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Abstract: Fluvial sedimentary archives have the potential to preserve a wide variety of palaeontological evidence, ranging from robust bones and teeth found in coarse gravel aggradations to delicate insect remains and plant macrofossils from fine-grained deposits. Over the last decade, advances in Quaternary biostratigraphy based on vertebrate and invertebrate fossils (primarily mammals and molluscs) have been made in many parts of the world, resulting in improved relative chronologies for fluviatile sequences. Complementary fossil groups, such as insects, ostracods and plant macrofossils, are also increasingly used in multiproxy palaeoclimatic and palaeoenvironmental reconstructions, allowing direct comparison of the climates and environments that prevailed at different times across widely separated regions. This paper reviews these topics on a regional basis, with an emphasis on the latest published information, and represents an update to the 2007 review compiled by the FLAG-inspired IGCP 449 biostratigraphy subgroup. Disparities in the level of detail available for different regions can largely be attributed to varying potential for preservation of fossil material, which is especially poor in areas of non-calcareous bedrock, but to some extent also reflect research priorities in different parts of the world. Recognition of the value of biostratigraphical and palaeoclimatic frameworks, which have been refined over many decades in the 'core regions' for such research (particularly for the late Middle and Late Pleistocene of NW Europe), has focussed attention on the need to accumulate similar palaeontological datasets in areas lacking such long research histories. Although the emerging datasets from these understudied regions currently allow only tentative conclusions to be drawn, they represent an important stage in the development of independent biostratigraphical and palaeoenvironmental schemes, which can then be compared and contrasted.

1 **QSR Special Issue: 20 years of Fluvial Archives Group** 

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Fossils from Quaternary fluvial archives: sources of 3

- biostratigraphical, biogeographical and palaeoclimatic evidence 4
- 5

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#### 15 Abstract

16 Fluvial sedimentary archives have the potential to preserve a wide variety of 17 palaeontological evidence, ranging from robust bones and teeth found in coarse gravel 18 aggradations to delicate insect remains and plant macrofossils from fine-grained 19 deposits. Over the last decade, advances in Quaternary biostratigraphy based on 20 vertebrate and invertebrate fossils (primarily mammals and molluscs) have been made 21 in many parts of the world, resulting in improved relative chronologies for fluviatile 22 sequences. Complementary fossil groups, such as insects, ostracods and plant 23 macrofossils, are also increasingly used in multi-proxy palaeoclimatic and 24 palaeoenvironmental reconstructions, allowing direct comparison of the climates and 25 environments that prevailed at different times across widely separated regions. This 26 paper reviews these topics on a regional basis, with an emphasis on the latest 27 published information, and represents an update to the 2007 review compiled by the 28 FLAG-inspired IGCP 449 biostratigraphy subgroup. Disparities in the level of detail 29 available for different regions can largely be attributed to varying potential for 30 preservation of fossil material, which is especially poor in areas of non-calcareous 31 bedrock, but to some extent also reflect research priorities in different parts of the 32 world. Recognition of the value of biostratigraphical and palaeoclimatic frameworks, 33 which have been refined over many decades in the 'core regions' for such research 34 (particularly for the late Middle and Late Pleistocene of NW Europe), has focussed 35 attention on the need to accumulate similar palaeontological datasets in areas lacking 36 such long research histories. Although the emerging datasets from these understudied 37 regions currently allow only tentative conclusions to be drawn, they represent an 38 important stage in the development of independent biostratigraphical and 39 palaeoenvironmental schemes, which can then be compared and contrasted. 40

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# 45 **Figures:**

46 Fig. 1 Map showing regions, major river catchments and key fossil sites mentioned in47 the text.

48 Fig. 2 Idealized tranverse section through the Lower Thames terrace sequence, east of

49 London, including details of Mammal Assemblage Zones (after Bridgland and

50 Schreve, 2001), updated to include biostratigraphically significant invertebrate

- 51 species and archaeological data.
- 52 Fig. 3 Correlation of Hoxnian fluvial sequences in the Lower Thames with the

53 stratotype at Hoxne (after White et al., 2013). The occurrence of the 'Rhenish' suite of

54 freshwater molluscs is critical to linking the Swanscombe and Clacton sequences and

55 thence to the pollen stratigraphy of the type-Hoxnian (sub-stages labelled). Units for

56 which there is clear palaeontological or sedimentological evidence for the prevailing

- 57 climate are colour-coded, and tentative correlations with the marine oxygen isotope
- 58 stages (MIS) of the deep-sea record are also shown.

59 **Fig. 4** Idealized cross-sections through the Middle Trent, Lower Trent and Witham

60 terrace sequences (modified from Bridgland et al., 2015 and White et al., 2017),

61 showing MIS correlation and biostratigraphically significant species.

62 **Fig. 5** Schematic diagrams showing the progressive appearance of forest land snail

63 species during three interglacials in northern France: La Celle (MIS 11), Caours (MIS

5e) and St Germain-le-Vasson (MIS 1). Modified from Limondin-Lozouet and Preece(2014).

66 **Fig. 6** Synthetic curves showing the development of ecological groups of molluscs

67 during MIS 11 at the site of La Celle, northern France, and a comparison with

68 malacological successions from other western European MIS 11 tufa sequences. The

69 appearances of critical species at La Celle that permit correlation with other

70 sequences are highlighted (after Limondin-Lozouet et al., 2015)

Fig. 7 Biostratigraphical scheme for Eastern European mammalian faunas (after
 Markova, 2007).

73 **Fig. 8** Correlation chart showing faunal changes, the hominin fossil record and

- significant technological changes in the Indian subcontinent, mainland and insular SE
   Asia (after Mishra et al., 2010).
- 76 Fig. 9 Pleistocene chronology of the Great American Biotic Interchange (GABI),
- showing details of major faunal exchanges between North and South America that

began at around 2.8 Ma (modified from Woodburne, 2010). Marine oxygen isotope

- 79 stages after Lisiecki and Raymo (2005).
- 80 **Fig. 10** Biostratigraphical significance of four species of glyptodont
- 81 (*Neosclerocalyptus*) in the Pleistocene of South America (modified from Zurita et al.,
  82 2009a).
- **Fig. 11** Schematic diagram showing temporal occurrence of megafaunal taxa at
- 84 Darling Downs, Australia, in relation to the widely-accepted hypothetical megafaunal
- 85 extinction 'window' and the approximate timing of human arrival on the continent
- 86 (after Price et al., 2011).

### 87 **1. Introduction**

88 The study of fossil assemblages recovered from fluvial deposits is well-established as 89 an important element of multi-disciplinary Quaternary research, providing a basis for 90 regional relative dating frameworks (e.g. Gliozzi et al., 1997; Schreve, 2001a; 91 Schreve and Bridgland, 2002; Bridgland et al., 2004; Markova, 2007; Megirian et al., 92 2010) and for detailed palaeoclimatic and palaeoenvironmental analyses (e.g. Coope, 93 2010; Schreve and Candy, 2010; Candy et al., 2010, 2015, 2016; Kahlke et al., 2011; 94 Limondin-Lozouet et al., 2010; Rule et al., 2012; White et al., 2013). The 95 contribution of the Fluvial Archives Group (FLAG) to these research areas over the 96 last 20 years has been considerable: two UNESCO-sponsored International 97 Geological Correlation Programme (IGCP) projects, entitled 'Global correlation of 98 Late Cenozoic fluvial deposits' (IGCP 449) and 'Fluvial sequences as evidence for 99 landscape and climatic evolution in the Late Cenozoic' (IGCP 518), ran under the 100 auspices of FLAG between 2000 and 2007 (Bridgland et al., 2007; Westaway et al., 101 2009). The former included a thematic biostratigraphy subgroup, which compiled 102 data on faunal assemblages from fluvial sequences in different parts of the world, 103 resulting in the publication of a review of progress in faunal correlation of Late 104 Cenozoic fluvial sequences (Schreve et al., 2007); following the conclusion of these 105 IGCP projects, data has continued to accumulate as part of the ongoing efforts of 106 members of FLAG (Cordier et al., 2017).

107

108 This paper provides updated reviews, organized on a regional basis, of advances in 109 Pleistocene vertebrate and invertebrate biostratigraphy made during the 20-year life of

110 FLAG; as such, it is intended to be a companion to the report of the IGCP 449

111 biostratigraphy subgroup (Schreve et al., 2007), which remains a benchmark review

112 of biostratigraphical frameworks derived from fluvial archives around the world. As

- 113 was the case with that paper, much of the most detailed work has been undertaken in
- regions such as NW and Central Europe, which have enjoyed long traditions of

115 Pleistocene palaeontological research, particularly for the late Middle and Late

Pleistocene (Fig. 1). This review also extends its scope beyond that of its predecessors to include considerations of the palaeoenvironmental, palaeoecological

and biogeographical information that can be derived from fossil assemblages, and the

- potential for Pleistocene fluvial archives to enhance knowledge of long-termQuaternary climate change.
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# 122 Insert Figure 1 hereabouts

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124 Fluvial archives have several advantages over other types of terrestrial sedimentary 125 sequences that commonly preserve fossils (such as those from lake basins or caves), 126 the most significant of these being the potential chrono-stratigraphical control provided by river terraces, which have great value as regional templates for the 127 128 terrestrial Quaternary record (Bridgland, 2000, 2006; Antoine et al., 2007; Bridgland 129 and Westaway, 2008a, 2014; Bridgland et al., 2004, 2006, 2017; Mishra et al., 2007; 130 Vandenberghe, 2015). The nature of rivers also means that Pleistocene fluvial 131 sequences have the potential to preserve a variety of plant and animal fossils derived 132 from terrestrial, freshwater and estuarine environments, recording concurrent changes 133 in a diverse range of palaeoenvironmental settings. Lacustrine sequences, although 134 undoubtedly sources of more continuous and higher-resolution longer-timescale 135 records than rivers, usually represent only the deepest part of a lake basin from which 136 the longest sediment sequences can be obtained; such records are therefore often

137 lacking in fossils indicative of local terrestrial environments (with the notable 138 exception of wind-borne pollen) and are (usually) isolated from marine influences. 139 Conversely, cave sequences, which are often significant repositories of vertebrate 140 fossils (particularly mammals), usually lack pollen and invertebrate assemblages. 141 Fluvial archives therefore have an unrivalled capacity to provide insights into 142 changing Quaternary climates and environments through time, due to the wide range 143 of fossil types available (which include pollen, plant macrofossils, mammals, 144 molluses, ostracods and insects), coupled with the chronological control provided by 145 river terraces. Together, these proxies can be used to generate local biostratigraphical 146 frameworks, which can then be extended into other regions and into different 147 depositional contexts outside river valleys. Moreover, where fluvial sequences interdigitate with other sediment types, such as glacigenic or estuarine-marine 148 149 deposits, biostratigraphical evidence can be used to constrain the age of regional 150 glacial and sea-level histories (e.g. Bridgland et al., 1999; Matoshko et al., 2004; 151 White et al., 2010, 2013, 2017; Vis et al., 2010), which can then provide further 152 stratigraphical ('event stratigraphy') markers. It should be noted, of course, that this potential can only be realized in areas conducive to the preservation of fossils, and 153 154 that fluvial sequences are subjected to the same taphonomic complications known to 155 affect other Quaternary sedimentary archives.

156

157 Much of the recent palaeontological research summarized here has been undertaken 158 as part of multidisciplinary projects, often driven by archaeologically-motivated 159 research questions (see Chauhan et al., 2017). In Europe and Asia, these have often 160 focussed on the timing of hominin occupations relative to changing Pleistocene 161 climate and environments (e.g. Parfitt et al., 2005, 2010; Antoine et al., 2015, 2016; 162 Maddy et al., 2015; Peretto et al., 2015), whereas further afield, in regions such as 163 North America and Australia, attention has been more focussed on potential human 164 impacts, particularly on megafaunal populations (e.g. Prideaux et al., 2010; Prescott et al., 2012; Sandom et al., 2014; Cooper et al., 2015; Stuart, 2015). The potential for 165 fossils to provide important chronological and palaeoenvironmental frameworks 166 167 within which to interpret the archaeological record has also inspired research in 168 (initially) less promising regions of the world that have previously received little 169 attention, such as the deserts of the Arabian Peninsula (e.g. Groucutt et al., 2015; 170 Stimpson et al., 2015, 2016). The resulting palaeontological data have the potential to 171 be interpreted on a variety of scales, ranging from considerations of changing climatic 172 conditions during a single interglacial in a given region (e.g. Candy et al., 2016) to 173 comparisons between different interglacials (e.g. Limondin-Lozouet and Preece, 174 2014) and even identification of patterns at the marine oxygen isotope substage level 175 (Schreve, 2001b; Candy and Schreve, 2007; Ashton et al., 2008; White et al., 2013). 176 Key to such studies have been important advances in the application of independent 177 geochronological methodologies to Quaternary fluvial archives (reviewed in detail by 178 Rixhon et al., 2017). Advances in radiometric dating techniques, such as optically 179 stimulated luminescence (OSL), electron spin resonance (ESR) and uranium-series 180 have provided crucial chronological control (e.g., Rittenour, 2008; Kock et al., 2009; Voinchet et al., 2015;). In tectonically-active areas, the presence of interbedded 181 182 volcanic deposits can provide further opportunities for constraining the age of the 183 fluvial sediments through the application of argon-argon dating (e.g. Maddy et al., 184 2012; Marra et al., 2016). In addition, amino-acid racemization dating (AAR) based 185 on the calcitic opercula of the freshwater gastropod genus Bithynia has been shown to 186 be more reliable than previous AAR methodologies, allowing the development of a

robust aminostratigraphy for the British Quaternary with potential for application in
other regions (Penkman et al., 2007, 2011, 2013; cf. Westaway, 2009).

