

1 Chironomid- and pollen-based quantitative climate reconstructions 2 for the post-Holsteinian (MIS 11b) in Central Europe

3 Tomasz Polkowski¹, Agnieszka Gruszczynska^{1,7}, Bartosz Kotrys², Artur Górecki³, Anna Hrynowiecka⁴,
4 Marcin Żarski⁵, Mirosław Błaszkiwicz¹, Jerzy Nitychoruk⁶, Monika Czajkowska¹, Stefan Lauterbach^{1,8},
5 Michał Słowiński¹

6 ¹Institute of Geography and Spatial Organization Polish Academy of Sciences, Warsaw, 00-818, Poland

7 ²Polish Geological Institute – National Research Institute, Szczecin, 71-130, Poland

8 ³Institute of Botany, Jagiellonian University, Cracow, 30-387, Poland

9 ⁴Polish Geological Institute – National Research Institute, Gdańsk, 80-328, Poland

10 ⁵Polish Geological Institute – National Research Institute, Warsaw, 00-975, Poland

11 ⁶Pope John Paul II State School of Higher Education, Biała Podlaska, 21-500, Poland

12 ⁷Faculty of Physics and Earth System Sciences, Leipzig University, Linnéstraße 5, 04103 Leipzig,
13 Germany

14 ⁸GFZ Helmholtz Centre for Geosciences, Section 4.6 – Geomorphology, Working Group Terrestrial
15 Climate Archives, 14473 Potsdam, Germany

16

17 *Correspondence to:* Tomasz Polkowski (tomasz.polkowski@twarda.pan.pl)

18 **Abstract.** Investigating climatic and environmental changes during past interglacials is crucial to improve our understanding
19 of the mechanisms that govern changes related to current global warming. Among the numerous proxies that can be used to
20 reconstruct past environmental and climatic conditions, pollen allows quantitative reconstructions of annual, warmest month
21 and coldest month air temperatures as well as precipitation sums, and Chironomidae larvae are widely used to infer past
22 summer air temperature. Chironomidae have mostly been used for reconstructing Holocene and Late Weichselian summer
23 temperatures whilst there are only four sites in Europe with chironomid-based summer air temperature reconstructions for the
24 Late Pleistocene and no such records for any Middle Pleistocene warm period as of the writing of this paper. In this study we
25 present the first quantitative palaeoclimate reconstruction for the post-Holsteinian (Marine Isotope Stage - (MIS) 11b) in
26 Central Europe based on both pollen and fossil chironomid remains preserved in palaeolake sediments recovered from Krępa,
27 southeastern Poland. Besides being used for the palaeoclimatic reconstruction, pollen analysis provides the biostratigraphic
28 framework and a broader perspective of climate development at the end of Holsteinian Interglacial. Fossil Chironomidae
29 assemblages at Krępa consist mainly of oligotrophic and mesotrophic taxa (e.g. *Corynocera ambigua*, *Chironomus*
30 *anthracinus*-type) while eutrophic taxa (e.g. *Chironomus plumosus*-type) are less abundant. The chironomid-based summer
31 temperature reconstruction indicates July air temperatures between 15.3 and 20.1°C during the early post-Holsteinian. In any

case, results from Krępa prove that conducting Chironomidae analysis is feasible for periods as early as the Middle Pleistocene, improving our understanding of the mechanisms that control present-day climatic and environmental changes.

1 Introduction

Earth's history is characterised by repeated climate fluctuations, which had not been influenced by humans until the Holocene (marine isotope stage (MIS) 1), the most recent interglacial period. This offers the opportunity to compare natural climatic changes in the past with current ones in order to assess anthropogenic impact on the present climate. With respect to human impact during the Holocene, the so-called "Anthropocene" is widely debated across various scientific disciplines though its exact timing, and the actual dimension of human influence on the environment are still debated (Brondizio et al., 2016). Holocene environmental archives, such as lake, palaeolake and ocean sediments provide material for comprehensive palaeoecological analyses. The sensitivity of some groups of organisms in these archives to changing hydrological or climatic conditions allows reconstruction of past events that directly affected the abundance or structure of the communities (Battarbee, 2000). Species, which are characterised by narrow ecological preferences, such as air temperature, water chemistry or water depth, are used for certain palaeoenvironmental reconstructions (Juggins and Birks, 2012). Many ecological parameters can be reconstructed using different proxies. For example, foraminiferas can be used to reconstruct ocean pH (Foster and Rae, 2016; Roberts et al., 2018), pollen can provide information about vegetation changes (Ralska-Jasiewiczowa et al., 2004; Kupryjanowicz et al., 2018) and can be used to reconstruct past human activity (Chevalier et al., 2020) or past climate conditions (e.g. Rylova and Savachenko, 2005; Hrynowiecka and Winter, 2016). Head capsules of chironomids can serve as the basis for summer air temperature reconstructions (Eggermont and Heiri, 2012) and for assessing the trophic state or pH of freshwater ecosystems (Płóciennik, 2005).

In general, palaeoecological and palaeoclimatological reconstructions record human impact on the environment from the Iron Age (Dumayne-Peaty, 1998; Szal et al., 2014). However, these reconstructions neither provide unequivocal information about air temperature changes nor allow the relative contribution of natural and human drivers to be distinguished. To gain a deeper understanding of the present human impact on climate and environment, it is therefore essential to investigate natural climate variability and environmental changes during past warm periods prior to any anthropogenic impact. In this regard, a particularly suitable targets are interglacial periods, e.g. Holsteinian Interglacial (or Mazovian Interglacial in Poland), which is commonly estimated to have lasted from 423 to 395 ka BP, thus corresponding to MIS 11c (Lauer and Weiss, 2018; Lauer et al., 2020; Fernández Arias et al., 2023). Holsteinian Interglacial is considered the analogue of the Holocene in terms of astronomical parameters (eccentricity, precession, insolation), climatic conditions and greenhouse gases levels (Koutsodendris et al., 2010; Yin and Berger, 2012; Kleinen et al., 2016). To date, there are only a few chironomid-based reconstructions of climatic and ecological conditions for the Middle and Late Pleistocene in Europe available (Engels et al., 2008; Bolland et al., 2021; Ilyashuk et al., 2022; Lapellegerie et al., 2024; Rigterink et al., 2024), but none for the Holsteinian Interglacial and the time thereafter. Hence, knowledge about climatic conditions at this time is mainly derived from pollen data, e.g. from the Praclaux

64 maar in southern France (Reille and de Beaulieu, 1995), Tenaghi Philippon in north-eastern Greece (Tzedakis et al., 2006;
 65 Ardenghi et al., 2019), Lake Ohrid on the North Macedonian-Albanian border (Kousis et al., 2018), Lake Fucino in central
 66 Italy (Vera-Polo et al., 2024) and marine core from ODP site 976 in the Alboran Sea (Sassoon et al., 2023, 2025) . In Central
 67 Europe, high-resolution MIS 11 pollen records are available from the Ossówka palaeolake in eastern Poland (Nitychoruk et
 68 al., 2005, 2018; Bińka et al., 2023) as well as from Nowiny Żukowskie in eastern Poland (Hrynowiecka and Winter, 2016)
 69 and Dethlingen in northern Germany (Koutsodendris et al., 2010). Also notable is Bilhausen in central Germany, which
 70 provided a pollen record for the so-called Bilshausen Interglacial, which might correspond to MIS 11 or MIS 13 (Kühl and
 71 Gobet, 2010). In Northern Europe, there are even fewer records covering MIS 11 e.g. the record from Hoxne in eastern England
 72 (Horne et al., 2023) where temperature reconstructions were performed using chironomids (e.g., (Brooks, 2006), ostracods
 73 (Horne, 2007) and beetle remains (Atkinson et al., 1987).

74 The contemporary state of knowledge on MIS 11 has been reviewed by Candy et al. (2014). Climate conditions in Central
 75 Europe were in general temperate at that time (Nitychoruk et al., 2018), but vegetation reconstructions suggest warmer and
 76 more humid conditions compared to the Holocene climatic optimum (Hrynowiecka and Winter, 2016). Two major climatic
 77 oscillations have so far been documented during the Holsteinian Interglacial, the Older Holsteinian Oscillation (OHO) and the
 78 Younger Holsteinian Oscillation (YHO). The OHO occurred around 418 ka BP (Koutsodendris et al., 2010, 2012; Górecki,
 79 2023) and is clearly connected to a rapid cooling as indicated by the disappearance of temperate vegetation (mostly *Picea*-
 80 *Alnus* forests) and the spread of pioneer tree taxa including *Betula*, *Pinus* and *Larix* (Koutsodendris et al., 2010, 2012; Candy
 81 et al., 2014; Hrynowiecka and Pidek, 2017; Górecki et al., 2022). Although the OHO has been described at multiple sites
 82 across northern Europe (Koutsodendris et al., 2012), it has so far been identified in few southern European sites (Kousis et al.,
 83 2018; Sassoon et al., 2023, 2025). In contrast to the OHO, the YHO occurred around 400 ka BP within the climatic optimum
 84 of the Holsteinian Interglacial (*Carpinus-Abies* phase) and was apparently not connected to a significant cooling (Górecki et
 85 al., 2022). Records from Germany and eastern Poland suggest a sudden regression of *Carpinus* from forest communities
 86 (Koutsodendris et al., 2010; Hrynowiecka et al., 2019; Górecki et al., 2022). Particularly in Poland a rapid spread of *Abies*
 87 with an admixture of *Corylus* is observed, with *Taxus* also found in southern sites (Górecki et al., 2022), suggesting that
 88 temperature was not limiting the growth of *Carpinus*.

89 The climate during Holsteinian Interglacial (MIS 11c) was characterised by relatively stable warm and moist conditions with
 90 global temperatures approximately 1.5–2 °C above the pre-industrial level (Masson-Delmotte et al., 2010). Raymo and
 91 Mitrovica (2012) and Muhs et al. (2012) suggest sea level was possibly 6–13 m higher than present in this period. This can be
 92 partially attributed to the melting of the Greenland Ice Sheet (Robinson et al., 2017), as pollen and palaeoDNA data suggest
 93 the existence of spruce forests in Greenland at this time (Willerslev et al., 2007; de Vernal and Hillaire-Marcel, 2008).

94 In Europe, warm and wet oceanic climate conditions extended far to the east as evidenced by the presence of *Taxus* and *Abies*
 95 pollen at sites in Lithuania (Kondratiene and Gudelis, 1983), Belarus (Mamakowa and Rylova, 2007), and western Ukraine
 96 (Łanczont et al., 2003; Benham et al., 2016), whilst modern distribution limits of these taxa are estimated further to the west
 97 (Benham et al., 2016). Evidence from several terrestrial records from Eurasia suggests that the MIS 11c climate was highly

98 complex, with pronounced climate variability occurring on both centennial and millennial timescales (Koutsodendris et al.,
 99 2010; Prokopenko et al., 2010; Tzedakis, 2010; Oliveira et al., 2016; Tye et al., 2016; Górecki et al., 2022). Holsteinian
 100 Interglacial was followed by gradual cooling period (MIS 11b) which resulted in annual temperature decline and forest
 101 contractions (Tzedakis et al., 2006; Kousis et al., 2018; Hrynowiecka et al., 2019; Sassoon et al., 2025).
 102 The pollen succession of the Holsteinian Interglacial in Poland is characterised by a dominance of *Picea-Alnus* at first , then
 103 *Carpinus* and *Abies*, as well as a significant proportion of *Taxus*. Thermophilic taxa also occur frequently, examples including:
 104 *Pterocarya*, *Celtis*, *Juglans*, *Ilex*, *Carya*, *Parrotia*, *Buxus*, *Vitis*, *Brasenia*, *Trapa*, and *Azolla* (Janczyk-Kopikowa, 1991).
 105 Temperature reconstructions based on the indicator species method suggest the warmest period was the *Carpinus-Abies* phase,
 106 with estimated temperatures of 0–3 °C in January and 21–26 °C in July. This, along with high precipitation created a suitable
 107 environment for the spread of rare warmth-adapted taxa (Krupiński, 1995; Hrynowiecka and Winter, 2016). However,
 108 palaeotemperature reconstructions from Dethlingen (Koutsodendris et al., 2012) suggest slightly lower temperatures in
 109 Western Europe for both January (-2.2 ± 3.1 °C) and July (17.8 ± 2.1 °C). Pollen-based temperature reconstruction from Lake
 110 Ohrid (SE Europe) indicates higher January (MTCO) maximum (4.4 °C) (Kousis et al., 2018).
 111 MIS 11b brought the AP percentages decrease in Central Europe (Hrynowiecka et al., 2019). Lake Ohrid pollen record reveals
 112 the domination of *Pinus* and plant open communities at the time, with Poaceae and *Artemisia* species included (Kousis et al.,
 113 2018). ODP Site 976 pollen-based climate reconstructions shows annual temperature drop to around 10 °C and summer
 114 temperature to 20 °C (Sassoon et al., 2025).
 115 The warm phase of the Holsteinian Interglacial was also confirmed by oxygen isotope analyses on endogenic lake carbonates
 116 (Nitychoruk et al., 2005) and snail shells (Szymanek, 2018). These showed significant changes in climatic conditions
 117 throughout the Holsteinian Interglacial, during which, continental and maritime influences intertwined in Central Europe.
 118 Continental influences resulted in a shortened vegetation period with long winters, whilst the opposite occurred under maritime
 119 influence, i.e. the vegetation period was significantly longer, temperatures were milder and precipitation rates were higher,
 120 also reflected by the appearance of stenothermal plant species (Nitychoruk et al., 2005). Aiming at improving the knowledge
 121 about climate variability at the demise of the Holsteinian Interglacial, we present the first quantitative climate reconstructions
 122 for the post-Holsteinian in Central Europe, based on chironomid and pollen analyses. The aim of analysing this post-interglacial
 123 period is to investigate temperature and vegetation changes and to determine if climate at the time was considerably cooler
 124 than today. This choice was also dictated by Chironomidae head capsules' presence in post-Holsteinian section of the core
 125 (unlike the Holsteinian part). In addition, we discuss the potential of chironomid analysis for palaeoecological study of
 126 Quaternary sediments as well as the challenges for chironomid analysis arising from both the evolution and interchanging
 127 adaptations to species ecological preferences and the preservation of fossil remains.

128 **2 Study site and methods**

129 **2.1 Study area**

130 The Krępa palaeolake sediment succession (51°37'53.2''N, 22°18'38.1''E, 146 m asl.) is located in SE Poland, near the city
131 of Kock, approximately 120 km southeast of Warsaw (Fig. 1). It is under influence of humid continental climate (Dfb) in terms
132 of the Köppen-Geiger climate classification (Peel et al., 2007). Average annual temperature for this region is ~ 8.6 °C, with
133 July mean temperature of ~ 19 °C and January mean temperature of ~ -1 °C, while average annual precipitation is ca. 600 mm
134 (Ustrnul et al., 2021). Geomorphologically, it is situated in the central-eastern part of the North European Plain behind the
135 maximum extent of the Saalian glaciation (Marks et al., 2018) and the sediment core analysed in this paper was obtained on a
136 moraine plateau related to this ice sheet. Holsteinian Interglacial deposits in the area were first identified by Jesionkiewicz
137 (1982) during cartographic work for the 1:50 000 Detailed Geological Map of Poland (DGMP; Sheet 676 - Kock) (Drozd and
138 Trzepla, 2007). On the moraine plateau, the interglacial deposits are found under a thin cover of moraine deposits, whereas at
139 the slopes of the nearby Wieprz River valley, they are exposed directly on the surface. This study's material was obtained from
140 a sediment core that was drilled at Krępa in 2015, using a Geoprobe drilling device (Górecki, 2023).

141 The basal part of the 23.8-m-long sediment core that was recovered from the Krępa sediment succession in 2015 (Fig. 1)
142 consisted of a 2-m-thick layer of light greyish brown sandy clays with a large number of rock fragments (unit 1), which is
143 interpreted as till. As indicated by its stratigraphic position and its petrographic characteristics (Drozd and Trzepla, 2007), this
144 till was likely accumulated during the Elsterian glaciation (Sanian 2 glaciation in Poland), which is considered to correspond
145 to MIS 12. Directly above the till, a 0.6-m-thick layer of laminated sandy silts and sandy-clayey silts is present (unit 2). These
146 sediments are interpreted as the result of glaciolimnic sedimentation in a relatively shallow water body between blocks of dead
147 ice during the recession of the Elsterian ice-sheet. The glaciolimnic sediments of unit 2 gradually turn into a carbonate gyttja
148 with small interlayers of carbonatic-minerogenic gyttja (unit 3), which was most likely deposited in the profundal zone of an
149 already relatively deep lake. Between 1187 and 760 cm core depth, non-carbonatic organic-minerogenic gyttjas are found with
150 mineral content generally increasing towards the top of the core (unit 4). The limnic sediments of unit 4 are interpreted to
151 reflect the gradual shallowing of the lake due to continuous sediment infilling. At the same time, the systematic increase in
152 mineral components in the sediments most likely reflects increased denudation and erosion in the catchment, possibly favoured
153 by reduced vegetation cover in response to a climatic shift towards colder conditions. The gyttja sequence of unit 4 is overlain
154 by a 1.9-m-thick layer of clays (unit 5), which probably represent accumulation in a periglacial lake. The following 1.1-m-
155 thick layer of fine- to medium-grained sands (unit 6) as well as the overlying 3.1-m-thick layer of rhythmically laminated
156 sandy silts (unit 7) are interpreted as proglacial sediments (units 6 and 7) of the transgressing Early Saalian (MIS 6) ice sheet.
157 Above this, the profile is capped by a 1.5-m-thick layer of sandy morainic till with rock fragments (unit 8) related to Saalian
158 glaciation.

