The evolution of the benthic foraminiferal assemblages from a 60 m core drilled in the Quaternary lagoon of Xeraco (Valencia, Spain) is studied. Two moments of warm climate, high sea-level and maximal marine influence in the lagoon have been identified. The first one at the base of the core, punctuated by highly frequent changes during the Upper Pleistocene, and the other one in the upper end of the series during the Holocene. The presence of a middle core section with low diversity and a nearly total absence of marine foraminifers is interpreted as a regressive interval of cold climate conditions linked to a glacial stage.

Keywords: Foraminifera, Quaternary, palaeoenvironments, Xeraco, Valencia.

Se estudia la evolución temporal de las asociaciones de foraminíferos bentónicos obtenidas de las muestras de un sondeo de 60 m de longitud realizado en la laguna costera cuaternaria de Xeraco (Valencia, España). Los cambios en estas asociaciones han permitido identificar dos momentos de máxima influencia marina, clima cálido y nivel del mar elevado. El primero de ellos en la base del sondeo y con presencia de cambios frecuentes en cuanto a esta influencia durante el Pleistoceno Superior, y otro superior con máxima influencia marina durante el Holoceno. La presencia de un tramo intermedio en que las especies de foraminíferos marinos están prácticamente ausentes, se identifica como una época de retroceso del nivel del mar, ligada a un enfriamiento global por glacialismo.

Palabras clave: Foraminíferos, Cuaternario, paleoambientes, Xeraco, Valencia.
1. INTRODUCTION

Although the Lower Pleistocene is poorly represented in the Valencian Community (some deposits possibly belonging to this epoch have been preserved in cliff sections, see Fumanal, 1995), the Middle Pleistocene (Marine Isotopic Stages 9 to 6) has been found in tectonically stable areas, such as in Xàbia (Usera et al., 1990a; Fumanal et al., 1991, 1993; Usera & Mateu, 1995; García-Forner, 1997) or Oliva-Pego (Viñals, 1996; García-Forner, 1997; Torres et al., 2014), where the stratigraphic record of the subsurface deposits shows two marine positive pulses alternating with continental sedimentation. The Upper Pleistocene (Marine Isotopic Stages MIS-5 to MIS-2) is easily identified in all the coastal sectors in Xàbia (Fumanal et al., 1993; Viñals et al., 1993; García-Forner, 1997), Pego (Dupré et al., 1988; Mateu, 1989; Viñals et al., 1989; García-Forner, 1997) and Elx (Blázquez, 2005; Blázquez & Usera, 2010). The Eemian interglacial corresponds to a high sea level whose deposits are preserved in littoral facies in low coasts and in dunes located next to the cliffs (Fumanal, 1995). Marine Isotopic Stages (MIS) 4, 3 and 2 are generally regressive with predominating continental, swamp or eolian facies, with some positive pulses (Martrat et al., 2004).

The current littoral is established during the Holocene. The Flandrian maximum in the Valencian coast took place around 6000 years BP, when the sea invaded a previously emerged continental surface. Some relict beaches have been identified in areas where the coastline was located several kilometres inland from its current position, such as the Oliva-Pego marsh. In other cases, like in Moraira, Xàbia or Peníscola, the inland advance of the coastline seems to have been less important (Fumanal & Viñals, 1989; Viñals et al., 1991).

Finally, the bars that close the current coastal lagoons formed in the upper Holocene. In some cases they can be found next to the Pleistocene ones as in Peníscola (Usera et al., 2006), Torreblanca (Usera et al., 1996a), Benicàssim (Usera et al., 1996b), l’Albufera de València (Sanjaume, 1985; Usera et al., 1990b; Sanjaume & Carmona, 1995; Rosselló, 1995; Santisteban et al., 2009; Marco-Barba et al., 2013; Carmona et al., 2016), l’Albufereta d’Alacant (Blázquez & Ferrer, 2012; Ferrer & Blázquez, 2012) and Elx (Blázquez, 2005; Blázquez & Usera, 2010).

