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1 Late Holocene ecological shifts and chironomid-inferred summer temperature

2 changes reconstructed from Lake Uddelermeer, the Netherlands

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13 Abstract

14 This paper presents a Late-Holocene chironomid-inferred July-air temperature record from a core 15 obtained from Lake Uddelermeer in the Netherlands. A core interval, which dates from 2500 to 400 cal. yr. BP, 16 was analysed at multi-decadal resolution for organic content, pollen, spores and NPPs (Non Pollen 17 Palynomorphs), and chironomid head capsules. These proxies indicate that, from 2500 to 1140 cal. yr. BP, the 18 lake was mesotrophic and sustained a Littorellion, while the chironomid assemblage was dominated by littoral 19 species associated with macrophytes. At 1140 cal. yr. BP a shift in the lake ecology occurred from low-nutrient to 20 high-nutrient conditions dominated by algae. This shift might be linked to a concurrent increase in human impact 21 and is reflected in the chironomid assemblage by increases in eurytopic taxa, which are resistant to disturbances. 22 Shifts in the chironomid record between 2500 and 1140 cal. yr. BP do not coincide with changes in lake ecology 23 and are presumably driven by climate change. Using a Norwegian-Swiss calibration dataset as a modern 24 analogue, we produced a chironomid-inferred temperature (C-IT) reconstruction. This reconstruction compares 25 well to other regional temperature reconstructions in timing and duration with a Roman Warm period between 26 2240-1760 cal. yr. BP, a Dark Age Cold Period starting at 1760 cal. yr. BP and the Medieval Climate Anomaly 27 beginning at 1280 cal. yr. BP. The C-IT record indicates a temperature drop of 1.5°C from the Roman Warm 28 Period to the Dark Age Cold Period. Findings improve knowledge of the first millennium AD in NW Europe, which 29 was characterised by changes in landscape, vegetation, society and climate.

30

31 Keywords

first millennium AD; non-biting midges; July air temperature; Dark Age Cold Period; Roman Warm Period; North western Europe

34

35 **1** Introduction

Many studies have shown that past climate fluctuations might be temporally linked to cultural shifts (Tinner *et al.* 2003; Büntgen *et al.* 2011; McCormick *et al.* 2012a; Büntgen *et al.* 2016). During the first millennium AD two large cultural transitions occurred in NW Europe: the decline of the Roman Empire and the later transition to medieval kingdoms. Recent studies (e.g. Büntgen *et al.* 2016b; Helama *et al.* 2017a; Helama *et al.* 2017b) have shown that climate in NW Europe during the first millennium was not as stable as previously assumed or as 41 evidenced from global climate reconstructions (Rasmussen *et al.* 2006; Vinther *et al.* 2006; Wanner *et al.* 2008).

- 42 Riechelmann and Gouw-Bouman (2019) indicate the presence of a Roman Warm Period (RWP; 1950-1700 cal.
- 43 yr. BP), a Dark Age Cold Period (DACP; 1700-1250 cal. yr. BP) and a Medieval Warm Period or Medieval Climate
- 44 Optimum (MCA; 1250-950 cal. yr. BP) in NW Europe (table 1). Other climate reconstructions from the Northern
- 45 Hemisphere also demonstrate the occurrence of significant climatic shifts during this time interval (e.g. Larsen *et*
- 46 *al.* 2008; Ljungqvist 2010; Wanner *et al.* 2011; Gräslund and Price 2012; McCormick *et al.* 2012b; Büntgen *et al.*
- 47 2016b; Toohey *et al.* 2016; Helama *et al.* 2017a). The identification of similar warm and cold periods in numerous
- proxy records throughout NW Europe indicates a regional climate trend (Helama *et al.* 2017a; Riechelmann and
 Gouw-Bouman 2019).

50 Following the collapse of the Roman Empire a decrease in population density, from approximately 1680-51 1450 cal. yr. BP (AD 270 to AD 500), was seen in the Netherlands, this time period is also known as the Migration 52 Period. This was followed by an exponential population growth from 1350 cal. yr. BP (AD 600) onward when the 53 medieval kingdoms emerged (Pierik and van Lanen 2017; Groenewoudt and van Lanen 2018; Pierik et al. 2018) 54 (table 1). Not only do these cultural shifts occur contemporaneously with climatic changes, they also are 55 concurrent with large-scale landscape and vegetation changes in the Netherlands (Jansma et al. 2014; Pierik 2017). In the Rhine-Meuse delta flooding frequency and intensity increased between 1700 cal. yr. BP and 1100 56 57 cal. yr. BP and multiple avulsions are placed within this time frame (Stouthamer and Berendsen 2000; Erkens 58 2009; Toonen 2013; Cohen et al. 2016; Pierik et al. 2017). In the Dutch coastal area increased storm surges are 59 observed from 1350 cal. yr. BP onward (Knol 1993; Vos and Van Heeringen 1997; Pierik et al. 2016). 60 Contrastingly, a decrease in aeolian activity is visible during the Roman period (1950-1550 cal. yr. BP) in the 61 Pleistocene cover-sand areas and an increase in driftsand activity took place at the end of the first millennium 62 around 1050 cal. yr. BP (Pierik et al. 2018). Vegetation development during the first millennium AD is characterised by a reforestation phase between 1850 cal. yr. BP and 1250 cal. yr. BP. Although timing and 63 64 magnitude vary, this reforestation trend is recognised at most palynological study sites in the Netherlands and 65 neighbouring Germany (e.g. Teunissen 1990; Bunnik 1999; Kalis et al. 2008; Litt et al. 2009; Dörfler et al. 2012). 66 Although all climate, landscape, vegetation and population changes during the first millennium AD are 67 concurrent and some relation among them seems indisputable, the exact influence of each of these factors 68 remains unknown. It is therefore essential to obtain detailed and well-dated evidence of these changes to 69 disentangle the cause and effect of human impact, climate, landscape and vegetation change during the first 70 millennium AD.

Yet, detailed climate data from the Netherlands for the first millennium AD are at present not available. This is probably due to the scarcity of archives suitable to study for this important time interval. Pristine natural archives such as peat bogs and lake fills are often no longer present due to peat excavation, artificial drainage and other human disturbances. Lake Uddelermeer, located in a cover-sand area in the central part of the Netherlands, is a unique site as it contains an undisturbed sediment profile covering the last ca. 14,000 years (Engels *et al.* 2016).

77 Chironomids, or non-biting midges, are an excellent proxy for the reconstruction of past summer 78 temperatures because they have a short life cycle and are sensitive to changes in temperature (Brooks et al. 79 2007; Telford and Birks 2011; van Asch 2012; Heiri et al. 2014). They have mainly been used to study Lateglacial 80 temperature variability but have been proven to be good proxy for Holocene temperatures as well (Axford et al. 81 2009; Millet et al. 2009; Brooks et al. 2012; Nazarova et al. 2013). However, as a result of human interference in 82 the environment, chironomid-inferred temperature reconstructions for the Late Holocene can be problematic. 83 Even small-scale human impact such as Prehistoric farming has been shown to impact chironomid communities 84 and distort the climate signal (Taylor et al. 2013; Taylor et al. 2017a; Taylor et al. 2017b). Therefore, a multi proxy 85 approach is recommended as a method for Holocene climate reconstructions, to infer possible human impact on

the record, and to disentangle the effects of natural processes such as climate change from anthropogenic

87 impacts on the landscape (McKeown and Potito 2016; Taylor *et al.* 2018).

The aims of this study are fourfold: (1) to present the first chironomid record from the Netherlands
covering the first millennium AD on a multi-decadal scale, (2) to determine, using the sedimentological,
palynological and chironomid records, to what extent human impact influenced the lake ecology and whether

91 temperature is the main environmental driver of changes in the chironomid assemblage, (3) to assess whether

92 absolute chironomid-inferred summer-temperature values are realistic and, (4) to assess whether the chironomid-

93 inferred temperature reconstruction shows evidence for temperature change during known phases of changing94 climate.

95

96 2 Area description

97 Lake Uddelermeer is a pingo remnant located on the Veluwe (52°14'48"N; 5°45'40"E; 24-27 m a.s.l.)
98 (Fig. 1). The pingo remnant is situated at the head of a periglacial valley, in the cover-sand area between the ice
99 pushed ridges of Ermelo (max. elevation ca. 50 m a.s.l.) and Apeldoorn (max. elevation ca. 100 m a.s.l.) (Fig. 1)
100 and in the subsurface an impermeable clay layer is present. These boundary conditions are typical for the
101 occurrence of pingos in the Netherlands due to the presence of a ground water gradient and/or pressure. This is
102 evidenced by the presence of another pingo remnant in this area: Lake Bleekemeer, which does not contain a
103 complete Late Holocene lake infill due to peat cutting (Polak 1959).

Lake Uddelermeer is oval shaped, 200-300 metres in diameter and has a current maximum water depth of 1.3 104 105 metres (Engels et al. 2016). The total lake sediment infill reaches a maximum depth of 15.6 metres of which 12.9 106 metres covers the Holocene. The lake is fed by groundwater and a small brook is draining the south-side of the 107 lake (Staverdense or Leuvenumse beek). The area directly surrounding the lake is slightly elevated, limiting inflow 108 of surface water runoff into the lake (Fig. 1). The lake is currently eutrophic and directly surrounding the lake is an 109 alder carr and a fringe of wetlands with reeds. The regional vegetation consists mainly of mixed oak and birch 110 forest, grasslands and agricultural fields. Average July temperature for the period 1901-2017 in de Bilt (ca. 50 km SW of Lake Uddelermeer) is 16.95 °C (1901-1950 = 16.4 °C) (Royal Netherlands Meteorological Institute; 111 112 www.knmi.nl)

113 Archaeological finds in the vicinity of the lake indicate the presence of human activities from the Middle 114 Mesolithic onward (Groenewoudt et al. 2006). Neolithic (7250-3950 cal. yr. BP; 5300-2000 BC) burial mounds 115 from the Trechterbeker culture (5350-4850 cal. yr. BP; 3400-2900 BC) are present at a distance of ca. 500 metres 116 from the lake. There is also evidence of charcoal production in the direct surroundings of the lake during the 9th and 10th centuries (Kraanen and Pape 1965). Situated next to the lake is a medieval structure dating to the late 117 118 10th century (Heidinga 1987). This so-called Huneschans is a medieval ring fort (ringwalburg) which is 119 approximately 100 metres in diameter, and is surrounded by a circular earthen wall and an outer ditch. The 120 earthen wall is not fully circular, with an opening present on the side of the lake. The wall currently does not reach 121 the lake itself, indicating that the water level of the lake was up to 1.7 metres higher during the time of 122 construction of the fort (10th century; Engels et al. 2016). Earlier palynological studies by Engels et al. (2016), 123 Bohncke (1999), Sohl (1983) and Polak (1959) show the first evidence of human activities at the beginning of the 124 Subboreal at 5000 cal. yr. BP, which is concurrent with the construction of the burial mounds encountered in the 125 area (Groenewoudt et al. 2006). Pollen records covering this period show an increase of Corylus and Pteridium 126 indicating open forests as well as the occurrence of human indicators such as Plantago and Cerealia. From 4000 127 cal. yr. BP onward the human-induced deforestation trend, which initiated around 5000 cal. yr. BP, increased.

