

Palaeoecological potential of phytoliths from lake sediment records from the tropical lowlands of Bolivia

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- 11 Abstract

12 Phytolith analysis is conventionally an archaeo-botanical tool used to study past human activity using 13 material from excavations or soil pits. However, phytolith analysis also has potential as a 14 palaeoecological tool, to reconstruct vegetation changes through periods of climatic change and human influence. To study phytoliths from lake sediment alongside pollen requires an understanding 15 16 of phytolith taphonomy in lakes. Theoretical models suggest phytoliths represent more local 17 vegetation at smaller spatial scales than pollen from lake sediments, but this has not been tested 18 empirically in the Neotropics. This paper compares pollen and phytolith assemblages from the same 19 lake sediment surface sample, from a suite of lakes of different sizes across different vegetation 20 types of lowland tropical Bolivia. We find three factors driving phytolith composition in lakes: 21 taphonomy, lake size and phytolith productivity. By comparing phytolith assemblages with pollen 22 assemblages, we find that they provide different taxonomic information and generally complement 23 each other as palaeo-vegetation proxies. We also demonstrate empirically that pollen assemblages 24 in lake samples represent a larger catchment area than phytolith assemblages. Our findings suggest

25 that phytoliths can be particularly useful in providing local-scale vegetation histories from large

26 lakes, to complement the regional-scale vegetation histories provided by pollen data.

27

28 Keywords

29 Phytoliths, pollen, taphonomy, lake records, tropical, South America

30 1. Introduction

31 Phytolith analysis is conventionally an archaeo-botanical tool to study past human activity, used by 32 archaeologists studying material from excavations and/or soil pits. However, phytolith analysis also 33 has the potential, as a palaeoecological tool, to reconstruct vegetation changes through periods of 34 climatic change and/or human influence. There are two key areas where phytoliths can provide 35 additional information beyond the conventional vegetation reconstruction proxy of fossilised pollen: 36 taxonomic information and spatial information. While the taxonomic benefits of phytolith analysis 37 are reasonably well studied, the spatial scale phytolith records represent is less certain, particularly 38 when analyses are conducted on typical palaeoecological samples taken from lake sediment. 39 There are many taxonomic benefits of phytolith analysis for palaeoecologists. For example, phytolith 40 analysis can differentiate sub-families of Poaceae and genera of Cyperaceae, neither of which is 41 currently possible from pollen analysis, and which can be particularly helpful in identifying different 42 herbaceous habitats such as forest understorey, savannahs, and semi-aquatic lacustrine vegetation. 43 Further taxonomic advantages of phytoliths include the identification of Heliconiaceae, a key 44 disturbance indicator (Piperno, 2006), as well as other economically useful taxa unidentifiable by 45 their pollen, such as squash (Cucurbita) (Bozarth, 1987) and rice (Oryzoideae) (Hilbert et al., 2017), 46 which can provide important insights into past human land-use and human-environment 47 interactions. Furthermore, soil surface sample work by Dickau et al. (2013) and Watling et al. (2016) 48 have demonstrated that several neotropical ecosystems (humid evergreen forest, palm forest, semi49 deciduous dry forest, seasonally inundated savannah and terra firme savannah) can be

50 differentiated based solely on the phytolith assemblage from soil samples.

51 The combination of pollen and phytolith analyses should therefore provide additional, 52 complementary palaeoecological information. As pollen does not preserve well in soils, this can be 53 achieved by sampling a combination of soil samples (for phytolith analysis) and lake records (for 54 pollen analysis). However, this combination is complicated by the different spatial scales and 55 temporal resolution provided by soil versus lake sediment records. The temporal resolution of 56 palaeo-records from the soil column is typically significantly lower than that of lake sediment 57 records due to bioturbation of soil via plant roots, insects and other animal disturbances which move 58 material through the soil profile (Butler, 1995; Gabet et al., 2003). The spatial scale represented by 59 soil samples is generally much smaller than that of lake records, as lakes act as sinks for microscopic 60 particles from the surrounding vegetation transported via wind or water (Bennett and Willis, 2002). 61 To avoid these complications, as phytoliths preserve well in lake sediments as well as soils, both 62 analyses can be conducted on lake sediment which would enable the same temporal resolution to 63 be achieved for both proxies, enabling direct comparison of pollen and phytolith assemblages and 64 providing truly complementary taxonomical and spatial information. This approach has been applied 65 to several Late Quaternary neotropical lake records, including La Yaguada, Panama, where Late 66 Pleistocene cooling was identified by phytoliths from montane forest taxa such as Magnolia and 67 Chrysobalanaceae (Bush et al., 1992; Piperno et al., 1990) and Monte Oscuro, Panama, where a 68 wetter Holocene climate and human disturbance were identified based on pollen and phytoliths 69 records (Piperno and Jones, 2003). At Lagunas Granja and San José in lowland Bolivia, pollen and 70 phytolith analyses revealed late Holocene Pre-columbian land use (Carson et al., 2015; Whitney et 71 al., 2013). In these studies phytoliths are interpreted as representing a more local spatial scale than 72 pollen, based on a theoretical understanding of phytolith deposition and transport, backed up in the 73 latter two studies by comparisons with shoreline vegetation inventories.

74 Integration of pollen and phytolith records from lake sediments requires an understanding of their 75 respective taphonomy in lakes. Piperno proposed a theoretical model of phytolith representation in 76 lakes (Piperno, 2006, 1990) whereby phytolith source area depends largely on the lake site 77 characteristics. For example, in stream-fed lakes in areas with high precipitation phytoliths have the 78 potential to be transported long distances via soil erosion and runoff over land and into streams. In 79 open, frequently burnt environments they can be transported up to 2000 km by wind. However, in 80 closed basins surrounded by dense forest, Piperno (2006) proposes that phytoliths are likely to 81 represent only shoreline vegetation.

82 The only published empirical study on phytolith input to lakes (Aleman et al., 2014) was conducted 83 on three small lakes in ecosystems of central Africa: savanna, forest-savanna mosaic, and forest 84 (0.03, 0.36, 0.14 km² respectively). Aleman found that the proportion of forest cover surrounding 85 the lake and the number of large fires (producing ash clouds) were the main factors influencing 86 phytolith catchment area. However, there is uncertainty over the extent to which the findings from 87 these small lakes are representative of much larger lakes, several km in diameter, which are 88 common throughout the tropics. In addition, these phytolith records were not compared with other 89 vegetation proxies with an estimable source area, such as pollen, and only one sample was taken 90 from each lake. Furthermore, the role of fire, wind and water in phytolith taphonomy in the drier, 91 frequently burnt environments of Aleman's study area is likely to differ from the wet environments 92 of the Neotropics.

This paper therefore aims to explore the potential value of phytoliths as a complementary proxy to
 pollen for enhancing the palaeoecological information that can be obtained from lake sediments
 in tropical lowland Bolivia.

96 Specifically, this paper will address two questions:

97 I. Can humid evergreen forest, semi-deciduous dry forest and seasonally-inundated savannah
98 ecosystems be differentiated based on phytolith assemblages from Bolivian lakes?

99 II. What are the key factors driving the composition of phytolith and pollen assemblages in100 these lakes, and how do they differ between proxies?

101 The approach of this study is to compare pollen and phytolith assemblages from the same lake 102 sediment surface sample, from a suite of lakes of different sizes across different vegetation types of 103 lowland tropical Bolivia. For some of the larger lakes, samples were analysed from multiple core sites 104 within the lake to enable analysis of the spatial variation in phytolith assemblages across the lake 105 compared to pollen. These assemblages were compared through ordination analyses (PCA), and 106 potential driving factors in their differentiation were identified through constrained ordination 107 (RDA). Comparison between these phytolith and pollen assemblages provides a useful reference 108 point for palaeoecologists, drawing out potential areas where phytoliths from lake sediments can 109 provide additional, useful information.

110

111 2. Study area and site selection

112 The study area in this paper is lowland Bolivia. This region encompasses a wide variety of vegetation 113 types, as it is the transitional zone between the humid evergreen forests of the Amazon in the north 114 and semi-deciduous dry forests and savannah to the south. The distribution of these ecosystems is 115 controlled at the broad scale by climate, as precipitation decreases towards the south of the study 116 area. At a finer scale their distribution is controlled by geology and edaphic variables, with acidic 117 soils of Pre-Cambrian bedrock supporting terra firme forests, alkaline soils supporting semi-118 deciduous forests, and clay-rich basins supporting seasonally-inundated ecosystems (Fig. 1). 119 [Figure 1 – Map of lake sites and vegetation surveys plotted against vegetation classifications 120 adapted from WWF terrestrial ecoregions (Olsen et al 2001) using ArcGIS 10.5.1. Lake sites are 121 shown by red circles. Vegetation inventories and surveys are shown by black circles (IL = Vegetation 122 survey around Lagunas Isirere and Limoncin, AC2 = Acuario 2 forest plot inventory, LF1 = Los Fierros

123 1 forest plot inventory). Inset map of South America.]

124

Our study sites encompass three regions: (2.1) Chiquitania-Pantanal, (2.2) the Beni basin, and (2.3)
terra firme humid evergreen forests on the Pre-Cambrian shield (Fig. 2). Lake sites were chosen from
within each region to represent a range of lake sizes (Table 1).

128 [Table 1 - Summary of lake site characteristics, including references to the papers originally

129 publishing some of the pollen and phytolith records.]

