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1 **Ecosystem Shifts at two Mid-Holocene Tipping Points in the alpine Lake Son Kol**
2 **(Kyrgyzstan, Central Asia)**

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12 **Abstract**

13 Tipping points can be defined as critical ecosystem thresholds that start self-enforced
14 dynamics pushing systems into new stable states. Many lake ecosystems of arid Central
15 Asia are sensitive to hydrological changes as they are located at the intersection of the
16 influence of the dry Siberian Anticyclone and the relatively humid mid-latitude Westerlies,
17 and their sediment records can be used to study past tipping points. We studied subfossil
18 chironomid remains preserved in a ca. 6000-year-long sediment record from the Central
19 Asian lake Son Kol (Central Kyrgyzstan) to reconstruct past ecosystem dynamics. Our
20 results show abrupt transitions from a chironomid fauna dominated by macrophyte-
21 associated, salinity-indicating taxa, to a vegetation-independent fauna, and subsequently to
22 a macrophyte-associated, freshwater-indicating fauna. A comparison of the chironomid-
23 based environmental reconstruction to other proxy indicators from the same record suggests
24 a phase of increased Westerly strength starting about 4900 cal. yr BP. This increase led to
25 enhanced precipitation and sediment fluxes into the lake, which in turn led to high turbidity
26 levels and consequently to a macrophyte collapse causing abrupt changes in the chironomid
27 fauna. At 4300 cal. yr BP, a weakening of the Westerlies in combination with higher lake
28 levels led to lower turbidity and ultimately to the recovery of the macrophyte population and
29 associated changes in the chironomid assemblage. These two sequences of events show
30 how the occurrence of a gradual change in an external trigger (Westerlies) can trigger a
31 cascade of within-lake processes (turbidity, macrophyte density) and may ultimately lead to
32 an abrupt reorganisation of the ecosystem (chironomid fauna), providing models for tipping
33 points.

34

35 **Keywords:** Holocene, Chironomids, Westerlies, Paleoecology, Macrophytes

36

37 Introduction

38 Originating in social sciences (Grodzins, 1958), the term “tipping point” has been used in
39 ecological discussions to describe the situation when ecosystem changes reach a threshold
40 from which self-enforcing dynamics lead the system into a new stable state (van Nes et al.,
41 2016). In lakes, these regime shifts are particularly common when climatic or anthropogenic
42 influence causes major hydrological and/or ecological changes (Andersen et al., 2009; Lees
43 et al., 2006; Randsalu-Wendrup et al., 2014, 2016). Understanding tipping point dynamics is
44 of particular interest for the future, because many systems will reach such thresholds under
45 continuing climate change (Lenton et al., 2008). In order to clearly describe the processes
46 around the tipping points, we differentiated between triggers, actors pushing the ecosystem
47 to the threshold, and drivers, actors of the self-enforcing feedback-loop.

48 Regions characterized by the limitation of an important climatic factor, such as frost-free
49 days or precipitation are particularly sensitive to changes in climate and therefore excellent
50 for investigating tipping points. For many ecosystems in arid Central Asia, changes in
51 precipitation and water availability are of crucial importance. As precipitation is mainly
52 provided by the mid-latitude Westerlies (Aizen et al., 1997, 2001), even minor fluctuations of
53 their interplay with dry air masses from the Siberian High can lead to major differences in the
54 water budget. Therefore, reconstructing ecosystem dynamics of arid Central Asia plays a
55 key role in understanding the dynamics of these two atmospheric circulation systems (Cheng
56 et al., 2012) as well as possible responses of freshwater ecosystems to climate forcing. The
57 central Kyrgyz lake Son Kol is situated in the transition zone between the two wind systems
58 (Aizen et al., 1997, 2001), and therefore provides an excellent opportunity to study their past
59 interactions.

60 Lake sediments typically contain a range of proxies, such as diatoms, ostracods and
61 chironomids (Alivernini et al., 2018; Battarbee, 2000; Brooks 2006; Schwalb et al. 1998), that
62 can be studied to reconstruct ecosystem development (Beer et al., 2007; Mischke and
63 Wünnemann, 2006; Schwarz et al., 2017). Previous studies from Son Kol reported the
64 development from a saline to a freshwater lake including major hydrological changes caused
65 by moister climate conditions ca. 5000–4500 cal. yr BP (Huang et al., 2014; Lauterbach et
66 al., 2014; Mathis et al., 2014; Paction et al., 2014; Schwarz et al., 2017). The sediment core
67 examined in this study was first investigated by Lauterbach et al. (2014) using
68 sedimentological, (bio)geochemical, isotopic, and palynological evidence to reconstruct
69 phases of enhanced allochthonous sediment input through snowmelt, reflecting increased
70 winter snowfall brought by episodically intensified Westerlies in Central Kyrgyzstan during
71 the last 6000 years. For the same record, Schwarz et al. (2017) combined diatom, ostracod
72 and stable isotope data derived from the Son Kol sediment record and identified climate-
73 driven organism responses. Both studies showed ecosystem-wide abrupt shifts at ca. 5000
74 and 4300 cal. yr BP. Diatoms, however, were occasionally poorly or not continuously
75 preserved in the sediment, especially from 4900 to 4300 cal. yr BP, therefore leaving final
76 conclusions about internal processes affecting the ecosystem open.

77 Subfossil remains of chironomid larvae (Diptera: Chironomidae) are preserved in lake
78 sediments due to their high chitin concentration. They can be especially valuable where
79 other bioindicators such as diatoms are poorly preserved due to alkaline conditions (Brooks,
80 2006; Caballero et al., 2003; Flower, 1993). They have been widely used to reconstruct past

81 environmental conditions, including summer air temperatures (Brooks and Birks, 2000;
82 Zhang et al., 2017), water depth (Engels et al., 2012), salinity (Chen et al., 2009) and pH
83 (Charles et al., 1987; Plank, 2010). Furthermore, the association of different chironomid taxa
84 with specific habitats, such as submerged macrophyte vegetation, can give an insight into
85 ecosystem changes beyond the physicochemical conditions (Motta and Massaferrò, 2019).

86 In this paper, we present the results of detailed chironomid analysis applied to the Holocene
87 record of Son Kol. Combining our new results with existing data on past environmental
88 change at the site (Lauterbach et al., 2014; Schwarz et al., 2017) allows us to study
89 ecosystem dynamics across the past 6000 years. We specifically aim to (1) reveal
90 ecosystem shifts, (2) identify the cascade which generated the observed abrupt shifts,
91 including both triggers and drivers, as well as (3) define tipping points and distinguish them
92 from other changes

93 **Study Site**

94 Son Kol (41° 50' N, 75° 10' E) is the second largest lake of Kyrgyzstan, with a surface area
95 of ca. 273 km² (Fig. 1). The relatively shallow lake (max. water depth ca. 13 m) is situated at
96 3016 m a.s.l. in the central Tian Shan. It is surrounded by mountain ranges peaking at 3800-
97 4000 m a.s.l., forming a ca. 1130 km² catchment area (Academy of Science of the Kyrgyz
98 SSR, 1987; Shnitnikov, 1980). Catchment geology is composed of Cambro-Ordovician
99 granitoids, carboniferous sedimentary rocks and Permian granitoids, sedimentary rocks and
100 tuffs. The plains around the lake consist of eroded Quaternary sediments (De Grave et al.,
101 2011). The main water sources are precipitation, snowmelt runoff and groundwater inflow.
102 The lake is drained by one outlet at its eastern end (Lauterbach et al., 2014).