189

# 190 2. NW and Central Europe

191 A substantial body of research has been undertaken over the last decade in this 192 important region, particularly in southern Britain and northern France. This is partly a 193 feature of preservation potential; the unequal global distribution of fossiliferous 194 fluvial sediments corresponds closely with that of calcareous bedrock outcrops, which 195 promote fossil preservation, limiting detailed biostratigraphical research to regions 196 dominated by limestone and chalk. Calcareous tufas have provided an important 197 additional source of data (Dabkowski, 2014), albeit from sediment sequences usually 198 outside major fluvial systems. Tufa sequences have been recorded across Europe 199 (Pentecost, 1995; Sancho et al., 2015), but are particularly well represented in 200 northern France and Britain, where they have mainly been attributed to MIS 11, MIS 201 5e and the Holocene (Dabowski, 2014; Limondin-Lozouet and Preece, 2014). The 202 Triassic Muschelkalk outcrop in Germany also gives rise to notable occurrences of 203 calcareous spring deposits formed on former subaerial floodplains, although these are 204 generally more lithified than their Ango-French counterparts and have thus generally 205 been termed travertines. Important multiple travertines are interbedded with the 206 terrace deposits of the Ilm at Weimar (Schreve and Bridgland, 2002) and the Wipper 207 at Bilzingsleben (Mania, 1995). Although tufa and travertine deposits tend to be 208 highly localized, they can preserve fossils that rarely survive in other fluvial 209 sedimentary settings; this wide variety of palaeontological data is therefore 210 particularly suitable for both biostratigraphy and palaeoclimatic reconstructions. As 211 well as molluscs and vertebrates, plant remains (sometimes in the form of imprints of 212 rapidly-encrusted leaves or fruits) are also common, although pollen is rarely well preserved (Dabkowski, 2014). The calcareous nature of tufas and travertines also 213 214 means that they are suitable for a range of geochemical analyses (e.g. Dabkowski et 215 al. 2012, 2015), and appropriate for radiometric dating using U-series techniques (e.g. Candy and Schreve, 2007; Sierralta et al., 2010). 216

- 217
- 218 2.1 Britain

219 Ouaternary palaeontology in Britain has benefitted from government funding of 220 multidisciplinary research projects through bodies such as English Heritage (now 221 Historic England) and schemes such as the Aggregates Levy Sustainability Fund 222 (ALSF). The English Rivers Palaeolithic Survey (TERPS), which commenced in 223 1991, resulted in an important baseline archive for Pleistocene archaeological 224 research (Wymer, 1999), but provided only basic information pertaining to the 225 palaeontological evidence that accompanied Palaeolithic assemblages. More detailed 226 reviews of regional Pleistocene fossil records were subsequently provided by projects 227 funded by the ALSF, such as the Trent Valley Palaeolithic Project (TVPP), which 228 conducted an exhaustive review of both the palaeontological and archaeological records of the English Midlands (Schreve, 2007; Bridgland et al., 2014), and the 229 230 Medway Valley Palaeolithic Project (MVPP), which conducted similar research in 231 Kent (Briant et al., 2012; Chauhan et al, this volume). These projects were able to 232 study fossiliferous localities that lacked archaeological material, adding valuable 233 palaeontological and palaeoclimatic data to the underlying TERPS dataset. An 234 overview of all the ALSF-funded projects was provided by White (2016). Additional 235 data was accumulated during the lifetime of the Ancient Human Occupation of

Britain Project (AHOB), funded by the Leverhulme Trust, which underook research at
 numerous fluvial localities (Ashton et al., 2011; see http://ahobproject.org/database).

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239 The data accumulated by these research projects have continued to test and reinforce 240 the biostratigraphical model developed by Schreve (2001a; Fig. 2), which proposed a succession of Mammal Assemblage-Zones (MAZs) representing post-Anglian (MIS 241 242 12) interglacials. An important aspect of recent research has been the recognition of 243 greater climatic complexity recorded in terrestrial interglacial sequences, leading to a 244 renewed appreciation of the potential ability of multi-proxy palaeontological analyses 245 to discern marine oxygen isotope substages (e.g. Schreve, 2001b; Ashton et al., 2008; 246 Koutsodendris et al., 2010, 2011; White et al., 2013; Candy et al., 2014, 2016). In 247 Britain and northern Europe, this has been especially relevant to sequences attributed to the MIS 11 Hoxnian/Holsteinian Interglacial (e.g. Preece et al., 2007; Ashton et al., 248 249 2008; Candy et al., 2014), which is particularly well represented in the terrestrial 250 record.

251

# 252 2.1.1 The Lower Thames

253 The terrace deposits of the River Thames, particularly in its lower reach, represent 254 one of the most important terrestrial archives of the Middle and Late Pleistocene in 255 the world. In addition to key MIS 9 and MIS 7 sites published earlier in the lifetime 256 of FLAG (Schreve et al., 2007 and references therein), new data have been obtained 257 from several important Lower Thames localities over the last decade, including several MIS 11 sites: Dierden's Pit, Swanscombe (White et al., 2013), Southfleet 258 259 Road, Swanscombe (Wenban-Smith, 2013), East Hyde, Tillingham (Roe, 2001; White, 2012) and Clacton-on-Sea (White, 2012). Further work has also been 260 undertaken on the MIS 9 sequence at Purfleet (Bridgland et al., 2013) and other MIS 261 262 9 localities downstream in eastern Essex (Roe and Preece, 2011; Roe et al., 2009, 263 2011), providing further insight into the differentiation of MIS 11 and MIS 9 in the 264 British terrestrial record (Bridgland et al., 2001; Thomas, 2001; Roe et al., 2009). Over the wider Thames valley, mammalian fossils have assisted with the correlation 265 266 of the Upper and Middle Thames terraces (Bridgland and Schreve, 2009).

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269

# 268 Insert Figure 2 hereabouts

270 At Swanscombe, the biostratigraphical and palaeogeographical significance of the 271 'Rhenish' suite of freshwater molluscs, an important feature of the MIS 11 faunal 272 succession in the Lower Thames, has been firmly established by new evidence from 273 Dierden's Pit (White et al., 2013). The Swanscombe sequence, together with its 274 downstream correlatives at East Hyde and Clacton, indicates that the six 'Rhenish' 275 species did not colonize the Thames simultaneously, but appeared in a distinctive 276 sequence that can be tied to particular stages in the development of the vegetation. At 277 both Swanscombe and Clacton, 'Rhenish' taxa are largely absent from the earliest 278 fluvial deposits, with the pioneer species Pisidium clessini and Theodoxus danubialis 279 first appearing during pollen zone Ho II. These, together with *Belgrandia marginata*, 280 become established at the onset of Ho III, just before the first evidence for estuarine 281 conditions (in the form of brackish indicator species) appears at both sites. They are 282 followed by Corbicula fluminalis, which appears during Ho IIIb, and Borysthenia 283 naticina, which is absent from Clacton but appears in significant numbers slightly 284 later than Corbicula at both Swanscombe and East Hyde (White et al., 2013). The 285 timing of the first appearance of the final 'Rhenish' species, *Viviparus diluvianus*, is

286 less clear-cut, since it is a rare component of the Swanscombe fauna (Kerney, 1971) 287 and is only represented by derived shells at Clacton. However, at East Hyde it 288 appears to be present from early in the sequence, suggesting it was established in the 289 lower Thames by at least pollen zone Ho IIIa (White et al., 2013). It has therefore 290 been possible, on the basis of multiple strands of fossil evidence, to correlate MIS 11 291 fluvial sequences in the Lower Thames and to establish the timing of sea-level change 292 and the migration of the estuarine environment during that interglacial (White et al., 293 2013; Fig. 3). This has shown that sea-levels rose relatively late in the interglacial, 294 during Ho III, with a significant period during which the Thames was confluent with 295 continental rivers such as the Scheldt; this was presumably also true of other fluvial 296 systems flowing into the southern North Sea basin. This palaeogeographical situation 297 allowed distinctive groups of invertebrate and vertebrate species (including humans) 298 to colonize Britain (White et al., 2013).

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# 300 Insert Figure 3 hereabouts

302 Other significant recent advances in knowledge of the Lower Thames sequence have 303 arisen from work undertaken as part of developer-funded archaeological appraisal, 304 particularly that related to construction of the high-speed railway connection to the Channel Tunnel (e.g. Bridgland et al., 2013; Wenban-Smith 2013) and within the 305 306 remit of work funded by the ALSF. The construction of the Channel Tunnel Rail 307 Link led to the unexpected discovery of an elephant-butchery site, preserved within 308 deposits of a south-bank Thames tributary at Southfleet Road, Swanscombe (Wenban-309 Smith et al., 2006; Wenban-Smith, 2013). The carcass of an extinct straight-tusked 310 elephant (Palaeoloxodon antiquus) was surrounded by an undisturbed scatter of flint 311 tools (Wenban-Smith, 2015). Fossil assemblages recovered from the Southfleet Road 312 sequence included vertebrates, molluscs, ostracods and pollen; the absence of 313 'Rhenish' molluscs indicates that the sequence pre-dates the Middle Gravels at 314 Swanscombe and is therefore probably equivalent to the Lower Loam at Barnfield Pit, a conclusion consistent with the presence of Clactonian artefacts (Wenban-Smith, 315 316 2013). The vertebrate assemblage is strikingly similar to those from Swanscombe 317 (Basal Gravel-Lower Loam) and Clacton, both of which have been assigned to the Swanscombe MAZ (Schreve, 2001a). 318

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Work at the important MIS 9 sequence at Purfleet, undertaken as part of various developer-funded projects, allowed assessment of the palaeontological assemblages from the site, together with new analyses of stable isotopes and geochronology (Bridgland et al., 2013). This work confirmed the attribution of the interglacial to

324 MIS 9 through the application of OSL and AAR and contributed new

- 325 palaeoecological information. The MIS 9 interglacial has also been studied at various
- 326 sites downstream in eastern Essex (Roe and Preece, 2011; Roe et al., 2011), allowing
- enhancement of knowledge of the evolution of the highstand MIS 9e estuary in theLower Thames valley. Further south, the MVPP produced amino acid racemization
- data from several fossiliferous localities in eastern Essex, including the MIS 11
- 330 sequences at Clacton-on-Sea, East Hyde, Bradwell Hall, the MIS 9 sequences at
- 331 Shoeburyness, Cudmore Grove, East Wick and the Last Interglacial (MIS 5e) site at
- 332 East Mersea. The famous Levallois site at Baker's Hole, located in the Ebbsfleet
- 333 Valley, NW Kent, has also been the subject of recent research following the
- 334 construction of the Channel Tunnel Rail Link and Ebbsfleet International station
- 335 (Scott et al., 2010; Wenban-Smith, 2014). This locality, dated to MIS 7, represents the

- last surviving remnant of what was (before quarrying) a wide area containing
- 337 Levalloisian lithic remains in undisturbed primary context, associated with
- assemblages of large mammals, molluscs, small vertebrates and ostracods; publication
- 339 of detailed biostratigraphical and palaeoenvironmental analyses is in progress.
- 340

# 341 2.1.2 The River Trent and its tributaries

342 The Trent, Britain's third longest river, had until recently received considerably less 343 attention than other British fluvial systems. Significant new insights into the origin 344 and evolution of the Trent system have resulted from the Trent Valley Palaeolithic 345 Project (TVPP), which conducted an exhaustive review of the geological, 346 palaeontological and archaeological records of the English Midlands (Howard et al., 347 2007; White et al., 2007, 2010; Bridgland et al., 2014, 2015). The dating of the Trent 348 terraces has been underpinned by biostratigraphical data, together with 349 aminostratigraphic and OSL dating programmes and uplift modelling (Penkman et al., 350 2011, 2013; Bridgland et al., 2014; Westaway et al., 2015; Westaway, 2017). 351 Pleistocene fossil assemblages from the Trent catchment are both less common and 352 less well known than equivalent records from the aforementioned Lower Thames sites 353 and adjacent catchments, such as the Warwickshire-Worcestershire Avon. The upper 354 and middle reaches of the Trent include few calcareous rocks, fossils being preserved 355 there only under exceptional circumstances, such as within the Allenton Terrace 356 deposits south-west of Derby (Bridgland et al., 2014) and the floodplain gravels at Whitemoor Haye (Schreve et al., 2013). Both of these sites are located at the 357 confluences of tributary rivers with the main Trent. At Allenton, calcareous gravels 358 359 enriched by Carboniferous limestone clasts from upstream in the tributary Derwent 360 valley are responsible for the localized preservation of vertebrate assemblages that include hippopotamus, on the basis of which these deposits have been attributed to 361 362 MIS 5e (Bridgland et al., 2014). The younger Whitemoor Haye locality, at the Tame-363 Trent confluence, is situated within low-lying 'floodplain terrace' deposits, dated by 364 radiocarbon and OSL to around 41–43 k cal a BP, placing them within the Middle 365 Devensian (MIS 3; Schreve et al., 2013). This site is prone to waterlogging, another 366 effective means of preserving organic remains; palaeotemperature reconstructions, 367 based on beetle and chironomid assemblages, have indicated mean July temperatures 368 of +8 to +11 °C and mean January temperatures of between -22 and -16 °C (Schreve 369 et al., 2013).

