



1 **The climate and vegetation of Europe, North Africa and the**  
2 **Middle East during the Last Glacial Maximum**  
3 **(21,000 years BP) based on pollen data**

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15

16 **Abstract.** Pollen data represents one of the most widely available and spatially-resolved sources of information  
17 about the past land cover and climate of the Last Glacial Maximum (21,000 years BP). Previous pollen data  
18 compilations for Europe, the Mediterranean and the Middle East however have been limited by small numbers of  
19 sites and poor dating control. Here we present a new compilation of pollen data from the region that improves on  
20 both the number of sites (63) and the quality of the chronological control. Data has been sourced from both public  
21 data archives and published (digitized) diagrams. Analysis is presented based on a standardized pollen taxonomy  
22 and sum, with maps shown for the major pollen taxa, biomes and total arboreal pollen, as well as quantitative  
23 reconstructions of forest cover and winter, summer and annual temperatures and precipitation. The reconstructions  
24 are based on the modern analogue technique (MAT) with a modern calibration dataset taken from the latest  
25 Eurasian Modern Pollen Database (~8000 samples). A site-by-site comparison of MAT and Inverse Modelling  
26 methods shows little or no significant difference between the methods for the LGM, indicating that no-modern-  
27 analogue and low CO<sub>2</sub> conditions during the LGM do not appear to have had a major effect on MAT transfer  
28 function performance. Previous pollen-based climate reconstructions based on MAT show a much colder and  
29 drier climate for the LGM than both Inverse Modelling and climate model simulations, but our new results suggest  
30 much greater agreement. Differences between our latest MAT reconstruction and those in earlier studies can be  
31 largely attributed to bias in the small modern calibration dataset previously used. We also find that quantitative  
32 forest cover reconstructions show more forest than that previously suggested by biome reconstructions, but less  
33 forest than that suggested by simple percentage arboreal pollen, although uncertainties remain large. Overall, we  
34 find that LGM climatic cooling/drying was significantly greater in winter than in summer, but with large site to  
35 site variance that emphasizes the importance of topography and other local factors in controlling the climate and  
36 vegetation of the LGM.

37



38 **1 Introduction**

39

40 During the Last Glacial Maximum (LGM) ~21,000 years BP (Mix et al., 2001), the climate, vegetation and  
41 landscape of Europe and its surrounding regions were very different than today. Scandinavia and a large part of  
42 the British Isles were covered by a single ice sheet, with separate ice sheets covering the Alps and Pyrenees, while  
43 many smaller and lower mountainous areas were also glaciated. As a result of this global build-up of ice on land,  
44 sea levels were around 120 meters lower than today, resulting in the retreat of Atlantic and Mediterranean  
45 coastlines and the emergence on land of the English Channel and North Sea basin. Falling sea levels also led to  
46 the disconnection of the Black Sea from the Mediterranean, and a subsequent drop in Black Sea water levels as  
47 evaporation exceeded inflow. On land, permafrost and periglacial processes occurred immediately to the south of  
48 the Scandinavian ice sheet, while the massive discharge of glacial clays and sands provided material to be  
49 redeposited by the wind as belts of loess across northern France, Benelux, Germany and central Europe. Under  
50 these cooler and drier climatic conditions, forests are thought to have retreated to the relative shelter of Southern  
51 Europe and the Mediterranean, while relatively unproductive steppe and boreal scrub dominated the region north  
52 of the Alps.

53

54 This traditional view of the LGM has been established for many years, but many details concerning the climate  
55 and vegetation of the LGM remain debated. Much of this debate concerns information derived from the pollen  
56 record, which represents one of the most widely available and spatially-resolved sources of information  
57 concerning LGM vegetation and climate, and the primary terrestrial proxy used to evaluate climate models in the  
58 Palaeo-model Inter-comparison Project (PMIP) (Bartlein et al., 2011; Harrison et al., 2014).

59

60 For example, climate model simulations continue to indicate a climate that is less cold and more humid than  
61 pollen-based reconstructions (Jost et al., 2005). These pollen climate reconstructions are similar to those based on  
62 glaciological modelling (Allen et al., 2008a). On the other hand, the pollen-based reconstructions that show the  
63 greatest disagreement with climate models have themselves been criticized for not considering the possible effect  
64 of low atmospheric CO<sub>2</sub> on the physiological relationship between plants and climate (Ramstein et al., 2007).  
65 Pollen-climate reconstructions based on inverse modelling that account for these low CO<sub>2</sub> effects show less  
66 cooling and drying and consequently greater agreement with climate models (Ramstein et al., 2007; Wu et al.,  
67 2007).

68

69 Further data-model discrepancies have also been highlighted concerning LGM vegetation cover. Earlier pollen  
70 synthesis studies, especially those that applied the biomisation method (Elenga et al., 2000) give the impression  
71 that non-glaciated areas of LGM Europe were dominated by treeless steppe, while vegetation models driven by  
72 climate model simulations indicate large areas of forest and woodlands (Binney et al., 2017; Kaplan et al., 2016;  
73 Velasquez et al., 2021). The apparent data-model discrepancy associated with steppe has led to the suggestion  
74 that early humans, which are not included in vegetation models, could have reduced the forest cover with only a  
75 relatively moderate use of fire because of the cold climate and slow speed of vegetation recovery (Kaplan et al.,  
76 2016). This debate is important because of studies that have shown the sensitivity of the climate system to  
77 vegetation boundary conditions during the LGM (Ludwig et al., 2017; Velasquez et al., 2021). This suggests that



78 accurate knowledge of the vegetation cover during the LGM is a necessary prerequisite to understanding the role  
79 of other influences on the climate system at this time.

80

81 More recent pollen and macrofossil studies from eastern Central Europe have shown that at least in this region  
82 there existed areas of open boreal forest and woodland with some temperate broadleaf species (Kuneš et al., 2008;  
83 Willis and Van Andel, 2004). The evidence of forest, and particularly elements of temperate broadleaf forest,  
84 north of the Alps has come to represent a challenge to the traditional view that forest species only survived the  
85 LGM in sheltered refugia far to the south of the Fenoscandian ice sheet and close to the moderating influence of  
86 the Mediterranean Sea. The presence of micro-refugia north of the Alps is important because it would represent a  
87 very different baseline for understanding the later rate and route of plant migrations under the rapid warming that  
88 occurred during the Late Glacial to Holocene transition (Douda et al., 2014; Giesecke, 2016; Krebs et al., 2019;  
89 Nolan et al., 2018), as well as understanding patterns of present-day genetic diversity (Normand et al., 2011;  
90 Svenning et al., 2008).

91

92 Much of this debate has been informed by an increasing number of LGM pollen studies from an ever-broader  
93 geographical area, and especially from an increasing number of studies from north of the Alps. Nevertheless, the  
94 synthesis of these studies into a single narrative is made difficult by several factors, for instance: different  
95 taxonomic definitions, pollen percentages calculated from non-standardized pollen sums, and quantitative  
96 analyses such as climate reconstructions that are based on different training sets and methodologies. This has led  
97 to some modelling studies ignoring the pollen record completely, on the basis that data from the LGM is too scarce  
98 (Janská et al., 2017). Where standardized methods have been applied to multiple LGM pollen records, poor dating  
99 control has resulted in the inclusion of many records that may not actually be from the selected LGM time window.  
100 This is particularly important because the  $21 \pm 2.0$  ka time slice commonly used to represent the LGM period in  
101 PMIP data-model comparisons and other synthesis studies (MARGO members, 2009; Bartlein et al., 2011) occurs  
102 immediately after the glacial maxima in the Alps, which occurs around 26-23 ka (Heiri et al., 2014; Spötl et al.,  
103 2021), and is therefore likely to be represented by a different vegetation and climate.

104

105 For example, of the 18 European pollen records used in the PMIP benchmarking dataset (Bartlein et al., 2011),  
106 10 fall into the worst class ('poor') in the COHMAP chronological quality classification scheme if relative dating  
107 such as pollen correlation is excluded. More recent synthesis studies have also relied heavily on records from the  
108 European Pollen Database (EPD) which currently has 116 records with samples of LGM age (as of June 2022).  
109 Many of these records however are based on chronologies that are considered reliable for the Holocene (Giesecke  
110 et al., 2014), but have large uncertainties for the LGM as a result of 1) excessive extrapolation back in time from  
111 Holocene age dates, 2) the use of pollen correlation or other relative dating despite poorly defined regional  
112 biostratigraphy, or 3) the inappropriate use of radiocarbon dates contaminated with old carbon. We found that 104  
113 of these 116 EPD records (Neotoma, 2021) fall into the worst class ('poor') in the COHMAP chronological quality  
114 classification.

115

116 Here we address these problems using a new synthesis of LGM pollen records from throughout Europe, the  
117 Mediterranean and the Middle East (EurMedMidEst) based on rigorous quality control criteria. Records were



118 compiled from an extensive review of public databases and archives, and the scientific literature. Pollen records  
119 were selected according to the robustness of their chronological control around the PMIP LGM time-window ( $21$   
120  $\pm 2$  ka), and combined into a single dataset based on a harmonized taxonomy and standardized pollen sum. The  
121 dataset was then analysed so that standardised maps could be produced to show the distribution of the major pollen  
122 taxa, biomes and total arboreal pollen at the LGM. In addition, quantitative reconstructions of forest cover as well  
123 as winter, summer and annual temperatures and precipitation were undertaken using the Modern Analogue  
124 Technique (MAT), utilizing the latest Eurasian Modern Pollen Database II calibration dataset. These climate  
125 reconstructions are compared and evaluated against previous LGM pollen-climate reconstructions, as well as  
126 reconstructions based on other proxies. The dataset and results are fully documented and the complete data files  
127 are provided in the supplementary information.

128

## 129 **2 Methods**

130

### 131 **2.1 Pollen Data**

132

133 LGM fossil pollen data from Europe and bordering regions including North Africa and the Middle East were  
134 selected and collated into a single standardized project database. This data was sourced from the EPD/Neotoma  
135 database (Williams et al., 2021), the Pangaea data archive, publications in scientific journals, and from the original  
136 authors. We selected LGM pollen sites/data according to strict quality control criteria. Where possible, primary  
137 raw pollen counts were used where this was available. Where the original electronic data was not available, the  
138 data was digitized from the published diagram. Overall, 35 out of 63 records were digitized, while the rest of the  
139 data consisted of raw pollen counts (Table 1).

140

141 The distribution of sites included in our study reflects the distribution of suitable archives, with fewer records  
142 available from climatically or environmentally challenging regions (Fig. 1). High rates of erosion and a drier and  
143 colder climate during the LGM reduced the number of suitable anoxic sediment sinks for pollen preservation,  
144 especially in Central Europe between the Scandinavian and Alpine ice sheets. Lakes sites are the most numerous  
145 archive and tend to be located in the more sheltered and topographically favourable regions of Southern Europe  
146 and the Mediterranean. Peat is the next most important archive, followed by alluvial and colluvial sediments, as  
147 well as cave sites, the later also often being known for their archaeological significance. Sites located at the ice  
148 margins that appear to be under the ice reflect uncertainties in the location of the ice margin both in time and  
149 space during the LGM, as well as the fact that the selected time window for this study ( $21 \pm 2$  ka) is later than the  
150 maximum ice advance in some regions (Hughes and Gibbard, 2015). For completeness, we also include marine  
151 records which have the advantage of more continuous deposition and often better dating over the LGM period,  
152 but which are prone to taphonomic biases compared to terrestrial records. These biases are discussed later in this  
153 section.

154

155 LGM pollen records were selected according to a number of quality control criteria, but primary amongst these  
156 was the existence of sufficient independent chronological control points to accurately identify samples that would  
157 fall within the  $21 \pm 2$  ka BP time-slice of interest. We have used all of the samples within this time frame where



158 the samples have been available in electronic form, else we have used the sample closest to the target time (21 ka  
159 BP). For records taken from the EPD we have used the latest Bayesian age-depth models where these were  
160 available (Giesecke et al., 2014), otherwise we have used the dates and chronology proposed by the original  
161 authors. We classified chronologies according to the COHMAP chronological quality scheme for the LGM period  
162 (Anderson et al., 1988; Yu and Harrison, 1995), which classifies record quality from 1-6 depending on whether a  
163 date falls within 2000 14C years (or less) of the time being assessed, or whether bracketing dates fall within 6000  
164 and 8000 14C years (or less) about the time being assessed (Table. 2). Chronologies based on dates that fall outside  
165 of these limits fall into COHMAP class 7, and are regarded as ‘poorly dated’ with respect to the LGM. Importantly,  
166 we have only included radiometric and other absolute dates (such as varves) in this assessment, and have excluded  
167 dates based on correlation with regional pollen records. These pollen-based stratigraphic dates have been widely  
168 used in previous LGM studies, but do not include estimates of uncertainty and are generally regarded as unreliable  
169 at this time given the sparsity of well dated pollen sites and samples on which to base any correlation (Giesecke  
170 et al., 2014).

171

172 All records that were classified as poorly dated (COHMAP class 7) were subsequently excluded from our analysis.  
173 This has meant that many of the pollen records used in previous studies were excluded, including 10 of the 18  
174 LGM records used by PMIP and associated studies in Europe (Bartlein et al., 2011; Elenga et al., 2000; Jost et al.,  
175 2005; Peyron et al., 1998; Wu et al., 2007; Cleator et al., 2020). We also excluded 104 of the 116 records in the  
176 EPD with samples that fall within our LGM time window. Many of these EPD pollen records have been used in  
177 more recent studies, although the exact record (EPD site #Entity) is often not stated. We estimate that we have  
178 excluded 16 of the 17 European sites used by Binney et al. (2017) (this study only included sites above latitude  
179 40N), 5 of the 6 European sites used by Allen et al. (2010), 28 of the 33 sites used by Cao et al. (2019) and 27 of  
180 the 71 sites used by Kaplan et al. (2016).

181

182 Other quality control criteria were also used in the selection of LGM pollen records. Published pollen diagrams  
183 that only included a small part of the terrestrial pollen assemblage, or only presented summary taxa, were  
184 excluded. Records were also excluded where the dating information was incomplete, for instance where  
185 radiocarbon dating uncertainties were not published or where it was not possible to determine if the date shown  
186 was in calibrated or uncalibrated radiocarbon years.

187

188 The modern pollen data for the climate and tree cover reconstructions were sourced from the latest version 2 of  
189 the Eurasian Modern Pollen Database (Davis et al., 2020), which is managed as part of the EPD. The EMPD2  
190 includes 8133 modern pollen samples from across the Palearctic biogeographic region from Europe to the far East  
191 of Asia. The taxa from both the fossil and modern pollen data were consolidated into 120 of the most commonly-  
192 occurring terrestrial taxa types. This taxa list was designed to be compatible with the biomisation scheme used in  
193 our study (Peyron et al., 1998; Tarasov et al., 2000) and that used in the Holocene mapping study of Brewer et al.  
194 (2017). The count of *Larix* was amplified by a factor of 10 due to its low pollen representation (Binney et al.,  
195 2017).

196

197 **2.2 Biomisation**



198

199 We converted pollen assemblages to biomes based on the European biomisation scheme of Peyron et al (1998),  
200 which in turn is based on Prentice et al. (1996). The method is described in detail in Collins et al. (2012). We  
201 expanded the number of taxa included in the biomisation procedure proposed by Peyron et al (1998) to include  
202 taxa from the Northern Eurasian biomisation procedure of Tarasov et al. (1998). The inclusion of additional  
203 Northern Eurasian taxa reflects recent evidence that modern analogues of LGM vegetation occur in parts of Siberia  
204 (Magyari et al., 2014a). The biomisation procedure (Prentice et al. 1996) assigns each taxa to a plant functional  
205 type (PFT) and calculates a score for each of these PFT's based on the sum of the square root of the percentage  
206 of each of the taxa included in that PFT. To reduce the influence of long-distance transport, taxa below 0.5% are  
207 removed at the start of the procedure. Each biome is then assigned one or more PFT's and a score for each biome  
208 is calculated as the sum of the associated PFT scores. The biome with the highest score is then viewed as the  
209 dominant biome. Where the highest score is the same for more than one biome, the dominant biome is decided  
210 based on a hierarchy of unique PFT's. Peyron et al. (1998) also included a procedure for distinguishing warm and  
211 cold steppe biomes based on re-assigning certain steppe PFT's according to the presence or otherwise of PFT's  
212 indicative of cold or warm conditions. Following the Biome6000 project (Elenga et al., 2000) and Allen et al.  
213 (2010), we did not apply this additional procedure and present only the merged steppe biome. In summary, the  
214 biomisation procedure categorised 39 arboreal pollen taxa and 39 non-arboreal taxa into 22 plant functional types  
215 (PFT's), which were then combined into 12 biomes.