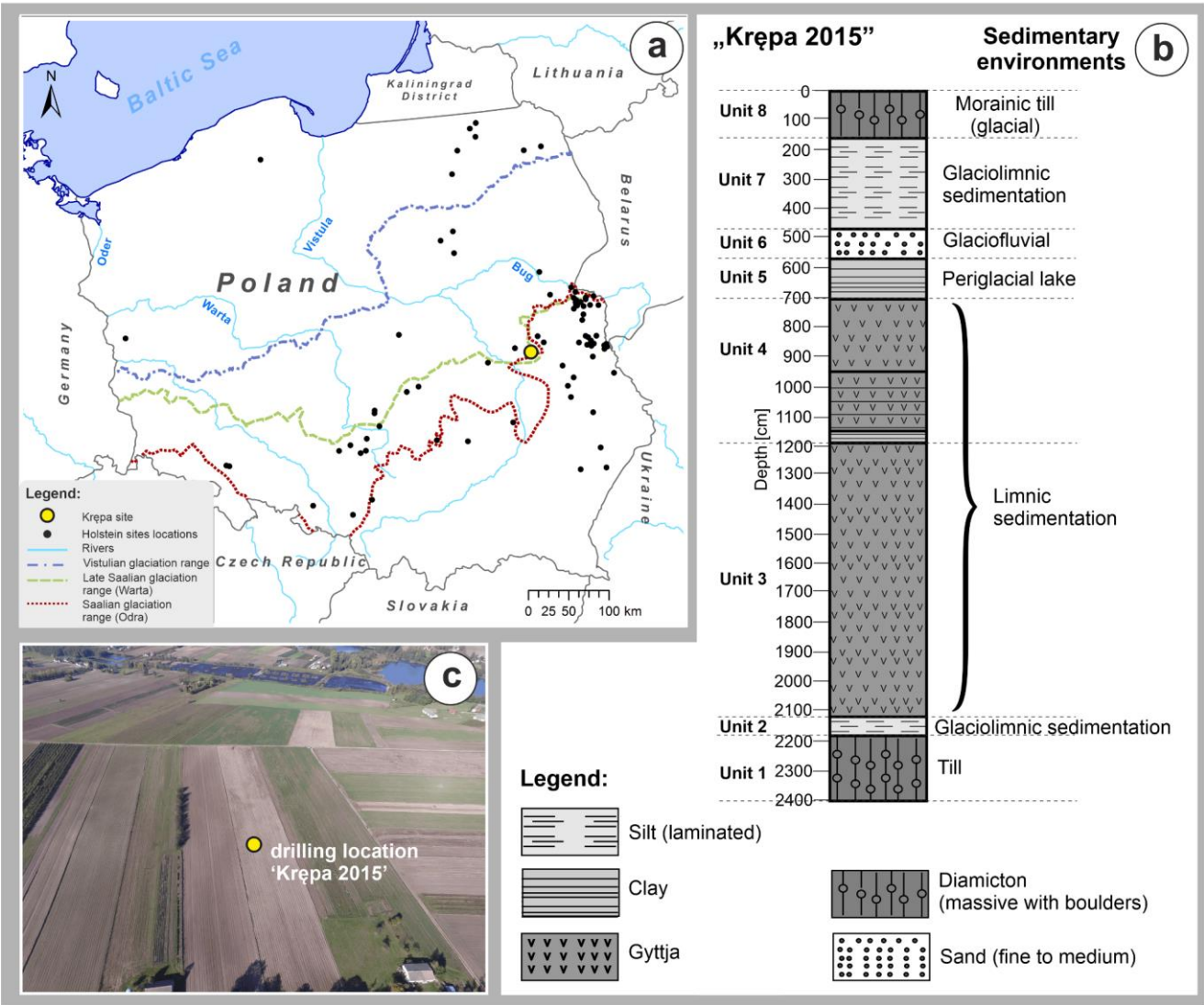


Figure 1: (a) Location of selected sites with deposits from the Holsteinian Interglacial in Poland with the Krepa site indicated by the big yellow dot. Glaciation ranges are based on Źarski et al. (2024), Pochocka-Szwarc et al. (2024) and Marks (2023). (b) Lithological profile of the Krepa sediment succession and (c) location of the drilling site (picture M. Źarski).

2.2 Chironomidae analysis

Initially, 79 sediment samples of 1 cm³, taken between 800 and 2160 cm depth at 5-40 cm intervals, were investigated for the presence of Chironomidae head capsules. However, only 30 of them (965-1155 cm depth) simultaneously contained more than 0-2 individuals. Chemical preparation followed Brooks et al. (2007). The precipitate was initially heated with KOH. The wet sediment was then passed through 212 µm (to remove larger sediment particles) and 100 µm mesh sieves and subsequent residues were treated in an ultrasonic bath for 3 sec. The processed sediment was subsequently examined under a stereomicroscope (Zeiss Axio Lab A1) at 25× magnification. Chironomid head capsules from each sample were picked and mounted in Euparal. In case of damaged head capsules, individuals were counted as one if more than half of a body was preserved. Identification of chironomid head capsules followed Wiederholm (1983), Schmid (1993), Klink and Moller Pillot (2003), Brooks et al. (2007) and Andersen et al. (2013). Ecological preferences of identified taxa are based mainly on Brooks et al. (2007), Brundin (1949), Brodersen and Lindegaard (1999a) and Saether (1979).

Preliminary tests of sample preparation avoided the use of chemicals and included soaking the samples in water for a long time instead to reduce mechanical stress exerted to the head capsules during sample sieving as much as possible. Nevertheless, intact head capsules could not be extracted from some sediment samples even when using this gentle way of sample preparation, likely because of the already highly compacted sediment. As small numbers of head capsules may hinder palaeoecological and palaeoclimatic reconstructions, it was therefore partly necessary to combine samples (see below) or to increase the volume of the analysed sediment material (some samples were even as large as 20 cm³).

Chironomidae subfossil larvae were obtained from a total of 30 samples from the gyttja sediments (unit 4 on Fig. 1). Samples that contained fewer than 50 head capsules were merged except for a solitary sample at 1000 cm core depth. For 5 samples the required number of 50 head capsules was obtained and the remaining 24 samples were merged into seven clusters. After merging, sample clusters at 975 cm, 1080 cm, 1120 cm and 1125 cm core depth still did not reach 50 head capsules, but nonetheless, these samples and the one from 1000 cm core depth were included in the reconstruction as preliminary results seemed credible in terms of obtained temperature values.

2.3 Chironomid-based mean July air temperature reconstruction

In order to reconstruct mean July air temperatures ($T_{\text{Jul-Ch}}$) from the Krępa chironomid assemblage, the Swiss-Norwegian-Polish (SNP) training set (Kotrys et al., 2020) was used as this covers a higher temperature span than other available European training sets (e.g. the Finnish, Russian, Swiss-Norwegian training sets) (Kotrys et al., 2020). The SNP training set includes 357 lakes, 134 taxa, covers a temperature range between 3.5 and 20.1 °C. Weighted averaging-partial least squares transfer function (WA-PLS) was used for performing the reconstruction. The RMSEP for this combined training set is 1.39°C, and the R^2 is 0.91 (Kotrys et al., 2020). Detrended Correspondence (MinDC) was also calculated. The temperature reconstruction was carried out using the C2 (v. 1.6) software (Juggins, 2007).

197 The lowest number of head capsules used for the T_{jul-Ch} reconstruction was 5 individuals at 1070 cm core depth whereas the
198 highest number was 78 at 985 cm core depth. After merging, the total number of samples used for the T_{jul-Ch} reconstruction
199 was 13.

200 **2.4 Pollen analysis**

201 The Krępa sediment core obtained in 2015 was sampled for palynological analyses at 5-cm intervals between 770 and 2180
202 cm depth, totaling 281 samples. A volume of 1 cm³ was collected from organic sediments (peat, gyttja), while minerogenic
203 sediments (clays, silts, sands) were sampled with a volume of 3 cm³ due to the anticipated low pollen grain concentration.
204 Samples were further processed following the standard methodology outlined by Erdtman (1960) with modifications such as
205 the use of HF (Berglund and Ralska-Jasiewiczowa, 1986). Prior to laboratory processing, a *Lycopodium* tablet (Lund
206 University, batch number 100320201, 20,408±543 spores per tablet) was added to each sample to determine the absolute
207 sporomorph concentration (Stockmarr, 1971). Pollen grains were counted using a ZEISS Axio Imager A2 light microscope.
208 Palynomorphs were identified using pollen keys and atlases (Beug, 1961; Stuchlik, 2001, 2002, 2009; Lenarczyk, 2014), as
209 well as online resources (PalDat, 2000; NPP Database, Shumilovskikh et al., 2022). For most samples, counts were conducted
210 up to a sum of 500 pollen grains from arboreal (AP) and non-arboreal (NAP) plants. However, samples from glacial sediments
211 with low palynomorph concentration were counted up to a sum of 300 pollen grains only. Percentages were calculated based
212 on the sum of pollen grains from trees and shrubs (AP), as well as herbaceous plants, and dwarf shrubs (NAP). The results of
213 the palynological analysis are depicted in a simplified pollen diagram (Fig. 3) that was plotted using R Studio with the package
214 riojaPlot (Juggins, 2022). Local Pollen Assemblage Zones (LPAZ) were established using the CONISS cluster analysis
215 function within riojaPlot and were visually adjusted if necessary.

216 **2.5 Pollen-based climate reconstructions**

217 Climate variables reconstructed using pollen data include mean annual air temperature (TANN), mean annual precipitation
218 (PANN), mean temperature of the warmest month (MTWA), and mean temperature of the coldest month (MTCO). All
219 reconstructed climatic factors were based on modern data sourced from the Northern Hemisphere database compiled by
220 Herzschuh et al. (2023a, b). Two reconstruction approaches were applied: the Modern Analog Technique (MAT; (Overpeck
221 et al., 1985; Guiot, 1990) and Weighted Averaging Partial Least Squares regression (WA-PLS; (ter Braak et al., 1993; ter
222 Braak and Juggins, 1993). In the MAT approach, the best number of analogues (k) was chosen by comparing model
223 performance (RMSE and R²) across k values from 1 to 10. This analysis indicated that using k = 7 nearest analogues minimised
224 prediction error, and thus 7 analogues were used in the final MAT reconstructions. For WA-PLS model selection, including
225 the determination of the optimal number of components, was based on predictive accuracy assessed through leave-one-out
226 (LOO) cross-validation and supported by randomization tests, following the methodology outlined by Chevalier et al. (2020).
227 Based on these criteria, a four component WA-PLS was adopted. For each reconstruction model, the coefficient of
228 determination (R²) and root mean square error (RMSE) were calculated to evaluate model performance. To express the

uncertainty in the fossil climate reconstructions, we calculated standard errors of prediction (SEP) and depicted them as error bars in the figures. In the WA-PLS approach, sample-specific SEP were obtained via a bootstrapping implemented in the rioja package (Juggins, 2022). For the MAT model we used the cross-validated RMSE as a uniform error estimate for the fossil MAT reconstructions. Modern pollen data used in the reconstructions were sourced from the Northern Hemisphere database compiled by Herzschuh et al. (2023a, b). To enhance spatial relevance, the modern dataset was geographically filtered to include only samples within a 3000 km radius of the fossil site. This geographic filtering yielded a regional calibration set of 4955 modern pollen samples, out of the original global dataset. From the fossil pollen dataset, only taxa present in at least 50% of the samples and reaching at least 1% pollen value at least once were included. Additionally we ensured taxonomic consistency between the modern and fossil pollen data by harmonizing taxa names and then removing taxa with zero abundance in the filtered modern set. After this filtering, 10 pollen taxa remained in common between the modern calibration set and the fossil record (primarily major arboreal and herb taxa such as *Larix*, *Betula*, *Pinus*, *Salix*, *Picea*, *Juniperus*, *Artemisia*, Asteraceae, Poaceae, and Amaranthaceae). Using only these common taxa helps avoid noise from spurious taxa and improves model robustness. All data processing and modeling were carried out in R (RStudio), making use of the analogue and rioja packages for calibration and reconstruction. The pollen-based reconstructions were restricted to the interval of the succession where chironomid remains were also present and were performed on 44 samples.

3. Results and interpretation

3.1 Ecological reconstruction based on Chironomidae assemblages from the Krępa site

In general, the chironomid assemblages preserved in the Krępa sediments are dominated by two species *Corynocera ambigua* and *Chironomus anthracinus*-type. *Corynocera ambigua* is a species often described as cold-adapted oligotrophic (Fjellberg, 1972; Pinder and Reiss, 1983; Walker and Mathewes, 1988; Brooks et al., 2007; Luoto et al., 2008; van Asch et al., 2012), inhabiting shallow lakes in arctic and subarctic regions, though it is also found in eutrophic lakes (Halkiewicz, 2008; Kotrys et al., 2020). Adults of this species are not able to fly, and breed on the water surface when the temperature reaches approximately 7-8 °C (optimum 13.7 °C). Mothes (1968) concluded that *Corynocera ambigua* larvae develop in autumn and winter, but not during summer. The decline in their numbers may be due to growth of filamentous algae in summer. Larvae of *Corynocera ambigua* are eurythermic, while the pupae are cold-stenothermic (Brundin, 1949). They only reproduce at low temperatures and inhabit water bodies with a maximum depth of approximately 25 m. The abundance of *Corynocera ambigua* has been shown to be correlated with charophyte contents (Brodersen and Lindegaard, 1999b). Although this species does not feed on charophytes, their presence may increase the number of diatoms and stabilise the trophic status and water clarity (Forsberg, 1965; Blindow, 1992). *Corynocera ambigua* live in dendritic tubes, its main food source being diatoms/algal detritus. (Fjellberg, 1972; Boubee, 1983).

This species has been recorded during cold episodes or glacial periods, at sites in England (Bedford et al., 2004), Norway (Velle et al., 2005), Poland (Płóciennik et al., 2015), and the Baltic region (Hofmann and Winn, 2000). However, *Corynocera*

261 *ambigua*, cannot be considered merely a cold species. Some authors believe that its occurrence depends on high oxygen content
 262 in the water (Brodersen and Lindegaard, 1999a) and for other authors, it is a pioneer species that appears first after glacial
 263 retreat, similarly to *Chironomus anthracinus*-type (e.g. Heiri and Millet, 2005; Ilyashuk et al., 2005, 2013, 2022; Gandouin et
 264 al., 2016). Luoto and Sarmaja-Korjonen (2011) suggest this is how the species adapts to existing climatic conditions. The
 265 locally observed decline in *Corynocera ambigua* numbers in the Krępa sediments could also be attributed to changes in lake
 266 productivity related to changes in the environment. For example, when the production of soil and trees increased, the number
 267 of this species has been found to decrease (Magny et al., 2006; Larocque-Tobler et al., 2009).
 268 *Chironomus anthracinus*-type occurs in various lake zones and is capable of surviving approximately 2–4 months of oxygen
 269 deficiency in water (Hamburger et al., 1994). It is a species which easily occupies niches that are inaccessible to others.
 270 According to some authors, it is a eutrophic (Kansanen, 1985; Brodersen and Lindegaard, 1999b) or cold-adapted species
 271 (Rohrig et al., 2004; Brooks et al., 2007; Płóciennik et al., 2011) and prefers soft, more organic sediments (McGarrigle, 1980).
 272 Therefore, the appearance of *Chironomus anthracinus*-type and *Glyptotendipes pallens*-type in the Krępa sediment may
 273 indicate the onset of eutrophication. Both *Chironomus anthracinus*-type and *Corynocera ambigua* are found in stratified lakes
 274 (e.g., Saether, 1979; Heiri, 2004). As we observe in our record, both species are relatively resistant to unfavourable
 275 environmental conditions, so possess a wide range of conditions in which they can occur.
 276 Lower part of the sediment sequence (2180 cm-1160 cm) is almost completely devoid of Chironomid remains, except few
 277 badly preserved *Chironomus anthracinus*, *Chironomus plumosus* and *Glyptotendipes pallens* head capsules at 2000 cm, 1680
 278 cm, and 1205-1190 cm depths. Head capsules are recorded again at 1155 cm-1122.5 cm depths – mostly *Corynocera ambigua*,
 279 *Chironomus anthracinus*, *Chironomus plumosus* and *Glyptotendipes pallens* (Fig. 2). 1072.5-1122.5 cm part predominantly
 280 contains cold-adapted species such as *Corynocera ambigua* and freeze-resistant species such as *Glyptotendipes pallens*-type
 281 and *Glyptotendipes severini*-type, which are often associated with algae and diatoms or mine leaves (Tarkowska-Kukuryk,
 282 2014). 1022.5-1072.5 cm depth range is characterised by species highly resistant to difficult environmental conditions, such
 283 as *Chironomus anthracinus*-type, *Corynocera ambigua* and *Glyptotendipes pallens*-type. From 1022.5 cm to 967.5 cm depth
 284 there are several species observed. . This is the part most abundant in Chironomidae head capsules, with over 40 individuals
 285 per sample on average and maximum 78 head capsules at 985 cm sample. Species composition during this part is dominated
 286 by *Corynocera ambigua*, *Chironomus anthracinus*-type, *Chironomus-plumosus*-type and *Prosilocerus lacustris*-type.
 287 Additionally, some *Tanytarsus glabrecens*-type head capsules appear – this species was almost unseen in remaining sections.
 288 Between 967.5 cm and 877.5 cm depth, the number of chironomid head capsules started to decline above 965 cm depth, with
 289 only single unidentified Chironomidae head capsules at 955 cm and 950 cm. . . In subsequent section (877.5-765 cm) the
 290 number of Chironomidae is very low – only 2 *Chironomus plumosus*-type individuals were identified. Even *Corynocera*
 291 *ambigua*, abundant in previous sections, disappears.
 292

