



Holocene climate variability, vegetation dynamics and fire regime in the central Pyrenees: the Basa de la Mora sequence (NE Spain)



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ABSTRACT

High resolution multiproxy data (pollen, sedimentology, geochemistry, chironomids and charcoal) from the Basa de la Mora (BSM) lake sequence (42° 32' N, 0° 19' E, 1914 m a.s.l.) show marked climate variability in the central southern Pyrenees throughout the Holocene. A robust age model based on 15 AMS radiocarbon dates underpins the first precise reconstruction of rapid climate changes during the Holocene from this area. During the Early Holocene, increased winter snowpack and high snowmelt during summer, as a consequence of high seasonality, led to higher lake levels, a chironomid community dominated by non-lacustrine taxa (*Orthocladinae*) related to higher inlet streams, and a forested landscape with intense run-off processes in the watershed. From 9.8 to 8.1 cal ka BP, climate instability is inferred from rapid and intense forest shifts and high fluctuation in surface run-off. Shifts among conifers and mesophytes reveal at least four short-lived dry events at 9.7, 9.3, 8.8 and 8.3 cal ka BP. Between 8.1 and 5.7 cal ka BP a stable climate with higher precipitation favoured highest lake levels and forest expansion, with spread of mesophytes, withdrawal of conifers and intensification of fires, coinciding with the Holocene Climate Optimum. At 5.7 cal ka BP a major change leading to drier conditions contributed to a regional decline in mesophytes, expansion of pines and junipers, and a significant lake level drop. Despite drier conditions, fire activity dropped as consequence of biomass reduction. Two arid intervals occurred between 2.9 and 2.4 cal ka BP and at 1.2–0.7 cal ka BP (800–1300 AD). The latter coincides with the Medieval Climate Anomaly and is one of the most arid phases of the Holocene in BSM sequence. Anthropogenic disturbances were small until 700 AD, when human pressure over landscape intensified, with *Olea* cultivation in the lowlands and significant deforestation in highlands. Colder and unfavourable weather conditions during the second part of the Little Ice Age caused a temporary cease of high-land management. The most intense anthropogenic disturbances occurred during the second half of 19th century. Last decades are characterized by recovery of the vegetation cover as a result of land abandonment, and lowered lake levels, probably due to higher temperatures.

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1. Introduction

Long-term climate evolution during the Holocene has been strongly modulated by orbitally-forced insolation trends which determine heat distribution throughout the planet. In the northern

Hemisphere, summer insolation sets limits on the position and strength of the Inter Tropical Convergence Zone (ITCZ), which controls the position of the north-hemisphere cell atmospheric system (Wanner and Brönnimann, 2012). In particular, the location of the Azores High and the Iceland Low pressure centres determines the latitudinal position and intensity of the North Atlantic westerlies and the storm tracks, which largely govern rainfall distribution in the Western Mediterranean area (Greatbatch, 2000; Marshall et al., 2002). During the Early Holocene, the maximum

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summer insolation in the Northern Hemisphere led to a rapid northward displacement in the ITCZ and its associated rain belt (Fleitmann et al., 2007). This northern position of the ITCZ was responsible for bringing moisture to the current world-largest desert in North Africa (Sahara and Sahel) (deMenocal et al., 2000). As the summer insolation decreased the ITCZ displaced southward, the monsoon system weakened and in south-western Europe the climate followed a general trend to an increasingly aridity since the Mid Holocene that led to decreased lake levels (Magny et al., 2007, 2011; Valero-Garcés and Moreno, 2011) and major shifts in the vegetation composition (Fletcher and Zielhofer, 2013; Roberts et al., 2011).

However, beyond this general climate trend, many recent studies have documented the existence of rapid climate variability during the Holocene (Bond, 1997, 2001; Mayewski, 2004). Although the nature and mechanisms of these abrupt climate changes still remain unclear, weakening in the thermohaline circulation as consequence of meltwater inputs in the North Atlantic has been recognized as one of the most important triggers (Renssen et al., 2007; Wanner et al., 2008). Furthermore, fluctuations in solar activity have also been responsible for climate shifts (Wanner et al., 2011). These short-living episodes of climate variability had a large impact over most of Europe, as it has been recorded in many continental palaeoclimate archives as lacustrine sediments (Magny et al., 2007), glacial deposits (Davis et al., 2009), and pollen records (Bordon et al., 2009; Magyari et al., 2012).

Holocene climate reconstructions for the North Atlantic region involve mainly changes in temperature (Brooks and Birks, 2001). However, in the Mediterranean area Holocene variability is mostly related to changes in water availability as it is documented in vegetation distribution (Jalut et al., 2009; Sadori et al., 2011), lake levels (Magny et al., 2011) and stalagmite growth (Fleitmann et al., 2007; Spötl et al., 2010).

The Iberian Peninsula climate integrates subtropical, Mediterranean and Atlantic influences due to its geographical location between the Mediterranean Sea and the Atlantic Ocean (Lionello et al., 2006). Moreover, the Iberian Peninsula has proven to be particularly sensitive to short-term climate shifts during the Holocene (Moreno et al., 2012a). Lakes experienced noteworthy variations in response to precipitation and evaporation shifts during the Holocene (Valero-Garcés et al., 2000; González-Sampériz et al., 2008; Martín-Puertas et al., 2008; Morellón et al., 2009). Changes in sea surface temperatures (Cacho et al., 2001) and deepwater formation (Frigola et al., 2007) in the Western Mediterranean show a fast response to changes in the North Atlantic. Other Iberian continental records highlight a large Holocene variability. For example, the isotope record in the Kaite Cave stalagmite (Domínguez-Villar et al., 2008) reflects variations in the amount of precipitation related to North Atlantic dynamics and fluctuations in palaeoflood activity of Tagus River, in Central Spain have been related to changes in prevailing atmospheric circulation patterns (Benito et al., 2003). Although vegetation is a very good indicator of past climate variability, there are only a few high-resolution pollen studies from the Iberian Peninsula (e.g. Jiménez-Moreno and Anderson, 2012; Fletcher and Zielhofer, 2013), documenting the fast response of vegetation to abrupt climate changes (decadal- to centennial-scale) during the Holocene.

A recent study has proved the high-sensitivity of middle-latitude high mountain ranges in general, and the Pyrenees in particular, to current global warming, documenting a speeding up of replacement of cold-adapted plants by thermophilic species (Gottfried et al., 2012). Past climate changes during the Holocene should have also affected the flora and landscape of the Pyrenees. Furthermore, the southern slopes of the Pyrenees are not affected by Foehn winds, and the present climate is rather complex,

influenced by a progressive west-to-east decrease in precipitation, due to weakening of the Atlantic humid fronts inland. Thus, the southern Pyrenees experience both Atlantic and Mediterranean climate regimes within a relatively short distance of less than 450 km. The Pyrenean vegetation reflects these climate conditions, varying from humid-Atlantic forests, dominated by oak and beech, in the west, to Mediterranean forests, dominated by pine and drought-resistant taxa, in the central and eastern regions. Due to these particular geographical features the central Pyrenees play a key role in providing information about past E–W shifts of the boundary between both regimes as a result of shifts in the atmospheric components and, particularly, shifts in the westerlies strength.

In Western Europe, human disturbances in the landscape can be traced back to the Neolithic period and the climate signal is often masked by anthropogenic activities during the most recent times (Oldfield, 2005; Carrión et al., 2007). Discriminating anthropogenic from natural forcings in landscape evolution has been subject of much debate during recent years. High-altitude sites are more useful than low-altitude sites for detecting climate signals, since more inhospitable climate conditions limit intense human landscape intervention.

Here we present a paleo-environmental reconstruction of climate, vegetation and fire dynamics from a lacustrine sequence located in the central part of the southern Pyrenees: the Basa de la Mora sequence. This record stands out as one of the best climate archives to tackle questions concerning: i) how the Atlantic and Mediterranean regimes have progressed along the Holocene, ii) identification of rapid episodes of climate change, and iii) elucidation of high mountain land-use system during recent times.

2. Study area

2.1. Geological and geomorphological setting

Lake Basa de la Mora (BSM) (42° 32' N, 0° 19' E, 1914 m a.s.l.) is a small, shallow glacial lake located on the north-facing slope of the Cotiella Peak (2912 m a.s.l.), the highest summit of the Cotiella Massif in the central southern Pyrenees (Fig. 1a). The Cotiella Massif belongs to the homonymous nappe, located in the western part of the South Pyrenean Central Unit (Seguret, 1972).

The landscape surrounding the lake results from intense karstic and glacial activity. Lake Basa de la Mora occupies a glacial over-deepened basin enclosed by a frontal moraine (Belmonte, 2004) and surrounded by steep limestone walls. The catchment consists of Mesozoic limestones and sandy limestones affected by several thrust sheets (reverse faults). Triassic marl and evaporite formations crop out at the base of the thrust sheets, providing a hydrological seal for the lake and favouring localized surface drainage into the lake along some creeks. Triassic ophiolite formations in the watershed are the source of highly characteristic sediments (hematite and other Fe- mineral with high magnetic susceptibility) within the lake deposits.

The Basa de la Mora basin belongs to the watershed of the Cinca River, one of the main tributaries of the Ebro River. The lake has smooth margins, a relatively small watershed (209 ha) and a total lake surface of ca 3 ha. It is characterized by large seasonal water-level fluctuations: the maximum depth varies from ca 2.5–4.5 m seasonally. The lake is fed by precipitation, surface runoff, ephemeral creeks and several small springs located on the southern margin. Water losses take place through a surface outlet to the north and evaporation. The substrate, made up of non-permeable Triassic material, greatly restricts groundwater losses.

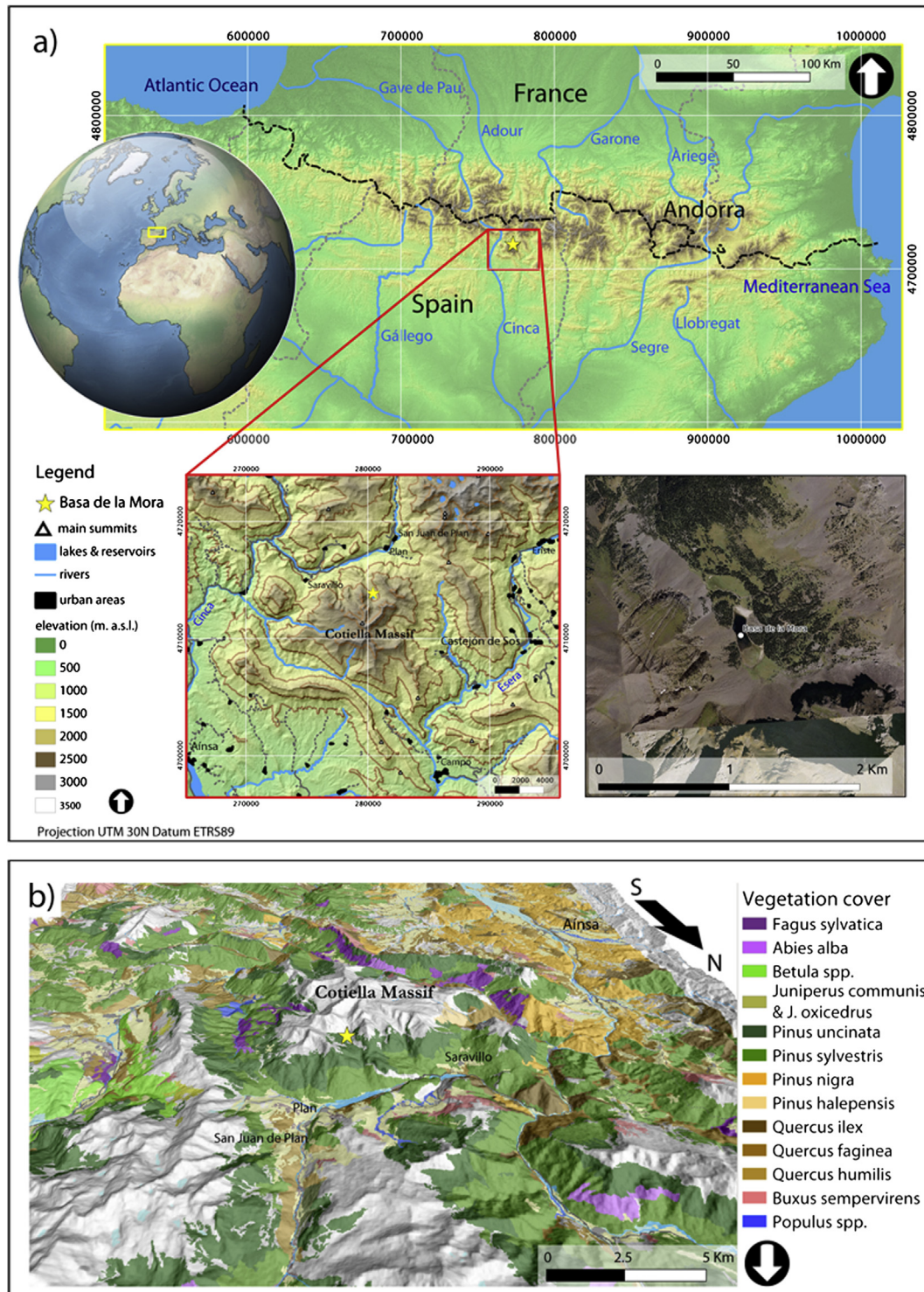


Fig. 1. a) Location map and orthophoto of Basa de la Mora Lake in central Pyrenees (Spain). b) 3D regional vegetation map. In order to better discern the topography, the North is plotted at the bottom of the figure.

