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Chironomid-based reconstruction of Lateglacial summer temperatures from the Ech palaeolake record (French western Pyrenees)

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1. Introduction

During the transition from the Pleniglacial to the early Holocene, the so-called Lateglacial period, Europe was characterised by high climatic variability. Several major, rapid and high amplitude climate changes occurred within ca. 3000 yr. The onset of the Lateglacial Interstadial was marked by a sudden warming at ca. 14,700 cal yr BP. This warm phase was abruptly interrupted at ca. 12,700 cal yr BP by a millennium-long cold reversal, known as the Younger Dryas. At ca. 11,700 cal yr BP, the onset of warm conditions corresponded to the beginning of the Holocene, the present interglacial. Super-imposed on this millennial-scale climate variability, the Lateglacial was punctuated by a series of centennial-scale events such as the Gerzensee and the so-called Lateglacial period, Europe was characterised by high climatic variability in the present context of global change. Within this framework, a spatial network of quantitative reconstructions of past climate changes is of major interest for assessing past climate gradients at the continental scale. Comparisons between reconstructions and model hindcasts can ultimately serve to improve climate model performance and forecasting.

Quantitative reconstructions of past climate changes are still unevenly distributed throughout Europe. Renssen and Isarin (2001) pointed out the lack of Lateglacial quantitative reconstructions for the southern part of the continent. Since then, new data were produced from Mediterranean sediments (e.g. Dormoy et al., 2009) and sediment records from lakes in the Southern Alps (Heiri et al., 2007a; Larocque and Finsinger, 2008; Ortu et al., 2008). Nevertheless, quantitative estimations of climate parameters remain rare for many areas of Europe and more specifically for the south-western part of the continent.

Among biological proxies from lake sediments, chironomid assemblages are viewed as one of the most promising climatic indicators (Battarbee, 2000). Most of the larvae of Chironomidae (Insecta: Diptera) live at the bottom of lakes. Chitinous remains (head capsules) produced during larval development are well preserved in sediment archives. Changes in past community composition through the understanding of the earth’s climate engine in the present context of global change. Within this framework, a spatial network of quantitative reconstructions of past climate changes is of major interest for assessing past climate gradients at the continental scale. Comparisons between reconstructions and model hindcasts can ultimately serve to improve climate model performance and forecasting.
time are reconstructed after extraction and identification of head capsules along sediment cores. In the pioneering work of Walker et al. (1991) followed by other studies in Europe (Brooks, 2006), temperature was identified as one of the key factors affecting chironomid assemblage distribution at a large geographical scale. A wide consensus exists that the large temperature changes registered during the Late-glacial had a distinct influence on chironomid assemblages of shallow lakes, although several other environmental and limnological factors may also have had a potential impact on the fauna (Walker, 2001). Thanks to the development of transfer functions based on the modern distribution of chironomids among a large set of lakes, summer temperature can be inferred from past changes in chironomid assemblages. The accuracy and reproducibility of this approach for Lateglacial temperature reconstructions is now broadly demonstrated (Brooks, 2006; Lang et al., 2010).

This study is the first attempt to reconstruct chironomid assemblages from a lake record in the Pyrenees (South-western France) and infer quantitatively summer temperatures during the Late-glacial in a region where such reconstructions are still lacking. To date, there is no transfer function available in the study area for inferring summer air temperature. This lack of local calibration data was circumvented by using a newly developed transfer function based on a large number of lakes in the northern and central Alps of Switzerland and in Norway (Heiri et al., 2011).

The aim of this work is to (i) unravel the response of lateglacial chironomid assemblages to climate changes in the Pyrenees, (ii) assess the applicability of a mixed Swiss–Norwegian transfer function to a Pyrenean fossil record and (iii) evaluate the consistency of the temperature reconstruction by means of comparison with other climate record from Europe and Greenland.

2. Study site

The mountain chain of the Pyrenees spreads from the Atlantic Ocean to the Mediterranean Sea (Fig. 1). This intermediate geographical position implies the existence of a strong W–E climatic and ecological gradient throughout the chain from the Atlantic climate prevailing in the western part to Mediterranean climate in the eastern zone. The study site (43° 04' 58.74 N; 0° 05' 38.53 W, 710 m a.s.l.) is located in the west-central Pyrenees (Fig. 1). The

![Fig. 1.](image-url)

Fig. 1. a) Location map of the Ech palaeolake and European sites used for comparisons in the discussion. 1: Lago Piccolo di Avigliana (Larocque and Finsinger, 2008), 2: Lago di Lavarone (Heiri et al., 2007a, b), 3: Maloja Pass (Ilyashuk et al., 2009), 4: Lac Lautrey (Heiri and Millet, 2005), 5: Gerzensee (Lotter et al., in press), 6: Egelsee (Larocque-Tobler et al., 2010), 7: Hijkermeer (Heiri et al., 2007a, b), 8: north-west England (Lang et al., 2010). b) Location map of the Ech palaeolake and some of the existing pollen records in the Pyrenees. 1: La Borde, 2: Ruisseau de Laurenti, 3: Estarrès, 4: Rubal, 5: Balèire, 6: Freychinède, 7: Barbazan, 8: Biscaye, 9: Lake Lourdes, 10: Monge, 11: Castet, 12: Le Moura, 13: Tramacastilla, 14: El Portalet, 15: La Pouretère. 1 to 13 from Jalut et al. (1992), 9 from Reille and Andrieu (1995), 14 from González-Samperiz et al. (2006), 15 from Aubert et al. (2004).
study area is characterised by a mountainous Oceanic climate with mean annual precipitation of 1200 mm (Météo France data, 1931–2006). The mean summer temperature for the site can be estimated to ca. 17.1 °C from the altitudinal correction of the meteorological data monitored at Lourdes (weather station located at 410 m a.s.l.) using a lapse rate of 0.5°/100 m elevation (Agusti-Panareda and Thompson, 2002). Regional vegetation consists of Atlantic-type oak forest mainly dominated by Quercus (Q. robur and Q. pyrenaica) with Corylus and Betula. The north-facing slopes (up to 900 m a.s.l.) are covered by a mixed beech-oak forest (Fagus sylvatica), which is the dominant vegetation type of the bioclimatic hill stage in the area (Dupias, 1985).

The col d’Ech peat bog is 200 m wide and 280 m long. It is a former small glacial lake which is located on a small plateau dammed by a recessional moraine at the south and by a 200 m high south-facing calcareous slope at the north.

3. Material and methods

In 2006, a total of 10 overlapping 1 m long sediment sections were retrieved using a Russian corer (120 mm in diameter) at the centre of the Ech peat bog. All sections were transferred to a PVC liner and kept at 6 °C in a cold-room.