103 Short temperate summers and long, cold winters with snow cover from November to April
104 characterize the local, high-alpine, continental climate. The average annual temperature is
105 -3.5 °C (average in January -20 °C, in July 10 °C), providing ice cover from October to late
106 April (Academy of Science of the Kyrgyz SSR, 1987; Lauterbach et al., 2014; Schwarz et al.,
107 2017; Shnitnikov, 1980). Due to the shallow water depth and strong winds, Son Kol is
108 polymictic and well oxygenated throughout the year, featuring water temperatures of 0–2 °C
109 in winter and ca. 16 °C in summer (Lauterbach et al., 2014). Today it is a freshwater lake
110 with conductivities ranging between 0.515 and 0.530 mS cm⁻¹ (Schwarz et al., 2017).

111 The Siberian Anticyclone in winter and the mid-latitude Westerlies in summer are the wind
112 systems that control the regional climate. Because the Westerlies are the main water source,
113 the annual precipitation of 500-600 mm is seasonally distributed, with only 20 % falling in
114 winter (Academy of Science of the Kyrgyz SSR, 1987; Aizen et al., 1997, 2001). The Son
115 Kol region is today located north of the monsoon influenced area and likely has been since
116 the mid-Holocene (Cheng et al., 2012; Winkler and Wang, 1993). The vegetation
117 surrounding the lake is characterized by alpine meadows, steppe-desert landscapes (Mathis
118 et al., 2014) as well as sedge marsh at the lake shores, followed by submerged
119 macrophytes, eg *Myriophyllum*, and algae on the lake bottom down to water depths of at
120 least 7–8 m (Lauterbach et al., 2014).

121 **[Insert Fig. 1]**

122

123

124 **Methods**

125 Two sediment cores of 121.5 cm (SONK_11_D1) and 166.0 cm (SONK_11_D2) length were
126 retrieved in the summer of 2011 from the south-eastern part of Son Kol (41°47'38"N,
127 75°11'49"E, 10.5 m water depth; Fig. 1), using an UWITEC gravity corer. The two sediment
128 cores were correlated using distinct lithological marker layers, resulting in a continuous
129 173.5 cm long composite sediment record (SONK_11_D1/2; Lauterbach et al., 2014). To
130 establish a chronology for the composite sediment record, 28 samples of terrestrial plant
131 macrofossil remains, bivalve shells, and bulk sediment were dated by accelerator mass
132 spectrometry (AMS) ¹⁴C dating at the Poznań Radiocarbon Laboratory. The final age-depth
133 model for the composite sediment core was established by Bayesian age modelling using a
134 *P Sequence* depositional model implemented in OxCal 4.1 (Bronk Ramsey, 2008, 2009). As
135 model input parameters we used 22 of the obtained (AMS ¹⁴C dates, which were calibrated
136 with the IntCal09 calibration data set (Reimer et al., 2009), as well as age information from
137 ¹³⁷Cs and ²⁴¹Am activity measurements obtained from the uppermost 10 cm of the composite
138 sediment record. According to the final age-depth model, the sediment sequence covers the
139 last 6000 years. A detailed description of the sedimentology of the composite core, as well
140 as comprehensive information about AMS ¹⁴C dating and age modelling, is provided by
141 Lauterbach et al. (2014).

142 In total, 118 samples were analysed for chironomid remains: 32 one-cm-thick samples were
143 collected from core SONK_11_D1 (12-98 cm composite depth); another 86 samples, each
144 0.5 cm thick, were retrieved from core SONK_11_D2 (99-173.5 cm composite depth).
145 Sample depths are given as middle depth in addition to the core abbreviation 11_D1/11_D2.
146 A minimum of 1 g sediment was analysed, with the exception of one sample (11-D2, 140.5
147 cm composite depth) that did not provide sufficient sediment. Sample preparation followed
148 Brooks et al. (2007). After solution in 10 % KOH and 20 minutes of heating up to 85 °C, the
149 sediment was rinsed through a 100 µm mesh sieve. Chironomidae head capsules (HC) as
150 well as Cladocera ephippia and Charophyta oogonia were hand-picked at 32 x magnification
151 and mounted in Euparal® mounting medium. Identification under 400 x magnification
152 followed Brooks et al. (2007), Rieradevall and Brooks (2001), and Bitušík and Hamerlík
153 (2014). The water content of samples was determined using freeze drying (two
154 measurements per sample). Dry weights of samples were subsequently used to determine
155 chironomid concentrations. Chironomid head capsules were hand-picked from freeze-dried
156 as well as from wet sediments and reported concentrations are based on the total number of
157 head capsules and the corresponding calculated dry weight. The datasets produced in this
158 study are uploaded to the data repository Pangaea
159 (<https://doi.pangaea.de/10.1594/PANGAEA.908301>).

160 If insufficient material was available to recover 50 head capsules (Heiri and Lotter, 2001;
161 Quinlan and Smol, 2001), adjacent samples were merged before further analyses. However,
162 even after merging it was not always possible to obtain a minimum count sum of 50 head
163 capsules. In the section from 90 to 120 cm composite depth, samples were included in
164 further analysis if the merged amount reached 40 head capsules. Below 130 cm composite
165 depth, samples with a head capsule count of at least 25 were accepted. Because of the
166 extremely low diversity, these samples were used despite the low count sum. Head capsules
167 of the genus *Chironomus* that missed parts of the lateral teeth necessary for further
168 identification were assigned to the two species groups *Chironomus anthracinus*-type and

169 *Chironomus plumosus*-type using the ratio of the identified head capsules prior to statistical
170 analysis. Head capsules of the genus *Tanytarsus* that missed mandibles, but possessed an
171 antennal pedastel without spur were listed as *Tanytarsus* “without spur”. Head capsules of
172 *Tanytarsus* that also missed the antennal pedestal were listed as *Tanytarsus* indet., but
173 included in the taxon *Tanytarsus* “without spur” for graphical representation. After excluding
174 rare taxa (those with less than 2 samples with abundances higher than 2 %), square root
175 transformed percentages of chironomid taxa were used to assign the samples to zones
176 following a cluster analysis (Everitt, 2011). The number of statistically significant zones was
177 determined with a broken-stick model (Bennett, 1996), and zonal boundaries are reported as
178 ages rounded to the nearest century. Principal component analysis was used to summarise
179 the data and to identify the most important taxa (ter Braak, 1983). Analyses were performed
180 with R (R Core Team, 2016) using the package ‘vegan’ (Oksanen et al., 2016). Results were
181 plotted using C2 (Juggins, 2014). We analysed the occurrence of regime shifts using the
182 sequential algorithm presented by Rodionov (2004, 2006). This method allows the statistical
183 detection of significant shifts time series based on a sequential t-test. We ran the algorithm
184 using both the sample scores on PCA axis 1 and 2 as input, and explored a range of
185 settings, in line with recommendations by Rodionov (2005). We here present the results for a
186 test run with a significance level of 0.1, a cut-off length of 14 and a Huber's weight parameter
187 of 1. Regime shifts were accepted as significant with a rate of change index of at least 0.95.