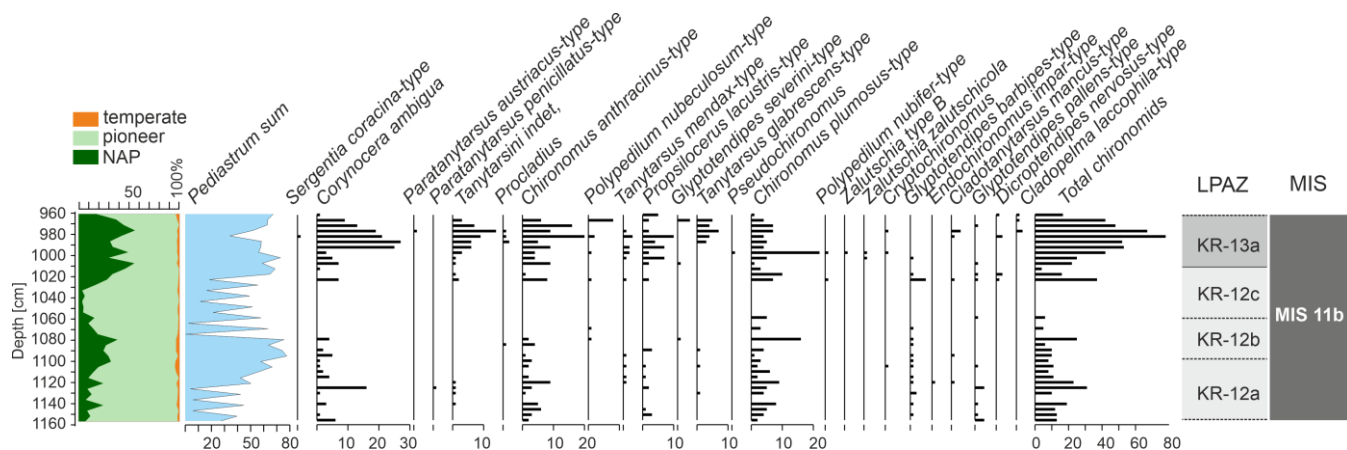


Figure 2: Stratigraphic diagram of the Chironomidae assemblages. Caption: Chironomidae species are presented as counted numbers of specimens.

3.2 July air temperature reconstruction based on Chironomidae assemblages from the Krępa site

Due to the low number of chironomid head capsules preserved in the Krępa sediments, a chironomid-based July temperature reconstruction was only possible for the uppermost part of the sediment core, encompassing the post-Holsteinian stadial that is most likely equivalent to MIS 11b. LPAZ KR-12a marks the onset of MIS 11b that directly follows the Holsteinian Interglacial. In this period, average July temperatures still ranged between 17 and 19 °C before rapidly dropping to about 16 °C and increasing again to 18-20 °C in LPAZ KR-12b. July temperatures remained at this level throughout LPAZ KR-12c, before significantly dropping to 15-17 °C in the middle of LPAZ KR-13a. Only at the end of LPAZ KR-13a, which is equivalent to the transition to the following interstadial that most likely corresponds to MIS 11a, July temperatures markedly increased again to about 20 °C.

Table 1: Temperature reconstruction from Chironomidae preserved in the Krępa sediments with reconstructed mean July air temperature (T_{jul-Ch}), error of the estimated T_{jul-Ch} , minimum dissimilarity between the chironomid assemblage in the Krępa sediments training set samples (MinDC), principal component analysis values (PCA) and corresponding LPAZ

Core depth (cm)	T_{jul-Ch}	error of est. (T_{jul-Ch})	MinDC	PCA	Number of Chironomidae head capsules	LPAZ
969	20.10	1.60	9.82830	-1.8144135	51	KR-13a
975	15.26	1.64	6.08105	-1.2383560	48	KR-12c
980	16.82	1.57	7.89471	-1.7518844	67	
985	17.23	1.59	8.37351	-1.4636110	78	KR-12b
990	15.93	1.70	7.35685	-1.9244674	52	
995	15.84	1.72	6.77137	-0.6709448	53	KR-12a
1000	18.77	1.52	8.27430	6.5934818	42	
1011	18.09	1.63	7.90763	0.4039345	51	
1022	19.25	1.50	7.06444	0.4114688	53	
1080	20.20	1.53	8.02666	0.5281182	52	
1102	18.55	1.52	8.95789	1.3132870	48	
1125	17.69	1.52	8.63666	-0.2629876	64	
1148	18.97	1.56	6.86405	-0.1236256	57	

According to the SNP training set-based reconstruction, 10 samples with good modern analogues remain below the 5 % percentile threshold (minDC), while 3 samples with average modern analogues have values above the 5 % percentile threshold ($6.08105 < \text{minDC} < 9.82830$). PCA values range between ~ -1.92 and 6.59 (Tab. 1).

3.3 Vegetation changes during the Early Saalian Glaciation at Krępa site

Initially, 14 Local Pollen Assemblages Zones (LPAZ) covering the end of MIS 12 and MIS 11 period were extracted. Post-holsteinian (MIS 11b) covers LPAZ from 12a to 13a.

LPAZ KR-12a (1122.5-1187.5 cm) - At the beginning of the zone, the development of *Pinus* forests with an admixture of *Picea* (up to 6 %) is observed. Low NAP percentages suggest a very dense vegetation. However, percentages of *Pinus* and other tree species gradually decrease, and open herbaceous communities appear. The end of the zone is associated with a decrease in the percentage of *Pinus* pollen. **Low number of Chironomidae head capsules (approximately 15 per sample). Dominance of *Chironomus anthracinus*-type (25 %) and *Corynocera ambigua* (16 %).**

LPAZ KR-12b (1072.5-1122.5 cm) - A further decrease in *Pinus* pollen is observed. At the end of the zone, the landscape was likely already dominated by open communities (NAP up to 40 %) and sparse *Pinus* forests. Dominance of *Corynocera ambigua* (24 %) and high contents of *Chironomus anthracinus*-type. **Disappearance of *Glyptotendipes pallens*-type and appearance of *Glyptotendipes severini*-type.**

LPAZ KR-12c (1022.5-1072.5 cm) - Initially, dense *Betula* forests with *Larix* as an admixture dominated the landscape. Subsequently, a rapid development of *Pinus* forests is observed. The end of the zone is associated with a sudden drop in the

percentage of *Pinus* pollen. **The number of Chironomidae declines. Dominant species are *Chironomus anthracinus*-type (17 %), *Corynocera ambigua* and *Glyptotendipes pallens*-type (13 %).**

LPZ KR-13a (967.5-1022.5 cm) - Initially, there was a significant opening in the vegetation, and herbaceous plants and shrubs dominated the landscape. In the middle of this zone, there was a temporary return of very sparse *Pinus* and *Betula* forests, followed by another expansion of herbaceous vegetation. The end of the zone is associated with an increase in *Betula* pollen. **Significant increase in the number of Chironomidae (on average 45 individuals per sample). Dominant species are *Corynocera ambigua* (approx. 29 %) and *Chironomus anthracinus*-type (18 %).**

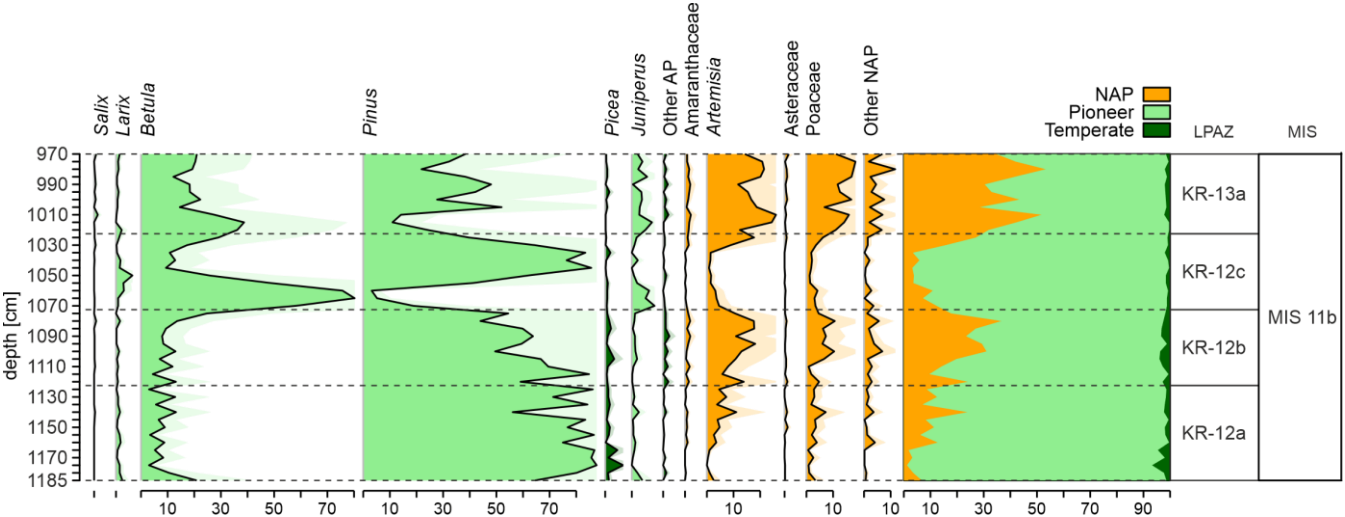


Figure 3: Simplified percentage pollen diagram from the Krępa 2015 sediment core on depth scale (cm) with zonation of the diagram.

3.4 Pollen-based climate reconstructions from the Krępa site

Pollen-based climate reconstructions from the Krępa sediment core reveal distinct climate variability throughout MIS 11b, reflecting stadial–interstadial transitions (Fig. 4). The two pollen-based methods show broadly similar trends across all zones, with MAT generally producing higher summer temperature values than WA-PLS except in KR-12c. Where chironomid data are available, pollen-based MTWA reconstructions reproduce similar patterns, with differences falling within their respective uncertainty ranges. Among the two pollen-based models, MAT generally corresponds better to the chironomid WA-PLS reconstructions, showing overall closer alignment in reconstructed summer temperatures.

WA-PLS reconstructions were somewhat less robust, especially for precipitation, while the TANN and MTWA estimates still showed moderate predictive ability (Tab. 2). Reconstructed MTWA from both pollen-based methods generally ranged between approximately 15°C and 19°C. The two pollen-based methods show similar trends across all zones, with MAT often producing slightly higher summer temperature values than WA-PLS.

During LPAZ KR-12a, MAT- and WA-PLS-derived MTWA averaged approximately 16.8°C, close to the chironomid-inferred mean of 18.3°C. In LPAZ KR-12b, both pollen methods indicate further warming (~18.7°C MAT, ~16.4°C WA-PLS), consistent with the chironomid estimate of 19.4°C, reflecting peak interstadial conditions. In LPAZ KR-12c, MTWA values dropped to ~15.1°C (MAT) and ~15.7°C (WA-PLS), indicating cooling during this interval. A moderate rebound is evident in LPAZ KR-13a, with MTWA increasing again to ~17.3°C (MAT) and ~15.3°C (WA-PLS), while the mean chironomid MTWA is 17.5°C.

TANN values generally followed the summer temperature trends, beginning with relatively warm conditions in LPAZ KR-12a (~3°C). A slight increase was observed in LPAZ KR-12b (~3.1°C), followed by cooling in LPAZ KR-12c (~1.2°C). In LPAZ KR-13a, a modest recovery occurred with TANN rising to around 1.57°C.

MTCO showed greater variability. Winters in LPAZ KR-12a and KR-12b were comparably cold, with MTCO values around -9.6°C and -11.7°C, respectively. LPAZ KR-12c showed slightly less severe winters (~-10.72°C). A more pronounced cooling occurred in LPAZ KR-13a, where MTCO reached around -13.2°C.

PANN reconstructions showed some uncertainty but generally ranged between 500 and 900 mm. LPAZ KR-12a was characterized by relatively high precipitation (~640 mm), followed by moderately high values in LPAZ KR-12b (~510 mm). A moderate increase occurred in LPAZ KR-12c (~580 mm). In LPAZ KR-13a, PANN remained lower, typically around 520 mm, suggesting continued reduction in annual precipitation.

lake, the water pH, and microhabitats. Furthermore, training sets are also available to reconstruct the historic water level, salinity or oxygen content of the studied water body (Lotter et al., 1997).

4.1.1 Possible difficulties in temperature reconstruction based on Chironomidae analysis during past interglacials

The basic principle of palaeoecological reconstructions is uniformitarianism, implying that processes taking place on Earth in the past were the same as today (Krzeminski and Jarzembowski, 1999). This, for example, allows temperature to be reconstructed based on fossil Chironomidae assemblages by assuming that a given species has the same habitat requirements as thousands or hundreds of thousands years ago. The oldest recorded chironomid remains date back to the Late Triassic, i.e. ~200 Ma BP (Krzeminski and Jarzembowski, 1999). Data from the MIS 11 Krępa sediments indicate a large difference in the number and state of preservation of chironomid remains compared to Holocene sites. Usually, at least 50 individuals per sample are required for robust reconstructions of the average July temperature, as smaller numbers of identified head capsules considerably increase the error range of the air temperature reconstruction. It is therefore commonly recommended to combine adjacent samples in case of low head capsule amounts (Heiri and Lotter, 2001). To enable selection of sites that could potentially yield chironomid-based palaeoenvironmental reconstructions, it is critical to analyse the factors that could limit the degree of preservation in chironomid remains, or cause a marked decrease/complete disappearance in the number of individuals.

Chironomidae inhabit all moist or aquatic habitats from moist wood to the ocean between the tropics and the Arctic. The high specialisation of individual species is thereby decisive for their common occurrence and their ability to survive under difficult environmental conditions. Among the features that allow specimens to succeed are: a short life cycle (in some cases only 8 days) (Reyes-Maldonado et al., 2021), osmoregulation, which enables survival in high-salinity waters (Kokkinn, 1986), or parthenogenesis, which implies a high efficiency of population reproduction, faster colonisation rate and high fertility (Lencioni, 2004; Nondula et al., 2004; Donato and Paggi, 2008; Orel and Semenchenko, 2019; Lackmann et al., 2020), as well as a short DNA chain (Gusev et al., 2010; Cornette et al., 2015). Some species are able to change food resources depending on the availability in their habitat (Tokeshi, 1995; Davis et al., 2003). Large lakes, such as the one that probably existed at Krępa (1) have a greater variety of habitats, thus being characterised by a larger biodiversity of Chironomidae (Allen et al., 1999; Heino, 2000; Tarr et al., 2005), and (2) are more resilient to extreme droughts and other extreme events. In contrast, small lakes with less diverse, isolated habitats exhibit reduced species diversity and dispersal (Roberts, 2003).

Despite the specialisation of chironomids, there are many conditions that limit the number of communities. One of the main factors limiting and determining the life processes of Chironomidae is temperature as each life stage is highly dependent on this factor. The development of eggs, larvae and pupae, nutrition and growth, the emergence of individuals and the ability to fly are all constrained by temperature maxima and minima, beyond which the given processes can no longer take place. Most groups can tolerate low sub-zero temperatures; the temperature below which the development of most species does not occur is -15°C (Walker and Mathewes, 1989; Plóciennik, 2005). At Krępa, however, our July temperature reconstruction indicates temperatures well above that threshold, so even in case of severe winters, Chironomids should have been able to develop

during the warmer periods of the year. Frost tolerance is highest in the Orthocladinae family and lowest in the Tanypodinae family (Danks, 1971). In the case of the Krępa sediments, species of both families were found (e.g. *Prosilocerus lacustris*-type and *Procladius* respectively) with Orthocladinae being more abundant than Tanypodinae (57 vs. 5 head capsules) with the highest number of head capsules being preserved during a period with relatively cool summers (15-17°C).

Another important factor causing the decline of Chironomidae populations is the lack of oxygen in the water, although this cannot be directly captured by palaeoreconstructions. Instead, low-oxygen conditions are generally only indicated by an abundant occurrence of organic matter in the sediment. Such increases in organic matter commonly increase bacterial respiration and result in oxygen deficiency in the profundal of water bodies (Charlton, 1980; Matzinger et al., 2010; Müller et al., 2012). Another factor limiting the preservation of chironomid head capsules in sediments are mechanical factors that cause damage to the head capsules. For example, Tanypodinae remains are, due to their large size, not very resistant to disintegration and the number of preserved capsules may therefore be smaller (Walker et al., 1984). This would be consistent with our finding of only 5 Tanypodinae individuals in the Krępa sediments across four different depths. The preservation of remains only from the 3rd and 4th larval stages is most likely related to the increased amount of chitin in these developmental stages, making remains of these stages more resistant to disintegration. The remains of Chironomidae may also not be preserved if accumulation rate is low and remains of species from shore habitats could be poorly preserved. However, studies confirm a positive relationship between biocenosis and thanatocoenosis (Iovino, 1975; Walker et al., 1984). The number of generations per year may also affect the abundance of Chironomidae, i.e. subfossils of multivoltine species can be overrepresented compared to bivoltine species, however, it is difficult to determine whether changes in species composition correspond with voltinism (Tokeshi, 1995).