A composite five metre long core was constructed from magnetic susceptibility and gamma density profiles acquired at high resolution (5 mm interval) using a Geotek Multi-Sensor Core device. The lithological profile is typical of a palaeolake basin gradually filled up by sediment and the subsequent development of a peat bog. Along the core, peat accumulation begins at 330 cm below the surface overlying lacustrine deposits consisting of silt, clay and coarse organic remains in changing proportions.

The whole sediment profile was sampled at 5 cm intervals for pollen analysis (30 samples). Some additional levels (3 samples) were analysed between 380 cm and 390 cm i.e. within a sediment unit corresponding to the Younger Dryas (see below). Standard methods involving NaOH (10%), HCl, HF (30%), and acetylsis treatment (5 min) were applied for pollen preparation. The pollen preservation was good and concentrations were always sufficient to count over 450 terrestrial pollen grains per slide. Pollen percentages are based on the pollen sum of arboreal (AP) and non-arboreal (NAP) pollen grains, excluding Alnus, Cyperaceae, aquatic taxa and fern spores.

Chironomid remains and Loss-On-Ignition (LOI) were analysed in the core section covering the Lateglacial and the early Holocene i.e. between 370 cm and 476 cm below the surface (as indicated by main pollen changes in reference to regional vegetation changes during the Late-glacial and early Holocene).

LOI was performed on wet samples taken continguously at 1 cm intervals between 369 and 422 cm and at 3 to 7 cm intervals corresponding to the Younger Dryas (see below). Standard methods involving NaOH (10%), HCl, HF (30%), and acetylsis treatment (5 min) were applied for pollen preparation. The pollen preservation was good and concentrations were always sufficient to count over 450 terrestrial pollen grains per slide. Pollen percentages are based on the pollen sum of arboreal (AP) and non-arboreal (NAP) pollen grains, excluding Alnus, Cyperaceae, aquatic taxa and fern spores.

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4. Results

4.1. Lithology and LOI

Between 360 and 500 cm, 6 sediment units (SU) were identified based on changes in lithology (i.e. texture, grain size and colour) and organic content of the sediment (Fig. 2):

At the lowest part of the record, SU 1 (500–485 cm) consisted of inorganic palgglacial blue clay. SU 2 (485–450 cm) was characterised by light grey silty-clay with low organic content (LOI<8%). Sediments between 450 and 420 cm (SU 3) consisted of grey-brown silt with a two samples, with a maximum relative abundance of more than 2%, were included for further analysis.

Stratigraphic diagrams of the relative abundance of pollen and chironomid taxa were produced using TView v. 2.0.2 (Grimm, 2004). The chironomid assemblage zones were defined from a stratigraphically constrained cluster analysis performed by Conis (Grimm, 2004). Ordinations (DCA, CA and CCA) were performed using ade4 (Chessel et al., 2004), the Vegan packages for R and Canoco for Windows version 4.52 (ter Braak and Smilauer, 2002).

Chironomid-inferred July air temperatures were reconstructed using the chironomid- temperature transfer-function described by Heiri et al. (2011). This transfer-function is based on a modern calibration dataset from the Alpine and Norwegian regions and Weighted Averaging-Partial Least-Squares regression (WA-PLS; ter Braak and Juggins, 1993; ter Braak et al., 1993). The overall Root Mean Square error of Prediction (RMSEP) of the transfer-function and estimated standard errors of prediction were calculated for the individual fossil samples using bootstrapping (9999 bootstrap cycles; Birks et al., 1990). Fossil assemblages with a squared $\chi^2$-distance to the most similar assemblage in the modern calibration dataset larger than the 2nd and the 5th percentile of all squared $\chi^2$-distances in the modern data were identified as samples with ‘no close’ and ‘no good’ analogue, respectively (Birks et al., 1990; Jones and Juggins, 1995; Heiri et al., 2003).

Fossil samples were added passively in a canonical correspondence analysis (CCA) of the modern calibration data with July air temperature as the only constraining parameter. Fossil samples with a residual distance to the first CCA axis larger than the 90th and 95th percentile of the residual distances of all the modern samples were identified as samples with a ‘poor fit’ and a ‘very poor fit’ with temperature, respectively (Birks et al., 1990). Chironomid taxa with a Hill’s N2 below 5 in the modern calibration dataset were considered to be rare in the modern dataset (Heiri et al., 2003). WA-PLS Hill’s N2 values, and squared $\chi^2$-distances were calculated with the programme C2 (Juggins, 2003) version 1.4.3. All calculations are based on square-root transformed percentage abundances.

Between 320 cm and 455 cm, 7 samples of terrestrial material (2) or bulk sediment (5) were submitted for AMS dating to the Poznan Radiocarbon Laboratory (Table 1). The 14C AMS dates were calibrated to years before present (cal yr BP) with Calib v. 6.0.1 (Stuiver and Reimer, 1993) using the Incal09 calibration curve (Reimer et al., 2009).

Table 1

<table>
<thead>
<tr>
<th>Depth (cm)</th>
<th>Radiocarbon date (BP)</th>
<th>Material</th>
<th>Calibrated age (2 sigma) (cal yr BP)</th>
<th>Median probability</th>
</tr>
</thead>
<tbody>
<tr>
<td>329.5</td>
<td>7340±40</td>
<td>Bulk</td>
<td>8020–8300</td>
<td>8134</td>
</tr>
<tr>
<td>380.5</td>
<td>9890±50</td>
<td>Bulk</td>
<td>11,200–11,600</td>
<td>11,292</td>
</tr>
<tr>
<td>390.5*</td>
<td>11,960±60</td>
<td>Bulk</td>
<td>13,650–13,990</td>
<td>13,819</td>
</tr>
<tr>
<td>390.5*</td>
<td>11,280±110</td>
<td>Charcoal</td>
<td>12,885–13,385</td>
<td>13,170</td>
</tr>
<tr>
<td>406.5*</td>
<td>12,730±80</td>
<td>Charcoal</td>
<td>14,625–15,585</td>
<td>15,093</td>
</tr>
<tr>
<td>420.5*</td>
<td>12,990±70</td>
<td>Bulk</td>
<td>15,120–16,320</td>
<td>15,610</td>
</tr>
<tr>
<td>454.5</td>
<td>14,550±160</td>
<td>Bulk</td>
<td>17,155–18,435</td>
<td>17,695</td>
</tr>
</tbody>
</table>
moderate organic content (LOI between 15 and 20%) and some coarse unidentifiable organic fragments probably of aquatic origin. This section was overlain by a layer (SU 4: 420–387 cm) of brown organic silt (LOI between 20 and 30%) with coarse unidentifiable organic aquatic remains. SU 5 (387–383 cm) was a thin layer of grey silty clay with moderate organic content (LOI<18%). Between 383 and 360 cm (SU 6) sediments consisted of dark-brown silt with a high organic content (LOI between 40 and 50%) and abundance of coarse organic remains.