188

189 **Results**

190 A total of 118 samples were analysed, containing on average 34.6 chironomid head
191 capsules (25.05 HC g⁻¹ dry weight). Out of these, 22 samples were excluded from further
192 analyses as they showed very low count sums. An additional 58 samples were merged into
193 13 samples to reach count sums suitable for further data exploration (Tab. S1). Fifty samples
194 remained after this data processing procedure, of which seven samples contained less than
195 50 head capsules. No samples reached sufficient head capsule counts below 150 cm
196 composite depth. We identified a total of 15 taxa in the record of Son Kol (Fig. 2). Five taxa
197 were identified as being rare.

198 The chironomid record was divided into four zones, covering the intervals 150–124, 124–
199 108, 108–92.5 and 92.5–12 cm, respectively (Fig. 2, Fig. S1). These correspond to the time
200 windows of 6000-4900, 4900-4300, 4300-3700 and 3700-500 cal. yr BP, respectively. The
201 lower two boundaries were statistically significant when compared to the a broken-stick
202 model (Fig. S2). The third boundary was visually established based on minor changes in the
203 chironomid record.

204

205 **Zone I: 6000–4900 cal. yr BP (150–124 cm)**

206 *Psectrocladius sordidellus*-type and *Cricotopus intersectus*-type dominate the assemblages
207 of Zone I. The zone is characterized by the highest fluctuations along the core, both seen in
208 between-sample fluctuations of dominant taxa, as well as in the head capsule concentration
209 record, which ranges from samples void of chironomids to 304 HC/g dry sediment, the
210 highest concentration of the record (Fig. 4). Charophyte oogonia and *Daphnia* ephippia
211 occur (0.2-132.3 and 0.4-3.4 g dry weight⁻¹, respectively) in this zone. The sample scores on
212 both PCA axis 1 and 2 are the lowest of the record and show a high variability.

213 **Zone II: 4900–4300 cal. yr BP (124–108 cm)**

214 In the second zone, the chironomid fauna is dominated by *Chironomus anthracinus*-type.
215 *Psectrocladius sordidellus*-type and *Cricotopus intersectus*-type, the two taxa that were
216 abundant during Zone I, disappeared almost completely at the onset of Zone II. Chironomid
217 concentrations show only minor fluctuations, and generally low values during Zone II. No
218 charophyte oogonia and only a single *Daphnia* ephippium were found in this zone. The
219 scores of the samples on PCA axis 1 start a continuous increase in Zone II, while the scores
220 on PCA axis 2 increase abruptly to reach the highest values of the record at the beginning of
221 Zone II. With the exception of a slight dip around 4650 cal. yr BP, they remain high until an
222 abrupt decrease occurs at the end of the zone.

223 **Zone III: 4300–3700 cal. yr BP (108–92.5 cm)**

224 After a second abrupt shift in the chironomid assemblages at ca. 4300 cal. yr BP *Tanytarsus*
225 “without spur”, *Tanytarsus gracilentus*-type and *Procladius* replaced *Chironomus*
226 *anthracinus*-type. *Procladius* is the dominant taxon during the first part of Zone III, but shows
227 declining abundances with time, whereas *Tanytarsus* “without spur” shows increasing
228 abundances, becoming the dominant taxon. Chironomid concentrations, while showing only
229 minor fluctuations, remain low at the zonal boundary, but start to increase at ca.
230 4000 cal. yr BP. While charophyte oogonia are still absent, *Daphnia* ephippia are present
231 (0.1–0.7 g dry sediment⁻¹) throughout Zone III. The sample scores on PCA axis 1 continue to
232 increase evenly, while the scores on PCA axis 2 are stable with values ranging between
233 those of Zone I and Zone II.

234 **Zone IV: 3700–500 cal. yr BP (92.5–12 cm)**

235 The uppermost zone represents the longest and most stable zone of the record, showing
236 dominance of *Tanytarsus gracilentus*-type and the more general morphotype *Tanytarsus*
237 “without spur”, accompanied by low percentages of *Procladius*. Chironomid concentrations
238 continue to increase across the zonal boundary and remain stable above 30 HC g⁻¹ dry
239 sediment after ca. 3500 cal. yr BP. *Daphnia* ephippia were found only in the topmost sample.
240 PCA scores remain at medium values throughout Zone IV.

241 Most of the variance of the chironomid data is explained by the first two PCA axes (in total
242 86.4 %), and three clusters of samples can be identified in a bi-plot showing PCA axis 1
243 and 2 (Fig. 3). *Tanytarsus gracilentus*-type and *Tanytarsus* “without spur” are most strongly
244 associated with PCA axis 1 (eigenvalue of 26.3, 58.1 % variance explained), while
245 *Psectrocladius sordidellus*-type, *Cricotopus intersectus*-type and *Chironomus anthracinus*-
246 type are most strongly associated with PCA axis 2 (eigenvalue of 12.8, 28.3 % variance
247 explained). Regime shift analysis revealed significant shifts at 4900, 4500–4300 and
248 3700 cal. yr BP, respectively. The exact timing of the shift at 4500–4300 cal. yr BP varied
249 between the two PCA axes (4500 cal. yr BP in the analysis using the axis 1 scores,
250 4300 cal. yr BP in the analysis using axis 2 scores) and was only significant in PCA axis 1.

251 **[Insert Fig. 2]**

252 **[Insert Fig. 3]**

253 **[Insert Fig. 4]**

254

255 **Discussion**

256 **Mid to late Holocene lake development**

257 Son Kol developed from a closed, saline lake at 6000 cal. yr BP to the modern open,
258 freshwater lake (Huang et al., 2014; Schwarz et al., 2017). Our chironomid record shows
259 four zones during this time-interval (> 4900, 4900–4300, 4300–3700 and < 3700 cal. yr BP).
260 These zones are similar to the four major zones previously recognized by Schwarz et al.
261 (2017), who analysed diatoms, ostracods and stable isotope analysis in the same sediment
262 sequence. The difference between the exact timings of the zonal boundaries established in
263 this study and those by Schwarz et al. (2017) is only minor (up to 150 years), and is
264 potentially due to differences in sampling density or proxy-response times.

265 **Zone I (> 4900 cal. yr BP)**

266 *Psectrocladius sordidellus*-type and *Cricotopus intersectus*-type were the most abundant
267 taxa in the first zone. Both these Orthoclaadiinae morphotypes include species adapted to
268 saline conditions (Plank, 2010; Zhang et al., 2007). Additionally, head capsule concentration
269 showed high fluctuations during Zone I. If lake levels were lower than today, the coring site
270 would have been closer to the shore and therefore more vulnerable to changing conditions
271 such as river influence connected to differences in the sediment regime, thus potentially
272 explaining the high variations observed in the chironomid concentrations. We therefore
273 suggest that Son Kol was a shallow, saline lake during Zone I.