216

### 217 **2.3 Quantitative climate reconstruction**

218

219 We reconstructed climate from pollen data based on a standard Modern Analogue Technique (MAT) that used  
220 PFT scores to match fossil samples with modern calibration pollen samples (as used by Davis et al., 2003). This  
221 is a similar approach to that used by Peyron et al. (1998) and Jost et al. (2005) who also applied pollen PFT scores  
222 to reconstruct LGM climate from pollen data, but who used a neural network technique which is a variant of the  
223 standard MAT (Chevalier et al., 2020). PFT scores have been used in previous large-scale European pollen-based  
224 climate reconstructions for the Holocene (Davis et al., 2003; Mauri et al., 2014, 2015), where performance was  
225 found to be better than the conventional approach based on individual taxa (eg Marsicek et al., 2018). A particular  
226 advantage of the PFT approach for the LGM is that it can help overcome problems associated with vegetation  
227 (pollen) assemblages that may have no modern analogue (Davis et al. 2003). This can be a problem during the  
228 LGM when the climate and environment could be expected to be very different from today, and when many taxa  
229 formed unusual vegetation assemblages as a result of their forced retreat to sheltered refugia locations. The  
230 problem of modern analogues is also addressed in our reconstruction by using the latest EMPD2 modern pollen  
231 dataset for calibration purposes. The EMPD2 provides a large number of potential modern analogues for many  
232 different LGM vegetation types and climates found today across the Palearctic region. PFT scores were calculated  
233 according to the methods outlined already in the Biomisation section, then normalized so that each sample was  
234 proportional to every other sample (Juggins and Birks, 2012).

235

236 The MAT method was applied using the Rioja program for R (Juggins, 2020). The modern calibration data was  
237 taken from the latest version 2 of the EMPD (as detailed earlier). The EMPD2 includes 8133 samples, which is



238 considerably larger than the modern datasets used in previous LGM pollen-based reconstructions. For instance,  
239 Peyron et al. (1998) used a calibration dataset of 683 samples, which was updated by Jost et al (2005) to include  
240 an additional 185 samples. These datasets were also mainly taken from the steppes of Kazakstan and Mongolia,  
241 while the EMPD2 covers a much wider area, spanning most of the Eurasian Palearctic region (Davis et al., 2020).  
242 The size and distribution of the modern training set in climate and vegetation space is important because in order  
243 for the method to work effectively, it is necessary to have samples representative of the likely vegetation and  
244 climate space that could be occupied by the fossil assemblage (Chevalier et al., 2020; Juggins, 2013).

245

246 A known problem with MAT is the role of spatial auto-correlation in providing unrealistically low estimates of  
247 uncertainty (Chevalier et al., 2020; Telford and Birks, 2009). This results from the fact that closely analogous  
248 modern pollen samples can also be located closely in physical space, and therefore in climate space. To reduce  
249 this problem it is possible to systematically exclude closely located modern samples from the analogue matching  
250 process, for instance, by excluding samples that fall within a certain spatial range (h-block filter) (Telford and  
251 Birks, 2009). While this approach can help, there are also three main problems associated with it. The first is error  
252 substitution, since removing samples also reduces the number of potential analogues, creating a different source  
253 of error that is not easy to categorise. Secondly, multiple samples taken from the same location are actually a  
254 strength of pollen training sets, since they are more likely to capture the full range of the assemblage diversity  
255 associated with a given climate. Thirdly, current methods that limit spatial range such as the h-block filter only  
256 do so on the horizontal axis, and do not consider the fact that samples can also be found at different elevations. In  
257 hilly or mountainous regions samples can therefore be excluded because they are closely located in horizontal  
258 space, but in fact they actually occupy very different climates and vegetation associations, contradicting the logical  
259 premise of the h-block filter. It was therefore decided not to apply this filter.

260

261 Climate reconstructions are presented as anomalies. These have been calculated with respect to modern climate  
262 at each core site location using WorldClim 2 (Fick and Hijmans, 2017), which was also used to assign the modern  
263 climate for the modern pollen samples in the transfer function (Davis et al., 2020).

264

#### 265 **2.4 Quantitative tree cover reconstruction**

266

267 It has long been recognized that the proportional representation of individual pollen taxa in a pollen assemblage  
268 does not necessarily reflect the proportion of land area covered by that taxa in the pollen source area surrounding  
269 the sample site. These differences can be caused by variations in pollen productivity, differential transport,  
270 deposition and preservation of pollen grains, and even the ease or otherwise of the identification of pollen grains  
271 themselves. This can make the interpretation of pollen taxa percentages difficult, even for relatively simple  
272 questions such as the proportion of forest to non-forest in the landscape.

273

274 There have been two main methods developed to account for this quantification problem, one using a physical  
275 modelling technique (PMT) based on estimates of pollen production for individual taxa (Gaillard et al., 2010),  
276 and the other using a MAT very similar to that used in pollen-climate reconstructions (Williams and Jackson,  
277 2003). Both approaches have been widely applied during the Holocene in Europe (Zanon et al., 2018), but we



278 know of no previous study that has applied either of these approaches to the LGM. The LGM presents a number  
279 of challenges, not least the problem of potential missing vegetation analogues, as well as low atmospheric CO<sub>2</sub>,  
280 which has been shown to influence pollen productivity (Leroy and Arpe, 2007).

281

282 Here we use the MAT to provide quantitative estimates of forest cover, following the approach of Zanon et al.  
283 (2018) who applied this method to the Holocene pollen record of Europe. We apply MAT in exactly the same  
284 way as for the climate reconstructions described earlier, including the use of PFT scores to match fossil and  
285 modern pollen samples. Instead of modern climate values, we assigned an estimate of modern forest cover to each  
286 of our modern pollen sites. To do this we use a high resolution (~100m) remote sensing dataset derived from  
287 satellite observations (Hansen et al., 2013). Zanon et al. (2018) have shown that the MAT calibrated in this way  
288 gives comparable results to the PMT approach in Europe, at least for the Holocene. One of the main differences  
289 however is that the PMT is designed to provide estimates of the proportions of different taxa, whereas the MAT  
290 (as applied here) is designed to provide estimates of the proportion of forest cover. Where the PMT can only  
291 reconstruct the proportion of forest forming trees, irrespective of their size, the MAT (following Zanon et al. 2018)  
292 is calibrated specifically to reconstruct forest composed of trees over 5m tall. This follows the FAO definition of  
293 forest as “land spanning more than 0.5 hectares with trees higher than 5 meters and a canopy cover of more than  
294 10 percent, or trees able to reach these thresholds in situ” (FAO Terms and definitions 2020  
295 <http://www.fao.org/3/I8661EN/i8661en.pdf>).

296

## 297 **2.5 Maps**

298

299 We present our results in the form of maps that include the main physiographic features of the LGM in the study  
300 area. The maps are based on the WGS84 projection. Coastlines reflect LGM sea level at 120m below present,  
301 while ice sheets are based on Ehlers et al. (2011). Modern national country boundaries are also included for  
302 reference.

303

## 304 **2.6 Marine pollen records**

305

306 We have included marine pollen records in our analysis for reasons explained below, but it is important that these  
307 records should be viewed with caution, particularly when used for biome and quantitative MAT reconstructions,  
308 and when compared with terrestrial records from different archives. Biomisation methods have been applied to  
309 individual marine pollen records (Combourieu Nebout et al., 2009), as well as multi-site synthesis studies such as  
310 the ACER project (ACER project members et al., 2017). However, marine records were specifically excluded  
311 from the Biome6000 project (Elenga et al., 2000). Similarly, quantitative climate methods have been applied to  
312 individual marine pollen records (Combourieu Nebout et al., 2009; Fletcher et al., 2010), as well as multi-site  
313 synthesis studies (Brewer et al., 2008). However, marine records have also been specifically excluded from other  
314 major pollen-climate studies (Cheddadi et al., 1996; Davis et al., 2003; Marsicek et al., 2018), as well as  
315 quantitative forest cover reconstructions (Zanon et al. 2018).

316





317 Discussion on the advantages and problems associated with marine records can be found elsewhere (Chevalier et  
318 al., 2020; Daniaux et al., 2019), but are reviewed briefly here where relevant to the methodologies applied in this  
319 study. Marine sedimentary records provide continuous and well dated pollen records for the LGM that are often  
320 lacking from many terrestrial regions, especially in arid areas with few alternative anaerobic sediment sinks.  
321 Conversely however, pollen source areas for marine sites may be many hundreds of kilometers from the coring  
322 site and may be liable to change through time in response to changes in distance to the coastline, rates of river  
323 discharge and ocean and atmospheric dynamics. This can theoretically give rise to changes in the vegetation  
324 shown in the pollen assemblage recorded at the marine site without any actual change in climate or other  
325 environmental pressure. The large and indeterminable source area of marine records also mean that it is difficult  
326 to apply quantitative MAT reconstruction methods, not least because the mean climate or forest cover of the  
327 source area is almost impossible to determine. In addition, the pollen record and the calibration dataset to which  
328 it is being compared are composed largely of terrestrial lakes and bog sites with much smaller and more  
329 homogeneous source areas. This creates a series of problems, the more obvious of which is the calculation of  
330 anomalies, since we cannot assume that the modern climate at the (marine) coring site location is representative  
331 of the (terrestrial) source area. In this study we have taken the closest point on land as the modern climate for the  
332 calculation of anomalies. The next problem is that the large source area may capture a combination of different  
333 vegetation types that is not going to be represented in a calibration dataset based on samples from terrestrial sites  
334 with much smaller source areas, for instance a mixture of coastal and mountain vegetation, or even vegetation  
335 from different continents (Magri and Parra, 2002). However, in our analysis we did not find any sample from a  
336 marine record that did not have a reasonable modern analogue in our training set (chord distance <0.3)(Huntley,  
337 1990), even though we did not adjust the pollen assemblage for the over-representation of Pinus in the marine  
338 pollen samples.

339  
340 Typically, the Pine component is excluded from the terrestrial pollen sum when calculating percentages for marine  
341 pollen samples, and in some cases Pine has been excluded entirely from the samples used in marine pollen-climate  
342 reconstructions (Combourieu Nebout et al., 2009). The problem with excluding Pinus is two-fold, the first is that  
343 Pinus often represents the main forest forming tree in the Koeppen Csb climate zone on the Atlantic coast where  
344 many marine sites are located (García-Amorena et al., 2007), as well as representing the most abundant tree taxa  
345 in Europe during the LGM (Figure A1c). The second problem is that the remaining terrestrial taxa often constitute  
346 a very small number of pollen grains in a typical marine pollen sample (<100 grains), which can result in pollen  
347 assemblages that are not based on a statistically stable count of the pollen sample (Chevalier et al., 2020).

348

### 349 **3. Results**

350

#### 351 **3.1 Vegetation & Biomes**

352

353 Results of the biomisation analysis shows that steppe (STEP) was the most common biome at the LGM across the  
354 study area, occurring at 36 out of 63 sites, indicating that the landscape was largely dominated by cool temperate  
355 grasslands across much of western Central Europe, central and eastern Mediterranean, as well as North Africa  
356 and the Middle East (Fig. 2). However, at the same time we also find that there were a significant number of sites



357 where we find woody and forest biomes occur, more particularly in southern and eastern Iberia, northern Italy and  
358 central eastern Europe. The most dominant of these forest and woody biomes are taiga (TAIG) in the north, and  
359 cool-mixed forest (COMX) and xerophytic woodlands (XERO) in the south.

360

361 As would be expected, the dominance of STEP biomes is generally reflected in low arboreal pollen percentages  
362 across the same areas/sites (Fig. 3 & 4). Exceptions to this rule can be found at marine sites such as [site #3] and  
363 [site #58] where STEP is reconstructed despite arboreal pollen percentages of 71 and 80 percent respectively. This  
364 apparent contradiction illustrates some of the idiosyncrasies of the biomisation method, especially when applying  
365 the method to marine pollen samples. In this case it is important to remember that the AP% is calculated from the  
366 sum of the percentages of each relevant taxa, but the score for each biome is the sum of the square root of the  
367 percentages of each of its constituent taxa. This results in biomes with taxa with large percentage values scoring  
368 proportionally smaller, and biomes with taxa with small percentage values scoring proportionally larger. For  
369 example, a single taxa at 50% has a square root of 7.07, but the sum of the square roots of 10 taxa each at 5% is  
370 22.36 even though the sum of the percentages is the same 50%. This effect can be particularly pronounced in  
371 marine pollen samples because they are usually dominated by a single taxa (Pinus) that forms a high percentage  
372 of the total assemblage. Since there are often more non-arboreal taxa than arboreal taxa in a pollen assemblage,  
373 the non-arboreal taxa can dominate in the biomisation process even if collectively their percentage of the  
374 assemblage is a lot less than the arboreal taxa, resulting in a non-arboreal biome such as STEP having the highest  
375 biome score.

376

377 The main arboreal biomes found at the LGM include Taiga (TAIG), Cool Mixed Forest (COMX), Cool Conifer  
378 Forest (COCO) and Xerophytic Scrub (XERO), with just a single occurrence of Cold Mixed Forest (COMX) and  
379 Warm Mixed Forest (WAMX). We do not record any Temperate Deciduous Forest (TEDE), Tundra (TUND) or  
380 Desert (DESE) biomes at any site at the LGM.

381

382 An alternative picture of LGM tree-cover is provided by the MAT reconstructions (Fig. 5). MAT performance  
383 statistics for tree cover are shown in table 3, based on an evaluation using the modern training set. This shows a  
384 relatively large root mean square error (RMSE) of 21.03. and an R2 of 0.52 that is not as good as for the MAT  
385 climate analysis, but overall the results are comparable with previous MAT tree cover studies (Zanon et al., 2018).  
386 In general, the MAT values (site average 34%) show forest-cover around 16% less than that suggested from AP%  
387 (site average 50%) (Fig. 3), although sites with very low AP% also show higher values based on MAT. These  
388 differences are consistent with comparisons between MAT and AP% in Zanon et al (2018), although it should be  
389 noted that uncertainties related to the MAT reconstructions are large ( $\pm 23\%$ ). Zanon et al (2018) found that the  
390 differences between MAT and AP% were greatest over Northern Europe and in Arctic and sub-Arctic climate  
391 regions that are likely to be comparable to many areas of Europe during the LGM. These regions today are  
392 associated with tree-forming taxa such as Birch that fail to grow to a height of 5m or more, developing only as  
393 shrubs or krummholz forms.

394

395 Pollen taxa percentages are shown in figure 6, and distribution maps of the 33 most common taxa are shown in  
396 the supplementary figures A1a-f. Of the 21 arboreal taxa, Pinus generally has the highest values and is the most



397 widespread, being present at all 63 sites. Other acicular arboreal taxa include *Juniperus*, which also has a wide  
398 distribution across EurMedMidEst although at lower values. The rest of the acicular arboreal taxa have more  
399 regional distributions. *Picea* is found mainly to the north of the study region, away from the Mediterranean, whilst  
400 *Abies* is generally found more to the south. *Larix* occurs only in the central European area including the northern  
401 edge of the Po plain just south of the Alps, whilst *Cedrus* is found mainly across south and west Europe in locations  
402 much further north than its Holocene and modern distribution which is confined mainly to Morocco and Lebanon  
403 (Collins et al., 2012). Temperate broadleaf arboreal taxa which also include cold-tolerant species such as *Betula*  
404 and *Salix* are relatively widely spread across the EurMedMidEst during the LGM, while less drought tolerant taxa  
405 such as *Alnus*, *Carpinus* and *Corylus* are found more to the south-west through to the north-east. Other temperate  
406 broadleaf arboreal taxa such as *Quercus* (deciduous) and *Ulmus* have a much more southern distribution, with  
407 *Fraxinus*, *Olea*, and *Quercus* (evergreen) being more prevalent in the south-west. In contrast, *Fagus* occurs more  
408 to centre and the east, while *Tilia* is found even in more northern locations of central Europe. The remaining  
409 arboreal taxa are more shrubby and drought adapted, with *Ephedra* and particularly *Ephedra fragilis* having a  
410 southern distribution, whilst the more cold adapted *Hippophae* being found even in the north of central Europe  
411 (similar to *Tilia*).

412

413 The main non-arboreal taxa generally indicate cool, dry and environmentally disturbed conditions across much of  
414 the EurMedMidEst. The most widely distributed taxon is *Poaceae*, which like *Pinus*, is found in all records. Other  
415 non-arboreal taxa with a widespread distribution include *Rubiaceae*, *Apiaceae* and *Asteraceae* (*Asteroideae*),  
416 while *Plantago*, *Cayophyllaceae*, *Brassicaceae* and *Asteraceae* (*Cichorioideae*) have a more southern and western  
417 distribution. *Thalictrum* can be found mostly at sites in the centre of the EurMedMidEst, along with  
418 *Helianthemum* which also extends to sites in the south-west. Other taxa such as *Chenopodiaceae* and *Artemisia*  
419 have a more southern distribution, reflecting their preference for drier and less cold climates.

420

### 421 3.2 Climate reconstruction evaluation

422

423 Evaluation of transfer function performance based on the modern training set is presented in table 3. This shows  
424 that root mean square error predicted (RMSEP) values were smallest for summer temperatures (2.21C), and largest  
425 for winter temperature (3.35C), with mean annual temperatures in between (2.28C). The weaker performance for  
426 winter temperatures largely reflects the much greater range of winter temperatures in the training set. In turn, this  
427 contributes to a better R2 performance for winter temperatures (0.91) than annual temperatures (0.9) and summer  
428 temperatures (0.81). Overall R2 performance for precipitation is weaker than for temperature, which is typical  
429 because of the higher spatial variability of precipitation compared to temperature. Summer precipitation has the  
430 strongest R2 performance (0.75) compared to winter and annual precipitation (both 0.69), as well as smaller  
431 RMSE values (52mm) than winter (78mm).

432

433 These results indicate good transfer function performance especially for temperature, and are comparable with  
434 those found in other continental scale pollen-climate studies (Bartlein et al., 2011). It is important to remember  
435 though that comparisons between studies can only be made with caution because results are often heavily



436 dependent on the nature of the modern pollen dataset used as the training set, which is not the same in all studies  
437 (Juggins, 2013).