4.2. Pollen-stratigraphy and Lateglacial climate events

The pollen stratigraphy between 500 and 360 cm is typical for the Lateglacial-early Holocene vegetation succession in the French Pyrenees as previously reconstructed by Jalut et al. (1988; 1992; 1996), Reille and Andrieu (1995) and Aubert et al. (2004). The boundaries of the main Lateglacial climate events can be defined from the pollen succession of the Ech record (Fig. 2).

In the lowest part of the sequence (i.e. during SU-1 and 2), pollen percentages are strongly dominated by *Artemisia* corresponding to steppe communities reconstructed in the Pyrenees before 14,500 14C BP (around 17,500 cal yr BP, Jalut et al., 1992). The presence of arboreal pollen (AP) such as those of *Pinus* (up to 30% during SU-1), *Quercus* and *Betula* is probably linked to isolated stands or long distance transport of tree pollen from refugium areas (Jalut et al., 1992).

During SU-3, the pollen record is marked by the increase in the relative abundance of *Juniperus* and the converse slight decrease in *Artemisia*. This is in close agreement with numerous pollen diagrams from the French Pyrenees which indicated the development of a landscape dominated by *Juniperus* after 14,500 14C BP (around 17,500 cal yr BP, Jalut et al., 1992). Sediment units 1 to 3 can then be correlated to the Oldest Dryas.

The beginning of SU-4 is characterised by the rapid rise in *Betula*, and the corresponding drop in *Juniperus* and *Artemisia* percentages, while Poaceae pollen remains abundant. The expansion of *Betula* is one of the key botanical events for the beginning of the Bølling in the area (Jalut et al., 1992). *Betula* and Poaceae were probably components of steppe or wooded steppe communities. In the second part (i.e. after 407 cm), the pollen diagram features a strong increase in *Pinus* relative abundance (up to 30%) and a parallel slight decrease in Poaceae percentages. This spread of *Pinus* is typical for the beginning of the Allerød (Jalut et al., 1992).

At the SU-4/SU-5 transition, the pollen diagram shows the decrease in *Pinus* and *Betula* percentages together with the increase in *Artemisia*, Poaceae and *Juniperus*. This re-advance in steppe elements and the opening up of the landscape is a characteristic feature of the Younger Dryas in the western French Pyrenees (Jalut et al., 1992; Aubert et al., 2004).

The onset of SU-6 is marked by a very sharp rise in AP with a strong dominance of *Quercus* pollen and the presence of *Pinus* and *Betula* while *Artemisia* percentages decrease. The increase in AP percentages and the expansion of *Quercus* indicate the beginning of the Holocene (Jalut et al., 1988; 1992).

4.3. Radiocarbon age reliability and age/depth model

In the Ech record, the coarse organic remains found within SU-3 to 6 consisted mainly of unidentifiable fibrous and translucent fragments and mosses. Despite the processing of a large number of sediment samples along the core (notably during the sieving for the chironomid analysis), unquestionably terrestrial remains (charcoal particles) were found in sufficient quantities for AMS radiocarbon dating in only two samples (Table 1). Other dates were produced from bulk sediment, implying a possible ageing effect due to the hard-water effect or allochtonous contamination by old carbon.
reliability of these radiocarbon dates must be carefully assessed under the light of the age determined for main lateglacial climate events and vegetation succession in the Pyrenees and in northwest Spain (Jalut et al., 1992; Reille and Andrieu, 1995; Allen et al., 1996; González-Samperíz et al., 2006; Muñoz-Sobrino et al., 2007).

The radiocarbon date obtained from 454.5 cm ([16,820–18,070] cal yr BP) is in agreement with the age of the spread of Juniperus which occurred after ca. 15,700 cal yr BP ([14,500 ¹⁴C BP) in the French Pyrenees (Jalut et al., 1992). At 420.5 cm, the radiocarbon date ([15,120–16,320] cal yr BP) seems slightly too old. Indeed, it just precedes the Oldest Dryas/Interstadial transition (at 419.5 cm) which was dated to 14,700 cal yr BP in the Spanish Pyrenees (González-Samperíz et al., 2006), and between ca. 14,500 cal yr BP ([12,425 ¹⁴C BP, Allen et al., 1996] and 14,800 cal yr BP (Muñoz-Sobrino et al., 2007) in northwest Iberia. The radiocarbon date obtained at 406.5 cm was rejected since it gave an aberrant age of 15,100 cal yr BP ([14,600–15,600] cal yr BP), which is several hundred years older than what would be expected for a sample from the Lateglacial Interstadial. At 390.5, two radiocarbon dates were obtained, a first from bulk material and a second from fine charcoal particles. The ages derived from bulk material (Table 1) imply a long break in sedimentation rate around 390 cm which is incompatible with both the sediment description and LOI; therefore this radiocarbon date was rejected. The age obtained from charcoal particles which occurred after ca. 17,500 cal yr BP (14,500 ¹⁴C BP) in the study area (Lake Lourdes, Reille and Andrieu, 1995), in the Spanish Pyrenees and in northwest Spain (González-Samperíz et al., 2006; Muñoz-Sobrino et al., 2007). Several studies showed that radiocarbon dates performed on charcoal particles can produce an ageing effect by ranges of several decades to centuries (e.g. Gavin, 2001). The two remaining radiocarbon dates (at 329.5 and 380.5 cm) are in agreement with ages found for the beginning of the Holocene in the study area (Lake Lourdes, Reille and Andrieu, 1995), in the Spanish Pyrenees and in northwest Spain (González-Samperíz et al., 2006; Muñoz-Sobrino et al., 2007).

A composite age-depth model using linear interpolation was built from the three accepted radiocarbon dates (Table 1) and ages estimated for the main Lateglacial transitions (Oldest Dryas/Interstadial, Interstadial/Younger Dryas) in the Pyrenees and northwest Iberia (Table 2 and Fig. 3).

4.4. The chironomid record

Chironomid head capsules were abundant in the whole record except for SU-1 where no remains were found. In 10 samples, counts were below the minimum of 45 remains usually recommended for palaeoenvironmental reconstruction (Heiri and Lotter, 2001). In only two samples (at 389.5 and 445.5 cm) counts were less than 20 remains (Fig. 4). A total of 46 taxa were identified in the Ech record, 34 of them met the minimum frequency criteria and were included in further analyses.