370 The most significant fossil assemblages from the Trent catchment have been 371 recovered from its lower reach, where numerous sites have now been recorded in the 372 Balderton-Southrey Terrace of the Trent and in the Fulbeck and Tattershall terraces 373 of its tributaries, the Witham and Bain, respectively (Bridgland et al., 2014, 2015; Fig. 374 4). A significant aspect of the late Middle and Late Pleistocene record in the Trent is the absence of deposits belonging to the MIS 11 and MIS 9 interglacials, which are 375 376 well represented in fluvial systems further south, most notably the Thames. This has 377 been attributed to extensive glaciation of the English Midlands during MIS 8 (White 378 et al., 2010, 2017; Bridgland et al., 2014). The oldest fossiliferous sediments known 379 from the Trent system are therefore those preserved within the Balderton-Southrey 380 terrace and its equivalents in the Witham and Bain valleys (Brandon and Sumbler, 381 1988, 1991; Schreve, 2007; Bridgland et al., 2014). The Balderton Terrace of the 382 Lower Trent is predominantly a cold-climate aggradation, from which characteristic 383 fossils such as Mammuthus primigenius (woolly mammoth), Coelodonta antiquitatis 384 (woolly rhinoceros) and Ovibos moschatus (musk ox) have been recovered (Brandon

and Sumbler, 1988, 1991; Bridgland et al., 2014). Earlier studies of these vertebrate 385 386 assemblages also recognised the presence of interglacial species (Brandon and Sumbler, 1988, 1991; Lister and Brandon, 1991), which were subsequently 387 established to have been derived from highly fossiliferous fine-grained deposits 388 389 preserved low down in the sequence, typically filling channels cut into the Lias Clay 390 bedrock or the lowermost part of the Balderton Formation (Bridgland et al., 2014). 391 The fossil assemblages from these basal channels are representative of interglacial 392 conditions and include molluscs, ostracods, small vertebrates, pollen, plant 393 macrofossils and insects (White et al., 2007; Bridgland et al., 2014). The best record 394 to date has come from Norton Bottoms, where significant exposures of fine-grained 395 organic-rich sediments were recorded between 2006 and 2008 (White et al., 2007; White, 2012; Bridgland et al., 2014). Amongst the large mammal remains was a 396 397 virtually complete cranium and horn cores of an adult aurochs (Bos primigenius), 398 together with well-preserved molluscan assemblages (some specimens retaining their 399 periostracum) that included the bivalves Corbicula fluminalis and Pisidium clessini 400 and the land snail *Candidula crayfordensis*, none of which are known in Britain after MIS 7 (see Bridgland et al., 2014). Insect assemblages from Norton Bottoms are also 401 402 a close match with those from British MIS 7 sites, lacking the exotic components that 403 characterize MIS 9 and MIS 5e (Coope, 2001; Murton et al., 2001; Green et al., 2006; Bridgland et al., 2014); these have provided mutual climatic range (MCR) estimates 404 for July temperatures (T<sup>max</sup>) of +16 to +18°C and January temperatures (T<sup>min</sup>) of -11 405 406 to +5°C (Bridgland et al., 2014). Complementary data derived from ostracod assemblages using the mutual ostracod temperature range (MOTR) method (Horne, 407 408 2007; Horne et al., 2012) have provided similar temperature estimates of  $T^{max} + 15$  to 409 +21°C and T<sup>min</sup> -4 and +3°C (Bridgland et al., 2014).

## 410 Insert Figure 4 hereabouts

411

412 Other sequences containing fine-grained sediments have been recorded in the Witham 413 Valley, notably from boreholes at Coronation Farm and Stainfield (Bridgland et al., 414 2014). These yielded molluscs, pollen, plant macrofossil and insect remains; the 415 molluscan assemblages contained no biostratigraphically-significant species, but the 416 presence of *Bithynia* opercula allowed AAR dating to be applied, which suggested an 417 age within MIS 7 (Penkman, 2007; Bridgland et al., 2014). In the valley of the River 418 Bain, a left-bank tributary of the Trent-Witham system, guarrying at Tattershall 419 Thorpe revealed interglacial sediments that contained fossils not found in deposits 420 attributed to MIS 5e (Holyoak and Preece, 1985 and references therein), providing 421 some of the earliest indications that both MIS 7 and MIS 5e were preserved in the 422 terrace deposits in that area. The Last Interglacial (Ipswichian, MIS 5e) is represented 423 in three parts of the Trent system, all of them areas where the main river is joined by a 424 significant tributary. In the Middle Trent, sites at the confluence of the River Derwent with the Trent, at Boulton Moor and Allenton, have yielded hippopotamus 425 426 fossils, together with beetles and plant remains (Arnold-Bemrose and Deeley, 1896; 427 Jones and Stanley, 1974, 1975). In the Witham Valley, hippopotamus was also found 428 at several sites in the vicinity of Fulbeck (Brandon and Sumbler, 1988, 1991; Howard 429 et al., 1999; Bridgland et al., 2014), and further downstream MIS 5e sediments 430 (without hippopotamus) were found at Tattershall Castle (Holyoak and Preece, 1985; 431 Bridgland et al., 2014).

As well as providing age constraints for the terrace deposits, the new recognition ofwidespread MIS 7 interglacial deposits in the Lower Trent has been critical in the

- 434 identification of a late Middle Pleistocene (post-Anglian-pre-Devensian) glaciation in
- 435 Britain during MIS 8, a considerable rarity globally. Indeed, widespread
- biostratigraphical evidence for MIS 7 deposits in Britain as far north as Bielsbeck in 436
- 437 East Yorkshire appear to exclude the possibility of an eastern British glaciation south
- of the River Humber during MIS 6 (White et al., 2010, 2017). No interglacial 438
- 439 sequences attributable to MIS 11 or 9 have been identified in the Trent system,
- 440 providing evidence (albeit negative) for widespread destruction of this part of the late
- Middle Pleistocene record (White et al., 2010, 2017). Dating evidence from the Trent 441
- 442 fluvial archive, in the form of biostratigraphy and aminostratigraphy, has been critical
- 443 in constraining the age of this glaciation, demonstrating the utility of fluvial records where they interdigitate with glacial deposits (see Cordier et al., this issue).
- 444
- 445

#### 446 2.1.3 The Fenland rivers

447 The rivers flowing into the basin of the Wash in eastern England were recently 448 studied as part of another ALSF project (the Fenland Rivers of Cambridgeshire 449 Project), which concluded in 2008 (Boreham et al., 2010). These included the 450 Witham (before the latest Pleistocene this was the lower Trent; see Bridgland et al., 451 2014, 2015), Welland, Nene, Great Ouse and Cam, all of which have vielded well-452 preserved fossil assemblages. Morphostratigraphical approaches to fluvial deposits 453 within the Fen Basin are extremely difficult to apply, since this is an area that has 454 undergone very little uplift, resulting in poor differentiation of the Pleistocene 455 terraces. Biostratigraphical approaches have also been attempted (Bridgland and 456 Schreve, 2001), although the precise correlations of several key interglacial deposits 457 in the Fen Basin with the marine oxygen isotope stage record remain controversial 458 due to mixed or inadequate biostratigraphical signals (Boreham et al., 2010). This is 459 largely due to the preservation of sediments representing more than one interglacial in 460 close proximity beneath a single terrace surface (Boreham et al., 2010). Research in the Peterborough area has revealed considerable complexity in the fluvial deposits 461 462 preserved there (Langford and Briant, 2004; White et al., 2010, 2016; Bridgland et al., 463 2014; Langford et al., 2014a, b). Once again, occurences of Corbicula fluminalis and 464 Hippopotamus amphibius have been key to distinguishing the Last Interglacial (Ipswichian, MIS 5e) from earlier interglacials in the Wash fluvial systems. The 465 resulting synthesis of data enabled common patterns in these archives to be discerned. 466 467 but also highlighted many significant differences between these rivers, all of which 468 have shared a downstream valley during periods of low sea-level.

469

### 470 2.2 France

Multidisciplinary research projects investigating fluvial archives in NW France have 471 472 largely focussed on the Somme and Seine river valleys. Over the last decade, several 473 important palaeontological datasets from these areas have provided significant 474 chronological and palaeoenvironmental information relevant to understanding 475 Pleistocene climates and human occupation (see Chauhan et al., this issue). The 476 French fluvial record is well-dated and includes several interglacial sequences 477 particulary rich in diverse fossil groups that allow palaeoclimatic comparisons with 478 nearby British records and other datasets at a European scale (e.g. Limondin-Lozouet 479 and Preece, 2014). Alongside mammalian evidence (see Auguste, 2009), molluscan 480 evidence remains a key element of French biostratigraphical schemes. The 481 Lateglacial-Holocene molluscan successions recorded at numerous fluvial sites have 482 been central to understanding the palaeoenvironmental significance of Pleistocene

- vegetational changes and climatic conditions (Limondin-Lozouet, 2011). During
   transitional climatic phases, the stages of recolonization by molluscan faunas show
- 484 transitional climatic phases, the stages of recolonization by molluscan faunas show 485 common features, such as the occurrence of specific taxa and well-defined biotope
- 486 successions. These malacological characteristics have been correlated with climatic
- 487 phases and botanical evolution. For example, in the early Holocene of NW Europe the
- 488 replacement of the landsnail *Discus ruderatus* by its congener *Discus rotundatus* has
- 489 been linked to the spread of *Corylus* (Preece and Day, 1994; Limondin-Lozouet et al.,
- 490 2005). The same faunal succession has also been identified within Pleistocene
- interglacial sequences at La Celle in the Seine Valley, dated to MIS 11, and at Caours
  in the Somme basin, dated to MIS 5e. Although pollen was not preserved at either of
- these sites, the first appearance of *D. rotundatus* can be used to identify the
  development of deciduous forest, replacing the pioneer open forest biotope
- 495 (Limondin-Lozouet, 2011; Fig. 5).
- 496
- 497 Insert Figure 5 hereabouts
- 498

# 499 2.2.1 The Somme and Nord Pas-de-Calais

500 Research undertaken over the last 20 years on the fluvial terraces and loess sequences 501 of the Somme Basin, and on the interactions between human populations and 502 changing environmental conditions, has adopted interdisciplinary approaches for the 503 analysis of Quaternary sequences and associated Palaeolithic sites (Antoine et al., 504 2003, 2007, 2010; Bahain et al., 2007, 2010; Bridgland et al., 2006). These studies, 505 mainly targeting fluvial sequences, have highlighted the impact of cyclic climatic 506 changes on sedimentation and river morphology, and especially the role of the 100 ka 507 climatic cycles for the last million years, giving rise to stepped terrace formation 508 (Antoine et al., 2007). Interglacial climatic optima are recorded by calcareous tufa 509 sequences, which are especially well preserved in the cases of MIS 11 and 5e.