130

131 [Figure 2 - Maps of lake sites and surrounding ecosystems created using ArcGIS 10.5.1. Panel A 132 shows the Chiquitania-Pantanal study region with Lagunas La Gaiba and Mandioré; Panel B shows 133 terra firme humid evergreen forests on the Pre-Cambrian shield study region with Laguna Chaplin; 134 Panel C shows the northern areas of the Beni basin study region with Lagunas Oricoré, La Luna and 135 Granja; and Panel D shows the southern areas of the Beni basin study region with Lagunas San José 136 and Limoncin. The locations of all surface samples are shown for each lake by black circles. 137 Vegetation classification of the study area, based on Landsat imagery, was provided by the Museo 138 de Historia Natural 'Noel Kempff Mercado', Santa Cruz, Bolivia, in 2015. Scale bars and latitude and 139 longitude are presented for each individual map panel.]

140

2.1. Chiquitania-Pantanal. This region encompasses the Chiquitano semi-deciduous dry forests. Our
study sites are at the eastern edge of the semi-deciduous dry forest on the Bolivia-Brazil border
where it meets the Pantanal wetlands of Brazil.

144 Semi-deciduous Chiquitano dry forest is a moderately diverse transitional forest type which grades

into humid evergreen forest to the north. Generally, dry forest is categorized by tree heights of 15-

146 20 m with emergents rarely exceeding 25 m. The canopy is less closed than humid evergreen forests,

147 allowing light to penetrate to the forest floor and develop relatively dense understory vegetation. A

148 key part of this understory is bamboo scrub, often characterised by *Guadua paniculata*. A key

dominant tree species is often *Anadenanthera colubrina* of the Fabaceae family (Killeen et al., 2006;

150 Killeen and Schulenberg, 1998) (Table 2). A detailed study of the vegetation of the eastern

151 Chiquitano forests around Lagunas La Gaiba and Mandioré can be found in Prance and Schaller

152 (1982).

[Table 2 – Vegetation inventory of Acuario 2, a 1-hectare vegetation plot just south of Noel Kempff
 Mercado National Park, gives a representative vegetation community composition for semi-

155 deciduous dry forest in south-west Amazonia. Inventory conducted by recording all taxa

representing >1% of the total number of stems >10 cm d.b.h. (Gosling et al., 2009).]

157 The Pantanal basin is a geographic depression surrounded by highlands which contains a floodplain 158 that houses the world's largest tropical wetland (~135, 000 km²). The vegetation of the Pantanal is a 159 heterogeneous mosaic of xeric, mesic and seasonally-inundated plant communities, controlled by 160 edaphic variables, topography and flooding. Seasonally-inundated savannahs are dominant in low 161 elevation areas, with higher elevation areas supporting inundation-tolerant gallery forests and semi-162 deciduous dry forests (Alho, 2005; Hamilton, 2002; Nunes da Cunha et al., 2007). Lagunas La Gaiba 163 and Mandioré have semi-deciduous dry forest to the west and south and the Pantanal wetlands to 164 the east and north:

2.1.1. Laguna La Gaiba is a large, shallow lake (~90 km², depth ~4-6 m) split into two sub-basins, the
deeper southern basin being largely surrounded by forest, and the shallow northern basin merging
into the Pantanal wetlands (Whitney et al., 2014). Core site 6 sits to the west of the south basin, core
site 10 towards the north of the south basin, and core site 16 within the northern basin (Fig. 2). Core
site numbers refer to those in the original publication of pollen and diatom records from this site
(Whitney et al., 2011).

171 2.1.2. Laguna Mandioré is a large, shallow lake (~152 km², depth ~4 m) forming a single continuous
172 basin. On the eastern shores dry forest grades into cerrado savannah as altitude increases (up to 846

m.a.s.l.) on the Amolar hill formation which separates the majority of the lake margin from the
Pantanal wetlands. Core site 2 is located in the southwest of the basin and core site 5 is located in
the northeast, close to the Amolar hills (Fig. 2).

176 **2.2. Beni basin.** This Amazonian sub-basin is formed by a bed of impermeable alluvial clays which 177 permit flooding during the wet season from November to March. The geomorphology of the Beni 178 leads to a dominance of seasonally-inundated savannah vegetation, with outcrops of humid 179 evergreen or seasonally-dry forest on islands of Pre-Cambrian Shield within the basin (Clapperton, 180 1993). The seasonally-inundated savannahs do not have a well-developed woody stratum but 181 maintain a short stratum of grasses, most commonly Pasalpum lineare, Leptocoryoheum lanatum, 182 Mesosetum sp., Sacciolepsis angustissima and Panicum parviflorum. Islands within the wetlands are 183 commonly formed by termite mounds which can raise an island 0.5-1.5 m above the flooded plain, 184 allowing forest species to colonise, such as Curatella americana and Davilla nitida (Dilleniaceae), 185 Casearia arborea (Salicaceae) and Tapiria guinanensis (Bignoniaceae) (Killeen and Schulenberg, 186 1998) (Table 3). The lake sites from within the Beni basin are Lagunas Oricoré, La Luna, Granja, 187 Limoncin and San José:

[Table 3 – Results of a qualitative vegetation survey ranking taxa as dominant, abundant, frequent or
occasional in coverage from the area surrounding Lagunas Limoncin and Isirere are presented to give
an example vegetation community composition for the Beni seasonally-inundated savannah. (Dickau
et al., 2013).]

2.2.1. Laguna Oricoré is a large lake (~10.5 km², depth 1-1.5 m) at the eastern edge of the Beni
basin, close to the geological boundary with terra firme humid evergreen forest on Pre-Cambrian
Shield bedrock. The lake is largely surrounded by seasonally-inundated savannah, with a small patch
of semi-deciduous dry forest near the northeastern shore (Fig. 2). It is situated 5 km south of the
evergreen forest boundary (Carson et al., 2014).

2.2.2 Laguna La Luna is a small lake (0.33 km², depth 2 m) at the eastern edge of the Beni basin, 5
km west of Laguna Oricoré. The lake is largely surrounded by seasonally-inundated savannah and
adjacent to a small (7.4 km²) forest island (Fig. 2) (Carson et al., 2016).

2.2.3 Laguna Granja is a small oxbow lake (0.071 km², depth 2 m) across the boundary from the Beni
 basin on the Pre-Cambrian Shield. The lake margins are dominated by riparian forest, blending into
 terra firme humid evergreen forest further away from the lake. To the east of the lake an area of
 ~0.3 km² has been cleared for cattle grazing (Fig. 2) (Carson et al., 2015).

204 **2.2.4 Laguna Limoncin** is a small lake (0.73 km², depth 0.9 m) in the south of the Beni basin,

surrounded by a mosaic of seasonally-inundated savannah and evergreen forest (Fig. 2).

206 **2.2.5 Laguna San José** is a large, shallow lake (14.3 km², depth 1 m) in the south of the Beni basin,

surrounded by a mosaic of seasonally-inundated savannah and evergreen forest. The lake is 4 km
from Laguna Limoncin and has a fringing strip of riparian forest <20 m wide. The core site is located
close to the northeast shoreline (Fig. 2) (Whitney et al., 2013).

210 **2.3. Terra firme humid evergreen forest**. This ecosystem is characterised by dense tall forest 211 comprised of trees reaching 45 m in height and 1.5 m in diameter, with a closed canopy and a low 212 density of understorey vegetation. The Moraceae family often dominates the evergreen forest 213 canopy, while palms (Arecaceae family) are also abundant. Emergent trees can include genera from 214 the Vochysiaceae, Fabaceae and Clusiaceae (Table 4). The rhizomatous species Phenakospermum 215 guianense (Strelitziaceae) is locally abundant and forms huge dense colonies (Killeen and 216 Schulenberg, 1998). The lake site from within terra firme evergreen forest is Laguna Chaplin. 217 [Table 4 – Vegetation inventory of Los Fierros 1, a 1-hectare vegetation plot within Noel Kempff Mercado National Park, gives a representative vegetation community composition for evergreen 218

forest in south-west Amazonia. Inventory conducted by recording all taxa representing >1% of the

total number of stems >10 cm d.b.h. (Gosling et al., 2005).]

2.3. Laguna Chaplin is a large, shallow, flat-bottomed lake (12.2 km², depth 2.5 m) within the Noel
Kempff Mercado National Park in eastern Bolivia, near the southern limit of Amazon humid
evergreen forest. Chaplin is surrounded by humid evergreen forest with a small fringe of palm
swamp and area of savannah wetland at the southwestern edge of the lake (Burbridge et al., 2004).
Core site 1 is close to the centre of the lake, core site 2 is towards the east of the lake, and core site
3 is close to the southern shore (Fig. 2).

227 3. Methods

228 3.1 Use of previously published datasets

This paper uses a number of previously published datasets in combination with new sample analyses(Table 1).

Field methods for all samples entailed long, overlapping sediment cores being retrieved using a stable floating platform and modified drop-hammer Livingston piston corer (Colinvaux et al., 1999) and shipped back to the UK in their core tubes. Surface sediments were taken using a 5-cm diameter Perspex[®] tube and piston to capture the uppermost unconsolidated sediments and were divided into consecutive 0.5 cm or 1.0 cm samples. These surface samples were stored in watertight plastic tubes. All samples were kept in cold storage at 4°C.

Standard pollen preparation protocols were followed for each of the previously published datasets, 237 238 details can be found in each paper (Burbridge et al., 2004; Carson et al., 2016, 2015, 2014; Whitney 239 et al., 2013, 2011, Plumpton et al., accepted). Chaplin 3 pollen counts are included in Figure 3 as 240 they are the only complete pollen counts we have for Chaplin. The pollen counts for Chaplin 1 and 2 241 have been summarised to show only the most abundant taxa, as described in Burbridge et al., 242 (2004). As the pollen assemblages from all three Chaplin surface samples are highly consistent with 243 each other (Burbridge et al 2004) we use the detailed pollen assemblage from Chaplin 3 to infer 244 likely pollen abundances at Chaplin 1 and 2 for taxa not included in the summary data e.g. 245 Arecaceae. Phytolith extraction was conducted using the wet oxidation methods as described in

- 246 Piperno (2006) for each of the previously published datasets, details can be found in each paper
- 247 (Carson et al., 2015; Whitney et al., 2013, Plumpton et al., accepted).