These processes have also been identified in other areas of the Mediterranean such as the Asperillo and Barbate-Meca cliffs and the estuaries of the rivers Guadalete, Tinto-Odiel, Guadalquivir and Guadiana in the Gulf of Cadiz, several rambla deposits in the Alboran coast or the beach-barrier systems of Roquetas in Almería province or Doñana in the Gulf of Cadiz (Goy et al., 2003; Zazo et al., 2008).

This work focuses on one of these littoral lagoons, the Quaternary albufera of Xeraco, which has already been analysed in some preliminary studies (García-Blázquez et al., 2008; Usera et al., 2012). Our goal is the reconstruction of the environmental evolution in this location of the Mediterranean coast basing on the study of its fossil foraminiferal assemblages. This comprises the information provided by both the different species about their habitats and several statistical indices about species richness and diversity and assemblage complexity.

2. MATERIAL AND METHODS

The studied material was recovered from a borehole drilled in the locality of Xeraco (Valencia, Spain) (Fig. 1), in the site known as Els Borrans, on one of the still remaining Pleistocene bars. Its precise geographical location is: 39º02’53” N and 00º11’06” W.

![Figure 1. Geographical location of the Xeraco borehole (simplified from García-Blázquez et al., 2008).](image)

The studied core was extracted by a rotary drilling procedure that produced a cylindrical section of sedimentary material with a 15 cm diameter. Sediments were recovered by introducing 3-m-long cylinders with a toothed drilling head at their end. Subsequently the corresponding 3-m-long cylindrical sections of sediments were extracted by hydraulic pressure, placed in specific boxes and transported to the laboratory to be prepared and studied.

In order to avoid any eventual contamination during the core extraction the external 1 cm of each sample surface was discarded. Samples for micropalaeontological analysis were taken from the core every 10 cm in some sections in order to obtain as much precise information as possible about even short duration environmental changes occurring in the basin. A total of 205 samples, each one with an initial weight of about 50 g before the laboratory preparation phase, were analysed.

Samples were dried and then disaggregated in 40–50 °C water. This non-aggressive technique intends to prevent...
the destruction of foraminiferal agglutinated tests with an organic lining. The sediment was wet sieved through a 0.063 mm mesh that prevents the loss of small-sized species, typical of restricted environments, and juveniles. Samples were then dried, split into fractions using a splitter and, when possible, at least 300 foraminiferal tests per sample (or otherwise all available tests) were picked, classified under a binocular microscope, fixed to micropalaentological slides and counted. Although 100 tests can be generally considered as the minimum sample size for further statistical analyses (Fatela & Taborda, 2002), we decided to include here also the nine samples containing 50–100 tests, since all contained few species and in most of them (with only one exception) more than 50% of the total sample was represented by a single taxon (Patterson & Fishbein, 1989). This inclusion did not alter the results or the conclusions in any significant way.

The use of different diversity indices is relatively frequent in environmental studies of benthic foraminifera (Murray, 1973, 1991, 2006). Besides the simple species richness (number of species in each sample), four additional diversity indices, based on different characteristics (Magurran, 1989), were employed here to characterise the diversity of the benthic foraminiferal assemblages and to track environmental changes along the stratigraphic sequence. Two of the measures, the Fisher $\alpha$ index (Fisher et al., 1943) and the Margalef richness index (Margalef, 1958), depend on both the number of species and the number of tests in the sample. The Fisher $\alpha$ index has been widely used in foraminiferal ecological studies but its determination requires the use of specific software or graphical plots and tables (e.g. Hayek & Buzas, 1997). In contrast, the Margalef index has been less employed but can be very easily calculated. On the other side, the Pielou’s equitability or evenness $J'$ index (Pielou, 1969) specifically reflects the distribution of tests among taxa and depends on the Shannon-Wiener H’ heterogeneity index (Shannon & Weaver, 1949), which also takes into account the number of species and has been very frequently used too in foraminifer studied studies (the formula with the natural logarithm was employed here).

Three environmental groups of benthic foraminifera (lagoonal, transitional and strictly marine), together with their predominant species and their relative abundance variations were considered. In addition, the proportion of dextral and sinistral-coiling forms of the largely abundant species Ammonia beccarii was calculated for every sample with more than 50 specimens of this species in order to check any eventual link to environmental changes (Longinelli & Tongiorgi, 1964).

