.

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1094 **Competing interests**

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1096 The authors declare that they have no conflict of interest.

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1124 **Appendix**

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1126 **Tab. A1a:** Correlation matrix between hydrological tracers and morphometric variables

	EC	pH	T	Ca ²⁺	Mg ²⁺	K ⁺	Na ⁺	Cl ⁻	SO ₄ ²⁻	NO ₃ ⁻	NH ₄ ⁺	δ ¹⁸ O	δ ² H
EC	1.00	0.73	-0.22	0.97	0.93	0.73	0.60	0.62	0.36	-0.13	-0.18	-0.80	-0.75
pH	0.73	1.00	-0.05	0.76	0.73	0.41	0.29	0.29	0.20	-0.15	-0.18	-0.47	-0.41
T	-0.22	-0.05	1.00	-0.24	-0.25	-0.32	0.06	0.05	-0.12	0.19	-0.16	0.34	0.33
Ca ²⁺	0.97	0.76	-0.24	1.00	0.94	0.66	0.43	0.48	0.40	-0.12	-0.18	-0.83	-0.78
Mg ²⁺	0.93	0.73	-0.25	0.94	1.00	0.72	0.36	0.37	0.26	-0.12	-0.18	-0.77	-0.73
K ⁺	0.73	0.41	-0.32	0.66	0.72	1.00	0.56	0.49	0.37	-0.01	0.05	-0.55	-0.51
Na ⁺	0.60	0.29	0.06	0.43	0.36	0.56	1.00	0.95	0.12	-0.06	-0.07	-0.31	-0.29
Cl ⁻	0.62	0.29	0.05	0.48	0.37	0.49	0.95	1.00	0.06	-0.06	-0.04	-0.35	-0.33
SO ₄ ²⁻	0.36	0.20	-0.12	0.40	0.26	0.37	0.12	0.06	1.00	0.03	-0.05	-0.33	-0.30
NO ₃ ⁻	-0.13	-0.15	0.19	-0.12	-0.12	-0.01	-0.06	-0.06	0.03	1.00	0.04	0.16	0.16
NH ₄ ⁺	-0.18	-0.18	-0.16	-0.18	-0.18	0.05	-0.07	-0.04	-0.05	0.04	1.00	0.14	0.16
δ ¹⁸ O	-0.80	-0.47	0.34	-0.83	-0.77	-0.55	-0.31	-0.35	-0.33	0.16	0.14	1.00	0.99
δ ² H	-0.75	-0.41	0.33	-0.78	-0.73	-0.51	-0.29	-0.33	-0.30	0.16	0.16	0.99	1.00
<i>d</i>	0.84	0.53	-0.34	0.87	0.82	0.59	0.33	0.37	0.35	-0.16	-0.11	-0.99	-0.96
E/I f	-0.80	-0.66	0.45	-0.83	-0.77	-0.59	-0.37	-0.30	-0.78	0.18	0.24	0.94	0.93
TC	0.96	0.75	-0.43	0.96	0.95	0.74	0.59	0.42	0.36	-0.17	-0.03	-0.88	-0.85
DOC	-0.12	-0.25	-0.05	-0.09	-0.19	0.26	0.00	-0.14	0.59	-0.03	0.50	-0.06	-0.08
DIC	0.96	0.81	-0.38	0.95	0.97	0.62	0.56	0.44	0.16	-0.16	-0.21	-0.82	-0.79
TN	-0.22	-0.30	-0.09	-0.21	-0.29	0.16	0.00	-0.09	0.44	-0.06	0.53	0.08	0.06
C/N	-0.13	-0.21	0.28	-0.10	-0.10	0.12	-0.10	-0.20	0.35	0.04	0.17	-0.07	-0.08
H	-0.67	-0.57	0.04	-0.69	-0.72	-0.55	-0.31	-0.28	-0.25	0.08	0.16	0.53	0.49
Z	0.09	0.20	0.17	0.06	0.18	0.06	-0.01	0.00	-0.18	0.08	0.01	0.01	0.05
S	0.51	0.50	-0.04	0.52	0.53	0.45	0.22	0.16	0.39	-0.10	-0.24	-0.33	-0.27
A	0.35	0.40	0.23	0.37	0.38	0.32	0.20	0.17	0.18	-0.06	-0.08	-0.24	-0.21
P	0.42	0.53	0.25	0.45	0.45	0.35	0.21	0.17	0.24	-0.07	-0.13	-0.29	-0.27
P/A	-0.48	-0.73	-0.28	-0.48	-0.49	-0.29	-0.29	-0.23	-0.40	0.00	0.21	0.10	0.02