248 3.2 New data analysis - laboratory methods – phytolith processing and identification

249 New phytolith extraction and analysis was conducted on lake surface sediments from Lagunas

250 Oricoré, La Luna, Chaplin (1 and 2) and Limoncin (Table 1). Phytolith extraction was not possible for

251 Chaplin 3 due to lack of sample material, so phytoliths were analysed from Chaplin 1 and 2 only.

Phytolith extraction was conducted using the wet oxidation method involving nitric acid heated to
90°C as described in Piperno (2006). For Laguna Chaplin, 3cc of wet sediment was sampled to allow

254 for fractionation during processing into "A" (<53 μm) and "C" (53-250 μm) fractions. For the other

sites, only 1cc of wet sediment was available from the surface sample horizon, due to intensive

256 previous study of these sites. These samples were not fractionated due to the small sediment

volume available. Non-fractionated and A fraction slides were counted at 400X magnification and a

258 minimum sum of 200 diagnostic phytoliths was counted for each sample. C fractions slides were

scanned at 100X, with identifications conducted at 400X magnification. All phytoliths with taxonomic

significance on the C fraction slides were counted.

261 Phytolith identification was carried out with reference to the University of Reading tropical phytolith 262 reference collection, which contains modern specimens from 152 species, and photographs of the University of Exeter phytolith reference collection, which contains over 500 modern neotropical 263 264 plant specimens. Published phytolith reference atlases were also consulted from the Neotropics, 265 tropical Africa, Asia and Australasia (Boyd et al., 1998; Dickau et al., 2013; Iriarte and Paz, 2009; 266 Kondo et al., 1994; Lu and Liu, 2003; Mercader et al., 2011, 2009; Piperno, 2006; D.R. Piperno and 267 Pearsall, 1998; Dolores R. Piperno and Pearsall, 1998; Runge, 1999; Wallis, 2003; Watling et al., 268 2016; Watling and Iriarte, 2013). Table 5 lists all of the phytolith types identified in this study, their 269 taxonomic association and codenames for PCA and RDA graphs.

270 [Table 5 – Phytoliths types identified with abundance >1%, their taxonomic association and PCA/RDA
271 codes.]

272 3.3 Numerical analysis

273 For statistical analysis, phytolith samples were not analysed as separate A and C fractions despite 274 Mandioré and Chaplin samples being fractionated during processing. In these two cases A and C 275 fraction sums were added together as laboratory error led to large number of small phytoliths being 276 present in the C fractions for Chaplin so samples were effectively not fractionated, and C fraction 277 counts were so low for Mandioré (3-6 total) that they could not have been analysed separately in 278 ordination analyses as a minimum count of 50 phytoliths is required for robust analysis (Dickau et al 279 2013). Furthermore, summing the A and C fractions for Mandioré and Chaplin enables comparison of 280 these records with all other lake sites studied which had insufficient sediment available for 281 fractionation.

282 Frequency plots for phytoliths and pollen taxa were created using version 1.7 of the C2 software 283 (Juggins, 2016). Ordination analyses were conducted and plotted using the vegan 2.5-2 package 284 (Oksanen et al., 2018) in R 3.4.1. Only taxa with >1% abundance were included in the analysis. 285 Abundances were square-root transformed (the Hellinger transformation) prior to analysis. 286 Detrended correspondence analysis (DCA) of summed counts demonstrated relatively short 287 environmental gradients in the dataset, therefore Principal Component Analysis (PCA) was chosen 288 for ordination of the phytolith and pollen assemblages, and Redundancy Analysis (RDA) for 289 constrained ordination. The environmentally constraining variables considered were: lake size, 290 distance of core site to shore, and average tree cover from local to regional scale (within 100, 1000, 291 5000, 10000 and 20000 m radius from the lake shore). Permutation tests were conducted on the 292 RDA model to assess the statisistcal significance of the environmental constraining variables as 293 predictors of the variation in pollen and phytolith assesmblages (Borcard et al., 2011; ter Braak and 294 Verdonschot, 1995). For both the pollen RDA and phytolith RDA, an ANOVA "by term" (i.e.

environmental variable) was conducted with 999 permutations in R 3.4.1 using vegan 2.5-2 (Oksanen
et al., 2018).

297 3.3.1 Spatial analyses - GIS

Tree cover data within 100, 1000, 5000, 10000 and 20000m radius from the lake shore were

299 extracted from the Hansen et al. (2013) dataset using the following method. The circumference of

- 300 each lake site was traced using Google Earth Pro geometry tools and these layer files were converted
- to shape files using QGIS 2.14.0. Buffer zones around each lake site were created using the buffer
- tool in ArcGIS 10.4 at distances of 100, 1000, 5000, 10000 and 20000m from the lake shorelines.
- 303 Tree cover data was then extracted from within each buffer zone and mean tree cover calculated
- using packages rgdal 1.4-4 (Bivand et al., 2019), raster 2.9-5 (Hijmans, 2019), and maptools 0.9-5
- 305 (Bivand and Lewin-Koh, 2019) in R 3.4.1. Lake area and distance of core site to shore were calculated
- 306 using Google Earth Pro geometry tools.
- **307** 4. Results

308 *4.1 Key trends in phytolith and pollen assemblages from each region*

- 309 4.1.1 Chiquitania-Pantanal semi-deciduous dry forest Lagunas Mandioré and La Gaiba
- 310 [Figure 3 Summary diagram of phytolith abundances from all lakes studied, presented as
- 311 percentage data. Vegetation surrounding the lakes has been classified into 3 ecosystem types: semi-

deciduous dry forest (Chiquitania-Pantanal), seasonally inundated forest-savannah (Beni basin), and

313 evergreen forest (PCS Humid Evergreen Forest).]

- 314
- 315 [Figure 4 Summary diagram of pollen abundance from all lakes studied, presented as percentage of
- terrestrial total. Vegetation surrounding the lakes has been classified into 3 ecosystem types: semi-
- 317 deciduous dry forest (Chiquitania-Pantanal), seasonally inundated forest-savannah (Beni basin), and
- evergreen forest (PCS Humid Evergreen Forest). Pollen data for Chaplin 1 and 2 is only available in

highly summarised format with data for a restricted number of taxa. Full pollen counts are availablefor Chaplin 3.]

321

322 The two surface samples from Laguna Mandioré (Fig. 2) show consistent phytolith assemblages with 323 high Poaceae phytolith total abundances (67-71%), with significant contributions from Bambusoid 324 (19-24%) and Panicoid types (22-29%) (Fig. 3). Arboreal phytoliths make up 22-26% of the 325 assemblage (Fig. 3). The three surface samples from Laguna La Gaiba (Fig. 2) show lower Poaceae 326 phytolith total abundances (50-62%), largely due to lower abundances of Panicoid types (6-13%) than the Laguna Mandioré samples (Fig. 3). Arboreal phytoliths comprise 18-35% of the assemblages 327 328 at Laguna La Gaiba, showing greater variation than the Laguna Mandioré samples. This is driven by 329 the higher aboreal phytolith abundance at La Gaiba 6 compared to La Gaiba 10 and 16, which are 330 more consistent with each other (Fig. 3).

331 Overall, both lake sites in the Chiquitania-Pantanal semi-deciduous dry forest have a high percentage 332 of Poaceae phytoliths (50-71% of total assemblage). A large proportion of this Poaceae total is 333 Bambusoid phytoliths at 17-24%. Cyperaceae phytolith abundance in the semi-deciduous forest 334 samples is the highest of the three ecosystem types studied ranging from 5-9% of total. Arboreal 335 phytolith totals are largely comprised of non-Arecaceae types. The most abundant arboreal taxa in 336 semi-deciduous dry forests, such as Fabaceae (including Anadenanthera) and Bignoniaceae 337 (including Tabebuia) (Table 2), cannot be differentiated to family or genus level using phytoliths 338 (Piperno, 2006). However, these taxa would contribute to the phytolith arboreal indicators such as 339 globular granulates and faceted elongates within the woody eudicot category (Fig. 3).

Pollen assemblages from the Chiquitania-Pantanal samples are reasonably consistent, showing the highest *Anadenanthera* (4%) and *Astronium* (3%) and lowest Moraceae/Urticaceae (0%) pollen abundances of all samples studied (Fig. 4). The total arboreal pollen abundance is constant across the samples at 18-29%, except La Gaiba 16 at 6%. This pattern broadly fits with the total arboreal

phytolith abundances where La Gaiba 16 is also low at 18%, but so are La Gaiba 10 and Mandioré 5
at 21% and 22% respectively (Fig. 3). The samples from Lagunas Mandioré and La Gaiba also show
the highest Poaceae pollen abundance of all samples studied at 42-56% (Fig. 4). These pollen results
are in contrast to the Poaceae phytolith abundances, which are similar between semi-deciduous dry
forest sites and Beni seasonally-inundated forest-savannah mosaic sites (Fig. 3). There is also
stronger variation in Cyperaceae pollen abundance, ranging from 7% to 34%, than Cyperaceae
phytolith abundances which range from 5% to 9%.