Available radiocarbon ($^{14}$C) datings published previously by García Blázquez et al. (2008) yielded Upper Pleistocene ages for three levels at 28.70 m, 18.50 m and 16.50 m core depth and Holocene ages for five additional levels between 13.90 m and 3 m core depth (Fig. 2).
estimated age throughout the core. Two selected diversity indices: Margalef richness and Shannon-Wiener diversity, with similar numerical ranges, have been employed.

3. RESULTS

The core reached a total depth of 60 m, but only the upper 50 m containing the Quaternary materials are shown in Figure 2. Its base consists of Miocene white marly limestones with root remains and bioturbation. Unconformably on these materials, the first Quaternary sediments can be found at 49 m depth. Along the core, successive environments with more or less marine influence are defined by their benthic foraminiferal assemblages and facies. Materials mostly consist of lagoonal silts and clays intercalated with increasingly predominant washover and eolian sand layers containing molluscs, ostracods and benthic foraminifers. A more detailed description is given in García-Blázquez et al. (2008).

A total of 205 samples, recovered from the upper 49 m of Quaternary sediments have been studied. The whole number of picked foraminiferal tests is 178066, that is, an average of 862 specimens per sample (Supplementary Data, Table 1: http://sepalontology.es/informacion-suplementaria/) which covers the occurrence of even the least abundant species in the different environments represented along the borehole. A total of 127 taxa have been identified. Five of them correspond to lagoonal species strongly related to restricted, continentally influenced environments; 15 species are associated with transitional zones with marine influence and the rest are clearly marine taxa (Murray, 1973, 1991).

In the first group the species are Ammonia tepida (Cushman, 1926), Cribrolphidium excavatum (Terquem, 1875), Deuterammina dublinensis Brönnimann & Whittaker, 1983, Haynesina germanica (Ehrenberg, 1840), Trichohyalus aguayoi (Bermúdez, 1935). The most abundant species in this group is C. excavatum (Fig. 3), with a relative abundance that averages 14.69%, followed by H. germanica with an average relative abundance of 10.71%. Both species show a wide range of variation (0.25–63.27% for C. excavatum and 0–49.15% for H. germanica), particularly in the lower half of the series, where they reach their maximum frequencies (C. excavatum at 37.30 m and H. germanica at 36.10 m). The rest of species are very rare although they are clear indicators of marginal and restricted environments.

The species in the transitional group are as follows: Ammonia beccarii (Linné, 1758), Aubignyna perlucida (Heron-Allen & Earland, 1913), Bolivina earlandi Parr, 1950, Bolivina striatula (Cushman, 1922), Bulimina elegantissima (d’Orbigny, 1839), Cornuspira involvens (Reuss, 1850), Cribrolphidium incertum Williamson, 1858, Elphidium clavatum Cushman, 1930, Elphidium granosum (d’Orbigny, 1826), Elphidium williamsoni Haynes, 1973, Haynesina depressula (Walker & Jacob, 1798), Haynesina sp., Quinqueloculina seminula (Linné, 1758), Pseudolachlanella eburnea (d’Orbigny, 1839), and Triloculina oblonga (Montagu, 1803). Among this group, the most abundant species is by far A. beccarii (Fig. 4), with an average relative abundance of 62.82% and an extremely wide variation range: 0.58–98.38%, particularly in the first 15 m of the core, whereas it becomes more constant in the central portion of the series (e.g. at 33.30 and 18.70 m). It occurs in large numbers both in the most marine influenced levels (e.g. at 46.80 m, 37.20 m or 36.90 m) and in the transitional ones (e.g. at 47.70 m, 40.70 m or 28.10 m). The rest of species are much less represented. Cribrolphidium incertum (average relative abundance: 0.93%) is only abundant in the central portion of the core where it reaches 32.97% at 28.10 m. Haynesina depressula (averaging 0.48%) attains maximum values at specific levels in the first half of the series
The rest of species, listed in Table 1 (Supplementary Data), are strictly marine. Many are miliolid taxa, among which the most abundant species are *Miliolinella subrotunda* (Montagu, 1803), which averages 0.86% and varies between 0 and 33.69%, *Pseudotriloculina rotunda* (Schlumberger, 1893), with a relative abundance ranging from 0 to 32.54% and averaging 1.20%, and *Triloculina schreiberiana* d’Orbigny, 1839 (average: 0.63%; range: 0–24.16%). In addition the *Rosalina* group, which mainly includes *Rosalina globularis* d’Orbigny, 1826 and *Rosalina mediterranensis* d’Orbigny, 1826, together with *Rosalina anglica* (Cushman, 1915) and *Rosalina bradyi* (Cushman, 1915) and *Rosalina williamsoni* (Chapman & Parr, 1932), is dominant in both the base (where it attains 21%) and the uppermost levels of the core, which show a strong marine influence (Fig. 5). Nevertheless, due to its absence in many samples, the whole group shows a low average relative abundance (0.78%) throughout the core. These species live fixed to leaves of marine phanerogams or to littoral rocky substrates. Most milioids are also associated to marine phanerogam meadows (Murray, 1973, 1991, 2006). The tests of these benthic marine taxa were included in the sediments deposited in the Xeraco area when the shoreline was closer.