- Human presence in the form of agriculture is continuous from 3450 cal. yr. BP onwards as indicated by a continuous Cerealia pollen curve (Engels *et al.* 2016).
- 130

131 Figure 1 around here

132

133 **3 Methods**

134 A 16-metre-long sediment core (ø 6 cm) was collected from the deepest part of the lake in April-May 135 2012 using a piston corer operated from an UWITEC coring platform. The lake infill was cored in 3 metre sections using overlapping cores, and coring tubes were subsequently cut to 1 metre intervals in the field to enable 136 137 transport. In the lab the core segments were cut in half lengthwise, where one part was completely preserved and 138 the other half was used for sampling. A continuous 1 cm sampling was carried out for Loss on Ignition (LOI) to 139 determine organic content following the protocol by Heiri et al. (2001). A total of 437 samples of ca. 1 cm³ were 140 dried at 105° C for 12 hours and subsequently placed in an oven at 550° for 4 hours. The samples were weighed 141 before and after each step, obtaining both the moisture as well as the organic carbon content. Pollen samples of 142 0.7 cm³ were collected using a 5 cm sampling resolution for the section 100-500 cm below the sediment/water 143 interface to obtain a high-resolution vegetation record capable of detecting the fast vegetation changes during the first millennium AD in detail. In total 70 pollen samples were prepared following the method described in Faegri 144 145 and Iversen (1989) and Moore et al. (1991). Identification of pollen, spores and Non Pollen Palynomorphs (NPPs) 146 was executed using a Zeiss microscope with 400x magnification and pollen and spores were identified using 147 Beug (2004), Moore et al. (1991) and Punt (1976; 1980; 1981; 1984; 1988; 1991; 1995; 2003). NPPs such as 148 algal remains and (fungal) spores were identified using van Geel et al. (2003), van Geel (1978) and van Geel 149 (1972). An upland pollen sum of 300 pollen grains, excluding alder and grasses, was counted per sample. This 150 pollen sum differs from the one used in Engels et al. (2016), as grasses and alder pollen are excluded from the 151 pollen sum in this study since reed land (included in the Poaceae curve) and alder carr occur locally.

152 The sediment record was previously analysed for chironomids at a coarse temporal resolution by Engels 153 et al. (2016). For the current research, additional chironomid samples were analysed to increase the temporal 154 resolution from the section 200-500 cm below the sediment surface to detect the short climatic phases during the 155 first millennium AD. From this section, a total of 31 samples were analysed at 5-25 cm resolution. In order to get a 156 better overview of preceding and subsequent trends, the lower temporal resolution results from Engels et al. 157 (2016) from the sections 100-200 cm and 500-550 cm are also included in this study. Sample preparation for 158 these samples is described in Engels et al. (2016). The additional samples (14 in total, sample size: 0.335 cm³) were added to a 5% KOH solution, heated to 90°C for 1 hour, and then passed successively through 212 µm and 159 160 90 µm mesh sieves. Chironomid head capsules were manually picked out of the sieving residues using a Bogorov 161 sorting tray under a dissection microscope (40x magnification) and subsequently mounted on permanent slides in 162 Euparal©. Chironomid head capsules were identified under a compound microscope (magnification 400x) 163 following keys by Wiederholm (1983), Moller Pillot (1984a); Moller Pillot (1984b), Rieradevall and Brooks (2001) 164 and Brooks et al. (2007). Information on chironomid ecology was derived from a.o: Moller Pillot and Buskens 165 (1990), Brooks et al. (2007), Moller Pillot and Klink (2009), Engels and Cwynar (2011), Luoto (2011), Engels et al. 166 (2012), Moller Pillot (2013) and Potito et al. (2014). The number of identified chironomid head capsules varied 167 between 65 and 146 per sample. A Principal Component Analysis (PCA) with Square-root-transformed 168 percentage-abundance data of the chironomid taxa was executed using the program C2 (Juggins, 2003). 169 The chironomid assemblages were subsequently used to quantitatively reconstruct mean July air 170 temperatures. This reconstruction was executed using a chironomid-temperature inference model, which is based 171 on a modern calibration dataset that consists of 274 lakes from Norway and the Alpine regions (Heiri et al. 2011).

172 This chironomid-climate calibration dataset spans a July air temperature range from 3.5 to 18.4°C (Heiri *et al.*

2011). For the model, a two component weighted averaging partial least-squares (WA-PLS) regression was used
since this approach produced inference models with the lowest error (Heiri *et al.* 2011). The selected model had a
bootstrapped root mean square error of prediction of 1.40°C after outlier deletion (Heiri *et al.* 2011). Bootstrapping
was used to estimate the sample-specific errors for the fossil samples.

177 We evaluated the robustness of the chironomid-inferred July air temperature reconstruction by 178 calculating the closest modern analogue of the fossil samples, as well as the goodness-of-fit to temperature (Birks 179 et al. 1990) and the cumulative percentage of rare or absent species from the calibration dataset. The closest 180 modern analogue was assessed based on squared Chi-squared distances using the program C2 (Juggins, 2003). Fossil samples with a distance to the closest analogue larger than the 2nd and 5th percentile of the distances of the 181 modern samples in the calibration dataset were classified as having 'no close' and 'no good' analogue, 182 183 respectively. Goodness-of-fit to temperature was assessed using a Canonical Correspondence Analysis (CCA) of 184 the modern samples with temperature as the only constraining variable. In this analysis, the fossil samples were 185 added passively. Fossil samples with a residual distance exceeding the 90th and 95th percentiles of the residual distances of the modern samples were identified as having a 'poor' and 'very' poor fit with temperature, 186 respectively. The CCA was calculated using CANOCO for Windows version 4.51 (ter Braak and Šmilauer 2012). 187 188 Both the pollen percentage diagram as well as the chironomid abundance diagram where produced using TILIA 189 (Grimm 1991-2015). Zonation of the pollen and chironomid diagrams differs from Engels et al. (2016) as a result 190 of the increased sampling resolution in this study and was established visually. The pollen diagram was 191 subdivided into zones based on variations in the main pollen taxa and zonal boundaries were verified using 192 CONISS (Grimm 1991-2015). The chronology of the core is based on the age-depth model of Engels et al. (2016), which is based on 26 samples for ²¹⁰Pb measurement for the top 65 cm of the core and 13 AMS 193 194 radiocarbon samples (excluding 3 outliers) throughout the rest of the core. The age-depth model was produced 195 using Bayesian modelling as included in Oxcal (Bronk Ramsey 2009). The radiocarbon samples contained 196 terrestrial macrofossils, wood fragments and charred material such as grass epidermis and charcoal (Engels et al. 197 2016). The age-depth chronology of the core section discussed in this paper (100 to 550 cm sediment depth) is 198 supported by six radiocarbon dates (excluding 1 outlier) and constrained at the top by the ²¹⁰Pb dates (Fig. 2). 199 The resultant age-depth model gives well-constrained age estimates and a nearly linear sedimentation rate, 200 confirming the reliability of the six dates included. All ages provided in this paper are rounded to the nearest five 201 year interval and are expressed in cal. yr. BP unless explicitly stated otherwise.

202

203 Figure 2 around here

204

205 4 Results and interpretation

206 4.1 Sedimentology

The investigated core interval consists entirely of gyttja (Fig. 3). The bottom part, 550 cm up to 220 cm 207 208 depth, consists of brown gyttja. From 220 cm upward this slowly shifts to a more greenish gyttja, indicating a 209 larger component of algae. The LOI-record follows this trend with values around 60% from 550 cm up to 460 cm 210 depth, followed by a more organic-rich sequence with values upward of 70%, until 270 cm depth. From 270 cm to 211 220 cm LOI values vary around 60% again and drop sharply to only 20% at 220 cm, which corresponds with the 212 shift to a higher algal content. The drop in organic content is probably the result of increased silica production 213 (a.o. diatom production) in the lake. Alternatively, an increase in sand input through increased aeolian activity 214 could explain this decrease. However, no visible sand grains were found during visual core inspection, suggesting

215 that an increase of silica production is the most likely reason for the observed decrease in LOI values. From 220 216 cm upwards, LOI values gradually increase, reaching 60% at 140 cm after which they gradually decrease to 30%.

217

- 218 Figure 3 around here
- 219

220 4.2 Palynology

221 Zone Pa (500-382 cm depth; 2420-1730 cal. yr. BP) is characterised by high values of Arboreal Pollen 222 (AP) and heather (Fig. 3). High values of deciduous tree species such as Quercus, Fagus and the occurrence of 223 Carpinus, Tilia and Ulmus suggest the presence of a mixed-deciduous forest in the vicinity of the lake. Fraxinus, 224 Corylus and Betula probably grew at the forest edge or in small bushes and thickets. Human presence in the area 225 around Lake Uddelermeer is indicated by high percentages of upland herbs with typical anthropogenic indicators 226 such as Rumex, Plantago lanceolata and Artemisia vulgaris and the presence of pollen grains of Cerealia, 227 indicating arable fields nearby. Heathlands are also present in the area, as evidenced from the large proportion of 228 Calluna. The lake was probably surrounded by an alder carr with grasslands in the vicinity and alder carr was 229 likely also present in the brook valley of the Leuvenumse beek to the south. The high values of both *Isoëtes* and 230 Littorella suggest relatively nutrient-poor and clear-water conditions in the lake itself. Isoëtes and Littorella are both adapted to low nutrient conditions, by a symbiosis with mycorrhiza to enable carbon uptake from the 231 232 sediment, and a slow growth rate, and are therefore typical inhabitants of oligotrophic lakes, where they most 233 often grow in water of 0 to 2 metres deep (Sand-Jensen 1978; Weeda et al. 1988; Engels et al. 2018). Isoëtes 234 can grow in water up to 4.5 metres deep and prefers a sandy subsoil (Sand-Jensen 1978; Farmer and Spence 235 1986). Most likely Littorella also grew at the shore of the lake, since this species only flowers when it is not 236 submerged (Weeda et al. 1988). Together, both species form the Littorellion which is a relatively stress tolerant 237 plant community (Rørslett and Brettum 1989). However, when nutrient availability increases, macrophytes and 238 algae can more easily inhabit the lake. As a result Littorella and Isoëtes receive less light and the Littorellion is 239 fast replaced by reedlands (Weeda et al. 2000). The nutrient-poor conditions in the lake are confirmed by the presence of Nymphaea alba which probably grew in the deeper parts of the lake. The algae Botryococcus, which 240 241 was present in Lake Uddelermeer continuously since the Late Glacial (Engels et al., 2016), can occur in a wide 242 range of environments but has a preference for oligotrophic lakes (Padisák et al. 1998). Engels et al. (2016) 243 reconstructed a water depth of around 2.5-3 metres during this time period, which is in line with the 2 metres 244 suggested by the aquatic taxa present in the lake. Spores of Glomus, a fungus often found in soils, is present in 245 the upper part of this zone. The occurrence of Glomus spores in lake sediments is indicative of soil erosion (van 246 Geel et al. 2003).