4.1.2 Beni seasonally-inundated savannah-forest mosaic – Lagunas Oricoré, La Luna, Granja, Limoncin
and San José.

353 Phytolith assemblages from the Beni basin lakes within seasonally-inundated savannah-forest 354 mosaic have consistently high grass and herb phytolith totals, between 75% and 85% of total 355 assemblage (Fig. 3). For most lake sites this total is dominated by Poaceae phytoliths, except Laguna 356 San José which has a significant contribution of herbs such as Heliconiaceae, Marantaceae and 357 Asteraceae (11%, 15% and 12% of total assemblage respectively). All other samples from the Beni 358 lakes contain low (<6%) abundances of phytoliths from these herbs, or they are absent (Fig. 3). 359 Across all samples from Beni lakes, the composition of the Poaceae phytolith total varies with 360 particularly high Panicoid phytolith abundance at Laguna Granja (51%), and high Bambusoid 361 phytolith abundance at Lagunas La Luna and Limoncin (23% and 20% respectively). Cyperaceae 362 phytoliths appear at low abundance in all samples from the Beni, except Laguna San José where they 363 are absent (Fig. 3). Total arboreal phytolith abundance is the lowest of all ecosystems studied, 364 ranging from 12% at Laguna Limoncin to 26% at Laguna Granja. Arecaceae phytoliths make up a 365 significant proportion of arboreal types at Lagunas Limoncin and San José, whereas woody eudicot 366 phytolith types dominate at Lagunas Oricoré, La Luna and Granja (Fig. 3). 367 Pollen assemblages from the Beni basin lakes show significant variation in total arboreal pollen 368 adundance, driven largely by the low abundance of Moraceae/Urticaceae pollen at Laguna Limoncin

369 (2%) and Laguna San José (5%), compared to 19% at Laguna Oricoré (Fig. 4). This pattern matches 370 that in the phytolith assemblages, where Laguna Limoncin has the lowest total arboreal phytolith 371 abundance at 12% (Fig. 3). The abundance of Arecaceae phytoliths at Lagunas Limoncin and San José 372 of 9% and 16% respectively, is not reflected in the pollen assemblage where Arecaceae pollen is 373 absent for both sites (Fig. 4). Cecropia pollen abundance is highest at Lagunas San José and Granja at 374 15% and 13% respectively (Fig. 4). There is signiciant variation in Cyperaceae pollen abundance with 375 particularly low levels at Oricoré and La Luna (10% and 6% respectively) compared to Limoncin and 376 Granja (39% and 31% respectively). Overall, the samples show more variation in total grass and herb 377 pollen than total grass and herb phytoliths, with Limoncin and Granja showing significantly higher 378 grass and herb pollen totals than the other samples (Fig. 4).

379 4.1.3 Terra firme humid evergreen forest – Laguna Chaplin

380 The two phytolith samples from Laguna Chaplin within terra firme evergreen forest are 381 differentiated from the samples from other ecosystems by the high abundance of Arecaceae 382 phytoliths (53-59% of total assemblage) (Fig. 3). Other arboreal phytolith types are present in similar 383 abundance to semi-deciduous forest assemblages, ranging from 16-22%. Grass and herb phytolith 384 abundances are the lowest of all samples studied at 19-30% of the assemblage, with low abundances 385 of all Poaceae (13-25%) and Cyperaceae (2%) phytolith types but particularly low abundances of 386 Bambusoid types (5-6%). The surface sample closer to the shore (Chaplin 2) shows a higher 387 abundance of Poaceae and lower arboreal phytoliths than the more central surface sample (Chaplin 388 1).

The complete pollen assemblage from Laguna Chaplin (Chaplin 3) shows the lowest Poaceae (4%) and highest total arboreal pollen (76%) abundance of all samples studied (Fig. 4), matching the pattern shown in the phytolith assemblages from Laguna Chaplin (Chaplin 1 and 2) (Fig. 3). Within this total arboreal figure, pollen abundance is made up largely of Moraceae/Urticaceae (48%) with

393 some *Cecropia* (9%) and *Alchornea* (5%) pollen. This assemblage is in contrast to the phytoliths,

394 where arboreal phytolith types are dominated by Arecaceae phytoliths (53-59%).

395 *4.2 PCA results*

[Figure 5 – PCA biplot for a) phytolith and b) pollen data. Axes are Principal Component 1 (PCA1) and Principal Component 2 (PCA2). The percentage in parentheses on these axes is the percentage of variance in the dataset explained by that principal component. Lake sites are coloured according to the ecosystem they represent: light green = semi-deciduous dry forest, yellow = Beni seasonallyinundated forest-savannah mosaic, dark green = humid evergreen forest. Taxa are presented in black. Codes for taxa names are presented in Tables 5 and 6 for phytoliths and pollen respectively.]

402

403 PCA of the phytolith dataset (Fig. 5) shows differentiation of the terra firme evergreen forest 404 samples (Chaplin 1 and 2) from all other samples studied in this paper, largely by PCA1. PCA1 405 explains 42% of the variance in the dataset and is driven largely by the abundance of Arecaceae 406 phytolith types (labelled Palm in Fig. 5, see Table 5 for full list of phytolith PCA codenames). PCA1 is 407 positively correlated with abundance of Arecaceae, Heliconiaceae and Marantaceae types and 408 negatively correlated with Poaceae types, such as Panicoid crosses, rondels and bulliforms (Fig. 5). 409 All other samples plot negatively on PCA1, except Laguna San José. PCA2 explains 20.5% of the 410 variance in the dataset and is correlated positively with Panicoid crosses and negatively with 411 Poaceae and Bambusoid bulliform phytolith types (Fig. 5). This axis separates sites with higher 412 Panicoid abundances, such as Laguna Granja which is at present surrounded by agricultural land and 413 Laguna Mandioré which is close to upland savannah, from sites with less Panicoid inputs such as 414 Laguna La Gaiba.

PCA of the pollen dataset (Fig. 5) shows clear differentiation of the terra firme evergreen forest
sample (Chaplin 3) from all other samples by PCA1, which is positively correlated with abundance of
Moraceae/Urticaceae pollen and also *Celtis* and *Isoetes* (see Table 6 for a list of pollen PCA

418 codenames). The two samples from Laguna Mandioré, within Chiqtuiania-Pantanal, are also clearly

419 differentiated from all other samples as they plots negatively on PCA1, correlated with abundance of

420 Poaceae, Anadenanthera and Eichhornia. Both Chaplin and Mandioré samples plot positively on

421 PCA2, correlated with low abundance of Cyperaceae, *Typha* and *Cecropia* pollen. The samples from

422 Laguna La Gaiba, within Chiquitania-Pantanal, plot negatively on PCA1 and PCA2, as do samples from

423 the Beni basin samples from Lagunas Limoncin, San José and Granja. Samples from Lagunas Oricoré

424 and La Luna are differentiated by plotting slightly positively on PCA1 and PCA2 (Fig. 5).

425 [Table 6 – Pollen taxa identified with abundance >1% and PCA/RDA codes.]

426 Laguna La Gaiba has three surface samples (6, 10, 16), Laguna Mandioré has two (2, 5) and Laguna 427 Chaplin has three (1, 2, 3). The phytolith PCA results generally show clustering by lake, with the two 428 Chaplin samples and the three La Gaiba samples plotting together (Fig. 5). Chaplin samples 1 and 2 429 are particularly closely grouped. For La Gaiba, samples 6, 10 and 16 are also closely clustered, but 430 with some overlap with lake sites from the Beni i.e. Oricoré. Mandioré samples 2 and 5 cluster 431 within the same quadrant of the phytolith PCA but are not closely grouped (Fig. 5). In comparison, 432 the pollen PCA results show tighter clustering of the Mandioré samples than the La Gaiba samples, 433 although both are clearly grouped (Fig. 5).

434 *4.3 RDA results*

435 [Figure 6 – RDA tri-plot for a) phytolith and b) pollen data. Axes are Redundancy Analysis 1 (RDA1) 436 and 2 (RDA2). The percentage in parentheses on these axes is the percentage of variance in the 437 dataset explained by that component. Environmental constraining variables are presented in blue: 438 lake area, distance of core site to lake shore, and tree cover within 100, 1000, 2000, 10,000 and 439 20,000 m of the lake shore. Lake sites are coloured according to the ecosystem they represent: light 440 green = semi-deciduous dry forest, yellow = Beni seasonally-inundated forest-savannah mosaic, dark green = humid evergreen forest. Taxa are presented in black. Codes for taxa names are presented in 441 442 Tables 5 and 6 for phytoliths and pollen respectively.]

444 The pollen RDA (Fig. 6) shows that lake area and distance of core site to shore are both strongly 445 negatively correlated with axes 1 and 2, explaining the differentiation of large lakes such as 446 Mandioré from smaller lakes such as Granja. Tree cover at 1,000 – 20,000 m is also negatively 447 correlated with axes 1 and 2, although more strongly with axis 2, particularly for tree cover at 20,000 448 m. Axis 2 represents the differentiation of highly forested sites such as Chaplin and Mandioré, from 449 sites in more open savannah-lake landscapes such as Limoncin, Granja, San Jose and to a lesser 450 extent La Gaiba. The results of the permutation test show that lake area, tree cover at 100 m and 451 tree cover at 20,000 m are the most statistically significant environmentally constraining variables 452 (p=0.001 for all three variables) (S1). Tree cover at 10,000 m and 5,000 m are also statistically 453 significant explanatory variables (p=0.008 and p=0.023 respectively) (S1).

The phytolith RDA (Fig. 6) shows that while tree cover at 1,000 - 20,000 m is positively correlated with axis 1, the strongest correlation with axes 1 and 2 is tree cover at 100 m. Lake area and distance to shore are both negatively correlated with axis 1. The results of the permutation test show that the most statistically significant environmentally constraining variable is tree cover at 100 m (p=0.001),

458 followed by tree cover at 5,000 m (p=0.002) (S2).