438

439 The evaluation of pollen-climate transfer function performance using the modern training set necessarily only  
440 indicates performance for the present-day climate and vegetation. We therefore undertook two additional tests of  
441 our MAT methodology to assess performance during the LGM. The first test was to compare our MAT results  
442 with previous pollen-climate reconstructions based on the same LGM sites but using different methods. These  
443 previous reconstructions include the neural-network methodology of Peyron et al. (1998) and Jost et al. (2005)  
444 which we call MAT-NN, as well as the Inverse Modelling approach by Wu et al. (2007) which we call INV.

445

446 We found 10 sites/records in our dataset which were also included in these earlier analysis (Fig. 7). All of the  
447 other sites used in these earlier studies were excluded from our study because of poor dating control. While these  
448 previous studies almost certainly shared the same pollen dataset, this dataset has never been placed in the public  
449 domain and no metadata provided other than the name of the site and the publication. In other words, it is not  
450 known if 1) the data represented a single sample or the mean of multiple samples within a time-window, 2) what  
451 the actual depth/age of those samples were or the actual sediment core in the case of multiple cores, 3) if the data  
452 was digitized or had a restricted taxa list (as was the case with the data from Huntley & Birks (1983), which is a  
453 likely source). While these aspects are unknown, it seems likely that the pollen data we used in our analysis was  
454 very similar if not identical in most cases, and in fact we found that the biomes reconstructed from our pollen  
455 dataset for these 10 sites are identical to the biomes reconstructed using the earlier pollen dataset (Elenga et al.,  
456 2000).

457

458 We compare our MAT with the MAT-NN and INV reconstructions in figure 7. On average across all 10 records,  
459 the MAT and INV methods give almost identical results for both anomalies of mean annual temperature (MAT -  
460 6.6C, INV -7.2C) and precipitation (MAT 158mm, INV 165mm). Uncertainties are also similar for both methods.  
461 In contrast, the MAT-NN method gives much cooler mean annual temperature anomalies (MAT-NN -13.9C) and  
462 drier precipitation anomalies (MAT-NN -474mm). On a site by site basis the MAT and INV methods show closer  
463 agreement for temperatures than precipitation, although precipitation has proportionally larger uncertainties. The  
464 reconstructions based on these two methods are close enough that the uncertainties overlap at all sites for both  
465 temperature and precipitation, except the precipitation reconstruction at Lac de Bouchet (site 25). The reason for  
466 this is not clear, but there could easily be differences with the pollen data analysed by Wu et al. (2007) in their  
467 INV reconstruction since the pollen record at Lac de Bouchet (Reille and de Beaulieu, 1988) includes multiple  
468 cores each with many different samples covering the LGM period.

469

470 The second evaluation of the MAT reconstruction method is based on comparison with a chironomid summer  
471 temperature record from Lago della Costa (site #34) in Northern Italy, analyzed by Samartin et al. (2016). We  
472 compare the chironomid record with our MAT reconstruction using pollen data that Samaratin et al (2016) also  
473 analysed from the same core. The results are presented in Fig. 8 and are shown as anomalies compared to the  
474 present day over our LGM time-window (19-23k BP). Our pollen-climate reconstruction is for JJA mean  
475 temperate, while the chironomid reconstruction is for July mean temperature, with the anomalies based on the



476 modern equivalent JJA and July mean temperatures respectively. The average anomaly values for all 8 samples  
477 reconstructed by the pollen-climate MAT are  $-10.2 \pm 3.5\text{C}$ , and for the chironomids  $-9.5 \pm 3.0\text{C}$ . This indicates  
478 that pollen and chironomid average summer temperature reconstructions are almost identical taking into account  
479 the overlapping uncertainties, while also showing a strong similarity on a sample-by-sample basis throughout the  
480 time-series.

481

### 482 **3.3 Climate reconstruction**

483

484 Reconstructed LGM temperatures indicate an overall mean annual cooling of  $-7.2 \pm 3.3\text{C}$ , with a greater cooling  
485 of around  $-9.3 \pm 4.5\text{C}$  in winter and  $-5.0 \pm 3.2\text{C}$  in summer (Fig. 9). All sites apart from Lake Van [62] in eastern  
486 Turkey show cooler temperatures at the LGM compared to modern (Fig. 10), and even at this site cooler conditions  
487 fall within the uncertainties. With greater cooling in winter compared to summer, the difference in temperature  
488 between winter and summer also increased (shown by positive anomalies) at most (but not all) sites (Fig. 10).  
489 This increase in continentality was around  $+4.2\text{C}$  on average across all sites (Fig. 9).

490

491 We reconstruct an overall decline in mean annual precipitation of around  $-91 \pm 270\text{mm}$  (-13%) at the LGM. Most  
492 of this decline is in winter ( $-38 \pm 90\text{mm}$ ) (-21%), while in summer a small increase is shown ( $10 \pm 57\text{mm}$ ) (6%),  
493 although uncertainties are large (Fig. 11). Compared to temperature there is significant seasonal and spatial  
494 variability in positive and negative precipitation anomalies (Fig. 12). Positive anomalies appear more predominant  
495 in eastern and southern Spain and in central eastern Europe in both summer and winter, while positive anomalies  
496 are found more generally in summer across sites in Southern Europe and the Mediterranean. These more positive  
497 summer anomalies also reflect a relative shift from winter to summer in the seasonality of precipitation in this  
498 region.

499

### 500 **4.0 Discussion**

501

502 Before we consider the results of our analysis it is important to provide some context in terms of European LGM  
503 geography and environment, which was very different from today (Fig. 1). Major ice sheets covered Scandinavia  
504 and much of the UK, the Alps, and the Pyrenees. Sea level was 120m lower, resulting in much of the North Sea  
505 and English Channel becoming dry land, and the European coastline extending over 100 km out into the Atlantic  
506 and Mediterranean, especially around the Bay of Biscay and Adriatic. The Black Sea was no longer connected to  
507 the Mediterranean, and was smaller with a water level around 100m lower than today (Genov, 2016). These  
508 changes in sea or water level had two main consequences, the first being that the marine sites were closer to land,  
509 and therefore closer to (low lying) terrestrial vegetation and (pollen carrying) river discharge points than they are  
510 today. The second consequence of lower seas levels is that terrestrial pollen sites were located further from the  
511 moderating effect of the ocean than they are today, resulting in a localised modification of the climate experienced  
512 by the site irrespective of regional or global changes.

513

514 The maps used in our analysis shows the maximum ice sheet at  $21\text{k} \pm 2\text{k}$  (Ehlers et al., 2011). The precise  
515 geographical location of the ice sheet is difficult to resolve at a fine spatial scale, however, which explains why



516 some sites close to the ice margin appear to be actually located under the ice (for example sites site #46 & site  
517 #54). The resolution of the map also shows the occurrence of permanent ice not only to the north and over the  
518 Alps, but also on many subsidiary areas of high ground across central and southern Europe, including areas such  
519 as the Pyrenees, Massif Central, Vosges and Carpathian Mountains. While global ice volume may have peaked  
520 ~21 ka individual ice sheets in Europe and other areas are known to have reached their maximum extent at  
521 different times (Hughes et al., 2016). The larger ice sheets are likely to have had a significant influence on regional  
522 climate and environmental conditions across Europe, but the smaller ice sheets had similar if more localized  
523 impacts as well. Surrounding each ice sheet would have been an unglaciated area of active peri-glacial processes  
524 and newly created and unstable ground. This would include outwash plains, impounded lakes and recently drained  
525 lake beds, seasonally and sporadically flooded areas, moraines, kettle holes and other glaciological and peri-  
526 glacial features. Soils in these areas would be non-existent or skeletal, and vegetation would find it difficult to  
527 obtain nutrients and water for survival, irrespective of the prevailing climatic conditions. Outside of these areas,  
528 permafrost is also likely to have been present, particularly north of the Alps (Vandenberghe et al., 2014), which  
529 would also act as an impediment to vegetation growth.

530

531 In terms of regional climate, the major ice sheets would have provided significant barriers to westerly atmospheric  
532 circulation, or even north-south circulation in the case of the Alps and Pyrenees. As well as representing a physical  
533 obstruction, the thermodynamic response of the atmosphere to these high, cold obstructions would have been to  
534 encourage the formation of areas of semi-permanent high pressure, similar to those found today for instance over  
535 the Greenland ice sheet. In addition, the Laurentide ice sheet located over North America would have generated  
536 downstream effects over Europe. These physical and thermodynamic effects would have affected the direction of  
537 storm tracks, as well as more local climatic effects commonly associated with ice sheets such as strong katabatic  
538 winds.

539

#### 540 **4.1 Vegetation Cover**

541

542 The nature and extent of forest cover during the LGM remains a matter of considerable debate. Vegetation models  
543 driven by LGM climate model simulations generally indicate extensive areas of boreal forest north of the Alps,  
544 and a mix of temperate and warm-temperate woodland to the south across southern Europe and much of the  
545 Mediterranean. Treeless areas such as steppe are mainly confined to those areas where it is also found today,  
546 namely inland Iberia, Ukraine, southern Russia and Turkey, while Tundra is found to the north close to the  
547 Scandinavian Ice Sheet (Allen et al., 2010; Cao et al., 2019; Prentice et al., 2011; Velasquez et al., 2021).

548

549 Evaluation of these vegetation-model simulations against data has been largely based on comparison with  
550 compilations of pollen-biome reconstructions (Prentice et al., 2011; Allen et al., 2010; Cao et al., 2019; Velasquez  
551 et al., 2021). Early studies were based on only a limited number of sites from southern Europe, and showed steppe  
552 at all sites in contradiction with model simulations (Elenga et al. 2000). More recent pollen compilations have  
553 included more sites especially to the north that have revealed a more mixed picture of vegetation cover, with forest  
554 biomes at some sites both south and north of the Alps that appear more consistent with model simulations (Binney  
555 et al., 2017; Cao et al., 2019). However, many of these pollen sites used in these studies were assigned an LGM



556 age based on poor or incorrect dating control, and likely date to MIS3, the Late-Glacial or even the Holocene.  
557 Nevertheless, based on our compilation of more securely dated LGM pollen sites, we also show a wider  
558 distribution of forest biomes particularly in Iberia, northern Italy and Central Europe, although with greater areas  
559 of steppe than suggested by the models over the remaining regions.

560

561 However, the interpretation of biome reconstructions requires care since the forest cover and vegetation  
562 composition may not be as clear as the dominant biome suggests. For instance, we find that steppe is still  
563 reconstructed as the dominant biome at some sites despite arboreal pollen forming 70-80% of the pollen  
564 assemblage. In addition, it is important to remember that pollen-biomes are based only on the proportion of taxa  
565 that can form forest and woodland, while these taxa may in fact exist only as shrubs or stunted krummholz forms  
566 in the challenging climate and environment of the LGM. Alternatively, similar conditions may favour low-lying  
567 non-arboreal taxa forms with poor pollen dispersion or even insect pollinated taxa forms that may be poorly  
568 represented in the pollen assemblage, giving greater prominence to arboreal taxa whose pollen may be the result  
569 of long-distance transport.

570

571 Quantitative MAT based reconstructions of forest cover can overcome some of these problems where they can be  
572 detected based on the composition of the pollen assemblage when compared with the modern land-cover. Chord-  
573 distance measurements of the match between fossil and modern pollen assemblages indicate good LGM analogues  
574 exist in our large Eurasian modern pollen dataset. The results of the MAT forest cover reconstruction indicates  
575 that forest cover was low but not entirely devoid of woodland in most areas, similar to the modern boreal forests  
576 of Siberia and consistent with a steppe-tundra-woodland mosaic proposed by many authors (e.g. Birks and Willis,  
577 2008; Willis and Van Andel, 2004). Uncertainties are large, but for comparison the MAT site-average of 33% is  
578 slightly less than the average today over the Boreal region of Europe (43%) and slightly more than the average  
579 today over Mediterranean region (27%) (Zanon et al. 2018).

580

581 By calculating the percentage of each of the taxa in each LGM pollen sample using a standardized pollen sum,  
582 we are able to make direct comparisons between different LGM pollen records and their taxa percentages (Figure  
583 6, A1). The results show a preponderance of boreal forest taxa to the north of the Alps, consistent with biome  
584 results mentioned earlier. Pinus is the most common forest forming taxa in this boreal zone, together with Picea,  
585 and including Larix to the east and Abies to the west. The occurrence of Betula and Juniperus also suggests  
586 shrubby elements consistent with arctic shrub-tundra, although high Poaceae and other herbaceous taxa such as  
587 Artemisia and Chenopodiaceae indicate more steppe than tundra. Other deciduous taxa found north of the Alps  
588 include cold tolerant generalists such as Corylus and Alnus, as well as low percentages of relatively thermophilous  
589 taxa in the east, such as Carpinus and Tilia.

590

591 These results are consistent with charcoal (Magyari et al., 2014a; Willis and Van Andel, 2004), malacological  
592 (Juričková et al., 2014), biomarkers (Zech et al., 2010) and genetic evidence (Stivrins et al., 2016; Willis and Van  
593 Andel, 2004) that the main forest region north of the Alps was in the eastern region of Central Europe around the  
594 Carpathian basin. This was also an area where cold and moisture sensitive deciduous taxa were also able to survive  
595 (Magyari et al., 2014), although evidence of temperate taxa found in the pollen record has yet to be supported by



596 charcoal and macrofossil records (Feurdean et al., 2014). Our pollen evidence indicates an open taiga or cool  
597 mixed forest that extended in central and eastern Europe to areas close to the Scandinavian and Alpine ice caps,  
598 as proposed by Willis and Van Andel (2004) and Huntley and Allen (2003), although whether this represents  
599 isolated pockets of forest or an extended open steppe-forest is difficult to determine (Kuneš et al., 2008). Even  
600 steppe or tundra areas in western Europe show a low but significant presence of the pollen of tree taxa at sites  
601 close to the ice sheets that are unlikely to be solely the result of long distance transport or reworking (Kelly et al.,  
602 2010). The presence of woodland in these areas is also supported by mammalian remains, for instance at Kents  
603 Cavern in SW England (Stewart and Lister, 2001).

604

605 Overall however, our results clearly show a much greater predominance of thermophilous and moisture sensitive  
606 deciduous taxa south of the Alps, particularly in Iberia and Northern Italy, where temperate broadleaf forests  
607 survived in sheltered refugia (Kaltenrieder et al., 2009). Most of these appear to be in hilly areas with the ability  
608 to generate orographic rainfall (Monegato et al., 2015), on south facing slopes to make the most of the sun's  
609 radiant energy and located above the valley floor to escape frost and flooding. We might also expect these areas  
610 to be sheltered from cold northerly winds, and benefit from relatively mild and moisture laden winds coming from  
611 the Mediterranean Sea. For instance, the presence of woodland and low glacier altitudes along the southern slopes  
612 of the Alps around the Po Valley and Trentino region is consistent with strong orographic rains generated by  
613 southerly and easterly winds that today can be generated by low pressure located south of the Alps in the Gulf of  
614 Genoa, and consistent with a southerly storm track around the Alps (Kehrwald et al., 2010; Luetscher et al., 2015).  
615 Generally, as might be expected, areas of forest reconstruct similar or increased precipitation compared to today,  
616 and areas of steppe indicate decreased precipitation (see next section).

617

618 Independent evidence of LGM vegetation is provided by archaeozoological data. This data supports the  
619 palynological evidence for the existence of forest and woodland refugia across the ice-free areas of Europe at  
620 latitudes north of the Alps. For instance, large vertebrates in these areas show patterns of extirpation and extinction  
621 in response to shifts in climate and vegetation cover that is different for different species, indicating a variety of  
622 environments and niches (Lister and Stuart, 2008; Stewart and Lister, 2001). As with the pollen record, the  
623 presence of temperate adapted large vertebrate taxa within the glacial landscape of Western Europe also suggests  
624 the existence of temperate "micro-refugia" (Stewart and Lister, 2001), consistent with suggestions that temperate  
625 arboreal taxa were not entirely extirpated from the region during the LGM (Magri, 2010). Further east, mammal  
626 assemblages indicate generalized loss of forest components in the East European Plain (Puzachenko et al., 2021)  
627 which is consistent with our data indicating low forest cover in this region. In other areas, evidence of the  
628 prevailing land cover at the LGM comes from studies of small vertebrate communities, which have a closer  
629 affinity to the prevailing environment than large vertebrates (López-García and Blain, 2020) that have the  
630 propensity to migrate large distances, often on a seasonal basis. These studies of small vertebrate assemblages  
631 also support the existence of temperate "micro-refugia" in France (Royer et al., 2016) and the existence of  
632 woodland components in many regions across Southern Europe including parts of Iberia (Bañuls-Cardona et al.,  
633 2014) Italy (Berto et al., 2019) and the Balkan Peninsula (Mauch Lenardić et al., 2018).

634





635 Other paleobotanical evidence also supports our land cover reconstruction. Schafer et al. (2016) suggest leaf wax  
636 patterns from palaeosols in Spain may indicate the presence of drought intolerant deciduous trees and more humid  
637 conditions during the LGM. Significantly, none of the pollen sites indicate that temperate broadleaf forests were  
638 dominant, and broadleaf temperate taxa always appear part of a mixed woodland together with cold or aridity  
639 adapted evergreen and needleleaf taxa, including typical Mediterranean taxa. This type of mixed vegetation  
640 probably extended to the Balkans where the hilly terrain and proximity to the Mediterranean would appear to have  
641 provided favourable climatic conditions, although we still lack LGM sites from this region. At sites in central and  
642 southern Italy and east through Greece and Turkey to the Middle East (and including North Africa), the vegetation  
643 appears drier with a greater prevalence of steppe. Only a site in Georgia in the Caucasian mountains indicates the  
644 presence of significant amounts of forest (mainly Pinus), a result that was also found by Tarasov et al. (2000), and  
645 probably linked to favourable orographic precipitation and proximity to the Black Sea.