1127 Note: electrical conductance (EC) is expressed in μS/cm, temperature (T) in °C, dissolved ions (Ca²⁺,
 1128 Mg²⁺, K⁺, Na⁺, Cl⁻, SO₄²⁻, NO₂⁻-NO₃⁻, NH₄⁺) in ppm, water stable isotopes (δ¹⁸O, δ²H) and deuterium excess
 1129 (*d*) in V-SMOW, total carbon (TC), dissolved organic carbon (DOC), dissolved inorganic carbon (DIC)
 1130 and total nitrogen (TN) in ppm, elevation (H), lake maximum depth (Z) and perimeter (P) in m, maximum
 1131 lake watershed slope (S) in % and area (A) in ha.

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Tab. A1b: Correlation matrix between hydrological tracers and morphometric variables (con't)

	<i>d</i>	E/I f	TC	DOC	DIC	TN	C/N	H	Z	S	A	P	P/A
EC	0.84	-0.80	0.96	-0.12	0.96	-0.22	-0.13	-0.67	0.09	0.51	0.35	0.42	-0.48
pH	0.53	-0.66	0.75	-0.25	0.81	-0.30	-0.21	-0.57	0.20	0.50	0.40	0.53	-0.73
T	-0.34	0.45	-0.43	-0.05	-0.38	-0.09	0.28	0.04	0.17	-0.04	0.23	0.25	-0.28
Ca ²⁺	0.87	-0.83	0.96	-0.09	0.95	-0.21	-0.10	-0.69	0.06	0.52	0.37	0.45	-0.48
Mg ²⁺	0.82	-0.77	0.95	-0.19	0.97	-0.29	-0.10	-0.72	0.18	0.53	0.38	0.45	-0.49
K ⁺	0.59	-0.59	0.74	0.26	0.62	0.16	0.12	-0.55	0.06	0.45	0.32	0.35	-0.29
Na ⁺	0.33	-0.37	0.59	0.00	0.56	0.00	-0.10	-0.31	-0.01	0.22	0.20	0.21	-0.29
Cl ⁻	0.37	-0.30	0.42	-0.14	0.44	-0.09	-0.20	-0.28	0.00	0.16	0.17	0.17	-0.23
SO ₄ ²⁻	0.35	-0.75	0.36	0.59	0.16	0.44	0.35	-0.25	-0.18	0.39	0.18	0.24	-0.40
NO ₃ ⁻	-0.16	0.18	-0.17	-0.03	-0.16	-0.06	0.04	0.08	0.08	-0.10	-0.06	-0.07	0.00
NH ₄ ⁺	-0.11	-0.24	-0.03	0.50	-0.21	0.53	0.17	0.16	0.01	-0.24	-0.08	-0.13	0.21
δ ¹⁸ O	-0.99	0.94	-0.88	-0.06	-0.82	0.08	-0.07	0.53	0.01	-0.33	-0.24	-0.29	0.10
δ ² H	-0.96	0.93	-0.85	-0.08	-0.79	0.06	-0.08	0.49	0.05	-0.27	-0.21	-0.27	0.02
<i>d</i>	1.00	-0.92	0.90	0.05	0.85	-0.11	0.05	-0.57	0.04	0.40	0.28	0.32	-0.20
E/I f	-0.92	1.00	-0.81	0.37	-0.81	0.49	-0.05	0.48	-0.07	-0.37	-0.22	-0.31	0.24
TC	0.90	-0.81	1.00	0.05	0.94	-0.05	0.01	-0.64	0.03	0.50	0.31	0.38	-0.27
DOC	0.05	0.37	0.05	1.00	-0.29	0.93	0.66	-0.04	-0.29	0.08	0.00	-0.12	0.54
DIC	0.85	-0.81	0.94	-0.29	1.00	-0.37	-0.21	-0.60	0.12	0.45	0.29	0.39	-0.40
TN	-0.11	0.49	-0.05	0.93	-0.37	1.00	0.40	0.05	-0.43	-0.03	-0.01	-0.10	0.60
C/N	0.05	-0.05	0.01	0.66	-0.21	0.40	1.00	-0.09	0.08	0.18	0.07	-0.03	0.03
H	-0.57	0.48	-0.64	-0.04	-0.60	0.05	-0.09	1.00	-0.26	-0.77	-0.46	-0.50	0.53
Z	0.04	-0.07	0.03	-0.29	0.12	-0.43	0.08	-0.26	1.00	0.30	-0.02	0.04	-0.40
S	0.40	-0.37	0.50	0.08	0.45	-0.03	0.18	-0.77	0.30	1.00	0.33	0.35	-0.52
A	0.28	-0.22	0.31	0.00	0.29	-0.01	0.07	-0.46	-0.02	0.33	1.00	0.91	-0.49
P	0.32	-0.31	0.38	-0.12	0.39	-0.10	-0.03	-0.50	0.04	0.35	0.91	1.00	-0.55
P/A	-0.20	0.24	-0.27	0.54	-0.40	0.60	0.03	0.53	-0.40	-0.52	-0.49	-0.55	1.00