510 At the Carrière Carpentier site at Abbeville, sediments (the 'White Marl') dated to 511 MIS 15 using ESR ( $584 \pm 48$  ka) and their stratigraphical position within the Somme

512 terrace system have vielded molluscs, large vertebrates and small mammals (Antoine

et al., 2015, 2016; Voinchet et al., 2015). A noteworthy occurrence within the small

514 vertebrate faunas are rodent molars tentatively attributed to *Arvicola* 

515 *cantiana/mosbachensis*. The presence of this taxon is indicative of a younger age in

the early Middle Pleistocene, i.e. younger than MIS19-17 (Antoine et al., 2016). The

517 large mammal fauna includes the Hundsheim rhinoceros (*Stephanorhinus* 

518 hundsheimensis), wild boar (Sus scrofa priscus) and red deer (Cervus elaphus),

519 representing the Cromerian faunal association known from West Runton and

520 Pakefield. When compared with British Cromer Forest-bed sites, the mammalian

521 fauna from Carrière Carpentier appears to be contemporaneous with Pakefield, more

522 recent than West Runton, but older than the Boxgrove fauna (Antoine et al., 2016).

- 523 The Carrière Carpentier sequence is within Alluvial Formation VII of the Somme
- system (Antoine, 1994, 2000) and is associated with two other sites located in the
  same terrace, Carrière Léon and Moulin Quignon (Bahain et al., 2016). Characteristic

526 freshwater mollusc species from this alluvial formation include *Tanousia* cf.

527 stenostoma), Borysthenia naticina and Bithynia troschelii, which form an assemblage

528 typical of Cromerian interglacial fluvial deposits in NW Europe (Meijer and Preece,

529 1996; Preece, 2001). Specimens of of *T*. cf. *stenostoma* from Moulin Quignon are

530 similar to those of Little Oakley (Bahain et al., 2016). This occurrence represents the

531 first record of this species in France; in Britain it is unknown after MIS 13.

532 Recent archaeological rescue excavations in the Somme valley have led to the 533 discovery of several alluvial sequences containing molluscan faunas (Locht et al., 534 2013). At Rue Boileau in Amiens, a sequence exposed in a pit adjacent to the famous 535 site of Saint Acheul includes a white silt horizon that has been correlated on 536 morphostratigraphical grounds with the Saint Acheul tufa; the latter has been dated to MIS 11 on the basis of a characteristic 'Lyrodiscus assemblage' of land snails, 537 538 supported by U/Th dating (Limondin-Lozouet and Antoine, 2006). The molluscan 539 fauna from Rue Boileau contains several critical species typical of this assemblage, 540 together with several xerophilous taxa (Cernuella virgata, Candidula unifasciata) 541 previously unknown from St Acheul (Locht et al., 2013). This assemblage has 542 therefore been tentatively attributed to a later phase of the interglacial, post-dating the 543 climatic optimum, when heavily-forested environments were being replaced by more 544 open habitats characterised by xerophilous land snails (Limondin-Lozouet et al., 545 2015).

546

547 The important fluvial sequence preserved at Caours represents the only Eemian (MIS 5e) interglacial sequence in the Somme basin (Antoine et al., 2006). Overlying a 548 549 periglacial gravel bed allocated to MIS 6 (Alluvial Formation I of the Somme 550 system), the fluvial fine-grained sequence consists of calcareous silts and tufa 551 deposits including thin organic layers, from which a wealth of palaeontological 552 material has been recovered, most notably non-marine molluscs. These have 553 provided a record of the glacial-interglacial transition between MIS 6 and MIS 5. 554 Cold-tolerant faunas similar to those of the Younger Dryas, including some boreo-555 alpine taxa (*Columella columella*), were replaced by numerous thermophilous taxa, 556 highlighting a hiatus corresponding to the very earliest part of the interglacial 557 (Limondin-Lozouet, 2011). Following this, the malacological succession indicates the 558 development of forest environments during an early part of MIS 5, followed by the 559 climatic optimum of MIS 5e and a subsequent cooling (Limondin-Lozouet and 560 Preece, 2014). Several species of Central European and Mediterranean origin (e.g. Daudebardia rufa, Ruthenica filograna, Platyla polita) occurred during the Eemian 561 562 climatic optimum and constitute the first record of a regional biostratigraphical marker for the last interglacial (Limondin-Lozouet and Preece, 2014; Fig. 5). In 563 564 addition, the Caours site has vielded archaeological material, providing an important 565 record of human occupation in northern France during the Last Interglacial. It has 566 now been identified as a butchery site, with lithic industries directly associated with mammalian remains (Auguste, 2009). A further important aspect of the Caours 567 568 sequence is evidence for palaeomagnetic reversal thought to represent the Blake 569 Event; identification of this geomagnetic excursion, which has been correlated with 570 the start of the continental Eemian Stage, indicates that the palaeoclimatic and 571 archaeological records from Caours post-date the MIS 5e interglacial peak (Sier et al., 572 2015).

573 To the north of the Somme basin, in the neighbouring region of Nord Pas-de-Calais, a 574 fluvial sequence was discovered in 2012 at Waziers (Hérisson et al., 2015). This 575 sequence comprises several peat layers overlying fluvial silts and sands, deposited in 576 a large meandering system. Preliminary geomorphological and palaeontological 577 observations, including the presence of loess covering the fluvial deposits and the peat 578 and the occurrence of assemblages of interglacial mammals (aurochs and red deer) 579 and aquatic molluscs (Belgrandia marginata, Anisus septemgyratus) suggested an 580 Eemian age for this sequence. This was corroborated by a minimum age of 103 581 +3.5/-3.4 ka obtained by U/Th dating of calcareous charophyte oogonia, extracted

from a fine-grained tufa layer directly underlying the peat (Hérisson et al., 2015). This
site is only the second to provide evidence of Neanderthal occupation during the
Eemian in Northern France.

585

586 Finally, the late MIS 5 fluvial sequence at Ailly-sur-Nove, located  $\sim$ 30 km south of Amiens, includes three Palaeolithic levels (Locht et al., 2013). The oldest lithic 587 588 industries and associated mammals (Bos primigenius, Equus sp.) lie at the top of a basal fluvial gravel attributed to the late Saalian/Eemian on the basis of its 589 590 geomorphological position within the terrace system. The upper part of the sequence, 591 consists of coarse gravels (also containing Palaeolithic artefacts), covered by a thin 592 calcareous tufa crust with facies typical of interglacial tufa deposits like those 593 described at Caours and likely to represent MIS 5e. Above this, the sediments consist 594 of fine grey silts capped by a calcareous mud containing two further archaeological 595 levels characterized by Levallois material. Molluscan faunas from these deposits are 596 'Arianta' assemblages typical of transitional climatic phases (Puisségur, 1976); 597 further evidence for deteriorating climatic conditions is provided by a decline in the 598 numbers of the aquatic snail Belgrandia marginata, a well-known thermophilous 599 species typical of Pleistocene interglacial sequences that occurs in abundance at 600 Caours (Antoine et al., 2006). The molluscan evidence indicates a transitional context from an interglacial phase to an early glacial episode, thought to represent the period 601 602 spanning MIS 5d to MIS 5a (Locht et al., 2013); this is the first regional 603 malacological record for this time period in France.

604

## 605 2.2.2 The Seine

606 New research undertaken at the site of Saint-Pierre-lès-Elbeuf has provided additional 607 evidence for the age and palaeontological content of the important sequence there 608 (Cliquet et al., 2009). Above the lowermost palaeosol (Elbeuf IV) lies a white sand 609 and a calcareous tufa from which a characteristic 'Lyrodiscus assemblage' was obtained: this fauna is a regional biostratigraphical marker correlated with the MIS 11 610 611 interglacial. Systematic malacological sampling within this unit has demonstrated a 612 high level of homogeneity within the molluscan population, which is largely 613 dominated by forest land snails (up to 80 % of the total shells). This implies that the tufa accumulated relatively rapidly during the optimum phase of MIS 11. New IRSL 614 615 dates obtained from the Elbeuf IV soil ( $475 \pm 38$  ka) and on the white sand ( $396 \pm 32$ 616 ka) confirm the MIS 11 correlation (Cliquet et al., 2009). 617 The Upper Seine valley preserves another significant tufa sequence at the site of La 618 Celle, which has been known for more than a century as an important source of non-

619 marine molluscs and plant macrofossil impressions. New research at this site has been undertaken in order to improve understanding of the palaeontological 620 assemblages and to provide additional chronological control, which has been 621 622 correlated with MIS 11 (Limondin-Lozouet et al., 2006, 2010). Dating evidence has 623 been provided by the geomorphological position of the site, within an old terrace of 624 the Seine deposited prior to the 'Nappe de Soucy' of the Yonne valley system dated at 350-300 ka (Limondin-Lozouet et al., 2006), together with the occurrence of land 625 snails characteristic of the 'Lyrodiscus fauna', known from several French and British 626 tufa sequences of MIS 11 age (Rousseau et al., 1992; Limondin-Lozouet and Antoine, 627

628 2006; Preece et al., 2007). Additional age control has now been provided by

629 radiometric dating (ESR/U-series on tooth enamel of horse and ESR quartz), which

630 have produced a mean age of ~400 ka (Bahain et al., 2010; Voinchet et al., 2015). 631 However, although this geochronological evidence indicates an age equivalent to MIS 11, the existing methods do not allow precise timing within the interglacial. The La 632 633 Celle tufa provides the longest known MIS 11 malacological succession in NW Europe, which has been used to generate detailed reconstructions of the development 634 of forest cover (Limondin-Lozouet et al., 2010, 2015) and as a biostratigraphical 635 636 standard against which other MIS 11 malacological successions can be compared (Limondin-Lozouet et al., 2015). Successive appearances of forest species, first from 637 638 an Atlantic corridor and later from central and southern Europe, allow 639 characterization of vegetational development. The initial immigrants indicate the 640 spread of closed habitats, while a peak in the diversity of thermophilous snails corresponds to the optimum phase of deciduous forest expansion. A subsequent 641 642 decrease in forest species, together with an expansion of hygrophilous taxa, indicates 643 the decline of closed canopy conditions. The occurrence of critical forest species, 644 especially those now extinct or occurring far beyond their modern ranges, provides a 645 framework within which molluscan successions from other tufa sites in northern 646 France and southern Britain can be understood (Fig. 6).

647

#### 648 **Insert Fig. 6 hereabouts**

649

650 Leaf impressions from La Celle have allowed the identification of 24 taxa, including 651 some Mediterranean plants such as Buxus, Ficus and Celtis (Limondin-Lozouet et al., 652 2010). Mammalian remains from La Celle (Auguste, 2009; Limondin-Lozouet et al., 653 2010) include Macaca sylvanus (Barbary macaque) and Hippopotamus amphibius; 654 the former last occurred in Britain during MIS 9, whereas the latter was absent from 655 Britain during MIS 11 (Schreve, 2001a); the occurrence of hippopotamus in northern France during MIS 11 is therefore of particular interest, perhaps indicating a 656 biogeographical barrier that prevented this species colonizing Britain at that time. 657

658

### 659 2.3 Germany and Belgium

660 One of the key sites in this region is the multiperiod locality at Schöningen, Germany, the evidence from which was summarized by Schreve et al. (2007). Debate on the age 661 662 of this sequence has continued over the last decade and was recently summarized by 663 Urban and Bigga (2015). As is the case in Britain, the occurrence of *Theodoxus danubialis* has been attributed biostratigraphical significance in Germany, where T. 664 665 serratiliniformis (= danubialis) has been suggested to indicate an MIS 11 age (e.g. 666 Meng and Wansa, 2005, 2008). This is by no means certain; in the middle Neckar 667 valley (SW Germany), an exposure of fluvial gravels at the site of Bietigheim-Bissingen, ~20 m above the modern River Enz, yielded a molluscan fauna including 668 669 Theodoxus serratiliniformis and Cochlostoma scalarinum saueri (Bibus and Rähle, 2003). On the basis of terrace stratigraphy and molluscan biostratigraphy, these 670 671 deposits were dated to MIS 13, although this age is not accepted by Meng (2007). 672 673 In the southern North Sea Basin the western coastal plain of Belgium preserves a

- 674 complex sequence of Pleistocene marine and fluvio-estuarine deposits. Recent
- 675 analysis of brackish and freshwater fossil assemblages (including ostracods, pollen
- and Foraminifera) recovered from boreholes in this region have indicated deposition 676
- 677 near the upper tidal limit of an estuary (Bogemans et al., 2016). Of biostratigraphical
- 678 and biogeographical significance is the occurrence, in the Zoutenaaie core, of a single
- 679 valve of the freshwater ostracod *Scottia browniana*, which is unknown in the nearby

680 British record after MIS 11 (Whittaker and Horne, 2009). Given that the oldest of the 681 Belgian channel fills are suggested to date to MIS 9 (Bogemans et al., 2016; cf.