The percentage of these strictly marine species is represented in Figure 6. As it can be observed, in the lowermost part of the borehole from 48.10 to 32.50 m depth, the assemblages of benthic foraminifera are subject to strong oscillations, with peak abundances of marine taxa at 46.80 m (45.55%), 45.60 m (48.92%), 36.90 m (42.39%) and 33.50 m (41.75%), and their total absence at 45 m, 44.80 m, 44.60 m, 44.40 m, 43.90 m, 41.10–41.50 m or 39.90–40.30 m. Their average relative abundance throughout the whole interval is 8.82%.

From 32.30 m to 18.70 m depth there is a nearly total lack of marine species (the maximum relative abundance is 1.72% at 31.50 m), but at 18.50 m and above marine taxa significantly increase again and reach the maximal frequencies along the borehole (70.97% at 9.90 m and 73.11% at 6.30 m depth).

The undoubtedly dominant species in Xeraco, *A. beccarii*, is very common in transitional and littoral environments and its coiling direction has been interpreted as the consequence of environmental changes (Longinelli & Tongiorgi, 1964; see also Guillem & Usera, 2004).
In Xeraco, sinistral-coiled individuals of *A. beccarii* are dominant throughout the borehole (Fig. 7). Dextral coiled specimens represent 30–40% of the *A. beccarii* tests in most of the samples and rarely depart significantly from this range. They reach minimal percentages at 44.70 m, 36.90 m, 28.30 m (when they show the minimal value in the borehole: 18.93%) and 25.30 m depth, whereas they peak at 47.90 m, 37 m, 33.30 m, 31.10 m, 17.70 m, 15.90 m (maximum value: 48.43%) and 11.30 m. Coiling direction variations do not seem to be linked here to environmental changes, since maximal and minimal values occur both in strictly lagoonal and in transitional or marine influenced moments. Similar percentage values were found by Usera *et al.* (1990c) in a Quaternary core in the locality of Pego (Alicante, Spain).

Some of the foraminiferal species recovered in the Xeraco borehole are shown in Figures 8 and 9.

### 3.1. Diversity

The Fisher $\alpha$ and the Margalef richness indices show an almost identical pattern throughout the core and are then described together.

#### 3.1.1. Fisher $\alpha$ and Margalef richness

In the lower part of the core and up to 32.10 m depth, strong oscillations can be observed (Fig. 10), with peaks (e.g., at 46.80 m, Fisher: 5.24 and Margalef: 3.84 or at 45.60 m, Fisher: 7.47, Margalef: 5.35) alternating with rather low values (e.g. at 43.90 m, Fisher: 0.37, Margalef: 0.28). After an interval with relatively low diversity between 41.50 m (Fisher: 0.48, Margalef: 0.39) and 37.60 m (Fisher: 0.76, Margalef: 0.64), generally higher values are recorded (mainly between 36.90 m and 32.70 m; e.g., at 33.50 m, Fisher: 5.90, Margalef: 4.18 or at 32.70 m, Fisher: 5.96, Margalef: 4.17). Diversity then drops and remains low from 32.10 m to 18.70 m. Fisher $\alpha$ never exceeds 1.5, whereas Margalef richness is always below 1.25, with an average for this interval of respectively 0.71 and 0.58. Both indices attain their minimum values at 24.50 m (Fisher: 0.35, Margalef: 0.26). In the upper part of the series strong fluctuations occur together with an increasing trend and the maximum values of the core are attained at 6.30 m core depth (Margalef: 7.11 and Fisher: 16.6, not shown in Fig. 10).