1143 Note: electrical conductance (EC) is expressed in μS/cm, temperature (T) in °C, dissolved ions (Ca²⁺,
1144 Mg²⁺, K⁺, Na⁺, Cl⁻, SO₄²⁻, NO₂⁻-NO₃⁻, NH₄⁺) in ppm, water stable isotopes (δ¹⁸O, δ²H) and deuterium excess
1145 (*d*) in V-SMOW, total carbon (TC), dissolved organic carbon (DOC), dissolved inorganic carbon (DIC)
1146 and total nitrogen (TN) in ppm, elevation (H), lake maximum depth (Z) and perimeter (P) in m, maximum
1147 lake watershed slope (S) in % and area (A) in ha.
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Tab. A2. Results of the **Wilcoxon signed-rank test** of individual conservative and non-conservative hydrological tracers using the 2-class lake typology as the categorical variable. Non-significant *p*-values are indicated in bold. Lakes in the 'recharge' zone are defined as lakes above an elevation of 282 m asl, whereas lakes in the 'discharge' zone are defined as lakes located at an elevation of less than 282 m asl.

Variable	Lakes in the recharge zone	Lakes in the discharge zone	Wilcoxon signed-rank test	
	Mean (SD)	Mean (SD)	V value	p-value
EC (µS/cm)	47.6 (53.5)	230.4 (88.4)	1275	<i>p</i> < 0.001
pH	6.7 (0.9)	7.9 (0.4)	1275	<i>p</i> < 0.001
T (°C)	17.0 (1.9)	16.5 (2.6)	1275	<i>p</i> < 0.001
Ca ²⁺ (ppm)	4.8 (6.6)	29.4 (10.7)	2138	<i>p</i> < 0.001
Mg ²⁺ (ppm)	1.1 (1.4)	7.1 (3.1)	1013	<i>p</i> < 0.001
K ⁺ (ppm)	0.4 (0.2)	1.0 (0.4)	699	0.559
Na ⁺ (ppm)	0.8 (1.9)	5.3 (7.9)	867	0.027
Cl ⁻ (ppm)	1.1 (3.9)	9.5 (15.0)	1572	0.024
SO ₄ ²⁻ (ppm)	0.9 (0.8)	6.6 (13.9)	1003	<i>p</i> < 0.001
NO ₂ ⁻ -NO ₃ ⁻ (ppb)	8.3 (25.0)	56.5 (297.6)	638	<i>p</i> < 0.001
NH ₄ ⁺ (ppb)	22.5 (6.4)	31.8 (28.6)	231	<i>p</i> < 0.001
δ ¹⁸ O (‰)	- 8.7 (1.4)	- 11.5 (1.6)	0	<i>p</i> < 0.001
δ ² H (‰)	- 78.2 (7.1)	- 90.6 (7.6)	0	<i>p</i> < 0.001
<i>d</i> (‰)	- 8.5 (4.5)	1.7 (5.5)	234	<i>p</i> < 0.001
E/I or f	0.6 (0.2)	0.2 (0.3)	1406	<i>p</i> < 0.001
TC (ppm)	11.4 (5.9)	30.4 (8.7)	666	<i>p</i> < 0.001
DOC (ppm)	5.9 (3.2)	6.2 (4.9)	666	<i>p</i> < 0.001
DIC (ppm)	5.5 (6.4)	24.2 (10.6)	653	<i>p</i> < 0.001
TN (ppm)	0.5 (0.2)	0.5 (0.3)	217	0.069
Atomic C/N	13.2 (3.3)	13.6 (4.0)	666	<i>p</i> < 0.001
Elevation (m)	287.7 (4.8)	275.1 (2.8)	1275	<i>p</i> < 0.001
Maximum depth (m)	12.4 (6.2)	13.2 (6.7)	1750	<i>p</i> < 0.001
Maximum lake watershed slope (%)	9.8 (6.9)	20.5 (5.7)	1273	<i>p</i> < 0.001