Two main biozones (Ecz-1 and 2) and 5 subzones (from Ecz-2a to Ecz-2-e) were recognised within the Ech chironomid stratigraphy from the cluster analysis (Fig. 4). A correspondence analysis (CA) was performed from the sample/taxa matrix (52 samples, 34 taxa). Major changes in sample scores along axis 1 and 2 (Fig. 4) corresponded to the main turn-over in chironomid assemblages as depicted by the cluster analysis. Characteristic taxa for each bionzone and sub-biozone were defined from the projection of the samples and the taxa in the plane of the first two axis of the CA (Fig. 5). Only taxa with relative contribution >15% to either axis one and/or two were considered and plotted.

4.4.1. Ecz-1

The samples of this biozone were clearly separated from the rest of the chironomid record in the CA biplot and were projected at the lower end of axis one. In descending order of relative abundance, the taxa characterising the bionzone are: Chironomus anthracinus-type, Tanytarsus lugens-type, Microspectra insignilobus-type, Orthocladius spp. and Psectrocladius sordidellus-type.

4.4.2. Ecz-2

As witnessed by the strong increase in sample scores along axis one, the Ecz-1/2 transition is marked by an important shift in the chironomid community. Dominant taxa of Ecz-1 sharply decline. Some of them like Microspectra insignilobus-type and Psectrocladius sordidellus-type definitively disappeared from the chironomid record. The biozone Ecz-2 is characterised by the appearance of a number of new taxa. Considered as a whole, samples of Ecz-2 have high scores along axis one and are characterised by Tanytarsus pallicicornis-type II, Polypedilum nubeculosum-type, Cladotanytarsus, Microtendipes, Glyptotendipes and Pseuderatomyia.
Fig. 4. Percentage diagram of the most common chironomid taxa during the Lateglacial, together with sample scores of axis one and two of a correspondence analysis. Dotted lines: stratigraphic boundaries of the main Lateglacial climate events, full lines: limits of chironomid biozones from the cluster analysis CONISS (Ecz: Ech chironomid Zone).
Samples of subzones 2a and 2c are plotted at the positive side of axis two. They are characterised by *Tanytarsus pallidicornis*-type II, *Glyptotendipes* and *Ablabesmyia*. Ecz-2c differs from 2a in the abundance of *Chironomus plumosus*-type (Figs. 4 and 5).

Samples of Ecz-2b and 2d, located at the negative end of the axis 2, are typified by high relative abundance of *Microtendipes*, and *Polypedilum nubeculosum*-type. Ecz-2d is further marked by slightly higher relative abundance of *Tanytarsus lugens*-type than Ecz-2b and the unique presence of *Tribelos* for the whole record. The samples of the biozone Ecz-2e are located at the high end of axis one. The assemblages are dominated by *Microtendipes*, *Tanytarsus pallidicornis*-type II and *Polypedilum nubeculosum*-type while *Tanytarsus lugens*-type and *Chironomus anthracinus*-type disappear.

### 4.5. Quantitative inference of temperatures

Chironomid-based temperature estimates were produced from the Ech record using a chironomid-temperature calibration dataset and transfer function based on 254 lakes sampled in Switzerland and Norway. The lakes cover a July air temperature gradient of 3.5–18.4 °C, a latitudinal gradient of 46.1–79.8°N, an altitudinal gradient of 5–2815 m a.s.l. and include lakes on acid, basic and carbonaceous bedrock. This dataset was produced by taxonomic amalgamation of two regional calibration datasets from Norway and Svalbard (Brooks and Birks, 2001; 2004) and from the Swiss Plateau, the Jura Mountains and the northern, central and southern Swiss Alps (Heiri and Lotter, 2005; 2010) and describes the distribution of 154 chironomid taxa in respect to mean July air temperature (Heiri et al., 2011). Since the dataset includes lakes in temperate lowland, alpine and arctic environments it can be expected that it includes most environmental conditions prevailing during the Lateglacial period in Central and Northern Europe and in southern European mountain regions.

A chironomid-based inference model for temperature was developed from this calibration dataset using WA-PLS regression with the number of useful WA-PLS components evaluated according to Birks (1998). This resulted in a two component WAPLS regression model. When evaluated in the modern environment using bootstrapping techniques the inference model predicts mean July air temperature based on chironomid assemblages with a RMSEP of 1.55 °C and a coefficient of determination of 0.84.

The transfer function reconstructed July air temperature variability during the Lateglacial (Fig. 6) that can be summarised as follows:

Prior to 14,700 cal yr BP, i.e. during SU 2 and 3 corresponding to the Oldest Dryas, the summer temperatures were the coldest for the whole record and fluctuated between 10 and 13 °C (mean: 11.7 °C). At the Oldest Dryas/Lateglacial Interstadial transition, the transfer function reconstructed an abrupt increase in July air temperatures from ca. 11 °C to ca. 16 °C. The temperature record displayed a high internal variability during the Lateglacial Interstadial. In a first step, temperature continued to increase to reach around 17.5 °C at ca. 14,000–14,200 cal yr BP. The following period was marked by a gradual cooling of around 1 °C interrupted by 3 short and rapid cold events of 1.5 °C magnitude at ca. 13,100 cal yr BP, 13,600 cal yr BP and 13,900 cal yr BP. At the Lateglacial Interstadial/Younger Dryas transition, inferred July air temperature shifted from 16.8 °C to values...
ranging from 14.8 °C to 15.6 °C (mean: 15.2 °C) during the Younger Dryas. At the beginning of the Holocene, July air temperature increased by about 2 °C to reach values around 17 °C.

Individual low counts (at 389.5 and 445.5 cm) did not seem to induce a major bias in the reconstruction since the inferred temperatures for the corresponding samples did not significantly differ from the adjacent levels and/or they were consistent within a trend of temperature changes. The sample specific standard errors of prediction (eSEP) for the temperature reconstruction were between 1.52 and 1.77 °C. All taxa found in the Ech sediment sequence were included in the calibration data set. Taxa rare in the modern data set were present in the Ech fossil record at low percentages ranging from 0 to 13.2% (Fig. 6). Samples of the Ech record generally have a good fit with temperature as estimated from the comparison between their residual distance to the first CCA axis with July air temperature as the only constraining factor and the 90th and the 95th percentiles of residual distances of modern samples to the first axis in a Canonical Correspondence Analysis used to determine samples with ‘very poor’ (0.95) and ‘poor’ fit (0.90) with temperature (see text for details). Stratigraphic boundaries of the main Lateglacial climate events are indicated (YD: Younger Dryas).

5. Discussion

The reliability of our Lateglacial summer temperature reconstruction must be carefully addressed for three main reasons:

(i) The chronology of the Ech record is based on a limited number of accepted radiocarbon dates and is mainly constrained by the palynostratigraphy. Even if this approach has given satisfying results in other studies (Heiri et al., 2007a, b; Lotter et al., in press), this can still possibly induce biases in the age assessment for some levels. More particularly, the age/depth relationship within the Lateglacial Interstadial is built from a linear interpolation. This implies a constant sediment accumulation rate, which may be regarded as unrealistic given the high environmental variability during this interval.