247 Zone Pb (382-255 cm depth; 1730-1270 cal. yr. BP) is characterised by increasing values of AP and 248 decreasing values of heather and to a lesser extent upland herbs (Fig. 3). The increase in AP can be mainly attributed to Quercus, Fagus and Carpinus. Anthropogenic indicators such as upland herbs (Rumex, Plantago 249 250 lanceolata and Artemisia vulgaris) and Cerealia decrease in this zone. Pollen grains of Cerealia are still present, 251 albeit in low abundances, and occasionally Secale cereale is found. These changes in the pollen assemblage 252 suggest that deciduous forests have expanded at the expense of heathlands and arable lands. We also observe 253 an increase of Alnus and a decrease of Poaceae suggesting that the local alder carrs are also expanding. The 254 basal cells of Nymphaeaceae (NPP type HdV-127) are decreasing sharply during this zone, although Nymphaea 255 alba is still present in the pollen record. Nuphar lutea, however, is nearly absent from zone Pb. The continued 256 presence of Isoëtes and Littorella suggests continued nutrient-poor and clear-water conditions in the lake. Pollen-257 percentage values of Littorella however, are slightly decreasing. The occurrence of Typha latifolia pollen during 258 this zone indicates the presence of reed lands at the lake fringe which possibly partly replaced the Littorellion.

- Zone Pc (255-220 cm depth; 1270-1140 cal. yr. BP) is a transitional zone which shows decreasing AP 259 260 values and increasing values of Cerealia and heather (Fig. 3). A mixed deciduous forest is still present but arable 261 lands and heathlands are fast expanding. This is also visible from the increase in anthropogenic indicators such 262 as Cerealia, Plantago lanceolata, a continuous curve of Secale cereale pollen, and an increase in the 263 occurrences of charcoal. The alder carr decreases in size and becomes more open with more grass or reed land, 264 as seen from the decreasing abundance of Alnus and increasing Poaceae values and the presence of Typha 265 angustifolia. Isoëtes and Littorella are still present in the pollen record indicating relatively stable conditions in the 266 lake through zone Pa, Pb and Pc. The water body must have remained clear enough throughout pollen zones Pa, 267 Pb and Pc to sustain the Littorellion.
- 268 Zone Pd (220-100 cm depth; 1140-320 cal. yr. BP) is characterised by further decreasing values of AP 269 and heather and increasing values of upland herbs and Cerealia (Fig. 3). The start of this zone is simultaneous 270 with the shift to green algal gyttja indicating a higher production in the lake. This is also evident from the rapidly 271 increasing values of algal remains such as Scenedesmus or Pediastrum. An increase in various algal species 272 from ca. 25% to over 100% occurs at 185 cm depth (1030 cal. yr. BP). In our record the abrupt increase in algal 273 remains coincides with increasing values of fungal spores of coprophilous fungi such as Sporormiella, Podospora 274 and Cercophora which are indicative for the presence of dung. The increase in algae also coincides with the 275 habitation phase of the Huneschans (late 10th cent.) and is most likely caused by increased nutrient input as a 276 result of human impact. Engels et al. (2018) however, found sediment age and water depth to be the most likely 277 explanatory variables for this ecosystem change. From 155 cm depth (1250-700 cal. yr. BP) onward Cannabis-278 type shows an increase from ca. 1% to ca. 35% which is likely the result of the retting of hemp in the lake to create hemp fibres for rope production. The practice of hemp retting in the Netherlands is known from the historic 279 280 record and also evidenced at Lake Uddelermeer from the high values of Chaetomium spores which are cellulose-281 decomposers and occur among others, on decaying herbaceous stems (van Geel et al. 2003).
- 282 Overall zone Pd indicates an increase in human activity, with higher values of Cerealia, Secale, Fagopyrum and 283 Sinapis-type and various herbs such as Rumex and Amaranthaceae. The presence of charcoal also increases 284 during this zone, which might be linked to the charcoal production in the area. There are fewer deciduous tree 285 species present indicating the disappearance of the closed and mixed deciduous forest, which are being replaced 286 by more open oak forest with hazel and ash. Arable lands are expanding while the heathlands are decreasing. 287 The alder carr is also disappearing and is being replaced by grassland. In the lake itself the eutrophic conditions 288 are evident from the disappearance of Littorella, Isoëtes and Nymphaea and the concurrent increase of Myriophyllum alterniflorum. The occurrence of this aquatic perennial is often linked to an increase in phosphate 289 290 (Roelofs et al. 1984). Myriophyllum alterniflorum grows in water up to 2 metres deep (Hannon and Gaillard 1997) 291 suggesting that water depth remained stable during the entire period, in line with results by Engels et al. (2016).
- 292

293 Figure 4 around here

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295 4.3 Chironomid record

Chironomids are well preserved and abundant throughout the record. Our record has an average count sum of 99 head capsules per sample (range 64-167). Most of the encountered chironomid taxa (e.g. *Lauterborniella, Parakiefferialla bathophila*-type) are commonly found in the littoral zone of meso- to eutrophic lakes and prefer warm or temperate conditions (Brooks *et al.* 2007, and references therein). Further, a number of taxa present in the record are often associated with macrophytes (e.g. *Ablabesmyia*). The chironomid diagram is divided into two zones Ch1 and Ch2, following major shifts in the chironomid assemblage, which correspond with the major lithological units in the core (Fig. 4). Chironomid zone Ch1 (550-220 cm; 2525-1140 cal. yr. BP) is dominated by *Lauterborniella*, Cladotanytarsus mancus-type and Parakiefferiella bathophila-type which are all typical inhabitants of littoral zones of lakes. In addition, *Lauterborniella* and *P. bathophila*-type are often associated with submerged vegetation (Moller Pillot 1984b; Brooks *et al.* 2007; Moller Pillot and Klink 2009; Moller Pillot 2013). *Lauterborniella* is found in the warmer lakes included in the Norwegian-Swiss calibration dataset (Heiri *et al.* 2011). Chironomid zone Ch1 is further subdivided based on smaller changes in the abundance of the chironomid taxa.

309 Subzone Ch1a (550-470 cm depth; 2525-2240 cal. yr. BP) is marked by relatively high values of 310 Endochironomus albipennis-type and Pseudochironomus and the presence of Limnophyes/Paralimnophyes and 311 Procladius. Both Pseudochironomus and E. albipennis-type occur in the littoral of lakes (Brooks et al. 2007; 312 Engels et al. 2012). E. albipennis-type is often associated with macrophytes (Moller Pillot 1984b) suggesting the 313 presence of aquatic vegetation in the lake. This is in line with the pollen record, which indicates the presence of a 314 Littorellion and Nymphaeaceae. Based on the presence or Littorella and Isoëtes, relatively nutrient-poor 315 conditions are expected in the lake. However chironomid taxa such as P. bathophila-type and Polypedilum 316 nubeculosum-type are typically found in eutrophic lakes (Moller Pillot 2013). Lauterborniella is often ascribed to 317 eutrophic conditions (Moller Pillot 2013) but this genus has been found in oligotrophic lakes as well (Brodin 1986). 318 Based on the combination of the presence of aquatic macrophytes that typically occur in oligo- to mesotrophic 319 conditions and chironomid taxa that typically occur in meso- to eutrophic conditions we conclude that the lake was 320 probably mesotrophic throughout zone Ch1. This conclusion is further supported by the occurrence of taxa such 321 as Pseudochironomus and Tanytarsus mendax-type, which are inhabitants of mostly mesotrophic lakes (Moller 322 Pillot 2013). Further, Limnophyes/Paralimnophyes, Lauterborniella, P. nubeculosum and Pseudochironomus are 323 most often found in shallow lakes (Brooks et al. 2007; Engels et al. 2012), which is in line with the presence of the 324 Littorellion and the estimated water depth of 2.5 to 3 metres by Engels et al (2016). C. mancus-type and P. 325 bathophila-type have a preference of lakes with a water depth less than 5 metres (Luoto 2011). The majority of species present in this zone such as E. albipennis, Procladius, C. mancus-type are acidophilic which is also in line 326 327 with the reconstructed nutrient-poor, clear-water conditions in the lake. In the calibration dataset, Procladius and 328 Limnophyes/Paralimnophyes are both found along a large temperature gradient, including colder lakes, resulting 329 in relatively low temperature optima for these taxa (Heiri et al. 2011).

330 The onset of subzone CH1b (470-390 cm depth; 2240-1760 cal. yr. BP) is defined by the appearance of 331 Parachironomus varus-type and an increase of Corynoneura edwardsi-type and further characterized by the 332 absence of Limnophyes/Paralimnophyes and Procladius. P. varus-type and C. edwardsi-type are both often 333 associated with macrophytes and have relatively high temperature optima in the Norwegian-Alpine calibration 334 dataset (Heiri et al. 2011) (Brooks et al. 2007). At the start of this subzone, E. albipennis-type reaches maximum 335 values, directly followed by a strong decline from 19.2 % to 1.4%. Lauterborniella also reaches maximum values 336 in this subzone. Pseudochironomus is still present, although at slightly lower abundances than in the previous 337 subzone. P. varus-type is acidophobic (Brooks et al. 2007) suggesting less acidic conditions which is in line with 338 the drop in Procladius and E. albipennis-type which are both acidophilic. Nevertheless, other acidophilic species 339 such as C. mancus-type, Psectrocladius sordidellus-type are still present. We therefore conclude that the shift in 340 the chironomid fauna associated with the Ch1a/Ch1b-transition does not reflect any significant changes in the 341 lake ecosystem but rather a reorganization of the fauna already present in the lake.

Subzone Ch1c (390-255 cm depth; 1760-1270 cal. yr. BP) is defined by maximum values of *P*. *bathophila*-type (35.7%) and is further marked by the reappearance of *Limnophyes/Paralimnophyes* and *Procladius* and an increase in *P. sordidellus*-type and *P. nubeculosum*-type. *Lauterborniella* gradually declines
from its maximum abundance of 35.2% to values of 15.3% during subzone Ch1c. *Pseudochironomus* and *P. varus*-type also reach lower abundances than in the previous subzone. Both *P. nubeculosum*-type and *P. sordidellus*-type typically occur in the littoral zone of temperate lakes (Engels *et al.* 2012; Luoto 2012), although

according to Moller Pillot (1984b) *P. nubeculosum*-type is found in lakes with a depth up to 18 m. The increase in *P. nubeculosum*-type, *P. bathophila*-type might suggest more nutrient-rich conditions in the lake. However,
oligotrophic and mesotrophic taxa such as *Heterotanytarsus* and *Pseudochironomus* are still present indicating
that conditions remained relatively nutrient poor.

