Environmental conditions at the Last Interglacial (Eemian) site Neumark-Nord 2, Germany inferred from stable isotope analysis of freshwater mollusc opercula

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9 Mollusc biogenic carbonates are valuable records of past environmental conditions. In particular, carbonate oxygen (δ^{18} O) and carbon (δ^{13} C) stable isotopes can be used to reconstruct different 10 11 physical and chemical parameters, according to the different genera used (marine, freshwater or 12 terrestrial). The Last Interglacial (early Eemian) palaeolake of Neumark-Nord 2 (NN2), Germany 13 provides an excellent example of a Neanderthal archaeological site with abundant freshwater 14 carbonate remains. As in other European contexts, one of the most abundant species is Bithvnia 15 tentaculata. In order to provide a robust regional baseline for the interpretation of the 16 archaeological data, this study includes a calibration phase on modern B. tentaculata opercula. The 17 results indicate that these calcitic structures are likely to be subjected to a growth 18 slowdown/cessation during summer, which influences their geochemistry, reflecting mainly the 19 water properties of the rest of the year. This modern calibration, together with the existing 20 palaeoenvironmental reconstructions developed for NN2 (e.g. pollen data), represents a valuable 21 opportunity to establish *B. tentaculata* opercula as reliable environmental proxies applicable to 22 several other freshwater contexts. The isotope data of the NN2 opercula, in agreement with the 23 pollen record, indicate that the major archaeological horizon was formed during a rather wet period and potentially in a semi-forested environment. However, human occupation occurred also during
drier phases at the site and within a wide temperature range, indicating the absence of restricted
environmental preferences by the local Neanderthal groups.

Keywords: Neanderthals; Last Interglacial; Middle Palaeolithic; Oxygen and carbon stable
isotopes; Bithynia tentaculata; Modern calibration

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Remains of freshwater molluscs, and in particular *B. tentaculata*, are abundant and well preserved throughout the Quaternary sedimentary record across Europe (Preece 2001). Recent advances in biomolecular techniques and biomineral understanding have resulted in the application of amino acid racemization analyses of these biominerals for dating (Penkman *et al.* 2008, 2011). However, only few studies have previously used these materials as palaeoenvironmental records (Filippi *et al.* 1997; Hammarlund *et al.* 1999; Anadón *et al.* 2006).

56 Given that the *B. tentaculata* habitat range includes closed and slow-running water bodies, 57 shells and opercula are often found in interglacial palaeolake infills (Anadón et al. 2006; Waghorne 58 et al. 2012; Milano & Szymanek 2019). Previous studies suggested that B. tentaculata opercula 59 constitute suitable environmental records and that their oxygen and carbon stable isotope 60 compositions are useful proxies for water temperatures and moisture regimes, respectively (Milano 61 & Szymanek 2019). For instance, oxygen incorporation by freshwater molluscs is influenced mainly by the water oxygen δ^{18} O and temperature (Anadón *et al.* 2006; Waghorne *et al.* 2012). 62 Furthermore, the carbonate δ^{13} C is influenced by the water DIC (dissolved inorganic carbon), 63 64 which, in turn, is controlled by water inflow and evaporation related to the moisture extent of the 65 environment (Leng & Marshall 2004 and references therein).

High-resolution climatic reconstructions of interglacial periods, and especially of the Last Interglacial, are valuable tools for the palaeoecological interpretation of Pleistocene humanenvironment interactions. Evidence of human activity during the Last Interglacial in central Europe has been attested to by the discovery of several sites in northern and central Germany such as Gröbern, Grabschütz, Weimar, Neumark-Nord, Rabutz, Lehringen and Stuttgart-Untertürkheim 71 (Roebroeks et al. 1992; Gaudzinki-Windheuser & Roebroeks 2011). Among them, Neumark-Nord 72 2 (NN2) stands out: the record includes not only abundant evidence for hominin activity 73 (Gaudzinski et al. 2014; Pop 2014), but it also contains several organic and inorganic materials 74 useful for palaeoenvironmental reconstructions, including pollen (Sier et al. 2011; Bakels 2014; 75 Wansa et al. 2014), sediment (Mücher 2014) and mammalian biominerals (Britton et al. 2019). 76 The aim of this study is to produce additional evidence for palaeoenvironmental conditions at NN2. 77 The abundance of *B. tentaculata* opercula throughout the *Hauptprofil* 7 (HP7) sequence, together with the existence of numerous other environmental proxies, represents a unique opportunity to 78 79 achieve such objective. In addition, the present study includes a modern calibration of B. tentaculata δ^{18} O and δ^{13} C based on a year-long collection and environmental monitoring at 80 81 Hundewasser (near Leipzig, Germany). This part of the study was designed to further refine our 82 understanding of the freshwater biogenic carbonates as environmental records.