459 5. Discussion

460 5.1 Differentiating ecosystems

Based on the phytolith assemblages of lake records studied here it is possible to differentiate humid evergreen forest from semi-deciduous forest and Beni seasonally-inundated savannah-forest mosaic sites, largely by the high abundance of Arecaceae phytoliths, but not possible to differentiate semideciduous forest from seasonally inundated savannah-forest mosaics (Fig. 5). This contrasts with the findings of Dickau et al. (2013) who were able to differentiate these ecosystems, based on soil samples from within Noel Kempff Mercado National Park (NKMNP) and the shorelines of Lagunas Limoncin and Isirere in the Beni Basin. There are two main differences between the phytolith

assemblages from lakes in this study and those in soil samples from the same ecosystems, as
described by Dickau et al. (2013). Firstly, in semi-deciduous forest sites our lake records contain less
Bambusoid phytoliths than Dickau's soil samples, with less consistent *Olyrae* and *Chusquea*phytoliths. Secondly, in seasonally-inundated savannah-forest mosaic sites the lake samples contain
less (or less consistent) Heliconiaceae, Asteraceae, and Marantaceae phytoliths than Dickau's soil
samples.

474 There are several possible reasons for the differences between the assemblages from the lakes 475 studied here and the soil samples studied by Dickau. Firstly, the soil samples studied in Dickau et al. 476 (2013) are from 1-hectare plots within closed canopy and continuous forest or savannah 477 ecosystems, while the lake samples studied here are surrounded by heterogeneous landscapes. For 478 example, the dry forest signal at Lagunas La Gaiba and Mandioré may be diluted by neighbouring 479 ecosystems in the catchment such as the seasonally-inundated savnannahs of the Pantanal wetlands 480 or the upland savannah of the Amolar hills (Fig. 2). Furthermore, the lake sites from the Beni basin 481 are spread across a large area (Fig. 1). Vegetation survey data from Lagunas Limoncin and Isirere 482 (Table 3) in the south of the Beni basin shows dominance of Marantaceae (Thalia genticulata) and 483 abundance of Heliconiaceae (Heliconia sp.), but similar surveys from Lagunas La Luna and Granja in 484 the north of the Beni basin show Heliconiaceae is not present at all and Marantaceae only at Laguna 485 La Luna (Carson et al., 2016, 2015). Therefore, key phytolith types that enabled the differentiation of 486 Beni samples by Dickau et al., (2013) are not present across all Beni sites. The heterogeneity of the 487 vegetation of the Beni basin may prevent these phytolith types from being used as indicators of Beni 488 phytolith assemblages. Additional lake sites need to be studied from within these ecosystems before 489 these findings can be generalised.

Secondly, the lack of fractionation of phytolith samples in this study may have contributed to the
differences in assemblages as large-sized phytoliths such as Asteraceae platelets, Marantaceae seed
and nodular phytoliths, and hairbases, tracheids and sclerids from woody eudicots would not have

been concentrated into the C-Fraction. Previous studies have found that the C-Fraction
differentiates neotropical ecosystems more easily than the A-Fraction (Dickau et al., 2013; Watling
et al., 2016). However, due to the small size of lake sediment samples, it is often not possible to
fractionate during phytolith processing. This is a key consideration for planning of future studies of
phytoliths from lake sediment. To ensure sufficient sediment is available, it may be necessary to
collect several replicate lake cores.

499 In comparison, the pollen assemblages from the lakes studied here differentiated all three 500 ecosystems (humid evergreen forest, seasonally inundated savannah-forest mosaic and semi-501 deciduous dry forest) (Fig. 4, 5). This finding corroborates pollen trap studies of 1-ha plots where 502 these three ecosystems were differentiated by their modern pollen assemblages (Gosling et al., 503 2009, 2005; Jones et al., 2011). Fossil pollen assemblages from lakes have also revealed changes 504 between these ecosystems across south-west Amazonia (e.g. Whitney, Mayle, et al. 2013; Carson et 505 al. 2015; Carson et al. 2016). Pollen can be used to differentiate a larger number of arboreal taxa 506 than phytoliths, such as Anadenanthera and Astronium pollen, which is indicative of dry forest 507 (Gosling et al., 2009), thus enabling differentiation of different forest types. However, for 508 understorey and herbaceous vegetation, the pollen record is significantly weaker, particularly for 509 Poaceae sub-families which are challenging to differentiate using pollen alone (Julier et al., 2016). 510 Herbaceous taxa can be more effectively differentiated by their phytoliths than by their pollen; e.g. 511 Poaceae sub-families such as bamboos and oryzoid (rice) types (Piperno, 2006). Identification of 512 these taxa from the phytolith record can assist with ecosystem differentiation as well as provide important archaeological information on human activities (Hilbert et al., 2017; Watling et al., 2018, 513 514 2017). Whilst in these lakes phytolith analysis alone was not be able to differentiate all three 515 ecosystems, it has added taxonomic value; e.g., the differentiation between understorey grass taxa 516 such as bamboos within semi-deciduous dry forest versus savannah grasses from the Panicoideae 517 sub-family. This distinction allows identification of changes in upland savannah compared to changes

518 in forest understorey at lake sites like Laguna Mandioré, which would not be possible from the

519 Poaceae pollen record alone.

520 5.2 Key factors driving the composition of phytolith and pollen assemblages in lakes

521 Some of the variation between our lake samples is driven by the surrounding ecosystem, but as the 522 PCA results show (Fig. 5), this does not account for all of the variation between samples. Other 523 factors are influencing the phytolith and pollen assemblages beyond the surrounding ecosystem. In 524 order to usefully interpret the fossil phytolith assemblages from these lakes, it is necessary to 525 understand these influences. Pollen assemblages are used as a reference point for comparison with 526 the phytolith assemblages, and to highlight the relative strengths of each proxy for palaeo-vegetation 527 reconstructions. This analysis has highlighted three key factors driving the composition of phytolith 528 assemblages in lakes: productivity, taphonomy, and lake size.

529 5.2.1 Productivity

530 As with pollen, different taxa produce different quantities of phytoliths (Piperno, 2006, 1985). 531 Arecaceae are high phytolith producers (Piperno, 2006) and are therefore overrepresented in the 532 phytolith record (Aleman et al., 2014; Bremond et al., 2005). The phytolith assemblages from Laguna Chaplin within evergreen forest show Arecaceae phytolith abundance to be 53-59%, in contrast to 533 534 the vegetation inventory from a 1-ha plot in evergreen forest in Noel Kempff Mercado National Park, TF-1 (Table 4), which shows that Arecaceae accounts for only ~9% of stems. This finding 535 536 corroborates Dickau et al (2013) who also found over-representation of Arecaceae phytolith types in 537 the soils sampled from terra firme evergreen forest, but at 30% of the assemblage. In a study of 538 surface soil samples in Acre state Brazil, Watling et al. (2016) found similar Arecaceae phytolith 539 abundances in evergreen forest, but 65% in palm forest. This finding puts the Arecaceae phytolith 540 abundances found at Chaplin more closely in-line with those from palm forest than humid evergreen forest. However, additional lakes from evergreen forest would need to be sampled to clarify 541 542 whether the super-abundance of Arecaceae phytoliths is a consistent pattern across humid

543 evergreen forest lake records, or a particular feature of Laguna Chaplin due to the narrow palm544 swamp fringe at the southern shoreline.

545 In stark contrast to the phytolith record, the modern pollen assemblage from Laguna Chaplin has 546 very low Arecaceae abundance (<3%). Therefore, Arecaceae is markedly over-represented in the 547 phytolith record (53-59%) relative to the TF-1 plot vegetation inventory from humid evergreen forest 548 in NKMNP, north-eastern Bolivia (9%), but is markedly underrepresented in the pollen record (<3%). 549 These relationships clearly have an important bearing on the interpretation of Arecaceae pollen and 550 phytolith fossil records from evergreen forest, especially in the context of the on-going debate over 551 whether high abundance of Arecaceae signifies pre-Columbian forest management (Rull and 552 Montoya, 2014; Watling et al., 2017).

553 5.2.2 Taphonomy

554 It is likely that different taphonomic processes are driving the deposition of phytoliths in soil and lake sediment settings. While the phytolith assemblage in soil is likely to be dominated by phytoliths 555 556 deposited directly from plants in situ, phytoliths in lake sediments will have been transported from 557 the surrounding vegetation. The theoretical model of phytolith transport into lakes proposed by 558 Piperno (2006) suggests that in dry, open environments with frequent burning they will be 559 transported by wind (Aleman et al., 2014), but in wet, closed environments they will be transported 560 by water run-off over land or via stream and river inputs to the lake. This theory therefore suggests 561 that in dry, open environments with frequent burning phytoliths may be transported long distances 562 by wind, similarly to pollen, and therefore represent large catchment areas. However, in wet 563 environments with closed-canopy forest vegetation, where transport is primarly via water, 564 phytoliths would be likely to represent more local vegetation, depending on precipitation and 565 flooding regimes. Closed-basin lakes fed largely by water run-off over land would be likely to present 566 highly local records, whereas lakes with stream and river inputs could collect phytoliths transported 567 from greater distances.

568 The lakes studied here are from humid environments of Amazonian Bolivia, where precipitation 569 ranges from 1000 - 2000 mm/year (Seiler et al., 2013) with forest vegetation in continuous or mosaic 570 distributions (Killeen et al., 2006; Killeen and Schulenberg, 1998; Whitney et al., 2013). The RDA 571 results demonstrate that local (100 m) tree cover is a stronger driver of variation in phytolith 572 assemblages between these lakes than extra-local (1,000 m) and regional (20,000 m) tree cover (Fig. 573 6). These results suggest that phytolith records from these lakes are predominantly representing 574 vegetation at the local scale (100 m from the lake shore), with only limited inputs from extra-575 local/regional sources (1,000 – 20,000 m of the lake shore). This finding supports Piperno's theory 576 that in wet environments phytoliths are transported by water, primarily over-land. Even in the 577 seasonally flooded landscape of the Beni basin and in lakes receiving an annual flood pulse from the 578 Paraguay river (McGlue et al., 2012), our results suggest the primary method of phytolith transport 579 into the lakes is likely to be local water run-off over land with the phytolith records largely 580 representing local vegetation within 100 m of the shore. Tree cover at 1,000 – 20,000 m was found 581 to be a weaker driver of variation in the phytolith assemblages than tree cover at 100 m, but 582 nevertheless still exerted an influence (Fig. 6). The influence of this 'extra-local' vegetation may be 583 due to the seasonal flooding regimes in the flat Beni and Chiquitano-Pantanal landscapes, whereby 584 phytoliths may be transported longer distances by flood waters.