352 The final subzone (Ch1d: 255-220 cm depth; 1270-1140 cal. yr. BP) is marked by higher values of E. 353 albipennis-type, while abundances of Lauterborniella and P. bathophila-type are lower than in the preceding 354 subzones. Taxa that appear or increase during this zone are Glyptotendipes pallens-type and Polypedilum 355 sordens-type which are both found in the warmer lakes in the training set. The most distinct feature of this 356 subzone, however, is the high abundance of T. mendax-type. This taxon consists of a large number of species 357 and is found in a range of environments (Brooks et al. 2007). The taxon also occurs along a wide temperature 358 range, but is absent from the coldest lakes (Heiri et al. 2011). T. mendax-type is sensitive to increases in 359 phosphate but can occur in all trophic states and prefers mesotrophic conditions. This species mostly lives in the 360 profundal zone and is most abundant in water over 5 metres deep. E. albipennis-type and P. bathophila-type have 361 similar nutrient preferences although P. bathophila-type is indicative for warmer conditions. The shifts that we observe in the chironomid fauna during zone Ch1 do not coincide with shifts in ecology of the lake as 362 reconstructed from the pollen and LOI records. We therefore suggest that the changes observed in the 363 chironomid record were potentially driven by external factors such as climate change rather than by changes in 364 365 within-lake processes.

366 The onset of chironomid zone Ch2 coincides with the transition of brown gyttja to green algal gyttja (220-367 100 cm depth; 1140-320 cal. yr. BP). Zone Ch2 is marked by increases or appearances of the chironomid taxa 368 Microtendipes pedellus-type, Cladopelma lateralis-type, Cricotopus laricomalis-type, Nanocladius branchicolis-369 type and the Tanypodinae Procladius and Ablabesmyia. Zone Ch2 also shows an increase in the total number of 370 chironomid taxa found, with a maximum of 25 taxa at 149 cm. Of the dominant taxa in zone Ch1, C. mancus-type 371 and P. bathophila-type are still present in zone Ch2, while Lauterborniella strongly declines towards the upper 372 part of this zone. In addition, the chironomid assemblage still includes a.o. E. albipennis-type, P. nubeculosum-373 type and T. mendax-type. Both the sedimentological and the palynological record indicate a strong shift in the 374 water quality of the lake during this time period, with more nutrient-rich and turbid conditions present after 1140 375 cal. yr. BP. This transition is reflected in the chironomid-species composition where most species with specific 376 habitat requirements such as the presence of aquatic macrophytes or low-nutrient levels decline in favour of 377 Procladius and Ablabesmyia. These latter two taxa are typically interpreted as generalists and are able to persist 378 in dynamic conditions (e.g. Vallenduuk and Moller Pillot 2007). Ablabesmyia prefers shallow, warm and still water 379 and is associated with macrophytes (Engels et al. 2012). Additionally, Procladius is sensitive to increases in 380 phosphate, can occur in acidified lakes and tolerates pollution well (Luoto 2011). M. pedellus-type is also often 381 encountered in dynamic environments and together with Ablabesmyia is found in shallow lakes of less than five 382 metres deep (Engels et al. 2008a; Engels et al. 2008b; Engels et al. 2012). Potito et al. (2014) found a relation with high agricultural cover and the occurrence of this chironomid taxon, which is in line with the low AP values 383 384 and high upland herbs and Cerealia pollen percentages in this zone. M. pedellus-type however, has a preference for mesotrophic conditions and a stable oxygen regime (Moller Pillot and Klink 2009), and Engels et al. (2018) 385 386 therefore suggested that this increase could be a relative increase, with other taxa decreasing faster under 387 unfavourable conditions than M. pedellus-type. Other taxa that indicate higher trophic conditions are e.g. G. 388 pallens-type, Chironomus anthracinus-type, Chironomus plumosus-type, C. lateralis-type, Dicrotendipes 389 nervosus-type. Overall, the combined proxy-records indicate a shift in the lake ecology from the clear-water 390 mesotrophic conditions of zone Ch1 to the eutrophic, turbid conditions of zone Ch2.

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394 4.4 Factors controlling the chironomid fauna composition of Lake Uddelermeer

395 The most important precondition to establish a C-IT reconstruction is that temperature fluctuations are 396 the main factor influencing the chironomid composition. This is especially important for Holocene temperature 397 reconstructions, since the amplitude of temperature fluctuations in e.g. the mid-latitudinal areas of the northern 398 Hemisphere, was small when compared to Late Glacial studies. Chironomid communities of shallow, temperate 399 lakes such as Lake Uddelermeer are especially sensitive to changes in the environment since dominant controls on the chironomid composition like hypolimnetic oxygen depletion and climatic extremes are lacking (Langdon et 400 al. 2006). Based on the sedimentological, palynological and chironomid record, conditions in the lake itself 401 402 remained stable with clear-water, nutrient-poor conditions during the interval between 2500 and 1140 cal. yr. BP 403 i.e. covering chironomid zone Ch1 and the pollen zones Pa, Pb and Pc. The strong eutrophication of the lake at 404 the onset of chironomid zone Ch2 is visible in the sediment core as a colour change, in the pollen record (here 405 zone Pd) as an increase in algal species, and in the chironomid record as an increase of generalists such as 406 Ablabesmyia and Procladius (Fig. 5). Together these changes indicate a significant ecosystem shift in Lake 407 Uddelermeer from mesotrophic clear-water conditions to more nutrient-rich and turbid conditions. This conclusion is in line with results by Engels et al. (2018), who reconstructed stable clear-water conditions in Lake 408 409 Uddelermeer from 6000 cal. yr. BP onward and an increase in nutrient availability at 1030 cal. yr. BP from 410 sedimentary pigments, chironomids and pollen. Engels et al. (2018) placed the change to turbid lake-water 411 conditions at 1030 cal. yr. BP, which corresponds to the maximum increase in algae abundances at 185 cm in the 412 pollen data. Our high-resolution record indicates that the first onset of change occurred from 1140 cal. yr. BP 413 onward, with increasing percentages of algae and fungal spores and decreasing pollen percentages of Littorella 414 and Isoëtes and a shift to brown-green gyttja (Fig.5).

The pollen record additionally indicates changes in the regional vegetation during the investigated 415 416 period, with a reforestation phase in zone Pb from 1730 till 1270 cal. yr. BP (Fig. 5). These regional changes in 417 forest cover did not result in significant changes in the aquatic vegetation in the lake during Ch1. It is possible that 418 the presence of a slightly raised area surrounding Lake Uddelermeer, as seen on the LIDAR elevation map (Fig. 419 1), limited inflow of surface water runoff into the lake from the surrounding area, limiting the impact of regional 420 changes in land use. Increased sediment and nutrient input into the lake as a result of agriculture and 421 deforestation are often the main actor in ecosystem changes in lakes. Potito et al. (2014) identified agricultural 422 land cover as one of the most dominant environmental variables influencing chironomid composition. 423 Nevertheless, chironomid inferred temperatures can still be valid from these lakes when agriculture remained stable during the investigated time period (Potito et al. 2014). From the pollen record the presence of agriculture 424 in the area was evident, albeit in low percentages which remained constant in pollen zones Pa and Pb and only 425 426 slightly increased during Pc. In addition, consistently high lake levels are reconstructed for the investigated period 427 (Engels et al. 2016), suggesting that changes in water level, which could also affect the chironomid assemblages 428 (Engels and Cwynar 2011), did not play a role either. We therefore conclude that environmental conditions in the 429 lake remained stable in the period between 2525 cal. yr. BP and 1140 cal. yr. BP and suggest that the chironomid 430 record could provide a reliable temperature reconstruction for this period. The chironomid fauna that was existent 431 during the formation of the upper samples (220-100 cm depth), obtained from the green algal gyttja (Ch2), was 432 strongly influenced by the local eutrophication process. These samples are therefore deemed unreliable for 433 temperature reconstruction purposes and will not be discussed. 434

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437 **4.5 Chironomid-inferred mean July air temperature reconstruction**

Reconstructed mean July air temperature values vary between 16.1 and 19.1°C with an average
reconstructed temperature of 17.6°C (Fig. 6). This places the reconstruction in the upper part of the temperature
range of the calibration dataset (3.5-18.4 °C) (Heiri *et al.* 2011). Sample-specific errors range between 1.4 and 1.6
°C.

442 A number of temperature fluctuations can be observed between 2525 and 1140 cal. yr. BP. At the start 443 of the investigated period (2525-2240 cal. yr. BP, or 550-470 cm depth), mean July air temperature values are low, with an average temperature of 17.4°C. For the period between 2240 and 1760 cal. yr. BP (470-390 cm), 444 445 slightly higher temperatures are reconstructed, with an average temperature of 18.6°C. Maximum reconstructed 446 temperatures of 19.1°C are reached at a depth of 464 cm (2197 cal. yr. BP). Slightly colder conditions, with an 447 average mean July air temperature of 17.1°C, prevailed in the period between 1760 and 1280 cal. yr. BP (390-448 255 cm). The coldest phase is reconstructed from 330-304 cm (1551-1465 cal. yr. BP) with an average mean July 449 air temperature of only 16.5°C. This is followed by slightly higher (average 18.3°C) values for the period between 450 1280 and 1140 cal. yr. BP (255-220 cm).

The down-core evaluation analyses suggest that most chironomid assemblages in the fossil record are classified as having both 'no close' or 'no good' modern analogue and a 'poor' or 'very poor' fit with temperature. Only two samples contain taxa which are not included in the calibration dataset, reaching values of maximum 2%. Therefore, even though the composition of the chironomid assemblages in the fossil record differs from the modern assemblages, the individual taxa in the fossil samples are well represented in the modern samples. WA-PLS can perform relatively well in no-analogue conditions as long as the majority of the chironomid assemblage is well represented in the calibration data set (Birks 1998).