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85 Study site: Neumark-Nord 2 (NN2)

86 Neumark-Nord is located ~ 35 km west of Leipzig in Saxony-Anhalt, Germany $(51^{\circ}19'28'' N,$ 87 11°53'56" E). The first excavations uncovered the Neumark-Nord 1 basin and subsequently, less 88 than 200 meters northeast, a second basin, Neumark-Nord 2, was discovered in 1996 (Fig. 1; 89 Gaudzinski-Windheuser et al. 2014). The NN2 basin infill revealed archaeological deposits related 90 to the Last Interglacial period, with the lake estimated to have started forming in the last part of 91 Saalian and to have lasted throughout the Eemian (Sier et al. 2011). In July 2007, the NN2 profile 92 HP7, located in the deep part of the infill, was sampled at 5-cm intervals with the deepest sample 93 at 595 cm from the top of the section. The lithostratigraphic succession was recorded by Mücher

94 (2014). The pollen record was documented along the entire HP7 profile and allowed a 95 reconstruction of the (local) vegetational change of the Eemian Interglacial (Sier et al. 2011; 96 Bakels 2014; Wansa et al. 2014). Numerous archaeological finds, including faunal and lithic 97 assemblages, were excavated from the margin of the basin in different units that are well correlated 98 to the HP7 profile. Based on multidisciplinary analyses of the finds, previous studies suggested 99 that the area around NN2 was used primarily for food processing and flint tool production (Kindler 100 et al. 2014; Pop 2014). Ongoing studies are focused on correlating the fine-grained archaeological 101 data from NN2 to the record obtained during rescue excavations of the neighbouring, much larger, 102 lake of Neumark-Nord 1 (e.g. Gaudzinski-Windheuser et al. 2018).

Along the NN2 HP7 stratigraphic succession freshwater molluscs dominated the mollusc fauna (Strahl *et al.* 2010; Kuijper 2014). Especially abundant were the species of *B. tentaculata* (used in this study), *Valvata piscinalis, Radix ovata, Anisus spirorbis, Gyraulus acronicus* and *Gyraulus crista*. The mollusc fauna indicated calm, shallow freshwater conditions (Kuijper 2014).

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109 Material and methods

110 Bithynia tentaculata and modern sampling

B. tentaculata (Linnaeus, 1758) is a prosobranch gastropod living in European slow-running rivers and lakes (Schäfer 1953; Frömming 1956). The snail has a short lifespan of about 1-3 years, with sexual maturity reached within the first 12 months of growth (Tashiro & Colman 1982; Richter 2001). It is characterized by a small shell (up to 13 mm) with five whorls and an upside-pointing operculum (Clarke 1981; Jokinen 1992; Welter-Schultes 2012). As for all molluses, the biomineralized structures are deposited throughout the animal's life. The shell is composed of
aragonite while the operculum is made of calcite (Filippi *et al.* 1997; Anadón *et al.* 2006).

For the modern component of this study, collection of live specimens was performed on a monthly basis for a period of one year from November 2017 to November 2018. Molluscs were collected from the small river Hundewasser in Lützschena near Leipzig, Germany (51°22'37.3"N, 12°16'50.7"E) ca. 30 km east of NN2 (Fig. 1). The molluscs (N = 39) were collected using small fishing nets at a depth of 50-100 cm. Juveniles and small specimens were avoided. On average, the opercula collected were 4.6 ± 0.2 mm long and 3.2 ± 0.1 mm wide.

The river is about 8 m wide and 1 m deep and it is characterized by extremely low current levels. Its banks are vegetated with abundant aquatic macrophytes. Water samples were collected and stored in 40-mL glass bottles with no headspace at ~ 4 °C until measurement. Air and water temperatures were measured during collection using a Voltcraft DT-300 thermometer.

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129 Sample preparation and analysis

The soft tissues of the modern molluscs were discarded immediately after collection. In order to remove foreign material, the carbonates were immersed in 3 vol % NaOCl for 2 hours and ultrasonically washed using deionized water for circa 3-5 minutes. After being air-dried for 24 hours, the opercula (N = 39) were individually crushed using an agate mortar and pestle.