646  
647 Comparison with LGM land cover from vegetation modelling studies driven by climate model simulations  
648 indicate a much wider presence of forest than that shown by the pollen data (Kaplan et al., 2016). Data-model  
649 agreement appears to be closest over eastern-central Europe where pollen indicates the presence of open Boreal  
650 forest, and over south-west Europe with the presence of cool mixed temperate forest, including broadleaf  
651 deciduous and thermophilous elements (Prentice et al., 2011; Allen et al., 2010; Cao et al., 2019; Velasquez et  
652 al., 2021). Nevertheless, agreement still appears to be weak over western-central Europe and Southern and Eastern  
653 Europe through to the Middle East, where pollen data continues to indicate widespread steppe. One proposed  
654 explanation for this data-model discrepancy has been the role of fire (including man-made fire) in maintaining  
655 forest openness, a factor influencing forest cover that is not included in most vegetation models (Kaplan et al.,  
656 2016). In the Carpathian basin Magyari et al. (2014a) noted that charcoal increased as forest cover declined,  
657 suggesting that wildfires played a role in decreasing forest cover during the LGM. Other studies have noted low  
658 levels of charcoal and therefore fires during the LGM, although these tend to be from steppe areas with low  
659 biomass and fuel availability (Connor et al., 2013; Kaltenrieder et al., 2009). Recent LGM vegetation simulations  
660 that include fire indicate much lower values of forest cover than those without fire over western central Europe,  
661 while forest remains in central eastern Europe (see figure 6 in Velasquez et al., 2021). This appears closer to the  
662 data, but the values are perhaps too low compared with our MAT reconstructions here (Figure 5).

663

#### 664 **4.2 Climate: Temperature**

665

666 The climate of the LGM is generally considered to have been cooler and drier than today, but data-model  
667 comparisons continue to highlight important discrepancies, not only in the degree of cooling and drying but also  
668 in their seasonal and spatial distribution. Data-model comparisons over Europe have mainly used quantitative  
669 pollen-based climate reconstructions, especially the Palaeo-model Intercomparison Project (PMIP/CMIP)  
670 (Bartlein et al., 2011; Harrison et al., 2015; Kageyama et al., 2006). These pollen reconstructions use many of the  
671 same compilations of LGM pollen data used in the biome reconstructions mentioned earlier (Elenga et al., 2000),  
672 and therefore suffer from the same problems of dating control, unclear provenance and a potentially limited taxa  
673 assemblages.

674



675 We identified 10 LGM pollen records where we could directly compare our MAT-based pollen-climate  
676 reconstruction with previous pollen-climate reconstructions based on Inverse Modelling (INV) (Wu et al., 2007),  
677 and the Neural Networks method which is a version of MAT (MAT-NN) (Peyron et al., 1998a). This comparison  
678 showed that our MAT reconstructions were very similar to the INV method, but not as cold or dry as the MAT-  
679 NN method. This has two main implications. The first is that our reconstructions indicate greater agreement with  
680 the results of climate model simulations since climate models indicate temperatures closer to the INV  
681 reconstructions (Latombe et al., 2018) than the MAT-NN reconstructions (Jost et al., 2005; Kageyama et al.,  
682 2006). The difference between our MAT and the MAT-NN reconstructions is likely the result of the modern  
683 calibration datasets used, since the MAT-NN reconstruction by Peyron et al. (1998a) was based on a considerably  
684 smaller number of samples taken mainly from the cold dry steppes of Kazakstan and Mongolia. The second  
685 implication is that the MAT method may not be significantly impacted by the effects of lower CO<sub>2</sub> (Cowling and  
686 Sykes, 1999; Prentice and Harrison, 2009; Williams et al., 2000) or indeed insolation changes during the LGM,  
687 since the MAT results are similar to those based on the INV method which specifically takes account of these  
688 non-climatic factors (Wu et al., 2007). This would also suggest that MAT could also work well for pollen-based  
689 climate reconstructions on longer glacial-interglacial timescales where insolation and CO<sub>2</sub> vary significantly from  
690 their modern values.

691

692 Comparison with LGM climate reconstructions based on other (non-pollen) proxies provides another way of  
693 evaluating our pollen-based reconstruction. We reconstruct both summer and winter temperatures and show that  
694 cooling in winter was greater than in summer at most sites, associated with an increase in continentality (increased  
695 temperature difference between summer and winter). A similar seasonal pattern of temperature change has also  
696 been shown in other studies that reconstruct both summer and winter LGM temperatures, including Prud'homme  
697 et al. (2016) using d18O analysis of earthworm calcite granules at Nussloch near the French-German border,  
698 Bañuls-Cardona et al. (2014) using faunal remains of small mammals at 4 locations in western Spain, and  
699 Ferguson et al. (2011) who examined seasonal temperature change using d18O and Mg/Ca analysis of limpet  
700 shells at Gibraltar in southern Spain. The increase in continentality at Nussloch (Prud'homme et al., 2016) was  
701 reconstructed at between 11.6 to 15.6 °C, comparable at the lower end with nearby pollen sites [site #28] 10.4 ±  
702 5.8 °C and [site #29] 7.9 ± 5.7 °C. The faunal sites in western Spain studied by Bañuls-Cardona et al. (2014) gave  
703 much reduced increases in continentality, but nevertheless similar to nearby pollen sites. For instance at Valdavara  
704 5.1 °C [site #3] 5.2 ± 3.1 °C, El Miron 1.2 °C [site #19] 5.1 ± 6.2 °C, El Portalon 0.9C [site #16] 2.8 ± 1.8 °C and  
705 Cueva de Maltrvieso 6.1C [site #2] 4.8 ± 3.4 °C. Further south at Gibraltar the limpet-based study of Ferguson et  
706 al. (2011) also shows a relatively small increase of 2 °C. The nearest pollen site [site #5] however shows a larger  
707 increase of 4.7 ± 2.3 °C, although differences could be expected given the different temporal resolution of annual  
708 laminae on mollusk shells compared to pollen assemblages that reflecting much slower changes in trees and other  
709 long-lived flora.

710

711 Summer temperatures were warm enough during the LGM over the Alpine areas that Swiss lakes were largely ice  
712 free in summer, while glacier ELA's around the time of the LGM suggest summers were -6.5 to -7.7 °C cooler  
713 compared to the LIA (Heiri et al., 2014). This cooling was similar to that found at Nussloch some 200km north  
714 of the Swiss border by Prud'homme et al. (2016), who reconstructed anomalies of -6 to -8 ± 4 °C from d18O



715 analysis of earthworm calcite granules (representing warm season May-September temperatures). Slightly less  
716 cooling was found close by at the nearby site of Achenheim where analysis of Mollusc assemblages gave summer  
717 (August) cooling estimates of  $-3.5$  to  $-6.5$  °C based on MAT (Rousseau, 1991), and  $-5.5$  to  $-9.5$  °C based on the  
718 Mutual Climatic Range method (Moine et al., 2002). These reconstructions appear somewhat cooler than nearby  
719 pollen sites [site #28]  $-1.4 \pm 3.6$  °C and [site #29]  $-2.7 \pm 5.1$  °C, although comparable with the pollen site [site  
720 #32]  $7.3 \pm 5.0$  °C 200 km further east. Similar differences also occur at the site of Les Echets on the western edge  
721 of the Alps where a Diatom based reconstruction of summer (July) temperatures (Ampel et al., 2010) indicated a  
722 greater cooling ( $-10.5$  to  $-11.5$  °C) than our pollen reconstruction [site #27] ( $-4 \pm 2.7$  °C). However, the authors  
723 caution that the results were based on poor analogues and rare taxa, as well as a small training set of only 90 lakes  
724 in Switzerland.

725

726 South of the Alps, other proxies show the opposite relationship with the pollen reconstructions. For instance, at  
727 Lago dela Costa in the Po valley, a summer (July) temperature chironomid reconstruction by Samartin et al. (2016)  
728 is around  $1-2$  °C less cool than the pollen reconstruction (JJA) for the same site [site #34]  $-11.4 \pm 2.7$  °C, although  
729 both reconstructions fall within their respective uncertainty ranges (Figure 8). In the Pindus Mountains in Greece,  
730 Hughes et al (Hughes et al., 2006) estimated LGM summer temperature anomalies of  $-7$  °C based on glacier  
731 modelling, which is comparable with that reconstructed at the nearest pollen site [site #51]  $-7.7 \pm 2.8$  °C. In Spain  
732 the analysis of small mammal remains by Bañuls-Cardona et al. (2014) shows similarly less cooling in summer  
733 or even warmer than present positive anomalies compared to the nearest pollen sites, such as Valdavara  $1.4$  °C  
734 [site #3]  $-2.3 \pm 2.8$  °C, El Miron  $-2.3$  °C [site #19]  $-5.7 \pm 5.4$  °C, El Portalon  $0.8$  °C [site #16]  $-2.6 \pm 1.1$  °C and  
735 Cueva de Maltrvieso  $-1.1$  °C [site #2]  $10.4 \pm 2.8$  °C. Further south at Gibraltar, the limpet-based study of Ferguson  
736 et al. (2012) suggests an anomaly of around  $-7$  °C, which is a greater cooling than the pollen reconstruction from  
737 this location [site #5]  $-1.3 \pm 2.2$  °C, although comparable with other pollen sites slightly further east.

738

739 Winter temperature reconstructions from non-pollen proxies show a similar pattern in relation to pollen  
740 reconstructions as for summer temperatures. North of the Alps at Achenheim, Prud'homme et al. (2016) use d18O  
741 on earthworm remains to reconstruct particularly cold winter anomalies of  $-17.6$  to  $-23.6$  °C compared to nearby  
742 pollen sites [site #28]  $-11.8 \pm 8.0$  °C and [site #29]  $-10.6 \pm 6.3$  °C. South of the Alps in Spain, the analysis by  
743 Bañuls-Cardona et al (2014) based on the remains of small mammals shows less cooling in winter compared to  
744 the nearest pollen sites, in particular Valdavara  $-3.7$  °C [site #3]  $-7.5 \pm 3.4$  °C, El Miron  $-3.5$  °C [site #19]  $-10.8$   
745  $\pm 7.0$  °C, El Portalon  $-0.1$  °C [site #16]  $-5.4 \pm 2.5$  °C and Cueva de Maltrvieso  $-7.2$  °C [site #2]  $15.2 \pm 4.0$  °C. And  
746 again, in southern Spain at Gibraltar, analysis of limpet shells by Ferguson et al (2011) suggests winter cooling  
747 of around  $-9$  °C while the pollen reconstruction suggests [site #5]  $-6.0 \pm 2.5$  °C, although sites further east indicate  
748 cooler conditions.

749

750 A number of additional proxies have also been used to reconstruct LGM mean annual temperature. Heyman et al.  
751 (2013) applied glacier mass balance modelling at sites located in the smaller mountain regions north of the Alps.  
752 These are generally slightly cooler than our pollen-based reconstructions at sites close to the Vosge Mountains -  
753  $12.7 \pm 2.0$  °C and Black Forest  $-11.4 \pm 2.3$  °C [site #29]  $-8.2 \pm 3.3$  °C, Bavarian Forest  $-10.7 \pm 2.2$  [site #32]  $-9.2 \pm$   
754  $1.2$  °C and Giant Mountains  $-8.5 \pm 1.8$  [site #46]  $-7.3 \pm 0.3$  °C. These values obtained by Heyman et al. (2013) are



755 warmer than Pud'homme et al. (2016) who estimated annual mean temperature anomalies of  $-15.1$  to  $19.1$  °C  
756 based on  $\delta 18\text{O}$  of earthworm calcite at the Nussloch site just north of the Vosge and Black Forest. The annual  
757 temperatures reconstructed by Heyman et al. (2013) are also around 2C warmer than Allen et al. (2008) who  
758 applied a similar, although simpler method to over 29 different mountainous regions across Europe that had been  
759 glaciated during the LGM. Since glacier mass balance is a function of both snowfall and temperature, these  
760 estimated temperatures vary according to estimated changes in precipitation. For instance, mean annual  
761 temperature estimates by Allen et al. (2008) are much cooler than reconstructed by pollen, with an average  
762 anomaly of  $-13.2$  °C for the 29 sites assuming a 40% reduction in precipitation, but this is reduced to  $-11.8$  °C  
763 assuming the same precipitation as modern. This compares with  $-7.2$  °C for our 63 pollen sites. The glacier mass  
764 balance modelling by Allen et al. (2008) assumes a seasonal distribution of precipitation that is similar to the  
765 present day, and does not consider increases in winter precipitation or mean annual precipitation above present  
766 day levels. Both of these are suggested by the pollen data in some regions, and both could explain glacier extent  
767 found during the LGM based on less extreme temperature anomalies more comparable with the pollen data.

768

769 To the east of the Alps in the Panonian basin, mean annual temperature anomaly estimates have been made from  
770 noble gas measurements on groundwater ranging from  $-2$  to  $-4$  °C (Stute and Deak, 1990) up to  $-9$  °C (Varsányi  
771 et al., 2011). These are similar to estimates ranging from  $-2$  to  $-9$  °C from oxygen isotope ratios from mammoth  
772 tooth enamel (Kovács et al., 2012) and are comparable with nearby pollen sites [site #50]  $-8.2 \pm 3.3$  °C and [site  
773 #52]  $-4.5 \pm 2.3$  °C. On a broader scale, Sanchi et al (2014) estimated LGM cooling in the Danube and Dneiper  
774 basins based on Lipid biomarkers in a core from the Black Sea and came up with similar mean annual temperature  
775 anomalies between  $-6$  to  $-10$  °C, which again are comparable with pollen sites from the region that range from  
776 [site #48]  $-10.5 \pm 4.1$  °C to [site #57]  $-4.3 \pm 5.8$  °C.

777

778 Further south and west, García-Amorena et al. (2007) reported mean annual temperature anomalies of  $-2.0$  to  $-$   
779  $11.3$  °C at LGM sites along the Portuguese coast, based on an indicator species method using plant macrofossils.  
780 This is similar to the closest marine pollen sites off the coast, which recorded values of [site #1]  $-10.5 \pm 4.6$  °C  
781 and [site #3]  $-5.3 \pm 2.9$  °C. Meanwhile, in the far east of the study area, Zaarur et al. (2016) estimated a mean  
782 annual temperature anomaly of around  $-3$  °C based on clumped isotope analysis of *Melanopsis* shells from LGM  
783 sediments in the Sea of Galilee. This limited cooling appears similar to the nearest pollen site [site #63] where we  
784 reconstruct a cooling of  $-2.2 \pm 4.6$  °C.

785

786 Reconstructions of LGM sea surface temperatures (SST's) provide yet another source of comparison with our  
787 terrestrial pollen-based reconstructions, although many of the physical processes controlling surface sea  
788 temperatures such as upwelling, surface mixing, surface currents, stratification and thermal inertia through the  
789 seasonal cycle, represent quite different processes to those controlling surface temperatures over land, particularly  
790 at the sub-regional scale. Nevertheless, the Atlantic coastal waters of Iberia and the waters throughout the  
791 Mediterranean Sea include many SST sites that lie in relative proximity to our terrestrial pollen-sites, allowing us  
792 to make a comparison at the largest scale. Within this area the MARGO database (MARGO Members, 2009)  
793 includes 13 Alkenone, 2 Mg/Ca and 41 Foraminifera based SST records of mean annual temperature, with the  
794 Foraminifera records also providing an additional 41 winter (JFM) and summer (JAS) SST estimates. We compare



795 the SST records with the 36 closest terrestrial pollen records which fall within a box of -11 to 35 degrees longitude  
796 and 32 to 43 degrees latitude containing all of the SST records. A simple site average indicates a mean annual  
797 SST anomaly of  $-5.5 \pm 1.0$  °C which is relatively close to the value of  $-7.2 \pm 3.4$  °C obtained from the terrestrial  
798 pollen sites [sites #1-4, 5, 7-24, 25, 26, 30, 35-38, 41, 47, 51, 53, 56-59]. Interestingly the inter-site variance  
799 (standard deviation of the reconstructed temperatures across all sites) is almost identical for the two datasets, 2.57  
800 °C for the SST sites and 2.63 °C for the pollen sites, despite representing very different environments, proxies and  
801 uncertainties. However, when we look at the seasonal temperature anomalies, we find very different results. Site  
802 averaged winter SST anomalies are  $-3.7 \pm 1.1$  °C compared to  $-9.3 \pm 4.2$  °C for winter temperatures from terrestrial  
803 pollen sites, while in summer the values are reversed,  $-7.0 \pm 0.8$  °C compared to  $-5.38 \pm 3.3$  °C respectively. This  
804 suggests that SST's experienced greater cooling in summer compared to winter, which is the opposite to that  
805 generally found in terrestrial seasonal temperature reconstructions throughout the region, although this is  
806 consistent with model simulations (Mikolajewicz, 2011).