Vanhoorne, 2003), it is possible that this fossil has been reworked; however, its 682

683 presence might also indicate that S. browniana became extinct in continental Europe

- later than in Britain. This might well be expected, since the fluctuating island status of 684
- Britain from MIS 12 onwards enhances turnover in faunal composition and 685 biogeographical differences.
- 686 687

#### 688 3. Eastern Europe and Russia

689 Faunal evidence in this region has been recovered from the significant fluvial archives 690 preserved in the valleys of the south-flowing rivers of the East European Platform, the 691 Dniester, Dnieper, Don and Volga, and has been key to unravelling their age and 692 evolution (Matoshko et al., 2002, 2004; Schreve et al., 2007; Bridgland and 693 Westaway, 2008, 2014). To the south-west of the Russian Plain, several fossiliferous 694 localities are known within the Prut and Danube basins, although these are 695 predominantly represented by lacustrine sequences (see Markova and van 696 Kolfschoten, 2012). Several important Middle Pleistocene faunal localities are 697 known from the Dnieper, mostly recovered from fluvial deposits assigned to Terrace 698 IV, correlated with the Likhvin Interglacial (=MIS 11) on the basis of pollen and 699 mammalian biostratigraphy (Markova, 2006; Markova and van Kolfschoten, 2012; 700 Fig. 7). Amongst the best studied of these are the mammal assemblages from Gunki, Pivikha and Chigirin (Fig. 7). The Gunki site has also yielded a molluscan fauna, 701 702 considered to represent the Early Euksinian (Markova and van Kolfschoten, 2012). In 703 the Don and Desna basins, the earliest small mammal faunas have been correlated 704 with the early Middle Pleistocene (Markova, 2007). Here, faunal assemblages have 705 been correlated with both interglacial and glacial stages (Fig. 7). The 706 biostratigraphical significance of several mollusc species from the Don Basin was 707 highlighted by Kondrashov (2007), who noted that the occurrence of species such as 708 Borysthenia intermedia, Lithoglyphus jahni and Viviparus fasciatus allows correlation 709 of the Don basin faunas with the early Middle Pleistocene Cromerian faunas of 710 Western Europe. In the Volga basin, a small mammal fauna that includes Arvicola 711 cantiana, Lagurus transiens-lagurus and Clethrionomys rufocanus has been described 712 from a fluvial sequence at Rybnaya Sloboda, at the mouth of the right-bank tributary Kama River, and has been attributed to the Likhvin Interglacial on biostratigraphical 713 714 grounds (Markova and van Kolfschoten, 2012; Fig. 7). Similar faunas have been 715 reported from sites in the middle and lower Volga, at Chernyi Yar and Spasskoe 716 (Markova and van Kolfschoten, 2012). To these can be added a considerable amount 717 of vertebrate, molluscan and vegetational data that has been collected since the 1960s 718 in the southern Urals region, summarized in a flurry of papers published since 2007 719 detailing the records of the Belava and Lemeza rivers (e.g. Chlachula, 2010; 720 Danukalova et al., 2007, 2008, 2009, 2011, 2016; Puchkov and Danukalova, 2009; 721 Yanina, 2013; Yakovlev et al., 2013). Changes to the fauna and flora in this area 722 appear to have been more muted than the adjacent northwestern territories due to the 723 absence of local glaciers during some cold stages (Danukalova et al., 2009).

#### 724 **Insert Figure 7 hereabouts**

725 Further afield, in Ukraine, mollusc and ostracod assemblages have been described 726 from a sequence of Early–early Middle Pleistocene fluvial gravels and loams exposed 728 2009). The freshwater mollusc assemblages from the basal fluvial gravels (Unit I)

729 include Theodoxus serratiliniformis (=danubialis) and Viviparus lungershauseni (=

730 fasciatus; Kondrashov, 2007). Also present was the ostracod Limnocythere

*tuberculata*, which is also indicative of the Early Pleistocene (Dykan, 2003). The

732 overlying loams of Unit III also contained *T. danubialis*, together with *Lithoglyphus* 

*neumayri*. The ostracod fauna from this upper unit contained species characteristic of

colder water bodies (Boguckyj et al., 2009). Palaeoclimatic research in Russia has
also been significantly advanced by the development of the QUINSIB database,

which contains details of over 600 fossil insect localities (Kuzmina, 2014). The

development of this resource has allowed large amounts of unpublished data and

738 published sources in Russian to be made widely accessible.

739

# 740 **4. Southern Europe and Iberia**

741

742 *4.1 Spain and Portugal*743

744 A surge of interest in Iberian fluvial systems followed the successful FLAG Biennial 745 Meeting at Castelo Branco in 2010, particularly with the development of improved 746 chronological frameworks for several fluvial systems in Spain and Portugal (such as 747 the Tagus/Tejo, Minho, Douro/Duero, Mondego and Guadiana rivers), which have 748 been considerably refined through the application of new luminescence techniques 749 (Cunha et al., 2008; Vis et al., 2008; Martins et al., 2010; Antón et al., 2012; Ramos et 750 al., 2012; Viveen et al., 2012, 2013; Carvalhido et al., 2014; Sancho et al., 2016). 751 However, with the exception of a few notable regions previously summarized by 752 Schreve et al. (2007), preservation of palaeontological material is generally poor 753 across the Iberian Peninsula and the terraces of these rivers have yielded relatively 754 little new fossil material (e.g. Cunha et al., 2012). Fluvial archives such as that 755 preserved in the Ter River basin, northeastern Spain, have proved difficult to date due 756 to the absence of fossiliferous deposits (Garcia, 2015). A notable recent discovery is the site of Barranc de la Boella (Catalonia, Spain), where vertebrate remains 757 758 (including a butchered elephant) were recovered in association with stone tools from 759 deposits of the Francolí river (Vallverdú et al., 2014). Dating of this site was based 760 on palaeomagnetic and cosmogenic determinations, supported by biostratigraphical 761 evidence derived from the vertebrate faunas, which included Mimomys savini and 762 Mammuthus meridionalis, suggesting a late Early Pleistocene age (Mosquera et al., 763 2015).

763 764

765 In central Spain, faunal assemblages have been recovered from the sites of Pinedo and 766 Cien Fanegas near Toledo, in the +25–30 m terrace of the River Tajo (Tagus), which 767 have also yielded Acheulian archaeology. These gravels been dated to between 290 and 220 ka using AAR and luminescence (pIR-IRSL) techniques, spanning a period 768 769 between MIS 9 and 7 (López-Recio et al., 2015). The gravel pit at Pinedo has yielded 770 fossils of straight-tusked elephant, together with hippopotamus, rhinoceros, deer, 771 horse and bovids (López-Recio et al., 2015). Elsewhere in the Middle Tajo, large 772 mammal assemblages have been recovered in association with stone tools from the 40 773 m terrace at Toledo: these include *Mammuthus trogontherii*. Equus caballus. 774 Hippopotamus amphibius, Megaloceros savini, Eliomys quercinus, Allocricetus 775 bursae, Microtus brecciensis and Apodemus sylvaticus, all characteristic of the 776 Middle Pleistocene (Sesé et al., 2000). Palaeoclimatic studies from this region

include reconstructions based on herpetofaunal assemblages from three Spanish

localities dated to MIS 11, including the fluvial deposits at Áridos-1, in the valley of
the River Jarama SE of Madrid, and the fluvio-lacustrine sequence at Ambrona (Blain
et al., 2015). These herpetofaunal assemblages include numerous taxa of high
environmental and climatic sensitivity, which can be used as valuable indicators of
palaeoclimate, through the application of MCR analyses (e.g. Martínez-Solano and
Sanchiz, 2005; Blain et al., 2008).

- 784
- 785 *4.2 Italy* 786
- 787 The Pleistocene vertebrate succession in the Italian peninsula is relatively well known 788 (e.g. Gliozzi et al., 1997), although much of this evidence has been recovered from 789 cave and fissure-fill sequences in karstic regions (e.g. Sardella et al., 2003; Pandolfi 790 and Petronio, 2011a, b; Pandolfi et al., 2013). The potential relationships between 791 these isolated cave sequences and local fluvial systems have not yet been explored in 792 detail, but research in other regions suggests that biostratigraphical schemes can be 793 usefully developed in this way (e.g. Yang et al., 2011; Bridgland et al., 2014, 794 Westaway, 2016). In the mountainous regions of northern and central Italy, fossil 795 assemblages have also been obtained from lacustrine sequences, with fluvio-deltaic 796 deposits sometimes represented (e.g. Girotti et al., 2003; Limondin-Lozouet et al., 2017). Arguably the best known Italian fluvial archives are those from the Tiber 797 798 River basin, the record for which spans much of the Pleistocene. The palaeo-Tiber 799 and its tributaries have been extensively studied in the area around Rome (Caloi et al., 800 1998; Di Stefano et al., 1998; Milli et al., 2004; Petronio et al., 2011). Recent work 801 on this system has been undertaken by Marra et al. (2014), who identified six 802 biochronological units (Slivia, Ponte Galeria, Isernia, Fontana Ranuccio, Torre in Pietra and Vitinia), spanning a period from ~600 ka to at least MIS 7, although this 803 804 scheme has been criticised (Sardella et al., 2015; cf. Marra et al., 2015).
- 805

806 In central Italy, fossiliferous fluvial sequences representing a period spanning the Early Pleistocene to the early Middle Pleistocene have been recorded in trenches and 807 808 boreholes at several localities in the Anagni Basin (Bellucci et al., 2012, 2014). The 809 important Coste San Giacomo locality, known since the late 1970s, has yielded a Villafranchian large mammal fauna, including taxa such as *Mammuthus meridionalis*, 810 811 Hippopotamus sp., Equus stononis, Gazella borbonica, Sus strozzi and Homotherium 812 sp. More recently, microfaunal assemblages obtained from the Coste San Giacomo 1 core, drilled in 2009, have allowed more detailed palaeoecological reconstructions to 813 814 be attempted (Bellucci et al., 2012, 2014). Analyses of small vertebrate, ostracod, 815 pollen and charophyte assemblages, together with sedimentological data, indicate an initially forested landscape giving way to an open alluvial plain fed by sand-bed rivers 816 (Bellucci et al., 2014). In addition, the ostracod assemblage contained sufficient 817 818 extant taxa to attempt summer and winter air temperature reconstructions using the 819 MOTR method, providing estimates of mean July temperatures of between 15°C and 820 22°C and mean January temperatures of between -5°C and 5.2°C. However, it should be noted that the assemblage used for these estimates was derived from 8 separate 821 822 assemblages recovered from a large section of the borehole (Bellucci et al., 2014) 823 and may therefore represent an averaging of a long period of time. Small vertebrate 824 assemblages included the biostratigraphically significant Early Pleistocene vole Mimomys pliocaenicus; this species, together with evidence from pollen 825 biostratigraphy and magnetostratigraphy, suggests a Gelasian age for the Coste San 826

Giacomo faunal unit, falling somewhere between 2.2 and 1.95 Ma (Bellucci et al.,2014).

829

# 830 5. The Levant and the Arabian Peninsula

831

832 Fossil preservation in the Levant is relatively poor, despite the widespread presence of 833 calcareous bedrock. It has been suggested that the formation of calcreted gravels, a 834 characteristic of such semi-arid regions (e.g. Candy et al., 2004, 2005), might be 835 responsible for the loss of faunal remains - in the Upper Orontes, for example, 836 cemented Pleistocene terrace deposits appear to have undergone repeated 837 decalcification and re-cementation, resulting in the weathering-out of calcareous clasts and, presumably, calcareous fossils (Bridgland et al., 2012). The general 838 absence of biostratigraphical data from fluvial sequences (although see Tchernov, 839 1981, 1994 for reviews of data from cave and lake sequences in Israel) is mitigated by 840 the presence of Pleistocene lava flows interbedded within the terrace sequences of 841 842 many of the Turkish and Syrian fluvial systems that can be dated using various radiometric techniques (Sharkov et al., 1998; Bridgland et al., 2007; Demir et al., 843 844 2007, 2012, Sevrek et al., 2008; Westaway et al., 2009; Maddy et al. 2012).