2.2. Climate and vegetation

The Pyrenees is a mountain range in south-western Europe that extends from the Atlantic Ocean in the west to the Mediterranean Sea in the east, leading to a diverse climate and plant community along a W–E transect. The precipitation in the Pyrenees results from two different mechanisms: precipitation in the east is linked

to cold fronts, while precipitation in the west comes from Atlantic frontal systems (Millán et al., 2005). The Atlantic influence extends as far as the Ordesa Valley (García-Ruiz et al., 2001), ca 150 km from the Atlantic coast and 22 km west of the BSM. Both systems are directly related to the North Atlantic Oscillation (NAO) that principally determines precipitation in western Europe (Trigo et al., 2002).

The climate of the study area is sub-Mediterranean with continental features. Rainfall (annual average = 1360 mm) peaks during spring and autumn, following the Mediterranean pattern (García-Ruiz et al., 1985). However, summers are not as dry as is typical of the Mediterranean because of frontal and convective precipitation which affects the mountainous areas in July and August. Mean air temperatures range from 0.5 to 15 °C between the coldest (January) and warmest (July) months, respectively.

The vegetation cover shows a characteristic contrast between south and north facing slopes: the southern slopes are characterized by Mediterranean-type components with sclerophyllous shrubland and evergreen *Quercus* communities, while the northern slopes have mixed conifer/deciduous taxa forests, including *Pinus sylvestris*, *Pinus uncinata*, *Abies alba*, *Betula alba*, *Corylus avellana*, *Fagus sylvatica*, *Quercus faginea* and *Quercus petraea* (Fig. 1b).

The elevational gradient between the valley bottoms and the Cotiella Peak, from 550 to 2900 m a.s.l., gives rise to an altitudinal distribution of vegetation, typical of mountain environments. Lowlands are occupied by crops and valley bottoms by riparian corridors (*Fraxinus excelsior*, *Populus* spp., and *Salix* spp.). Forests occur from the base of the foothills up to ~2000 m a.s.l. Below 1700 m a.s.l., the dominant species are determined by moisture availability and temperature range, mostly controlled by the slope orientation. From 1700 to 2000 m a.s.l. the forest is mainly composed of *P. uncinata* mixed with *Juniperus communis* shrubland and *Rhododendron ferrugineum* at the treeline. Above 2000 m a.s.l., steep rock formations and harsh climate prevent forest development, leading to a scrub-dominated landscape formed by dwarf junipers (*J. communis* sbsp. *nana*), and alpine grassland (*Nardus stricta*, *Festuca eskiae*, *Caricion davallianae* and *Cynosurus cristatus*). Lake Basa de la Mora (BSM) is located in the subalpine belt, near the treeline, so the vegetation surrounding the lake is alpine grassland, *P. uncinata* forest and *J. communis*–*R. ferrugineum* shrublands.

3. Methodology

The composite sequence of Basa de la Mora (BSM08-1A-1U) is based on two parallel cores retrieved from the deepest part of the lake in summer 2008. The longest core was taken with an Uwitec coring system and platform from the Pyrenean Institute of Ecology (IPE-CSIC). Two gravity cores were taken to recover the uppermost part of the sequence and the sediment/water interface. One of the short cores (BSM08-1A-1G) was sub-sampled every 1 cm in the field for ^{210}Pb and ^{137}Cs analyses and the other core (BSM08-1B-1G) was used to complete the upper part of the sequence. The cores were correlated applying sedimentological and geochemical criteria. The total length of the composite sequence is 12.10 m. An additional littoral core (BSM-2A-1U) was taken in order to compare lacustrine depositional environments.

The cores were split lengthwise into two halves, imaged with a DMT Core Scanner and analysed with a Geotek Multi-Sensor Core Logger (MSCL) at 5 mm intervals to characterize the sediment physical properties at the Limnological Research Center at the University of Minnesota (USA). Elemental geochemical composition was analysed using the Itrax XRF Core Scanner at the Large Lakes Observatory (LLO) at the University of Minnesota (USA) at 0.5 cm resolution using 30-s count times, 30 kV X-ray voltage, and an X-ray current of 20 mA. These measurements provide estimates of relative element concentrations. The cores were sub-sampled at 2 cm resolution for Total Organic Carbon (TOC) and Total Inorganic Carbon (TIC) and analysed with a LECO144DR elemental analyser at the IPE-CSIC laboratory of Zaragoza (Spain). Sedimentary facies were defined by macroscopic characteristics including colour, grain-size,

sedimentary structures, fossil content and by microscopic smear slide observations (Schnurrenberger et al., 2003) (Fig. 3). The sedimentological descriptions are supported by Scanning Electronic Microscopic (SEM) observations of selected samples made at the University of Zaragoza (Spain). Up to 11 samples representing the main facies were analysed for grain size distributions using a Malvern Laser Sizer, Mastersizer, 2000, after removing the organics by H_2O_2 and using a dispersant agent to disaggregate the samples. Additionally, 36 samples were analysed for their mineralogical content by X-Ray Diffraction using an automatic Siemens D-500 X-ray diffractometer: Cu ka, 40 kV, 30 mA and graphite monochromator. Identification and quantification of the different mineralogical species present in the crystalline fraction were carried out following a standard procedure (Chung, 1974). Sedimentary facies and physical properties (density, magnetic susceptibility) were also obtained for the littoral core (BSM-2A-1U).

Samples for pollen analyses were obtained every 5 cm on both the BSM08-1A-1U and BSM08-1B-1G cores. This record covers the whole sedimentary record, except the base of the sequence (1209–1165.5 cm depth) which was sampled at higher resolution (1 cm) since the sedimentation rate was extremely low (see below). Sediment samples were prepared following the standard protocol described by Faegri and Iversen (1964) or Moore and Webb (1978), with some modifications (Dupré, 1988) including HCl, and KOH and HF digestion, mineral-organic particles separation with Thoullet solution (2.0 g/cm³ density) and sieving with 212 and 10 µm mesh. *Lycopodium* spores in a known concentration were added in order to calculate the pollen concentration in the sediment and to test the laboratory procedures (Stockmarr, 1971). Pollen was identified using an optical microscope, with help of the reference collection of the IPE-CSIC and identification keys (Moore et al., 1991; Reille and Lowe, 1995). Counts were made to obtain a pollen sum, excluding aquatics and exotics, of at least 300 grains from a minimum of 20 taxa. The results have been plotted using PSIMPOLL 4.27 (Bennett, 2009).

Correlation analyses were made on smoothed data, after testing for normality (Shapiro–Wilk), using Pearson or Spearman correlation tests. Analysis have been performed by the R software package (Venables et al., 2008). Pairwise comparison was performed between MS and geochemical parameters, to help in the facies description, and then between MS (as a high-resolution sedimentological proxy) and the pollen data, to assess possible links between sedimentary changes and vegetation.

Sedimentary micro-charcoal particles were identified on pollen slides by optical microscopy. Only charcoal particles over 10 µm were counted and these were easily identified as black, angular and opaque particles (Clark, 1988). Charcoal influx (mm²/cm³) was estimated after Tinner and Hu (2003). No *Lycopodium* spores were found in some of the slides, so charcoal influx values were obtained by linear interpolation between the adjacent samples.

Chironomid samples were collected every 20 cm along the entire core, except at the top of the sequence (2.5–50 cm depth) where the sample interval was increased to 5 cm. The samples were processed following the standard procedure (Hofmann, 1986): 10% KOH digestion at 70° and 300 rpm for 20 min, followed by sediment sieving (90 µm). *Chironomidae* larvae head capsules were examined under stereo microscope using a Bolgorov tray, picked out manually and dehydrated in 96% ethanol, before being mounted ventral side upwards in Euparal® as permanent slides. Taxonomic identification was carried out using an optical microscope (Olympus CX41) at 40× magnification and Cell B Imaging Software for Life Science Microscopy (Olympus). The larval head capsules were identified to the lowest taxonomic level possible using several specialized guides (Wiederholm, 1983; Rieradevall and Brooks, 2001; Brooks et al., 2007).

The chronology of the sequence is based on 15 calibrated AMS radiocarbon dates from the long core BSM08-1A-1U and ^{137}Cs and ^{210}Pb dating from the short core BSM08-1B-1G (Fig. 2). Most of radiocarbon dates are based on terrestrial macrofossils and charcoal (Table 1). Bulk sediment and pollen concentrates were dated in the lowermost part of the sequence because of the paucity of organic remains. Dates have been calibrated using CALIB 6.0 software and the INTCAL09 curve (Reimer et al., 2009). The 2σ probability distribution interval was chosen. The age model was constructed by linear interpolation between the median ages of the probability distribution of adjacent calibrated dates. The $^{210}\text{Pb}_{\text{ex}}$ and ^{137}Cs activity in the upper samples was measured by gamma-ray spectrometry, using a high-resolution low-energy coaxial HPGe detector coupled to an amplifier. The chronology based on $^{210}\text{Pb}_{\text{ex}}$ was estimated by applying the constant rate of supply (CRS) model by Appleby (2001). The resulting age model provides a robust chronological framework for the high resolution paleo-environmental reconstruction presented in this work.

4. Results

4.1. Chronology

According to the age–depth model, the BSM sequence spans the last ca 16 cal ka BP (Table 1, Fig. 2). The two lowermost dates (at 11.98 and 12.06 m depth) are the only ones not based on terrestrial macrofossils (Table 1). When these two dates are included in the age model ($12,628 \pm 100$ and $15,828 \pm 600$ cal yr BP), they result in a change from consistently high sedimentation rates (1.2 mm/yr) between 0 and 11.67 m depth to extremely low rates (0.064 mm/yr) at the base (11.67–12.09 m depth). Given that these dates point out to the Lateglacial period, we attempted to characterize this zone by increasing the pollen sampling. However, the pollen record did not show changes indicative of the Last Glacial–interglacial transition (LGIT, see Section 4.3, zone BSM-0). Since there is no sedimentary evidence for a depositional hiatus, and no major change in the vegetation composition has been recorded, these two dates were not used in the final age model. It is possible that a reservoir effect is responsible for these samples being too old. The age model excluding the two basal dates indicates that the 11.67 m long record spans the last ca 9.8 cal yr BP (Fig. 2). The final age–depth model is based on 13 calibrated AMS radiocarbon dates, 11 on macrofossils and two on charcoal. The short core, that includes the most recent period, has been dated by ^{210}Pb and ^{137}Cs activities. Two well-defined ^{137}Cs peaks are recorded at the uppermost part of the sequence providing markers for the 1954–1959 and the 1963

maximum atmospheric nuclear bomb testing. The chronology based on $^{210}\text{Pb}_{\text{ex}}$ compares fairly well with the ^{137}Cs peaks (Fig. 2).

4.2. Sedimentary facies, geochemistry and lithological units

Six sedimentary facies were identified based on visual description, microscopic observations, grain-size data and mineralogical and geochemical composition (Table 2). The sediments consist of either: i) carbonate-poor (<2% TIC), with lower TOC and high MS, organized in laminated or banded intervals, or ii) carbonate-rich (2–7% TIC) with variable, but higher organic matter content (1–3%) and low magnetic susceptibility, arranged in massive to banded deposits. The grain-size data indicates finer (mode at 6–7 μm) and better-sorted sediments in the silicate-rich, carbonate-poor facies, and coarser and more poorly sorted material in the carbonate-rich sediments.