![Figure 6. Relative abundance (percentage) variation of foraminiferal tests belonging to strictly marine taxa along the Xeraco borehole.](image)

![Figure 7. Percentage variation of dextral-coiled tests in *Ammonia beccarii* along the Xeraco borehole.](image)
Figure 8. a) Textularia pseudogramen Cushman & Parr, 1937. b) Siphonaperta aspera (d’Orbigny, 1826). c) Adelosina longirostris (d’Orbigny, 1846). d) Quinqueloculina lamarckiana d’Orbigny, 1839. e) Triloculina trigonula (Lamarck, 1804). f) Quinqueloculina agglutinans d’Orbigny, 1839. g) Quinqueloculina cf. reticulata Karrer, 1862. h) Miliolinella circularis (Bornemann, 1855). i) Adelosina colomi (Le Calvez & Le Calvez, 1958). j) Quinqueloculina berthelotiana d’Orbigny, 1839. k) Quinqueloculina seminula Linné, 1758. l) Ammonia beccarii (Linné, 1758).
Figure 9. a) *Globulina gibba* (Deshayes, 1830). b) *Ammonia tepida* (Cushman, 1926). c) *Lobatula lobatula* (Walker & Jacob, 1798). d) *Elphidium macellum* (Fichtel & Moll, 1798). e) *Rosalina globularis* d’Orbigny, 1826. f) *Planorbulina mediterranensis* d’Orbigny, 1826. g) *Bolivina striatula* Cushman, 1922. h) *Cribroelphidium excavatum* (Terquem, 1875). i) *Haynesina germanica* (Ehrenberg, 1840). j) *Aubignyna perlucida* (Heron-Allen & Earland, 1913).
3.1.2. Shannon-Wiener $H'$

This index (Fig. 11) also shows high values in the lower part of the core from 46.80 m (2.23) to 32.90 m (2.17), but strongly oscillates, with minimal values at 44.80 m (0.34), at 43.50 m (0.30) or at 38.10 m (0.38) and peaks at 45.50 m (2.52), 43.30 m (2.37) or 33.50 m (2.28). Above 32.10 m depth the index shows both lower values and less variation (from 0.09 at 23.50 m to 1.32 at 27.30 m) but then it rises again and reaches its maximum towards the top of the borehole (2.73 at 9.90 m or 3.17 at 6.30 m).

3.1.3. Pielou J' or Equitability

Since this index is based on the Shannon index it follows the same general trend (Fig. 11). Minimal values are recorded at 43.50 m (0.19) or at 23.50 m (0.08), where A. beccarii is overwhelmingly dominant and occurs with only two other species. On the contrary, maximal values are noticed in moments of strong marine influence, at 44.60 m (0.97) or in the uppermost samples, at 9.90 m (0.86) or 6.30 m (0.89).

3.1.4. Species richness

Finally, we can evaluate the variation of the species richness (number of species) along the core (Fig. 12). Considering that higher species diversity is associated to stable environments and that it decreases in more isolated and unstable conditions (Margalef, 1958; Murray, 1973), the figure shows the varying influence of the marine realm. The number of mesolittoral and infralittoral species increases and reaches peak values of 36 (at 45.60 m), 35 (at 6.30 m), 34 (at 10.90 m) or 29 species (at 33.50 m), when the shoreline was presumably closer to the Xeraco area. Contrasting with them, several samples along the borehole show only 3 species, the lowermost one at 48.35 m, and particularly between 32.10 m and 18.70 m. The minimal value throughout the core (2 species at 8.10 m) is not significant since only 2 tests were recovered.