845

846 Beyond the Levant, in the interior of the now hyper-arid Arabian Peninsula,

847 significant fluvial archives are largely absent; however, fossils have been recovered

from lacustrine sequences in the Nefud and Rub' al Khali deserts (e.g. Thomas et al.,

849 1998; Groucutt et al., 2014; Stimpson et al., 2014, 2016). Most of these assemblages

have been dated to the Late Pleistocene (MIS 5e and younger) and Holocene,

- although in exceptional circumstances older assemblages have been preserved (see
  below, Stimpson et al., 2014, 2016). As new sites are discovered and recorded, there
  is increasing potential to develop biostratigraphical schemes linking Arabian lake
- 854 sites to Levantine fluvial systems.
- 855
- 856 5.1 The Levant

857 Productive new research in Syria, investigating the terraces of the River Orontes (Bridgland et al., 2003, 2012) and on the catchments of Mesopotamian rivers such as 858 859 the Euphrates and Tigris (Demir et al., 2007, 2008, 2012; Westaway et al., 2009) has 860 been curtailed in recent years due to ongoing conflicts in this war-torn region. The summary of research provided by Schreve et al. (2007) therefore remains largely up 861 862 to date, with the exception of a handful of recently-published papers detailing 863 research undertaken before 2009 (e.g. Bridgland et al., 2012). Fossiliferous 864 sequences are also relatively rare in accessible reaches of these rivers in Turkey (e.g. Demir et al., 2007, 2008, 2012; Westaway et al., 2009; Seyrek et al., 2014a, b). 865 866 Vertebrate faunas have provided important biostratigraphical evidence constraining the ages of the Orontes terraces (Bridgland et al., 2003, 2012; Bridgland and 867 868 Westaway, 2007; Mishra et al., 2007). Biostratigraphical evidence provided by the 869 vertebrate assemblages from Latamneh has necessitated the reattribution of the 870 Orontes QfIII terrace in the Middle Orontes to an age of 1.2–0.9 Ma, leading to a 871 revision of the age model for the Orontes terrace sequence (Bridgland et al., 2012; cf. 872 2003). Given this revision, it is no longer tenable to attribute the Middle Orontes terraces to formation in response to 100 ka Milankovitch climatic forcing (cf. 873 874 Bridgland and Westaway, 2008b). A revised model for the evolution of that reach of 875 the Orontes, resulting from this modification of the supposed age of the key 876 biostratigraphical marker, sees similarities between the incision history in the Hama – 877 Latamneh area and those determined from the Euphrates in its southern Turkish and 878 Syrian reaches, related to the crustal characteristics of the Arabian Platform 879 (Bridgland et al., 2017). Downstream from Latamneh, molluscan and ostracod faunas 880 were obtained from river-cliff locations at Karkour, alongside the Orontes channel as 881 it traverses the subsiding Ghab Basin, including the ostracod Cyprideis torosa. This 882 taxon is generally associated with brackish environments, developing noded valves in 883 salinities below ~5‰, but can also tolerate hypersaline conditions in lakes and water bodies prone to desiccation, which is presumably how it comes to be in the Ghab 884 885 sediments. The faunas here also include freshwater elements, both ostracods and 886 molluses, the latter including the large viviparid gastropod Apameaus apameae 887 (Bridgland et al., 2012), which is also recorded at the key comparator (biostratigraphically somewhat younger) Israeli locality at Gesher Benot Ya'agov, in 888 889 the Jordan Valley, where it is an index fossil for the definition of the 'upper 890 freshwater series' or 'Viviparus Beds' of the Benot Ya'agov Formation (Picard, 1963; 891 Tchernov, 1973; Goren-Inbar and Belitzky, 1989; Bar-Yosef and Belmaker, 2010). 892 The last appearance of *Apameus apamae* in the Jordan Valley was at ~ 240 ka, on the 893 basis of U-series dating (Kafri et al., 1983; Moshkovitz and Magaritz, 1987; Heller, 894 2007); its presence at two localities in the Ghab basin, Syria (Bridgland et al., 2012) 895 and at the site of Alaattin Köyü in Turkey (Seyrek et al., 2014) therefore suggests a 896 latest Early Pleistocene to late Middle Pleistocene age for these deposits.

897

# 898 5.2 The Arabian Peninsula

899 In recent years, the Arabian Peninsula (Saudi Arabia, Yemen, Oman and the United 900 Arab Emirates) has emerged as an important region for research into low-latitude 901 Quaternary environmental change (e.g. Petraglia, 2007; Parker, 2009; Armitage et al., 902 2011; Groucutt and Petraglia, 2012; Delagnes et al., 2012). Expansions of plant and 903 animal communities, including humans, into and through the Arabian interior have 904 occurred on several occasions, corresponding with humid climatic phases during the 905 Pleistocene, when savanna-type landscapes prevailed in what are now hyper-arid 906 regions (Vaks et al., 2007, 2013; Rosenberg et al., 2011, 2013; Breeze et al., 2015; 907 Jennings et al., 2015a, 2017; Parton et al., 2015a). However, the highly fragmentary 908 nature of Arabian terrestrial sequences, compounded by poor stratigraphical and 909 chronological control on associated palaeontological assemblages, has hampered the 910 construction of biostratigraphical and palaeoenvironmental frameworks.

911 A major hindrance to better regional understanding of the Arabian record is the lack 912 of perennial fluvial systems and their resulting sedimentary archives; evidence for 913 fluvial activity consists largely of poorly-dated gravels and alluvial deposits resulting 914 from ephemeral and highly seasonal catchments (for recent reviews see Breeze et al., 915 2015, 2016). The principal drainage across central Arabia consists of several 916 eastward-flowing wadis that presently carry water only seasonally. It is likely that 917 these systems were repeatedly activated during humid periods in the Pleistocene and 918 early Holocene (Powers et al., 1966; Chapman, 1971; Anton, 1984; Edgell, 2006), 919 although some are now choked in places by dunes (Holm, 1960), suggesting that they 920 have been inactive as continuous systems for a considerable period of time. At a 921 smaller scale, more localized alluvial fans have formed around the bases of steep-922 sided jebels and dykes, from which intermittent streams flowed during wet phases 923 (e.g. Parton et al., 2015b; Jennings et al., 2015b). It has been suggested that most of 924 the trans-Arabian wadi systems were incised in their current configurations by the

926 flows (Al-Sayari and Zötl, 1978; Anton, 1984). Younger terraces have also been dated 927 using early radiocarbon techniques (e.g. Jado and Zötl, 1984) but, given the 928 unreliability of these methods, these ages require verification. More recent dating 929 programmes have applied OSL and U-series techniques to fluvial deposits in south-930 central Arabia (e.g. Maizels, 1987, 1990; Blechschmidt et al., 2009; Mclaren et al., 931 2009; Parton et al., 2010, 2013; Rose et al., 2011; Sitzia et al., 2012; Atkinson et al., 932 2013), contributing to an increasingly detailed chronology for the Late Pleistocene 933 and Early Holocene. However, the general absence of substantial fluvial archives in 934 Arabia is reinforced by the fact that the most significant stratified archaeological 935 assemblages reported in recent years have been found in association with either 936 lacustrine sequences, representing a source of fresh water, or raw materials suitable 937 for stone tool production (e.g. Armitage et al., 2011; Delagnes et al., 2013; Hilbert et 938 al., 2014; Groucutt et al., 2015; Jennings et al., 2015b; Scerri et al., 2015).

939

# 940 6. Data from other regions941

Beyond the 'core regions' of NW and Central Europe, palaeontological datasets from
fluvial sequences are increasingly used as a basis for regional biostratigraphies and
palaeoenvironmental reconstructions. At the time of the last review (Schreve et al.,
2007), these consisted entirely of mammalian data; since then, significant new
information, including invertebrate and floral assemblages, have been published and
are included in the summaries below.

948

# 949 6.1 China and the Far East

950 In China, the Nihewan Formation, comprising fluvio-lacustrine sediments containing 951 abundant mammalian fossils, is widely distributed in the Nihewan Basin of northern 952 Hebei Province. This region has vielded the densest concentration of Early 953 Pleistocene Palaeolithic sites outside Africa (Dennell, 2013), together with fossil 954 assemblages (vertebrates, molluscs, pollen) that have long been considered to be of 955 Early Pleistocene age, although alternative ages ranging from late Pliocene to Middle 956 or Late Pleistocene have also been proposed (Zhao et al., 2010). ESR dating of quartz 957 obtained from the Majuangou and Banshan sites provided ages of between 1.70 and 958 1.35 Ma (Liu et al., 2014). Further south in the Three Gorges region, located in the transitional zone between the upper and middle reaches of the Yangtze (Changjiang) 959 960 River, several fossiliferous fluvial sequences have now been reported (Pei et al., 961 2013). Seven fluvial terraces  $(T_7 - T_1)$  have been identified and dated using ESR, TL, 962 OSL and radiocarbon techniques (Pei et al., 2013). The site at Jingshuiwan, assigned 963 to T<sub>2</sub> and dated to  $75.9 \pm 3.7$  to  $64.5 \pm 4.1$  ka using OSL, yielded an assemblage of 58 964 mammalian fossils including Stegodon orientalis (Pei et al., 2010). 965 966 In northwest Hunan Province, the terraces of the Suoxi River have been dated using

967 ESR and TL techniques, allowing them to be correlated with neighbouring karstic 968 cave systems (Yang et al., 2011). Although none of the dated localities in this study 969 proved to be fossiliferous, presumably due to the local sandstone bedrock, such 970 research highlights the potential for fluvial archives to provide chronological evidence 971 that can be linked to calcareous sequences with potential to provide palaeontological 972 data. Palaeoclimatic research in East Asia using methods such as beetle MCR 973 estimates has been hampered by a lack of knowledge of modern distributions of beetle 974 species and the sparse distribution of meteorological stations, especially in Siberia 975 (Shivake, 2014). Japan is exceptional in having conditions that have allowed

976 application of the Mutual Climatic Range (MCR) method, having a dense

977 meteorological observation network and higher quality distribution data for its extant

978 beetle fauna (Shivake, 2014). Experimental work at the site of Nojiriko, Nagano

979 Prefecture has provided the first MCR reconstructions from East Asia (Shiyake, 2014).

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- 981
- 982

### 983 6.2 South Asia

984 The Indian subcontinent has yielded a range of palaeoecological and 985 biostratigraphical data, in the form of vertebrate and invertebrate fossil assemblages, 986 pollen records and archaeological material, much of which has been recovered from 987 fluvial sediments of the Narmada, Godavari, Manjra, Son, Ghod, Krishna and 988 Mahanadi rivers (Chauhan, 2008). Most of these deposits have been assigned ages no 989 older than the Middle Pleistocene, although some formations in the Narmada Valley 990 might date to the Early Pleistocene (Tiwari and Bhai, 1997; Tiwari, 2001). The 991 application of modern dating techniques has shown that many of these sedimentary 992 units have the potential to be significantly older or younger than previously thought 993 (Chauhan 2008 and references therein), a situation that is also prevalent in the Levant 994 (Bridgland et al., 2012) and Arabia (e.g. Thomas et al., 1998, contra Stimpson et al., 995 2014, 2015). Until reliable chronological frameworks are available for these fluvial 996 archives, their usefulness in terms of regional biostratigraphy remains somewhat 997 limited.

998

999 The Siwalik deposits of northern India, Pakistan, Nepal and Myanmar represent one 1000 of the best-studied fluvial sequences in Asia. The Upper Siwaliks have been divided 1001 into three subdivisions, the youngest two of which (the Pinjor and Boulder 1002 Conglomerate Formations) represent most of the Ouaternary, spanning the period 1003 from 2.58 to 0.2 Ma (Prasad, 2001; Mishra et al., 2010). Mammalian fossils are 1004 adundant within the Pinjor Formation (2.58 to 0.6 Ma), which has yielded at least 98 1005 vertebrate species including *Elephas hysudricus*, *Stegodon insignis*, *Rhinoceros*, 1006 Sivatherium, Equus sivalensis, Bos acutifrons and Cervus palaeindicus, together with 1007 carnivores such as Canis pinjorensis, Crocuta felina and Pachycrocuta brevirostris 1008 (Nanda, 2002, 2008). However, the stratigraphic range of most of these species 1009 within the Pinjor Formation is unknown, limiting their biostratigraphical application 1010 (Dennell et al., 2008). Nevertheless, reconstructions of the palaeoecology of the 1011 fluvial landscapes within which the Pinjor Formation was deposited have been 1012 attempted (Dennell et al., 2008), including predator-prey interactions and the 1013 circumstances under which fossil assemblages were accumulated and buried.