The main factor influencing the preservation of Chironomidae remains is the content of CaCO₃, especially in moderately and strongly acidified lakes. This factor is often more important than pH, depth or time since the deposition of remains (Bailey et al., 2005). The microenvironment and the presence of organic matter are of great importance for the preservation of remains (Briggs and Kear, 1993; Sageman and Hollander, 1999). The faster mineralisation occurs, the better the preservation of any remains (Briggs and Kear, 1993; Park, 1995). Further factors reducing the abundance of chironomids are extreme temperatures, low nutrient levels, acidic waters, high Se concentrations (Del Wayne et al., 2018; Mousavi, 2002), the content of hydrogen sulphide during holomixis, as well as paludification of the lake (Takagi et al., 2005; Płóciennik et al., 2020).

The lack of oxygen in the sediment could have limited not only the number of Chironomidae but also the number of preserved head capsules in the sediment. In particular, chitin does not usually accumulate in anaerobic sediment, because it is more easily broken down by bacteria, effectively mineralising it into CH₄ and CO₂ (Wörner and Pester, 2019).

Chironomid species found in the Krępa sediments have a wide range of environmental conditions in which they occur. In particular, we observe dominance of species resilient to harsh conditions, such as the oxygen-deficiency-resistant *Chironomus anthracinus*-type, the eutrophic *Chironomus plumosus*-type (18.7 and 22.2 % of the total number of head capsules, respectively), as well as the cold-adapted *Corynocera ambigua* (25.7%) and the freeze-resistant *Prosilocerus lacustris*-type (7.5 %).

459 *Corynocera ambigua* is a species often described as cold-adapted oligotrophic (Fjellberg, 1972; Pinder and Reiss, 1983;
 460 Walker and Mathewes, 1988; Brooks et al., 2007; Luoto et al., 2008; van Asch et al., 2012), inhabiting shallow lakes in arctic
 461 and subarctic regions, though it is also found in eutrophic lakes (Halkiewicz, 2008; Kotrys et al., 2020) Adults of this species
 462 are not able to fly, and breed on the water surface when the temperature reaches approximately 7-8 °C (optimum 13.7 °C).
 463 Mothes (1968) concluded that *Corynocera ambigua* larvae develop in autumn and winter, but not during summer. The decline
 464 in their numbers may be due to growth of filamentous algae in summer. Larvae of *Corynocera ambigua* are eurythermic, while
 465 the pupae are cold-stenothermic (Brundin, 1949). They only reproduce at low temperatures and inhabit water bodies with a
 466 maximum depth of approximately 25 m. The abundance of *Corynocera ambigua* has been shown to be correlated with
 467 charophyte contents (Brodersen and Lindegaard, 1999b). Although this species does not feed on charophytes, their presence
 468 may increase the number of diatoms and stabilise the trophic status and water clarity (Forsberg, 1965; Blindow, 1992).
 469 *Corynocera ambigua* live in dendritic tubes, its main food source being diatoms/algal detritus. (Fjellberg, 1972; Boubee, 1983).
 470 This species has been recorded during cold episodes or glacial periods, at sites in England (Bedford et al., 2004), Norway
 471 (Velle et al., 2005), Poland (Płóciennik et al., 2015), and the Baltic region (Hofmann and Winn, 2000). However, *Corynocera*
 472 *ambigua*, cannot be considered merely a cold species. Some authors believe that its occurrence depends on high oxygen content
 473 in the water (Brodersen and Lindegaard, 1999a) and for other authors, it is a pioneer species that appears first after glacial
 474 retreat, similarly to *Chironomus anthracinus*-type (e.g. Heiri and Millet, 2005; Ilyashuk et al., 2005, 2013, 2022; Gandouin et
 475 al., 2016). Luoto and Sarmaja-Korjonen (2011) suggest this is how the species adapts to existing climatic conditions. The
 476 locally observed decline in *Corynocera ambigua* numbers in the Krępa sediments could also be attributed to changes in lake
 477 productivity related to changes in the environment. For example, when the production of soil and trees increased, the number
 478 of this species has been found to decrease (Magny et al., 2006; Larocque-Tobler et al., 2009).
 479 *Chironomus anthracinus*-type occurs in various lake zones and is capable of surviving approximately 2–4 months of oxygen
 480 deficiency in water (Hamburger et al., 1994). It is a species which easily occupies niches that are inaccessible to others.
 481 According to some authors, it is a eutrophic (Kansanen, 1985; Brodersen and Lindegaard, 1999b) or cold-adapted species
 482 (Rohrig et al., 2004; Brooks et al., 2007; Płóciennik et al., 2011) and prefers soft, more organic sediments (McGarrigle, 1980).
 483 Therefore, the appearance of *Chironomus anthracinus*-type and *Glyptotendipes pallens*-type in the Krępa sediment may
 484 indicate the onset of eutrophication. Both *Chironomus anthracinus*-type and *Corynocera ambigua* are found in stratified lakes
 485 (e.g., Saether, 1979; Heiri, 2004). As we observe in our record, both species are relatively resistant to unfavourable
 486 environmental conditions, so possess a wide range of conditions in which they can occur.
 487 *Chironomus plumosus*-type, also quite abundant in the sediment sequence, occurs in a wide range of habitats and is particularly
 488 resistant to anoxia (Saether, 1979; Brooks et al., 2007). Moreover, along with *Dicrotendipes nervosus*-type, this species is an
 489 indicator of progressive eutrophication (Brodersen and Lindegaard, 1999b)
 490 Both eutrophic and oligotrophic species, as well as warm- and cold-adapted species, occur in the Krępa sediments.
 491 The origin of the sedimentary basin at Krępa is difficult to interpret. Most sites with deposits from the Holsteinian Interglacial
 492 in this region of Poland are associated with tunnel valleys that formed during the Elsterian glaciation (Żarski et al., 2005;

493 Nitychoruk et al., 2006). However, these sites are usually located beyond the maximum extent of the Older Saalian glaciation
494 (Drenthe Stage in Germany; Odra glaciation in Poland; MIS 6) and thus, are subtly visible in the present surface morphology.
495 In the case of Krępa, these deposits have been covered by the Older Saalian glacial advance, resulting in the complete
496 transformation of the post-Elsterian landscape. Based on the geological cross section presented in the DGMP sheet 676 - Kock
497 (Drozd and Trzepla, 2007) and the distribution of interglacial deposits in the study area (Jesionkiewicz, 1982), it can be inferred
498 that the depression hosting the Krępa palaeolake was a relatively extensive kettle hole, formed during the recession of the
499 Elsterian ice sheet.

500 As there are obviously only very few habitats where no invertebrates occur, the absence of chironomid remains during most
501 of the Holsteinian Interglacial could be best explained by sediment-related disintegration and/or anoxic conditions at the
502 bottom of a relatively deep lake. Another reason for the lack of remains could be the mineralisation of chitin. This would be
503 in agreement with the parallel observed lack of cellulose remains from plants as well as with the very low number of
504 Tanypodinae head capsules. However, satisfyingly explaining the lack of chironomid remains in most of the interglacial lake
505 deposits requires further research and an in-depth comparison of our results with other lake sediments that lack chitinous
506 remains.

507 **4.1.2 Chironomid-inferred temperature reconstruction from the Krępa site in relation to pollen-based climate** 508 **reconstructions**

509 A chironomid-based July temperature reconstruction was only possible for the part of the Krępa sediment core that corresponds
510 to LPAZ KR-12 and early LPAZ KR-13, which most likely corresponds to MIS 11b. Chironomid-based July temperatures
511 during the early part of this interval (LPAZ KR-12a and LPAZ KR-12b) were probably still relatively high and stable, ranging
512 from 19 to 21 °C, but dropping rapidly in LPAZ KR-12c and LPAZ KR-13a to 15-17 °C. The following increase to ~20 °C at
513 the top of LPAZ KR-13a possibly reflects the transition into the post-Holsteinian interstadial that corresponds to MIS 11a.
514 This data indicates the July temperature maximum during the post-Holsteinian is consistent with the temperature range of the
515 SNP training set (3.5-20.0 °C) (Kotrys et al., 2020). Comparing MIS 11 to the Holocene, it is crucial to mention that insolation
516 patterns for both periods differ - MIS 11 was characterised by two insolation maxima, whilst there was only one (though more
517 distinct) during the Holocene (Rohling et al., 2010). In fact, summer temperature increase during MIS 11b might be explained
518 by increasing insolation.

519 In general, most Chironomidae remains in the Krępa sediments are found during cool periods, but are absent during warm
520 periods. In contrast, Chironomidae were most abundant in LPAZ KR-12, which roughly corresponds to MIS 11b, the first cold
521 phase after the Holsteinian Interglacial (Imbrie et al., 1984; Fawcett et al., 2011). To date, studies using subfossil Chironomidae
522 to reconstruct past climate conditions mainly focused on the Weichselian Late Glacial and the Holocene (Gandouin et al.,
523 2016; Nazarova et al., 2018; Druzhinina et al., 2020). As a result, there are very few chironomid-based July temperature
524 reconstructions for the Late and Middle Pleistocene older than 20 ka BP available (Gandouin et al., 2007; Samartin et al., 2016;
525 Pliikk et al., 2019; Ilyashuk et al., 2020; Bolland et al., 2021; Lapellegerie et al., 2024; Rigterink et al., 2024), and no studies

for the MIS 11 complex. In general, chironomid records from other sites and time intervals are characterised by a higher abundance and species diversity of Chironomidae, whilst at Krępa, Chironomidae occur only during the early glacial period following the Holsteinian Interglacial. A similar phenomenon has so far only been observed in the Laptev Sea region (Arctic Siberia), where Chironomidae also appear only in the cold period after the Eemian Interglacial, when the site was surrounded by wet grass-sedge shrub tundra period (Andreev et al., 2004). Assemblages from this site consist mostly of unidentified Tanytarsini individuals, eutrophic *Chironomus plumosus* and semi-aquatic taxa such as *Limnophyes/Paralimnophyes*, *Smittia* and *Paraphaenocladus*. The three species from the latter group were not identified at Krępa as opposed to *Chironomus plumosus* and Tanytarsini. Contrary to the patchy occurrence of chironomids, pollen-based climate reconstructions using MAT and WA-PLS provide continuous and robust records that have been successfully applied across various European regions and time periods (Mauri et al., 2015; Chevalier et al., 2020). During LPAZ KR-12a, pollen reconstructions indicate relatively stable and moderate summer temperatures. Additionally, PANN remains relatively high during this phase, suggesting consistently moist conditions supporting dense forest coverage. This is in agreement with the observed dominance of *Pinus* forests with some admixed *Picea* during this phase, reflecting more humid but not necessarily warmer conditions (Caudullo et al., 2016). The significant NAP increase LPAZ KR-12b suggests substantial forest decline, although the pollen-based MTWA reconstructions indicate relatively warm summers. This combination of ecological and climatic signals strongly suggests that the decline in forest cover was primarily driven by colder winter temperatures rather than summer thermal conditions. The pollen-based MTWA reconstruction confirms peak interstadial warmth in terms of summer temperatures are comparable to current mean July temperatures in Eastern Poland (Mauri et al., 2015; Kotrys et al., 2020; Gedminienė et al., 2025). Furthermore, the pollen-based TANN reconstruction also highlights peak interstadial warmth during LPAZ KR-12b, indicating overall favourable climatic conditions during the growing season. The pronounced increase in open-ground vegetation (NAP dominance) and herbaceous taxa thus likely reflects an ecological response to severe winter conditions, that restricted the establishment and survival of forest taxa, particularly those sensitive to extreme winter frosts (Körner and Paulsen, 2004; Harrington and Gould, 2015).

LPAZ KR-12c begins with pioneer *Betula-Larix* forests, reflecting a significant climatic shift towards colder and possibly drier conditions. The appearance of *Larix*, a cold-tolerant, light-demanding taxon adapted to short growing seasons and low temperatures, reinforces the interpretation of subarctic or boreal-like climate conditions. *Larix* is typically associated with northern coniferous forests, and reaches its distributional limits in areas with low winter temperatures and moderate precipitation (San-Miguel-Ayán et al., 2022). Gradually increasing pollen signals from *Pinus* indicate a modest rise in thermal conditions later within LPAZ KR-12c, but within a generally cool and moisture-limited climatic regime. The absence of chironomids during this interval corroborates the interpretation of sustained cooler and drier conditions. Chironomid assemblages are sensitive to environmental harshness, and under extremely cold or oligotrophic conditions, their production may be so low that remains are not preserved in sediment records (Eggermont and Heiri, 2012).

The gradual cooling indicated by our chironomid-based reconstruction during LPAZ KR-13a is consistent with the presence of sparse *Betula* forests at the onset of this zone. Pollen-based reconstructions suggest that MTWA remained relatively mild

(~17.3 °C MAT, ~15.3 °C WA-PLS), closely aligning with the chironomid-inferred mean $T_{\text{Jul-Ch}}$ value of ~17.5°C for this interval. Although the chironomid data exhibits a broader range (15–20°C), this variability falls within typical reconstruction uncertainties and does not suggest a fundamentally different climatic signal. Meanwhile, declines in T_{ann} and especially MTCO indicate cold-season severity remained the primary constraint on forest development (Nienstaedt, 1967; Körner and Paulsen, 2004; Harrington and Gould, 2015). In parallel, a reduction in P_{ann} further supports increasing climatic stress, potentially limiting moisture availability and forest resilience during this transitional phase (Körner and Paulsen, 2004).

The broader relevance of the climatic conditions reconstructed from Krępa pollen data is given by the comparison with other MIS 11 palaeotemperature reconstructions from Southern Europe (Fig. 5) (Rodrigues et al., 2011; Oliveira et al., 2016; Kousis et al., 2018; Ardenghi et al., 2019; Sassoon et al., 2023, 2025). These Mediterranean records indicate generally warm conditions during MIS 11b, punctuated by recurrent cooling and drying events. For instance, the Lake Ohrid from SE Europe record shows a transition from temperate deciduous to cold mixed forests, with T_{ann} dropping to ~2 °C and mean temperature of the coldest month below –8 °C during the coldest events, despite precipitation often remaining near or above 800–900 mm (Kousis et al., 2018). Meanwhile, records from the SW Mediterranean reflect similar climate oscillations, with Sassoon et al. (2025) documenting synchronous declines in T_{ann} and P_{ann} centered ~398 ka BP. Krępa record reflects relatively steady summer cooling alongside more marked declines in winter temperatures and moderately decreasing precipitation. Mediterranean vegetation is primarily water-limited, making it especially vulnerable to fluctuations in atmospheric moisture and reductions in winter rainfall (Giorgi, 2006; Lionello et al., 2006). Vegetation in Eastern Europe, however, is highly responsive to winter climate extremes. In particular, cold-season frost events, snow cover variability, and late-winter cold snaps affect plant performance, especially in temperate and continental zones (Kreyling, 2010; Camarero et al., 2022).

The lack of accurate absolute dating for terrestrial sediment sequences from the Holsteinian Interglacial makes it difficult to directly compare the results from Krępa to other MIS 11 sites. However, as there are a few quantitative temperature reconstructions based on pollen and biomarkers from other sites in Europe for the post-Holsteinian, a general comparison of temperature levels during this interval is feasible. For example, Tenaghi Philippon record indicates mild summer temperature drop to ~16 °C at the coolest period of MIS 11b (Ardenghi et al., 2019). Climatic fluctuations at another Mediterranean region site – ODP 986 at Alboran Sea – were not abrupt, especially during first half of MIS 11b. Initially summer temperature stayed above 20 °C, only at further stage decreasing to ~17 °C (Fig. 5) (Sassoon et al., 2025). Pollen analyses on marine sediments from the Iberian margin show similar climatic and ecological patterns for MIS 11b as observed at Krępa, namely repeated forest decline events. These were paralleled by reductions in sea surface temperature, although temperatures were still relatively high during most of MIS 11b – only about 1 °C below MIS 11c levels (Rodrigues et al., 2011; Oliveira et al., 2016). A similar pattern between still relatively high air temperature during early MIS 11b, and a temperature drop only during late MIS 11b is also seen in palynological data from Lake Ohrid in SE Europe (Kousis et al., 2018).