This transport via water is likely to have a bias towards smaller, lighter phytoliths over long distances, potentially reducing the number of large C Fraction sized phytoliths in lake records. This bias is also true for pollen transport within lakes, whereby larger pollen grains, such as maize, are preferentially deposited closer to the lake shore. This potential size bias in taphonomy may lead to fewer large C Fraction phytoliths in lake sediments, e.g. Asteraceae and Marantaceae which can be important for differentiating ecosystems (Dickau et al., 2013; Watling et al., 2016). Larger lakesediment samples may therefore be necessary for recovering sufficient C Fraction phytoliths.

592 This study has provided empirical evidence to support the hypothesis that pollen assemblages in 593 lake samples represent a larger catchment area than that of phytolith assemblages. This is 594 demonstrated by the pollen RDA results which show that the key environmental variables driving 595 the differences between pollen assemblages across lake sites in our study region are lake area and 596 tree cover at 20,000 m (Fig. 6). This suggests that pollen records most strongly represent vegetation 597 at a regional scale (20,000 m from lake shore). This is in alignment with studies of pollen taphonomy 598 and catchment area (Bunting et al., 2004; Sugita, 1994), which demonstrate that while other factors 599 such as pollen productivity, grain fall speed, atmospheric turbulence and wind speed have an 600 influence, pollen records from large lakes generally represent larger catchment areas. This is 601 supported by the co-correlation of lake area and tree cover at 1,000 - 20,000 m in the pollen RDA 602 plot (Fig. 6). Pollen records are therefore most strongly influenced by regional vegetation (20,000 m) 603 scales, while phytolith records are most strongly influenced by local (100 m) vegetation.

604

605 5.2.3 Lake Size

606 The lakes studied here encompass a wide range of sizes, from Laguna Granja at 0.071 km² to Laguna 607 Mandioré at 152 km² in area. The RDA results (Fig. 6) demonstrate that lake size does influence 608 phytolith assemblage composition, although it is not statistically significant at the 95 % confidence 609 level (p=0.059). By contrast, lake size has a statistically significant influence on the pollen 610 assemblage (p=0.001). The strong influence of local (100 m) vegetation on the phytolith assemblage 611 would suggest that, in larger lakes in particular, the assemblage will be strongly influenced by the 612 vegetation type near to the sample site. This can be seen in the within-lake variation in phytolith 613 assemblages from the large lakes Lagunas Mandioré, La Gaiba and Chaplin, reflecting differences in 614 local vegetation around the lake.

For example, Mandioré sample 5 is located in the north-east of the lake basin, close to the Amolar
upland savannah which contains a high proportion of Panicoid taxa. The phytolith record reflects this

617 with a higher Panicoid abundance and lower Bambusoid and arboreal abundance. Mandioré 2 is 618 located in the south-west of the basin surrounded by seasonally dry tropical forest, which is 619 reflected in the higher Bambusoid and arboreal phytolith abundance at this core site. At La Gaiba, 620 the differences in assemblage between core sites are related to the proximity to the Pantanal 621 wetlands. La Gaiba 6 is in the centre of the south basin, where the surrounding lake shores are 622 dominated by semi-deciduous forest. This sample position is reflected in the higher arboreal 623 phytolith abundances and lower grass and herb abundances. Towards the north end of the south 624 basin (La Gaiba 10) and into the north basin (La Gaiba 16), the semi-deciduous forest merges into the 625 Pantanal wetlands. These wetlands likely contribute to the higher grass and herb phytolith totals 626 found in these two samples. At Chaplin, the core site closer to the shore (Chaplin 2) shows a higher 627 abundance of Poaceae and lower arboreal phytoliths than the more central site (Chaplin 1), possibly 628 reflecting greater inputs from local vegetation, including an area of savannah marsh at the south 629 west shoreline.

630 For these three large lakes, the level of within-lake spatial variation is greater in the phytolith 631 assemblages than the pollen assemblages, as shown by the groupings in the PCA results (Fig. 5). 632 While some differentiation in the pollen assemblage can be seen at La Gaiba depending on proximity 633 to the Pantanal wetlands, it is not as strong as the variation in the phytolith assemblage (Fig. 4, 5). 634 This supports the finding that phytolith assemblages are more heavily influenced by local vegetation 635 than pollen assemblages. The different spatial scales that these two vegetation proxies represent 636 provide a valuable opportunity for enhancing the ecological detail that can be extracted from palaeoecological records. Not only do the two proxies provide complementary taxonomic 637 638 information, but they also provide vegetation information at complementary spatial resolution. A 639 sediment core from a large lake can therefore be used to reconstruct regional vegetation using the 640 pollen record, and local vegetation using the phytolith record.

641

642 6. Conclusions

Phytolith analysis of lake sediments, particularly when integrated with pollen analysis, has
demonstrated potential for enhancing the ecological detail in Neotropical palaeo-vegetation
reconstructions.

1. It is possible to differentiate evergreen forest from semi-deciduous forest and Beni seasonally
inundated savannah- forest mosaic using the phytolith assemblage from lake samples alone, largely
based on the abundance of Arecaceae phytolith types. It was not possible in this study to
differentiate semi-deciduous forest from seasonally inundated savannah- evergreen forest mosaics
from the Beni Basin using phytoliths alone.

2. Empirical analysis demonstrated that in the ecosystems studied here, phytolith assemblages in
lake sediment records are most strongly influenced by local (within 100 m of the lake shore)

vegetation and therefore likely to be transported via water run-off over land rather than long-

distance travel via wind. This leads to significant spatial variation in phytolith assemblages within

large lakes which have a diversity of vegetation types in their local catchment. Pollen and phytoliths

656 from lake sediment samples from large lakes therefore represent different spatial scales: pollen

657 represents extra-local or regional vegetation; phytoliths represent local or shoreline vegetation.

3. Phytoliths from lake sediment core samples can reveal the history of local vegetation. Therefore,
sediment cores from large lakes can be used to assess the regional vegetation using pollen, and the
local vegetation using phytoliths. However, while a pollen record from a single lake core is sufficient
to record the regional vegetation, multiple cores across large lakes are likely to be needed for
phytolith analysis to adequately capture spatial variation in local/shore-line vegetation around the
lake.

664 *6.1 Future work*

To build our understanding of the spatial scale phytoliths from lakes represent under different
 environmental conditions, further studies on phytolith taphonomy are needed. Mechanistic and

667 experimental studies of phytolith taphonomy would provide a solid basis for interpretation of 668 observations from field studies. Additionally, this study was limited to only three ecosystems, not 669 including upland terra firme savannah. The influence of long-distance dispersal by wind in open 670 savannah-dominated, drier environments may have a significant impact on the spatial scale the 671 phytolith records represent (Aleman et al., 2014). Also, the specific characteristics of the lake sites 672 chosen for this study will have affected the results. For example, the proximity of the Pantanal 673 wetlands to both semi-deciduous dry forest lake sites is likely to have influenced the results. 674 Furthermore, as only one lake - Chaplin - from within humid evergreen forest was studied, it is 675 difficult to generalise to all lake records within this ecosystem. For example, if a study area of humid 676 evergreen forest does not contain palms, it may not be possible to differentiate it from other forest types using phytoliths alone. Therefore, further studies including additional ecosystems and greater 677 678 numbers of lakes is needed to confirm these findings.

679

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- 863 192. https://doi.org/10.1016/j.palaeo.2011.05.012

864 List of Figures



- 866 Figure 1 Map of lake sites and vegetation surveys plotted against vegetation classifications adapted
- 867 from WWF terrestrial ecoregions (Olsen et al 2001) using ArcGIS 10.5.1. Lake sites are shown by red
- 868 circles. Vegetation inventories and surveys are shown by black circles (IL = Vegetation survey around
- 869 Lagunas Isirere and Limoncin, AC2 = Acuario 2 forest plot inventory, LF1 = Los Fierros 1 forest plot
- 870 inventory). Inset map of South America.



872 Figure 2 - Maps of lake sites and surrounding ecosystems created using ArcGIS 10.5.1. Panel A shows the Chiquitania-Pantanal study region with Lagunas La Gaiba and Mandioré; Panel B shows terra 873 874 firme humid evergreen forests on the Pre-Cambrian shield study region with Laguna Chaplin; Panel C 875 shows the northern areas of the Beni basin study region with Lagunas Oricoré, La Luna and Granja; 876 and Panel D shows the southern areas of the Beni basin study region with Lagunas San José and 877 Limoncin. The locations of all surface samples are shown for each lake by black circles. Vegetation 878 classification of the study area, based on Landsat imagery, was provided by the Museo de Historia Natural 'Noel Kempff Mercado', Santa Cruz, Bolivia in 2015. Scale bars and latitude and longitude are 879 880 presented for each individual map panel.

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891 percentage data. Vegetation surrounding the lakes has been classified into 3 ecosystem types: semi-

- deciduous dry forest (Chiquitania-Pantanal), seasonally inundated forest-savannah (Beni basin), and
- 893 evergreen forest (PCS Humid Evergreen Forest).





Figure 4 – Summary diagram of pollen abundance from all lakes studied, presented as percentage of
terrestrial total. Vegetation surrounding the lakes has been classified into 3 ecosystem types: semideciduous dry forest (Chiquitania-Pantanal), seasonally inundated forest-savannah (Beni basin), and
evergreen forest (PCS Humid Evergreen Forest). Pollen data for Chaplin 1 and 2 is only available in
highly summarised format with data for a restricted number of taxa. Full pollen counts are available
for Chaplin 3.