The opercula from NN2 were bleached, washed and crushed following the same protocol as the modern specimens. A total of 107 opercula were analysed from the NN2 collection. On average, three opercula were selected from 41 5-cm mollusc-rich intervals in the HP7 stratigraphic succession. The analysed opercula came from units (and archaeological levels): 5, 6 (NN2/3), 7, 8 (NN2/2b), 9 (NN2/2a), 10, 11 (NN2/1c), 17 (NN2/1a) and 18. 139 The stable isotope analyses of the carbonates were performed at two laboratories, the Iso-140 Analytical laboratory in Crewe, United Kingdom and the Department of Human Evolution, Max 141 Planck Institute for Evolutionary Anthropology in Leipzig (henceforth MPI-EVA), Germany. At 142 the Iso-Analytical laboratory, carbonate powder samples were digested in He-flushed borosilicate 143 exetainers using a water-free phosphoric acid. The released CO₂ gas was then measured in 144 continuous flow mode with a Europa Scientific 20-20 gas source isotope ratio mass spectrometer 145 (CF-IRMS). At MPI-EVA, samples were digested at 70 °C and the CO₂ gas was then measured 146 with a ThermoFisher MAT 253 Plus IRMS linked to a Kiel IV automated carbonate preparation 147 device. At Iso-Analytical, stable isotope ratios were calibrated against an IA-R022 calibrated NBS-19 ($\delta^{13}C = +1.95\%$; $\delta^{18}O = -2.20\%$), whereas at MPI-EVA, data were calibrated against an IAEA-148 603 calibrated Carrara marble ($\delta^{18}O = -1.64\%$; $\delta^{13}C = +1.87\%$). Results are reported in per mil 149 150 (‰) relative to the Vienna Pee-Dee Belemnite (VPDB) standard. At Iso-Analytical, the average 151 precision error (1 σ ; computed from eight injections per sample) was better than 0.05% for both 152 δ^{18} O and δ^{13} C. At MPI-EVA, the average precision error (1 σ ; computed from eight measurements 153 per sample) was better than 0.05‰ for δ^{18} O and 0.03‰ for δ^{13} C, and the long-term accuracy based on IAEA-603 samples measured blind (n = 268) was better than 0.05‰ for δ^{18} O and 0.03‰ for 154 $\delta^{13}C$. 155

156 The oxygen stable isotope analyses of the water collected on monthly basis from Hundewasser 157 were performed at the Iso-Analytical laboratory in Crewe, United Kingdom. The samples were 158 measured by continuous flow isotope ratio mass spectrometry using a Europa Scientific ANCA-159 GSL and 20-20 IRMS. The analyses were performed in duplicate by the equilibration technique. 160 The samples were measured against three reference standards. The first standard being IA-R063 161 with δ^{18} O v-smow = -0.41‰, the second IA-R065 with δ^{18} O v-smow = -33.57‰ and the third IA- 162 R064 with $\delta^{18}O_{V-SMOW} = -12.34\%$. All three standards are traceable to the primary reference 163 standards V-SMOW2 (Vienna-Standard Mean Ocean Water) and V-SLAP2 (Vienna-Standard 164 Light Antarctic Precipitation) distributed by the IAEA. All the modern isotope and environmental 165 data can be found in the Supplementary Information (Table S1).

In order to test the water isotopic equilibrium deposition of *B. tentaculata* opercula, the measured δ^{18} O values were compared to the expected δ^{18} O calculated using the following fractionation equation:

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$$T = 15.70 - 4.36 (\delta^{18}O_c - \delta^{18}O_w) + 0.12 (\delta^{18}O_c - \delta^{18}O_w)^2$$
 (1)

170 (Hays & Grossman 1991) where T = water temperature; $\delta^{18}O_c$ = calcite $\delta^{18}O_i$, $\delta^{18}O_w$ = water $\delta^{18}O_i$. 171 For the fossil material from NN2, to standardize the dataset and to enhance long-term isotope 172 trends, departures ("deviations" or "anomalies") in the values of the single specimens from the 173 $\delta^{18}O$ and $\delta^{13}C$ overall averages of the sequence were calculated. The $\delta^{18}O$ and $\delta^{13}C$ 2-sigma 174 standard deviations within the monthly modern datasets (0.7‰ for both isotope ratios) were used 175 as significance thresholds in the interpretation of the archaeological results. The isotope data from 176 NN2 can be found in the Supplementary Information (Table S2).

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179 Results

180 Modern water and carbonate data

181 At Hundewasser, the seasonal water temperatures fluctuated between 0.6 and 21.7 °C, with a 182 similar variation for the recorded air temperatures (from -4.2 to 21.2 °C; Fig. 2A,B). In association 183 with these variations, the water $\delta^{18}O(\delta^{18}O_w)$ showed a variance range of 1.6‰, with more negative 184 values in winter and more positive ones during summer (Fig. 2B). The opercula showed relatively 185 narrow seasonal variances in their isotope values of 0.7% for δ^{18} O and 0.9% for δ^{13} C (Fig. 2C). Significant offsets occur between measured and expected δ^{18} O monthly values (calculated using 186 187 Eq. (1)), with rather flat data distributions compared to the expected value distributions (Fig. 3A). 188 The largest offsets occurred during the summer months of June, July and August, suggesting a 189 growth slowdown/cessation during this time of the year (Fig. 3A). The expected average δ^{18} O 190 value $(-5.9\pm1.4\%)$ using the annual dataset resulted lower than the measured value $(-5.4\pm0.2\%)$; 191 Fig. 3B). This offset decreased (from 0.5% to 0.3%) when the period of slow growth was excluded 192 from the average calculations, with an expected δ^{18} O value of $-5.2\pm1.0\%$ and a measured δ^{18} O 193 value of $-5.5\pm0.3\%$ (Fig. 3B). Likewise, the average reconstructed water temperature using the 194 whole annual dataset was calculated to be 7.7±1.8 °C, about 2 °C colder than the measured one 195 (9.8±7.4 °C; Fig. 3C). By excluding the period of slow growth, the reconstructed water temperature 196 $(7.4\pm1.9 \text{ °C})$ and the measured one $(6.7\pm5.1 \text{ °C})$ showed an offset of 0.7 °C.