807

### 808 **4.3 Climate: Precipitation**

809

810 Few proxies apart from pollen provide quantitative reconstructions of precipitation during the LGM. Glacier mass  
811 balance modelling includes assumptions about precipitation in order to derive temperatures (Allen et al., 2008a),  
812 but neither is independent of the other. Hughes et al (2006) estimate from glacier modelling that mean annual  
813 precipitation during the LGM at sites in the Pindus mountains in Greece was around  $2300 \pm 200$ , which they  
814 consider to be similar to the present day ( $>2000$ mm). A small change in precipitation compared to modern values  
815 is also indicated by the nearest pollen site, which is around 47km to the south [site #51], and indicates a mean  
816 annual precipitation anomaly of  $-152 \pm 294$ mm, representing just 15% of the modern value. A larger reduction in  
817 mean annual precipitation of -45% (maximum) is reconstructed by García-Amorena et al. (2007) based on plant  
818 macrofossil remains from sites on the Portuguese coast. In comparison, the closest pollen sites record values  
819 which are a little lower, ranging from [site #1] -22% to [site #3] -34%. Further north in south-west Germany,  
820 Prud'homme et al. (2018) reconstructed mean annual precipitation from the delta 13C of earthworm calcite  
821 granules at Fussloch. They estimate a field site average of 333 (159-574) mm/yr at the LGM, which represents an  
822 anomaly of -503 mm/yr (-60%) relative to the modern precipitation of 836 mm/yr. This is comparable with the  
823 closest pollen site [site #29] with an anomaly of -540 mm/yr.

824

825 As with glaciers, lake levels reflect changes in moisture balance that includes the effects of both temperature (via  
826 evapotranspiration) and precipitation, rather than just precipitation. They also represent semi-quantitative data at  
827 best, with changes often described relative to the modern or other baseline. There are few lake level records  
828 available north of the Alps, but to the south, many records indicate high lake levels in areas such as Spain (Lacey  
829 et al., 2016; Moreno et al., 2012; Vegas et al., 2010), Italy (Belis et al., 1999; Giraudi, 2017), Greece and Turkey  
830 (Harrison et al., 1996; Reimer et al., 2009) and the Middle East (Kolodny et al., 2005; Lev et al., 2019). These  
831 lake records are also supported by evidence of higher river levels in Morocco (El Amrani et al., 2008). The cause  
832 of the higher lake levels has been the subject of some debate, since many pollen records (and especially early  
833 biome reconstructions) show steppe vegetation that would suggest aridity that appears incompatible with higher  
834 lake levels. Prentice et al. (1992) proposed that the co-existence of steppe vegetation and high lake levels could



835 be possible if precipitation increased outside of the summer growing season, while summers themselves were  
836 drier and cooler with decreased evaporation. However, the results of our analysis tend to indicate the opposite in  
837 regions with higher lake levels, with increased summer rainfall and decreased winter rainfall. In addition, the  
838 increase in summer precipitation was enough to compensate for the decrease in winter rainfall, leading to an  
839 overall increase in mean annual precipitation at many pollen sites in Spain and Greece for instance. This together  
840 with depressed temperatures and consequently decreased evaporation could explain the higher lake levels, whilst  
841 also limiting the growth of trees as a result of cooler temperatures and prolonged aridity outside of the summer  
842 season. Davis & Stevenson (2007) also note a differential hydrological response between summer and winter  
843 rainfall in the Mediterranean during the Holocene that may also provide an explanation. In this case sporadic  
844 summer storms may result in high rates of runoff that may fill run-off fed lakes, but low rates of soil moisture  
845 recharge that fails to benefit vegetation in the same way winter rainfall does.

846  
847 Overall, we reconstruct only a small reduction in precipitation during the LGM of around 91mm (13%) averaged  
848 over all sites, which is less than the ~200mm reduction based on the sites in the pollen-climate compilation used  
849 by PMIP (Bartlein et al., 2011). Since our precipitation reconstruction on average matches that of the INV  
850 reconstruction by Wu et al (REF), we can attribute much of the difference to the greater aridity shown in the  
851 MAT-NN reconstruction by Peyron et al and Jost et al (2005) (see figure 7). As with temperature, this is probably  
852 a reflection of the modern training set used in the MAT-NN reconstruction which is much smaller than our training  
853 set and is largely taken from the arid steppes of Kazakhstan and Mongolia. However, it is also important to  
854 recognize the significant spatial variability in precipitation, which means that a simple average of different sets of  
855 sites from different regions may not accurately reflect the change in LGM precipitation at the European scale.  
856 Nevertheless, one of the most consistent signals in our dataset is for an increase in summer precipitation over  
857 many areas of Southern Europe and the Mediterranean. This is also found in climate models, where it has been  
858 attributed to an increase in convection-driven precipitation, although the amount of precipitation generated by this  
859 mechanism varies significantly between models (Beghin et al., 2016). A more consistent response in models is  
860 for an increase in winter precipitation across Southern Europe and the Mediterranean related to a stronger and  
861 more southerly displaced jet stream, with winter precipitation also accounting for much of the change in mean  
862 annual precipitation (Beghin et al., 2016). Our reconstruction of winter precipitation however shows less support  
863 for this scenario with a more general decrease in winter precipitation apart from southern and eastern Iberia, and  
864 with summer precipitation generally more important in those sites that show an overall increase in mean annual  
865 precipitation. This may not necessarily contradict the models in terms of the strength and position of the winter  
866 jet stream, but may instead indicate that models over-estimate the amount of moisture being carried westward  
867 from the cold North Atlantic along the storm track, especially across the far northern Mediterranean. The increase  
868 in winter precipitation across southern and eastern Iberia is however entirely consistent with a strengthened and  
869 more southerly jet stream, which also brings increased winter precipitation to the region today as a result of  
870 blocking over northern Europe/Atlantic and a negative NAO (Vicente-Serrano et al., 2011).

871  
872 Other areas that show an increase in winter precipitation include pollen sites around the eastern end of the Alps.  
873 This is consistent with a recent study by Spötl et al (2021) who argued, on the basis of cryogenic carbonates  
874 preserved in a cave in Austria, that heavy winter (and autumn) precipitation was a significant factor in driving



875 LGM glaciation in the region. The seasonally specific nature of this precipitation is also supported by the same  
876 pollen sites, which do not show any increase in summer precipitation at this time.

877

## 878 **5.0 Conclusions**

879

880 We have reconstructed the climate and vegetation cover across Europe, North Africa and the Middle East at the  
881 time of the LGM based on 63 pollen records. These records were selected using strict quality control criteria, with  
882 particular attention paid to dating control, which led to the exclusion of many records that have been used in  
883 previous studies. This fully documented dataset represents the most chronologically precise and spatially resolved  
884 view of LGM climate and vegetation during the PMIP benchmarking time window at  $21 \pm 2$  ka. Nevertheless, it  
885 is important to recognize that there are still significant spatial gaps in pollen sites especially north of the Alps, the  
886 Balkans, Turkey and the Middle East, and we continue to have only a partial understanding of the LGM over  
887 these areas.

888

889 Distribution maps were created using a standardized pollen taxonomy and sum to enable direct comparison  
890 between sites/records. Pollen biomes and quantitative MAT estimates of tree cover have also been reconstructed  
891 allowing us to determine the relative proportion of forest and woodland cover. These results show that although  
892 steppe and tundra was extensive at the time of the LGM, areas of open forest also occurred in many regions,  
893 particularly (but not exclusively) in Iberia, northern Italy and Central Europe. These forest or woodland stands  
894 are likely to have been located in environmentally favourable areas, with good soils, elevated rainfall and shelter  
895 from cold, desiccating winds. In those areas where woodland existed, Boreal taxa generally dominated north and  
896 east of the Alps, while temperate and thermophilous (mainly drought adapted) taxa were generally confined to  
897 areas south of the Alps and around the Mediterranean. The temperate deciduous forests that compose the climax  
898 community in many areas of Europe today were displaced to the south and reduced to a partnership role with  
899 Boreal elements. Overall our new reconstruction indicates greater agreement with model land cover simulations,  
900 but models still appear to over-estimate the amount of forest and woodland over areas such as France and the  
901 Benelux, Greece, Turkey and the Far East.

902

903 We reconstructed the LGM climate of Europe based on a MAT in combination with a large modern pollen  
904 calibration dataset covering the Eurasian region. In a direct comparison we found no significant difference  
905 between LGM temperatures and precipitation reconstructed from pollen using MAT, and that reconstructed by  
906 Inverse Modelling. We therefore do not find MAT performance at the time of the LGM to be significantly  
907 impaired by non-climatic plant growth factors, such as atmospheric CO<sub>2</sub> concentration, that were significantly  
908 different compared to modern. Our results are also much closer to climate model simulations than previous studies  
909 using the Neural Network method (a variant on MAT), which suggested a climate that was much cooler and drier.  
910 The difference between our MAT reconstruction and the Neural Network reconstruction is probably attributable  
911 to the much smaller size and spatial bias of the modern calibration dataset that was used in this earlier analysis.

912

913 We also found little difference between our MAT reconstruction and a Chironomid-based summer temperature  
914 record using downcore sample by sample comparison, as well as records from a variety of other proxies based on



915 a more general site by site comparison at a regional scale. However, it is notable that some studies using glacier  
916 mass balance modelling methods indicate LGM temperatures that are much cooler than our pollen-based  
917 reconstruction as well as reconstructions based on other proxies. The reasons behind this are not clear, but our  
918 pollen-based results indicate higher than present precipitation in some areas that could explain low altitude LGM  
919 ELA's without the need for such cold temperatures. We also find that although our pollen-based reconstruction  
920 and those of SST's generally agree in terms of mean annual temperatures, SST's indicate greater cooling in  
921 summer compared to winter, while terrestrial records indicate greater cooling in winter compared to summer.  
922 These differences area also reproduced in climate models, and probably reflect the different processes driving  
923 seasonal temperature change in the terrestrial and marine domain.

924

925 Reconstructions of precipitation show large spatial and seasonal variability, but generally indicate less overall  
926 aridity than previously suggested from smaller scale studies which sampled less of the spatial domain. We find  
927 that in some regions of Southern Europe precipitation may actually have been greater than present, especially in  
928 summer, but also in winter in southern and eastern Iberia and around the southern slopes of the Alps. This may  
929 have important implications in understanding the development of LGM glaciation, which may be less a function  
930 of temperature than previously supposed. This could also help better explain the observed asynchronous nature  
931 of glaciation even within relatively small regions such as Europe, as a result of more localized controls on ice  
932 sheet development such as precipitation.

933

#### 934 **Code/Data availability**

935

936 All of the data shown in the figures together with the fossil and modern pollen datasets will be made available on  
937 pangaea.de once the review process has been completed and these datasets are therefore no longer subject to  
938 change.

939

#### 940 **Author contribution**

941

942 BASD designed the study, undertook the analysis and wrote the manuscript. MF and ER designed and prepared  
943 the maps. JOK and AB reviewed the manuscript and provided additional input.

944

#### 945 **Competing interests**

946

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

948

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950

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1637 Tables

Site	Site Name	Country/Ocean	Latitude	Longitude	Elevation	Site Type	Data Type	Samples	Source	Reference
1	MD95-2039 (M)	Atlantic	40.578333	-10.348333	-3381	Marine	Raw Count	21	EPD (E#1472)	Roucoux et al. 2005
2	SU81-18 (M)	Atlantic	37.77	-9.82	-3135	Marine	Raw Count	10	ACER	Turon et al. 2003
3	MD99-2331 (M)	Atlantic	41.15	-9.68	-2110	Marine	Raw Count	41	ACER	Naughton et al. 2006
4	Carn Morval	United Kingdom	49.926111	-6.313889	5	Lake	Digitised	1	Publication	Scourse 1991
5	Gorham Cave	Spain	36.132826	-5.347358	0	Cave	Digitised	1	Publication	Carrion et al. 2008
6	Dozmary Pool	United Kingdom	50.5347222	-4.5358333	265	Lake	Raw Count	32	Author	Kelly et al. 2010
7	Bajondillo	Spain	36.619722	-4.496389	20	Cave	Raw Count	1	EPD (E#1570)	Cortes-Sanchez et al 2011
8	Laguna del maar de Fuentillejo	Spain	38.937996	-4.0539	637	Lake	Digitised	1	Publication	Ruiz-Zapata et al. 2009
9	Padul-1	Spain	37.016338	-3.608503	785	Peat Bog	Digitised	13	Publication	Pons & Reille 1988
10	Padul-2	Spain	37.010833	-3.603889	726	Peat Bog	Digitised	1	Publication	Camuera et al. 2019
11	Cova di Carihuela	Spain	37.4489	-3.4297	1020	Cave	Digitised	1	Publication	Carrion 1992
12	Ifri El Baroud	Morocco	34.75	-3.3	539	Cave	Digitised	1	Publication	Poti et al. 2019
13	MD95-2043 (M)	Mediterranean	36.14	-2.621	-1841	Marine	Raw Count	7	ACER	Fletcher et al. 2008
14	San Rafael	Spain	36.773611	-2.601389	0	Peat Bog	Raw Count	2	EPD (E#574)	Pantaléon-Cano 1997
15	Siles	Spain	38.24	-2.3	1320	Lake	Digitised	1	Publication	Carrion 2002
16	Torreclilla de Valmadrid	Spain	41.4469444	-0.895	570	Colluvium	Digitised	1	Publication	Valero-Garces et al. 2004
17	Navarrés-1	Spain	39.1	-0.683333	225	Peat Bog	Raw Count	1	EPD (E#469)	Carrion & Dupré-Olivier 1996
18	Navarrés-2	Spain	39.1	-0.683333	225	Peat Bog	Raw Count	1	EPD (E#470)	Carrion & Dupré-Olivier 1996
19	Tourbiere de l'Estarres	France	43.0933	-0.3792	356	Lake	Digitised	1	Publication	Jalut et al. 1988
20	Cova de les Malladetes	Spain	39.058	-0.321	20	Cave	Digitised	1	Publication	Dupré Ollivier 1988
21	Lourdes	France	43.033333	-0.075	430	Lake	Digitised	15	Publication	Reille & Andrieu 1995
22	Lake Estanya	Spain	42.0333333	0.5333333	670	Lake	Digitised	1	Publication	Vegas-Villarubia et al. 2013
23	Freychinède	France	42.7833	1.4333	1350	Lake	Digitised	1	Publication	Jalut et al. 1992
24	Banyoles	Spain	42.133333	2.75	173	Lake	Raw Count	13	EPD (E#931)	Pérez-Obiol & Julia 1994
25	Lac du Bouchet B5	France	44.916667	3.783333	1200	Lake	Digitised	14	Publication	Reille & de Beaulieu 1988
26	MD99-2348 (103) (M)	Mediterranean	42.692778	3.841667	-296	Marine	Raw Count	41	EPD (E#1474)	Beaudouin et al. 2007
27	Les Echets G	France	45.9	4.93	267	Peat Bog	Digitised	136	ACER	de Beaulieu & Reille 1984
28	La Grotte Walou	Belgium	50.585278	5.536389	252	Cave	Digitised	1	Publication	Dambon 2011
29	Bergsee	Germany	47.572222	7.93638889	382	Lake	Digitised	1	Publication	Duprat-Qualid et al. 2017
30	Garaat El-Ouez	Algeria	36.818333	8.33333	45	Peat Bog	Raw Count	6	EPD (E#1501)	Benslama et al 2010
31	Pian del Lago	Italy	44.321561	9.485682	833	Lake	Digitised	1	Publication	Guido et al. 2020
32	Pilsensee	Germany	48.0267	11.1883	534	Lake	Digitised	1	Publication	Küster 1995
33	Orgiano	Italy	45.29	11.43	19	Peat Bog	Digitised	1	Publication	Paganelli 1996
34	Lago della Costa	Italy	45.2702778	11.7430556	7	Lake	Digitised	8	Publication	Kaltenrieder et al. 2009
35	Lagaccione	Italy	42.566667	11.85	355	Lake	Raw Count	7	ACER	Magri 1999
36	Lago Vico	Italy	42.3166667	12.1666667	510	Lake	Digitised	15	Publication	Magri & Sadori 1999
37	Stracciaccappa	Italy	42.13	12.32	220	Lake	Raw Count	2	ACER	Giardini 2007
38	Lago di Monterosi	Italy	42.2166667	12.4333333	237	Lake	Raw Count	1	Publication	Bonatti 1970
39	Venice	Italy	45.629523	12.654086	0	Peat Bog	Digitised	1	Publication	Miola et al. 2006
40	Azzano Decimo	Italy	45.8833	12.7165	10	Alluvial Fan	Raw Count	6	ACER	Pini et al. 2009
41	Valle di Castiglione	Italy	41.89	12.75	44	Lake	Raw Count	2	ACER	Follieri et al. 1989
42	Travesio	Italy	46.2	12.87	220	Lake	Digitised	1	Publication	Monegato et al. 2007
43	Orvenco	Italy	46.252088	13.169771	380	Alluvial Fan	Digitised	1	Publication	Monegato et al. 2007
44	Rio Doidis	Italy	46.12	13.19	152	Lake	Digitised	1	Publication	Monegato et al. 2007
45	Billerio	Italy	46.22	13.21	300	Lake	Digitised	1	Publication	Monegato et al. 2007
46	Kersdorf-Briesen	Germany	52.333704	14.269142	44	Lake	Digitised	1	Publication	Strahl 2005
47	Lago Grande di Monticchio	Italy	40.944444	15.6	1326	Lake	Raw Count	6	EPD (E#932)	Watts et al. 1996
48	Nagyomhós	Hungary	48.326944	20.436389	297	Peat Bog	Raw Count	14	Publication	Magyari et al. 1999
49	Safarka	Slovakia	48.8819444	20.575	600	Peat Bog	Digitised	1	Publication	Jankovska 2008
50	Fehér Lake	Hungary	46.45	20.65	86	Lake	Raw Count	10	Publication	Magyari et al. 2014
51	Ioannina	Greece	39.75	20.85	470	Peat Bog	Raw Count	20	ACER	Tzedakis et al. 2004
52	Kokad	Hungary	47.4027778	21.9286111	112	Peat Bog	Raw Count	2	Publication	Magyari et al. 2019
53	Lake Xinias	Greece	39.05	22.27	500	Lake	Raw Count	5	EPD (E#976)	Bottema 1979
54	Mickunai	Lithuania	54.722114	25.532218	143	Lake	Digitised	1	Publication	Satkunas & Grigienė 2012
55	Lake Sfânta Anna	Romania	46.1263889	25.880556	946	Lake	Digitised	1	Publication	Magyari et al. 2014
56	Megali Limni	Greece	39.1	26.3	323	Lake	Digitised	1	Publication	Margari et al. 2009
57	Straldzha	Bulgaria	42.630278	26.77	138	Peat Bog	Raw Count	3	Publication	Connor et l. 2013
58	MD01-2430 (M)	Turkey	40.796833	27.725166	-580	Marine	Digitised	1	Publication	Valsecchi et al. 2012
59	Lake Iznik	Turkey	40.433889	29.533056	88	Lake	Raw Count	7	EPD (E#714)	Miebach et al 2016
60	M72/5 628-1 (M)	Black Sea	42.1035	36.62383	-418	Marine	Raw Count	6	Pangaea (833387)	Shumilovskikh et al. 2014
61	Dziguta	Georgia	42.99	41.07	35	Peat Bog	Digitised	1	Publication	Arslanov et al. 2007
62	Lake Van LG	Turkey	38.667	42.669	1649	Lake	Raw Count	10	Pangaea (853779)	Pickarski et al. 2015
63	Lake Zeribar	Iran	35.533333	46.116667	1286	Lake	Raw Count	17	EPD (E#714)	van Zeist & Bottema 1977