458 Reconstructed mean July air temperatures in part, exceed the maximum mean July air temperature of 459 the modern samples in the calibration dataset of 18.4 °C. This might have affected the absolute values of the temperature reconstruction. The magnitude of the observed fluctuations (1.39-1.56 °C) is similar to the sample-460 461 specific error of the reconstructed temperatures. Nevertheless, the fluctuations are recorded in multiple samples 462 on a high temporal resolution, thus giving a consistent picture of these temperature changes. Moreover, observed 463 fluctuations approximately coincide with shifts in the chironomid assemblage. For example, during the two colder oscillations, taxa with relatively low optimum temperatures are present, such as Limnophyes/Paralimnophyes, 464 465 Procladius and P. sordidellus. Similarly, the sample with the highest reconstructed mean July air temperature 466 (464 cm) coincides with maximum abundances of E. albipennis-type, which is found in the warmer lakes of the calibration dataset (Heiri et al. 2011) suggesting that temperature is indeed the main driver of the chironomid 467 fauna in this part of the record. When we compare the reconstructed C-IT from Lake Uddelermeer to the average 468 469 July temperature of 17 °at de Bilt (50 km SW of Lake Uddelermeer) for the period 1901-2017 it seems that our 470 reconstructed temperatures are relatively high. Even during the DACP reconstructed temperatures are similar to 471 recent conditions (17 °C vs. 17.1°C) indicating that the record is likely overestimating the absolute temperature. 472 Potito et al. (2014) found that sites with agriculture in their catchment can overestimate temperature as a result of 473 increased production in the lake. Other chironomid records from NW Europe also indicate that temperature during 474 the DACP was at least 1°C lower than current temperature values (Northern England: Langdon et al. 2004; 475 Barber et al. 2013; Cairngorms UK: Dalton et al. 2005; northern Alps: Millet et al. 2009). Additionally, the 476 reconstructed temperature range is located in the upper ranges of the Norwegian/Alpine training set which could 477 result in so called edge-effects. It therefore seems likely that the absolute values of our C-IT reconstruction 478 overestimate past July air temperatures.

479

480 **5 Discussion**

481 Based on the chronology of the record, we can correlate the observed temperature oscillations in the record to climate fluctuations as recorded in the Northern Hemisphere (e.g. Ljungqvist 2010). The slightly higher 482 reconstructed temperatures in the period from 2240-1760 cal. yr. BP probably correspond with the RWP. The 483 484 subsequent cold phase (1760-1280 cal. yr. BP) can be correlated to the DACP. The warmer conditions 485 reconstructed for the upper part of the record (1280-1140 cal. yr. BP) can be linked to the MCA (Fig. 6). The C-IT 486 record shows that temperature during the Iron Age (till 2240 cal. yr. BP) was lower than during the RWP by an 487 estimated 1.2 °C. To evaluate the representability of these temperature trends in NW Europe we will first compare 488 our results to other C-IT records. In a second step we will compare our results to other available climate 489 reconstructions from NW Europe including detailed tree-ring records and global data compilations.

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491 Figure 7 around here

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5.1 Chironomid-inferred temperature reconstructions from NW Europe

494 To our knowledge only two other Late Holocene C-IT reconstructions, which have been identified as 495 representative NW-European temperature trends for the period discussed here, are available. Both of these sites 496 are located in Northern England: Talkin Tarn (Langdon et al. 2004) and Bigland Tarn (Barber et al. 2013). A RWP, 497 a distinct DACP and the MCA can be distinguished in the record from Bigland Tarn (Fig. 7). These phases might 498 also be visible in the record from Talkin Tarn, although the comparison with that record is less robust, due to the 499 lower sampling resolution and uncertain chronology of that site. The similarity between the three sites, both in 500 timing as well as amplitude of temperature change, suggests that the reconstructed temperature fluctuations in 501 Lake Uddelermeer record reflect a regional climate signal. The overall trend of the reconstruction from 502 Uddelermeer is very similar to that of Bigland Tarn and Talkin Tarn. The RWP is the warmest phase in the 503 Uddelermeer record with an increase in temperature of 1.2 °C compared to the Iron Age. This is similar to the 504 increase in the Bigland Tarn record of 1°C. The average temperature drop from the RWP to the DACP in the 505 Uddelermeer record is 1.5 °C whereas this shift in the Bigland Tarn record is slightly bigger with 2.2 °C. The MCA 506 forms the warmest period in the Bigland Tarn record with an average temperature of 13.7°C. The MCA in the 507 Uddelermeer is only partly present and as seen in the Bigland Tarn record, the first part is not the warmest part of 508 the period. When we compare absolute values of the independent reconstructions, it becomes apparent that 509 reconstructed temperatures from both sites in England are consistently lower by 4-5 °C than the reconstruction 510 from Lake Uddelermeer (Fig. 7). This could in part be the result of the more western location and higher elevation 511 of these sites. The current average July temperatures of both Bigland Tarn as Talkin Tarn are around 15°C, compared to the average July temperature of 17°C at Uddelermeer. Additionally, previous studies have shown 512 513 that the application of different chironomid-climate calibration datasets to the same fossil record will yield C-IT 514 records that are up to several degrees different in absolute reconstructed temperatures (e.g. Engels et al. 2014; 515 Bajolle et al. 2018). The temperature reconstructions from England are based on the Norwegian calibration 516 dataset, which has a lower maximum temperature (16 °C; Brooks and Birks 2000) than the extended and 517 combined Swiss-Norwegian dataset used for the reconstruction from Uddelermeer (18.4 °C; Heiri et al. 2011). 518 However, it is likely that a large part of the difference in absolute temperatures is the result of a temperature overestimation in the record from Lake Uddelermeer. 519

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521 Figure 8 around here

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523 5.2 Regional evidence from the NH for the RMP, DACP and MCA

524 When we compare the start and end dates of the climatic phases indentified in the record from 525 Uddelermeer to other temperature reconstructions and overviews from the Northern Hemisphere we observe that 526 overall there is a good match between the available records (Fig. 8). All datasets shown in Fig. 8 show a similar 527 starting date for the MCA, which is placed around 1300 cal. yr. BP. The start of the DACP shows slightly more 528 variation in timing with starting dates ranging between 1536-1760 cal. yr. BP. Taking into account all age 529 uncertainties it is likely that all records and overviews show the same simultaneous event. The start of the DACP 530 in Uddelermeer is the earliest of the available estimates with a suggested starting date of 1760 cal. yr. BP. The 500-year duration of the DACP in the Uddelermeer record is 150 years longer than the duration of the DACP in 531 532 the tree-ring record presented by Büntgen et al. (2011).

533 The latter part of the DACP in the tree-ring record of Büntgen et al. (2011) dated at 1414-1290 cal. yr. BP 534 (536-660 AD) was later termed the Late Antique Little Ice Age or LALIA (Büntgen et al. 2016). This LALIA shows 535 a strong temperature decline and is causally linked to a cluster of volcanic eruptions in 1414; 1410 and 1403 cal. 536 yr. BP (536, 540 and 547 AD). Büntgen et al. (2017) promote the substitution of the DACP by the LALIA. 537 However, the LALIA has not been identified in a large range of proxies and sites, and falls within the previously 538 defined DACP. Additionally, since the timing and the causality differ, with a link to volcanism for the LALIA and an unknown cause for the DACP, the LALIA cannot be used as a substitution of the DACP. It therefore seems more 539 540 appropriate to refer to this cold stage as a whole as the DACP (sensu Helama et al. 2017b; Helama et al. 2017a). 541 The Uddelermeer record does suggest a colder second part of the DACP which starts at 1550 cal. yr. BP and 542 lasts till 1450 cal. yr. BP. It is possible that this colder phase reflects the LALIA, although the chronology of the 543 cold period observed in our Uddelermeer record does not match the timing and duration observed in the tree ring record (Büntgen et al. 2017). More detailed and well-dated studies covering this time period would contribute to a 544 545 better understanding of this coldest phase of the DACP.

An earlier start and a longer duration of the DACP in the Netherlands, compared to the DACP recorded in the tree ring record, could be the result of different local climatic conditions. Nevertheless, a longer and earlier DACP is in line with both the Bigland Tarn record as well as the two overview studies by Ljunqvist (2009) and Helama (2017). Aside from the tree-ring data set, the Uddelermeer chironomid record is the most detailed temperature record reconstructing absolute temperature values during the Dark Ages Cold Period in for NW Europe.

552 The start of the RWP cannot be identified in all studies presented in Fig. 8. In the Bigland Tarn and 553 Uddelermeer records the start of the RWP is placed at 2365 cal. yr. BP and 2240 cal. yr. BP, respectively. Taking into account the error margins of the chronology of both studies, this could be a simultaneous event. In the tree-554 ring data set no clear RWP is evident, but the start of a warmer phase does coincide with the start of the RWP at 555 556 Uddelermeer. However, this warm period as identified in the tree ring record (Büntgen et al. 2016) is interrupted 557 by a colder phase. The reconstructed temperatures in Uddelermeer and Bigland Tarn are both stable during the 558 RWP and show no evidence of a brief colder period. The most probable explanation for this discrepancy are different climatic conditions in the Alps in contrast to the lower lying sites, as well as differences in sampling 559 560 resolution and proxy sensitivity. In the chironomid record from the northern French Alps by Millet et al. (2009), 561 which is not interpreted as a representative temperature reconstruction for the first millennium, also no clear RWP was visible. Different climatic conditions might also explain the differences in duration and the start of the DACP 562 563 across the European continent. Overall, the start and end dates of the different climatic phases fall within the 564 dating uncertainties and current results indicate that especially the MCA and DACP and to a lesser extent the 565 RWP can be considered as simultaneous events across NW Europe.

566

567 6 Conclusions

We presented the first high-resolution chironomid record for the late Subatlantic in the Netherlands from 568 569 Lake Uddelermeer. Our data shows that in the period 2500 to 1140 cal. yr. BP the lake was mesotrophic with clear-water conditions and sustained a Littorellion. From 1140 cal. yr. BP onward a shift in the water quality was 570 571 reconstructed with high-nutrient turbid conditions and a dominance of algal species. The conditions in the lake in 572 the period 2500-1140 cal. yr. BP do not signal large ecological changes or effects of human impact and thus 573 suggest that this record is suitable for a temperature reconstruction. In this C-IT record from Lake Uddelermeer, 574 we identified a RWP, a DACP and the MCA. The RWP is placed in the period from 2240-1760 cal. yr. BP, the DACP starts at 1760 cal. yr. BP and the MCA starts at 1280 cal. yr. BP. Our detailed summer temperature record 575 shows that the average temperature drops by 1.5°C from the RWP to the DACP. Overall the temperature 576 577 reconstructions appears to be too high, however, the trends, temperature fluctuations and timing of the cold and 578 warm periods in this records are in agreement with other records and reconstructions indicating that the 579 Uddelermeer record can be used as an representative record for temperature in the Netherlands during the first 580 millennium AD.