The first group of sediments (Facies 1, 2 and 3) are banded to laminated silicate and carbonate fine silts dominated by clay minerals (20–30%) and quartz (5–15%) with minor amounts of calcite (<25%) and with presence of hematite, pyrite and clinochlorite. Facies 3 has the highest MS, and relatively high carbonate content. Facies 1 and 2 are more silicate-rich, but Facies 2 is finer, with lower MS, better-defined lamination and higher TOC content than Facies 1. The second group (Facies 4, 5 and 6) is dominated by massive carbonates (ca 6% TIC; 60–80% calcite). Facies 5 and 6 have mottled textures and abundant gastropods, indicating littoral deposition. These facies dominate the littoral core (BSM-2A) almost entirely. Facies 4 has a higher TOC content (up to 3%) dominated by macrophyte and terrestrial remains. Facies 5 contains authigenic crystals of carbonate and gypsum, partially dissolved, pointing to deposition in ephemeral lake conditions with rapid fluctuations of lake level and salinity. Diatoms (pennate, benthic) only occur in the carbonate-rich Facies 5. Facies 6 has a slightly banded texture and lower TOC content than the other carbonate facies.

The BSM sequence has been divided into three main sedimentary units according to sedimentary facies, MS, TIC and TOC percentages and the mineralogical and geochemical composition (XRF) (Fig. 3).

- i) Unit 3 (1168–491 cm depth; 9800–5700 cal yr BP) corresponds to the lowermost part of the sequence and it is characterized by banded carbonate-poor sediments with high values of MS and relatively low TOC percentages (Facies 1, 2 and 3). TIC percentages and Ca, Sr and S values are low throughout Unit 3 while Si, K, Ti values (and particularly Fe and Mn) are high. The lowermost Sub-unit 3b (1168–690 cm

Table 1
AMS radiocarbon dates from core BSM08-1A-1U. Rejected dates are shown in italics.

(b) Lab code	Depth (cm)	Sample type	^{14}C age (yr BP)	Calibrated age, 2σ (yr cal BP)	Median probability (yr cal BP)
Poz-29744	60	Terrestrial macrorest	385 ± 30	426–507	456
Poz-35854	172	Terrestrial macrorest	1335 ± 30	1231–1304	1276
Poz-29745	230	Terrestrial macrorest	2100 ± 30	1995–2146	2072
Poz-35853	269	Terrestrial macrorest	2615 ± 30	2718–2777	2749
Poz-35852	337	Terrestrial macrorest	3200 ± 30	3368–3469	3419
Poz-35804	422	Terrestrial macrorest	3815 ± 35	4089–4299	4206
Poz-29743	502	Terrestrial macrorest	5185 ± 35	5893–6002	5942
Poz-35803	562	Terrestrial macrorest	5840 ± 40	6533–6745	6657
Poz-35802	677	Terrestrial macrorest	6450 ± 40	7288–7430	7367
Poz-29746	795	Charcoal	7330 ± 50	8014–8214	8125
Poz-35801	943	Terrestrial macrorest	7930 ± 50	8628–8983	8778
Poz-29747	1011	Charcoal	7950 ± 50	8640–8990	8817
Poz-29779	1167	Terrestrial macrorest	8780 ± 50	9581–9941	9798
<i>Poz-35856</i>	<i>1198</i>	<i>Bulk sediment</i>	<i>10,710 \pm 60</i>	<i>12,547–12,743</i>	<i>12,627</i>
152,235	1206	Pollen concentrates	$13,080 \pm 100$	15,181–16,476	15,828

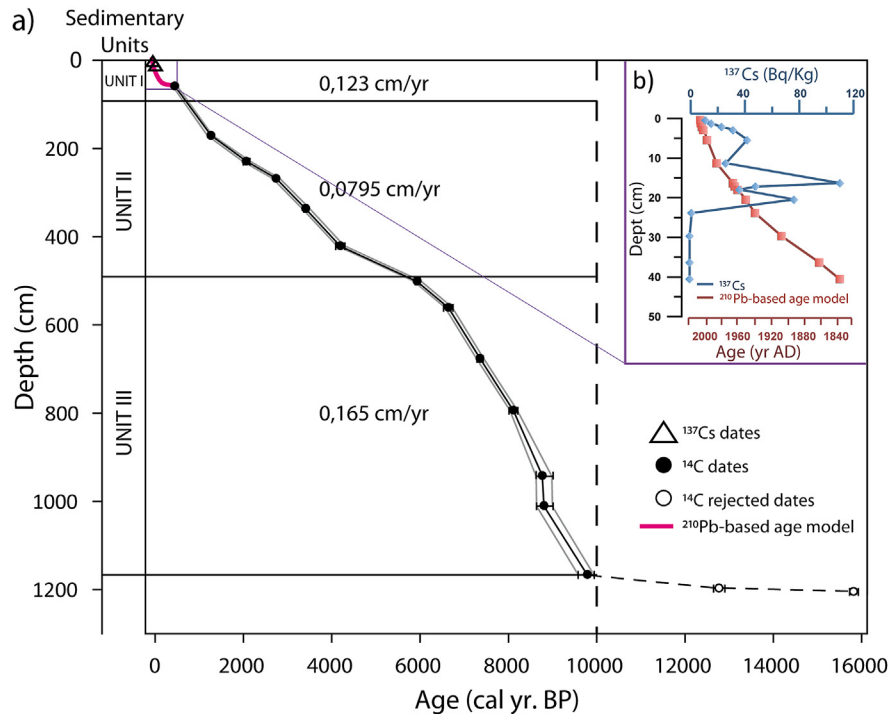


Fig. 2. a) Age–depth model for the composite sequence of Basa de la Mora Lake based on 15 AMS ^{14}C dates and ^{210}Pb and ^{137}Cs activity at top. b) ^{210}Pb -based age model and ^{137}Cs profile obtained for the top 50 cm.

depth, 9800–7450 cal yr BP) is composed of laminated Facies 1 and a thin interval of Facies 3. Magnetic Susceptibility (MS) reach the highest values of the sequence and are positively correlated with Mn (Table 3a). The high MS values are related to the presence of paramagnetic minerals eroded from ophite outcrops. Values of Ca and TIC are relatively low, but also display a strong positive correlation with MS. TOC percentages are the lowest in the sequence while TOC/N ratios are the highest. Sub-unit 3a (690–491 cm depth, 7450–5700 cal yr BP) is composed of Facies 2 and has finer lamination, lower MS and higher TIC and TOC values. Sub-

unit 3a MS values are still high but decrease progressively. MS is significantly positively correlated with Mn and Fe (Table 3a). Ca values are very low and not significantly correlated with MS. TOC percentages increase, showing a significant negative correlation with MS, while TOC/N ratios decreases.

- ii) Unit 2 (491–93 cm depth; 5700–680 cal yr BP) is made up of carbonate-rich Facies 5 and 6 with intercalations of organic-rich Facies 4. Thus, Unit 2, although highly variable, is characterized by the lowest values of MS and the highest content in TIC of the whole sequence. The high values of TIC in Unit 2

Table 2
Facies description and interpreted depositional environment of BSM sequence.

Facies	Facies description
<i>Clastic, laminated facies</i>	
1	Grey banded to laminated quartz and carbonate silts. Mostly composed by clay minerals (45%), calcite (17%) and, quartz (7%) and low organic matter (<1%). High MS (100 SI). Laminated intervals are composed of up to 1 cm thick couplets of (1) black, carbonate silty-sands with high quartz content, abundant hematites, chlorite and mafic minerals and occasional terrestrial and macrophyte remains and (2) grey carbonate silts with lower silicate minerals content and rare organic matter.
2	Dark grey laminated carbonate silts. Mineralogical composition similar to Facies 1, but better laminated higher organic content (1–2%) and lower MS (average 40 SI). Couplets composed of mm- thick laminae of (1) black, carbonate silty-sands with abundant terrestrial and macrophyte remains and (2) brown carbonate silts with less siliciclastic minerals and lower organic matter.
3	Light grey banded carbonate silts. Dominant carbonate content (TIC, X %; calcite, 40%); quartz (6%) and significant amounts of hematites, pyrite, clinoclhorite, other mafic. Low organic matter (1%). Very high MS (>150 SI).
Interpretation	Clastic dominated deposition in distal, deeper setting. Laminated facies reflect flooding episodes reaching the centre of the lake. More abundant carbonate (Facies 3) or organic matter (Facies 2) reflects changes in watershed and littoral environments.
<i>Carbonate and organic-rich facies</i>	
4	Black, massive, carbonate silts. Composition is dominated by calcite (45%), quartz (10%), clay minerals (10%) and organic matter (>2%) of terrestrial and macrophyte origin. Abundant pyrite and rare hematites. Low MS (25 SI). Occasional presence of pennate diatoms.
5	Light grey, massive, carbonate silts. Composition is dominated by calcite (70%), with relatively low quartz and clay minerals (7%) and organic matter (<2%); occasional pyrite and rare hematites. Low MS (25 SI). Organic matter is terrestrial, macrophyte and lacustrine origin. Mottling is common. Abundant gastropods and presence of pennate diatoms.
6	Light brown, banded, carbonate silts. Composition is dominated by calcite (30%), clay minerals (15%) and relatively low quartz (9%) and organic matter (<2%) mostly terrestrial and macrophyte remains.
Interpretation	Carbonate dominated deposition in littoral environments with higher carbonate and organic productivity (Facies 5) deeper, with more frequent anoxic conditions (Facies 4) and transitional (Facies 6).

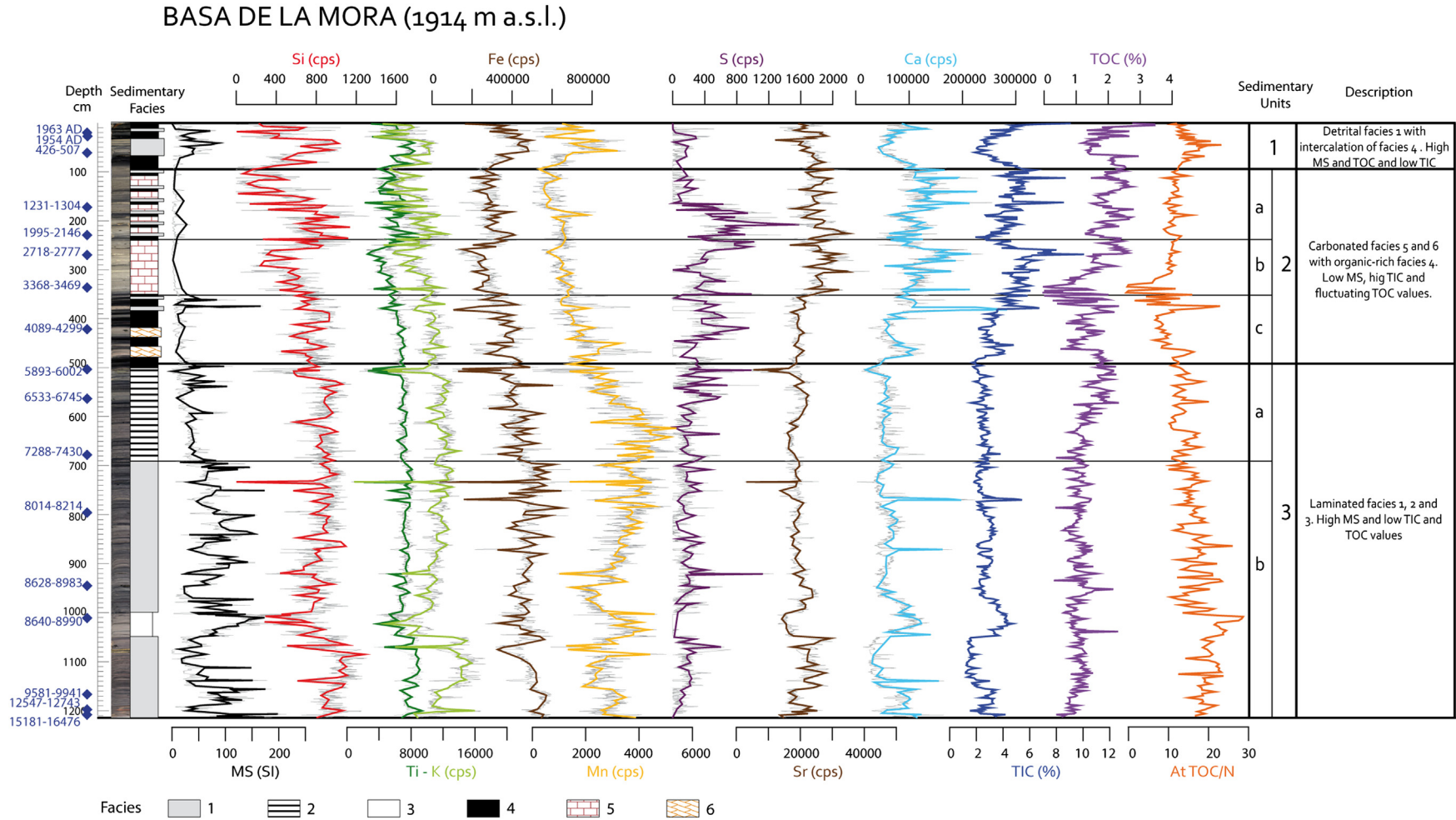


Fig. 3. Main sedimentological features, geochemical and physical properties of the Basa de la Mora sequence plotted in depth, indicating the location and results of radiocarbon dates, Facies 1 to 6 (see Table 2), identification and description of the Sedimentary units.