(ii) The transfer function used to infer summer air temperature from the Ech fossil chironomid record was based on modern calibration data from the Alps and Norway. The question therefore arises about the applicability of a mixed Swiss–Norwegian calibration data set to infer temperature from a Pyrenean record.

(iii) Apart from the Oldest Dryas/Interstadial transition, the magnitudes of changes in summer temperatures inferred from the Ech samples were low and remained in the range of the sample specific estimated standard errors of prediction of the inference model. In times of low amplitude temperature changes, local environmental and limnological factors may potentially control the chironomid response and lead to biases in temperature reconstructions (Velle et al., 2010).

Fig. 6. a) Chironomid-inferred July air temperatures from the Ech record plotted together with sample-specific estimated standard errors of prediction (eSEPs), b) nearest modern analogues for the fossil samples in the calibration dataset, c) goodness-of-fit of the fossil samples with temperature, and d) the percentage of identified chironomid head capsules rare in the calibration dataset (N2<5). Horizontal lines indicate the 2nd and 5th percentiles of all squared $\chi^2$-distances in the calibration dataset used to identify samples with no close (2%) and no good (5%) modern analogue, and the 90th and the 95th percentiles in residual distances of modern samples to the first axis in a Canonical Correspondence Analysis used to determine samples with ‘very poor’ (0.95) and ‘poor’ fit (0.90) with temperature (see text for details). Stratigraphic boundaries of the main Lateglacial climate events are indicated (YD: Younger Dryas).
In order to validate the temperature reconstruction derived from Ech chironomid record, the following discussion comprises three sub-sections dealing with (i) the applicability of the combined Swiss–Norwegian transfer function to a Pyrenean record; (ii) the controlling factors of the Lateglacial chironomid succession from Ech (iii) the comparison of the reconstructed temperature with other similar studies in Europe and the Greenland δ¹⁸O record.

5.1. Applicability of the transfer function

To date, there is no modern training set published from the Pyrenean massif. The transfer function used in this study is based on the distribution of chironomid taxa in the Swiss Alps and in Norway. The possibility of using a training set from one region to reconstruct climate in another region was previously addressed by Lotter et al. (1999) and Larroque-Tobler (2010). Both studies concluded that transfer functions from other regions can be applied, however the magnitude of changes should be taken with caution. Larroque-Tobler (2010) indicated that fossil record and modern calibration data must share similar taxa. A representation of 80% of the taxa from the sediment record in the modern calibration data was suggested as a minimum criterion for the applicability of any transfer function. This quantitative prerequisite is met by the Ech chironomid record: all the fossil taxa are represented in the transfer function and total percentages of rare taxa in the transfer function are less than 14% in any of the Ech samples.

From the beginning of the Interstadial to the early Holocene, the fossil samples of the Ech record had no close or good analogue (assessed using a χ²-distance computation) in the modern calibration data used for the transfer function. Equivalent poor analogue situations were encountered for significant portions of fossil records located in the same region as the modern calibration data used to develop the transfer function (e.g. Ilyashuk et al., 2009; Larroque-Tobler et al., 2010). However, WA-PLS regression tends to perform well under no-analogue conditions, as long as the fossil taxa of high numerical importance are well-represented in the modern calibration dataset (Birks, 1998). The applicability of the mixed Swiss–Norwegian transfer function to the Ech fossil record is further supported by the goodness-of-fit statistics as few samples are above the 0.90 and 0.95 percentile cut-levels set for samples with a poor and very poor fit with temperature (Fig. 6).

The other qualitative prerequisite for the applicability of the transfer function to the Ech fossil samples is a similar relationship of taxa with temperature between the Pyrenees and the modern calibration data. Lotter et al. (1999) found that the temperature optima of fossil taxa from North America and Europe were similar and highly correlated even from these two different biogeographical zones. Likewise, Heiri et al. (2011) demonstrate that the relationship of taxa to temperature is similar in the Norwegian and Swiss training sets used in the present study. Furthermore, in the early 70’s, Laville (1971) carried out an extensive study of chironomid assemblages in nine Pyrenean lakes distributed along an altitude gradient. He concluded that, at that time, the lakes studied shared 85% of their chironomid species with lakes in the Alps. Moreover, his results showed that relative distribution of taxa with respect to altitude was similar between the two regions. The only difference was the higher altitude reached by taxa in the Pyrenees. This finding was taken as evidence of an influence of temperature on the chironomid distribution at a broad scale in the Alps and in the Pyrenees since climate is warmer in the Pyrenean Massif than in the Alps at equivalent altitude. It also implies that individual taxa have similar temperature preferences, irrespective of their geographical origin, in the Alps or the Pyrenees.

In summary, both quantitative (representation of fossil taxa in the transfer function and goodness of fit statistic) and qualitative interpretations (equivalent relationship of taxa towards temperature in the Pyrenees and in Swiss and Norwegian lakes) support the applicability of the Swiss/Norwegian transfer function to the Pyrenean Ech record.

5.2. Controlling factors of the Ech chironomid succession during the Lateglacial

In lakes, chironomid assemblages are affected by a variety of environmental conditions interacting in complex ecological processes (Brodersen and Quinlan, 2006; Velle et al., 2010; Eggertmont and Heiri, 2011). In descending order of scale, from regional to ecosystem to microhabitat, the most commonly recognised influencing factors are temperature (Walker et al., 1991), trophic status (Brooks et al., 2001), oxygen conditions (Verneaux and Aleya, 1998), nature and amount of organic matter in the sediment (Brinkhurst, 1974; Verneaux and Aleya, 1998), and type of substrate (Mcgarigle, 1980; Palomaki, 1989). Inferring temperature changes from a chironomid sequence using a transfer function assumes that the chironomid assemblages respond, directly or indirectly, to temperature. In times of low amplitude temperature changes, the influence of temperature may be masked by the influence of other environmental factors on the chironomid response as suggested by Velle et al. (2010) for some Holocene Fennoscandian records. The Lateglacial was characterised by much stronger changes in summer temperatures throughout the continent and most published chironomid-inferred temperature records are regionally consistent (Lang et al., 2010). Nevertheless, even in lateglacial records, temperature reconstruction can occasionally be compromised by the influence of local parameters on the chironomid fauna, as highlighted in the study of a lowland Mediterranean lake that experienced water-level fluctuations (Millet et al., 2007).