581

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592 **References**

- 593 AHN Actueel Hoogtebestand Nederland. www.ahn.nl; www.nationaalgeoregister.nl.
- Archeologisch Basis Register (ABR) in Nationale Onderzoeksagenda Archeologie (NOaA). (2016) Rijksdienst
 Cultureel Erfgoed. https://archeologieinnederland.nl/bronnen-en-kaarten/noaa. Accessed 01-08-2018
 Axford Y., Geirsdóttir Á., Miller G.H., Langdon P.G. (2009) Climate of the Little Ice Age and the past 2000 years in
 northeast Iceland inferred from chironomids and other lake sediment proxies. Journal of Paleolimnology
- 598 41:7-24 doi:https://doi-org.proxy.library.uu.nl/10.1007/s10933-008-9251-1
- 599Bajolle L., Larocque-Tobler I., Gandouin E., Lavoie M., Bergeron Y., Ali A.A. (2018) Major postglacial summer600temperature changes in the central coniferous boreal forest of Quebec (Canada) inferred using
- chironomid assemblages. Journal of Quaternary Science 33:409-420 doi:https://doi.org/10.1002/jqs.3022
 Barber K., Brown A., Langdon P., Hughes P. (2013) Comparing and cross-validating lake and bog palaeoclimatic
 records: A review and a new 5,000 year chironomid-inferred temperature record from northern England.
 Journal of Paleolimnology 49:497-512 doi:https://doi.org.proxy.library.uu.nl/10.1007/s10933-012-9656-8
- Beug H.-J. (2004) Leitfaden der pollenbestimmung, für Mitteleuropa und angrenzende Gebiete. Fischer,
 Stuttgart:542 p.
- Birks H.J.B. (1998) Numerical tools in palaeolimnology-Progress, potentialities, and problems. Journal of
 Paleolimnology 20:307-332
- Birks H.J.B., Ter Braak C.J.F., Line J.M., Juggins S., Stevenson A.C. (1990) Diatoms and pH reconstruction. Phil
 Trans R Soc Lond B 327:263-278

612 Uddelermeer. Vrije Universiteit Amsterdam, Amsterdam 613 Brodin Y.W. (1986) The postglacial history of Lake Flarken, southern Sweden, interpreted from subfossil insect 614 remains. Internationale Revue der gesamten Hydrobiologie und Hydrographie 71:371-432 615 Bronk Ramsey C. (2009) Bayesian analysis of radiocarbon dates. Radiocarbon 51:337-360 Brooks S.J., Axford Y., Heiri O., Langdon P.G., Larocque-Tobler I. (2012) Chironomids can be reliable proxies for 616 617 Holocene temperatures. A comment on Velle et al. (2010). The Holocene 22:1495-1500 doi:https://doiorg.proxy.library.uu.nl/10.1177/0959683612449757 618 Brooks S.J., Birks H.J.B. (2000) Chironomid-inferred late-glacial and early-Holocene mean July air temperatures 619 for Kråkenes Lake, western Norway. Journal of Paleolimnology 23:77-89 620 Brooks S.J., Langdon P.G., Heiri O. (2007) The indentification and use of Palaeartic Chironomidae larvae in 621 622 palaeoecology. QRA Technical guide No. 10. Quaternary Research Association, London 623 Bunnik F.P.M. (1999) Vegetationsgeschichte der Lößbörden zwischen Rhein und Maas von der Bronzezeit bis in 624 die frühe Neuzeit., Utrecht University 625 Büntgen U., Myglan V.S., Ljungqvist F.C., McCormick M., Di Cosmo N., Sigl M., Jungclaus J., Wagner S., Krusic P.J., Esper J. (2017) Reply to 'Limited Late Antique cooling'. Nature Geoscience 10:243 626 627 Büntgen U., Myglan V.S., Ljungqvist F.C., McCormick M., Di Cosmo N., Sigl M., Jungclaus J., Wagner S., Krusic P.J., Esper J., Kaplan J.O., de Vaan M.A.C., Luterbacher J., Wacker L., Tegel W., Kirdyanov A.V. (2016) 628 629 Cooling and societal change during the Late Antique Little Ice Age from 536 to around 660 AD. Nature 630 Geosci 9:231-236 doi:https://doi.org/10.1038/NGEO2652 631 Büntgen U., Tegel W., Nicolussi K., McCormick M., Frank D., Trouet V., Kaplan J.O., Herzig F., Heussner K.U., 632 Wanner H., Luterbacher J., Esper J. (2011) 2500 years of European climate variability and human 633 susceptibility. Science 331:578-582 doi:https://doi.org/10.1126/science.1197175 634 Cohen K.M., Toonen W.H.J., Weerts H.J.T. (2016) Overstromingen van de Rijn gedurende het Holocene 635 Relevantie van de grootste overstromingen voor archeologie van het Nederlands rivierengebied. 636 Deltares 637 Dalton C., Birks H.J.B., Brooks S.J., Cameron N.G., Evershed R.P., Peglar S.M., Scott J.A., Thompson R. (2005) 638 A multi-proxy study of lake-development in response to catchment changes during the Holocene at 639 Lochnagar, north-east Scotland. Palaeogeography, Palaeoclimatology, Palaeoecology 221:175-201 640 doi:https://doi.org/10.1016/j.palaeo.2005.02.007 641 Dörfler W., Feeser I., van den Bogaard C., Dreibrodt S., Erlenkeuser H., Kleinmann A., Merkt J., Wiethold J. (2012) A high-quality annually laminated sequence from Lake Belau, Northern Germany: revised 642 643 chronology and its implications for palynological and tephrochronological studies The Holocene 22:1413-644 1426 doi: https://doi.org/10.1177/0959683612449756 645 Engels S., Bakker M.A.J., Bohncke S.J.P., Cerli C., Hoek W.Z., Jansen B., Peters T., Renssen H., Sachse D., van Aken J.M. (2016) Centennial-scale lake-level lowstand at Lake Uddelermeer (The Netherlands) indicates 646 647 changes in moisture source region prior to the 2.8-kyr event. The Holocene 26:1075-1091 doi:https://doiorg.proxy.library.uu.nl/10.1177/0959683616632890 648 Engels S., Bohncke S.J.P., Bos J.A.A., Heiri O., Vandenberghe J., Wallinga J. (2008a) Environmental inferences 649 650 and chironomid-based temperature reconstructions from fragmentary records of the Weichselian Early 651 Glacial and Pleniglacial periods in the Niederlausitz area (eastern Germany). Palaeogeography, 652 Palaeoclimatology, Palaeoecology 260:405-416 doi:https://doi.org/10.1016/j.palaeo.2007.12.005 653 Engels S., Bohncke S.J.P., Heiri O., Nyman M. (2008b) Intraregional variability in chironomid-inferred temperature 654 estimates and the influence of river inundations on lacustrine chironomid assemblages. Journal of 655 Paleolimnology 40:129-142 doi:https://doi-org.proxy.library.uu.nl/10.1007/s10933-007-9147-5

Bohncke S.J.P. (1999) Verslag betreffende het onderzoek naar de archiefwaarde van de sedimenten van het

611

- Engels S., Cwynar L.C. (2011) Changes in fossil chironomid remains along a depth gradient: evidence for
 common faunal thresholds within lakes. Hydrobiologia 665:15-38 doi:https://doi org.proxy.library.uu.nl/10.1007/s10750-011-0601-z
- Engels S., Cwynar L.C., Rees A.B.H., Shuman B.N. (2012) Chironomid-based water depth reconstructions: an
 independant evaluation of site-specific and local interference models. Journal of Palaeolimology
 doi:https://doi-org.proxy.library.uu.nl/10.1007/s10933-012-9638-x
- Engels S., Self A.E., Luoto T.P., Brooks S.J., Helmens K.F. (2014) A comparison of three Eurasian chironomid climate calibration datasets on a W-E continentality gradient and the implications for quantitative
 temperature reconstructions. Journal of Paleolimnology 51:529-547 doi:https://doi-
- 665 org.proxy.library.uu.nl/10.1007/s10933-014-9772-8
- Engels S., van Oostrom R., Cherli C., Dungait J.A.J., Jansen B., van Aken J.M., van Geel B., Visser P.M. (2018)
 Natural and anthropogenic forcing of Holocene lake ecosystem development at Lake Uddelermeer (The
 Netherlands). Journal of Paleolimnology 59:329-347 doi:https://doi-
- 669 org.proxy.library.uu.nl/10.1007/s10933-017-0012-x
- Erkens G. (2009) Sediment dynamics in the Rhine catchement-Quantification of fluvial response to climate
 change and human impact. Netherlands Geographical Studies 388
- Faegri K., Iversen J. (1989) Textbook of Pollen Analysis, revised by Faegri K, Kaland PE, Krzywinski K. J Wiley,
 New York
- Farmer A.M., Spence D.H.N. (1986) The growth strategies and distribution of isoetids in Scottish freshwater
 lochs. Aquatic Botany 26:247-258
- 676 Gräslund B., Price N. (2012) Twilight of the gods? The 'dust veil event' of AD 536 in critical perspective Antiquity 677 86:428-443 doi:10.1017/S0003598X00062852
- 678 Grimm E.C. (1991-2015) Tilia., 2.0.41 edn.,
- Groenewoudt B.J., Schut P.A.C., van der Heijden F.J.G., Peeters H., Wispelwey M.H. (2006) Een inventariserend
 veldonderzoek bij de Hunneschans (Uddel, Gelderland): nieuwe gegevens over de steentijdbewoning bij
 het Uddelermeer en een beknopt overzicht van de onderzoeksgeschiedenis van de Hunneschans:
 Rapportage Archeologische Monumentenzorg 143.
- Groenewoudt B.J., van Lanen R.J. (2018) Diverging decline. Reconstructing and validating (post-)Roman
 population trends (AD 0-1000) in the Rhine-Meuse delta (the Netherlands). Post Classical Archaeologies
 8:30
- Hannon G.E., Gaillard M.-J. (1997) The plant-macrofossil record of past lake-level changes. Journal of
 Paleolimnology 18:15-28
- Heidinga H.A. (1987) Medieval Settlement and Economy North of the Lower Rhine: Archeology and History of
 Kootwijk and the Veluwe (the Netherlands). Van Gorcum Ltd,
- Heiri O., Brooks S.J., Birks H.J.B., Lotter A.F. (2011) A 274-lake calibration data-set and inference model for
 chironomid-based summer air temperature reconstruction in Europe. Quaternary Science Reviews
 30:3445-3456 doi:https://doi.org/10.1016/j.quascirev.2011.09.006
- Heiri O., Brooks S.J., Renssen H., Bedford A., Hazekamp M., Ilyashuk B., Jeffers E.S., Lang B., Kirilova E.,
 Kuiper S. (2014) Validation of climate model-inferred regional temperature change for late-glacial
 Europe. Nature communications 5:4914 doi:https://doi.org/10.1038/ncomms5914
- Heiri O., Lotter A.F., Lemcke G. (2001) Loss on ignition as a method for estimating organic and carbonate content
 in sediments: reproducibility and comparability of results. Journal of Paleolimnology 25:101-110
- Helama S., Jones P.D., Briffa K.R. (2017a) Dark Ages Cold Period: A literature review and directions for future
 research. The Holocene 27:1600-1606 doi:https://doi.org/10.1177/0959683617693898
- 700 Helama S., Jones P.D., Briffa K.R. (2017b) Limited Late Antique cooling. Nature Geoscience 10:242