Table 3a

Correlation values between Magnetic Susceptibility and other geochemical parameters in the different sedimentary units.

	Unit 1 (0–93 cm)		Unit 2 (93–491 cm)		Sub-unit 3a (491–690 cm)		Sub-unit 3b (690–1168)	
	MS		MS		MS		MS	
	<i>r</i>	<i>p</i>	<i>r</i>	<i>p</i>	<i>r</i>	<i>p</i>	<i>r</i>	<i>p</i>
Si	0.726	<0.001	0.546	<0.001	0.146	0.148	–0.346	<0.001
Ti	0.699	<0.001	0.688	<0.001	0.280	0.005	–0.388	<0.001
Mn	0.688	<0.001	0.545	<0.001	0.543	<0.001	0.451	<0.001
Fe	0.806	<0.001	0.582	<0.001	0.643	<0.001	0.162	0.013
Ca	–0.660	<0.001	–0.564	<0.001	0.179	0.075	0.671	<0.001
TIC	–0.700	<0.001	–0.591	<0.000	0.404	<0.001	0.689	<0.001
TOC	–0.746	<0.001	–0.409	<0.001	–0.609	<0.001	–0.494	<0.001

(up to 8%) are related to precipitation of authigenic carbonates. Sr and S elements increase considerably in this unit. TOC percentages also vary greatly during this period but, in general, they are relatively high and increase upwards. Relatively low TOC/TN values (<12) indicate the dominance of lacustrine organic matter (Meyers, 2003). Si, Ti, Fe and Mn show parallel trends to MS (Table 3a). Unit 2 can be subdivided into three sub-units, following the facies association. Thus, BSM 2c (491–350 cm depth; 5700–3540 cal yr BP) is constituted by the alternation of cm-thick intervals of Facies 4 and 6 and displays an upward TIC increase (up to 8%). TOC percentages are highly variable but generally low (1–2%). BSM 2b (350–240 cm depth; 3540–2200 cal yr BP) represents a 1 m-thick interval of Facies 5 with the highest TIC, Ca and calcite values and the lowest TOC and MS of the sequence (Fig. 3). Higher Sr values occur as a result of more abundant biogenic aragonite. Finally, BSM 2a (240–93 cm depth; 2200–700 cal yr BP) comprises rhythmic sequences of about 20 cm-thick composed of thin layers of Facies 1->, Facies 4-> Facies 5 (detrital- organic-carbonate).

- iii) Unit 1 (93–0 cm depth; 698 cal yr BP–2007 AD) comprises carbonate-poor Facies 1 and organic-rich Facies 4. As a consequence, all geochemical indicators show high variability. Facies 1 lamination is less well defined than in Unit 3. MS values increase again and show strong positive correlation with Si, Ti, Mn and Fe, while the correlation with Ca and TIC and TOC is strongly negative (Table 3a). TOC/N ratios increase at the base of the unit and decrease towards the top: TOC percentages show the opposite pattern.

4.3. Pollen and charcoal data

The pollen record can be divided into six zones (BSM-0 to BSM-V: Fig. 4). In BSM-I to BSM-V (9.8 ka cal BP–present), the 5 cm-resolution pollen analyses provide a temporal resolution of 22–150 years per sample. Statistical results for pairwise comparison between vegetation and geochemical parameters are shown in Table 3b. The maximum number of charcoal particles counted was 3098, with a mean of 307 and a SD of 453. The patterns of charcoal influx are consistent with the pollen zones.

BSM-0 (1209–1167.5 cm depth; before 9800 cal yr BP)

This zone is characterized by scarce representation of the herbaceous component (NAP) and particularly the steppe taxa group (*Artemisia*, *Chenopodiaceae*, *Helianthemum*, *Plantago*, *Rumex*, which rarely exceed 5–10%), and abundant representation of arboreal pollen (AP), dominated by conifers (mainly *Pinus*) and deciduous forest taxa (*Betula*, *Corylus*, *Alnus*, *Salix*, *Ulmus*, *Populus*, *Acer*, *Fraxinus*, *Fagus*, *Tilia* and deciduous *Quercus*), with values around 25–30% (Fig. 4). Representation of Poaceae and aquatics (*Cyperaceae*,

Ranunculus, *Myriophyllum* and *Potamogeton*) in this zone is not significantly different to the rest of the sequence. This pollen spectrum is not consistent with a pre-Holocene deposit as would be inferred from the two dates (15.8 cal ka BP and 12.6 cal ka BP) from this interval. These spectra, together with the lack of sedimentological evidences for a hiatus, indicate that these dates are too old. Both pollen and sedimentological data suggest these are Holocene sediments, but given the absence of chronological control the record from this zone is not considered further in this study.

BSM-I (1167.5–815 cm depth; 9800–8200 cal yr BP)

Arboreal pollen varies between 60 and 80% of the total pollen abundance, and in some cases it exceeds 85%. *Pinus* is the main arboreal taxon, but deciduous taxa are well represented by *Betula*, *Corylus* and deciduous *Quercus*, with some significant fluctuations in *Betula*. *Juniperus* is also present with percentages above 6%. Evergreen *Quercus* and Mediterranean shrubs (*Pistacia*, *Rhamnus*, *Phillyrea*, *Buxus*, *Sambucus*, *Ephedra fragilis* and *Ephedra distachya*) are present in relatively low but continuous percentages. The first *Tilia* is recorded at 870 cm depth (8500 cal yr BP); this timing is consistent with other records from the region (Montserrat-Martí, 1992; González-Sampériz et al., 2006; Miras et al., 2007; Pélachs et al., 2007). Poaceae dominates the herbaceous stratum, while the abundance of *Helianthemum* significantly declines and *Artemisia* decreases in importance. *Myriophyllum* is the dominant aquatic. A significant change is found towards the end of the zone (860–815 cm depth; 8400–8200 cal yr BP) characterized by a sharp decline in *Betula*, *Corylus* and deciduous *Quercus*, the virtual disappearance of Other Mesophytes (Fig. 4) and the total absence of *Tilia*. *Pinus* increases to its maximum in the whole sequence, reaching 75%, and *Helianthemum* reappears at this time. This is a phase of high variability in fire activity, although charcoal counts are very low. *Pinus* and *Juniperus* show a positive correlation with MS within this zone, while *Betula*, *Corylus*, *Q. faginea*, evergreen *Quercus* and *Myriophyllum* are negatively correlated with MS (Table 3b). Thus, MS is correlated negatively with moisture-adapted and temperate taxa, but positively with more drought-resistant taxa such as *Pinus* and *Juniperus*.

BSM-II (815–491 cm depth; 8200–5700 cal yr BP)

After the short, abrupt vegetation change previously described, forest contracts slightly but there is considerable compositional variability. *Pinus* decreases to 35% and *Juniperus* is also highly reduced in abundance. Deciduous taxa, mainly *Betula*, *Corylus* and deciduous *Quercus*, show large and more continuous expansion reaching their maximum values in the sequence. *Tilia* reappears and is constantly present at moderate levels throughout the zone. Evergreen *Quercus* declines to its minimum values, while Mediterranean Shrubs fluctuate in abundance. The first isolated

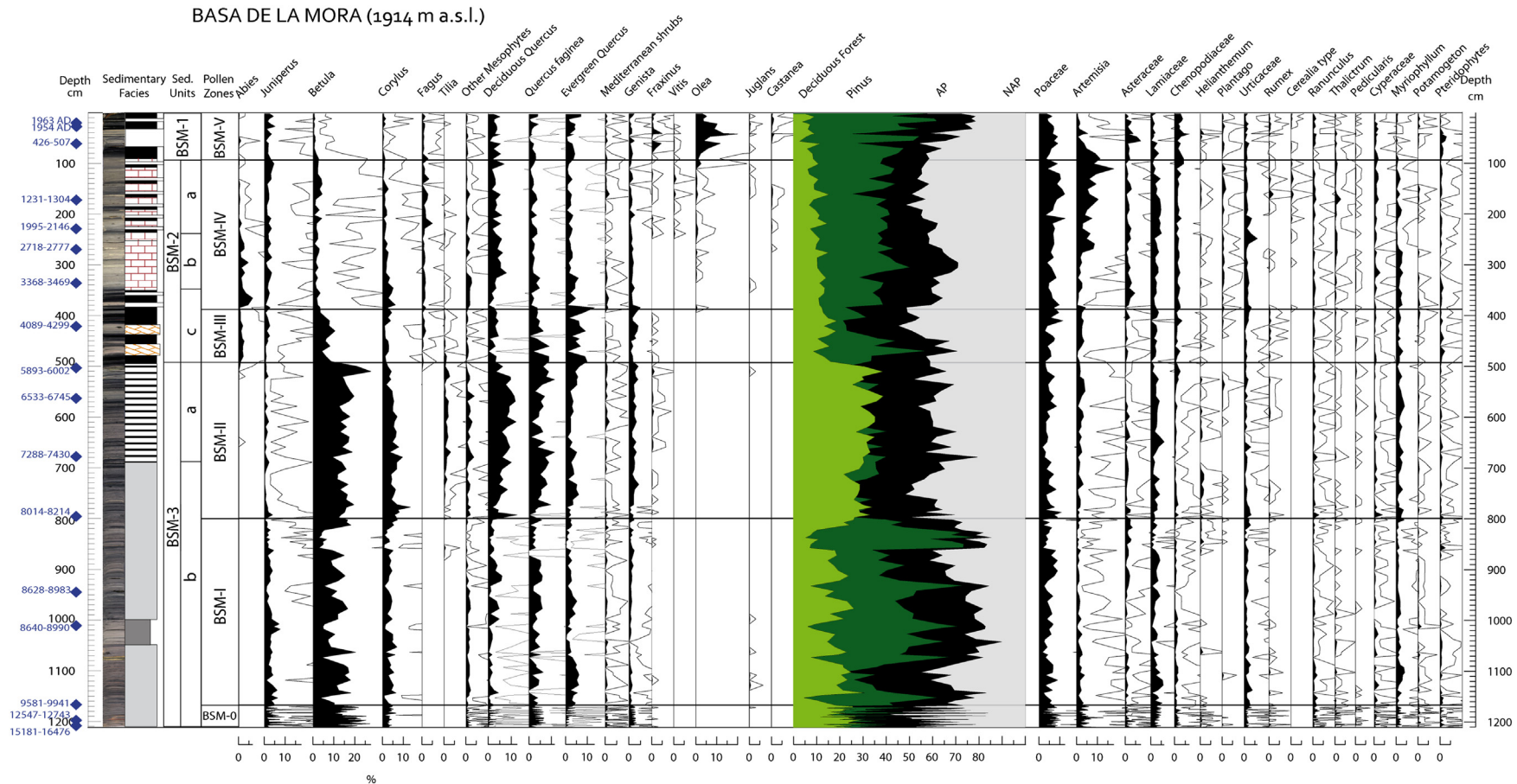


Fig. 4. Pollen diagram of selected taxa from Basa de la Mora sequence, plotted in depth. Other Mesophytes curve groups *Alnus*, *Salix*, *Ulmus*, *Populus* and *Juglans* pollen types; Mediterranean shrubs groups *Pistacia*, *Rhamnus*, *Phillyrea*, *Buxus*, *Sambucus*, *Ephedra fragilis* and *E. distachya* pollen types; and Deciduous forest curve groups *Betula*, *Corylus*, *Fagus*, *Tilia*, deciduous *Quercus* and *Other Mesophytes* pollen types. As usually, AP includes all the arboreal taxa (trees and shrubs) and NAP the herbaceous component excluding aquatics and ferns.

Table 3b
Correlation between MS and pollen taxa in the different pollen zones.