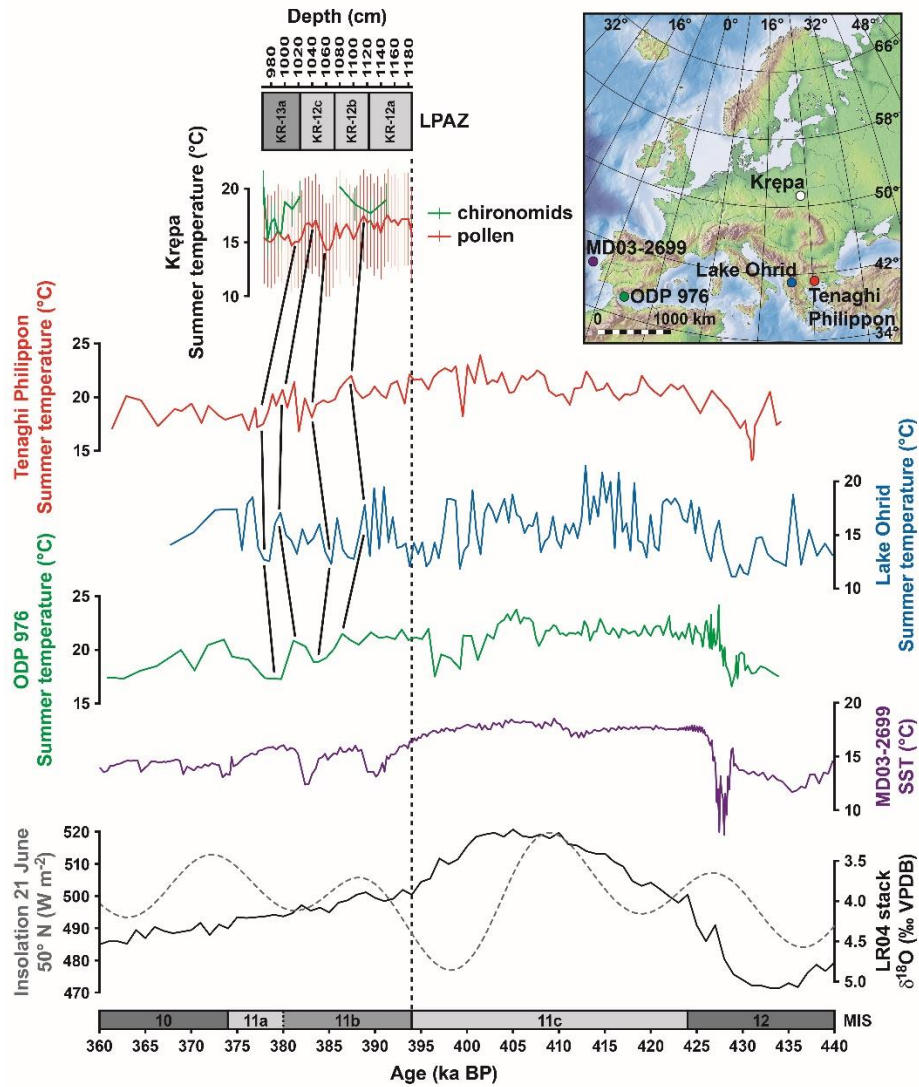


Figure 5: Comparison of (top to bottom) the Marine Isotope Stage (MIS) 11b pollen- and chironomid-based summer temperature reconstructions from Krępa, a summer temperature reconstruction based on branched glycerol dialkyl glycerol tetraethers (brGDGTs) from Tenaghi Philippon, Greece (Ardenghi et al., 2019), a pollen-based summer temperature reconstruction from Lake Ohrid, Balkan Peninsula (Kousis et al., 2018; Kountsodendris et al., 2020), a pollen-based summer temperature reconstruction from ODP Site 976, Alboran Sea (Sassoon et al., 2025), and a biomarker-based (Uk'37) sea surface temperature (SST) reconstruction from marine core MD03-2699, Iberian margin (Rodrigues et al., 2011). The LR04 $\delta^{18}O$ stack (solid black line; Lisiecki and Raymo, 2005) and the 21 June insolation at 50° N (approximate latitude of Krępa; dashed grey line; Laskar et al., 2004) are provided as a palaeoclimatic context. The timing of the MIS boundaries 12/11c and 11a/10 is given according to Lisiecki and Raymo (2005); the timing of the MIS boundaries 11c/11b and 11b/11a is tentative. The insert map shows the locations of the individual proxy records.

In line with our chironomid-based July temperature reconstruction from Krępa, these results show that the temperature decline at the demise of the Holsteinian Interglacial was not abrupt, and at least summer temperatures likely remained at a relatively high level for several thousand years. The general summer temperature variability that is seen in the Krępa record throughout the post-Holsteinian, i.e. the initial drop during the early MIS 11b, the following increase and the more pronounced decrease

during late MIS 11b, as well as the marked increase at transition into MIS 11a, closely resembles vegetation and sea surface temperature variability at the Iberian margin, and may indicate a substantial impact of insolation variability (Rodrigues et al., 2011; Oliveira et al., 2016).

Conclusion

This study presents the first combined chironomid- and pollen-based palaeoclimatic reconstruction for the post-Holsteinian i.e. MIS 11b, offering a new perspective on climate variability in Eastern Europe during this period. The results highlight the complementarity and reliability of both proxies, as pollen-based MAT and WA-PLS reconstructions show strong internal consistency and correspond well with chironomid-inferred summer temperatures where data is available. The summer temperatures range from 15 to 19 °C and between 15 and 20 °C for the pollen- and chironomid-based reconstruction respectively. This indicates colder summers compared to present times for most of the post-Holsteinian period. The pollen-based MAT reconstructions exhibit particularly high predictive skill, especially for temperature variables. The analysed part of the Krępa sediment record reveals a progressive shift towards a more continental climate throughout MIS 11b. This is reflected by gradually cooling summers, increasingly severe winters, and a decline in annual precipitation. These climatic trends coincide with marked vegetation changes, including forest retreat and a rise in herbaceous taxa during colder phases. To date, the vast majority of studies addressing terrestrial palaeoclimate variability during the Middle Pleistocene relies on pollen analysis. However, this does not imply a complete lack or low abundance of Chironomid-inferred reconstruction in sites other than Holocene. Moreover, they may prove to be a priceless source of knowledge on temperature, considering potential differences between pollen and Chironomid-inferred records. By comparing the results from different sites, it will be possible to identify the factor(s) that influenced the preservation of Chironomidae subfossil remains. Ultimately, this study underscores the value of multi-proxy approaches in palaeoclimate reconstruction, particularly for pre-Holocene periods. Chironomids show significant potential as a summer temperature proxy in older sediments, as long as preservation conditions are favourable.

Financial support

The research was funded by the National Science Centre project, "Novel multi-proxy approaches for synchronisation of European palaeoclimate records from the Holstein interglacial". Project no. 2019/34/E/ST10/00275.

Authors contribution

TP proposed the idea of the main text, and contributed to the figures. TP,AGr and AG wrote the original draft version of the manuscript. BK performed the chironomid-inferred summer temperature reconstruction. AG performed the pollen-inferred climate reconstructions. MŻ collected and described the core in the field. AG, AH, and MS analysed the pollen data. TP, AGr and MS analysed the chironomid data. TP, AGr, AG,, SL, MB and MS did the visualisations (graphs and maps). AG, AH, MŻ,

636 MB, JN, MC, SL and MS reviewed the paper. All authors have made substantial contributions to the submission of this
637 manuscript.

638 **Competing interests**

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

640 **References**

- 641 Allen, A. P., Whittier, T. R., Kaufmann, P. R., Larsen, D. P., O'Connor, R. J., Hughes, R. M., Stemberger, R. S., Dixit, S. S.,
642 Brinkhurst, R. O., Herlihy, A. T., and Paulsen, S. G.: Concordance of taxonomic richness patterns across multiple assemblages
643 in lakes of the northeastern United States, *Can. J. Fish. Aquat. Sci.*, 56, 739–747, 1999.
- 644 Andersen, T., Sæther, O., Cranston, P., and Epler, J.: 9. The larvae of Orthocladiinae (Diptera: Chironomidae) of the Holarctic
645 Region – Keys and diagnoses. In: Andersen, T. Cranston, P.S. & Epler J.H. (Sci. eds): The larvae of Chironomidae (Diptera)
646 of the Holarctic region – Keys and diagnoses., *Insect Syst. Evol. Suppl.*, 66, 189–386, 2013.
- 647 Andreev, A. A., Grosse, G., Schirmmeister, L., Kuzmina, S. A., Novenko, E. Yu., Bobrov, A. A., Tarasov, P. E., Ilyashuk, B.
648 P., Kuznetsova, T. V., Krbetschek, M., Meyer, H., and Kunitsky, V. V.: Late Saalian and Eemian palaeoenvironmental history
649 of the Bol'shoy Lyakhovsky Island (Laptev Sea region, Arctic Siberia), *Boreas*, 33, 319–348, [https://doi.org/10.1111/j.1502-](https://doi.org/10.1111/j.1502-3885.2004.tb01244.x)
650 [3885.2004.tb01244.x](https://doi.org/10.1111/j.1502-3885.2004.tb01244.x), 2004.
- 651 Ardenghi, N., Mulch, A., Koutsodendris, A., Pross, J., Kahmen, A., and Niedermeyer, E. M.: Temperature and moisture
652 variability in the eastern Mediterranean region during Marine Isotope Stages 11–10 based on biomarker analysis of the Tenaghi
653 Philippon peat deposit, *Quat. Sci. Rev.*, 225, 105977, <https://doi.org/10.1016/j.quascirev.2019.105977>, 2019.
- 654 van Asch, N., Kloos, M. E., Heiri, O., de Klerk, P., and Hoek, W. Z.: The younger dryas cooling in northeast Germany: summer
655 temperature and environmental changes in the Friedländer Große Wiese region, *J. Quat. Sci.*, 27, 531–543,
656 <https://doi.org/10.1002/jqs.2547>, 2012.
- 657 Atkinson, T. C., Briffa, K. R., and Coope, G. R.: Seasonal temperatures in Britain during the past 22,000 years, reconstructed
658 using beetle remains, *Nature*, 325, 587–592, <https://doi.org/10.1038/325587a0>, 1987.
- 659 Bailey, J. V., Cohen, A. S., and Kring, D. A.: Lacustrine Fossil Preservation in Acidic Environments: Implications of
660 Experimental and Field Studies for the Cretaceous–Paleogene Boundary Acid Rain Trauma, *PALAIOS*, 20, 376–389,
661 <https://doi.org/10.2110/palo.2003.p03-88>, 2005.
- 662 Battarbee, R. W.: Palaeolimnological approaches to climate change, with special regard to the biological record, *Quat. Sci.*
663 *Rev.*, 19, 107–124, 2000.
- 664 Bedford, A., Jones, Richard. T., Lang, B., Brooks, S., and Marshall, J. D.: A Late-glacial chironomid record from Hawes
665 Water, northwest England, *J. Quat. Sci.*, 19, 281–290, <https://doi.org/10.1002/jqs.836>, 2004.
- 666 Benham, S., Durrant, T., Caudullo, G., and de Rigo, D.: *Taxus baccata* in Europe: distribution, habitat, usage and threats, 2016.

667 Berglund, B. E. and Ralska-Jasiewiczowa, M.: Pollen analysis and pollen diagrams, Handb. Holocene Palaeoecol.
668 Palaeohydrology, 455, 484–486, 1986.

669 Beug, H.-J.: Leitfaden der Pollenbestimmung für Mitteleuropa und angrenzende Gebiete, No Title, 1961.

670 Bińka, K., Szymanek, M., and Nitychoruk, J.: Climate and vegetation changes recorded in the post Holsteinian lake deposits
671 at Ossówka (eastern Poland), Acta Geol. Pol., 341–354, 2023.

672 Blindow, I.: Decline of charophytes during eutrophication: comparison with angiosperms, Freshw. Biol., 28, 9–14,
673 <https://doi.org/10.1111/j.1365-2427.1992.tb00557.x>, 1992.

674 Bolland, A., Kern, O. A., Allstädt, F. J., Peteet, D., Koutsodendris, A., Pross, J., and Heiri, O.: Summer temperatures during
675 the last glaciation (MIS 5c to MIS 3) inferred from a 50,000-year chironomid record from Füramoos, southern Germany, Quat.
676 Sci. Rev., 264, 107008, <https://doi.org/10.1016/j.quascirev.2021.107008>, 2021.

677 Boubee, J. a. T.: Past and present benthic fauna of Lake Maratoto with special reference to the chironomidae, The University
678 of Waikato, 1983.

679 ter Braak, C. J. F. and Juggins, S.: Weighted averaging partial least squares regression (WA-PLS): an improved method for
680 reconstructing environmental variables from species assemblages, Hydrobiologia, 269, 485–502,
681 <https://doi.org/10.1007/BF00028046>, 1993.

682 ter Braak, C. J. F., H., van D., Juggins, S., and Birks, H. J. B.: Calibration of modern diatom-climate regressions for
683 palaeoenvironmental reconstruction, in: Proceedings of the European Palaeoclimate and Man Symposium, European
684 Palaeoclimate and Man Symposium, Stuttgart, 1993.

685 Briggs, D. E. G. and Kear, A. J.: Fossilization of Soft Tissue in the Laboratory, Science, 259, 1439–1442,
686 <https://doi.org/10.1126/science.259.5100.1439>, 1993.

687 Brodersen, K. P. and Lindegaard, C.: Classification, assessment and trophic reconstruction of Danish lakes using chironomids,
688 Freshw. Biol., 42, 143–157, <https://doi.org/10.1046/j.1365-2427.1999.00457.x>, 1999a.

689 Brodersen, K. P. and Lindegaard, C.: Mass occurrence and sporadic distribution of *Corynocera ambigua* Zetterstedt (Diptera,
690 Chironomidae) in Danish lakes. Neo- and palaeolimnological records, J. Paleolimnol., 22, 41–52,
691 <https://doi.org/10.1023/A:1008032619776>, 1999b.

692 Brondizio, E. S., O'Brien, K., Bai, X., Biermann, F., Steffen, W., Berkhout, F., Cudennec, C., Lemos, M. C., Wolfe, A., Palma-
693 Oliveira, J., and Chen, C.-T. A.: Re-conceptualizing the Anthropocene: A call for collaboration, Glob. Environ. Change, 39,
694 318–327, <https://doi.org/10.1016/j.gloenvcha.2016.02.006>, 2016.

695 Brooks, S. J.: Fossil midges (Diptera: Chironomidae) as palaeoclimatic indicators for the Eurasian region, Quat. Sci. Rev., 25,
696 1894–1910, <https://doi.org/10.1016/j.quascirev.2005.03.021>, 2006.

697 Brooks, S. J., Langdon, P. G., and Heiri, O.: The identification and use of Palaearctic Chironomidae larvae in palaeoecology,
698 Quat. Res. Assoc. Tech. Guide, i-vi,1, 2007.

699 Brundin, L.: Chironomiden und andere bodentiere der südschwedischen urgebirgsseen: Ein beitrag zur kenntnis der
700 bodenfaunistischen charakterzüge schwedischer oligotropher seen, Fiskeristytelsen, 1949.

701 Butler, M. G.: A 7-year life cycle for two *Chironomus* species in arctic Alaskan tundra ponds (Diptera: Chironomidae), *Can.*
702 *J. Zool.*, 60, 58–70, <https://doi.org/10.1139/z82-008>, 1982.

703 Camarero, J. J., Sánchez-Salguero, R., Sangüesa-Barreda, G., Lechuga, V., Viñegla, B., Seco, J. I., Taïqui, L., Carreira, J. A.,
704 and Linares, J. C.: Reply to the letter to editor regarding Camarero et al. (2021): Overgrazing and pollarding threaten Atlas
705 cedar conservation under forecasted aridification regardless stakeholders' nature, *For. Ecol. Manag.*, 503, 119779,
706 <https://doi.org/10.1016/j.foreco.2021.119779>, 2022.

707 Candy, I., Schreve, D. C., Sherriff, J., and Tye, G. J.: Marine Isotope Stage 11: Palaeoclimates, palaeoenvironments and its
708 role as an analogue for the current interglacial, *Earth-Sci. Rev.*, 128, 18–51, <https://doi.org/10.1016/j.earscirev.2013.09.006>,
709 2014.

710 Caudullo, G., Tinner, W., and De Rigo, D.: *Picea abies* in Europe: distribution, habitat, usage and threats, 2016.

711 Charlton, M. N.: Hypolimnion Oxygen Consumption in Lakes: Discussion of Productivity and Morphometry Effects, *Can. J.*
712 *Fish. Aquat. Sci.*, 37, 1531–1539, <https://doi.org/10.1139/f80-198>, 1980.

713 Chevalier, M., Davis, B. A. S., Heiri, O., Seppä, H., Chase, B. M., Gajewski, K., Lacourse, T., Telford, R. J., Finsinger, W.,
714 Guiot, J., Kühn, N., Maezumi, S. Y., Tipton, J. R., Carter, V. A., Brussel, T., Phelps, L. N., Dawson, A., Zanon, M., Vallé, F.,
715 Nolan, C., Mauri, A., de Vernal, A., Izumi, K., Holmström, L., Marsicek, J., Goring, S., Sommer, P. S., Chaput, M., and
716 Kupriyanov, D.: Pollen-based climate reconstruction techniques for late Quaternary studies, *Earth-Sci. Rev.*, 210, 103384,
717 <https://doi.org/10.1016/j.earscirev.2020.103384>, 2020.

718 Cornette, R., Gusev, O., Nakahara, Y., Shimura, S., Kikawada, T., and Okuda, T.: Chironomid Midges (Diptera,
719 Chironomidae) Show Extremely Small Genome Sizes, *Zoolog. Sci.*, 32, 248–254, <https://doi.org/10.2108/zs140166>, 2015.