- Jansma E., Gouw-Bouman M.T.I.J., Van Lanen R.J., Pierik H.J., Cohen K.M., Groenewoudt B.J., Hoek W.Z.,
 Stouthamer E., Middelkoop H. (2014) The Dark Age of the Lowlands in an interdisciplinary light: people,
 landscape and climate in The Netherlands between AD 300-1000. European Journal of Post-Classical
 Archaeologies (PCA 4):471-476
- Kalis A.J., Karg S., Meurers-Balke H., Teunissen-van OOrschot H. (2008) Mensch und Vegetation am Unteren
 Niederrhein währen der Eisen- und Römerzeit. In: Müller M, Schalles H-J, Zieling N (eds) Colonia Ulpia
 Traiana, Xanten und sein Umland in römischer Zeit, Xantener Berichte, Geschichte der Stadt Xanten.,
 vol 1. pp 31-48
- 709 Knol E. (1993) De Noordnederlandse kustlanden in de Vroege Middeleeuwen., VU University Amsterdam
- 710 Kraanen C.J.M., Pape J.C. (1965) De bodemgesteldheid van de omgeving van het Uddelermeer. STIBOKA,
- Langdon P.G., Barber K.E., Lomas-Clarke S.H. (2004) Reconstructing climate and environmental change in
 northern England through chironomid and pollen analyses: evidence from Talkin Tarn, Cumbria. Journal
 of Palaeolimology 32:197-203
- Langdon P.G., Ruiz Z., Brodersen K.P., Foster I.D.L. (2006) Assessing lake eutrophication using chironomids:
 understanding the nature of community response in different lake types. Freshwater biology 51:562-577
 doi:https://doi.org/doi:10.1111/j.1365-2427.2005.01500.x
- Larsen L.B., Vinther B.M., Briffa K.R., Melvin T.M., Clausen H.B., Jones P.D., Siggaard-Andersen M.L., Hammer
 C.U., Eronen M., Grudd H., Gunnarson B.E., Hantemirov R.M., Naurzbaev M.M., Nicolussi K.C.L. (2008)
 New ice core evidence for a volcanic cause of the A.D. 536 dust veil. Geophysical Research Letters
 35:n/a-n/a doi:https://doi-org.proxy.library.uu.nl/10.1029/2007GL032450
- Litt T., Schölzel C., Kühl N., Brauer A. (2009) Vegetation and climate history in the Westeifel Volcanic Field
 (Germany) during the past 11 000 years based on annually laminated lacustrine maar sediments.
 Boreas 38:679-690 doi:https://doi.org/10.1111/j.1502-3885.2009.00096.x
- Ljungqvist F.C. (2009) Temperature proxy records covering the last two millennia: a tabular and visual overview.
 Geografiska Annaler: Series A, Physical Geography 91:11-29 doi:https://doi.org/10.1111/j.1468 0459.2009.00350.x
- Ljungqvist F.C. (2010) A new reconstruction of temperature variability in the extra-tropical northern hemisphere
 during the last two millennia. Geografiska Annaler, series A:339-351
- Luoto T.P. (2011) The relationship between water quality and chironomid distribution in Finland—a new
 assemblage-based tool for assessments of long-term nutrient dynamics. Ecological Indicators 11:255 262 doi:https://doi.org/10.1016/j.ecolind.2010.05.002
- Luoto T.P. (2012) Intra-lake patterns of aquatic insect and mite remains. Journal of Paleolimnology 47:141-157
 doi:https://doi-org.proxy.library.uu.nl/10.1007/s10933-011-9565-2
- McCormick M., Büntgen U., Cane M.A., Cook E.R., Harper K., Huybers P., Litt T., Manning S.W., Mayewski P.A.,
 More A.F.M., Nicolussi K., Tegel W. (2012) Climate Change during and after the Roman Empire:
 Reconstructing the Past from Scientific and Historical Evidence. Journal of Interdisciplinary History
 43:169-220
- McKeown M., Potito A.P. (2016) Assessing recent climatic and human influences on chironomid communities
 from two moderately impacted lakes in western Ireland. Hydrobiologia 765:245-263 doi:https://doi org.proxy.library.uu.nl/10.1007/s10750-015-2418-7
- Millet L., Arnaud F., Heiri O., Magny M., Verneaux V., Desmet M. (2009) Late-Holocene summer temperature
 reconstruction from chironomid assemblages of Lake Anterne, northern French Alps. The Holocene
 19:317-328 doi:https://doi.org/10.1177/0959683608100576
- Moller Pillot H.K.M. (1984a) De Larven der Nederlandse Chironomidae (Diptera): Orthocladiinae sensu lato.
 Nederlandse Faunistische Mededelingen 1:1-175

746 Moller Pillot H.K.M. (1984b) De larven der Nederlandse Chironomidae (Diptera); (Inleiding, Tanypodinae & 747 Chironomini). Nederlandse Faunistische Mededelingen 1:1-277 748 Moller Pillot H.K.M. (2013) Chironomidae Larvae of the Netherlands and adjacent lowlands: biology and ecology 749 of the chironomini. KNNV publishing, Moller Pillot H.K.M., Klink A.G. (2009) Chironomidae larvae. Biology and Ecology of the Chironomini KNNV 750 751 Publishing, Zeist 752 Moore P.D., Webb J.A., Collison M.E. (1991) Pollen analysis. Blackwell scientific publications, 753 Nazarova L., de Hoog V., Hoff U., Dirksen O., Diekmann B. (2013) Late Holocene climate and environmental changes in Kamchatka inferred from the subfossil chironomid record. Quaternary Science Reviews 754 755 67:81-92 doi:https://doi.org/10.1016/j.guascirev.2013.01.018 756 Padisák J., Krienitz L., Scheffler W., Koschel R., Kristiansen J., Grigorszky I. (1998) Phytoplankton succession in 757 the oligotrophic Lake Stechlin (Germany) in 1994 and 1995 Hydrobiologia 369:179-197 758 Pierik H.J. (2017) Past human-landscape interactions in the Netherlands: Reconstructions from sand belt to 759 coastal-delta plain for the first millennium AD. vol 139. Utrecht studies in earth sciences. Utrecht 760 University, 761 Pierik H.J., Cohen K.M., Stouthamer E. (2016) A new GIS approach for reconstructing and mapping dynamic late 762 Holocene coastal plain palaeogeography. Geomorphology 270:55-70 doi:http://dx.doi.org/10.1016/j.geomorph.2016.05.037 763 764 Pierik H.J., Stouthamer E., Cohen K.M. (2017) Natural levee evolution in the Rhine-Meuse delta, the Netherlands, 765 during the first millennium CE. Geomorphology 295:215-234 766 doi:https://doi.org/10.1016/j.geomorph.2017.07.003 767 Pierik H.J., van Lanen R.J. (2017) Roman and early-medieval habitation patterns in a delta landscape: The link 768 between settlement elevation and landscape dynamics. Quaternary International 769 doi:https://doi.org/10.1016/j.guaint.2017.03.010 770 Pierik H.J., Van Lanen R.J., Gouw-Bouman M.T.I.J., Groenewoudt B., Wallinga J., Hoek W.Z. (2018) Controls on 771 late Holocene drift-sand dynamics: the role of people and climate on inland aeolian activity in the 772 Netherlands The Holocene 28:1361-1381 doi:https://doi.org/10.1177/0959683618777052 773 Koomen A., Maas G. (2004) Geomorfologische Kaart Nederland (GKN); Achtergronddocument bij het 774 landsdekkende digitale bestand. Alterra. 775 Polak B. (1959) Palynology of the "Uddeler Meer". A Contribution to our Knowledge of the Vegetation and of the 776 Agriculture in the Northern Part of the Veluwe in Prehistoric and Early Historic Times. Acta botanica 777 neerlandica 8:547-571 778 Potito A.P., Woodward C.A., McKeown M., Beilman D.W. (2014) Modern influences on chironomid distribution in 779 western Ireland: potential for palaeoenvironmental reconstruction. Journal of Paleolimnology 52:385-404 780 doi:https://doi-org.proxy.library.uu.nl/10.1007/s10933-014-9800-8 781 Punt W., Blackmore S., Clarke G.C.S. (1976) The Northwest European pollen flora 1. Elsevier, Amsterdam 782 Punt W., Blackmore S., Clarke G.C.S. (1980) The Northwest European pollen flora 2. Elsevier, Amsterdam 783 Punt W., Blackmore S., Clarke G.C.S. (1981) The Northwest European pollen flora 3. Elsevier, Amsterdam 784 Punt W., Blackmore S., Clarke G.C.S. (1984) The Northwest European pollen flora 4. Elsevier, Amsterdam 785 Punt W., Blackmore S., Clarke G.C.S. (1988) The Northwest European pollen flora 5. Elsevier, Amsterdan=m 786 Punt W., Blackmore S., Clarke G.C.S. (1991) The Northwest European pollen flora 6. Elsevier, 787 Punt W., Blackmore S., Clarke G.C.S. (1995) The Northwest European pollen flora 7. Elsevier, 788 Punt W., Blackmore S., Hoen P.P., Stafford P.J. (2003) The Northwest European pollen flora 8. Elsevier, 789 Amsterdam