274 This chironomid-based palaeoenvironmental inference is in agreement with Schwarz et al.
275 (2017) who infer a shallow, saline environment based on high conductivities reconstructed
276 using a diatom-conductivity-transfer-function (8.2 mS cm^{-1}) as well as the dominance of the
277 halophile ostracod species *Eucypris mareotica* (Fig. 4). The occurrences of single freshwater
278 diatom valves opposing the otherwise dominating salinity indicators was interpreted as a
279 result of freshwater inflow (Schwarz et al., 2017) which matches our interpretation of a
280 shorter distance to the shore and thus higher influence of the tributaries.

281 **Zone II (4900–4300 cal. yr BP)**

282 Both cluster analysis and regime shift analysis show the occurrence of a statistically
283 significant boundary at 4900 cal. yr BP. This result is consistent between regime shift
284 analyses using both PCA axis 1 and axis 2 scores, and tests run using different statistical
285 settings. The onset of Zone II at 4900 cal. yr BP is characterised by an abrupt increase in
286 *Chironomus anthracinus*-type, a chironomid morphotype which can occur under a range of
287 environmental conditions. Due to the large range of habitat conditions suitable for
288 *C. anthracinus*-type, it is hard to provide an ecological interpretation of the environmental
289 conditions during Zone II based on the chironomid data alone. A chironomid-based
290 interpretation of palaeoenvironmental conditions during Zone II is further hampered by the
291 fact that the generalist genus *Procladius* (Vallenduuk and Moller Pillot, 2013) is the only
292 other chironomid taxon that occurred in abundances over 10 % during Zone II. The
293 comparison of the chironomid fauna of Zone II to the other zones shows striking differences
294 in the habitat preferences of the dominant taxa. In contrast to the sediment-bound *C.*
295 *anthracinus*-type that was dominant during Zone II (Moller Pillot, 2013a), the morphotypes
296 (*Cricotopus intersectus*-type, *Psectrocladius sordidellus*-type and *Tanytarsus gracilentus*-

297 type) that were abundant during Zones I and III are all associated with macrophyte
298 vegetation (Brodersen et al., 2001; Ives et al., 2008; Lindegaard et al., 1979; Moller Pillot,
299 2013b). We suggest that the disappearance of these macrophyte-associated taxa in favour
300 of the sediment-bound *C. anthracinus*-type that (Moller Pillot, 2013a) at the onset of Zone II
301 could have been caused by a decline of the aquatic macrophyte vegetation. Alternatively,
302 the strong could be explained by other factors such as low oxygen conditions (Moller Pillot,
303 2013a), eutrophication (Meriläinen et al., 2000), or heavy metal pollution (Mocq et al., 2018).
304 However, such conditions are unlikely to have occurred in the relatively shallow and
305 oligotrophic Son Kol, which even at present is not under high anthropogenic pressure.
306 Furthermore, a high pH resulting from e.g. increased fluvial input of carbonate rocks from the
307 catchment could also have potentially led to high abundances of *C. anthracinus*-type (Moller
308 Pillot, 2013a). However, as *Tanytarsus gracilentus*-type, the chironomid taxon dominating
309 the following zones, is also adapted to high pH values, a shift to higher pH values is unlikely
310 to be the only driver of the abrupt ecosystem shifts observed at the start and end of Zone II.

311 The ostracod assemblages were dominated by *Limnocythere inopinata*, a pH-independent
312 generalist as well (Caballero et al., 2003; Kulköylüoğlu, 2005), and a lack of sufficient
313 preserved diatom remains only confirmed conditions harsh for aquatic life, but did not
314 provide further insight into the environmental conditions at that time (Schwarz et al., 2017;
315 Fig. 4). However, some information can be gained from the geochemical analyses by
316 Lauterbach et al. (2014), in particular from the total organic carbon (TOC) content, indicating
317 the lowest productivity of the entire record during this interval, and the $\delta^{15}\text{N}$ values, which
318 showed the highest values during this zone (Fig. 4). While there are several reasons for high
319 $\delta^{15}\text{N}$ values in general, eg denitrification processes in an anoxic hypolimnion, increased
320 aquatic production or high evaporation, increased input of terrestrial organic material,
321 brought by fluvial input due to increased precipitation, hence increased Westerly strength,
322 was proposed as the only plausible cause for Son Kol in this case (Lauterbach et al., 2014).
323 The sediment input, combined with high wind velocities during this phase of increased
324 Westerly strength, might have resulted in high turbidity, which in turn could explain the
325 inferred low lake-internal productivity. High turbidity could also explain the low diatom
326 concentrations during Zone II found by Schwarz et al. (2017), because the low availability of
327 sunlight could have led to low diatom production and the dispersed sediment particles might,
328 in particular under high pH, have damaged the diatom valves. Increased turbidity could have
329 caused a decline of macrophyte vegetation and could therefore explain the dominance of
330 *C. anthracinus*-type in Zone II (Blindow et al., 2002; Ibelings et al., 2007) and would further
331 explain the low TOC values in Zone II.

332 We therefore argue that increased turbidity between 4900 and 4300 cal. yr BP could have
333 hampered macrophyte growth, which might have further increased turbidity (eg through
334 wave-driven mobilization of now more unconsolidated lake bottom sediments) until this self-
335 enforcing process led to a collapse of macrophyte vegetation. Scheffer (1993) showed, even
336 though driven by eutrophication instead of sediment input and strong wind velocities, how
337 the interplay of turbidity and macrophyte growth can lead to abrupt shifts between turbid and
338 clear stable states. Indicators for the clear state were in particular Charophyta and
339 Cladocera (Scheffer et al., 1993, 2003; van den Berg et al., 1998), whose remains were
340 concordantly found in Zone I, but were absent in Zone II (Fig. 4). The observed delay in the
341 shift of bioindicators compared to that in $\delta^{15}\text{N}$ supports the hypothesis of alternative stable

342 states, because the ecosystem pressure applied by fluvial input and high wind speeds
343 needed to add up to trigger the cascade which then abruptly changed the ecosystem state.
344 The opposing effect of decreasing Westerly strength triggering the cascade of decreasing
345 turbidity followed by macrophyte recolonization leading to even lower turbidity (Ibelings et al.,
346 2007; Scheffer et al., 2003; van Nes et al., 2016) might have driven the second abrupt
347 change.

348 Especially noticeable is the comparison of TOC and $\delta^{15}\text{N}$ to the scores of the chironomid
349 based PCA axis 2. The pattern of all three graphs, even though reversed for TOC,
350 throughout Zone II is very similar, showing an abrupt increase, a dip in the middle of the
351 zone, and an abrupt decrease, the timing of increase and decrease however, is delayed in
352 the PCA axis 2 (Fig. 4). We interpret the difference between the timing of allochthonous
353 input indicated by the $\delta^{15}\text{N}$ values, and the changes seen in the bioindicators to reflect a
354 delayed response of the lake ecosystem to the change in strength of the Westerlies.