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198 Oxygen isotopes of B. tentaculata opercula from NN2

Compared to the average opercula δ^{18} O values of modern specimens, the archaeological δ^{18} O 199 200 values were from 2.5% to 6.5% higher. Along the HP7 profile, the geochemical composition of 201 *B. tentaculata* opercula showed a certain degree of variation. The average δ^{18} O value varied among 202 the different stratigraphic units (Fig. 4A), with the most negative values recorded in units 6, 10 203 (respectively $-1.7\pm1.1\%$ and $-2.3\pm1.1\%$) and 18 (with only two specimens; $-2.9\pm2.0\%$) and the 204 most positive values recorded in units 5 and 17 (resp. $0.8\pm0.6\%$ and $0.9\pm1.9\%$). Data in units 6, 7, 8 and 17 recorded the largest δ^{18} O variances due to the large spread among the opercula (Fig. 205 4A). In unit 6, the high variance was due to the particularly low δ^{18} O values of the opercula at 545-206 207 550 cm and to the particularly high values of the specimens at 550-555 cm (Fig. 5A). In unit 7, the spread of the data was likely affected by the discontinuous sampling of opercula for the unit. Within unit 8, high δ^{18} O values occurred in the lower part but they decreased toward lower values in the central part of the unit and then they showed a tendency towards high values in the final parts of the unit (Fig. 5A). Furthermore, unit 17 showed a clear tendency toward lower values in the upper part (Fig. 5A).

When considering the deviations from the profile δ^{18} O average, some data resulted to be smaller 213 214 than the threshold of 0.7% (based on the 2σ standard deviations within the monthly modern 215 dataset). Only deviations above this threshold were considered significant for the 216 palaeoenvironmental reconstructions and therefore further discussed (Fig. 6A). Warm (positive) 217 anomalies were observed at the end of unit 5, at the beginning of unit 7 and 8 (NN2/2b) and, to a 218 lesser degree within unit 11 (NN2/1c). Exceptionally positive anomalies were observed at the 219 beginning of unit 17 (NN2/1a). Cold (negative) anomalies were observed at the end of unit 6 220 (NN2/3), 7, 8 (NN2/2b) and, to a lesser degree in unit 9, at the end of unit 10 and in the central 221 part of unit 11 (NN2/1c). A rather large negative anomaly was also observed at the beginning of 222 unit 18 (Fig. 6A).

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224 *Carbon isotopes of B. tentaculata opercula from NN2*

As with oxygen isotope composition, the archaeological δ^{13} C values were from 5.6‰ to 6.8‰ higher than the values of modern specimens. The average δ^{13} C values varied among the different units (Fig. 5B), with the most negative values recorded in units 8 and 10 (resp. -6.1±0.9‰ and -5.4±1.4‰) and the most positive values recorded in units 7, 11 and 17 (resp. -4.5±1.0‰, -4.1± 1.2‰ and -4.5±1.94‰). Likewise for δ^{18} O data, large δ^{13} C variances were recorded in units 8 and 17 (Fig. 5B). As for unit 8, the variance was primarily associated with exceptionally high δ^{13} C

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values at depth 430-435 cm, contrasting with the rest of the unit characterized by lower values. In unit 17, the δ^{13} C showed a tendency toward lower values in the uppermost parts. The same trend was visible within unit 9 (Fig. 5B).

234 When considering the significant deviations (>0.7‰) from the profile δ^{13} C average (Fig. 6B),

dry (positive) anomalies were observed at the end of unit 5, at the beginning of units 6 (NN2/3), 7

and 17 (NN2/1a), in unit 11 (NN2/1c). Wet (negative) anomalies were observed at the beginning

of units 5, 6 (NN2/3) and 7, through most of unit 8 (NN2/2b) and at the end of unit 17 (NN2/1a).