	BSM III and BSM IV (93–491 cm)		BSM II (491–815 cm)		BSM I (815–1168 cm)	
	MS		MS		MS	
	r	p	r	p	r	p
<i>Pinus</i>	0.453	0.003	0.444	0.006	0.464	0.001
<i>Juniperus</i>	0.351	0.023	–	–	0.339	0.021
<i>Betula</i>	–0.573	<0.001	–	–	–0.517	<0.001
<i>Corylus</i>	–	–	–	–	–0.292	0.049
<i>Tilia</i>	–	–	–0.537	0.002	–	–
Dec. <i>Quercus</i>	–	–	–0.528	0.001	–	–
<i>Quercus</i> fag.	–0.401	0.009	–	–	–0.373	0.018
Ever. <i>Quercus</i>	–0.378	0.014	–	–	–0.505	<0.001
<i>Artemisia</i>	–	–	–0.433	0.007	–	–
Cyperaceae	–	–	0.414	0.017	–	–
<i>Myriophyllum</i>	–	–	–	–	–0.592	<0.001

appearance of *Abies* occurs at 646 cm (7200 cal yr BP). The NAP is mainly composed by *Poaceae*, *Artemisia* and *Lamiaceae*, as in the rest of the record. Aquatic plants are well represented by Cyperaceae, *Pedicularis*, *Ranunculus* and *Potamogeton*, although *Myriophyllum* is dominant and reaches its highest values in the sequence. Deciduous *Quercus* and *Tilia* abundances show a strong negative correlation with MS (Table 3b). There is an increasing trend of fire activity, although the variability is high.

BSM-III (491–389 cm depth; 5700–3900 cal yr BP)

The beginning of this zone is characterized by a steep decline in deciduous forest taxa, mainly *Betula* (abruptly reduced by nearly 60%) and deciduous *Quercus*. In contrast, *Pinus* expands rapidly and *Juniperus* and evergreen *Quercus* increase slightly. *Fagus* appears for the first time, chronologically fitting the regional expansion (Montserrat-Martí, 1992; Pla and Catalán, 2005). The base of the zone is characterized by the permanent presence of *Abies* in the area, after its initial appearance shortly before. *Poaceae*, *Artemisia*, *Lamiaceae* and *Chenopodiaceae* are still the main NAP taxa and *Rumex* rises. No significant changes are recorded on the aquatic component except a decrease in *Myriophyllum* and a short-term disappearance of *Potamogeton* at the base of the zone. The conifer/mesophyte ratio is inverted at the top of the zone, just before the transition from Sub-unit 2c into Sub-unit 2b. Fire activity reaches a maximum towards the end of this zone.

BSM-IV (389–93 cm depth; 3900–700 ca yr BP)

The beginning of this zone is characterized by a change in forest composition. *Pinus* recovers and becomes the dominant arboreal taxon, *Abies* reaches its maximum abundance and *Betula* exhibits its minimum values. *Juniperus* and evergreen *Quercus* increase, but *Corylus* and Other Mesophytes only experience a slight increase. *Tilia* decreases progressively and disappears at top of the zone. In contrast, *Fagus* reaches its highest levels, at a time consistent with other records from the region (Pla and Catalán, 2005; Pérez-Obiol et al., 2012). A sudden and abrupt rise of *Artemisia* and further decrease in mesophyte taxa accompany the *Pinus*-dominant landscape. The NAP, of which *Poaceae* and *Artemisia* constitute the main elements, accounts for 40% of the pollen sum. There are two peaks of *Artemisia* in this zone, the youngest of which (when *Artemisia* reaches its maximum value in the whole sequence) coincides with the disappearance of *Abies* and *Tilia*. The aquatic component is markedly reduced in abundance, with low values of *Myriophyllum* and the absence of *Potamogeton* during the most of the zone contrasting with an increase in Cyperaceae. Cultivated taxa like *Olea*,

Vitis, *Castanea* and *Cerealia* type appear more continuously. Although there are some marked peaks of *Pinus* in this zone, the general trend is for relatively stable pine forest during the last phase of sedimentary Sub-units 2b and 2a. An abrupt decrease in charcoal concentration lasting several centuries was followed by a new abrupt increase in fire activity at the end of the zone.

BSM-V (93–0 cm depth; 700 cal yr BP-present, 1250–2008 cal AD)

This zone is characterized by important changes in both pollen and sedimentological records (Unit 1). The most relevant feature is the increase in *Olea* and *Fraxinus*. *Pinus* increases up to the 70%, but with very short episodes of where abundance is much lower (40%). The expansion of pine is coincident with the decline of *Abies*, *Betula*, *Corylus* and Other Mesophytes. Deciduous *Quercus* and, especially evergreen *Quercus* increase in abundance in the topmost part of the sequence. The NAP is still dominated by *Poaceae*, but *Artemisia* drops dramatically while *Asteraceae* and *Chenopodiaceae* reach their maximum values. *Myriophyllum* becomes less important and Cyperaceae dominates the aquatic assemblage. Variations in MS at this time are not correlated with vegetation composition changes. Fire activity is very high during most of the zone, but ceases in the top part of the record.

4.4. Chironomids

A total of 6422 chironomid head capsules were picked up, individually mounted and identified from 71 samples of the core BSM08-1A. Total chironomid biodiversity was represented by 18 taxa (up to 9 taxa per sample), belonging to three chironomid subfamilies: *Tanytarsinae*, *Orthoclaadiinae* and *Chironominae*. *Tanytarsus* gr. *lugens* was the most abundant all through the core, followed by *Procladius*, *Chironomus* and *Paratanytarsus*. *Chironomus* or *Paratanytarsus* are not shown in the diagram (Fig. 5) because they are present through the entire sequence and show no clear pattern of changes through the Holocene. The chironomid assemblage indicates that the lake has been always relatively shallow and oligotrophic, although relatively rich in organic matter. Quantitative analysis of the *Chironomidae* allows the sequence to be divided into 4 zones:

CHZ-1: Chironomid Zone 1 (1168.5–491 cm depth; 9895–5700 cal yr BP)

Low densities characterize this zone. *Tanytarsus* gr. *lugens* abundance is relatively low although with some fluctuations. *Procladius* reaches its maximum relative abundance within the core (30–60%), whereas *Pentaneurini* tribe appears through the entire zone although with a highly fluctuating distribution. The *Orthoclaadiinae* tribe is quite diverse, with an early representation of *Psectrocladius* gr. *limbatellus* and *Corynoneura* and a moderate representation of *Orthoclaadiinae* indet. (5–7%), which include several taxa related to water runoff and seepages (e.g. *Smittia*).

CHZ-2: Chironomid Zone 2 (491–357 cm depth; 5700–3600 cal yr BP)

The *Tanytarsinae* subfamily taxa (*Procladius* and *Pentaneurini*) is reduced in abundance, whereas the abundance of *Tanytarsus* gr. *lugens* increase and remains relatively high values throughout the zone (50–60%). Density values increase, although 3 samples from the base of the zone were almost sterile.

CHZ-3: Chironomid Zone 3 (357–56 cm depth; 3600–350 cal yr BP)

High densities occur, although they decrease towards the top of the zone. The main difference from the previous zone is the presence of *Psectrocladius* gr. *limbatellus* throughout the zone with relatively high abundances (up to 20%). *Procladius* reaches relatively high abundance (10–20%), although it does not reach previous values.

CHZ-4: Chironomid Zone 4 (56–0 cm depth; 350 cal yr BP–present; 1600–2008 AD)

The uppermost zone is characterized by a strong increase of *Psectrocladius* gr. *limbatellus*, together with *Pentaneurini* and *Corynoneura*, and a reduction in *Tanytarsus* gr. *lugens*. Density values particularly of *Procladius*, fluctuate, although its abundance is similar to the previous zone.

5. Discussion

5.1. The Early Holocene: strong Mediterranean influence and high climate variability (sub-unit 3b, BSM-I, CHZ 1, 9800–8150 cal yr BP)

During the Early Holocene, the Atlantic regions of Iberia were dominated by deciduous broadleaf trees (Muñoz Sobrino et al., 2005, 2007; Moreno et al., 2011) while the Mediterranean, mountain and inland areas were covered mainly by dense pine forest (Franco-Múgica et al., 2000, 2001; Carrión et al., 2010; Rubiales et al., 2010; Morales-Molino et al., 2012). The southern Pyrenees record both climate regimes in a relative small area: the Atlantic climate to the west and the Mediterranean climate to the east. These particular geographical features led to some marked differences in plant communities between the two regions at the onset of the Holocene. Increasing humidity was much pronounced in the Atlantic-influenced area, with a large expansion of mesophytes (Montserrat-Martí, 1992; González-Sampériz et al., 2006), while pine was the main tree taxon in the Mediterranean-influenced region (Miras et al., 2007; Pérez-Obiol et al., 2012). This suggests a stronger W–E precipitation gradient in the southern Pyrenees at the onset of the Holocene, with stronger influence of humid fronts in the west and persistent summer drought in the east.

In the BSM sequence, located at the modern transition between the Atlantic and Mediterranean climate regimes, the Early Holocene is characterized by the dominance of conifers over mesophytes (BSM-I) (Fig. 5). High values of pines and *Juniperus* reflect a continental Mediterranean-climate influence during this period. The fire regime is not characterized by either frequent or virulent fires, probably because of fuel limitation as pine-dominated forests are less flammable than broadleaf woodlands. The dominance of *Pinus* over deciduous taxa suggests the existence of extreme seasonal temperatures and marked summer drought during the Early Holocene. However, deposition of carbonate-poor laminated Facies 1 and 3 indicates permanent and relatively high lake levels with abundant sediment delivery by run-off. High values of MS are related to the presence of paramagnetic minerals eroded from ophite outcrops and are consistent with high-energy transport to the lake. High correlation between MS and Ca and TIC is indicative of the detrital origin of carbonate minerals and supports high erosion rates during this period. The high abundance of non-lacustrine *Orthocladiinae* taxa, related to inlet streams, in this zone supports the idea of increased runoff due to high rainfall. The *Procladius* genus has been reported to be important in the Early Holocene in other European regions (Heiri et al., 2003) and its high abundance is consistent with higher lake levels because it inhabits fine sediments in the profundal zones of lakes (Saether, 1979; Prat et al., 1992).

The Early Holocene maximum in seasonality in the Northern Hemisphere may have been responsible for particularly cold winters and hot summers. In the southern Pyrenees, this would have led to increased snow accumulation in winter and subsequent large snowpack melt during the warmer summer months leading to higher run-off. Evapotranspiration and low precipitation during summer drought periods would be largely compensated by increased melting water, leading to higher lake levels. The negative correlation between moisture-adapted taxa and MS supports the idea that run-off would be likely linked to melt processes rather than direct precipitation. Furthermore, positive correlation between MS and drought-resistant taxa such as *Juniperus* and *Pinus* confirms that run-off is related to increased continentality during this period.

The relatively dry and cold Early-Holocene climate of the Basa de la Mora (BSM) is in agreement with many studies from western Europe (Leira and Santos, 2002; Bjune et al., 2005) and North America (Shuman et al., 2001; Zhao et al., 2010), which have inferred a cooler and drier climate probably related to weakened ocean conveyor circulation as the rapid, global increase in temperature provoked large input of freshwater from the Laurentide sheet into the North Atlantic, weakening Labrador Sea deep convection (Kaplan and Wolfe, 2006; Renssen et al., 2009, 2012).

Superimposed on the long-term insolation-driven climate trend, the BSM sequence shows significant short-term (sub-millennial) shifts in pollen percentages and sedimentological features during the Early Holocene. Such shifts occurred at 9.7, 9.3, 8.8 and 8.3 cal ka BP and are mainly characterized by short-term expansion of pine, accompanied by large reductions in all deciduous taxa but most particularly in *Betula*, implying a substantial reduction in humidity. The highest MS values of the whole sequence are also recorded during these events, indicating that these periods are characterized by particularly intense run-off and sediment delivery from the catchment (Fig. 5). Cold and relatively humid winters with large amount of snow accumulation, and the subsequent snowpack melt and runoff, could be responsible for increased erosion in the catchment. This interpretation is supported by the sharp and discontinuous presence of rheophilous and non-strictly lacustrine chironomid taxa during these short-events. Low percentages of TOC and low TOC/N ratio also point to reduced vegetation in the catchment (Fig. 5). Phases of reduced forest may be due to a downward displacement of the treeline, supporting the occurrence of cooler temperatures. These events were as short-lived periods of drier and cooler conditions. Sedimentary phases with particularly high sedimentation rates associated with arid conditions have been recognized in the Central Ebro Basin complex during this period (Sancho et al., 2008; Gómez-Paccard et al., 2013). The strong response of the vegetation and hydrology at BSM indicates that climate instability was characteristic of the Early Holocene. Similar evidences for Early Holocene climatic oscillations have been widely recognized throughout the North Atlantic region (O'Brien et al., 1995; Alley et al., 1997; Bond, 1997, 2001; Mayewski, 2004; Frigola et al., 2007).