355 **Zone III (4300–3700 cal. yr BP)**

356 The zonal boundary at 4300 cal. yr BP is statistically significant when compared to a broken
357 stick model, and regime shift analysis furthermore suggests the occurrence of an abrupt shift
358 between 4500 and 4300 cal yr BP, although the exact timing of the shift varies between
359 different model runs. The onset of Zone III is characterised by a decrease in *C. anthracinus*-
360 type in favour of *Procladius* and *Tanytarsus gracilentus*-type. Both these chironomid taxa
361 indicate mesohaline conditions (Plank 2010), and the abundant presence of the free
362 swimming genus *Procladius* (Vallenduuk and Moller Pillot, 2013) and the macrophyte-
363 associated *T. gracilentus*-type (Ives et al., 2008; Lindegaard et al., 1979) during Zone III
364 could reflect a re-established clear water state with abundant macrophyte vegetation.
365 Zone III is furthermore characterized by low chironomid head capsule concentrations, which
366 increased throughout the zone. The increase in head capsule concentrations is
367 accompanied by an increase in the relative abundance of *Tanytarsus gracilentus*-type. We
368 therefore interpret this increase in concentrations as a result of increasing productivity of
369 *T. gracilentus*-type rather than as a decrease of productivity of *Procladius*. This development
370 might have been connected to a stabilization of the conditions connected to the opening of
371 the lake system.

372 The chironomid-based inference of a shift toward less saline conditions is in line with the
373 diatom-based conductivity reconstruction (1.5 mS cm^{-1}) and the dominance of the ostracod
374 species *Candona neglecta* (Schwarz et al., 2017, Fig. 4). The reduced salinity can be
375 interpreted as a result of a lake level rise during the phase of increased Westerly strength.

376 **Zone IV (< 3700 cal. yr BP)**

377 The zonal boundary between chironomid Zones III and IV is not statistically significant, but
378 was visually established to highlight the minor changes observed in the chironomid diagram
379 around 3700 cal. yr BP. From this time onward, *Tanytarsus gracilentus*-type strongly
380 dominated the chironomid assemblages (average abundances 92.4 %). Abundances of *T.*
381 *gracilentus*-type that exceed 90 % are currently only observed in the fresh alkaline (pH of
382 8.1-10) Icelandic lake Mývatn (Ives et al., 2008; Opfergelt et al., 2004). The chironomid
383 results therefore suggest alkaline, freshwater to mesohaline conditions from ca. 3700
384 cal. yr. BP onward.

385 The chironomid-based interpretation is in line with the results by Schwarz et al. (2017) who

386 reconstructed freshwater conditions in a deeper, open lake. The proportion of planktonic
387 diatoms in this zone was very high and its increase around the zone boundary (Fig. 4) was
388 the main signal that allowed the interpretation that the lake level rose and the lake system
389 opened. Whereas our chironomid record shows little variability from 3700 cal. yr BP onward,
390 Schwarz et al. (2017) identified four separate zones based on changes in the diatom and
391 ostracod communities of the lake. We hypothesise that the strong dominance of
392 *T. gracilentus* obscured the registration of environmental changes that affected the diatom
393 and ostracod assemblages.

394 **Tipping points**

395 Many definitions exist concerning the term “tipping point”. We use the definition of an abrupt
396 change triggered by minor changes tipping the system over a threshold where self-enforcing
397 processes lead the system to another stable state (Lenton et al., 2008; van Nes et al., 2016).
398 In our study we found two abrupt changes, both showing shifts to alternative stable system
399 states that lasted for more than 500 years.

400 The main environmental trigger resulting in the first abrupt change at 4900 cal. yr BP was
401 the increased influence of the Westerlies causing increased water and sediment input into
402 the lake as well as high wind velocities. These factors led to increased turbidity followed by
403 macrophyte decline. While such an increased influence of the Westerlies at Son Kol has
404 been found several times during the last 6000 years (Lauterbach et al., 2014; Schwarz et al.,
405 2017), it only once led to an ecosystem-wide abrupt change. A probable explanation is that
406 only in this case the environmental pressure of turbidity was high enough to trigger a decline
407 of macrophyte vegetation extensive enough to tip the system into the self-enforcing cascade
408 of further increasing turbidity leading to the complete collapse of macrophyte vegetation.
409 Alternate lake states coupled to macrophyte vegetation and turbidity have been observed in
410 the context of eutrophication and subsequent ecosystem recovery in Lake Veluwe,
411 Netherlands (Ibelings et al., 2007; Scheffer 1993, 2003), and a similar process under
412 westerly wind induced turbidity is plausible (Fig. 5).

413 The opposite mechanism took place at 4300 cal. yr BP, when the Westerlies influence
414 declined forming an environmental trigger that ultimately led to an abrupt ecosystem change.
415 The lower amounts of precipitation associated with the decreasing influence of Westerlies
416 led to less water and sediment input into the lake and, combined with lower wind velocities,
417 resulted in lower turbidity in the water column. At one point the resulting environmental
418 pressure on the submerged plant vegetation was low enough to allow the recolonization of
419 the lake, tipping the system over the threshold into the self-enforcing cascade of increasing
420 vegetation cover leading to lower turbidity until a clear water state was reached once more.

421 In both cases, we see environmental triggers, namely changes in the intensity of Westerlies,
422 leading to environmental pressure starting self-enforced within-lake cascades driven by
423 turbidity and macrofauna disappearance or expansion. These sequences are in line with the
424 criteria of tipping points as suggested by Lenton et al. (2008) and van Nes et al. (2016). Our
425 interpretation is further supported by the delayed timing of the ecosystem change compared
426 to the shift in $\delta^{15}\text{N}$ -values, because these values display the environmental pressure applied
427 to the system and the delayed response of the ecosystem demonstrates how the pressure
428 had to rise until it crossed the tipping point starting the abrupt ecosystem reaction.

429 **[Insert Fig. 5]**

430 **Comparison against other records**

431 Environmental changes between about 4900 and 4500 cal. yr. BP have been reconstructed
432 for many different Central Asian archives. The underlying conditions characterizing these
433 changes though, are different. While most results indicate cooling, the precipitation signal
434 reveals a more heterogeneous spatio-temporal pattern, strongly depending on the locality. In
435 many Tibetan lakes, the cooling in this interval was associated with dry conditions
436 (Doberschütz et al., 2014; Ma et al., 2019; Morinaga et al., 1993; Shi et al., 2017; Xu et al.,
437 2019a). A similar picture was found in several studies from the Southern Altai (Li et al., 2011;
438 Wang and Zhang, 2019; Xu et al., 2019b), as well as Southern Mongolia (Felauer et al.,
439 2012). In contrast, a shift to wetter conditions was found in studies of the Southern (Zhang et
440 al., 2016, 2018), Western and Northern Altai mountains (Ilyashuk and, Ilyashuk, 2007;
441 Zhang and Feng, 2018) and the Tian Shan (Beer et al., 2007; Huang et al., 2015).