Since our main objective was to infer the lateglacial temperature from chironomids, the study site was chosen in order to maximise the climate influence on the chironomid community. In deep lakes, the relationship between subfossil chironomid assemblages and climate can be complicated by several interacting factors (e.g. Verbruggen et al., 2011): (1) the strong influence on the profundal community of oxygen conditions, food sources and quantity, and trophic functioning, (2) the bathymetric distribution of living larvae from the littoral to the deep zone, and (3) the variable redeposition of littoral remains in the deepest zones. In small and shallow systems, such as the Ech palaeolake, air temperature is expected to be an important, direct or indirect, influencing factor for distribution and abundance of chironomid taxa (Eggertmont and Heiri, 2011). Furthermore, in these lakes the chironomid head capsules of the surface layers are to some extent redeposited and mixed and this reduces the spatial variation of living larva within the lake (Brodersen and Lindeggard, 1997; Van Hardenbroek et al., 2011). Thus a single sample taken in the middle of the basin may be considered to be representative for the fauna of the entire lake.

Despite the study of a record which was a priori considered suitable for temperature reconstruction, the temperature influence on the chironomid succession in the Lateglacial should also be assessed with reference to known ecological preferences of the main taxa involved. The most striking feature of the Ech chironomid record is the distinct change of chironomid assemblages between Ec1-1 and Ec2-2 corresponding to the Oldest Dryas/Lateglacial Interstadial transition. During the Oldest Dryas, the chironomid community was dominated by Tanytarsus lugens-type, Chironomus anthracinus-type and Microspectra insignilobus-type, taxa commonly found in modern lakes between 1500 and 2000 m in the Alps (Heiri and Lotter, 2010). In the Pyrenees, Laville (1971) studied the distribution of chironomid species among nine lakes from 2100 to 2400 m a.s.l. The species included Tanytarsus bathophilus (Tanytarsus lugens fossil morphotype) and Microspectra contracta (Microspectra insignilobus fossil morphotype), which were recognised as cold stenothermic species from their distribution in shallow cold lakes.

In order to validate the temperature reconstruction derived from Ech chironomid record, the following discussion comprises three sub-sections dealing with (i) the applicability of the combined Swiss–Norwegian transfer function to a Pyrenean record; (ii) the controlling factors of the Lateglacial chironomid succession from Ech (iii) the comparison of the reconstructed temperature with other similar studies in Europe and the Greenland δ¹⁸O record.
The chironomid fauna of Ecz-2, characterised by taxa such as *Tanytarsus pallidicornis*-type II, *Polypedilum nubeculosum*-type, *Glyptotendipes* and *Microtendipes* was analogous to that of lowland and intermediate-altitude modern lakes in the Alps (Heiri and Lotter, 2010). Compared to taxa characterising the Oldest Dryas, taxa of Ecz-2 have warmer optima in modern training sets (e.g. Heiri et al., 2011). During Ecz-2, in addition to the influence of temperature, some local factors may have influenced individual taxa. For example, the increase in *Glyptotendipes* may have been linked to the growth of aquatic macrophytes (Pinder and Reiss, 1983; Klink and Moller Pillot, 2003), *Microtendipes* may have been favoured by the increase in sediment grain size and a higher proportion of coarse organic fragments (McGarrigle, 1980), and *Polypedilum nubeculosum*-type and *Chironomus plumosus*-type may have responded to the increase in organic matter in the sediment (Maitland, 1979; Wolfram, 1996; Millet et al., 2007). Nevertheless, the above mentioned taxa all have high temperature optima in both the Norwegian and Swiss calibration sets and so the overall influence of temperature change on the shift of the whole chironomid assemblage from Ecz-1 to Ecz-2 remains unquestionable even if local parameters may also account for some of the variability in particular taxa. Indeed, goodness-of-fit analysis shows that all but a few of the fossil samples in the entire sequence have a strong response to temperature.

The distribution of samples and taxa along axis 2 of the CA plot (Fig. 5) suggests that the chironomid response within Ecz-2 (from Ecz-2a to Ecz-2e) is more subtle compared to the major taxonomic turnover at the Ecz-1/Ecz-2 transition. There is no clear relationship between taxon scores along axis 2 and their temperature optimum. This implies that the differences in the chironomid assemblages do not reflect different temperature conditions. Sample scores along axis 2 were also not correlated to LOI values during Ecz-2. These results suggest that multiple interacting factors probably controlled the chironomid succession depicted by the sample trajectory in the CA. In this context, trying to disentangle respective influence of temperature, habitat, and organic matter is difficult. Nevertheless, the high relative abundance of *Tanytarsus lugens*-type (up to 20%) and correspondingly lower percentages of lowland taxa such as *Chironomus plumosus*-type, *Polypedilum* and *Tanytarsus pallidicornis*-type during Ecz-2d, Ecz-2b and at the beginning of Ecz-2a suggest the prevalence of colder conditions during these phases.

The overall control of temperature on the assemblage changes is well established for the Oldest Dryas/Interstadial transition, but is more difficult to assess throughout the subbiozones of Ecz-2 when multiple factors probably interacted in complex processes. Therefore, the inference results will be further validated by comparison with other Lateglacial climate records.

### 5.3. Comparison of the Ech temperature reconstruction with other European chironomid records and the Greenland δ18O record

The chironomid-based temperature reconstruction from Ech displays the main lateglacial climate trends expected for the North Atlantic region as inferred by glacial and continental records. The succession through a cold Oldest Dryas (equivalent to GS-2 in NGRIP), a warmer Lateglacial Interstadial (GI-1 in NGRIP), a cold reversal during the Younger Dryas (GS-1 in NGRIP) and a warming at the beginning of the Holocene is clearly shown in the Ech record (Fig. 7). Furthermore, at the centennial-scale, the climate variability suggested by the Ech chironomid record is also in good agreement with the δ18O variations in the Greenland ice cores during the Lateglacial Interstadial (Fig. 7). The reconstruction of summer temperature changes derived from the Ech record can be compared to other available chironomid-based temperature reconstructions throughout Europe.

#### 5.3.1. The Oldest Dryas/Interstadial transition

During the final part of the Oldest Dryas, the summer temperatures reconstructed at Ech were 2–3 °C lower than inferred from existing chironomids records in the Southern Alps (Table 3a,b). The temperature inferences from the Ech chironomids are in agreement

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**Fig. 7.** Chironomid-inferred July air temperatures from Ech compared with the NGRIP δ18O record (Rasmussen et al., 2006). Age estimates for individual chironomid samples were assigned from the age-depth model.
with the reconstructions derived from palaeobotanical data in the French and Spanish Pyrenees which indicate July temperature around 10.7 °C in the region at the elevation of Ech (Renssen and Isarin, 2001).