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1640 Table 1. List of selected sites

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Site	Site Name	COHMAP Quality	COHMAP							Upper 14C	Upper Cal. BP	Lower 14C	Lower Cal. BP
			< 17k	18k	19k	20k	21k	22k	23k				
1	MD95-2039 (M)	3C								14830±80	18166±269	19950±210	23883±374
2	SU81-18 (M)	2C								17510±270	20952±404	21250±280	25420±441
3	MD99-2331 (M)	2C								16170±130	19325±303	19770±170	23682±336
4	Carn Morval	4C									18600±3700	21500±890/800	25862±1127
5	Gorham Cave	4D										18440±160	22053±341
6	Dozmary Pool	2C								14568±129	17569±523	18325±216	21769±602
7	Bajondillo	1C									18701±2154		
8	Laguna del maar de Fuentillejo	5D								16540±90	19847±308		
9	Padul-1	3D								18300±300	21821±412	19100±160	22922±308
10	Padul-2	1D										17450±539	21082±539
11	Cova di Carihuela	2C								15700±220	18958±280	21430±130	25659±226
12	Ifri El Baroud	2D								17296±87	20761±293		
13	MD95-2043 (M)	2C								15440±90	18533±294	18260±120	21951±335
14	San Rafael	3D								9980±60	11464±133	16860±120	20083±292
15	Siles	2D								17030±80	20345±351		
16	Torrecilla de Valmadrid	2D								17100±85	20456±366		
17	Navarrés-1	4D								18360±195	22001±353	20700±295	24664±411
18	Navarrés-2	5D								5150±50	5881±85	16000±	19144±
19	Tourbiere de l'Estaries	1C								17150±250	20522±470	18970±160	22847±317
20	Cova de les Malladetes	5D								16300±1500	19686±1723		
21	Lourdes	4D								18510±130	22121±130	20025±175	23952±355
22	Lake Estanya	5D									9498±50		19184±251
23	Freychinède	3C								14800±800	17912±856	21300±760	25615±1030
24	Banyoles	4C									19878±100		27862±3000
25	Lac du Bouchet B5	2C								15350±350	18513±435	19200±300	23006±384
26	MD99-2348 (103) (M)	1D								17660±60	21065±310	19350±90	23111±271
27	Les Echets G	1C								17530±270	20970±407	18030±250	21704±473
28	La Grotte Walou	1D										21200±700	21200±700
29	Bergsee	2D										17780±90	21244±306
30	Garaat El-Ouez	2C								16010±320	19200±801		21260±320
31	Pian del Lago	2D											
32	Pilsensee	6D								15860±250	19073±290		
33	Orgiano	2D								17760±160	21221±373	19290±520	23141±621
34	Lago della Costa	2C								15400±150	18484±330	19285±160	23052±302
35	Lagaccione	2C								16080±450	19369±527	20615±940	24746±1201
36	Lago Vico	3C								14385±140	17541±272	20500±230	24430±376
37	Stracciaccappa	4C								12060±130	14093±281	19745±820	22675±955
38	Lago di Monterosi	2D								17040±350	20398±544		
39	Venice	5D										18640±100	22277±336
40	Azzano Decimo	2D								18000±300	21637±529	21025±245	25179±449
41	Valle di Castiglione	3C								14220±145	17443±270	20300±700	24266±842
42	Travesio	5D										18780±200	22483±406
43	Orvenco	2D								17760±160	21221±373	19290±520	23141±621
44	Rio Doidis	5D										18860±190	22390±373
45	Billerio	3D										18165±200	21872±382
46	Kersdorf-Briesen	1D										17622±94	21183±356
47	Lago Grande di Monticchio	2C									20204±		24014±
48	Nagymohos	2C								14246±144	17361±425	18159±247	21735±622
49	Safarka	3D										18287±1512	21912±1781
50	Feher Lake	1D								17715±250	21190±463	19911±81	23841±313
51	Ioannina	3C								15330±140	18420±312	20760±230	24748±330
52	Kokad	5D								14326±63	17433±443	16280±90	19685±538
53	Lake Xinias	6C								11150±130	13049±160	21390±430	25671±648
54	Mickunai	1D									21000±2200		
55	Lake Sfanta Anna	1D								17626±96	20955±432		
56	Megali Limni	6D								19072±237	22906±340		
57	Straldzha	6C								14696±65	18022±364	23653±114	28580±390
58	MDO1-2430 (M)	4C								12050±75	14904±324	18310±380	21746±968
59	Lake Izniik	7D								16910±100	19515±115		
60	M72/5 628-1 (M)	2C								16835±85	18490±	19495±90	21280±
61	Dziguta	4C								12990±160	15839±483	20560±880	24666±1126
62	Lake Van LG	2C									18590±62		23290±596
63	Lake Zeribar	4C								13650±160	16610±399	22000±500	26462±880

COHMAP chronological quality classification:

- 1C: Bracketing dates within 2000 14C (2360 Cal.) yr interval about the time being assessed
- 2C: Bracketing dates, one within 2000 14C (2360 Cal.) yr and the second within 4000 14C (4682 Cal.) yr of the time being assessed
- 3C: Bracketing dates within 4000 14C (4682 Cal.) yr interval about the time being assessed
- 4C: Bracketing dates, one being within 4000 14C (4682 Cal.) yr and the second being within 6000 14C (7490 Cal.) yr of the time being assessed
- 5C: Bracketing dates within 6000 14C (7490 Cal.) yr interval about the time being assessed
- 6C: Bracketing dates, one within 6000 14C (7490 Cal.) yr and the second within 8000 14C (9681 Cal.) yr of the time being assessed
- 7C: Poorly dated
- 1D: Date within 250 14C (206 Cal.) yr of the time being assessed
- 2D: Date within 500 14C (684 Cal.) yr of the time being assessed
- 3D: Date within 750 14C (975 Cal.) yr of the time being assessed
- 4D: Date within 1000 14C (1123 Cal.) yr of the time being assessed
- 5D: Date within 1500 14C (1881 Cal.) yr of the time being assessed
- 6D: Date within 2000 14C (2360 Cal.) yr of the time being assessed
- 7D: Poorly dated

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Table 2. Chronological control



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	RMSE	R2
<b>TANN</b>	2.28	0.9
<b>TDJF</b>	3.35	0.91
<b>TJJA</b>	2.21	0.81
<b>PANN</b>	224.94	0.69
<b>PDJF</b>	78.51	0.69
<b>PJJA</b>	52.49	0.75
<b>Tree Cover</b>	21.03	0.52

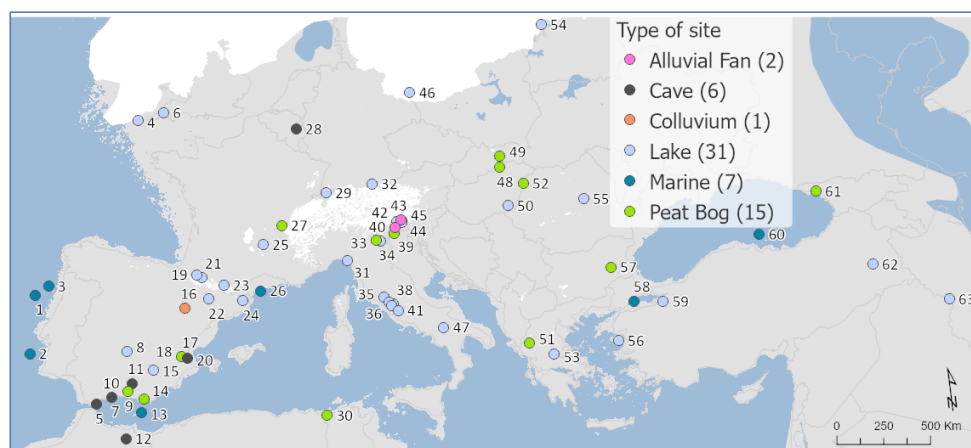
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Table 3. MAT performance statistics based on the modern pollen sample training set. This includes Mean Annual Temperature and Precipitation (TANN and PANN), Mean Winter Temperature and Precipitation (TDJF and PDJF) and Mean Summer Temperature and Precipitation (TJJA and PJJA).



1677 **Figures**

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1681 Figure 1. Site locations and archives (Site numbers are as shown in Table 1)

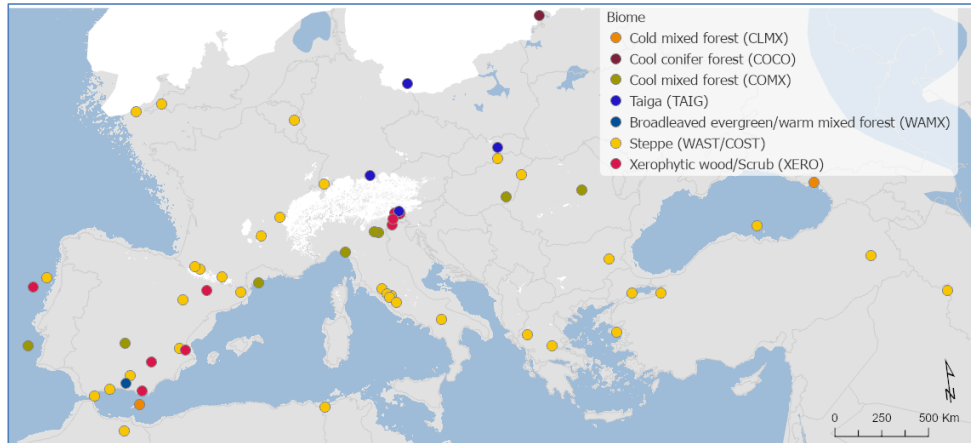
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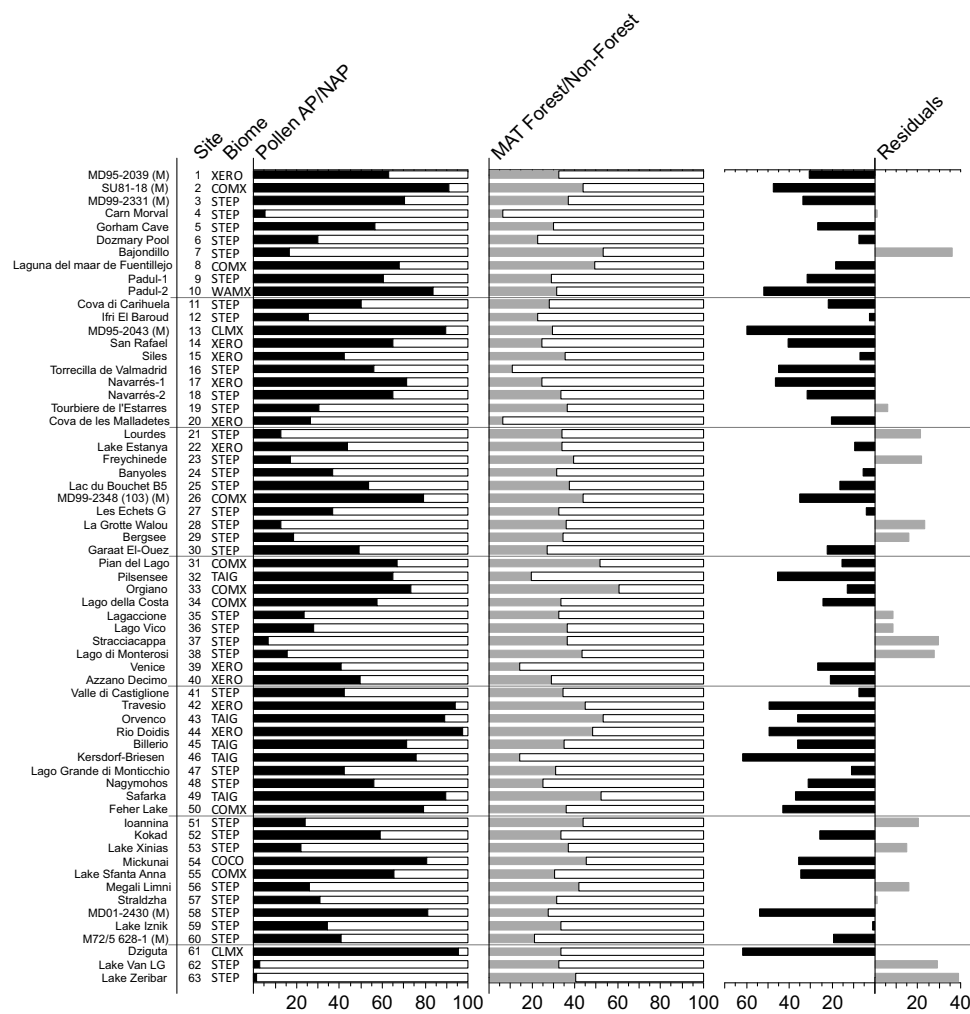
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1688 Figure 2. Pollen biomes

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1693 Figure 3. Pollen biomes (see figure 2 for key), Arboreal Pollen (AP) % forest cover, MAT % forest cover and

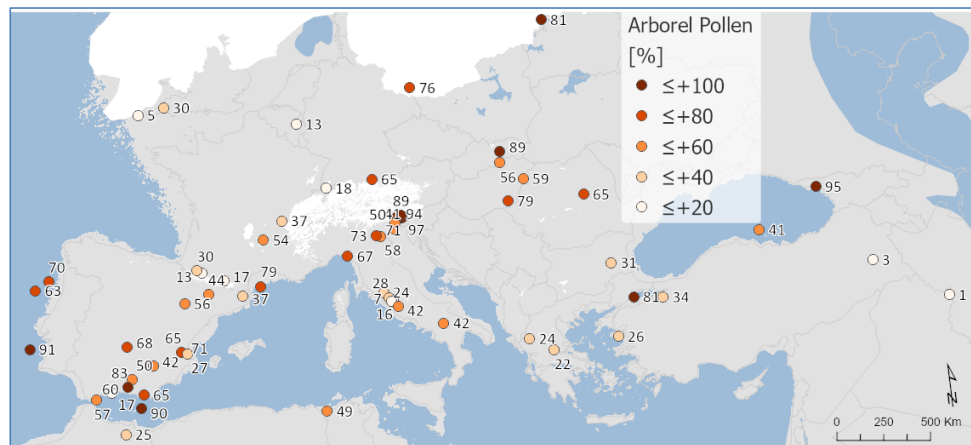
1694 residuals (AP % compared to MAT Forest %)

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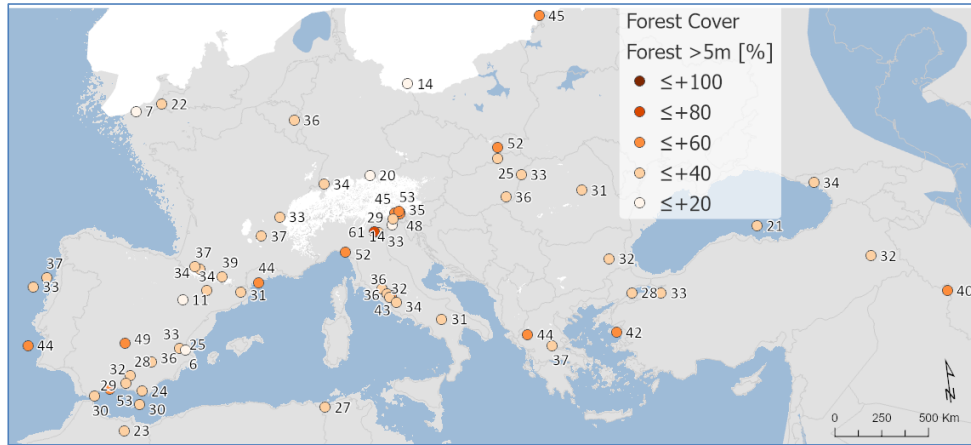
1700 Figure 4. Arboreal Pollen (AP) % forest cover