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240 Discussion

241 Modern calibration of B. tentaculata

242 The results of modern *B. tentaculata* specimens demonstrated the potential of the opercula δ^{18} O 243 values to be proxies for water temperature, with an error of less than 1 °C. Due to the small size of 244 the opercula, the sampling technique used did not ensure the right spatial resolution to resolve 245 seasonal isotopic fluctuations and therefore sub-annual temperature reconstructions. Other 246 analytical techniques, such as SIMS (Secondary Ion Mass Spectrometry) and SHRIMP (Sensitive 247 High Resolution Ion Microprobe), would be needed to achieve a higher temporal resolution of the 248 data. Although the δ^{18} O seasonal variability in the opercula was limited, it was possible to identify 249 a period of potential growth slowdown/cessation during the warmest months of summer (Fig. 3A). 250 Generally, in bivalves, the main shell growth coincides with the warm season (Richardson 2001). 251 However, several studies previously reported cases of summer growth cessation (Ramón 2003; 252 Milano et al. 2017). Waghorne et al. (2012) investigated the growth of B. tentaculata opercula 253 from southern Great Britain and they estimated the growing season to be between May and 254 October. The different growing season recorded here is likely related to the animal's exposure to 255 different environmental factors such as food availability, temperature and water conditions 256 (Witbaard et al. 1997). Furthermore, B. tentaculata is known to spawn between May and July 257 (Richter 2001). The observed slow-down in opercula formation can be a result of energy 258 reallocation from growth to reproduction, which was previously observed in several other mollusc species (Sato 1995; Purroy et al. 2018). Our results indicated that B. tentaculata δ^{18} O values 259 260 reflected well the water temperature throughout the year with the exception of about three months 261 during summer. When applied to palaeoenvironmental studies, this temporal window has to be 262 taken into account.

Although the modern data are rather promising, the application of the opercula δ^{18} O as 263 264 quantitative palaeothermometer for fossil remains has to be considered with caution. The 265 superficial δ^{18} O value of freshwater bodies is known to reflect the rainfall δ^{18} O value, which varies 266 primarily with latitude and altitude (Craig 1961; Bowen & Wilkinson, 2002). At a specific 267 location, the rainfall δ^{18} O is mainly controlled by air temperature and, in the tropical regions, by 268 the amount of precipitation (Dansgaard 1964). However, the size and type of water body significantly affects the relationship between the water δ^{18} O and rainfall δ^{18} O (Darling *et al.* 2003). 269 In fossil contexts, in which the water δ^{18} O value is not known, a quantitative temperature 270 271 reconstruction is not advisable, especially when applied to lacustrine environments. Open and 272 large lakes have different hydrological properties compared to small and closed ones, whose $\delta^{18}O$ 273 water can be significantly affected by evaporative processes (Darling *et al.* 2003). For this reason, 274 only qualitative temperature reconstructions are recommended (Waghorne et al. 2012). Our results 275 from NN2 confirmed that hydrological characteristics significantly influence water and carbonate 276 stable isotope signature. The δ^{18} O difference observed between modern and archaeological

277 specimens was too large to be simply explained by changes in water temperature, suggesting that 278 the stable isotope composition of the NN2 lake was rather different than the modern Hundewasser. A previous rainfall δ^{18} O value estimation based on the oxygen isotope analysis of *Equus* tooth 279 enamel from NN2 showed similar values to today's average, with δ^{18} O precipitation values 280 281 calculated from main find level NN2/2b being -9.1±1.1 ‰, for example (Britton et al. 2019). This 282 indicates that the differences observed in the opercula was likely related to fractionation processes 283 in the lake water itself. Considering the small size of the lake, NN2 might have been affected by 284 evaporative processes inducing the lake water and therefore the carbonates to have higher δ^{18} O 285 values. The discrepancy between the two types of environments (NN2 and Hundewasser) could 286 have been reduced by collecting modern specimens from a water body similar to NN2. However, 287 several field inspections and personal communications with local malacologists, revealed that the 288 modern distribution of *B. tentaculata* in Saxony is largely confined in small rivers and channels. 289 Therefore, a modern collection from lakes proved unfeasible.

290 As for the δ^{18} O values, the δ^{13} C modern variability was low (Fig. 2). Biogenic carbonate δ^{13} C 291 values in lacustrine environments are known to be mainly related to the δ^{13} C values of the water 292 DIC and therefore water inflow and evaporation processes (Milano & Szymanek 2019 and 293 references therein). In particular, when evaporation exceeds water inflow, the water DIC δ^{13} C is 294 more enriched, with consequential higher values in the biogenic carbonate δ^{13} C (Leng & Marshall 295 2004 and references therein). Previous works on *B. tentaculata* suggest that the δ^{13} C of this species 296 can be used as an evaporation/inflow proxy (Hammarlund et al. 2003; Anadón et al. 2006; Milano 297 & Szymanek 2019). Hundewasser is a flowing water body with percolation of groundwater 298 recharge. This, together with the small opercula size and consequent isotope signal averaging, 299 could explain the limited variability of the opercula δ^{13} C values throughout the year. On the other

300 hand, closed hydrological systems, especially when small, are significantly more sensitive to evaporative processes. As for oxygen, δ^{13} C values of specimens from NN2 were significantly 301 302 higher than the Hundewasser opercula, supporting the hypothesis that the local water isotope 303 composition would have been significantly different than modern running systems. However, 304 records from evaporation-sensitive settings such as NN2 might offer important insights into the 305 humidity variations at a local scale. For the δ^{18} O values, the larger sensitivity to evaporation may 306 bias the interpretation of the isotopic signal purely based on water temperature. Therefore, the 307 temperature cannot be reconstructed in a quantitative way and the results have to be considered 308 with caution.