3.2. Radiometric datings

García-Blazquez et al. (2008) and Usera et al. (2012) originally provided the radiocarbon datings ($^{14}$C) shown...
in Figure 2. The oldest ones at 28.70 m (> 46912 yr BP), 18.50 m (> 45240 yr BP) or at 16.50 m (> 44760 yr BP) correspond to the Upper Pleistocene whereas the one at 13.90 m depth (10500 ± 121 cal yr BP) and above (5714 ± 123 cal BP; 5320–5060 cal yr BP; 2240–2000 cal yr BP or 1870–1690 cal yr BP) are clearly Holocene in age.

Figure 12. Species richness (number of foraminiferal species) variation along the Xeraco borehole.

3.3. Age model

The age depth model (Fig. 13) strongly suggests there is a probable hiatus or nearly non-deposition period between 16.50 m (dated as 44760 yr BP) and 16.30 m core depth, where García-Blázquez et al. (2008) located the Bølling-Allerød (dated as ~13000–15000 yr BP, 14000 yr BP in this model), which would encompass the uppermost part of the MIS 3 and most of the MIS 2 stage, including the Last Glacial Maximum, LGM). The intervals below and above this hiatus show notably similar average sedimentation rates: ~1.03 m/kyr for the 43.50 m–16.50 m section and ~1.09 m/kyr for the 16.30 m–3 m interval.

Figure 13. Age depth model for the Xeraco core based on data obtained from García-Blázquez et al. (2008). Black points with error bars: calibrated radiocarbon; blue triangles: minimum non-calibrated ages; red squares: Bølling-Allerød and assumed limits MIS-5–MIS-4 and MIS-4–MIS-3.

3.4. Diversity versus estimated age

The temporal variation of diversity along the core is represented in Figure 14. The Margalef richness and the Shannon-Wiener diversity indices show a very similar pattern, with strong oscillations in the lower part of the series (enlarged in Fig. 14b). Remarkable peaks at (estimated) 78108 yr BP (Margalef: 3.84; Shannon: 2.23), 75523 yr BP (Margalef: 5.35; Shannon: 2.35) or 71431 yr BP (Margalef: 3.30; Shannon: 2.07) and 70569 yr BP (Margalef: 3.23; Shannon: 2.37) alternate with low values, e.g. at 77246 yr BP (Margalef: 0.59; Shannon: 1.01), 71862 yr BP (Margalef: 0.29; Shannon: 0.72) or 71000 yr BP (Margalef: 0.56; Shannon: 0.30). This is followed by a relatively long period between approximately 68000 yr BP and 59000 yr BP of low diversity (with the Margalef index only exceeding 1.5 in two samples). High values are recorded between 57000 yr BP and 51000 yr BP (a maximum is attained at 52746 yr BP, Margalef: 4.18; Shannon: 2.28) and afterwards again an interval of relatively low diversity around 46000–45000 yr BP (e.g. Margalef: 0.30; Shannon: 0.09 at 46057 BP). After a last increase in diversity (Margalef: 2.59; Shannon: 1.41 at 45240 yr BP), the supposed non-deposition period is displayed as a data gap between 44000 and 14000 yr BP. Above this, the uppermost part of the Pleistocene and the Holocene record is visibly fragmentary, but reflects a rapid increase of diversity around 6000–5000 yr BP.
4. DISCUSSION

The identified species along the core seem to point to two moments of maximal marine influence, determined by the abundance of the strictly marine species group together with a high number of species at both the lower and the uppermost (above 18.50 m) parts of the sequence. Nevertheless, from 48.35 m to 32.10 m, these episodes of close proximity of the shoreline are frequently punctuated by intervals in which there is a total lack of open marine species (e.g. at 45 m, 44.20 m or at 40.50-39.10 m), possibly representing short episodes of a lagoonal environment with only a slight drainage connection to the sea. The assemblage of species indicative of transitional environments is well defined in the middle of the borehole, where *A. beccarii, H. germanica* or *C. incertum* are more abundant. These levels with scarce occurrence of marine species might be interpreted as a phase with a more distant shoreline and with the brackish lagoon still somewhat influenced by the sea, as it is indicated by the presence of *A. perlucida* and *B. elegantissima*.