Figure 5 – PCA biplot for a) phytolith and b) pollen data. Axes are Principal Component 1 (PCA1) and
Principal Component 2 (PCA2). The percentage in parentheses on these axes is the percentage of
variance in the dataset explained by that principal component. Lake sites are coloured according to
the ecosystem they represent: light green = semi-deciduous dry forest, yellow = Beni seasonally-

907 inundated forest-savannah mosaic, dark green = humid evergreen forest. Taxa are presented in

908 black. Codes for taxa names are presented in Tables 5 and 6 for phytoliths and pollen respectively.





911 Figure 6 – RDA tri-plot for a) phytolith and b) pollen data. Axes are Redundancy Analysis 1 (RDA1)

and 2 (RDA2). The percentage in parentheses on these axes is the percentage of variance in the

913 dataset explained by that component. Environmental constraining variables are presented in blue:

914 lake area, distance of core site to lake shore, and tree cover within 100, 1000, 2000, 10,000 and

- 915 20,000 m of the lake shore. Lake sites are coloured according to the ecosystem they represent: light
- 916 green = semi-deciduous dry forest, yellow = Beni seasonally-inundated forest-savannah mosaic, dark
- green = humid evergreen forest. Taxa are presented in black. Codes for taxa names are presented in
- 918 Tables 5 and 6 for phytoliths and pollen respectively.
- 919
- 920 List of Tables

921 Table 1 - Summary of lake site characteristics, including references to the papers originally publishing922 some of the pollen and phytolith records.

Lake	No. samples	Region	Ecosystem	Area of lake /km²	Pollen analysed by	Phytoliths analysed by
Mandioré	2	Chiquitania- Pantanal	Semi-deciduous forest/Pantanal wetlands	152	Plumpton et al., accepted	Plumpton et al., accepted
La Gaiba	3	Chiquitania- Pantanal	Semi-deciduous forest/Pantanal wetlands	90	Whitney et al., 2011	Plumpton et al., accepted
Oricoré	1	Beni Basin	Seasonally- inundated savannah-forest mosaic	10.5	Carson et al., 2014	This study
La Luna	1	Beni Basin	Seasonally- inundated savannah-forest mosaic	0.33	Carson et al., 2016	This study
Limoncin	1	Beni Basin	Seasonally- inundated savannah-forest mosaic	0.73	Whitney, unpublished	This study

San José	1	Beni Basin	Seasonally-	14.3	Whitney et	Whitney et
			savannah-forest		al., 2015	al., 2015
			mosaic			
Granja	1	Beni basin/	Seasonally-	0.071	Carson et	Carson et
		Terra firme	inundated		al., 2015	al., 2015
		evergreen	savannah-forest			
		forest on PCS	mosaic /Terra			
			firme evergreen			
			forest			
Chaplin	2	Terra firme	Terra firme	12.2	Burbridge	This study
		evergreen	evergreen forest		et al., 2004	
		forest on PCS				

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926 Table 2 – Vegetation inventory of Acuario 2, a 1-hectare vegetation plot within Noel Kempff

927 Mercado National Park, gives a representative vegetation community composition for semi-

928 deciduous dry forest in south-west Amazonia. Inventory conducted by recording all taxa

929 representing >1% of the total number of stems >10cm d.b.h. (Gosling et al., 2009).

Family	Species	% of total stems
Fabaceae — Caes.	Caesalpinia floribunda Tul.	11.72
Bignoniaceae	Tabebuia roseo-alba (Ridley) Sandwith	7.42
Fabaceae — Mim.	Anadenanthera colubrina (Vell.) Brenan	7.03
Flacourtiaceae	Casearia gossypiosperma Brig.	6.25
Combretaceae	Combretum leprosum Mart.	5.66
Arecaceae	Orbignya phalerata Mart.	3.71
Fabaceae — Caes.	Bauhinia rufa (Bong.) Steud.	3.32
Rubiaceae	Simira cordifolia (Hook. f.) Steyerm.	2.93
Boraginaceae	Cordia alliodora (Ruíz and Pavón) Oken	2.73
Sterculiaceae	<i>Guazuma ulmifolia</i> Lam.	2.73
Apocynaceae	Aspidosperma cylindrocarpon Müll. Arg.	2.34
Sterculiaceae	<i>Sterculia apetala</i> (Jacq.) Karsten	2.15
Flacourtiaceae	<i>Casearia arborea</i> (Rich.) Urban	1.95
Rhamnaceae	Rhamnidium elaeocarpum Reisseck	1.95
Arecaceae	Scheelea princeps (Mart.) Karsten	1.76
Malvaceae	Chorisia integrifolia Ulbr.	1.76
Meliaceae	Cedrela fissilis Vell.	1.76

Sapindaceae	Dilodendron bipinnatum Radlk.	1.76
Anacardiaceae	Spondias mombin L.	1.56
Bignoniaceae	Arrabidea spicata Bureau and K. Schum	1.37
Euphorbiaceae	Sebastiana huallagensis Croizat	1.37
Tiliaceae	Apeiba tibourbou Aubl.	1.37
Malvaceae	Pseudobombax marginatum (A. StHil.) Robyns	1.17
Fabaceae — Pap.	Machaerium villosum Vogel	1.17
Malpighiaceae	Dicella macroptera A. Juss.	1.17
Fabaceae — Pap.	Machaerium acutifolium Vogel	0.98
Tiliaceae	Triumfetta grandiflora Vahl	0.98
TOTAL		80.08

930

931 Table 3 – Results of a qualitative vegetation survey ranking taxa as dominant, abundant, frequent or

932 occasional in coverage from the area surrounding Lagunas Limoncin and Isirere are presented to give

933 an example vegetation community composition for the Beni seasonally-inundated savannah. (Dickau

934 et al., 2013).

Family	Species	Abundance classification (Soto 2010)
Marantaceae	Thalia geniculata	Dominant (>50%)
Fabaceae	Inga stenopoda	Dominant (>50%)
Fabaceae	Erythrina fusca	Abundant (20-50%)
Cyperaceae	Cyperus gigantus	Abundant (20-50%)
Heliconiaceae	Heliconia sp.	Abundant (20-50%)
Typhaceae	Typha dominguensis	Frequent (10-20%)
Moraceae	Ficus sp.	Frequent (10-20%)
Cannaceae	Canna glauca	Occasional (2-10%)
Urticaceae	Cecropia sp.	Occasional (2-10%)

- 936 Table 4 Vegetation inventory of Los Fierros 1, a 1-hectare vegetation plot within Noel Kempff
- 937 Mercado National Park, gives a representative vegetation community composition for evergreen
- 938 forest in south-west Amazonia. Inventory conducted by recording all taxa representing >1% of the
- total number of stems >10cm d.b.h. (Gosling et al., 2005).

Family	Species	% of total
		stems
Strelitziaceae	Phenakospermum guianensis Aubl.	13.07
Moraceae	Pseudolmedia laevis (Ruiz and Pav.) J. F. Macbr.	7.64

Arecaceae	Euterpe precatoria Mart.	6.91
Rubiaceae	Capirona decorticans Spruce	6.29
Vochysiaceae	<i>Qualea paraenesis</i> Ducke	6.04
Vochysiaceae	Erisma uncinatum Warm.	4.19
Elaeocarpaceae	Sloanea eichleri K. Schum.	3.45
Moraceae	Pseudolmedia macrophylla Trécul	3.33
Rubiaceae	Amaioua guianensis Aubl.	3.08
Hippocrateaceae	Cheiloclinium cognatum (Miers) A. C. Sm.	2.84
Euphorbiaceae	Hyeronima oblonga (Tul.) Müll. Arg.	2.47
Arecaceae	Socratea exorrhiza (Mart.) H. L. Wendl.	2.1
Moraceae	Pourouma guianensis Aubl.	1.85
Melastomataceae	Miconia pyrifolia Naudin	1.6
Moraceae	Brosimum acutifolium subsp. obovatum (Ducke) C. C. Berg.	1.6
Lythraceae	Physocalymma scaberrimun Pohl	1.11
Melastomataceae	indet. 3	1.11
Lauraceae	Nectandra sp. 2	0.99
Melastomataceae	Miconia multiflora Cogn.	0.99
Melastomataceae	Miconia sp. 3	0.99
Moraceae	Helicostylis tomentosa (Poepp. and Endl.) Rusby	0.99
TOTAL		72.63

940

- 941 Table 5 Phytoliths types identified with abundance >1%, their taxonomic association and PCA/RDA
- 942 codes.