442 The regional differences in precipitation changes during this interval were attributed to the
443 influence of the dominating wind system: under Monsoon influence, the trend is connected to
444 dry conditions, whilst under Westerlies influence it is related to humid conditions (Rao et al.,
445 2019). Our Son Kol reconstruction, which shows a wet phase from 4900 to 4300 cal. y. BP
446 and continuing humid conditions thereafter, fits the model of a dominant influence of the
447 Westerlies (Huang et al., 2014; Mathis et al., 2014; Schwarz et al., 2017).

448 While the time window 4900-4300 cal. yr BP covers both abrupt changes identified in our
449 study, there are not many records from Central Asia that also show an environmental
450 change around 3700 cal. yr BP, where a third, more gradual, transition is observed in the
451 Son Kol record. Han et al. (2019) found evidence for major sand storms in the Tarim Basin
452 ca. 3500 cal. yr BP, connecting it to a southward shift of the Westerlies. While a connection
453 to changes in the Westerlies regime might have partly influence the lake system at this time,
454 we consider local effects, possibly the opening of the lake system, as the more likely cause
455 of the Son Kol ecosystem transition at ca. 3700 cal. yr BP.

456 **Conclusions**

457 We produced a high-resolution palaeoecological and palaeoenvironmental record for Lake
458 Son Kol. The main aim of this study was to assess the occurrence and ecosystem cascades
459 characterizing tipping points as well as their distinction from other environmental changes,
460 and our results show prominent ecosystem shifts at two tipping points (4900 and 4300 cal. yr
461 BP) and one gradual shift (3700 cal. yr BP) in the development of the central Kyrgyzstan
462 lake Son Kol from a saline to a freshwater lake. These results are in line with previous
463 studies on diatoms and ostracods (Schwarz et al., 2017), as well as geochemical analyses
464 (Lauterbach et al., 2014). We postulate that a phase of increased Westerly strength causing
465 increased water and sediment input, in combination with high wind velocities to be the trigger
466 of both ecosystem-wide tipping points. As driver we postulate the self-enforcing cascade of
467 increasing turbidity and declining submerged macrophyte cover at the first and the opposite
468 process at the second tipping point (Fig. 5).

469 These conditions are represented by abrupt changes from the macrophyte-associated
470 salinity indicators *Cricotopus intersectus*-type and *Psectrocladius sordidellus*-type over the
471 sediment-bound *Chironomus anthracinus*-type to the again macrophyte associated alkaline
472 but fresh-mesohaline conditions indicating *Tanytarsus gracilentus*-type. While salinity plays

473 an important role distinguishing the chironomid, ostracod and diatom taxa before 4900 and
474 after 4300 cal. yr BP, it cannot explain the assemblages in-between. The gradual change
475 (3700 cal. yr BP) reflects the transition from a closed to an open lake system and is
476 represented by increasing chironomid head capsule concentrations and *Tanytarsus*
477 *gracilentus*-type abundances and fits the increase of planktonic diatoms recognized by
478 Schwarz et al. (2017).

479 Our study demonstrates in detail how ecosystems can shift in reaction to an external trigger
480 and how self-enforcing cascades can generate ecosystem-wide tipping points.

481

482

483 **Acknowledgements**

484 We thank Martina Stebich for data contribution and discussion. We thank two anonymous
485 reviewers and editor professor Vivienne Jones for their constructive and critical reviews of
486 the manuscript that helped to improve the manuscript.

487 **Declaration of conflicting interests**

488 The authors declare that there is no conflict of interest.

489 **Funding**

490 **The authors disclosed receipt of the following financial support for the research of**
491 **this article:** This work was supported by the German Federal Ministry of Education and
492 Research (BMBF) through funding the projects “CADY – Central Asian Climate Dynamics”
493 (03G0813) and “CAHOL – Holozäne Klimaschwankungen in Zentralasien“ (03G0864).

494

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727

728

729 **Supplementary**

730

731 **Tab. S1: Excluded and merged samples due to low head capsule concentration**

732

733 Excluded samples:

734 123, 123.5, 124, 140, 140.5, 141, 141.5, 142, 142.5, 143, 143.5, 144, 144.5, 145, 145.5, 146,
735 152.5, 157.5, 163, 165, 167.5, 170 cm

736

737 Merged samples:

738 91-95 cm	(5 samples, total count sum 53.5 head capsules)
739 96-100 cm	(6 samples, total count sum 40 head capsules)
740 100.5-105 cm	(10 samples, total count sum 41 head capsules)
741 105.5-107.5 cm	(5 samples, total count sum 42 head capsules)
742 108-110.5 cm	(5 samples, total count sum 67 head capsules)
743 111.5-112 cm	(2 samples, total count sum 52.5 head capsules)
744 112.5-113.5 cm	(3 samples, total count sum 64 head capsules)
745 114-115.5 cm	(4 samples, total count sum 57 head capsules)
746 116-118 cm	(5 samples, total count sum 56 head capsules)
747 121.5-122.5 cm	(3 samples, total count sum 53 head capsules)
748 130.5-132 cm	(4 samples, total count sum 32.5 head capsules)
749 146.5-147 cm	(2 samples, total count sum 26 head capsules)
750 148.5-150 cm	(4 samples, total count sum 29 head capsules)

751

752 [Insert Fig. S1]

753 [Insert Fig. S2]

754

755 **Figure captions**

756 Fig. 1: Left: Bathymetric map of Lake Son Kol (isobaths with corresponding water depths in
757 meters below lake level; modified after Academy of Science of the Kyrgyz SSR, 1987) and
758 relief of the surrounding area (elevations in meters above sea level). The coring site is
759 indicated by a white circle (modified after Schwarz et al., 2017). Right: Topographic maps of
760 Central Asia (top panel) and Kyrgyzstan with neighboring countries (CHN – China, KZ –
761 Kazakhstan, UZ – Uzbekistan, TJ – Tajikistan) and the location of Son Kol (bottom panel).
762 The relief map of Kyrgyzstan is based on the CGIAR-CSI SRTM 90 m (3 arcsec) digital
763 elevation data (Version 4) of the NASA Shuttle Radar Topography Mission (Jarvis et al.,
764 2008).

765 Fig. 2: Summary diagram showing chironomid taxa (in % of the head capsule sum), sample
766 count sums (samples < 50 head capsules) and chironomid head capsule concentration
767 against depth (cm), with a secondary age-scale (cal. yr BP) plotted for comparison. The
768 width of the chironomid sample bars corresponds to the thickness of the (merged) samples.
769 Zonation based on chironomid assemblages. Samples below 150 cm are excluded due to
770 their low count sums.

771 Fig. 3: Principal components analysis(PCA) biplot of abundant chironomid taxa (arrows) and
772 samples (circles). The variance explained by the first and second axis is 58.1 and 29.3 %,
773 respectively. Blue circles mark the chironomid assemblage zones.

774 Fig. 4: Summary diagram of the paleoenvironmental development of Son Kol. From left to
775 right: picture of the sediment record, $\delta^{15}\text{N}$ - and TOC-values (Lauterbach et al., 2014),
776 selected ostracod taxa, the planktonic-benthic diatom ratio and the diatom-inferred
777 conductivity record (Schwarz et al., 2017); PCA axes 1 and 2 scores, presence of
778 Charophyte oogonia and *Daphnia ephippia* (this study). Horizontal lines in the PCA-records
779 indicate statistically significant regime shifts (see text for further information). Data plotted
780 against depth (cm) with a secondary age-scale (cal. yr BP) shown for comparison. Zonation
781 based on chironomid assemblages. Samples below 150 cm are excluded due to their low
782 count sums.