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1706 Figure 5. Modern Analogue Technique (MAT) % forest cover

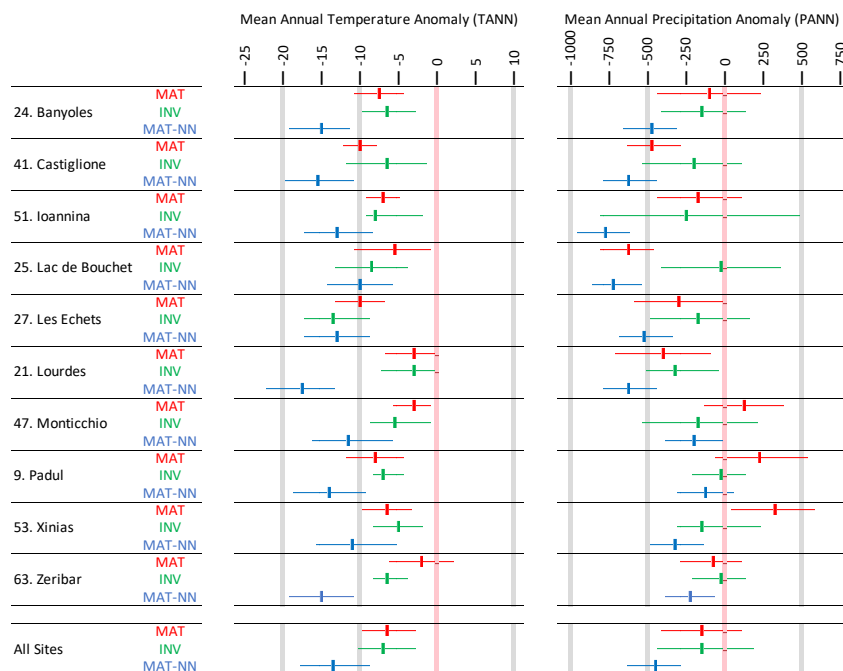
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1715 Figure 7. A site-by-site comparison between LGM pollen-climate reconstructions based on Modern Analogue

1716 Technique MAT (this study), neural-networks MAT-NN (Peyron et al., 1998), and Inverse Modelling INV (Wu

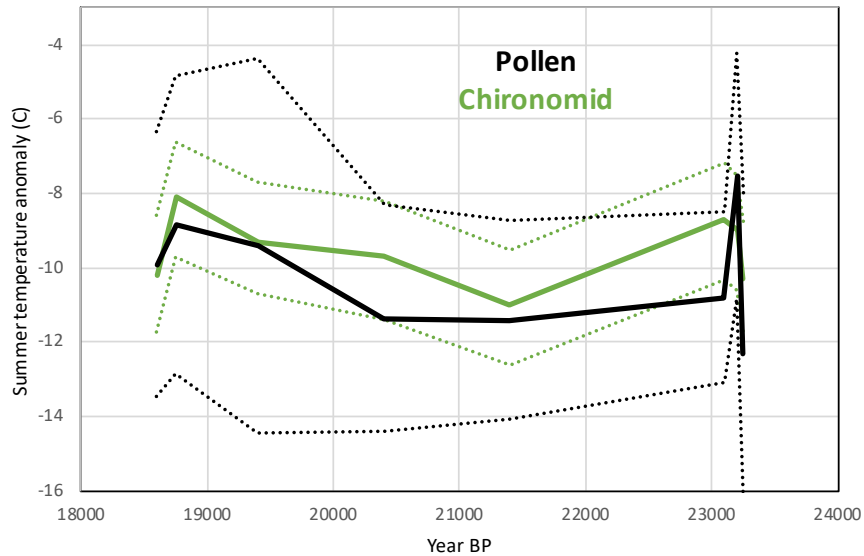
1717 et al., 2007). The results show that MAT and INV give similar climate reconstructions, but MAT-NN is

1718 significantly cooler/drier.

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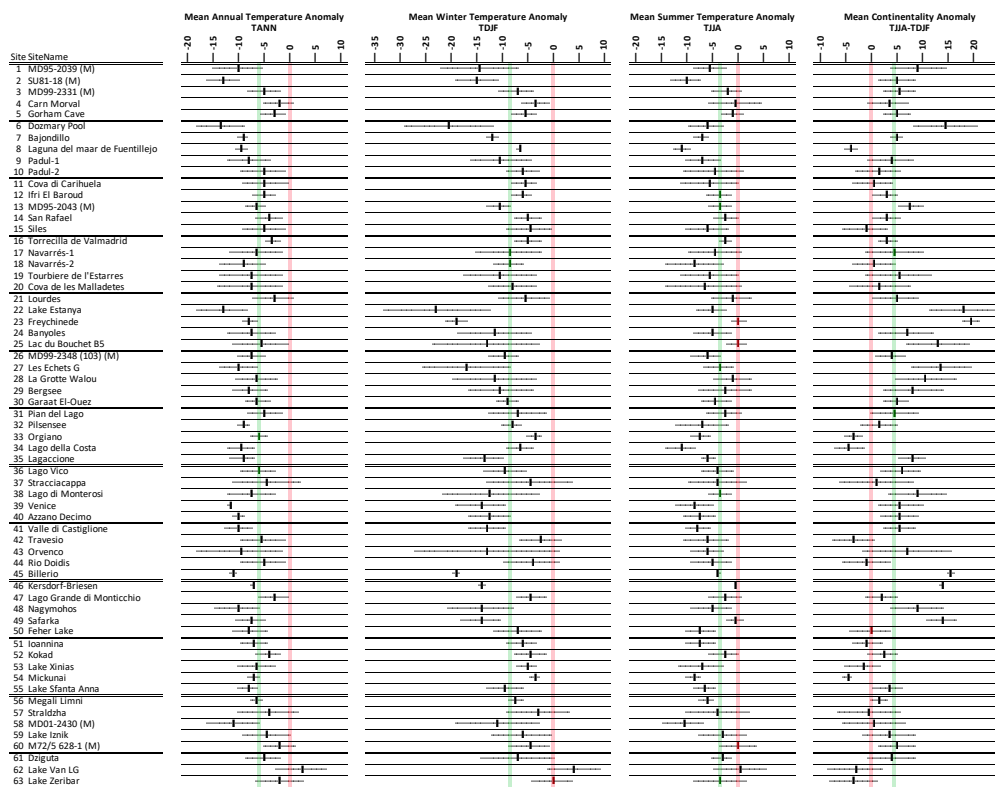


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Figure 8. Comparison between LGM pollen-climate MAT and chironomid summer temperature reconstructions at Lago della Costa, Italy (chironomid reconstruction and pollen data from Samartin et al., 2016). Dash lines show uncertainties.



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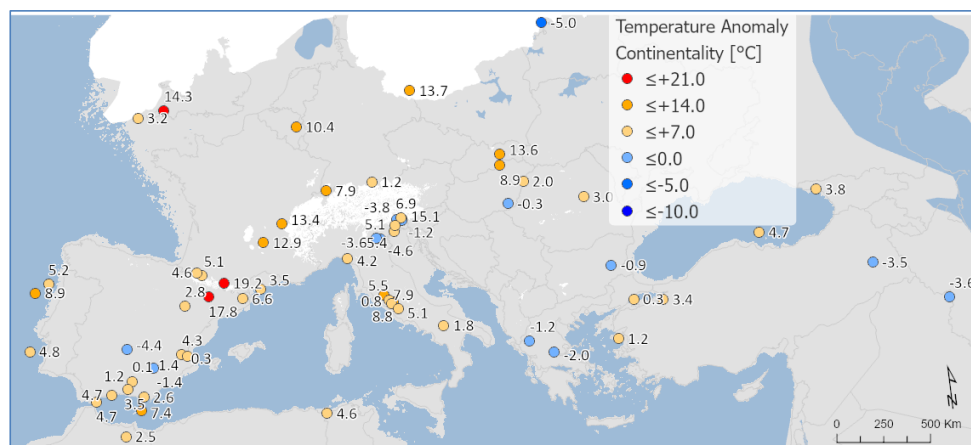
1733 Figure 9. Pollen-based MAT reconstructions for LGM annual, winter and summer temperature anomalies  
 1734 (uncertainties represent one standard deviation). Continentality represents the difference in temperature between  
 1735 summer and winter, with positive anomalies indicating an increase in the temperature difference between  
 1736 summer and winter. All values are expressed as anomalies compared with the present day. The green line indicates the  
 1737 mean for all the sites.

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1745 Figure 10. Maps of pollen-based MAT reconstructions for LGM annual, winter and summer temperature

1746 anomalies (as shown in figure 9). Continentality represents the difference in temperature between summer and

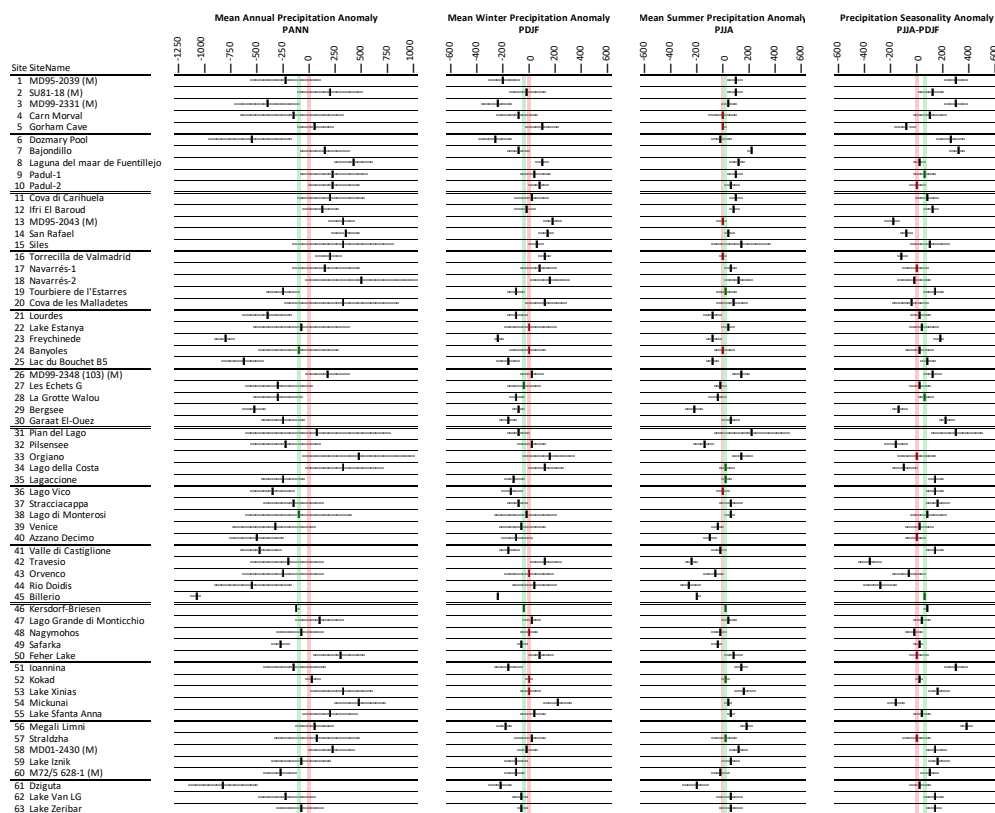
1747 winter, with positive anomalies indicating an increase in the temperature difference between summer and

1748 All values are expressed as anomalies compared with the present day.

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1753 Figure 11. Pollen-based MAT reconstructions for LGM annual, winter and summer precipitation anomalies

1754 (uncertainties represent one standard deviation). Seasonality represents the difference in precipitation between

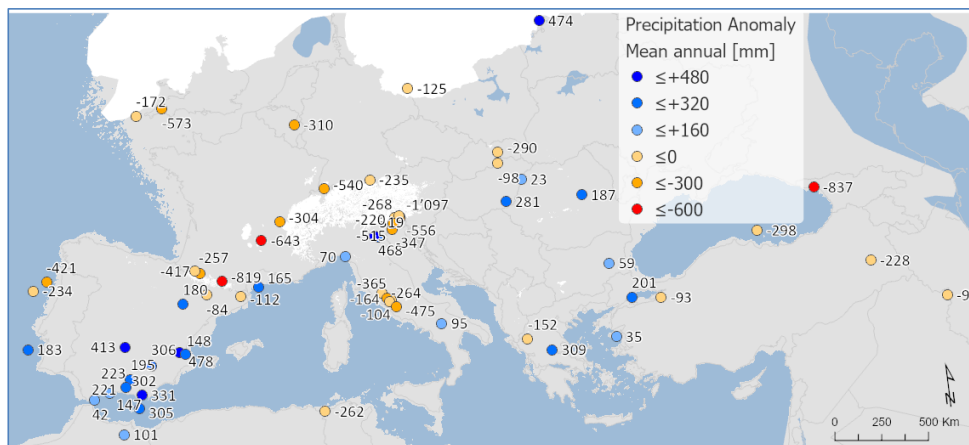
1755 summer and winter, with positive anomalies indicating an increase in summer precipitation compared to winter.

1756 All values are expressed as anomalies compared with the present day. The green line indicates the mean for all

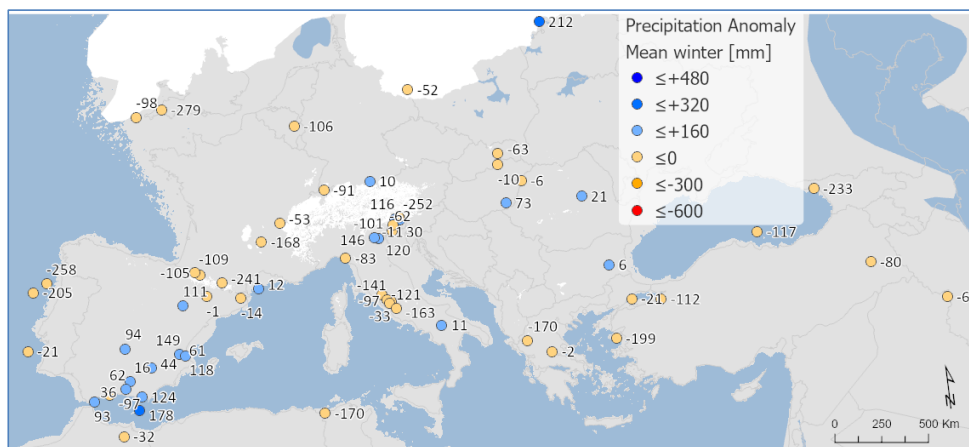
1757 the sites.



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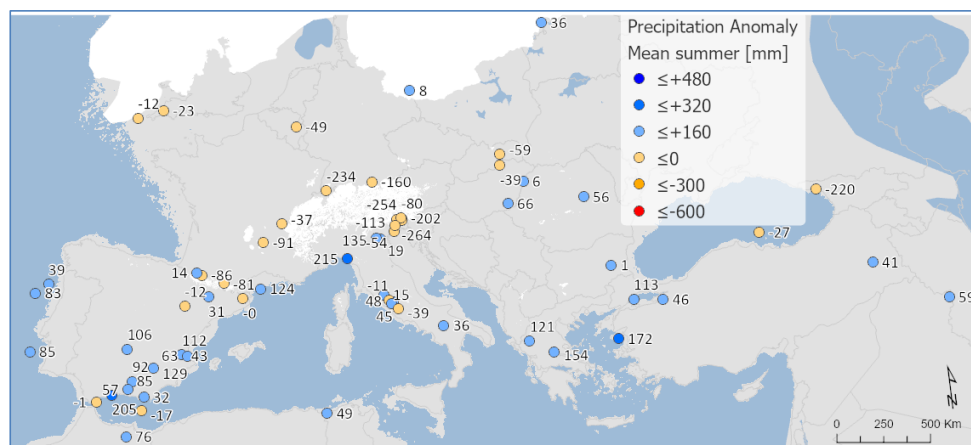


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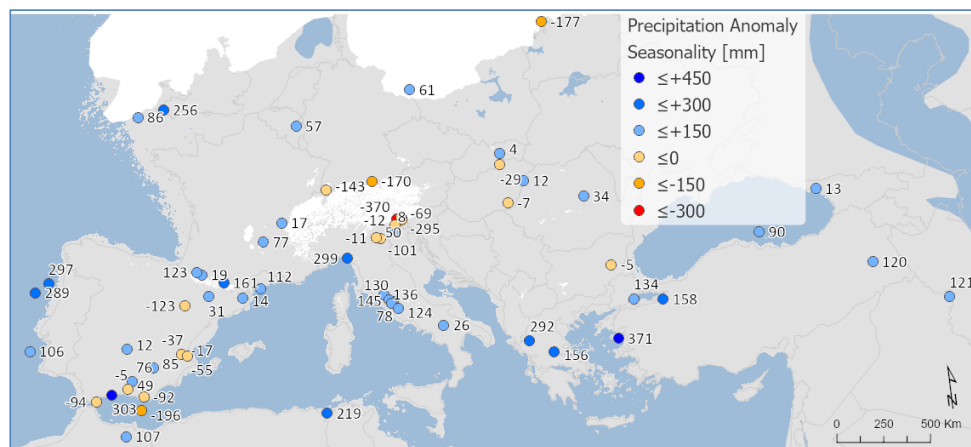