The first Early Holocene cold event is recorded just at the beginning of the BSM sequence at 9.8–9.7 cal ka BP. Since the BSM record starts at 9.8 cal yr BP, we suggest that this may be coincident with the short-lived 9.95 ka cold anomaly detected in the NGRIP record (Rasmussen et al., 2007). The impact of this anomaly has been previously noted in the western Mediterranean as a phase of forest decline (Fletcher et al., 2010b), as in BSM sequence. A global event centred in 9.3 ka cal BP has been widely recorded in many sequences from the North Atlantic and Europe (Haas et al., 1998; Rasmussen et al., 2007; Fletcher et al., 2013). In the BSM sequence, this interval coincides with an expansion of pine forest and decline in mesophyte taxa but there is no sedimentological change. The

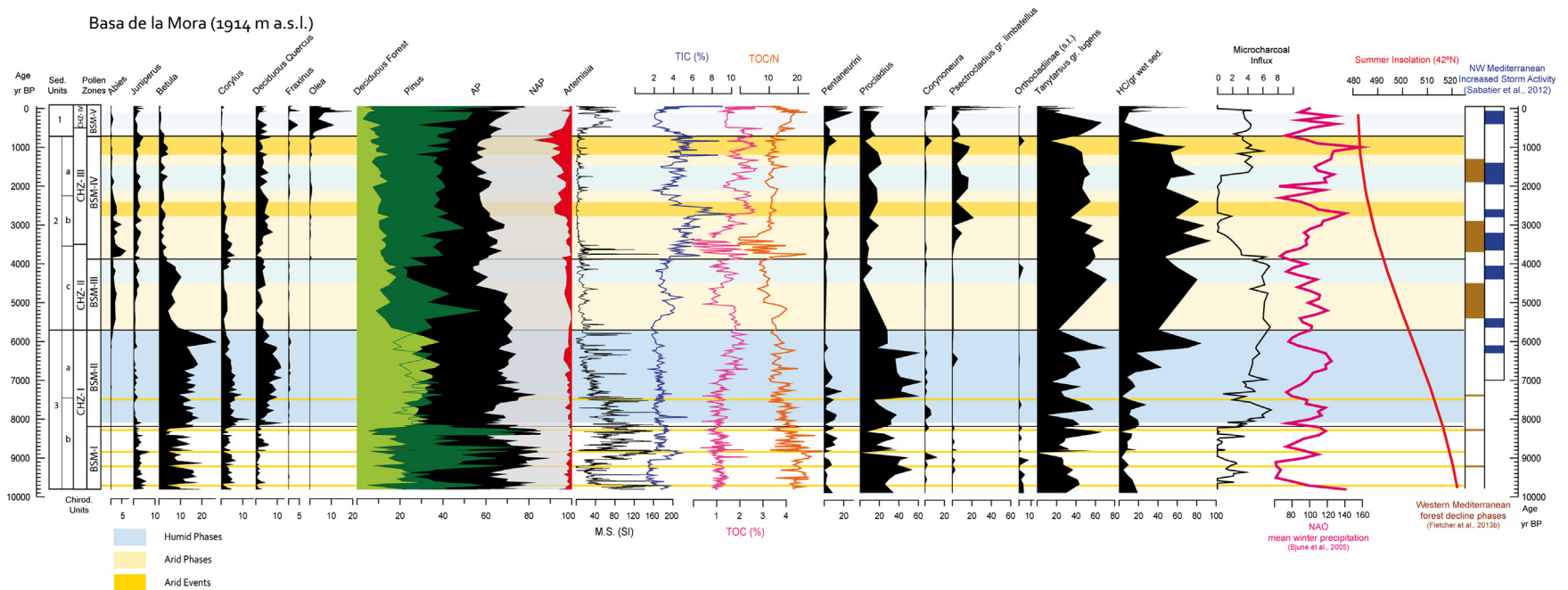


Fig. 5. Diagram plotted in age, including selected pollen taxa, geochemical parameters, chironomid taxa and microcharcoal influx curves of Basa de la Mora sequence compared to NAO summer insolation curve for latitude 24°N, regional phases of deforestation (Fletcher et al., 2013) and phases of increased storm activity (Sabatier et al., 2012) in the Western Mediterranean. Note: Orthocladinae (s.t.) means sum of rheophilous (see page taxa). Blue horizontal bars represent humid phases whereas yellow and orange bands represent arid phases and further arid events respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

next cold and arid event occurs at 8.8 ka cal BP. In BSM sequence, this event is resulted in major shifts in vegetation and sediment deposition and the apparent disappearance of chironomids. This phase coincides with the only occurrence of Facies 3 and the high TOC/TN ratios characteristic of this unit suggest a well-vegetated watershed, dominated by *Pinus*. The 8.8 ka cal BP cool event is reported in the Arctic by Ebbesen et al. (2007) but has not previously been reported in southern Europe.

The next event is recorded at 8.3 ka cal BP. This is the most remarkable vegetation shift in the BSM record, with *Pinus* reaching its highest values and *Betula* dropping to its minimum. Taking into account the age–depth model uncertainties for this period (8300 ± 100 cal yr BP), this event could be synchronous with the 8.2 ka cool event (Alley and Agustsdottir, 2005; Rasmussen et al., 2007), triggered by a large freshwater discharge from former glacial Lake Agassiz into the North Atlantic Ocean, causing a reduction the Atlantic Meridional Overturning Circulation (AMOC) (Hoffman et al., 2012). The high-resolution study carried out in BSM sequence for this period indicates a minimum timing of 150 years and maximum of 200 years for the 8.2 ka event. This timing agrees with the precise characterization of the 8.2 ka event obtained from trapped air in a Greenland ice core (GISP2) (Kobashi et al., 2007). The abrupt increase in pine in BSM matches the spread of *Pinus* recorded in the Alps (Blarquez et al., 2009), Switzerland (Tinner and Lotter, 2001) and northern Spain (Muñoz Sobrino et al., 2007), suggesting a widespread impact in mountain/alpine regions. The 8.2 event is widely recorded in the north-eastern of the Iberian Peninsula, where human settlements located in a particular harsh region of the Central Ebro basin moved towards more humid areas during this interval (González-Sampérez et al., 2009).

The rapid response of the vegetation to these short climate shifts, related to changes in the North Atlantic, seems to be amplified in the BSM sequence because of its ecotonal location for some species. The highly responsive nature of the vegetation record highlights the climate sensitivity of high altitude transitional areas to environmental changes, as previously demonstrated for the central Pyrenees during the Lateglacial period in El Portalet sequence (González-Sampérez et al., 2006).

5.2. The Mediterranean “climatic optimum” (sub-unit 3a, BSM-II, CHZ-I, 8100–5700 cal yr BP)

The Mid-Holocene is the period with the greatest forest development in Europe, when treeline moved upward and reached its maximum elevation in most mountain regions (David, 1993; Ali et al., 2003; Carnelli et al., 2004; Ortu et al., 2008; Favilli et al., 2009; Talon, 2010; Cunill et al., 2011; Magyari et al., 2012). In northern Europe, forest expansion is related to higher summer temperature (Davis et al., 2003; Bjune et al., 2005; Nesje et al., 2006), while in southern Europe this is an interval of increased humidity (Carrion et al., 2010; Colanese et al., 2010; Spötl et al., 2010; Stoll et al., in press).

There is a marked shift in the vegetation composition after ca 8.2 ka BP in the BSM sequence (Fig. 5). *Betula*, *Corylus* and deciduous *Quercus* became the dominant AP elements, *Tilia* and other mesophytes were present, and conifers declined to their minimum values, with pine oscillating between 20 and 30% and juniper between 2 and 3% (Fig. 5). This assemblage is very different from that of a dense conifer community near the lake (Court-Picon et al., 2005). The high values of *Betula* (up to 26%) in the BSM sequence compare fairly well with similar high values recorded in the Pyrenean sequence of El Portalet peatbog (González-Sampérez et al., 2006), located at 1802 m a.s.l., Lake Burg (Pèlachs et al., 2007), located at 1821 m a.s.l. or Tramacastilla lake, at 1682 m a.s.l., where birch accounted for 40% of the total pollen

(Montserrat-Martí, 1992). The similarity between these sequences indicates that *Betula* grew at higher altitude, in the upper part of the montane belt and probably reaching the subalpine belt. The rise of birch and the consequent drop of pine at BSM could result from either an increase in annual precipitation or reduced evaporation, as a consequence of decreased continentality, favouring water-demanding taxa. High charcoal values indicate increased regional fire activity. An increase in moisture does not necessarily imply reduced fire activity; the expansion of mesophytes, which are more flammable than most mountain pines (Blarquez and Carcaillet, 2010), provides high amounts of fuel at an altitudinal zone normally devoid of large forest to be burnt. Only minimal changes in summer climate or lightning would be required to promote large and virulent fires, leaving a sizeable imprint in the charcoal record. In addition *Betula* is a pioneer taxa that spreads well after fire disturbances (Blanco-Castro et al., 1997; Morales-Molino et al., 2012). This pattern has been also found in El Portalet sequence (Gil-Romera et al., submitted for publication) and in many other Holocene records from the European mountains (Tinner et al., 1999; Colanbaroli et al., 2008; Vannière et al., 2008) as well as in current patterns of fire occurrence (Pausas and Paula, 2012).

The interval from 8100 to 5700 cal yr BP was characterized by stable environmental conditions in the BSM catchment, as inferred from the stable vegetation composition and the lack of marked decreases in any tree taxon despite the high fire activity. Sedimentological and geochemical indicators indicate a stable, relatively deep lacustrine environment. The laminated nature of Facies 2 is consistent with high lake level and the activity of several inflow streams. The finer grain size of Facies 2, in comparison to laminated Facies 1 and 3, indicates even higher lake levels. Low values of TIC and Ca suggest dilute water, and the lack of a significant correlation between Ca and MS indicates that delivery of carbonates from the catchment through run-off was negligible. Moreover, the decrease in MS and TOC/N along with the increase in TOC suggests a more vegetated environment that would limit the erosive effect of precipitation. High and constant *Myriophyllum* values and the chironomid association also reflect a well-established, deeper lacustrine environment; as genus *Procladius* presents its highest abundances along the sequence and *Tanytarsus* gr. *lugens* is also important in the chironomid assemblage. Moreover, the increase of littoral and macrophyte-related taxa such as *Corynoneura* or *Pentaneurini* tribe (e.g. Brodersen et al., 2001) during this period, reflects the greater development of aquatic vegetation in the lake favoured by milder climate conditions.

The Mid-Holocene warmer conditions occurred when the flux of meltwater from the Laurentide ice sheet stopped and deep convection in the Labrador Sea led to enhanced transport heat over the Atlantic-influenced area (Renssen et al., 2009, 2012). Increased meridional circulation in the North Atlantic as the Laurentide sheet waned could bring warmer condition to the Iberian Peninsula. Changes in the SST and shifts in insolation triggered reorganization of the atmosphere circulation and strengthened meridional atmosphere circulation. A northward shift of the monsoon system and its associated rainfall belt gave rise to particularly humid conditions in the Sahara and Sahel (deMenocal et al., 2000; Hély et al., 2009). Enhanced westerlies could bring increased summer humidity over the Iberian Peninsula, as inferred from the spread of broad-leaf taxa in this region (Carrion et al., 2001a,b; Carrion, 2002; Pantaleón-Cano et al., 2003).

Although this interval (8.2–5.7 cal ka BP) is the most humid period recorded at BSM, high MS and a slight expansion of *Pinus* indicates a short-lived arid event around 7.5 cal ka BP. However, mesophytes only decrease slightly suggesting this interval was less pronounced than in previous arid intervals. This event is broadly

coincident with the arid 7.4 event detected in southern Spain (Jalut et al., 2000) that has been related to the emergence of the Neolithic in southern Iberia (Cortés Sánchez et al., 2012), and also correlates with a phase of forest decline detected in the western Mediterranean (Fletcher et al., 2010a). In the central southern Pyrenees, this phase does not represent a dramatic change in moisture supply and vegetation recovers rapidly.