783 Fig. 5: Schematic representation of the Son Kol ecosystem response to a) an increase in the
784 strength of the Westerlies at 4900 cal. yr BP and b) a decrease in strength of the Westerlies
785 at 4300 cal. yr BP. We propose that this change in the strength of the Westerlies can be
786 seen as an external trigger that subsequently led to changes in turbidity, which themselves
787 acted as a driver affecting the macrophyte population of the lake, and, ultimately, the
788 ecosystem state.

789 Fig. S1: Cluster analysis based on square-root transformed percentages of non-rare
790 chironomid head taxa.

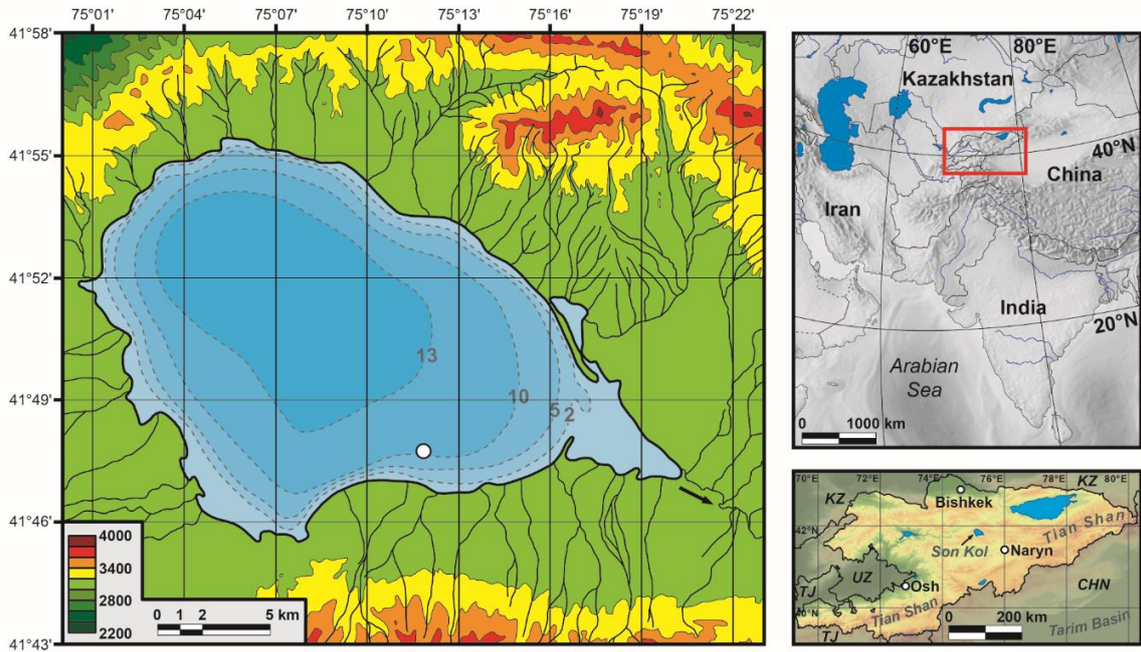
791 Fig. S2: Broken-stick analysis based on square-root transformed percentages of non-rare
792 chironomid taxa.