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310 Palaeoenvironmental reconstruction of NN2

B. tentaculata opercula excavated from NN2 (HP7) indicated that, within the Eemian Interglacial, variations in the local environmental conditions occurred. Two important issues have to be considered when interpreting the opercula NN2 isotopic record: (i) the opercula, as all biogenic minerals, offer snapshots of past environmental conditions, and (ii) the environmental signal refers to the average annual conditions, summer excluded.

In the final phase of unit 5 (corresponding to the *Pinus-Betula* pollen assemblage zone PAZ II) an increase in temperature and a tendency toward more arid conditions occurred (Fig. 6). Micromorphological analysis of this laminated sedimentary unit indicated the presence of nonturbulent water during its colluvial deposition (Mücher 2014). These features, in agreement with our results, may relate to periods with less intense rainwash.

At the end of PAZ II, the abundance of *Pinus* increased and *Corylus* started to appear (Bakels 2014). In unit 6 (PAZ III), *Pinus* declined and *Quercus* started to dominate the vegetation. In addition, *Ilex* and various deciduous trees appeared, suggesting the occurrence of a maritime
temperate climate (Bakels 2014). However, the high percentage of herb pollen indicated that,
despite the oceanic climatic influence, the environment was still rather open (Fig. 6C; Pop *et al.*2015). The *B. tentaculata* record agreed well with the pollen, indicating a wet climate with periods
of more arid conditions (Fig. 6B). Temperatures showed a decrease toward the end unit 6 (Fig.
6A).

The few B. tentaculata data of unit 7 were not sufficient to achieve a robust environmental 329 330 interpretation during this phase. According to the pollen record, this part of the succession (end of 331 PAZ III and start of PAZ IV) began with a steady rise in *Corylus* and its subsequent dominance. 332 In general, PAZ IV was characterized by semi-open landscapes with rather scattered forests and 333 some steppe elements (Bakels 2014). During PAZ IVa1 Ulmus reached its maximum abundance 334 and algae of the genus Spyrogyra overgrew in the lake waters. Furthermore, the percentage of AP 335 (arboreal pollen) versus NAP (non-arboreal pollen) increased (Bakels 2014). This indicated that the climate became more humid, as supported by the decreasing trend in the opercula δ^{13} C values 336 337 throughout unit 7 into unit 8 (Fig. 6B). A similar trend was visible for the δ^{18} O values, suggesting 338 a decrease in temperature (Fig. 6A).

In the first part of unit 8, arboreal species were still dominant (Fig. 6C) and *B. tentaculata* recorded rather high temperatures and humid conditions. In the second part of unit 8, temperatures dropped and a dry event was recorded (only visible at depth 430-435 cm; Fig. 6 A,B). Simultaneously, *Corylus* decreased and non-arboreal vegetation became more abundant (Bakels 2014). It is possible that the temperature switch toward cooler conditions may have triggered the decline of the arboreal species. At the beginning of unit 9, the temperatures increased slightly and the humidity level decreased (Fig. 6A,B). From this unit onwards, laminated deposits were

346 replaced by massive deposits with occasional soil formation (Pop et al 2015). In this phase, the 347 abundance of Corylus remained low (Bakels 2014). Very few mollusc remains were retrieved from 348 PAZ IVb1 (second half of unit 9 and unit 10), suggesting a lake dry up event. Likewise, the shells 349 from unit 11 (PAZ IVb2) indicated that substantial aridity occurred during this period (Fig. 6B), 350 coupled with a rather variable temperature regime (Fig. 6A). Simultaneously, a drop in arboreal 351 vegetation (mainly Corvlus) was identified by the pollen analyses, together with an increase of 352 steppe-like vegetation (Bakels 2014). Compared to units without aquatic shell remains (e.g. units 353 12 to 16), unit 11 (and units with an aridity signal) indicates that the climate was rather dry but it 354 still allowed the basin to contain water. This hypothesis was also supported by the presence of 355 water plants, Lemna pollen and Spirogyra (Pop et al. 2015, Kuiper 2014). Unit 17 was characterized by the highest δ^{18} O values of the whole sequence, indicating that the temperatures 356 357 were significantly higher than before, especially in the lower portions corresponding to the last part of PAZ V. With the start of PAZ VI/VII (end of unit 17), the δ^{18} O values showed a significant 358 359 decrease, indicating the occurrence of a cooling trend. A similar tendency was observed in the δ^{13} C 360 values, going from less humid to more humid conditions at the end of unit 17 and beginning of 361 unit 18. The observed elevated temperatures during PAZ V agreed with the reconstruction based 362 on pollen by Kühl & Litt (2003) and the bulk carbonate δ^{18} O record from Gröbern (an Eemian lake 363 \sim 70 km north-east to NN2) by Litt et al. (1996). Indeed, these studies suggested that the 364 temperatures reached a maximum during PAZ IVb and they remained rather stable during PAZ V. 365 Interestingly, the drier conditions observed in the *Bithynia* record from unit 11 (PAZ IVb2) do 366 not seem to be reflected in the pollen sequence of NN1 (Neumark-Nord 1), the larger lake located 367 about one hundred meters away from NN2 (Seifert 1990). Previous studies hypothesized that the 368 absence of such environmental change at NN1 might have been related to either a very small-scale