1014

1015 Two younger faunal horizons, roughly equivalent to the late Middle and Upper 1016 Pleistocene respectively, are preserved on the Indo-Gangetic plain in India and in Sri 1017 Lanka (Nanda, 2008; Mishra et al., 2010), although these have yielded only 16 and 26 1018 mammalian taxa, respectively. These 'post-Siwalik' faunas suggest that a large 1019 proportion of the Upper Siwalik Pinjor fauna became extinct during the Middle 1020 Pleistocene. Because the stratigraphical relationships between these numerous fluvial 1021 localities remain unclear, it has not yet been possible to propose a testable 1022 biostratigraphical framework for the region. However, a relative chronostratigraphic, 1023 biostratigraphic and archaeological correlation for the Narmada Basin formations has been proposed (Badam, 2007; Patnaik et al., 2009). At Hathnora, assemblages of 1024 vertebrates (including herpetiles and fish), molluscs, ostracods, charophytes and 1025

- 1026 pollen, in association with archaeological material, have been reported (Patnaik, 2000; 1027 Patnaik et al., 2009). Freshwater molluscs have also been recovered from the sites at 1028 Bhedaghat and Devakachar (Hirdepur Formation), including 18 species of gastropods 1029 and bivalves (Kotlia and Joshi, 2011). Several lines of evidence suggest that 1030 Southeast Asia was dominated by a mosaic of savannah, open woodland and 1031 evergreen forest throughout much of the Pleistocene, leading to suggestions that this 1032 region served as a refugium for hominins and other mammal species during glacial 1033 periods (Louys and Turner, 2012)
- 1034

1035 Well-dated fossiliferous sequences in Southeast Asia are relatively rare and are 1036 predominantly preserved in caves, although the syntheses described here include some data from fluvial deposits. Broad characteristics of the extinctions of 1037 1038 Pleistocene large vertebrates in this region were recently reported by Louys et al. 1039 (2007), together with consideration of the palaeoenvironmental requirements of many 1040 of these species (Louys and Meijaard, 2010). Many species which became extinct 1041 appear to have been endemic to specific areas, with others suffering severe range reduction before their eventual demise. Members of the latter group include 1042 1043 proboscideans (Stegodon and Palaeloxodon), the pygmy hippopotamus 1044 (Hexaprotodon), the orangutan (Pongo), hyenas (Crocuta and Hyaena), the giant 1045 panda (Ailuropoda), tapirs (Tapirus and Megatapirus), rhinoceroses (Rhinoceros), 1046 and the giant Asian ape, Gigantopithecus. The loss of these species is likely to have 1047 been the result of a combination of climatic changes (Louys et al., 2007) and human 1048 impacts (Corlett, 2007). Unlike other regions which experienced megafauna 1049 extinctions, such as South America (see below), eustatic changes in sea level in 1050 Southeast Asia seems to have been an important factor (Louys et al., 2007).

1051

1052 On the island of Java, Indonesia, fluvial deposits of the Solo River have recently 1053 yielded both vertebrate fossils and archaeological material. An extensive survey of 1054 the region around Matar, close to the site at Ngandong (where fossils of Homo erectus 1055 were recovered in the early 1930s), revealed the presence of at least three river 1056 terraces; palaeontological and archaeological assemblages were recovered from the 1057 First and Second terraces (Fauzi et al., 2016). Tentative comparisons with other 1058 Pleistocene faunal localities on Java suggest that the Matar assemblage is younger 1059 than the Middle Pleistocene Kedungbrubus Fauna but older than the Punung Fauna, 1060 since it contains several species (such as Stegodon trigonocephalus, Bubalus paleokarabau, Bibos paleosondaicus and Hexaprotodon sivalensis) that occur in the 1061 1062 former but not in the latter (Fauzi et al., 2016). The Punung Fauna, the type locality 1063 for which is Punung Cave, Indonesia, has been dated to 128±15 and 118±3 ka using 1064 luminescence and U-series techniques (Westaway et al., 2007). Further work is 1065 required in order to understand fully the regional biostratigraphy.

1065

# 1067 Insert Fig. 8 hereabouts

1068

# 1069 6.3 North and South America

A significant proportion of Pleistocene palaeontological research on these continents
has been driven by a desire to understand the reasons for megafaunal extinctions (e.g.
Faith et al., 2009; Haynes, 2009; Scott, 2010; Doughty et al., 2013) and the timing of
the first human colonization of the Americas (e.g. Goebel et al., 2008; Rothammer
and Dillehay, 2009; Pitblado, 2011), as well as dynamics between the two (e.g. Gill et
al., 2009). Fluvial archives are also potential sources of palaeontological evidence

1076 relevant to debates surrounding the Great American Biotic Interchange (GABI), a 1077 period of faunal exchanges between North and South America. The isolation and 1078 faunal endemism of South America ended with the formation of the Panamanian land 1079 bridge and the onset of the GABI, with the first of four major faunal exchanges now 1080 thought to coincide with the base of the Pleistocene at  $\sim 2.6$  Ma (Reguero et al., 2007; 1081 Woodburne, 2010). Further major pulses, at 1.8, 0.7 and 0.125 Ma (Fig. 8) appear to 1082 have coincided with periods of lower sea-level, leading to the suggestion that 1083 expanded coastal regions, cooler climates and associated changes to local flora 1084 promoted animal dispersals at these times (Woodburne, 2010).

1085

# 1086 Insert Fig. 9 hereabouts

1087 1088 There are few Pleistocene sites in South America from which well-dated fossil 1089 material has been recovered from secure stratigraphical contexts; correlations between 1090 localities have therefore frequently been based on the biostratigraphical scheme 1091 proposed for the fossiliferous beds of the Pampean region of Argentina, which has 1092 been periodically updated over the last two decades (e.g. Cione and Tonni, 1999, 1093 2001, 2005). The Ouaternary mammal faunas of South America have also been 1094 reviewed from time to time by several authors (see Prado and Alberdi, 2009 and 1095 references therein). Major phases of mammalian dispersal occurred during the 1096 Pleistocene Ensenadan and Lujanian biochrons (Fig. 9), which have also been dated 1097 on the basis of the Pampean sequence (Cione and Tonni, 1999, 2001, 2005). 1098 Biostratigraphical significance has been ascribed to four species of Neosclerocalyptus 1099 (Glyptodontidae) which occurred during the Pleistocene in the Pampean region of 1100 Argentina (Zurita et al., 2009a). Two of these, Neoclerocalyptus pseudornatus and N. ornatus are indicative of the Ensendan Stage, the former occurring between 1.07 and 1101 0.98 Ma and the latter between 0.98 and 0.40 Ma (Fig. 9). Two other species, N. 1102 1103 gouldi and N. paskoensis, are thought to reperesent the Bonaerian and Lujanian 1104 stages, respectively (Zurita et al., 2009a; Fig. 9). This biostratigraphical scheme has 1105 therefore become a standard with which other South America sequences are now 1106 routinely compared (e.g. Lopes et al., 2010; Tonni et al., 2009; Beilinson et al., 2015). 1107

1108 In Bolivia, the Tarija Valley preserves some of the richest Pleistocene mammal 1109 localities in South America, although the dating of these assemblages remains unclear (Coltorti et al., 2007; Tonni et al., 2009; Zurita et al., 2009b). A Middle Pleistocene 1110 1111 age has been suggested on the basis of mammalian biostratigraphy (e.g. Tonni et al, 1112 2009 and references therein), although they have also been attributed to a much 1113 younger period on the basis of radiocarbon dating ( $\sim 44 - 21$  ka BP). A similar lack of 1114 reliable dating for fossiliferous sites has hampered the bio- and chronostratigraphical 1115 correlation of South American Pleistocene faunas elsewhere; in Brazil, Lopes et al. 1116 (2010) published ESR dates of between 226 and 34 ka for a vertebrate assemblage 1117 from Chuí Creek; this broad age range, spanning a time period encompassing the final 1118 part of MIS 7 until MIS 4, is probably due to the reworked nature of the Chuí Creek 1119 vertebrate assemblages (Pereira et al., 2012). In Venezuela, strata such as the Taima 1120 Taima fossil bed and fluvial sites at Muaco, Cucuruchú, and Quebrada Ocando have

1121 been dated on the basis of biostratigraphy (Carlini et al., 2008).

In Uruguay, the fossiliferous beds of the Sopas Formation have yielded vertebrate
faunas, together with freshwater molluscs, plant remains and other trace fossils
(Ubilla et al., 2004, 2009, 2016; Ubilla and Martínez, 2016). The mammalian

- assemblage includes extinct taxa, such as the capybara *Neochoerus aesopi*, the
- 1126 glyptodont Neuryurus rudis, and two extinct species of deer (Antifer ultra and
- 1127 *Morenelaphus brachyceros*). This assemblage has been correlated on the basis of
- biostratigraphy with the Lujanian Stage (Late Pleistocene–Early Holocene) of the
- 1129 Pampean region of Argentina; this is supported by radiocarbon AMS dates from the
- 1130 vertebrate material range from  $39,900 \pm 1,100$  to  $33,560 \pm 700$  BP (cal 45,389 to
- 1131 42,025 ya) and TL/OSL ages derived from the sediments range from
- 1132  $71,400 \pm 11,000$  to 27,400  $\pm$  3,300, indicating a MIS 3 age for the fauna (Ubilla et al.,
- 1133 2016). Palaeoenvironmental evidence suggests that the Sopas formation represents 1134 open habitats, savannahs and woodlands (Ubilla et al., 2016).
- 1135

# 1136 Insert Fig. 10 hereabouts

1137 1138 *6.4 Australia* 

1139 In Australia, Quaternary palaeoenvironmental research based on fluvial archives has a 1140 long pedigree in two important regions of the continent: the arid interior of the Lake Eyre Basin, where fluvial, lacustrine and aeolian sequences representing the last ~300 1141 1142 ka are preserved (recently reviewed in detail by Habeck-Fardy and Nanson, 2014), 1143 and the extensive meandering river systems of southeastern Australia, particularly the 1144 well-dated terraces of the Lachlan and Macquarie rivers in the Murray-Darling Basin 1145 (Kemp and Spooner, 2007; Yonge and Hesse, 2009; Kemp and Rhodes, 2010). 1146 Faunal and floral responses to Pleistocene climate change in these regions remain less 1147 well understood, primarily due to the rarity of stratified fossil assemblages and 1148 significant issues with directly dating fossil material (Price et al., 2013; Westaway et 1149 al., 2017). As is the case in North America, much research in Australia has been 1150 focussed on the extinction of megafaunal species during the Late Pleistocene; 1151 approximately 96 % of the large mammal fauna was extinct by  $\sim$ 45 ka, a period 1152 broadly concurrent with human colonisation, although the extent to which these 1153 extinctions can be directly related to human activity remains a source of considerable 1154 debate (e.g. Koch and Barnosky, 2006; Prideaux et al., 2007, 2010; Price et al., 2015; 1155 Dortch et al., 2016; Johnson, 2016; Johnson et al., 2016; Westaway et al., 2017; Fig. 1156 11). Overhunting and the burning of the landscape by people, with a corresponding 1157 abrupt reduction in plant diversity, have been suggested as causal factors (Porch and 1158 Kershaw, 2010; Rule et al., 2012), but significant climate change in the period 1159 between 50 and 46 ka, resulting in a major shift to more arid conditions and a 1160 corresponding drop in water levels in Lake Eyre and Lake Frome, is also considered 1161 to have played an important role (Murphy et al., 2011; Cohen et al., 2012, 2015; 1162 Sakaguchi et al., 2013).