At Ech, the temperature increase (ca. 6 °C) at the Oldest Dryas/Interstadial transition was almost twice as high as in chironomid records from southern Europe (Table 3b) and Switzerland (Egelsesse, Larocque–Toberle et al., 2010; Gerzensee, Lotter et al., in press) and higher than the French Jura (Lake Lautrey, Heiri and Millet, 2005) where the amplitude of change in chironomid-inferred temperature was around 3–4 °C (Table 3b). An increase in summer temperature of around 6 °C in south–western France is consistent with the reconstructions and simulations carried out by Renssen and Isarin (2001). The Ech record provides new insights into the existence of a strong West–east gradient in July air temperature during the Oldest Dryas from the Atlantic margin (colder) to the Mediterranean regions (warmer) along latitudes 44–45°N. During the Oldest Dryas, the Ech July air temperatures were probably influenced by the cold conditions prevailing over the North Atlantic Ocean. Indeed, during the Oldest Dryas the summer and winter sea–ice margins were inferred to be at approximately 60°N and 45°N, respectively (Ruddiman and McIntyre, 1981; Renssen et al., 2002). Over the sea ice, the air was strongly cooled and consequently it was a potential source of cold air during summer for the west European coast.

5.3.2. The Lateglacial Interstadial

The temperatures reconstructed during the Interstadial were in the same range as in the existing chironomid-based reconstructions from the Southern Alps (Table 3a).

After the sharp rise corresponding to the Oldest Dryas/Interstadial transition, the inferred summer temperature continued increasing with a gentler slope, passing from 16 °C at the onset of the Interstadial to 17.5 °C at ca. 14,100 cal yr BP. Given the uncertainties of the Ech record chronology, the hypothesis of a step-wise temperature reconstruction must be carefully addressed and supported by comparison with other climate records.

This seems similar to the two-step warming evidenced in some European oxygen isotope (Ammersee; von Grafenstein et al., 1999; Hawes water; Marshall et al., 2002) and chironomid records (Lake Lautrey, Heiri and Millet, 2005). Nevertheless, the significance of this summer temperature increase at the beginning of the Interstadial still remains an open question. Indeed, this trend differs from the Greenland oxygen isotope records (Rasmussen et al., 2006) and from some European records (e.g. Eicher and Siegenthaler, 1976; Lotter et al., 1992) where the Oldest Dryas/Interstadial transition is marked by a single major abrupt warming shift in climate proxies. Apart from effective change in climate, the reconstruction of a gradual increase in temperature could potentially be explained by a time lag in the response of Ech chironomid assemblages to the warming at the Oldest Dryas/Interstadial transition. However, this temperature increase was not due to the appearance of new warm-adapted taxa but corresponded to a progressive decrease of taxa with cold optima (e.g. Tanytarsus lugens-type) and the corresponding increase in taxa with warmer optima, such as Tanytarsus pallidicornis-type II (Fig.4). This suggests the influence of continuous changing environment and/or climate influencing chironomid succession rather than an immigration lag of warm-adapted chironomid taxa. Indeed, the winged adults are dispersed widely by the wind and so chironomids can rapidly colonise lakes. Furthermore, the Ech Palaeolake was a mountain lake and it is expected that most temperate chironomid taxa were already present in the adjacent lowlands during the Oldest Dryas. Another alternative hypothesis would be the influence of other factors than temperature on chironomid succession at the beginning of the Interstadial. The trend in the temperature reconstruction (increase to reach around 17.5 °C at ca. 14,000–14,200 cal yr BP) in the early Interstadial seems to follow closely an increase in organic matter in the sediment (Fig. 2). This trend in organic matter might originate from an increase in allochthonous inputs following soil and vegetation development on the watershed combined with a rise in autochthonous production following the abrupt warming of the onset of the Interstadial. Since organic matter is an important controlling factor for chironomid assemblages in lakes (Verneaux and Aleya, 1998), the increase in organic matter might have provoked some changes in chironomid assemblages indirectly linked with climate and this may have influenced the temperature reconstruction.

From ca. 14,100 cal yr BP onwards, a gradual decrease in the inferred summer temperature of ca. 1 °C until the end of the Interstadial is in accordance with numerous climate records including oxygen isotopes from Greenland (Rasmussen et al., 2006) and lake sediments (Lotter et al., 1992; Marshall et al., 2002), and temperature reconstructions from pollen (Peyron et al., 2005) and chironomids in England (Lang et al., 2010), Norway (Brooks and Birs, 2000), south-western Alps (Larocque and Finsinger, 2008) and Switzerland (Larocque–Toberle et al., 2010).

This gradual cooling throughout the Interstadial is punctuated by three short-duration cold events. They are broadly similar in magnitude (ca. 1.5 °C), which lies in the range of the error of prediction of the inference model (Fig. 6). Nevertheless, the concordance in the timing and amplitude of these events with climate changes reconstructed from other records in Europe highlights the sensitivity of the Ech temperature reconstruction. The short cooling events dated at ca. 13,900 cal yr BP and at ca. 13,100 cal yr BP may respectively correlate to the GI-1d event in the Greenland event stratigraphy (Lowe et al., 2008), equivalent to the Agelisse Oscillation on the Swiss Plateau (Lotter et al., 1992), and to the GI-1b event (Lowe et al.,

### Table 3a

<table>
<thead>
<tr>
<th>Site</th>
<th>Altitude</th>
<th>Latitude</th>
<th>Longitude</th>
<th>Oldest Dryas</th>
<th>Interstadial</th>
<th>Younger Dryas</th>
<th>Early Holocene</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ech palaeolake</td>
<td>710 m a.s.l</td>
<td>43°04'N</td>
<td>0°5'W</td>
<td>11 °C</td>
<td>17.5–16.8 °C</td>
<td>15.2 °C</td>
<td>17 °C</td>
</tr>
<tr>
<td>Lago Piccolo di Avigliana (1)</td>
<td>365 m a.s.l</td>
<td>45°03'N</td>
<td>7°23'E</td>
<td>14 °C</td>
<td>16.7–15.5 °C</td>
<td>14 °C</td>
<td>16.5 °C</td>
</tr>
<tr>
<td>Lago di Lavaronne (2)</td>
<td>100 m a.s.l</td>
<td>45°56'N</td>
<td>11°15'E</td>
<td>13 °C</td>
<td>16.2–17.6 °C</td>
<td>15.6 °C</td>
<td>18.1–18.7 °C</td>
</tr>
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</table>