369 local change or to a bias introduced by the small size of the NN2 lake (Bakels 2014). Indeed, small 370 lakes may catch pollen from a restricted area around the lake itself (Tauber 1967). Furthermore, droppings from animals coming to the lake might have altered the composition of the pollen, 371 372 enriching the consumed plant species (Bakels 2014). However, the aridity signal from the mollusc 373 opercula confirms that the pollen data were robust and the animal droppings did not significantly 374 affect the vegetation reconstruction. The absence of the Corvlus drop in NN1 might be related to 375 the different sizes of the two basins. Being a much smaller lake than NN1, NN2 may possibly have 376 been more sensitive to changes in temperature and humidity levels. Therefore, the interpretation 377 of pollen and opercula data from HP7 at NN2 might describe fairly localized environmental 378 conditions and its application to a broader regional level might be challenging.

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381 Implications for Neanderthal presence at NN2

382 Traces of human presence at NN2 were found from unit 6 to unit 17 (Fig. 6; Table 1; Gaudzinski-383 Windheuser et al. 2014; Pop 2014). Our reconstruction indicated that the local environmental 384 conditions changed within these phases, suggesting that Neanderthal groups were present at NN2 385 under varying temperature and humidity regimes. This correlates with the results from the oxygen 386 isotope analysis of horse tooth enamel from the main archaeological find levels in a recent study 387 (Britton et al. 2019). The major archaeological assemblage, with the highest concentration of finds, 388 was NN2/2b (unit 8; PAZ IVa2). The reconstructions based on *B. tentaculata* opercula indicated 389 NN2/2b was the most humid phase of the whole succession (Fig. 6B). High humidity potentially 390 supported the expansion of forested areas, suggesting that semi-closed environments may have 391 occurred during this period. In support of our data, the percentage of arboreal pollen, (especially

392 in the first half of unit 8), was particularly high compared to non-arboreal pollen (Bakels 2014). 393 This also correlates with evidence for niche partitioning between horses and bovids in this phase, 394 which suggested a mosaic and semi-forested environment (Britton et al. 2012). Besides NN2/2b, 395 less dense archaeological assemblages were found at various different stratigraphic depths (Fig. 6; 396 Table 1). The corresponding temperature estimations ranged from very warm (NN2/1a), cold 397 (NN2/3), average (NN2/2a) and variable (NN2/1c; Fig. 6A; Table 1). Similarly, the humidity range 398 was rather broad with very dry (NN2/1c), average (NN2/2a) or with variable conditions (NN2/3, 399 NN2/1a; Fig. 6B; Table 1). Virtually no B. tentaculata opercula were recovered from levels 400 NN2/1b and NN2/2c. Therefore, their respective palaeoenvironmental reconstructions are missing. 401 Our results suggested that Neanderthals were present both during the occurrence of relatively 402 dense forests, semi-open environments, as well as more open steppe-like landscapes around the 403 basin. Our data did not indicate any evident environmental preference that might have affected the 404 presence of Neanderthals during the Eemian Interglacial here, which is in agreement with similar 405 recent studies at Neumark (Britton et al. 2019). The character of the presence of Neanderthal 406 groups in central Europe during the Last Interglacial has been the topic of several previous studies 407 (Gamble 1987; Roebroeks et al. 1992; Gaudzinski 2002; Richter 2006; Wenzel 2007; Gaudzinski-408 Windheuser & Roebroeks 2011). In the past, some researchers argued that Neanderthals might 409 have had limited adaptive abilities to survive successfully in fully forested interglacial conditions 410 (Roebroeks & Tuffreau 1999). However, in Central Europe, Neanderthals show a high flexibility 411 in subsistence strategies in a variety of habitats (Gaudzinski-Windheuser & Roebroeks 2011 and 412 references therein; Gaudzinski-Windheuser et al. 2018), indicating that they were highly 413 responsive to climatic and environmental shifts (Gaudzinski 2002, 2004; see also Antoine et al.

414 2006, and Sier *et al.* 2015 for comparable evidence from north western France). These inferences415 are in line with the results obtained by this new proxy for palaeoenvironmental reconstructions.