720 Danks, H. V.: OVERWINTERING OF SOME NORTH TEMPERATE AND ARCTIC CHIRONOMIDAE: II.
721 CHIRONOMID BIOLOGY, *Can. Entomol.*, 103, 1875–1910, <https://doi.org/10.4039/Ent1031875-12>, 1971.

722 Davis, S., Golladay, S. W., Vellidis, G., and Pringle, C. M.: Macroinvertebrate Biomonitoring in Intermittent Coastal Plain
723 Streams Impacted by Animal Agriculture, *J. Environ. Qual.*, 32, 1036–1043, <https://doi.org/10.2134/jeq2003.1036>, 2003.

724 Del Wayne, R. N., Herrmann, S. J., Sublette, J. E., Melnykov, I. V., Helland, L. K., Romine, J. A., Carsella, J. S., Herrmann-
725 Hoesing, L. M., Turner, J. A., and Heuvel, B. D. V.: Occurrence of Chironomid Species (Diptera: Chironomidae) in the High
726 Se-78 Concentrations and High pH of Fountain Creek Watershed, Colorado, USA, *West. North Am. Nat.*, 78, 39–64,
727 <https://doi.org/10.3398/064.078.0106>, 2018.

728 Donato, M. and Paggi, A. C.: *Polypedilum parthenogeneticum* (Diptera: Chironomidae): a new parthenogenetic species from
729 *Eryngium* L. (Apiaceae) phytotelmata, *Aquat. Insects*, 30, 51–60, <https://doi.org/10.1080/01650420701829633>, 2008.

730 Drozd, M. and Trzepla, M.: *Objaśnienia do Szczegółowej mapy geologicznej Polski 1:50 000*, Arkusz Kock (676), 2007.

731 Druzhinina, O., Kublitskiy, Y., Stančikaitė, M., Nazarova, L., Syrykh, L., Gedminienė, L., Uogintas, D., Skipityte, R.,
732 Arslanov, K., Vaikutienė, G., Kulkova, M., and Subetto, D.: The Late Pleistocene–Early Holocene palaeoenvironmental
733 evolution in the SE Baltic region: a new approach based on chironomid, geochemical and isotopic data from Kamyshevoye
734 Lake, Russia, *Boreas*, 49, 544–561, <https://doi.org/10.1111/bor.12438>, 2020.

735 Dumayne-Peaty, L.: Human Impact on the Environment during the Iron Age and Romano-British Times: Palynological
 736 Evidence from Three Sites near the Antonine Wall, Great Britain, *J. Archaeol. Sci.*, 25, 203–214,
 737 <https://doi.org/10.1006/jasc.1997.0205>, 1998.
 738 Eggermont, H. and Heiri, O.: The chironomid-temperature relationship: expression in nature and palaeoenvironmental
 739 implications, *Biol. Rev.*, 87, 430–456, <https://doi.org/10.1111/j.1469-185X.2011.00206.x>, 2012.
 740 Engels, S., Bohncke, S. J. P., Bos, J. A. A., Brooks, S. J., Heiri, O., and Helmens, K. F.: Chironomid-based palaeotemperature
 741 estimates for northeast Finland during Oxygen Isotope Stage 3, *J. Paleolimnol.*, 40, 49–61, [https://doi.org/10.1007/s10933-](https://doi.org/10.1007/s10933-007-9133-y)
 742 [007-9133-y](https://doi.org/10.1007/s10933-007-9133-y), 2008.
 743 Erdtman, G.: Pollen walls and angiosperm phylo-geny., 1960.
 744 Fawcett, P. J., Werne, J. P., Anderson, R. S., Heikoo, J. M., Brown, E. T., Berke, M. A., Smith, S. J., Goff, F., Donohoo-
 745 Hurley, L., Cisneros-Dozal, L. M., Schouten, S., Sinninghe Damsté, J. S., Huang, Y., Toney, J., Fessenden, J., WoldeGabriel,
 746 G., Atudorei, V., Geissman, J. W., and Allen, C. D.: Extended megadroughts in the southwestern United States during
 747 Pleistocene interglacials, *Nature*, 470, 518–521, <https://doi.org/10.1038/nature09839>, 2011.
 748 Fernández Arias, S., Förster, M. W., and Sirocko, F.: Rieden tephra layers in the Dottinger Maar lake sediments: Implications
 749 for the dating of the Holsteinian interglacial and Elsterian glacial, *Glob. Planet. Change*, 227, 104143,
 750 <https://doi.org/10.1016/j.gloplacha.2023.104143>, 2023.
 751 Fjellberg, A.: Present and late weichselian occurrence of *Corynocera ambigua* zett.(Diptera, Chironomidae) in Norway, *Nor.*
 752 *Entomol Tidsskr*, 1972.
 753 Forsberg, C.: Nutritional Studies of *Chara* in Axenic Cultures, *Physiol. Plant.*, 18, 275–290, [https://doi.org/10.1111/j.1399-](https://doi.org/10.1111/j.1399-3054.1965.tb06890.x)
 754 [3054.1965.tb06890.x](https://doi.org/10.1111/j.1399-3054.1965.tb06890.x), 1965.
 755 Foster, G. L. and Rae, J. W. B.: Reconstructing Ocean pH with Boron Isotopes in Foraminifera, *Annu. Rev. Earth Planet. Sci.*,
 756 44, 207–237, <https://doi.org/10.1146/annurev-earth-060115-012226>, 2016.
 757 Gandouin, E., Ponel, P., Andrieu-Ponel, V., Franquet, É., Beaulieu, J.-L. de, Reille, M., Guiter, F., Brulhet, J., Lallier-Vergès,
 758 É., Keravis, D., Grafenstein, U. von, and Veres, D.: Past environment and climate changes at the last Interglacial/Glacial
 759 transition (Les Échets, France) inferred from subfossil chironomids (Insecta), *Comptes Rendus Géoscience*, 339, 337–346,
 760 <https://doi.org/10.1016/j.crte.2007.03.002>, 2007.
 761 Gandouin, E., Rioual, P., Pailles, C., Brooks, S. J., Ponel, P., Guiter, F., Djamali, M., Andrieu-Ponel, V., Birks, H. J. B.,
 762 Leydet, M., Belkacem, D., Haas, J. N., Van der Putten, N., and de Beaulieu, J. L.: Environmental and climate reconstruction
 763 of the late-glacial-Holocene transition from a lake sediment sequence in Aubrac, French Massif Central: Chironomid and
 764 diatom evidence, *Palaeogeogr. Palaeoclimatol. Palaeoecol.*, 461, 292–309, <https://doi.org/10.1016/j.palaeo.2016.08.039>, 2016.
 765 Gedminienė, L., Spiridonov, A., Stančikaitė, M., Skuratovič, Ž., Vaikutienė, G., Daumantas, L., and Salonen, J. S.: Temporal
 766 and spatial climate changes in the mid-Baltic region in the Late Glacial and the Holocene: Pollen-based reconstructions,
 767 *CATENA*, 252, 108851, <https://doi.org/10.1016/j.catena.2025.108851>, 2025.
 768 Giorgi, F.: Climate change hot-spots, *Geophys. Res. Lett.*, 33, <https://doi.org/10.1029/2006GL025734>, 2006.

769 Górecki, A.: Plant succession as an indicator of climatic oscillations during Masovian Interglacial (MIS 11c), PhD Thesis,
770 2023.

771 Górecki, A., Żarski, M., Drzewicki, W., Pleśniak, Ł., Zalewska-Gałosz, J., and Hrynowiecka, A.: New climatic oscillations
772 during MIS 11c in the record of the Skrzyńka II site (Eastern Poland) based on palynological and isotope analysis, *Quat. Int.*,
773 632, 4–20, <https://doi.org/10.1016/j.quaint.2021.09.017>, 2022.

774 Guiot, J.: Methodology of the last climatic cycle reconstruction in France from pollen data, *Palaeogeogr. Palaeoclimatol.*
775 *Palaeoecol.*, 80, 49–69, [https://doi.org/10.1016/0031-0182\(90\)90033-4](https://doi.org/10.1016/0031-0182(90)90033-4), 1990.

776 Gusev, O., Nakahara, Y., Vanyagina, V., Malutina, L., Cornette, R., Sakashita, T., Hamada, N., Kikawada, T., Kobayashi, Y.,
777 and Okuda, T.: Anhydrobiosis-Associated Nuclear DNA Damage and Repair in the Sleeping Chironomid: Linkage with
778 Radioresistance, *PLOS ONE*, 5, e14008, <https://doi.org/10.1371/journal.pone.0014008>, 2010.

779 Halkiewicz, A.: *Corynocera ambigua* (Insecta, Diptera) subfossils occurrence in recent sediments of four shallow Polesie lakes,
780 in: *Annales Universitatis Mariae Curie-Skłodowska*, 31, 2008.

781 Hamburger, K., Dall, P. C., and Lindegaard, C.: Energy metabolism of *Chironomus anthracinus* (Diptera: Chironomidae) from
782 the profundal zone of Lake Esrom, Denmark, as a function of body size, temperature and oxygen concentration, *Hydrobiologia*,
783 294, 43–50, <https://doi.org/10.1007/BF00017624>, 1994.

784 Harrington, C. A. and Gould, P. J.: Tradeoffs between chilling and forcing in satisfying dormancy requirements for Pacific
785 Northwest tree species, *Front. Plant Sci.*, 6, 120, 2015.

786 Heino, J.: Lentic macroinvertebrate assemblage structure along gradients in spatial heterogeneity, habitat size and water
787 chemistry, *Hydrobiologia*, 418, 229–242, <https://doi.org/10.1023/A:1003969217686>, 2000.

788 Heiri, O.: Within-lake variability of subfossil chironomid assemblages in shallow Norwegian lakes, *J. Paleolimnol.*, 32, 67–
789 84, <https://doi.org/10.1023/B:JOPL.0000025289.30038.e9>, 2004.

790 Heiri, O. and Lotter, A. F.: Effect of low count sums on quantitative environmental reconstructions: an example using subfossil
791 chironomids, *J. Paleolimnol.*, 26, 343–350, <https://doi.org/10.1023/A:1017568913302>, 2001.

792 Heiri, O. and Millet, L.: Reconstruction of Late Glacial summer temperatures from chironomid assemblages in Lac Lautrey
793 (Jura, France), *J. Quat. Sci.*, 20, 33–44, <https://doi.org/10.1002/jqs.895>, 2005.

794 Herzschuh, U., Böhmer, T., Li, C., Chevalier, M., Hébert, R., Dallmeyer, A., Cao, X., Bigelow, N. H., Nazarova, L., Novenko,
795 E. Y., Park, J., Peyron, O., Rudaya, N. A., Schlütz, F., Shumilovskikh, L. S., Tarasov, P. E., Wang, Y., Wen, R., Xu, Q., and
796 Zheng, Z.: LegacyClimate 1.0: A dataset of pollen-based climate reconstructions from 2594 Northern Hemisphere sites
797 covering the last 30 kyr and beyond, *Earth Syst. Sci. Data*, 15, 2235–2258, <https://doi.org/10.5194/essd-15-2235-2023>, 2023a.

798 Herzschuh, U., Böhmer, T., Chevalier, M., Hébert, R., Dallmeyer, A., Li, C., Cao, X., Peyron, O., Nazarova, L., Novenko, E.
799 Y., Park, J., Rudaya, N. A., Schlütz, F., Shumilovskikh, L. S., Tarasov, P. E., Wang, Y., Wen, R., Xu, Q., and Zheng, Z.:
800 Regional pollen-based Holocene temperature and precipitation patterns depart from the Northern Hemisphere mean trends,
801 *Clim. Past*, 19, 1481–1506, <https://doi.org/10.5194/cp-19-1481-2023>, 2023b.

802 Hofmann, W. and Winn, K.: The Littorina Transgression in the Western Baltic Sea as Indicated by Subfossil Chironomidae
 803 (Diptera) and Cladocera (Crustacea), *Int. Rev. Hydrobiol.*, 85, 267–291, [https://doi.org/10.1002/\(SICI\)1522-2632\(200004\)85:2/3<267::AID-IROH267>3.0.CO;2-Q](https://doi.org/10.1002/(SICI)1522-2632(200004)85:2/3<267::AID-IROH267>3.0.CO;2-Q), 2000.
 804
 805 Horne, D. J.: A Mutual Temperature Range method for Quaternary palaeoclimatic analysis using European nonmarine
 806 Ostracoda, *Quat. Sci. Rev.*, 26, 1398–1415, <https://doi.org/10.1016/j.quascirev.2007.03.006>, 2007.
 807 Horne, D. J., Ashton, N., Benardout, G., Brooks, S. J., Coope, G. R., Holmes, J. A., Lewis, S. G., Parfitt, S. A., White, T. S.,
 808 Whitehouse, N. J., and Whittaker, J. E.: A terrestrial record of climate variation during MIS 11 through multiproxy
 809 palaeotemperature reconstructions from Hoxne, UK, *Quat. Res.*, 111, 21–52, <https://doi.org/10.1017/qua.2022.20>, 2023.
 810 Hrynowiecka, A. and Pidek, I. A.: Older and Younger Holsteinian climate oscillations in the palaeobotanical record of the
 811 Brus profile (SE Poland), *Geol. Q.*, 61, 723–737, doi: 10.7306/gq.1358, <https://doi.org/10.7306/gq.1358>, 2017.
 812 Hrynowiecka, A. and Winter, H.: Palaeoclimatic changes in the Holsteinian Interglacial (Middle Pleistocene) on the basis of
 813 indicator-species method – Palynological and macrofossils remains from Nowiny Żukowskie site (SE Poland), *Quat. Int.*, 409,
 814 255–269, <https://doi.org/10.1016/j.quaint.2015.08.036>, 2016.
 815 Hrynowiecka, A., Źarski, M., and Drzewicki, W.: The rank of climatic oscillations during MIS 11c (OHO and YHO) and post-
 816 interglacial cooling during MIS 11b and MIS 11a in eastern Poland, *Geol. Q.*, 63, 375–394, doi: 10.7306/gq.1470,
 817 <https://doi.org/10.7306/gq.1470>, 2019.
 818 Ilyashuk, E. A., Ilyashuk, B. P., Hammarlund, D., and Larocque, I.: Holocene climatic and environmental changes inferred
 819 from midge records (Diptera: Chironomidae, Chaoboridae, Ceratopogonidae) at Lake Berkut, southern Kola
 820 Peninsula, Russia, *The Holocene*, 15, 897–914, <https://doi.org/10.1191/0959683605hl865ra>, 2005.
 821 Ilyashuk, E. A., Ilyashuk, B. P., Kolka, V. V., and Hammarlund, D.: Holocene climate variability on the Kola Peninsula,
 822 Russian Subarctic, based on aquatic invertebrate records from lake sediments, *Quat. Res.*, 79, 350–361,
 823 <https://doi.org/10.1016/j.yqres.2013.03.005>, 2013.
 824 Ilyashuk, E. A., Ilyashuk, B. P., Heiri, O., and Spötl, C.: Summer temperatures and lake development during the MIS 5a
 825 interstadial: New data from the Unterangerberg palaeolake in the Eastern Alps, Austria, *Palaeogeogr. Palaeoclimatol.*
 826 *Palaeoecol.*, 560, 110020, <https://doi.org/10.1016/j.palaeo.2020.110020>, 2020.
 827 Ilyashuk, E. A., Ilyashuk, B. P., Heiri, O., and Spötl, C.: Summer temperatures and environmental dynamics during the Middle
 828 Würmian (MIS 3) in the Eastern Alps: Multi-proxy records from the Unterangerberg palaeolake, Austria, *Quat. Sci. Adv.*, 6,
 829 100050, <https://doi.org/10.1016/j.qsa.2022.100050>, 2022.
 830 Imbrie, J., Hays, J. D., Martinson, D. G., McIntyre, A., Mix, A. C., Morley, J. J., Pisias, N. G., Prell, W. L., and Shackleton,
 831 N. J.: The orbital theory of Pleistocene climate: support from a revised chronology of the marine $\delta^{18}\text{O}$ record, 1984.
 832 Iovino, A. J.: Extant chironomid larval populations and the representativeness and nature of their remains in lake sediments.,
 833 Indiana University, 1975.
 834 Janczyk-Kopikowa, Z.: The Ferdynandów Interglacial in Poland, *Geol. Q.*, 35, 71–80, 1991.
 835 Jesionkiewicz, P.: Nowe stanowisko interglacjału mazowieckiego w Krępie koło Kocka, *Geol. Q.*, 26, 423–430, 1982.

836 Juggins, S.: C2: Software for ecological and palaeoecological data analysis and visualisation (user guide version 1.5), Newctle.
837 Tyne Newctle. Univ., 77, 680, 2007.

838 Juggins, S.: riojaPlot: Stratigraphic diagrams in R, package version (0.1-20), 2022.

839 Juggins, S. and Birks, H. J. B.: Quantitative Environmental Reconstructions from Biological Data, in: Tracking Environmental
840 Change Using Lake Sediments: Data Handling and Numerical Techniques, edited by: Birks, H. J. B., Lotter, A. F., Juggins,
841 S., and Smol, J. P., Springer Netherlands, Dordrecht, 431–494, https://doi.org/10.1007/978-94-007-2745-8_14, 2012.