The diversity and richness indices seem to confirm this interpretation since they reach their maximal values at the base and the uppermost part of the sequence and show high variability in the lower part up to 32.10 m depth, which supports the environmental instability in this Upper Pleistocene interval. The upper levels up to 18.90 m depth, with low diversity and equitability, probably reflect a more stable lagoon environment still subject to restricted conditions that favour the occurrence of a reduced number of species very well adapted to annual climatic fluctuations in the basin. Both richness and diversity increase again towards the top of the series with a new phase of strong marine influence during which the lagoon finally dries off and is replaced by eolian dunes and beach sediments.

4.1. Diversity versus estimated age interpretation

It should be kept in mind that, except for the radiocarbon ages in the upper part of the core, the rest of age assignments are only estimations. Due to the uncertainties of the age depth model, any inference based on it must be taken with caution.

Nevertheless, it can be deduced that the strong diversity changes of the lower part of the core reflect millennial scale climatic oscillations (probably Dansgaard-Oeschger stadials and interstadials) with cold periods, during which the Xeraco lagoon was isolated from the sea, alternating with warm periods, in which the lagoon was closer to the

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Figure 14. a) Margalef richness and Shannon-Wiener H’ diversity variation versus estimated age along the whole Xeraco core. b) Enlargement of the previous plot showing the lower part of the core.
coastline and more prone to marine influence, via washover events that would transport marine foraminiferal tests, or even temporal breaks of the bar, thus transforming the lagoon in an enclosed bay. Several abrupt cooling and warming changes have been recorded in the interval 80000–70000 yr BP (during the MIS-5a substage) in the western Mediterranean, such as stadials AS-21 and AS-20 or interstadials AI-20 and AI-19 (Martrat et al., 2004), which might correspond to the wide oscillations observed in the lower part of the core. The prolonged interval of low diversity that follows, between the estimated ages 68000 yr BP and 59000 yr BP, approximately matches the relatively cold MIS-4, in which the Heinrich 6 (HE-6) event has been shown to have had a discernible effect on both planktonic and benthic foraminiferal assemblages in the Alborán Sea (Pérez-Folgado et al., 2003, 2004; Pérez-Martin et al., 2006). The subsequent interval of prolonged high diversity between the estimated ages 57000 yr BP and 51000 yr BP indicates a stable relatively warm period that possibly corresponds to the Interstadials AI-14 and AI-13, as recorded in several Mediterranean studies (e.g. Martrat et al., 2004; Sprovieri et al., 2012; Incarbona et al., 2013). The next low diversity interval represented by a relatively long core section (from 32.10 m to 18.70 m depth; Figs 10-12) suggests a distant shoreline, between approximately 50000 yr BP and 45000 yr BP, and particularly around 46000 yr BP, which is coincident with the Heinrich 5 (HE-5) cold spell, whose effects have been recorded in the western Mediterranean by several authors (Pérez-Folgado et al., 2003; Martrat et al., 2004; Sierro et al., 2003; Frigola et al., 2008).

The Holocene sequence is incomplete (Fig. 14) but the diversity increase shows the growing marine influence associated to the development of the Flandrian transgression around 6000-5000 yr BP until the establishment of beach facies and eolian dunes in the uppermost part of the series.

4.2. Comparison with other areas

Several studies have treated the Quaternary evolution of foraminifera from coastal restricted environments in the western Mediterranean area, but many are limited to the Holocene. In the Iberian Peninsula, this is the case of the Ebros delta (Cearreta et al., 2016), the Vilanova i La Geltrú peat bog (Calzada, 1970), the Peníscola marsh (Usera et al., 2006), the Torreblanca lagoon (Colom, 1959; Collado & Robles, 1983; Usera et al., 1996a), the Albufera of València (Robles et al., 1985; Carmona et al., 2016) or the Albufereta of Alacant (Blázquez & Ferrer, 2003; Ferrer et al., 2005). Although Upper Pleistocene sediments have been preserved in the Albufera of Alcúdia (Mallorca), the studied foraminiferal assemblages are Holocene in age (Viñals & Mateu, 1999).