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418 Conclusions

419 The present study presents the reconstruction of palaeoenvironmental conditions occurring during 420 the Last Interglacial at Neumark-Nord 2 (NN2), Germany. Oxygen and carbon stable isotopes of 421 B. tentaculata opercula were used as environmental proxies. Modern specimens were analysed for 422 calibration purposes and the results suggest that the opercula are generally formed throughout the 423 year, with a minor slowdown during summertime. Therefore, the water properties (i.e. temperature 424 and moisture regimes), encoded in the opercula bulk isotope signal in form of δ^{13} C and δ^{18} O values, 425 encompasses annual averages with potential exclusion of the warmest period. The isotope data 426 from the NN2 opercula indicate the local conditions during the Last Interglacial were characterized 427 by alternations between warm and cold conditions as well as successions of phases with different 428 humidity levels, although indicate that the absolute values measured were influenced by local 429 hydrological conditions in the lake. The major evidence of human activity, related to Neanderthals, 430 coincides with a humid phase and potentially with a semi-closed type of vegetation. However, 431 human presence was detected in six additional levels within NN2 which were characterized by 432 rather variable environmental conditions. In the light of the results presented in this study, and 433 other recent analyses, we conclude that the local Neanderthal groups in NN2 area did not manifest 434 a clear environmental preference, and instead were active across a wide range of temperature and 435 humidity conditions.

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- 626 Figure and table captions



Fig. 1 Location of the modern *B. tentaculata* collection site (Hundewasser: 51°22'37.3" N,
12°16'50.7" E) and the archaeological site of Neumark-Nord 2 (NN2: 51°19'28" N, 11°53'56" E).



631 **Fig. 2** A. Air temperatures (open circles) measured at Hundewasser and modelled monthly rainfall 632 δ^{18} O values (grey triangles and secondary y-axis) obtained from OIPC. B. Water temperature 633 (black circles) and δ^{18} O values (grey triangles and secondary y-axis) measured throughout the

annual collection in Hundewasser. C. Modern *B. tentaculata* opercula (N = 37) δ^{18} O (open circles)

and δ^{13} C values (black circles and secondary y-axis). Error bars indicate standard deviation.



Fig. 3 Modern *B. tentaculata* δ^{18} O data. A. Monthly opercula δ^{18} O values measured throughout 637 the annual collection in Hundewasser (open triangles) and the respective expected values (circles) 638 639 calculated using Eq. (1). The highlighted area represents the potential period of 640 cessation/slowdown of the opercula growth. B. Average measured and predicted values of the 641 opercula δ^{18} O calculated using the inverse function of Eq. (1). For this purpose, the calculations were carried out using both the whole annual dataset (i.e. 13 measured water δ^{18} O values and 642 643 temperatures) and the partial dataset excluding the period of slow growth (i.e. using 10 water and 644 temperature datapoints except the data from June, July and August). C. Average measured and 645 predicted (Eq. (1)) values of the water temperature using the whole annual dataset and the partial 646 dataset excluding the period of slow growth. Error bars indicate standard deviation.



Fig. 4 Overviews of the stable oxygen (A) and carbon isotopes (B) data obtained on *B. tentaculata*opercula from NN2, grouped by stratigraphic units (18 to 5).



Fig. 5 Raw stable oxygen (A) and carbon isotope (B) data obtained on *B. tentaculata* opercula throughout the NN2 stratigraphic sequence. The bars above the two graphs represent the subdivision of the succession in 14 units (18 to 5). The units in grey contain archaeological findings.



Fig. 6 Stable oxygen (A) and carbon isotope (B) anomalies throughout the different stratigraphic
units (18 to 5) and PAZ (Pollen Assemblage Zones; VII to II). Grey areas mark the significance
level of 0.7‰ representing the annual isotopic variability. Data outside these envelopes were

659 considered significant for the palaeoenvironmental interpretation. C. Pollen diagram from Pop &

Bakels (2015) showing the percentages of arboreal pollen (AP; using the three key taxa of *Pinus*,

661 *Corylus* and *Carpinus*) versus non-arboreal pollen (NAP; including grasses and sedges). *Q-U-C* =

662 *Quercus-Ulmus-Corylus*.

663

Table 1. Find levels at NN2 with corresponding palaeoenvironmental conditions reconstructed from *B. tentaculata* geochemistry record.

Find level	Corresponding unit	Finds m ⁻²	Total finds	Palaeoenvironmental reconstruction	
NN2/3	6	68	271	cold	variable
NN2/2c	7	11	2296	n/a	n/a
NN2/2b	8	255	125,322	warm than cold	very humid
NN2/2a	9	3	1501	average	average
NN2/1c	11	37	8811	variable	very dry
NN2/1b	13	17	1250	n/a	n/a
NN2/1a	17	1	4	very warm	dry than wet

664

665 Table S1. Modern environmental data and *B. tentaculata* isotope values during the annual
666 collection conducted at Hundewasser. SD = standard deviation.

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668

669 Table S2. B. tentaculata opercula isotope values throughout the HP7 profile in NN2. SD =

670 standard deviation.