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1173 **Tab. A3:** Results of the **Wilcoxon signed-rank test** of conservative and non-conservative hydrological
 1174 tracers using the 3 class lake typology as the categorical variable. Non-significant *p*-values are indicated
 1175 in bold. Lakes in the 'recharge' zone are defined as lakes above an elevation of 282 m asl and non-
 1176 alkaline, lakes in the 'seepage' zone are defined as lakes above an elevation of 282 m asl and alkaline
 1177 whereas lakes in the 'discharge' zone are defined as lakes located at an elevation of less than 282 m asl.

Variable	Recharge lakes	Seepage lakes	Discharge lakes	Wilcoxon signed-rank test	
	Mean (SD)	Mean (SD)	Mean (SD)	V value	p-value
EC (µS/cm)	20.5 (11.9)	135.9 (60.8)	233.2 (92.5)	1275	<i>p</i> < 0.001
pH	6.2 (0.7)	7.8 (0.3)	7.9 (0.4)	1275	<i>p</i> < 0.001
T (°C)	17.1 (2.0)	16.4 (1.7)	16.5 (2.8)	1275	<i>p</i> < 0.001
Ca ²⁺ (ppm)	1.4 (1.4)	16.8 (8.6)	29.5 (11.2)	1705	<i>p</i> = 0.002
Mg ²⁺ (ppm)	0.4 (0.3)	3.5 (1.7)	7.2 (3.3)	927	0.005
K ⁺ (ppm)	0.5 (0.3)	0.4 (0.1)	1.0 (0.4)	3	<i>p</i> < 0.001
Na ⁺ (ppm)	0.4 (0.7)	1.9 (2.9)	5.6 (8.3)	876	0.009
Cl ⁻ (ppm)	0.4 (1.3)	3.3 (6.5)	9.9 (15.7)	832	0.004
SO ₄ ²⁻ (ppm)	0.7 (0.6)	1.7 (1.1)	7.0 (14.6)	486	0.145
NO ₂ ⁻ -NO ₃ ⁻ (ppb)	78.0 (347.3)	0.0 (0.0)	8.8 (25.6)	1	<i>p</i> < 0.001
NH ₄ ⁺ (ppb)	33.9 (32.9)	25.4 (0.0)	22.6 (6.5)	0	<i>p</i> < 0.001
δ ¹⁸ O (‰)	-8.5 (1.3)	-9.7 (1.6)	-11.6 (1.7)	0	<i>p</i> < 0.001
δ ² H (‰)	-77.4 (6.8)	-82.1 (7.8)	-90.8 (8.0)	0	<i>p</i> < 0.001
<i>d</i> (‰)	-9.6 (3.7)	-4.5 (5.4)	1.9 (5.7)	129	<i>p</i> < 0.001
E/I or f	0.7 (0.3)	0.6 (0.2)	0.2 (0.2)	218	<i>p</i> < 0.001
TC (ppm)	8.7 (4.0)	19.9 (6.6)	30.8 (9.4)	666	<i>p</i> < 0.001
DOC (ppm)	6.7 (3.5)	3.9 (1.2)	6.7 (5.2)	649	<i>p</i> < 0.001
DIC (ppm)	2.0 (1.3)	15.9 (7.2)	24.1 (11.5)	644	<i>p</i> < 0.001
TN (ppm)	0.6 (0.2)	0.4 (0.1)	0.5 (0.3)	2	<i>p</i> < 0.001
Atomic C/N	14.1 (3.5)	11.1 (1.4)	14.0 (4.3)	666	<i>p</i> < 0.001
Elevation (m)	287.0 (4.6)	286.7 (7.6)	275.1 (3.0)	1275	<i>p</i> < 0.001
Maximum depth (m)	12.0 (6.6)	12.4 (5.7)	13.5 (6.8)	1701	<i>p</i> < 0.001
Maximum lake watershed slope (%)	10.2 (7.4)	11.3 (6.7)	19.9 (6.7)	1272	<i>p</i> < 0.001