5.3. The end of the Middle Holocene: transitional phase (sub-unit 2c, BSM-IV, CHZ-II, 5700–3900 cal yr BP)

The evolution of the landscape in southern Europe from 6 ka (or even earlier) onwards has been widely assumed to be influenced by both climate and human forcings (Oldfield and Dearing, 2003; Vannièr et al., 2008; Roberts et al., 2011; Sadori et al., 2011). Many palynological studies show a clear increase of anthropogenic indicators from the Middle Holocene, pointing to an intensification of human activities and a subsequent change in the vegetation composition related to forest clearance for pastures and agriculture fields (Jalut et al., 2009). However, some of these taxa are naturally found in xeric Mediterranean ecosystems (De Beaulieu et al., 2005) and this makes it difficult to discriminate between climate and anthropogenic forcings. The spread of xeric vegetation across the Mediterranean region during Middle-Holocene does not necessarily imply anthropogenic degradation of the landscape (Collins et al., 2012). In addition, fire activity in Mediterranean areas increased significantly at this time and its impact on vegetation composition has to be taken into consideration (Colombaroli et al., 2008, 2010; Vannièr et al., 2008, 2011). Increased fire activity can result from anthropogenic activities but also reflects the climatic shift towards arid conditions (Carrión et al., 2001a, 2010; Fernández et al., 2007; Fletcher and Goñi, 2007; González-Sampériz et al., 2008; Morellón et al., 2008; Jalut et al., 2009; Corella et al., 2010; Anderson et al., 2011). The expansion of heliophytes (like *Artemisia*, *Chenopodiaceae*, *Asteraceae*, *Rumex*, *Plantago*, *Poaceae*, and Mediterranean taxa similar to *Cerealia*.) observed during this period is favoured by increased fire, increased aridity, and anthropogenic activity. Overall, the complex changes found in Mediterranean areas at the end of the Mid-Holocene are not necessarily related to intense human pressure, but could equally well be explained by the trend towards drier conditions.

There is a sharp change in the vegetation cover and sedimentological features in the BSM sequences at 5.7 cal yr BP. The pollen record in BSM-IV is characterized by a pronounced increase in pine and decrease in mesophytes, mainly *Betula*, in combination with a rise in *Juniperus*, deciduous and evergreen *Quercus* and heliophytes (*Artemisia* and *Chenopodiaceae*). The replacement of mesophytes by conifers suggests a change from humid to drier conditions or, at least, a significant shift in the seasonal distribution of the precipitation since reduced summer rainfall is unfavourable to the broad-leaf taxa. The sedimentary shift is defined by an increase in carbonates, indicating lower lake levels (Sub-unit 2c). Lower values of MS suggested reduced sediment transport as consequence of lower run-off and inflow streams, which in turn indicates reduced precipitation or meltwater inputs. The decrease in allochthonous sediments is reflected in lowered sedimentation rates and deposition of carbonate Facies 6, which reflects high carbonate productivity in a littoral setting with low and fluctuating water level. The decline in *Myriophyllum* is consistent with a reduction in water level (Figs. 4 and 5). Moreover, the sharp decrease in *Procladius* and the near disappearance of non-lacustrine *Orthocladiinae* taxa also indicates reduced runoff and stream inflow during this period. The increase in chironomid abundances, mainly *Tanytarsus*, could indicate increased decomposition rates in the sediments.

Both biological and sedimentological indicators are consistent with a trend to increased aridity and a persistent arid phase between 5.6 and 4.6 cal ka BP (Fig. 5). Similar vegetation changes have been recognized in other Pyrenean sequences (Pèlach et al., 2007), in southern Spain (Jiménez-Moreno and Anderson, 2012) and in Mediterranean records (Carrión et al., 2010). Fletcher et al. (2013) have identified a major phase of deforestation in the Western Mediterranean during this period. The coincidence between lowered lake levels and forest decline supports the idea of climate as the main forcing. A major climate shift has been recognized in many other regions at this time, including the end of wet conditions in the Sahara between 6 and 5.5 cal ka BP (deMenocal et al., 2000; Kröpelin et al., 2008), and lake-level and vegetation changes indicating drier conditions in eastern North America (Shuman et al., 2001; Menking et al., 2012; Zhao et al., 2010). The similarities in climate changes between such different geographic areas during the Mid-Holocene suggest broad-scale changes in the coupled ocean-atmosphere circulation. This large-scale and synchronous climate shift may be related to changes in global atmospheric circulation. The weakened summer insolation in North Hemisphere led to a southward shift in the Inter Tropical Convergence Zone (ITCZ) and thus, the summer Asian monsoon also weakened considerably (Wanner and Brönnimann, 2012). Readjustment of these two main climatic system drivers led to the establishment of similar conditions to present atmospheric teleconnections (ENSO) since ca 5.5 ka (Wanner et al., 2008; Carré et al., 2012; Fletcher and Moreno, 2012). Southward movement of the ITCZ favoured southward shift of the sub-tropical North Atlantic high pressure and led to increased summer aridity in the Iberian Peninsula (González-Sampériz et al., 2008; Morellón et al., 2009; Carrión et al., 2010; Corella et al., 2010; Valero-Garcés and Moreno, 2011). As the North Atlantic high-low pressure system moved away, westerlies became weaker and lost their capacity to penetrate inland.

A change towards wetter conditions is observed in the BSM sequence between 4.5 and 3.9 cal ka BP, marked by increased abundance of mesophytes, and the recovery of *Betula* and deciduous *Quercus* values (Fig. 5). This humid period corresponds well with a phase of increased storm activity recorded in the Gulf of Lion (Sabatier et al., 2012), suggesting stronger and southward migration of the westerlies. However, the total AP decreases during this phase. This reduction of the arboreal pollen in the BSM sequence occurs at the same time as the first deforestation phase recognized in the Pyrenean sequence of Tramacastilla at ca 4000 BP (Montserrat-Martí, 1992). However, no other indicator of anthropogenic pressure was found during this period in the BSM sequence suggesting that the vegetation shift was mainly climate driven. The high regional fire activity detected during this period is the culmination of a previous trend. Although there was an initial dry phase when fire occurrence was linked to the presence of pine forest, higher charcoal influx values during this subsequent humid phase are linked with the spread of mesophyte forest. The fact that fire is high during both humid and arid spells, reflects on the one hand more permanent drying conditions than any time before in the Holocene leading to frequent fire-conducive conditions coupled with relatively high fuel availability from mesophyte vegetation, and on the other hand, the strengthening of fire activity during any interval of mesophyte forest expansion when fire-conducive conditions occur.

5.4. The Late Holocene: aridity crisis (sub-units 2b–2a, BSM-VI, CHZ-III, 3700–700 cal yr BP)

Complex societies developed across the Mediterranean during the Late Holocene and human pressure on the landscape intensified and expanded (Carrión et al., 2007; Bal et al., 2011; Finné et al.,

2011; Magyari et al., 2012). High altitude palaeoenvironmental records, where anthropogenic activities would have been limited due to both severe weather and difficult access, provide an opportunity to isolate the climate signal influencing vegetation evolution in recent times (Pérez-Sanz et al., 2011).

The BSM sequence reveals a well forested landscape during most of the late Holocene (AP abundance around 70%, BSM-IV), indicating negligible anthropogenic pressure until ca 1150 cal yr BP, when the first evidence of forest management is found. The trend towards increased aridity that started during the Mid-Holocene transition intensified considerably at 3700 cal yr BP. The pollen record (BSM-V) is characterized by a sharp fall of *Betula* and the disappearance of birch from this area. The expansion of conifers (*Pinus* and *Juniperus*, which reaches its maximum proportions of the whole record), indicates either a reduction in annual mean precipitation or a significant change in the seasonal distribution of precipitation (Franco-Múgica et al., 2000). The *Pinus* expansion in BSM is coeval with an expansion in other high altitude Pyrenean sites (Pèlachs et al., 2011), which suggests it is more likely to be controlled by changed climate than by human action. At ca 2900 cal yr BP, *Artemisia* starts to spread rapidly and *Myriophyllum* decreases strongly (BSM-V). Traditionally, the *Artemisia* expansion has been explained by an increase in pastoral activity during the Late Holocene. However, modern values of *Artemisia* rarely reach 2% even though there is moderate pastoral activity in the BSM area. Given that there is no evidence for major deforestation at the time of the *Artemisia* expansion, it seems unlikely that this represents an interval of more intense anthropogenic activity than today. Nor is the *Artemisia* expansion synchronous with the presence of coprophilous fungi, indicative of intensive pastoral land use (López-Merino et al., 2010). This suggests the *Artemisia* expansion at the Basa de la Mora site indicates a climatically-induced expansion of dry steppe. There is evidence for a period of intensified aridity across the Mediterranean at around 2900–2400 cal yr BP (Jalut et al., 2000).

The deposition of carbonate-rich massive Facies 5, characterized by the presence of authigenic calcite crystals, gastropods, pennate diatoms and mottling textures, indicative of bioturbation, provides evidence for lowered lake levels and the development of a larger palustrine area (Fig. 5) at the time of the expansion of dry steppe. Facies 5 characterizes most of littoral core BSM-2A-1U, supporting our interpretation of the depositional environment. The presence of partially dissolved authigenic crystals of calcite and gypsum in Facies 5 suggests the lake was ephemeral and may have desiccated at times. The strong negative correlation between MS and TIC indicates that decreased runoff, and thus reduced external water supply into the lake, led to increased concentration of the lake water and authigenic carbonate precipitation. Furthermore, the negative correlation between MS and drought-resistant taxa such as *Pinus* and evergreen *Quercus* and the positive correlation between MS and *Betula* strengthen the link between lack of run-off and precipitation deficit. Intercalation of organic Facies 4 supports the development of a palustrine area with high accumulation of organic matter. In addition, the high percentages of TOC and low TOC/N ratio indicate increased lacustrine productivity, consistent with shallower conditions. This expansion of littoral areas is consistent with the very high abundance of Cyperaceae while *Myriophyllum* values remain relatively unchanged. The higher percentages (up to 20%) of *Psectrocladius* gr. *Limbatellus* than in previous zones also indicates an increase lacustrine productivity, as this genus is associated with productive environments and/or littoral areas with abundance of biofilm primary production on stones or macrophytes (Rieradevall et al., 1999; Brodersen et al., 2001).

There is no charcoal in the BSM between 3.2 and 1.5 cal ka BP. An interval of two millennia without fire is highly unusual as fire

activity is registered in most southern European sequences during this time (Tinner et al., 2005; Vescovi et al., 2007; Vannièrè et al., 2008; Colombaroli et al., 2010). Arid pulses could prevent forest development at high altitudes and, therefore, limiting charcoal production through fires but, considering the absence of any other clear biotic or abiotic indicators, it seems more likely that the lack of microcharcoal is linked to taphonomical issues affecting charcoal preservation during oxic periods and/or short sub-aerial exposure events (Facies 5).

There is a common pattern to the evolution of vegetation across the Western Mediterranean (including southern Iberia, northern Africa and Italy) during this interval. A general phase of forest decline has been recorded in marine record MD95-2043 from the Alborán Sea between 3.7 and 2.9 cal ka BP (Fletcher et al., 2013). In Zoñar sequence, low values of AP (<10%) and an expansion of steppe taxa occurred between 4 and 2.9 cal ka BP (Martín-Puertas et al., 2008). At Sierra de Gádor (Carrión et al., 2003), *Pinus* and evergreen oak expand at the expense of deciduous *Quercus* after 3940 cal yr BP. In Sierra de Baza, there was a replacement of mesophytic by more xeric taxa around 3800 cal yr BP (Carrión et al., 2007), while in El Cañizar de Villarquemado, mesophytes and deciduous *Quercus* decreased and steppe herbs increased between 4000 and 3800 cal yr BP. A similar pattern has been recorded in Italian sequences, with an expansion of sclerophyllous taxa between 3.9 and 3.4 ka (Sadori et al., 2010). These changes can all be attributed to both drier climate conditions and human activities, especially considering that several civilizations collapsed at ca 4000 cal yr BP (i.e., Akkadians: Cullen et al., 2000).

Peaks in *Artemisia* and high TIC percentages in BSM record mark two periods of increased aridity at 2.9–2.4, and at 1.2–0.7 cal ka BP (800–1300 AD). Both episodes are characterized by high TIC and TOC percentages and low TOC/TN ratios suggesting high precipitation of carbonates and high bioproductivity and content of autochthonous organic matter. These episodes are separated by a relative humid period between 2.1 and 1.5 cal ka BP. The arid phase between 2.9 and 2.4 ka cal BP is synchronous with a dry episode recorded in both western (Ferrio et al., 2006; Aguilera et al., 2012) and eastern Iberia, that led to a prominent decline in deciduous *Quercus* pollen in the Amposta sequence (Pérez-Obiol et al., 2011). Increased water level can be inferred from the significant reduction of TIC percentages between 2.1 and 1.5 cal ka BP. An episode of more humid conditions has been recognized in Iberia (Martín-Puertas et al., 2008, 2009; Corella et al., 2010; Currás et al., 2012), coinciding with the Iberian civilization and the Roman occupation and thus is called the Iberian–Roman Humid Period (IRHP). The NW Mediterranean region also registers an intensification of rainfall reflected by higher storm activity in the Gulf of Lion (Sabatier et al., 2012). However Fletcher et al. (2013) report another phase of forest decline in Western Mediterranean at this time (Fig. 5). Since wetter conditions should have positively affected forest development in the Mediterranean, where water is the greatest limiting factor, it is possible that depletion in tree mass could be related in some areas of Iberia to higher land use by the Romans (García-Bellido, 1985). However, we do not observe great exploitation of the subalpine belt at BSM suggesting that the vegetation composition, which runs in parallel with sedimentological features, is still primarily controlled by climate.