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794

795 **Figures**

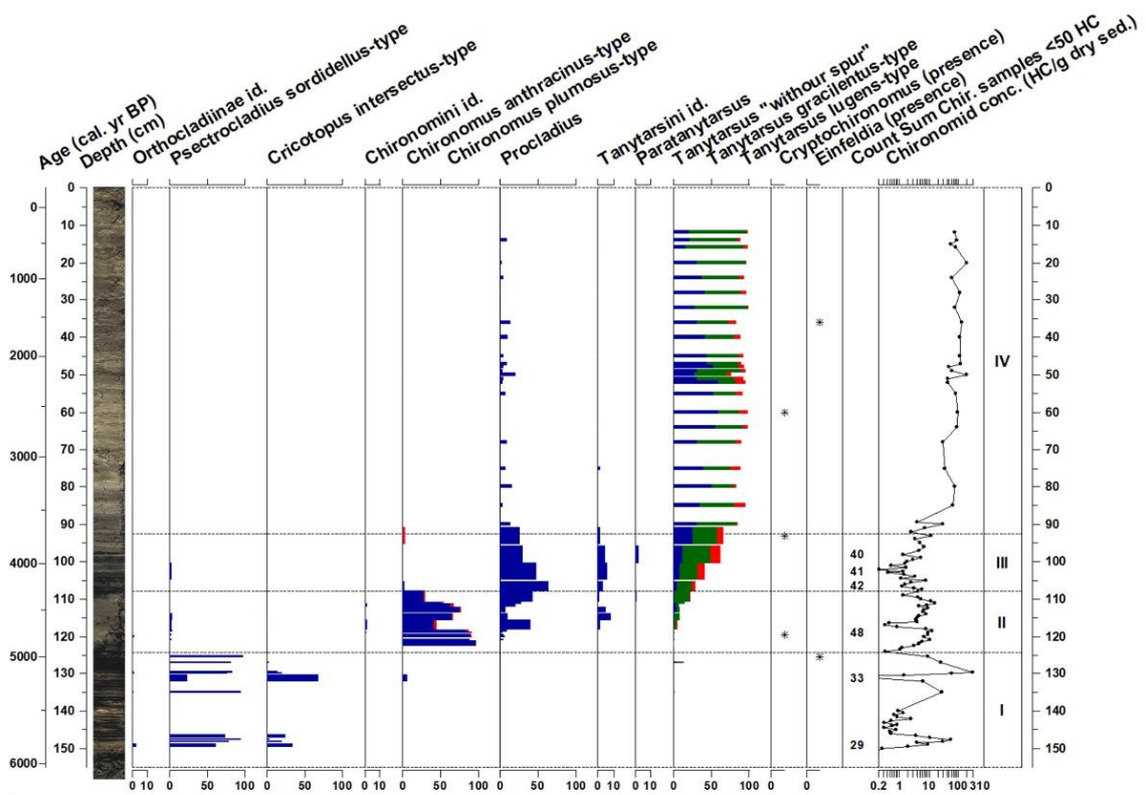
796 **Fig 1**



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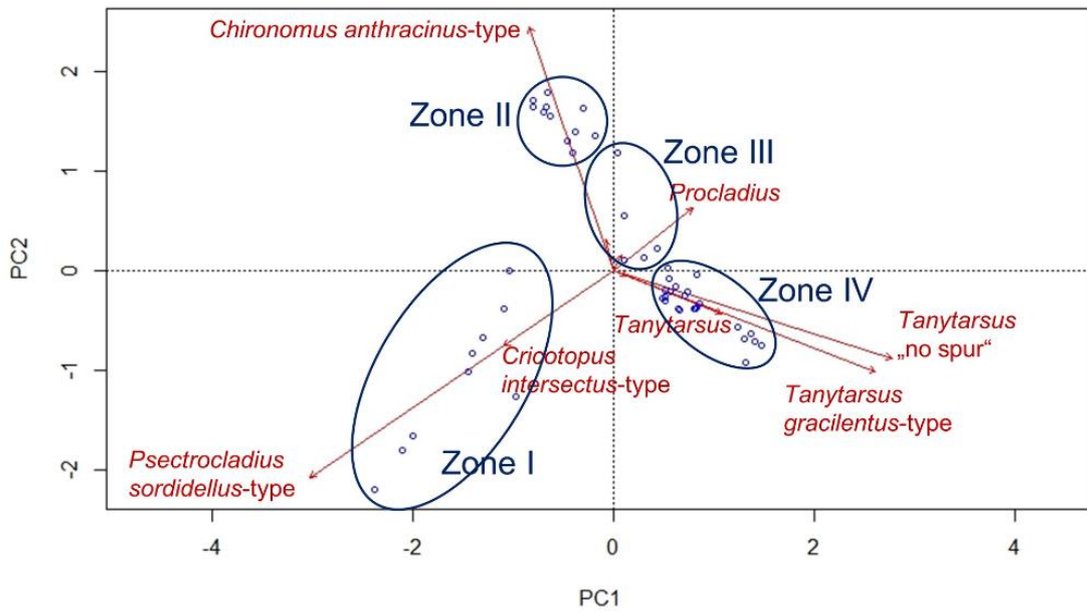
799 **Fig 2**



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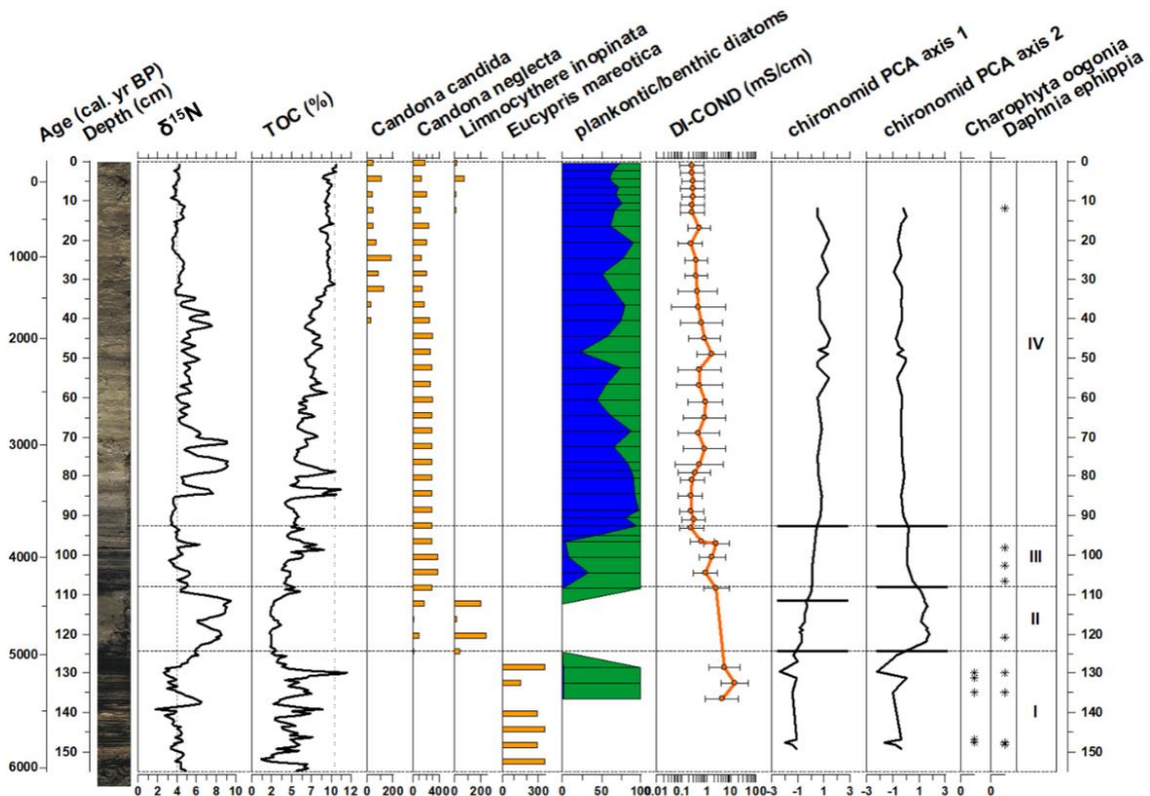
802 **Fig 3**



803

804

805 Fig 4

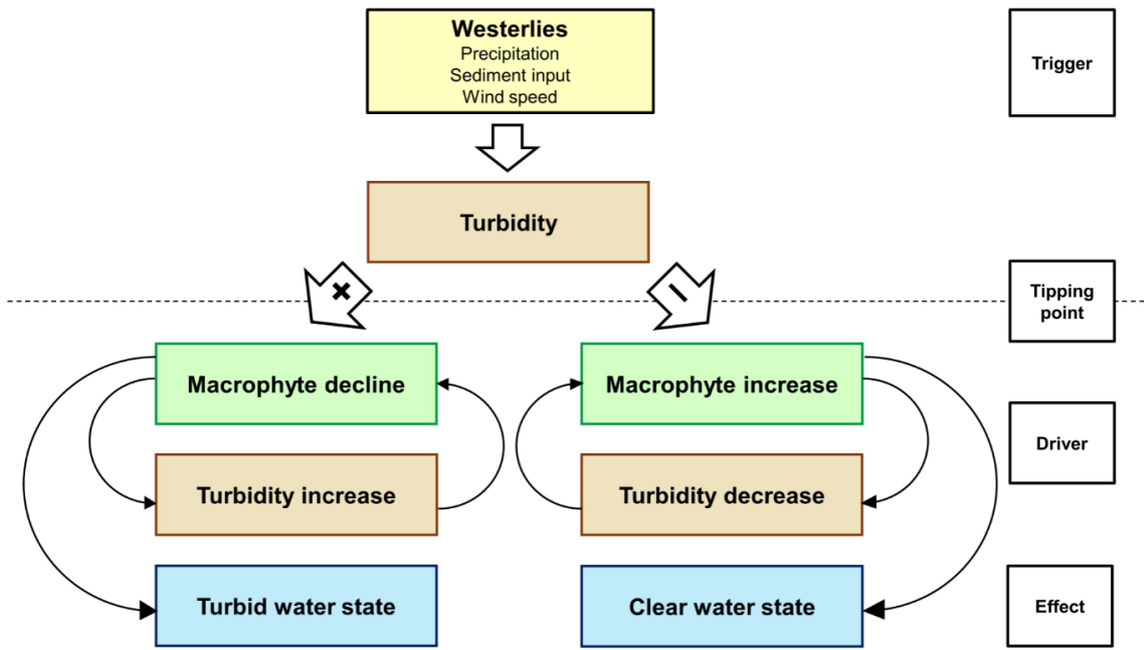


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808 Fig 5

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