1163

1164 In the Lake Eyre Basin, fossils of 21 megafaunal mammal species have now been 1165 recorded (Webb, 2008, 2009), including a giant wombat-like marsupial (Diprotodon), 1166 giant short-faced kangaroo (Procoptodon goliah), a large flightless bird (Genyornis 1167 newtoni), giant goanna (Varanus priscus) and the 'marsupial lion' (Thylacoleo 1168 *carnifex*). In addition, smaller mammal species have been recovered from the upper 1169 Katipiri Formation (MIS 6-4), including the Southern brown bandicoot (Isoodon 1170 obesulus), the extinct Eastern hare wallaby (Lagorchestes leporides), the Pale field rat 1171 (Rattus tunneyi) and the Western grey kangaroo (Macropus fuliginosus); the fossil 1172 occurrences of the three extant species are well beyond their modern distributions 1173 (Webb, 2009). However, the lack of chronological control for much of this fossil 1174 material means that the first and last appearances of many mammal species in the

1175 wider Lake Eyre Basin remain poorly understood. Better biostratigraphical evidence 1176 has been forthcoming in eastern Australia, where vertebrate and molluscan 1177 assemblages (44 taxa in total) have been recovered from the Darling Downs, within fluvial deposits of the Kings Creek (Price and Sobbe, 2005; Price et al., 2011). 1178 1179 Palaeoenvironmental analyses have indicated that a mosaic of local habitats, including 1180 vine thickets, scrublands and open grasslands, prevailed during the late Pleistocene; 1181 increasing aridity led to a contraction of the more wooded environments in favour of 1182 grassland. Dating of the Kings Creek sequences using a variety of techniques (OSL, 1183 U-series and radiocarbon) has shown that individual megafaunal species responded 1184 independently to climatic and environmental change, revealing a more complex 1185 staggered extinction pattern in southeastern Austrial prior to the arrival of humans (Price et al., 2011; Fig. 11). 1186

1187 1188

Insert Fig. 11 Australian biostrat chart

1189 1190

# 1191 **7. Synthesis**

1192 1193 Regional biostratigraphies based on mammalian and molluscan assemblages continue 1194 to contribute significantly to the dating and correlation of Pleistocene fluvial deposits 1195 on a global scale. However, the quality of these datasets varies widely, depending on 1196 the potential for preservation of fossils in a given region and the priority given to 1197 palaeontological and biostratigraphical research. The period since the 2007 review of 1198 regional biostratigraphies has seen a steady accumulation of data in many parts of the 1199 world. Those based on mammalian and molluscan assemblages, in particular, have 1200 continued to be tested and developed, especially in NW Europe, and clearly have a 1201 significant role to play in the dating and correlation of Quaternary deposits. Biostratigraphical frameworks are still predominantly based on mammals and other 1202 1203 vertebrates, which is not surprising considering the robust nature of their fossils. 1204 Molluscs are also commonly used. Other groups, such as reptiles, amphibians, 1205 ostracods and insects are still only infrequently used as a dating tools (e.g. Gleed-1206 Owen, 1988, 1999, Coope, 2001; Griffiths, 2001; Whittaker and Horne, 2009; 1207 Borodin et al., 2013), but can provide invaluable complementary information as 1208 palaeoclimatic indicators. The relative rarity of non-mammalian vertebrate remains, 1209 which has been noted in fluvial sequences in regions such as Europe (Holman, 1998; 1210 Roe et al., 2009), Australia (Price and Sobbe, 2005) and Africa (Stoetzel et al., 2012) 1211 is usually attributed to taphonomic biases caused by the relative fragility of these 1212 fossils, although only a few studies specifically addressing this issue have been 1213 published (e.g. Pinto Llona and Andrews, 1996, 1998).

1214

1215 The predominance of work in the cooler temperate regions is in large part a result of 1216 the common preservation there of vertebrate and molluscan faunas in fluvial contexts 1217 that coincide with calcareous settings. In the warmer temperate Mediterranean region 1218 limestone abounds but dryland soil processes have led to reprecipitation of calcareous 1219 cements that seems to have coincided with the destruction of fossils. It is probably no 1220 accident that the key late Early Pleistocene fauna site of Latamneh in the Orontes in 1221 Syria is in one of the few reaches of that river with terrace gravels that are not 1222 calcareously cemented (Bridgland et al., 2012). The only other Orontes sites with 1223 significant faunas are preserved in the Ghab, where the presence of fossils might owe 1224 much to waterlogging in this fluvio-lacustrine subsiding basin (see Section 5).

1225 1226 There is an increasing need for research to address the validity of inferred patterns of 1227 faunal (including hominin) exchange between neighbouring regions. These have the 1228 potential to address important questions about the timing and nature of 1229 palaeoenvironmental change in response to climate change, by identifying key drivers 1230 of dispersal and the waxing and waning of biogeographical barriers. Of critical 1231 importance when comparing adjacent regions is the accuracy of chronological 1232 frameworks, which are required to establish whether significant palaeoenvironmental 1233 changes and faunal turnovers occured synchronously in different regions, or whether 1234 leads and lags can be identified. It is often not possible to detect diachroneity in 1235 faunal change due to the lack of precision in radiometric methods, especially in the 1236 Early and Middle Pleistocene. Improved dating is thus required to determin patterns 1237 of north-south exchange between refugia (in the Northern Hemisphere), an issue of further relevance in respect of Latamneh, which has been suggested as a possible 1238 1239 cold-stage assemblage that includes taxa represented in interglacials further north 1240 (Bridgland et al., 2012).

1241

1242 Detailed analyses are also important when it comes to reconstructing

1243 palaeoenvironmental change through longer fluvial sequences, with the potential to 1244 detect smaller-scale climatic oscillations, potentially correlateable with oxygen isotopic substages (e.g. Schreve, 2001b; White et al., 2013). These problems are well 1245 1246 known to Quaternary biostratigraphers and palaeoecologists. The value of 1247 biostratigraphical and palaeoecological data has led to such studies becoming an 1248 increasingly common element of multi-disciplinary research; such projects are often 1249 driven primarily by archaeological research questions. Evidence corroborating and 1250 strengthening established biostratigraphical frameworks (Schreve, 2001a, b) has been 1251 obtained, together with substantial new understanding of the palaeoclimatic and 1252 palaeoecological significance of numerous plant and animal species that occur 1253 commonly within fossil assemblages (e.g. Candy et al., 2012, 2015, 2016).

1254

1255 Certain taxa appear to have been extremely widespread at particular times during the 1256 Pleistocene, only for their ranges to fragment in response to subsequent environmental pressures. The distribution of the hippopotamus (Hippopotamus 1257 1258 *amphibius*) has long had significance in Britain, where it is a distinctive marker-fossil 1259 for the Last Interglacial (MIS 5e) (Sutcliffe, 1964; Currant and Jacobi, 2001; Schreve, 1260 2001a, 2009). Similarly, distinctive freshwater mollusc species such as those of the 1261 'Rhenish suite' characteristic of the MIS 11 Thames were clearly able to disperse 1262 rapidly across large parts of Europe. These two examples illustrate the potential for 1263 fluvial systems to connect otherwise separate biogeographic provinces, linking regional biostratigraphic schemes at certain times. Terrestrial species, for which large 1264 1265 rivers are potential barriers to dispersal, are less likely to be able to profit from fluvial 1266 connectivity. However, in the case of the 'Lyrodiscus' fauna that characterises 1267 molluscan faunas in Britain and northern France during MIS 11, some continuity in 1268 woodland habitat was clearly required to allow this biome to develop.

1269

Some of the most significant patterns in the palaeontological record are evident at the extremities of regions and in differences between continental and island records. For example, in NW Europe the well established British biostratigraphical record differs from that of its nearest continental neighbours; both regions benefit from excellent

1274 preservation of fossils and well-dated fluvial stratigraphies, allowing direct

- comparison of the faunas and floras of Pleistocene interglacials. The occurrence of
  hippopotamus in Britain and on the near-continent continues to be a point of
  difference, for example. This taxon occurs in abundance in Britain during the Late
  Pleistocene only during the Last Interglacial (MIS 5e, Ipswichian), whereas in
  northern France it has been found in assemblages dated to MIS 11 and MIS 7.
- 1279

1281 Palaeoclimatic reconstructions based on multiple fossil proxies have proved 1282 particularly useful for understanding interglacials, especially on the northern fringes 1283 of Europe. This sort of research has only been possible in regions where long, 1284 detailed sedimentary sequences are preserved. In Britain, much of this evidence has 1285 been derived from fossils of a diverse range of thermophilous flora and fauna that 1286 now occur in more southerly parts of continental Europe, or even further afield. At a 1287 basic level, qualitative measures of enhanced warmth (or cold) can be shown by the 1288 relative abundances of thermophilous or cryophilous species within fluvial deposits. 1289 More precise palaeotemperature estimates can be obtained from quantitative methods 1290 based on palaeoecological proxies, which now have a long pedigree within 1291 Quaternary research in northwest Europe (e.g. Atkinson et al., 1987; Zagwijn, 1996; 1292 Horne et al., 2012). Such approaches employ the known climatic tolerances, or the 1293 mapped climatic range, of extant plant or animal species in order to infer the warmest 1294 (Tmax) and coldest (Tmin) temperatures under which a given assemblage of species 1295 could survive (Candy et al., 2010). Fossils of these indicator species (or assemblages 1296 of groups of species) within Pleistocene fluvial deposits can thus be used as proxies 1297 for the palaeotemperature regime that prevailed at the time the deposit accumulated; 1298 within longer sequences, it is also possible to infer significant climatic changes from 1299 the fossil record.

1300 Consideration of the ecological preferences of mollusc and ostracod species is 1301 important and can bolster arguments from faunal comparison data that can otherwise 1302 seem circular or overly reliant of negative (absence) evidence. An interesting 1303 observation from recent years has been the occurrence of halophytic taxa in warm-1304 climate fluvio-lacustrine settings, where evaporation enhances salinity. This was 1305 noted in the reach of the Orontes in northern Syria, where the aforementioned 1306 subsiding Ghab basin is crossed (Bridgland et al., 2012). At that land-locked locality 1307 there is a low potential for confusion with an estuarine environment, but in more 1308 coastal settings the co-occurrence of certain species could raise problems for 1309 environmental interpretation.

131013118. Conclusions

1312 Since the last major review of biostratigraphical data derived from Quaternary fluvial 1313 archives in 2007, significant new research on Pleistocene fossil assemblages has been 1314 undertaken on almost every continent. Established regional biostratigraphical 1315 frameworks have been challenged, modified and, in many cases, strengthened by the 1316 new data generated by this work. Over the 20-year lifespan of FLAG, much of the 1317 evidence summarized here has been obtained from fluvial sequences in the NW 1318 European 'hotspot' that includes Britain and France, together with important archives 1319 from Germany, Iberia and Italy and Eastern Europe. Research in Europe has included 1320 the reporting of new sites, dating or re-dating of known localities to improve 1321 chronological control, and the development of high-resolution palaeoenvironmental 1322 reconstructions based on fossil assemblages and geochemical analyses. There have 1323 also been important advances in understanding of the palaeoclimatic and

palaeoecological significance of numerous plant and animal species that occur
commonly within Pleistocene fossil assemblages; in combination with improved
chronological frameworks, these provide important evidence for changing climate and
environments through time.

1328

1329 In the Levant and South Asia, where the archaeological record has driven the research 1330 agenda, the recognition of the utility of fossil assemblages as chronological and 1331 palaeoevironmental tools has led to them being increasingly studied, despite 1332 considerable taphonomic difficulties. The Arabian Peninsula, in particular, has been 1333 the subject of an increased focus of studies relating to hominin dispersals out of 1334 Africa, but because well preserved fluvial deposits are largely absent in this arid 1335 region, fossils have mainly been derived from lacustrine and cave sequences. The 1336 potential for developing biostratigraphical correlation schemes, linking the Arabian 1337 record to better-preserved fluvial archives in neighbouring regions such as the Levant, is therefore an exciting possibility for future research. 1338

1339

1340 Europe and the Middle East have in common long-timescale records of human 1341 occupation; beyond these areas, in the Americas and Australia, human impacts came 1342 much later and have been studied mainly in the context of human contributions to the 1343 extinctions of Pleistocene megafaunas. The chronology of the Great American Biotic 1344 Interchange, together with refinements in South American biostratigraphical schemes 1345 in countries such as Brazil, Argentina and Bolivia continue to be major areas of 1346 research. Similarly, in Australia, frameworks of faunal extinctions have been 1347 developed. Australia has an apparently impoverished Middle and Late Pleistocene 1348 biostratigraphical record compared with other parts of the world, and many of the 1349 species that went extinct during this period are poorly dated. Archaeological sites 1350 attesting to potential human impact are rare and detailed ecological information for 1351 most extinct megafauna is lacking. As a result, the processes leading to megafaunal 1352 extinction remain unclear, although the weight of evidence points to a direct human 1353 impact as a major cause of extinction.

1354

Although it is now possible to generate reliable radiometric dates for river terrace sequences in many parts of the world, enabling their correlation with the globally applicable marine oxygen isotope record, in some regions this remains extremely difficult. Where such robust chronological frameworks exist, providing fossils are also well preserved, it is possible to explore more detailed patterns in the occurrences of plant and animal species during the Pleistocene.

1361

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1 Numbers indicate the oxygen isotope stages to which these have been attributed



**Cold-climate gravels** 

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# Highlights

- Research over the last decade into biostratigraphical data recorded in fluvial archives is reviewed
- Consideration of biogeography and palaeoclimatic evidence is also included
- This emphasizes the disparities in research priorities and fossil preservation on a global scale