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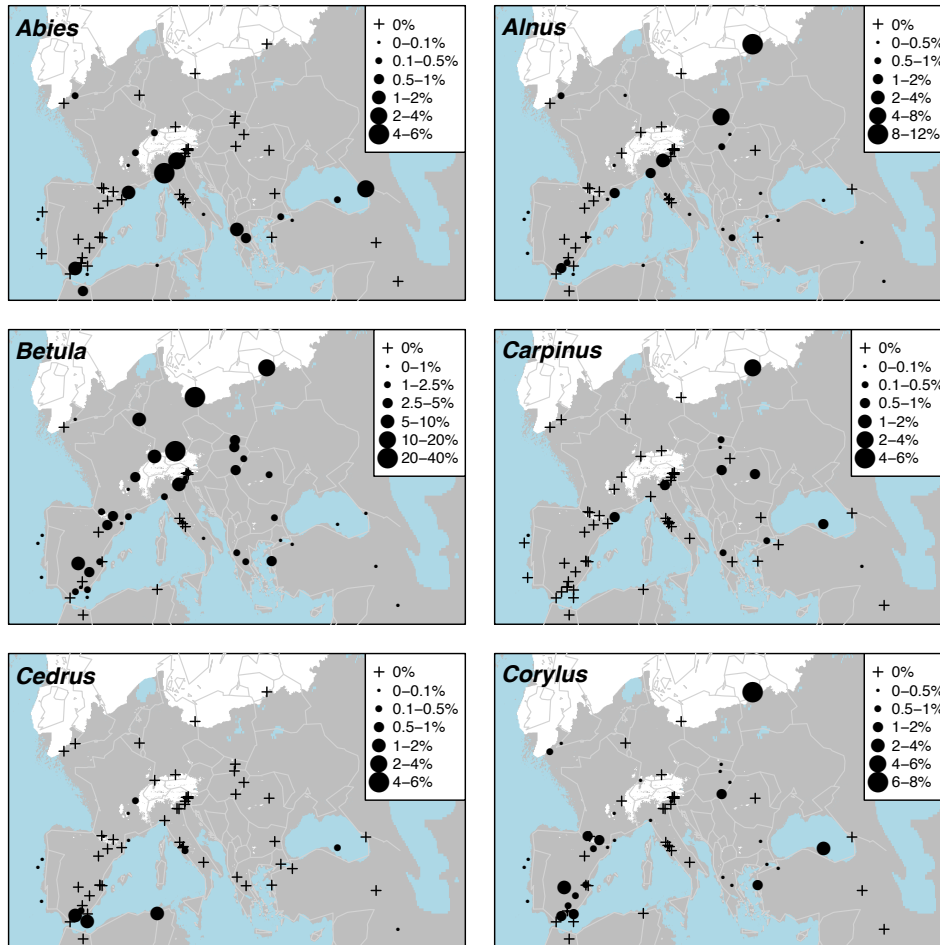
Figure 12. Maps of pollen-based MAT reconstructions for LGM annual, winter and summer precipitation anomalies (as shown in figure 11). Seasonality represents the difference in precipitation between summer and winter, with positive anomalies indicating an increase in summer precipitation compared to winter. All values are expressed as anomalies compared with the present day.



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1770 Appendix

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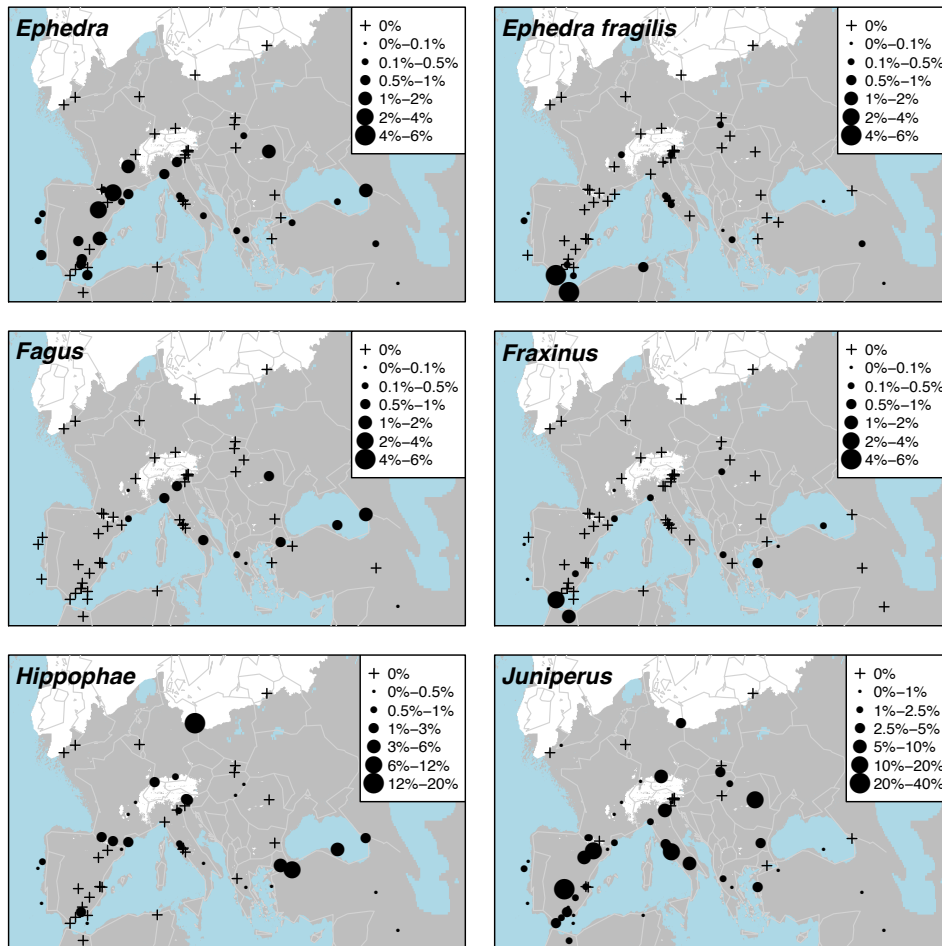


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1774 Figure A1a. Percentage maps of *Abies*, *Alnus*, *Betula*, *Carpinus*, *Cedrus* and *Corylus*

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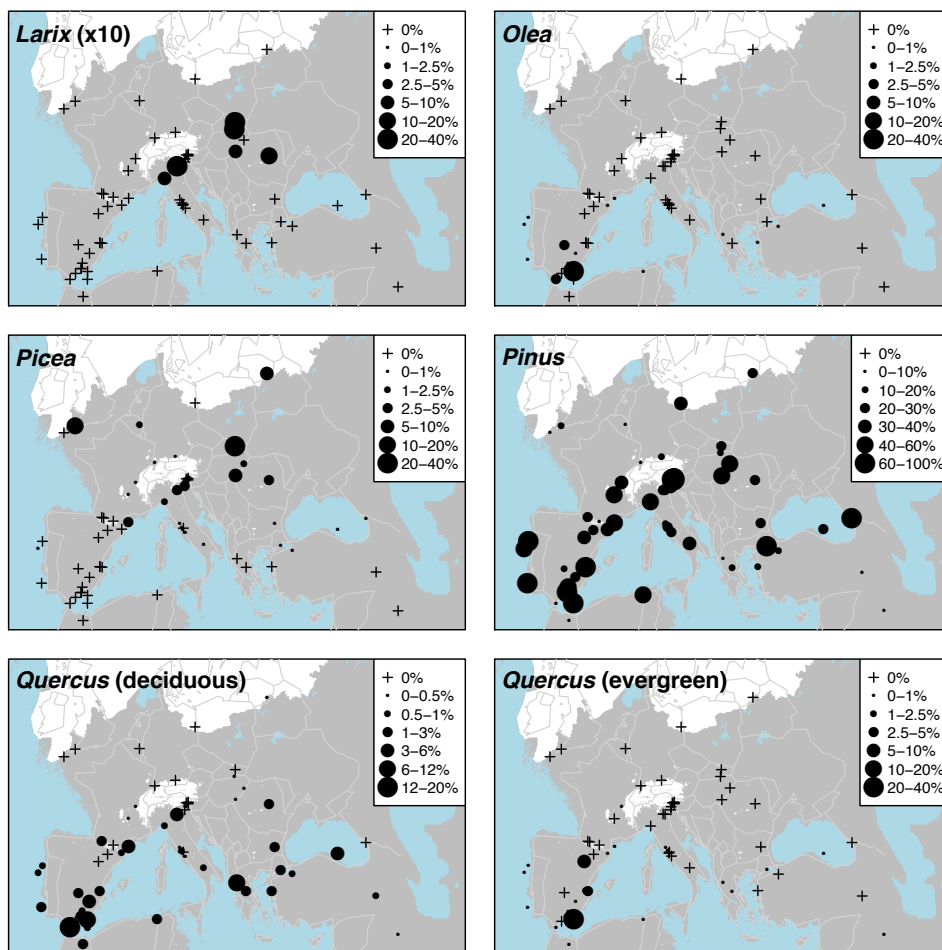


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1778 Figure A1b. Percentage maps of *Ephedra*, *Ephedra fragilis*, *Fagus*, *Fraxinus*, *Hippophae* and *Juniperus*

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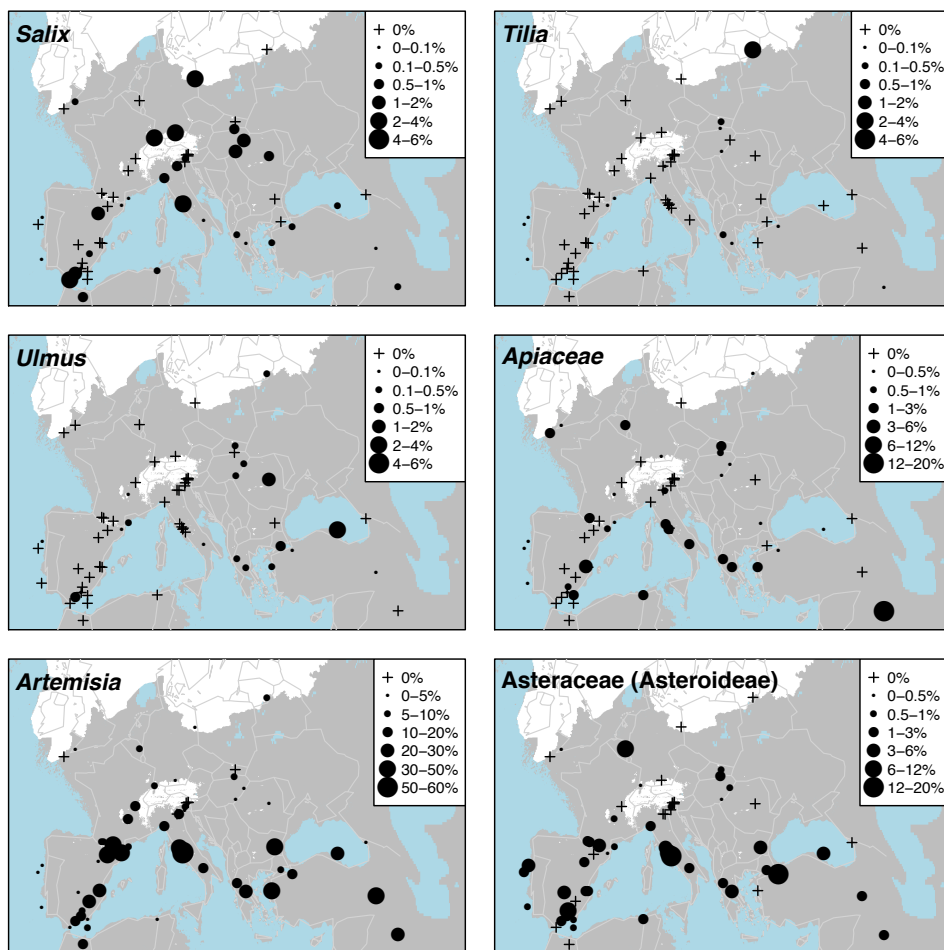
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1782 Figure A1c. Percentage maps of *Larix* (x10 exaggeration), *Olea*, *Picea*, *Pinus*, *Quercus* (deciduous) and *Quercus*

1783 (evergreen)

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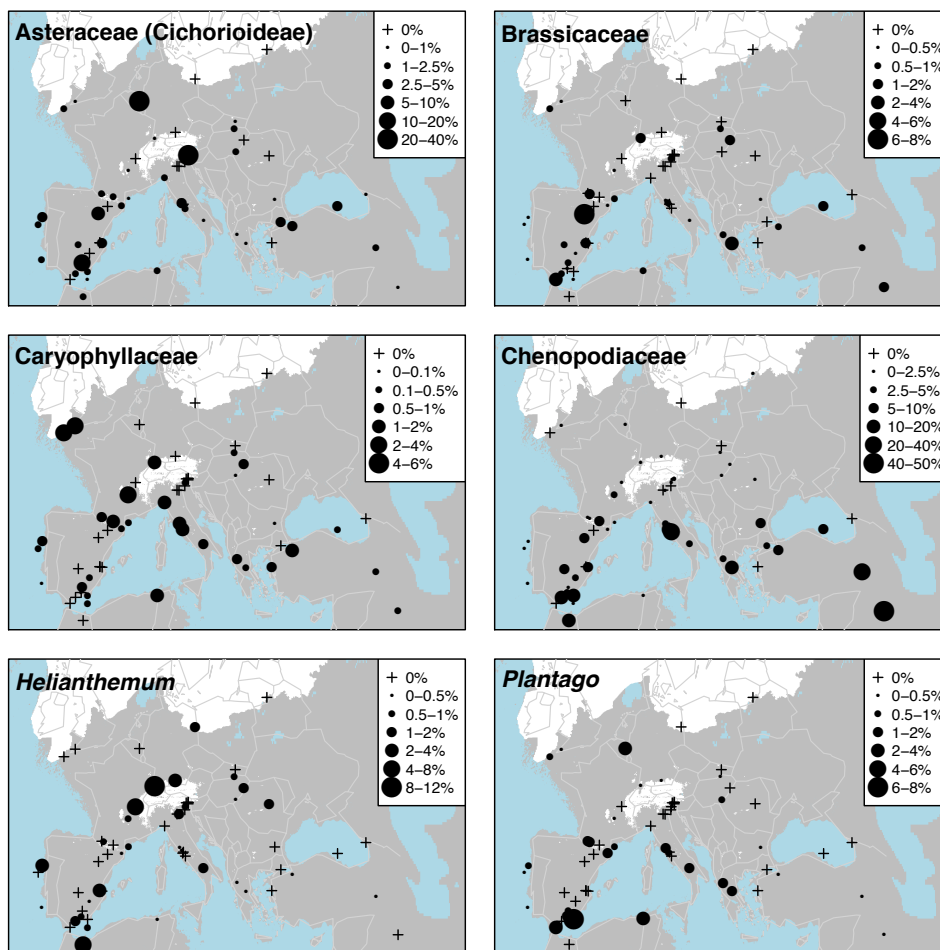


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1787 Figure A1d. Percentage maps of *Salix*, *Tilia*, *Ulmus*, *Apiaceae*, *Artemisia* and *Asteraceae* (*Asteroideae*)





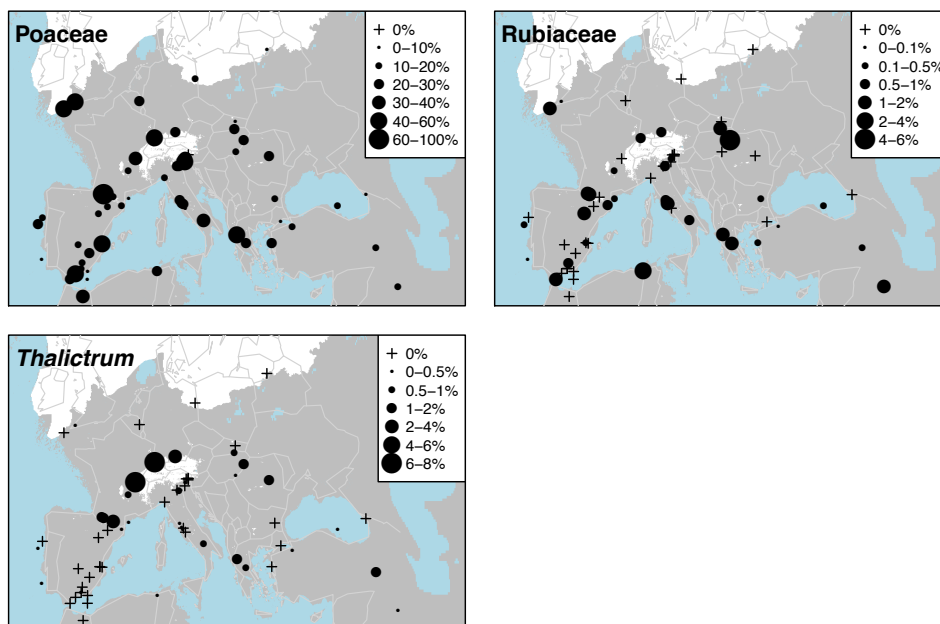
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1790 Figure A1e. Percentage maps of Asteraceae (Cichorioideae), Brassicaceae, Caryophyllaceae, Chenopodiaceae,

1791 Helianthemum and *Plantago*

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1795 Figure A1f. Percentage maps of Poaceae, Rubiaceae and *Thalictrum*

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