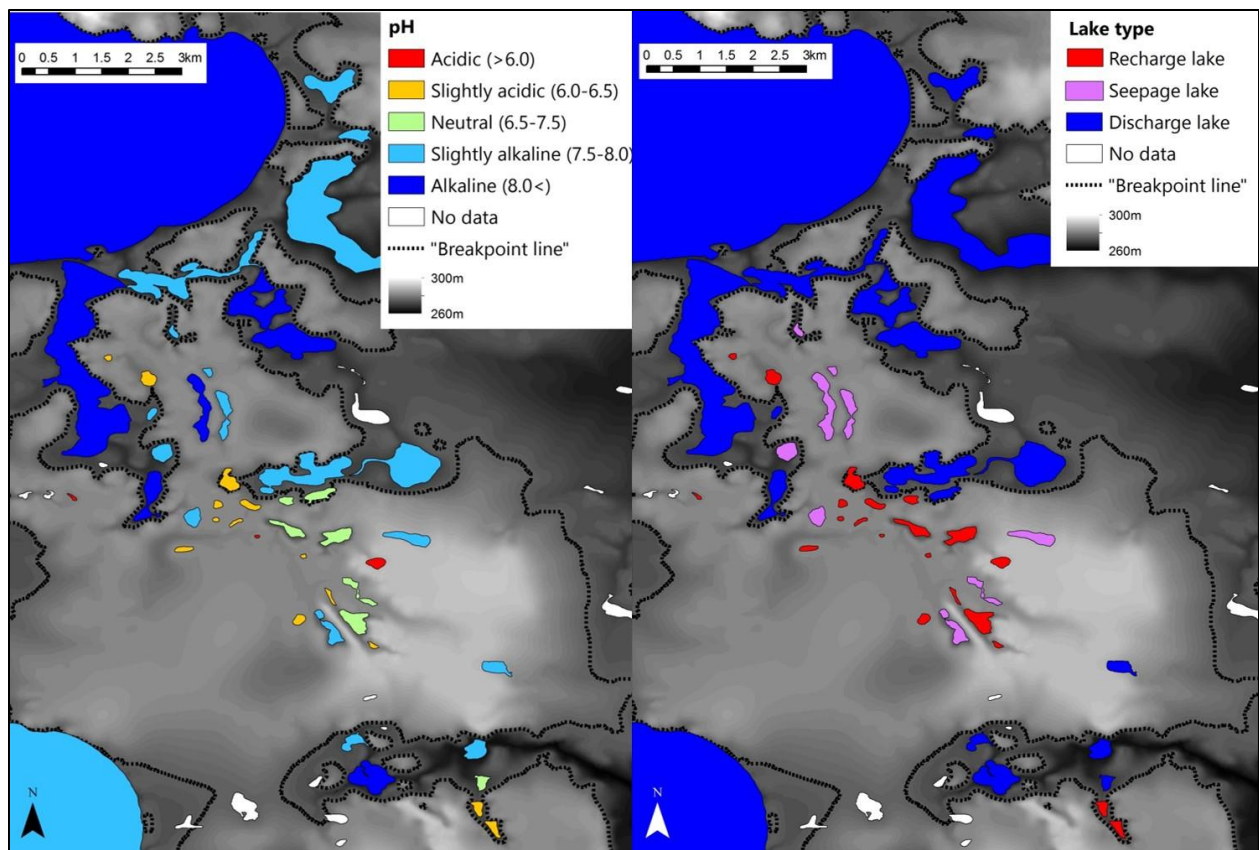
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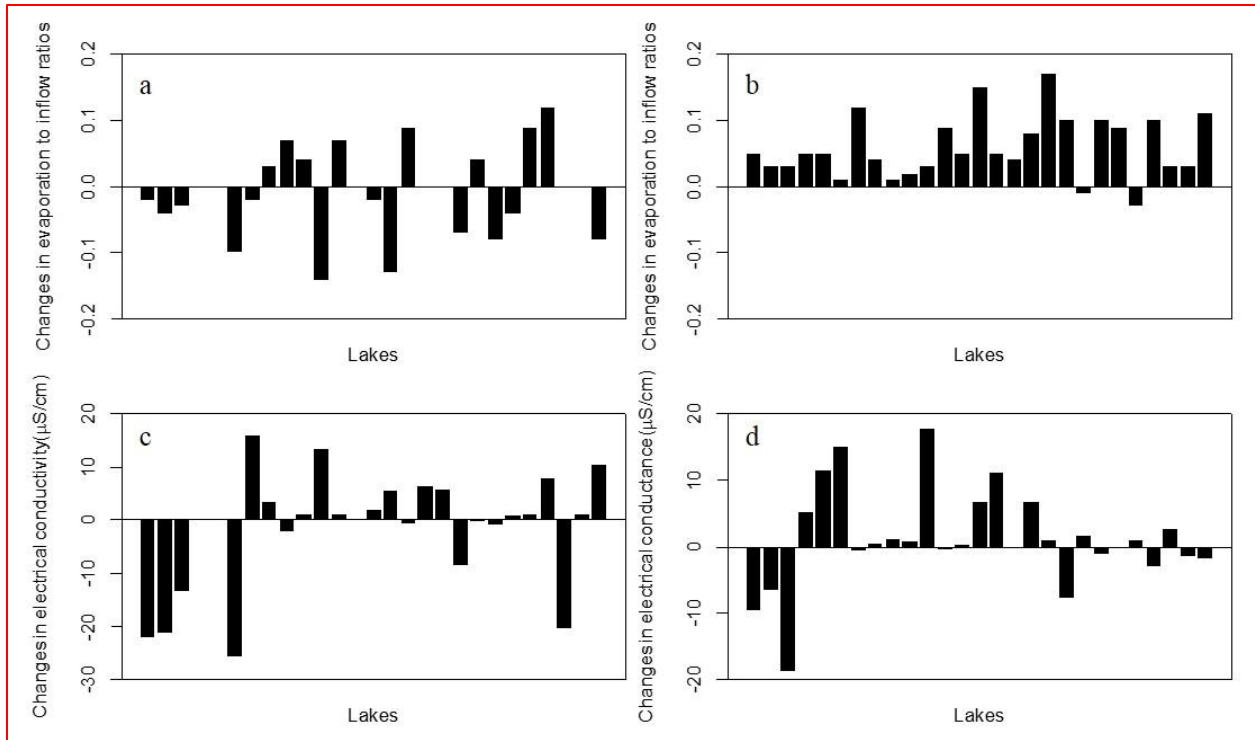
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 1184 **Fig. A1:** Spatial depiction between elevation and lake-water pH (a) and lake type (b). The elevation of the
 1185 breakpoint line (284 m) is shown as a dashed line

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Fig. A2: Changes in E/I or f between June 2013 and June 2014 (E/I or f 2014 – E/I or f 2013) (a) and changes in E/I or f between August 2014 and June 2014 (E/I or f Aug – E/I or f Jun) (b), and changes in specific conductance between June 2013 and June 2014 (EC 2014 – EC 2013) (c) and changes in electrical conductance between August 2014 and June 2014 (EC Aug – EC Jun) (d). Lakes are organized along an elevation gradient (lower to higher elevation).