790 Rasmussen S.O., Andersen K.K., Svensson A.M., Steffensen J.P., Vinther B.M., Clausen H.B., Siggaard-791 Andersen M.L., Johnsen S.J., Larsen L.B., Dahl-Jensen D., Bigler M., Röthlisberger R., Fischer H., 792 Goto-Azuma K., Hansson M.E., Ruth U. (2006) A new Greenland ice core chronology for the last glacial 793 termination. Journal of Geophysical Research: Atmospheres 111:n/a-n/a doi:10.1029/2005jd006079 794 Riechelmann D., Gouw-Bouman M.T.I.J. (2019) Climate during the first millennium AD in NW Europe: a review of climate reconstructions from terrestrial archives. Quaternary Research 91:111-131 795 796 doi:https://doi.org/10.1017/qua.2018.84 797 Rieradevall M., Brooks S.J. (2001) An identification guide to subfossil Tanypodinae larvae (Insecta: Diptera: Chrironomidae) based on cephalic setation. Journal of Paleolimnology 25:81-99 798 Roelofs J.G.M., Schuurkes J.A.A.R., Smits A.J.M. (1984) Impact of acidification and eutrophication on 799 800 macrophyte communities in soft waters. II. Experimental studies. Aquatic Botany 18:389-411 801 Rørslett B., Brettum P. (1989) The genus Isoëtes in Scandinavia: an ecological review and perspectives. Aquatic 802 Botany 35:223-261 803 Sand-Jensen K. (1978) Metabolic adaptation and vertical zonation of Littorella uniflora (L.) Aschers. and Isoetes 804 lacustris L. Aquatic Botany 4:1-10 805 Sohl H. (1983) A palaeo-ecological investication of the Late Glacial and Holocene lake sediments of the 806 Uddelermeer (The Netherlands): methods and some provisional results. Quaternary studies in Poland 807 9:547-571 808 Stouthamer E., Berendsen H.J.A. (2000) Factors Controlling the Holocene Avulsion History of the Rhine-Meuse 809 Delta (The Netherlands). Journal of Sedimentary Research 70:1051-1064 doi:10.1306/033000701051 810 Taylor K.J., McGinley S., Potito A.P., Molloy K., Beilman D.W. (2018) A mid to late Holocene chironomid-inferred 811 temperature record from northwest Ireland. Palaeogeography, Palaeoclimatology, Palaeoecology 505:274-286 doi:https://doi.org/10.1016/j.palaeo.2018.06.006 812 813 Taylor K.J., Potito A.P., Beilman D.W., Ghilardi B., O'Connell M. (2013) Palaeolimnological impacts of early prehistoric farming at Lough Dargan, County Sligo, Ireland. Journal of Archaeological Science 40:3212-814 3221 doi:https://doi.org/10.1016/j.jas.2013.04.002 815 816 Taylor K.J., Potito A.P., Beilman D.W., Ghilardi B., O'Connell M. (2017a) Impact of early prehistoric farming on chironomid communities in northwest Ireland. Journal of Paleolimnology 57:227-244 817 818 doi:https://doi.org/10.1007/s10933-017-9942-6 819 Taylor K.J., Stolze S., Beilman D.W., Potito A.P. (2017b) Response of chironomids to Neolithic land-use change 820 in north-west Ireland. The Holocene 27:879-889 doi:https://doi.org/10.1177/0959683616675935 821 Telford R.J., Birks H.J.B. (2011) A novel method for assessing the statistical significance of quantitative 822 reconstructions inferred from biotic assemblages. Quaternary Science Reviews 30:1272-1278 823 doi:https://doi.org/10.1016/j.guascirev.2011.03.002 824 ter Braak C.J.F., Šmilauer P. (2012) Canoco reference manual and user's guide: software for ordination, version 825 5.0. Microcomputer power, Teunissen D. (1990) Palynologisch onderzoek in het oostelijk rivierengebied: een overzicht. In: Mededelingen van 826 827 de afdeling Biogeologie van de Discipline Biologie van de Katholieke Universiteit van Nijmegen. p 16 828 Tinner W., Lotter A.F., Ammann B., Conedera M., Hubschmid P., van Leeuwen J.F.N., Wehrli M. (2003) Climatic 829 change and contemporaneous land-use phases north and south of the Alps 2300 BC to 800 AD. 830 Quaternary Science Reviews 22:1447-1460 doi:https://doi.org/10.1016/S0277-3791(03)00083-0 Toohey M., Krüger K., Sigl M., Stordal F., Svensen H. (2016) Climatic and societal impacts of a volcanic double 831 832 event at the dawn of the Middle Ages Climatic Change 136:401-412 833 Toonen W.H.J. (2013) A Holocene flood record of the Lower Rhine. Utrecht University

- Vallenduuk H.J., Moller Pillot H.K.M. (2007) Chironomidae larvae of the Netherlands and adjacent lowlands.
 KNNV Publishing,
- van Asch N. (2012) Environmental response to Lateglacial climate change-Reconstructions of temperature and
 vegetation changes in northwest Europe., Utrecht University
- van Geel B. (1972) Palynology of a section from the raised bog 'Wietmarscher moor', with special reference to
 fungal remains. Plant Biology 21:261-284
- van Geel B. (1978) A palaeoecological study of Holocene peat bog sections in Germany and the Netherlands,
 based on the analysis of pollen, spores and macro-and microscopic remains of fungi, algae,
- 842 cormophytes and animals. Review of Palaeobotany and Palynology 25:1-120
- van Geel B., Buurman J., Brinkkemper O., Schelvis J., Aptroot A., van Reenen G., Hakbijl T. (2003)
 Environmental reconstruction of a Roman Period settlement site in Uitgeest (The Netherlands), with
 special reference to coprophilous fungi. Journal of Archaeological Science 30:873-883
 doi:https://doi.org/10.1016/S0305-4403(02)00265-0
- Vinther B.M., Clausen H.B., Johnsen S.J., Rasmussen S.O., Andersen K.K., Buchardt S.L., Dahl-Jensen D.,
 Seierstad I.K., Siggaard-Andersen M.L., Steffensen J.P., Svensson A., Olsen J., Heinemeier J. (2006) A
 synchronized dating of three Greenland ice cores throughout the Holocene. Journal of Geophysical
 Research: Atmospheres 111 doi:10.1029/2005jd006921
- Vos P.C., Van Heeringen R.M. (1997) Holocene geology and occupation history of the Province of Zeeland
 Mededelingen Nederlands Instituut voor Toegepaste Geowetenschappen TNO 59:5-109
- Wanner H., Beer J., Bütikofer J., Crowley T.J., Cubasch U., Flückiger J., Goosse H., Grosjean M., Joos F.,
 Kaplan J.O., Küttel M., Müller S.A., Prentice I.C., Solomina O., Stocker T.F., Tarasov P., Wagner M.,
 Widmann M. (2008) Mid- to Late Holocene climate change: an overview. Quaternary Science Reviews
 27:1791-1828 doi:http://dx.doi.org/10.1016/j.quascirev.2008.06.013
- Wanner H., Solomina O., Grosjean M., Ritz S.P., Jetel M. (2011) Structure and origin of Holocene cold events.
 Quaternary Science Reviews 30:3109-3123
- Weeda E.J., Schaminée J.H.J., Van Duuren L. (2000) Atlas van Plantengemeenschappen in Nederland. deel 1:
 Wateren, moerassen en natte heiden. KNNV Uitgeverij, Utrecht
- Weeda E.J., Westra R., Westra C., Westra T. (1988) Nederlandse oecologische flora, wilde planten en hun
 relaties. deel 3. Hilversum
- Wiederholm T. (1983) Chironomidae of Holarctic region: keys and diagnoses. Part 1. Larvae Entomol Scand
 Suppl 19:1-457







Green gyttja Brown-green

gyttja

Brown gyttja











Figure 1 (a) Location of Lake Uddelermeer; (b) Lidar elevation model of the central Netherlands (AHN; www.ahn.nl) and (c) the area directly surrounding Lake Uddelermeer. Yellow star indicates location core. Numbers indicate sites discussed in the text. 1 Ice pushed ridge Garderen-Ermelo max. 50 m A.S.L.; 2 Ice pushed ridge Apeldoorn max. 100 m A.S.L.; 3 Lake Uddelemeer; 4 Lake Bleekemeer; 5 brook Leuvenemse beek; 6 medieval ringfort (ringwalburg) (Koomen & Maas, 2004; Heidinga, 1987). For visibility purposes (c) is plotted with a different colour range, only a relative elevation legend is shown for both maps.

Figure 2Bayesian modelled age-depth model of core section 50 - 650 cm depth for centralCore E from Lake Uddelermeer (figure adapted from Engels et al. (2016)). The 95% probabilitydistribution range is shown in grey, calibrated C¹⁴ ages and 95% probability ranges are shown asblack squares with error bars.

Figure 3 Percentage diagram of selected pollen, spores and NPPs from Lake Uddelermeer showing regional (A) and local (B) vegetation development in and around Lake Uddelermeer. All curves are shown with an additional fivefold exaggeration. Diagram plotted on sediment depth (cm) with a secondary scale (cal. yr. BP) plotted for comparison. Sedimentological and Loss-on-Ignition (represented as organic content% in 3A) profiles are plotted to the left of the pollen curves. The presence of charcoal (indicated with '+' present; '++' present in large quantities) is plotted to the right of the pollen curves.

Figure 4 Percentage diagram showing all chironomid taxa and the chironomid count sum (head capsules/ sample) for Lake Uddelermeer. Diagram plotted on sediment depth (cm) with a secondary scale (cal. yr. BP) plotted for comparison. The sedimentological profile is plotted to the left of the chironomid curves. To the right the scores of the chironomid samples on the first axis of a PCA performed with square-root-transformed percentage data is shown.

Figure 5 Selected percentage curves of pollen, NPPs and chironomid taxa as indicators for regional vegetation change, lake ecology and chironomid assemblage. Diagram plotted on sediment depth (cm) with a secondary scale (cal. yr. BP) plotted for comparison. The sedimentological profile is plotted to the left of the curves.

Figure 6 Chironomid inferred July air temperature (CI-T) reconstructions from Lake Uddelermeer with sample specific error estimates. Curves plotted on age (cal. yr. BP) with a secondary scale of sediment depth (cm) plotted for comparison. Warm (red) and cold (blue) periods for the period 2550 - 1140 cal. yr. BP are indicated; the part of the record which is not included in the reconstruction is shown in grey. To the right of the reconstruction, the distance of the fossil sample to the nearest modern analogues in the Norwegian/Alpine training set (squared chi²-distance (sq.x²d.): the vertical lines indicate the 2nd and 5th percentiles of all squared chi²-distances in the Norwegian/Alpine training set and are defined here as 'no close' and 'no good' analogues respectively (after Birks et al. 1990); the goodness-of-fit of the fossil samples to temperature (squared residual distance; sq.res.d): the vertical lines indicate the 90th and 95th percentiles in residual distances of the modern samples to the first axis in a constrained CCA and are defined here as 'poor' and 'very poor' fit respectively (e.g. Birks et al. 1990); and the cumulative percentage

(Frequency %) of fossil chironomids that are absent (%absent) from the modern calibration set are given

Figure 7 CI-T reconstructions for Lake Uddelermeer (this paper); Bigland Tarn (Barber et al. 2013) and Talkin Tarn (Langdon et al. 2004) plotted on a cal. yr. BP timescale. Calibrated AMS radiocarbon dates and their 2 sigma error ranges are plotted to the right of each temperature reconstruction.

Figure 8 Warm (red) and cold (blue) periods in NW European temperature reconstructions for the period 1000 - 2500 cal. yr. BP (grey is not identified/available). IA: Iron Age; RWP: Roman Warm Period; DACP: Dark Age Cold Period; MCA: Medieval Climate Anomaly. A: CI-July air temperature Uddelermeer (this paper); B: CI-July air temperature Bigland tarn UK(Barber et al. 2013) also included in overview record D; C:Tree ring inferred JJA temperature for the Alps (Büntgen et al. 2011) also included in overview records D, E and F; D:Compilation various archives from NW Europe summer - winter and annual temperature (Riechelmann and Gouw-Bouman 2019); E:Compilation various archives N Hemisphere summer - winter - annual temperature (Ljungqvist 2009); F: compilation various archives globally temperature (Helama et al. 2017a). For the records A and C zone boundaries were placed where the temperature curve crossed the mean. When available, age uncertainty (minimum and maximum of the age model) is plotted with black lines.