842 Kansanen, P. H.: Assessment of pollution history from recent sediments in Lake Vanajavesi, southern Finland. II. Changes in
843 the Chironomidae, Chaoboridae and Ceratopogonidae (Diptera) fauna, Ann. Zool. Fenn., 22, 57–90, 1985.

844 Kleinen, T., Brovkin, V., and Munhoven, G.: Modelled interglacial carbon cycle dynamics during the Holocene, the Eemian
845 and Marine Isotope Stage (MIS) 11, Clim. Past, 12, 2145–2160, <https://doi.org/10.5194/cp-12-2145-2016>, 2016.

846 Klink, A. G. and Moller Pillot, H.: Chironomidae larvae: key to the higher taxa and species of the lowlands of Northwestern
847 Europe., 2003.

848 Kokkinn, M. J.: Osmoregulation, salinity tolerance and the site of ion excretion in the halobiont chironomid, Tanytarsus
849 barbitarsis Freeman, Mar. Freshw. Res., 37, 243–250, <https://doi.org/10.1071/mf9860243>, 1986.

850 Kondratiene, O. and Gudelis, W.: Morskie osady plejstocénskie na obszarze Pribałtyki, Przegląd Geol., 31, 497, 1983.

851 Körner, C. and Paulsen, J.: A world-wide study of high altitude treeline temperatures, J. Biogeogr., 31, 713–732,
852 <https://doi.org/10.1111/j.1365-2699.2003.01043.x>, 2004.

853 Kotrys, B., Płóciennik, M., Sydor, P., and Brooks, S. J.: Expanding the Swiss-Norwegian chironomid training set with Polish
854 data, Boreas, 49, 89–107, <https://doi.org/10.1111/bor.12406>, 2020.

855 Kousis, I., Koutsodendris, A., Peyron, O., Leicher, N., Francke, A., Wagner, B., Giaccio, B., Knipping, M., and Pross, J.:
856 Centennial-scale vegetation dynamics and climate variability in SE Europe during Marine Isotope Stage 11 based on a pollen
857 record from Lake Ohrid, Quat. Sci. Rev., 190, 20–38, <https://doi.org/10.1016/j.quascirev.2018.04.014>, 2018.

858 Koutsodendris, A., Müller, U. C., Pross, J., Brauer, A., Kotthoff, U., and Lotter, A. F.: Vegetation dynamics and climate
859 variability during the Holsteinian interglacial based on a pollen record from Dethlingen (northern Germany), Quat. Sci. Rev.,
860 29, 3298–3307, <https://doi.org/10.1016/j.quascirev.2010.07.024>, 2010.

861 Koutsodendris, A., Pross, J., Müller, U. C., Brauer, A., Fletcher, W. J., Kühl, N., Kirilova, E., Verhagen, F. T. M., Lücke, A.,
862 and Lotter, A. F.: A short-term climate oscillation during the Holsteinian interglacial (MIS 11c): An analogy to the 8.2 ka
863 climatic event?, Glob. Planet. Change, 92–93, 224–235, <https://doi.org/10.1016/j.gloplacha.2012.05.011>, 2012.

864 Kreyling, J.: Winter climate change: A critical factor for temperate vegetation performance, Ecology, 91, 1939–1948,
865 <https://doi.org/10.1890/09-1160.1>, 2010.

866 Krupiński, K. M.: Taxus in plant communities of the Mazovian interglacial age in Central Europe and its climatostratigraphical
867 consequences, Bull. Pol. Acad. Sci. Earth Sci., 43, 1995.

868 Krzeminski, W. and Jarzembowski, E.: Aenne triassica sp.n., the oldest representative of the family Chironomidae [Insecta:
869 Diptera], Pol. Pismo Entomol., 68, 1999.

870 Kühl, N. and Gobet, E.: Climatic evolution during the Middle Pleistocene warm period of Bilshausen, Germany, compared to
871 the Holocene, *Quat. Sci. Rev.*, 29, 3736–3749, <https://doi.org/10.1016/j.quascirev.2010.08.006>, 2010.

872 Kupryjanowicz, M., Nalepka, D., Pidek, I. A., Walanus, A., Balwierz, Z., Bińka, K., Fiłoc, M., Granoszewski, W., Kołaczek,
873 P., Majecka, A., Malkiewicz, M., Nita, M., Noryśkiewicz, B., and Winter, H.: The east-west migration of trees during the
874 Eemian Interglacial registered on isopollen maps of Poland, *Quat. Int.*, 467, 178–191,
875 <https://doi.org/10.1016/j.quaint.2017.08.034>, 2018.

876 Lackmann, A. R., McEwen, D. C., and Butler, M. G.: Evidence of parthenogenetic populations from the *Paratanytarsus*
877 *laccophilus* species group (Diptera: Chironomidae) in the Alaskan Arctic, *CHIRONOMUS J. Chironomidae Res.*, 48–58,
878 <https://doi.org/10.5324/cjcr.v0i33.3478>, 2020.

879 Łanczont, M., Pidek, I. A., Bogucki, A., Wieliczekiewicz, F., and Wojtanowicz, J.: Kluczowy profil interglacjału
880 mazowieckiego w Krukienicach na międzyrzeczu Sanu i Dniestru (Ukraina), *Przegląd Geol.*, 51, 597–608, 2003.

881 Lapellegerie, P., Millet, L., Rius, D., Duprat-Oualid, F., Luoto, T., and Heiri, O.: Chironomid-inferred summer temperature
882 during the Last Glacial Maximum in the Southern Black Forest, Central Europe, *Quat. Sci. Rev.*, 345, 109016,
883 <https://doi.org/10.1016/j.quascirev.2024.109016>, 2024.

884 Larocque-Tobler, I., Grosjean, M., Heiri, O., and Trachsel, M.: High-resolution chironomid-inferred temperature history since
885 ad 1580 from varved Lake Silvaplana, Switzerland: comparison with local and regional reconstructions, *The Holocene*, 19,
886 1201–1212, <https://doi.org/10.1177/0959683609348253>, 2009.

887 Lauer, T. and Weiss, M.: Timing of the Saalian- and Elsterian glacial cycles and the implications for Middle – Pleistocene
888 hominin presence in central Europe, *Sci. Rep.*, 8, 5111, <https://doi.org/10.1038/s41598-018-23541-w>, 2018.

889 Lauer, T., Weiss, M., Bernhardt, W., Heinrich, S., Rappsilber, I., Stahlschmidt, M. C., von Suchodoletz, H., and Wansa, S.:
890 The Middle Pleistocene fluvial sequence at Uichteritz, central Germany: Chronological framework, paleoenvironmental
891 history and early human presence during MIS 11, *Geomorphology*, 354, 107016,
892 <https://doi.org/10.1016/j.geomorph.2019.107016>, 2020.

893 Lenarczyk, J.: The Algal Genus "pediastrum Meyen"(chlorophyta) in Poland, W. Szafer Institute of Botany, Polish Academy
894 of Sciences, 2014.

895 Lencioni, V.: Survival strategies of freshwater insects in cold environments, 2004.

896 Lionello, P., Malanotte-Rizzoli, P., Boscolo, R., Alpert, P., Artale, V., Li, L., Luterbacher, J., May, W., Trigo, R., Tsimplis,
897 M., Ulbrich, U., and Xoplaki, E.: The Mediterranean climate: An overview of the main characteristics and issues, in:
898 *Developments in Earth and Environmental Sciences*, vol. 4, edited by: Lionello, P., Malanotte-Rizzoli, P., and Boscolo, R.,
899 Elsevier, 1–26, [https://doi.org/10.1016/S1571-9197\(06\)80003-0](https://doi.org/10.1016/S1571-9197(06)80003-0), 2006.

900 Lotter, A. F., Birks, H. J. B., Hofmann, W., and Marchetto, A.: Modern diatom, cladocera, chironomid, and chrysophyte cyst
901 assemblages as quantitative indicators for the reconstruction of past environmental conditions in the Alps. I. Climate, *J.*
902 *Paleolimnol.*, 18, 395–420, <https://doi.org/10.1023/A:1007982008956>, 1997.

903 Luoto, T. P. and Sarmaja-Korjonen, K.: Midge-inferred Holocene effective moisture fluctuations in a subarctic lake, northern
 904 Lapland, *Boreas*, 40, 650–659, <https://doi.org/10.1111/j.1502-3885.2011.00217.x>, 2011.

905 Luoto, T. P., Nevalainen, L., and Sarmaja-Korjonen, K.: Multiproxy evidence for the ‘Little Ice Age’ from Lake Hamträsk,
 906 Southern Finland, *J. Paleolimnol.*, 40, 1097–1113, <https://doi.org/10.1007/s10933-008-9216-4>, 2008.

907 Magny, M., Aalbersberg, G., Bégeot, C., Benoit-Ruffaldi, P., Bossuet, G., Disnar, J.-R., Heiri, O., Laggoun-Defarge, F.,
 908 Mazier, F., Millet, L., Peyron, O., Vannière, B., and Walter-Simonnet, A.-V.: Environmental and climatic changes in the Jura
 909 mountains (eastern France) during the Lateglacial–Holocene transition: a multi-proxy record from Lake Lautrey, *Quat. Sci.*
 910 *Rev.*, 25, 414–445, <https://doi.org/10.1016/j.quascirev.2005.02.005>, 2006.

911 Mamakowa, K. and Rylova, T. B.: The interglacial from Korchevo in Belarus in the light of new palaeobotanical studies, *Acta*
 912 *Palaeobot.*, 47, 2007.

913 Marks, L.: Quaternary stratigraphy of Poland–current status, *Acta Geol. Pol.*, 73, 307–340, 2023.

914 Marks, L., Karabanov, A., Nitychoruk, J., Bahdasarau, M., Krzywicki, T., Majecka, A., Pochocka-Szwarc, K., Rychel, J.,
 915 Woronko, B., Zbucki, Ł., Hradunova, A., Hrychanik, M., Mamchuk, S., Rylova, T., Nowacki, Ł., and Pielach, M.: Revised
 916 limit of the Saalian ice sheet in central Europe, *Quat. Int.*, 478, 59–74, <https://doi.org/10.1016/j.quaint.2016.07.043>, 2018.

917 Masson-Delmotte, V., Stenni, B., Pol, K., Braconnot, P., Cattani, O., Falourd, S., Kageyama, M., Jouzel, J., Landais, A.,
 918 Minster, B., Barnola, J. M., Chappellaz, J., Krinner, G., Johnsen, S., Röthlisberger, R., Hansen, J., Mikolajewicz, U., and Otto-
 919 Bliesner, B.: EPICA Dome C record of glacial and interglacial intensities, *Quat. Sci. Rev.*, 29, 113–128,
 920 <https://doi.org/10.1016/j.quascirev.2009.09.030>, 2010.

921 Matzinger, A., Müller, B., Niederhauser, P., Schmid, M., and Wüest, A.: Hypolimnetic oxygen consumption by sediment-
 922 based reduced substances in former eutrophic lakes, *Limnol. Oceanogr.*, 55, 2073–2084,
 923 <https://doi.org/10.4319/lo.2010.55.5.2073>, 2010.

924 Mauri, A., Davis, B. A. S., Collins, P. M., and Kaplan, J. O.: The climate of Europe during the Holocene: a gridded pollen-
 925 based reconstruction and its multi-proxy evaluation, *Quat. Sci. Rev.*, 112, 109–127,
 926 <https://doi.org/10.1016/j.quascirev.2015.01.013>, 2015.

927 McGarrigle, M. L.: The Distribution of Chironomid Communities and Controlling Sediment Parameters in L. Derravaragh,
 928 Ireland, in: *Chironomidae*, edited by: Murray, D. A., Pergamon, 275–282, [https://doi.org/10.1016/B978-0-08-025889-](https://doi.org/10.1016/B978-0-08-025889-8.50043-X)
 929 [8.50043-X](https://doi.org/10.1016/B978-0-08-025889-8.50043-X), 1980.

930 Mousavi, S. K.: Boreal chironomid communities and their relations to environmental factors: the impact of lake depth, size
 931 and acidity, *Boreal Chironomid Communities Their Relat. Environ. Factors Impact Lake Depth Size Acidity*, 7, 63–75, 2002.

932 Muhs, D. R., Pandolfi, J. M., Simmons, K. R., and Schumann, R. R.: Sea-level history of past interglacial periods from
 933 uranium-series dating of corals, Curaçao, Leeward Antilles islands, *Quat. Res.*, 78, 157–169,
 934 <https://doi.org/10.1016/j.yqres.2012.05.008>, 2012.

935 Müller, B., Bryant, L. D., Matzinger, A., and Wüest, A.: Hypolimnetic Oxygen Depletion in Eutrophic Lakes, *Environ. Sci.*
 936 *Technol.*, 46, 9964–9971, <https://doi.org/10.1021/es301422r>, 2012.

937 Nazarova, L. B., Subetto, D. A., Syrykh, L. S., Grekov, I. M., and Leontev, P. A.: Reconstructions of Paleoecological and
 938 Paleoclimatic Conditions of the Late Pleistocene and Holocene according to the Results of Chironomid Analysis of Sediments
 939 from Medvedevskoe Lake (Karelian Isthmus), *Dokl. Earth Sci.*, 480, 710–714, <https://doi.org/10.1134/S1028334X18060144>,
 940 2018.

941 Nienstaedt, H.: Chilling requirements in seven *Picea* species, *Silvae Genet.*, 16, 65–68, 1967.

942 Nitychoruk, J., Bińka, K., Hoefs, J., Ruppert, H., and Schneider, J.: Climate reconstruction for the Holsteinian Interglacial in
 943 eastern Poland and its comparison with isotopic data from Marine Isotope Stage 11, *Quat. Sci. Rev.*, 24, 631–644,
 944 <https://doi.org/10.1016/j.quascirev.2004.07.023>, 2005.

945 Nitychoruk, J., Bińka, K., Ruppert, H., and Schneider, J.: Holsteinian Interglacial=Marine Isotope Stage 11?, *Quat. Sci. Rev.*,
 946 25, 2678–2681, <https://doi.org/10.1016/j.quascirev.2006.07.004>, 2006.

947 Nitychoruk, J., Bińka, K., Sienkiewicz, E., Szymanek, M., Chodyka, M., Makos, M., Ruppert, H., and Tudryn, A.: A
 948 multiproxy record of the Younger Holsteinian Oscillation (YHO) in the Ossówka profile, eastern Poland, *Boreas*, 47, 855–
 949 868, <https://doi.org/10.1111/bor.12308>, 2018.

950 Nondula, N., Marshall, D. J., Baxter, R., Sinclair, B. J., and Chown, S. L.: Life history and osmoregulatory ability of
 951 *Telmatogeton amphibius* (Diptera, Chironomidae) at Marion Island, *Polar Biol.*, 27, 629–635, [https://doi.org/10.1007/s00300-](https://doi.org/10.1007/s00300-004-0619-z)
 952 004-0619-z, 2004.

953 Oliveira, D., Desprat, S., Rodrigues, T., Naughton, F., Hodell, D., Trigo, R., Rufino, M., Lopes, C., Abrantes, F., and Goni,
 954 M. F. S.: The complexity of millennial-scale variability in southwestern Europe during MIS 11, *Quat. Res.*, 86, 373–387,
 955 <https://doi.org/10.1016/j.yqres.2016.09.002>, 2016.

956 Orel, O. and Semenchenko, A.: Morphological description and DNA barcodes of adult males of *Tanytarsus heliomesonctios*
 957 Langton, 1999 (Diptera, Chironomidae) in northeast of Russia, *Zootaxa*, 4686, <https://doi.org/10.11646/zootaxa.4686.1.6>,
 958 2019.

959 Overpeck, J. T., Iii, T. W., and Prentice, I. C.: Quantitative Interpretation of Fossil Pollen Spectra: Dissimilarity Coefficients
 960 and the Method of Modern Analogs, *Quat. Res.*, 23, 87–108, [https://doi.org/10.1016/0033-5894\(85\)90074-2](https://doi.org/10.1016/0033-5894(85)90074-2), 1985.

961 Park, L. E.: Geochemical and Paleoenvironmental Analysis of Lacustrine Arthropod-Bearing Concretions of the Barstow
 962 Formation, Southern California, *PALAIOS*, 10, 44–57, <https://doi.org/10.2307/3515006>, 1995.

963 Peel, M. C., Finlayson, B. L., and McMahon, T. A.: Updated world map of the Köppen-Geiger climate classification, *Hydrol.*
 964 *Earth Syst. Sci.*, 11, 1633–1644, <https://doi.org/10.5194/hess-11-1633-2007>, 2007.

965 Pinder, L. C. V. and Reiss, F.: The larvae of Chironominae (Diptera : Chironomidae) of the Holarctic region-Keys and
 966 diagnoses, *Larvae*, 1983.

967 Pliikk, A., Engels, S., Luoto, T. P., Nazarova, L., Salonen, J. S., and Helmens, K. F.: Chironomid-based temperature
 968 reconstruction for the Eemian Interglacial (MIS 5e) at Sokli, northeast Finland, *J. Paleolimnol.*, 61, 355–371,
 969 <https://doi.org/10.1007/s10933-018-00064-y>, 2019.

970 Plóciennik, M.: Zastosowanie subfosylnych szczątków ochotkowatych (Diptera: Chironomidae) w badaniach nad
 971 paleoklimatem i rekonstrukcją zmian w środowisku, *Kosmos*, 54, 401–406, 2005.