Other authors have reported the occurrence of Upper Pleistocene coastal lagoon environments with foraminiferal assemblages. In a 90 m core drilled in Tuscany (Italy) Carboni et al. (2010) reported sediments with foraminiferal assemblages typical of a restricted lagoon (dominated by *Ammonia parkinsoniana* (d’Orbigny, 1839) together *Cribroelphidium poeyanum* (d’Orbigny, 1839), *H. depressula* *H. germanica* tentatively ascribed by the authors to a slight sea-level rise during MIS-3, which perhaps might be related to the interval of high diversity recorded in Xeraco between 36.90 m and 32.70 m depth (Figs 10-12) and estimated as 57-51 kyr (Fig. 14).

In the Pego Quaternary lagoon (about 20 km SE from Xeraco) four different cores have been studied (Dupré et al., 1997; Torres et al., 2014). Two transgressive episodes, tentatively dated as MIS-5, have been recognized, with marine sediments (including foraminifera tests) transported into the lagoon, alternating with phases characterised by an enclosed lagoon environment with freshwater or brackish conditions, in which case foraminifers like *A. beccarii*, *H. germanica* or *C. excavatum* are predominant. After events that would transport marine foraminiferal tests, the establishment of a peat bog, dated as early Holocene, the Flandrian transgression (around 5000 yr BP) is also recorded in Pego. The lack of more chronologic precisions for the Pleistocene part of the series precludes further comparisons with the Xeraco core.

In the Xàbia marsh (located about 40 km SE from Xeraco) up to five cores were studied by Fumanal et al. (1993), some of which reach the Middle Pleistocene (MIS-9), and show alternating littoral and lagoonal facies with their characteristic foraminiferal fauna together with continental facies. The Upper Pleistocene transgressive phase (MIS-5) is recorded and characterised by marine stenoaline foraminifera (including planktonic forms, not recorded in Xeraco), after which stages MIS-4, MIS-3 and MIS-2 are poorly differentiated and are followed by Holocene lagoonal deposits with brackish foraminifera and increasing marine influence (see also Viñals et al., 1993; Garcia-Forner, 1997).

Middle and Upper Pleistocene transgressive and regressive movements of the shoreline are also preserved in the ancient coastal lagoon of Elx (up to MIS-12). Blázquez (2005) and Blázquez & Usera (2010) recorded different MIS-5 substages in three studied cores (starting with a brackish lagoon phase corresponding to MIS-5d in core Pinet). If the age ascriptions of the lower part of the Xeraco core were correct, its observed wide fluctuations of diversity, indicating the close proximity of the shoreline, would correspond to the shoreface bed deposited in the Salinas core (Units IV and V) in Elx and attributed by Blázquez & Usera (2010) to substage MIS-5a. Above this transgressive episode, no sedimentary record of
stages MIS-4, 3 or 2 has been detected in Elx. A brackish lagoon subject to an increasing marine influence and dated as Holocene is finally replaced, as in Xeraco, by beach (shoreface/foreshore) deposits.

5. CONCLUSIONS

The study of the foraminiferal assemblages and the autoecological characteristics of some of the occurring species suggest that this area of the Mediterranean coast was first subject to warm climate conditions, with high sea level and a shoreline close to the borehole location, and that the conditions of a lagoon subjected to a strong marine influence and frequent washover events, with some millennial scale environmental fluctuations, persisted up to the 32.10 m depth. Afterwards the assemblage data show a sudden drop in diversity and a nearly total absence of littoral species. These conditions continued up to the 18.70 m depth and are interpreted as a lagoon environment in colder/glacial climatic conditions with low sea level and a more distant shoreline. Finally the diversity increase and the reappearance of littoral foraminifera in the upper part of the sequence indicate a new sea level rise which, according to the radiocarbon datings, can be attributed to the Flandrian transgression, already in a new interglacial warm phase, in which the Xeraco lagoon is finally replaced by beach deposits.

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[All historical references concerning original descriptions of the different taxa cited in the text or in Table 1 have been included as Supplementary Data in a separate list: (http://sepaleontologia.es/informacion-suplementaria/)].

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