### Table 3b

<table>
<thead>
<tr>
<th>Site</th>
<th>Altitude</th>
<th>Latitude</th>
<th>Longitude</th>
<th>OD/LI</th>
<th>LI/YD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ech palaeolake</td>
<td>43°04'N</td>
<td>0°5'W</td>
<td>+5/+6 °C</td>
<td>-1.5 °C</td>
<td></td>
</tr>
<tr>
<td>Lago Piccolo di Avigliana (1)</td>
<td>45°03'N</td>
<td>7°23'E</td>
<td>+3 °C</td>
<td>-1.5 °C</td>
<td></td>
</tr>
<tr>
<td>Lago di Lavaronne (2)</td>
<td>45°56'N</td>
<td>11°15'E</td>
<td>+3 °C</td>
<td>-2 °C</td>
<td></td>
</tr>
<tr>
<td>Maloja Pass (3)</td>
<td>46°24'N</td>
<td>9°41'E</td>
<td>-3.5 °C</td>
<td>3.5 °C</td>
<td></td>
</tr>
<tr>
<td>Lake Lautrey (4)</td>
<td>46°35'N</td>
<td>5°51'E</td>
<td>+4/+5 °C</td>
<td>-2.5/-3.5 °C</td>
<td></td>
</tr>
<tr>
<td>Gerzensee (5)</td>
<td>46°50'N</td>
<td>7°33'E</td>
<td>+2/+3 °C</td>
<td>-1.5 °C</td>
<td></td>
</tr>
<tr>
<td>Egelsesse (6)</td>
<td>47°11'N</td>
<td>8°35'E</td>
<td>+3/+4 °C</td>
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<tr>
<td>Hijkerveer (7)</td>
<td>52°53'N</td>
<td>6°29'E</td>
<td>-4 °C</td>
<td>-1.5 °C</td>
<td></td>
</tr>
<tr>
<td>North-west (8)</td>
<td>54°09'N</td>
<td>2′30&quot;</td>
<td>-3.8/-4.5 °C</td>
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</table>
During the Younger Dryas, the temperatures inferred from the Ech chironomids were similar to other previous summer temperature reconstructions from sites located on the southern slope of the Alps (Table 3a). The temperature inferences for the early Holocene (17 °C) lie also in the range of these previous reconstructions (Table 3a). For both the Interstadial/Younger Dryas and Younger Dryas/Holocene transitions, changes in inferred-temperatures (i.e. 1.5–2 °C and 2.5 °C, respectively) were close to those reconstructed from chironomids at Lago di Lavarone and Lago Piccolo di Avigliana (Table 3a). The magnitudes of these temperature changes were smaller than those recorded at higher latitude in Europe (e.g. in northwest England, Lang et al., 2010, Western Norway, Brooks and Birks, 2000). The Ech temperature reconstruction agrees with the other available chironomid-based summer temperature reconstructions to indicate a continuous south to north gradient in the amplitude of the Younger Dryas cooling (Table 3b). These findings are also in close agreement with climate simulations and reconstructions presented in Renssen and Isarin (2001) which showed a clear N–S gradient in the magnitude of temperature changes associated to the “11.7 ka transition”. This points to the absence of local interference into large-scale climate forcing factors (linking insolation, ocean and atmosphere), contrary to the Oldest Dryas where sea ice apparently strongly influenced local climate along Atlantic coast.

6. Conclusion

This paper presents the first study of Lateglacial chironomids in the Pyrenees, a region where quantitative climate reconstructions are still rare. Temperature change was found to be the main controlling factor for the Ech chironomid assemblage during the Lateglacial even if assemblage changes may also have been mediated by other local conditions. This implies that temperature reconstruction based on changes in chironomid assemblages was justified. The feasibility of quantitative reconstruction using a merged Swiss–Norwegian training set was supported by the good error and prediction statistics of the model and the similarity in ecological preferences of the main taxa between Norway, the Alps and the Pyrenees.

The model reconstructed July air temperatures between 10 and 13 °C during the Oldest Dryas, a two-step increase in temperatures to 16 and 17.5 °C during the first part of the Interstadial, followed by a decrease to 16.5 °C by the end of the Interstadial. Inferred temperatures decreased to 15–15.5 °C during the Younger Dryas, and reached ca. 17 °C at the beginning of the Holocene. Superimposed on the changes in temperature corresponding to the major lateglacial events, climate variability at a centennial-scale is further suggested by the Ech record.

During the Interstadial, the Ech record features three centennial-scale coolings of ca. 1.5 °C amplitude. The older (ca. 13,900 cal yr BP) may correlate with the GI-1d event in the Greenland event stratigraphy (Lowe et al., 2008), equivalent to the Aegelsee Oscillation (Lotter et al., 1992). The younger short-lived cold oscillation (at ca. 13,100 cal yr BP) may be related to the GI-1b event (Lowe et al., 2008), equivalent to the Gerzensee Oscillation (Lotter et al., 1992). The third cold reversal, ca. 13,600 cal yr BP, may correlate with a short anomaly in δ18O in the Greenland ice-cores registered at a similar age within GI-1c (Fig. 7). Similarly, a rapid cooling event placed between GI-1b and GI-1d was also evidenced from chironomids in northwest England (LG-3 in Lang et al., 2010) but with a weaker reconstructed decrease in July air temperature (ca. 0.6–0.8 °C). These three cooling events during the Interstadial are also apparent in the chironomid-inferred temperature record from Whittig Bog in southeast Scotland (Brooks and Birks, 2001).

5.3.3. The Younger Dryas and the Early Holocene

During the Younger Dryas, the temperatures inferred from the Ech chironomids may correlate with a third cold reversal of ca. 13,600 cal yr BP, i.e. between the two previous centennial-scale cold events GI-1b and GI-1d (Fig. 7). It may be correlated with a short anomaly in δ18O in the Greenland ice-cores registered at a similar age within GI-1c (Fig. 7). Similarly, a rapid cooling event placed between GI-1b and GI-1d was also evidenced from chironomids in northwest England (LG-3 in Lang et al., 2010) but with a weaker reconstructed decrease in July air temperature (ca. 0.6–0.8 °C). The decrease of 1.5 °C in July air temperature between 0.8 and 2.1 °C for these two events (reviewed in Ilyashuk et al., 2009; Lang et al., 2010) may correlate with the GI-1d event in the Greenland event stratigraphy (Lowe et al., 2008), equivalent to the Aegelsee Oscillation (Lotter et al., 1992). The decrease of 1.5 °C in July air temperature inferred from the Ech chironomid record is in the upper range of previous published records and extends the available data-set describing climate change at the end of the last glaciation (EUCLIM) headed by O. Heiri and A.F. Lotter, the Nederlandse Organisatie voor Wetenschappelijk Onderzoek, Aard- en Levenswetenschappen (NWO-AWL grant 818.01.001), the Communauté de Travail des Pyrénées and the project ANPYR (FEDER/National Park of the Western Pyrenees) headed by D. Galop.

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