Phytolith type	Association	References	PCA/RDA code
Bilobates	Panicoideae	1–6	PanBilob
Polylobates	Panicoideae	1–6	PanPolyb
Crosses	Panicoideae	5–12	PanCross
Squat saddles	Chloridoideae	2, 3, 5, 6, 13	Chloro
Aristida bilobates	Aristidoideae	5, 6	Arist
Rondels	Poaceae	2, 3, 5, 6	Rond
Rondeloid/saddeloid	Bambusoideae	5	BambRond
Collapsed saddles	Bambusoideae	5–7, 14, 15	BambCSaddle
Tall saddles	Bambusoideae	6	BambTSaddle
Bilobates (blocky)	Bambusoideae	6	BambBilob
Crosses (blocky)	Bambusoideae	5–12	BambCross
Chusquoid bodies	Bambusoideae	5,6	BambChusquoid
Two-spiked crown bodies	Bambusoideae	5	BambCrown
Chusquea bodies	Bambusoideae	5,6	BambChusquea
Oryzeae scooped bilobates	Oryzeae	1, 16	OryzBilob
Oryzeae scooped crosses	Oryzeae	1, 16	OryzCross
Olyreae bodies	Olyreae	5, 17	BambOlyra
Bulliforms	Poaceae	6, 7	Bull

				-
Bulliforms (bamb)	Bambusoideae	35	BambBull	
Cyperaceae cones	Cyperaceae	13, 20–23	CypCone	
Scirpus achene	Cyperaceae	20	CypScir	
Cyperus/Carex achene	Cyperaceae	20	СурСур	
Heliconia troughed body	Heliconia	6, 18, 19	Hel	
Marantaceae globular nodular	Marantaceae	23	MarSpher	
Marantaceae seed	Marantaceae	23	MarSeed	
Strelitziaceae druse	Strelitziaceae	18	Strel	
Echinate globular/hat	Arecaceae	6, 13, 17, 18, 24,	Palm	
		25		
Echinate irregular platelet	Celtis	17, 28	Celtis	
Globular granulate	Woody eudicot	6, 26, 27	GlobGran	
Faceted elongate	Woody eudicot	6	Arbor	
Terminal tracheid	Woody eudicot	6, 7	TermTrach	
Asteraceae platelets	Asteraceae	6, 7, 32	Ast	
Vesicular infillings	Woody eudicot	29, 30	VesFill	
Scooped globular	Pteridophyte, <i>Trichomanes</i>	31	PterGlob	

943 References: 1. (Metcalfe 1960); 2. (Twiss et al. 1969); 3. (Brown 1984); 4. (Fredlund & Tieszen 1994);

944 5. (Piperno & Pearsall 1998); 6. (Piperno 2006); 7. (Piperno 1988); 8. (Piperno 1984); 9. (Pearsall

- 945 1978); 10. (Pearsall 1982); 11. (Pearsall & Piperno 1990); 12. (Iriarte 2003); 13. (Kondo et al. 1994);
- 946 14. (Lu 1995); 15. (Lu et al. 2006); 16. (Chaffey 1983); 17. (Watling & Iriarte 2013); 18. (Tomlinson
- 947 1961); 19. (Prychid et al. 2003); 20. (Ollendorf 1992); 21. (Honaine et al. 2009); 22. (Metcalfe 1971);
- 948 23. (Wallis 2003); 24. (Runge 1999); 25. (Bozarth et al. 2009); 26. (Amos 1952); 27. (Scurfield et al.
- 949 1974); 28. (Bozarth 1992); 29. (Stromberg 2003); 30. (Strömberg 2004); 31. (Mazumdar 2011).
- 950 Table 6 Pollen taxa identified with abundance >1% and PCA/RDA codes.

Family	Genus or species	PCA/RDA code
Amaranthaceae	Alternanthera	AmarAlt
Amaranthaceae	Amaranthus/Chenopodiaceae	AmarAma
Amaranthaceae	Gomphrena	AmarGom
Anacardiaceae	Astronium	AnacAst
Anacardiaceae	Schinopsis	AnacSch
Anacardiaceae	Spondias	AnacSpo
Anacardiaceae	Tapirira	AnacTap
Annonaceae	Annona	AnnonAnn
Apocynaceae	Prestonia	ApoPres
Araliaceae	Didymopanax	AralDid
Arecaceae	undiff.	PalmUnd

Arecaceae	Astrocaryum	PalmAst
Arecaceae	Copernicia	PalmCop
Arecaceae	Mauritia	PalmMaur
Arecaceae	Sygarus	PalmSyg
Asteraceae	Mikania-type	AstMik
Asteraceae	undiff.	AstUnd
Asteraceae	Ambrosia-type	AstAmb
Bignoniaceae	Jacaranda	BigJac
Bromeliaceae	undiff.	Brom
Burseraceae	Bursera-type	BurBurs
Cannabaceae	Celtis	CanCelt
Cannabaceae	Trema	CanTrem
Combretaceae/Melastomataceae	undiff.	CombMelUnd
Melastomataceae	Miconia	CombMelMic
Cyperaceae	undiff.	Сур
Dilleniaceae	Curatella americana	DillCur
Erythroxylaceae	Erthroxylum	EryEryth
Euphorbiaceae	Acalypha	EuphAca
Euphorbiaceae	Alchornea	EuphAlch
Euphorbiaceae	Hura-type	EuphHura
Euphorbiaceae	Sapium	EuphSap
Euphorbiaceae	Asparisthium	EuphAsp
Fabaceae	Copaifera	FabCopa
Fabaceae	Macrolobium	FabMacr
Fabaceae	Apuleia leiocarpa	FabApul
Fabaceae	Dalbergia/Macherium	FabDalMach
Fabaceae	Erythrina	FabEryth
Fabaceae	Senna-type	FabSen
Fabaceae	undiff.	FabUnd
Fabaceae	Acacia	FabAcac
Fabaceae	Anadenanthera	FabAnad
Fabaceae	Inga	FabInga
Fabaceae	Mimosa	FabMimo
Lamiaceae	Hyptis	LamHyp
Lamiaceae	<i>Vitex</i> -type	LamVit
Malpighiaceae	Byrsonima	MalpBrys
Malpighiaceae	"periporate"	MalpPeri
Malvaceae	<i>Bytternia-</i> type	MalvBytt
Malvaceae	undiff.	MalvUnd
Malvaceae	<i>Guazuma</i> -type	MalvGuaz
Meliaceae	Cedrela/Trichilia	MeliCedTri
Moraceae/Urticaceae	undiff.	MorUrtUnd
Moraceae	Brosimum	MorBros
Moraceae	Ficus	MorFic
Moraceae	Helicostylis	MorHeli
Moraceae	Maclura	MorMacl

Moraceae	Maquira	MorMaq
Moraceae/Urticaceae	Pourouma/Sorocea	MorUrtPourSor
Moraceae	Pseudolmedia	MorPsued
Urticaceae	Cecropia	UrtCecr
Myrtaceae	undiff.	Myrt
Phytolaccaceae	Gallesia	PhyGal
Phyllanthaceae	Amanoa	PhylAma
Piperaceae	Piper	PipPiper
Poaceae	undiff.	Роас
Polygonaceae	Symmeria	PolySym
Polygonaceae	Triplaris	PolyTrip
Rubiaceae	Borreria "pericolporate"	RubBorrPeri
Rubiaceae	Borreria latifolia	RubBorrLat
Rubiaceae	<i>Borreria</i> "undiff."	RubBorrUnd
Rubiaceae	Faramea	RubFar
Rubiaceae	Uncaria	RubUnc
Rubiaceae	undiff.	RubUnd
Saliaceae	undiff.	Sali
Sapindaceae	undiff.	SapinUnd
Sapindaceae	Dilodendron	SapinDilo
Sapindaceae	Talisia	SapinTal
Sapotaceae/Melastomataceae	undiff.	SapotMel
Sapotaceae	Pouteria	SapotPout
Solanaceae	undiff.	Solan
Ulmaceae	Ampeloera-type	UlmAmp
Ulmaceae	Phyllostylon	UlmPhyll
Vitaceae	Cissus	VitCis
Vochysiaceae	Vochysia	VocVochy
"Unknowns"	"Unknowns"	Unkno
Alismataceae	Sagittaria	AlisSagg
Alismataceae	Echinodorus	AlisEchin
Pontederiaceae	Eichhornia	PontEich
Polygonaceae	Polygonum	PolyPolyg
Selaginellaceae	Selaginella	SelSelag
Typhaceae	Typha	TypTypha
Isoetes	undiff.	Isoetes
Onagraceae	Ludwigia	OnagLud
Fern	Parkeriaceae	PterPark
Fern	Polypodium	PterPoly

951

952 <u>Supplementary Information</u>

953 S1 – Results of permutation test on RDA results for pollen.

```
Permutation test for rda under reduced model
954
955
      Terms added sequentially (first to last)
      Permutation: free
956
      Number of permutations: 999
957
958
959
      Model: rda(formula = pollen.trim.sq ~ LakeArea + DistanceShore + treecover
      100m + treecover1000m + treecover5000m + treecover10000m + treecover20000m
960
      , data = pollen.env)
961
                      Df Variance
962
                                         F Pr(>F)
                                            0.001 ***
                            7.8826 11.6825
963
      LakeArea
                        1
964
      DistanceShore
                        1
                            1.4076
                                    2.0861
                                            0.067
                                            0.001 ***
965
      treecover100m
                            3.2239
                                    4.7780
                       1
                                            0.043 *
966
      treecover1000m
                       1
                            1.6132
                                    2.3909
                                            0.035 *
967
      treecover5000m
                       1
                            1.8427
                                    2.7309
      treecover10000m
                                            0.015 *
968
                       1
                            2.2946
                                    3.4007
                                            0.001 ***
969
      treecover20000m
                       1
                            4.8210
                                    7.1450
970
      Residual
                        3
                            2.0242
971
      Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
972
973
974
      S2 – Results of permutation test on RDA results for phytoliths.
975
      Permutation test for rda under reduced model
      Terms added sequentially (first to last)
976
977
      Permutation: free
978
      Number of permutations: 999
979
      Model: rda(formula = phyto.trim.sq ~ LakeArea + DistanceShore + treecover1
980
981
      00m + treecover1000m + treecover5000m + treecover10000m + treecover20000m,
982
      data = phyto.env)
                      Df Variance
983
                                        F Pr(>F)
                            2.6529 2.4699
984
      LakeArea
                       1
                                           0.059 .
985
      DistanceShore
                        1
                            2.5158 2.3422
                                           0.063
                                           0.001 ***
986
      treecover100m
                       1
                            6.1033 5.6822
                                           0.059
987
      treecover1000m
                       1
                            2.3299 2.1692
                                           0.003 **
988
                       1
                            4.9024 4.5642
      treecover5000m
989
                       1
                                           0.098 .
      treecover10000m
                            2.1535 2.0050
990
                       1
      treecover20000m
                            1.5547 1.4474
                                           0.201
991
      Residual
                        4
                            4.2964
992
      Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
993
994
```