972 Plóciennik, M., Self, A., Birks, H. J. B., and Brooks, S. J.: Chironomidae (Insecta: Diptera) succession in Żabieniec bog and
 973 its palaeo-lake (central Poland) through the Late Weichselian and Holocene, *Palaeogeogr. Palaeoclimatol. Palaeoecol.*, 307,
 974 150–167, <https://doi.org/10.1016/j.palaeo.2011.05.010>, 2011.

975 Plóciennik, M., Kruk, A., Michczyńska, D. J., and Birks, J. B.: Kohonen Artificial Neural Networks and the IndVal Index as
 976 Supplementary Tools for the Quantitative Analysis of Palaeoecological Data, *Geochronometria*, 42,
 977 <https://doi.org/10.1515/geochr-2015-0021>, 2015.

978 Plóciennik, M., Pawłowski, D., Vilizzi, L., and Antczak-Orlewska, O.: From oxbow to mire: Chironomidae and Cladocera as
 979 habitat palaeoindicators, *Hydrobiologia*, 847, 3257–3275, <https://doi.org/10.1007/s10750-020-04327-6>, 2020.

980 Pochocka-Szwarc, K., Żarski, M., Hrynowiecka, A., Górecki, A., Pidek, I. A., Szymanek, M., Stachowicz-Rybka, R.,
 981 Stachowicz, K., and Skoczylas-Śniaz, S.: Mazovian Interglacial sites in the Sosnowica Depression and the Parczew-Kodeń
 982 Heights (Western Polesie, SE Poland), and their stratigraphic, palaeogeographic and palaeoenvironmental significance, *Geol.*
 983 *Q.*, 68, 68: 18-doi: 10.7306/gq.1746, <https://doi.org/10.7306/gq.1746>, 2024.

984 Prokopenko, A. A., Bezrukova, E. V., Khursevich, G. K., Solotchina, E. P., Kuzmin, M. I., and Tarasov, P. E.: Climate in
 985 continental interior Asia during the longest interglacial of the past 500 000 years: the new MIS 11 records from Lake Baikal,
 986 SE Siberia, *Clim. Past*, 6, 31–48, <https://doi.org/10.5194/cp-6-31-2010>, 2010.

987 Ralska-Jasiewiczowa, M., Nalepka, D., and Goslar, T.: Some problems of forest transformation at the transition to the
 988 oligocratic/ *Homo sapiens* phase of the Holocene interglacial in northern lowlands of central Europe, *Veg. Hist.*
 989 *Archaeobotany*, 13, 71–71, <https://doi.org/10.1007/s00334-003-0027-2>, 2004.

990 Raymo, M. E. and Mitrovica, J. X.: Collapse of polar ice sheets during the stage 11 interglacial, *Nature*, 483, 453–456,
 991 <https://doi.org/10.1038/nature10891>, 2012.

992 Reille, M. and de Beaulieu, J.-L.: Long Pleistocene Pollen Records from the Praclaux Crater, South-Central France, *Quat.*
 993 *Res.*, 44, 205–215, <https://doi.org/10.1006/qres.1995.1065>, 1995.

994 Reyes-Maldonado, R., Marie, B., and Ramírez, A.: Rearing methods and life cycle characteristics of *Chironomus* sp. Florida
 995 (Chironomidae: Diptera): A rapid-developing species for laboratory studies, *PLOS ONE*, 16, e0247382,
 996 <https://doi.org/10.1371/journal.pone.0247382>, 2021.

997 Rigterink, S., Krahn, K. J., Kotrys, B., Urban, B., Heiri, O., Turner, F., Pannes, A., and Schwalb, A.: Summer temperatures
 998 from the Middle Pleistocene site Schöningen 13 II , northern Germany, determined from subfossil chironomid assemblages,
 999 *Boreas*, bor.12658, <https://doi.org/10.1111/bor.12658>, 2024.

1000 Roberts, J., Kaczmarek, K., Langer, G., Skinner, L. C., Bijma, J., Bradbury, H., Turchyn, A. V., Lamy, F., and Misra, S.:
 1001 Lithium isotopic composition of benthic foraminifera: A new proxy for paleo-pH reconstruction, *Geochim. Cosmochim. Acta*,
 1002 236, 336–350, <https://doi.org/10.1016/j.gca.2018.02.038>, 2018.

1003 Roberts, M.: Investigation into the population dynamics and key life history characteristics of non-biting midge (Diptera:
 1004 Chironomidae) which can be utilised to improve nuisance control at Lake Joondalup, Western Australia, Theses Honours,
 1005 2003.

1006 Robinson, A., Alvarez-Solas, J., Calov, R., Ganopolski, A., and Montoya, M.: MIS-11 duration key to disappearance of the
 1007 Greenland ice sheet, *Nat. Commun.*, 8, 16008, <https://doi.org/10.1038/ncomms16008>, 2017.

1008 Rodrigues, T., Voelker, A. H. L., Grimalt, J. O., Abrantes, F., and Naughton, F.: Iberian Margin sea surface temperature during
 1009 MIS 15 to 9 (580–300 ka): Glacial suborbital variability versus interglacial stability, *Paleoceanography*, 26,
 1010 <https://doi.org/10.1029/2010PA001927>, 2011.

1011 Rohling, E. J., Braun, K., Grant, K., Kucera, M., Roberts, A. P., Siddall, M., and Trommer, G.: Comparison between Holocene
 1012 and Marine Isotope Stage-11 sea-level histories, *Earth Planet. Sci. Lett.*, 291, 97–105,
 1013 <https://doi.org/10.1016/j.epsl.2009.12.054>, 2010.

1014 Rohrig, R., Beug, H., Trettin, R., and Morgenstern, P.: Subfossil chironomid assemblages as paleoenvironmental indicators in
 1015 lake faulersee (Germany), *Stud. Quat.*, 117–127, 2004.

1016 Rylova, T. and Savachenko, I.: Reconstruction of palaeotemperatures of pleistocene interglacial intervals of Belarus from
 1017 palynological evidences, *Pol. Geol. Inst. Spec. Pap.*, Vol. 16, 2005.

1018 Saether, O. A.: Chironomid communities as water quality indicators, *Ecography*, 2, 65–74, <https://doi.org/10.1111/j.1600->
 1019 [0587.1979.tb00683.x](https://doi.org/10.1111/j.1600-0587.1979.tb00683.x), 1979.

1020 Sageman, B. B. and Hollander, D. J.: Cross correlation of paleoecological and geochemical proxies: A holistic approach to the
 1021 study of past global change, *Spec. Pap. Geol. Soc. Am.*, 332, 365–384, <https://doi.org/10.1130/0-8137-2332-9.365>, 1999.

1022 Samartin, S., Heiri, O., Kaltenrieder, P., Köhl, N., and Tinner, W.: Reconstruction of full glacial environments and summer
 1023 temperatures from Lago della Costa, a refugial site in Northern Italy, *Quat. Sci. Rev.*, 143, 107–119,
 1024 <https://doi.org/10.1016/j.quascirev.2016.04.005>, 2016.

1025 San-Miguel-Ayanz, J., de Rigo, D., Caudullo, G., Houston Durrant, T., and Mauri, A.: European atlas of forest tree species.
 1026 Publication Office of the EU, Luxembourg, 2022.

1027 Sassoon, D., Lebreton, V., Combourieu-Nebout, N., Peyron, O., and Moncel, M.-H.: Palaeoenvironmental changes in the
 1028 southwestern Mediterranean (ODP site 976, Alboran sea) during the MIS 12/11 transition and the MIS 11 interglacial and
 1029 implications for hominin populations, *Quat. Sci. Rev.*, 304, 108010, <https://doi.org/10.1016/j.quascirev.2023.108010>, 2023.

1030 Sassoon, D., Combourieu-Nebout, N., Peyron, O., Bertini, A., Toti, F., Lebreton, V., and Moncel, M.-H.: Pollen-based climatic
 1031 reconstructions for the interglacial analogues of MIS 1 (MIS 19, 11, and 5) in the southwestern Mediterranean: insights from
 1032 ODP Site 976, *Clim. Past*, 21, 489–515, <https://doi.org/10.5194/cp-21-489-2025>, 2025.

1033 Schmid, P. e.: Random patch dynamics of larval Chironomidae (Diptera) in the bed sediments of a gravel stream, *Freshw.*
 1034 *Biol.*, 30, 239–255, <https://doi.org/10.1111/j.1365-2427.1993.tb00806.x>, 1993.

1035 Shumilovskikh, L. S., Shumilovskikh, E. S., Schlütz, F., and van Geel, B.: NPP-ID: Non-Pollen Palynomorph Image Database
 1036 as a research and educational platform, *Veg. Hist. Archaeobotany*, 31, 323–328, <https://doi.org/10.1007/s00334-021-00849-8>,
 1037 2022.

1038 Stockmarr, J.: 1971: Tablets with spores used in absolute pollen analysis, *Pollen et Spores* 13, 615-621, 1971.

1039 Stuchlik, L.: Atlas of pollen and spores of the Polish Neogene, W. Szafer Institute of Botany, Polish Academy of Sciences,
 1040 2001.

1041 Stuchlik, L.: Atlas of Pollen and Spores of the Polish Neogene: Gymnosperms, W. Szafer Institute of Botany, Polish Academy
 1042 of Sciences, 2002.

1043 Stuchlik, L.: Atlas of pollen and spores of the Polish Neogene. 3. Angiosperms (1), W. Szafer Inst. of Botany, Polish Academy
 1044 of Sciences, 2009.

1045 Szal, M., Kupryjanowicz, M., Wyczółkowski, M., and Tylmann, W.: The Iron Age in the Mrągowo Lake District, Masuria,
 1046 NE Poland: the Salet settlement microregion as an example of long-lasting human impact on vegetation, *Veg. Hist.*
 1047 *Archaeobotany*, 23, 419–437, <https://doi.org/10.1007/s00334-014-0465-z>, 2014.

1048 Szymanek, M.: Elemental geochemistry of freshwater snail shells: palaeolimnology of a Holsteinian (MIS 11) deposit from
 1049 eastern Poland, *Boreas*, 47, 643–655, <https://doi.org/10.1111/bor.12283>, 2018.

1050 Takagi, S., Kikuchi, E., Doi, H., and Shikano, S.: Swimming Behaviour of Chironomus acerbiphilus Larvae in Lake Katanuma,
 1051 *Hydrobiologia*, 548, 153–165, <https://doi.org/10.1007/s10750-005-5196-9>, 2005.

1052 Tarkowska-Kukuryk, M.: Spatial distribution of epiphytic chironomid larvae in a shallow macrophyte-dominated lake: effect
 1053 of macrophyte species and food resources, *Limnology*, 15, 141–153, <https://doi.org/10.1007/s10201-014-0425-4>, 2014.

1054 Tarr, T. L., Baber, M. J., and Babbitt, K. J.: Macroinvertebrate community structure across a wetland hydroperiod gradient in
 1055 southern New Hampshire, USA, *Wetl. Ecol. Manag.*, 13, 321–334, <https://doi.org/10.1007/s11273-004-7525-6>, 2005.

1056 Tokeshi, M.: Life cycles and population dynamics, in: *The Chironomidae: Biology and ecology of non-biting midges*, edited
 1057 by: Armitage, P. D., Cranston, P. S., and Pinder, L. C. V., Springer Netherlands, Dordrecht, 225–268,
 1058 https://doi.org/10.1007/978-94-011-0715-0_10, 1995.

1059 Tye, G. J., Sherriff, J., Candy, I., Coxon, P., Palmer, A., McClymont, E. L., and Schreve, D. C.: The $\delta^{18}\text{O}$ stratigraphy of the
 1060 Hoxnian lacustrine sequence at Marks Tey, Essex, UK: implications for the climatic structure of MIS 11 in Britain, *J. Quat.*
 1061 *Sci.*, 31, 75–92, <https://doi.org/10.1002/jqs.2840>, 2016.

1062 Tzedakis, P. C.: The MIS 11 – MIS 1 analogy, southern European vegetation, atmospheric methane and the “early
 1063 anthropogenic hypothesis,” *Clim. Past*, 6, 131–144, <https://doi.org/10.5194/cp-6-131-2010>, 2010.

1064 Tzedakis, P. C., Hooghiemstra, H., and Pälike, H.: The last 1.35 million years at Tenaghi Philippon: revised chronostratigraphy
 1065 and long-term vegetation trends, *Quat. Sci. Rev.*, 25, 3416–3430, <https://doi.org/10.1016/j.quascirev.2006.09.002>, 2006.

1066 Ustrnul, Z., Wypych, A., Marosz, M., Biernacik, D., Czekierda, D., Chodubska, A., Wasielewska, K., Kusek, K., and
 1067 Kopaczka, D.: Climate of Poland 2021, IMGW-PIB, n.d.

1068 Velle, G., Brooks, S. J., Birks, H. J. B., and Willassen, E.: Chironomids as a tool for inferring Holocene climate: an assessment
 1069 based on six sites in southern Scandinavia, *Quat. Sci. Rev.*, 24, 1429–1462, <https://doi.org/10.1016/j.quascirev.2004.10.010>,
 1070 2005.

1071 Vera-Polo, P., Sadori, L., Jiménez-Moreno, G., Masi, A., Giaccio, B., Zanchetta, G., Tzedakis, P. C., and Wagner, B.: Climate,
 1072 vegetation, and environmental change during the MIS 12-MIS 11 glacial-interglacial transition inferred from a high-resolution
 1073 pollen record from the Fucino Basin of central Italy, *Palaeogeogr. Palaeoclimatol. Palaeoecol.*, 655, 112486,
 1074 <https://doi.org/10.1016/j.palaeo.2024.112486>, 2024.

1075 de Vernal, A. and Hillaire-Marcel, C.: Natural Variability of Greenland Climate, Vegetation, and Ice Volume During the Past
 1076 Million Years, *Science*, 320, 1622–1625, <https://doi.org/10.1126/science.1153929>, 2008.

1077 Walker, I. R. and Mathewes, R. W.: LATE-QUATERNARY FOSSIL CHIRONOMIDAE (DIPTERA) FROM HIPPA LAKE,
 1078 QUEEN CHARLOTTE ISLANDS, BRITISH COLUMBIA, WITH SPECIAL REFERENCE TO CORYNOCERA ZETT.,
 1079 *Can. Entomol.*, 120, 739–751, <https://doi.org/10.4039/Ent120739-8>, 1988.

1080 Walker, I. R. and Mathewes, R. W.: Chironomidae (Diptera) remains in surficial lake sediments from the Canadian Cordillera:
 1081 analysis of the fauna across an altitudinal gradient, *J. Paleolimnol.*, 2, 61–80, <https://doi.org/10.1007/BF00156985>, 1989.

1082 Walker, I. R., Fernando, C. H., and Paterson, C. G.: The chironomid fauna of four shallow, humic lakes and their representation
 1083 by subfossil assemblages in the surficial sediments, *Hydrobiologia*, 112, 61–67, <https://doi.org/10.1007/BF00007667>, 1984.

1084 Wiederholm, T.: Chironomidae of the holarctic region. Keys and diagnoses. Part 1 : larva, *Ent Scand Suppl*, 19, 1–457, 1983.

1085 Willerslev, E., Cappellini, E., Boomsma, W., Nielsen, R., Hebsgaard, M. B., Brand, T. B., Hofreiter, M., Bunce, M., Poinar,
 1086 H. N., Dahl-Jensen, D., Johnsen, S., Steffensen, J. P., Bennike, O., Schwenninger, J.-L., Nathan, R., Armitage, S., de Hoog,
 1087 C.-J., Alfimov, V., Christl, M., Beer, J., Muscheler, R., Barker, J., Sharp, M., Penkman, K. E. H., Haile, J., Taberlet, P., Gilbert,
 1088 M. T. P., Casoli, A., Campani, E., and Collins, M. J.: Ancient Biomolecules from Deep Ice Cores Reveal a Forested Southern
 1089 Greenland, *Science*, 317, 111–114, <https://doi.org/10.1126/science.1141758>, 2007.

1090 Wörner, S. and Pester, M.: The Active Sulfate-Reducing Microbial Community in Littoral Sediment of Oligotrophic Lake
 1091 Constance, *Front. Microbiol.*, 10, <https://doi.org/10.3389/fmicb.2019.00247>, 2019.

1092 Yin, Q. Z. and Berger, A.: Individual contribution of insolation and CO₂ to the interglacial climates of the past 800,000 years,
 1093 *Clim. Dyn.*, 38, 709–724, <https://doi.org/10.1007/s00382-011-1013-5>, 2012.

1094 Żarski, M., Nita, M., and Winter, H.: Nowe stanowiska interglacjalne w rejonie dolin Wilgi i Okrzejki na Wysoczyźnie
 1095 Żelechowskiej (Polska południowo-wschodnia), *Przegląd Geol.*, 53, 137–144, 2005.

1096 Żarski, M., Hrynowiecka, A., Górecki, A., Winter, H., and Pochocka-Szwarc, K.: The maximum extent of the Odranian
 1097 Glaciation (Saalian, MIS 6) in the South Podlasie Lowland (SE Poland) in the light of sites with lacustrine deposits of the
 1098 Mazovian Interglacial, *Acta Geol. Pol.*, e14–e14, <https://doi.org/10.24425/agp.2024.150006>, 2024.

1099