The second arid period recorded in BSM sequence matches the well-known Medieval Climate Anomaly (MCA: 900–1300 AD), a period of aridity recognized in most of south-western Europe (Seager et al., 2007; Mann et al., 2009) which led to notable agro-economic crisis in medieval societies. In Spain, it resulted in a major water deficit leading to lower lake levels and expansion of thermophytes and steppe taxa (Moreno et al., 2012b). In the BSM sequence, this phase coincides with the first signal of deforestation,

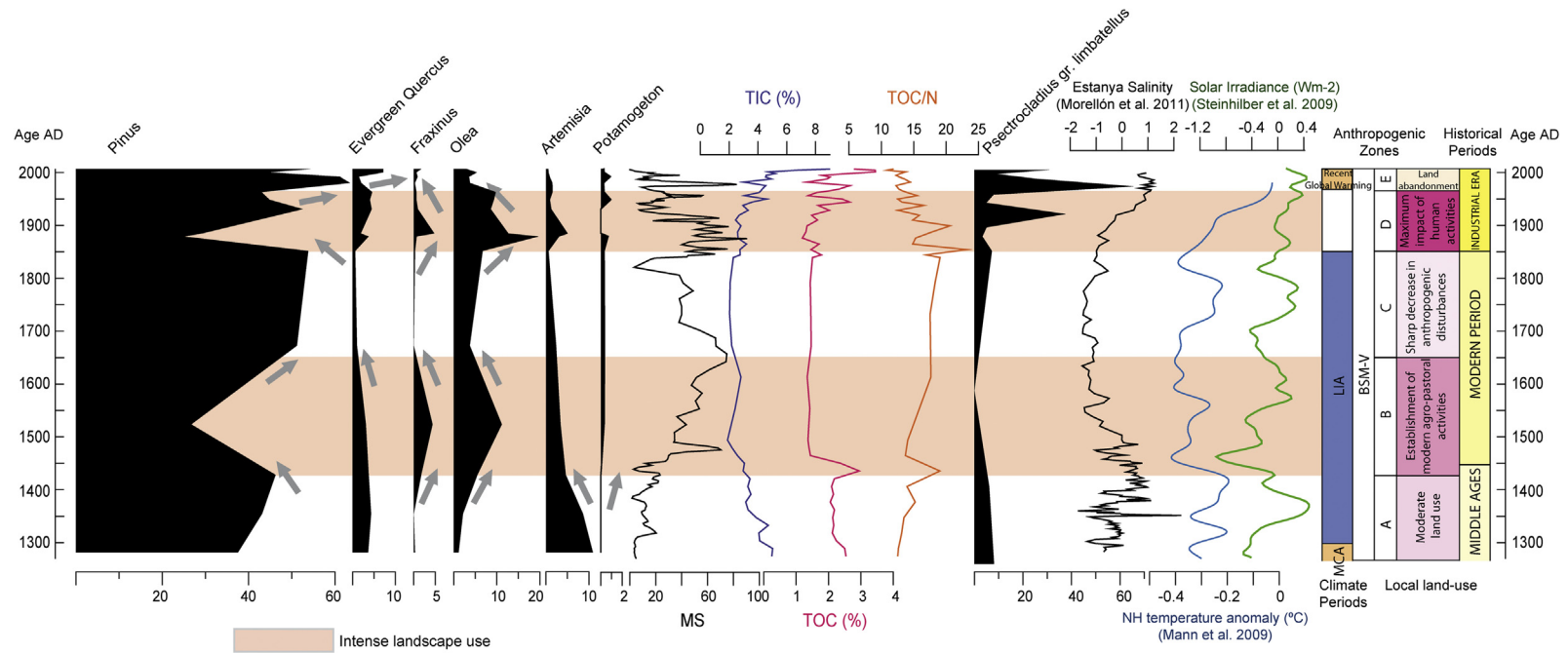


Fig. 6. Comparison of selected curves (pollen – *Pinus*, Evergreen *Quercus*, *Fraxinus*, *Olea*, *Artemisia*, *Potamogeton*–, geochemical proxies–MS, TIC, TOC, TOC/N– and chironomids – *Psectrocladius gr. limbatellus*–) from Basa de la Mora sequence with global and regional records (Estanya salinity (Morellón et al., 2011); NH temperature reconstruction (Mann et al., 1999) and Solar Irradiance (Steinhilber et al., 2009)) for the last 750 years, indicating the main climate and historical periods and the interpretation of local land use. Bands in rose mark the intense periods of anthropogenic activities. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

indicated by abrupt decreases in pine percentages. Charcoal influx increased ca 1700 cal BP, most likely because of either warmer conditions or strengthened regional fire activity in the lowlands.

Both episodes of depleted water availability correspond with maxima in reconstructed North Atlantic Oscillation (NAO) indexes (Fig. 5). This indicates that there is a fast response of palaeoenvironmental changes in the BSM record to changes in the North Atlantic. The persistence of a positive NAO index during 2.9–2.4, and at 1.2–0.7 cal ka BP, led to maximum winter precipitation in Scandinavia and to minimum winter precipitation in the Iberian Peninsula (Trouet et al., 2009).

5.5. The last 8 centuries (unit 1, BSM-V, CHZ-IV, 700 cal yr BP–present)

In contrast to most Pyrenean studies that indicate intensified human disturbance during at least the last two millennia (Riera et al., 2004; Guiter et al., 2005; Pélachs et al., 2011), the effects of anthropogenic pressure are only detected in the BSM sequence during the last 700 cal yr BP (Pérez-Sanz et al., 2011) (Fig. 6, BSM-V-A). As seen in Figs. 4–6, the increase in *Olea* marks an expansion of agricultural practises in the lowlands (Cañellas-Boltà et al., 2009) whereas large, short-term reductions in *Pine* indicate phases of deforestation and expansion of grazing lands at higher altitudes (Fig. 6, BSM-V-B). Parallel to *Olea*, *Fraxinus* also spreads. *Fraxinus* has traditionally been used in the region for hedgerows (Gómez and Fillat, 1981). Its parallel expansion to *Olea* marks the regional establishment of modern and intense agro–pastoral activities. The drop in *Artemisia* synchronous with clear evidence of increasing anthropogenic pressure in the highlands supports the idea that *Artemisia* is not an indicator of human activities in the BSM sequence.

The expansion of *Olea* and *Fraxinus* ceased, and deforestation temporarily stopped, between 1600 and 1850 AD coinciding with the second half of the Little Ice Age. This interval is characterized by the coldest conditions in the southern Pyrenees (González-Trueba et al., 2008; Morellón et al., 2012). A sharp decrease in evergreen *Quercus* coincides with these colder conditions. The rapid recovery of pine after intervals of deforestation emphasizes the fact that human disturbance at high altitudes was not strong and climatic conditions were the main determinant of vegetation changes. High values of MS and strong negative correlation with TIC during this period (Fig. 6; BSM-V-C) indicate increased sediment delivery to the lake and decreased carbonate productivity, both indicative of higher lake levels and increased runoff. The abundance of allocthonous organic matter, shown by low TOC and high TOC/N ratios, also supports the inference of high sediment delivery from the catchment. Fire activity was high for most of this period, confirming the occurrence of either regional fires linked to husbandry or local fires correlated with the occasional pine deforestation (Lasheras-Álvarez et al., 2013). Although it is difficult to distinguish between human and climate-induced fires in this period, all other records indicate an intensification of anthropogenic activities after 700 yrs BP. A general decrease in temperature coinciding with the Little Ice Age (LIA i.e. 1300–1850 AD) has been recorded throughout Europe. Higher storm activity occurred in the NW Mediterranean (Sabatier et al., 2012) (Fig. 5) while stronger climatic variability has been recognized in Iberia, although generally cold and humid conditions dominated (Benito et al., 2003; Moreno et al., 2008, 2012b; Valero-Garcés et al., 2008; Morellón et al., 2012).

A significant expansion of *Olea* associated with a marked phase of deforestation of the pine forest occurred right after the LIA (1880 AD) (Fig. 6, BSM-V-D). The Industrial Revolution in the 17th century brought major advances in agricultural techniques that resulted in

increased efficiency and production and led to increased supply of food and raw materials. As result of the improvement of the agricultural sector the population rose and demographic pressure in the southern Pyrenees increased up to its maximum at the end of 19th and the early 20th century (García-Ruiz and Valero-Garcés, 1998).

After 1960 AD pine forest recovered, AP increased up to 65% and there was a reduction in trees (*Olea*, *Fraxinus*) related to anthropogenic activities (Fig. 6, BSM-V-E). During the mid-20th century, social and economic changes in Spain forced population to migrate from villages into cities as the industrial sector developed. In Spain, and more specifically in the southern Pyrenees, mass migration took place in the last third of the 20th century, resulting in abandonment of the rural lands and gradual recovery of forests (Lasanta-Martínez et al., 2005). We observe a steep drop in fire activity during this phase, most likely as consequence of rural abandonment (Fig. 6, BSM-V-E). Geochemical proxies suggest a decrease of average lake level during the last 50 years. TIC percentages reach the highest values of the entire sequence, exceeding the values recorded during the MCA. Particularly high bioproductivity is shown by high TOC values, along with TOC/TN ratios and an increase of macrophyte-related taxa, such as *Corynoneura* and *Pentaneurini*, and especially *Psectrocladius* gr. *limbatellus*. Increases in bioproductivity in the recent period may be linked to the presence of cow stockbreeding near the lake. However, stockbreeding has taken place in this area at least since the last century (Lucio, 1982) but the increase in bioproductivity only occurs during the last 30 years. One possible explanation is that enhanced bioproductivity during the last decades reflects increased water temperatures. A global warming trend has been widely recognized over recent decades (IPCC, 2007) and an increase in temperature is also evident in the Mediterranean area (Brunetti et al., 2004; Vargas-Yáñez et al., 2008; Camuffo et al., 2010) and in north-eastern Spain (El Kenawy et al., 2012). Climate change in the Mediterranean area involves not only increased temperature but often decreased precipitation. A decrease in snowpack depth, snow cover and direct precipitation has been detected in the southern Pyrenees during the most recent period (López-Moreno, 2005; López-Moreno and Stähli, 2008). The recent drop in level at Basa de la Mora could be linked to the reduction in water availability in the southern Pyrenees, while the increase in bioproductivity could be related to the occurrence of warmer waters. The impact of the recent climate conditions on the lake sediments confirms the high sensitivity and rapid response of Basa de la Mora record to short-term climate shifts.

6. Conclusions

The multi-proxy sequence of Basa de la Mora (BSM) has recorded significant climate variability during the last ca 10 cal ka BP. Consistent shifts in vegetation, fire activity, depositional environments and aquatic communities throughout the sequence can be correlated with other regional and global reconstructions.

Higher seasonality between 10 and 8.2 cal ka BP caused high snow accumulation in winter and subsequent melt during warmer summers resulted in high lake levels. As a consequence of this high seasonal contrast, *Pinus* spread while mesophytes were restricted to watercourses. High climate instability during this period is illustrated by the occurrence of four short arid intervals at 9.7, 9.3, 8.8 and 8.3 cal ka BP, each characterized by a decrease in mesophytes and increased runoff. The most intense event occurred at 8.3 ± 0.1 cal ka BP, when vegetation diversity and abundance dropped to a minimum.

The most humid period in BSM sequence occurred between 8.2 and 5.7 cal ka BP. During this period, mesophytes expanded,

conifers retreated and the highest lake level was recorded. As a consequence of increasing biomass, fire activity also intensified.

The end of the Mid-Holocene marks the transition from a significant Atlantic influence (before ca 5.7 cal ka BP) into a typical Mediterranean climate with summer drought. A long-term trend towards increasing aridity, with decreasing lake levels and decreasing abundance of mesophytes started at 5.7 cal ka BP and intensified after ca 3.9 cal ka BP. During this period and until 700 cal yr BP human exploitation of the subalpine belt was minor and the vegetation composition was primarily controlled by climate.

The BSM record shows that the central Pyrenees are particularly sensitive to climate changes due to its geographical position between the Mediterranean and the Atlantic climate regimes.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at <http://dx.doi.org/10.1016/j.quascirev.2013.05.010>.

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