

Abstract

Landwards of a MIS5 bar, a borehole core (SRA) was analyzed to establish the relationship between the lagoonal record and the raised beach deposits in the surroundings of the Antas river mouth and to reconstruct the Pleistocene palaeoenvironmental evolution of the southern Mediterranean coast of the Iberian Peninsula. 63 samples were recovered for amino acid racemization dating, 86 samples for sedimentological and paleontological determination, 37 samples for pollen identification and 54 for biomarker analysis. AAR revealed that the borehole record contains MIS11, MIS6 and MIS5 deposits, the latter extensively represented. During the end of MIS6 and MIS5, a sand barrier developed and created a shallow lagoon with alternating terrestrial inputs this process being common in other Mediterranean realms. Litho- and biofacies allowed the identification of distinct paleoenvironments through time, with the presence of a lagoonal environment alternating with alluvial fan progradation. Biomarkers indicated constant input from terrestrial plants, together with variable development of aquatic macrophytes. The palynological content allowed the reconstruction of the paleoclimatological conditions during MIS6 and 5, with evidence of seven scenarios characterized by alternating arid and relatively humid conditions.

1 Introduction

In many areas of the Mediterranean coast – Spain, Morocco, Tunisia – the current morphology appears as a series of cliffs where Alpine tectonics-linked parallel ranges reach the sea alternating with lagoons and wetlands. Some of the Quaternary coastal deposits of the Spanish Mediterranean realm have been studied, mostly focusing on raised marine deposits and addressed faunal remains and radiometric ages (Goy et al., 1986; Hillaire-Marcel et al., 1986; Causse et al., 1993; Zazo, 1999; Zazo et al., 2003; Dabrio et al., 2011) and amino acid chronology (Hearty, 1986, 1987; Hearty et al., 1986; Torres et al., 2000a, 2010a; Ortiz et al., 2004a). Unfortunately, raised marine

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(Fig. 1), which can be age-correlated in spite of they were subjected to strong neotectonic processes (Torres et al., 2000a, 2006, 2010a, 2013).

In fact, in the immediate neighborhood of the SRA, a stripe representing an ancient barrier system (MIS5) and running parallel to the present day coastline is clearly visible in Fig. 1. This stripe consist of gravel and coarse sands that are strongly carbonate-cemented, and should protect the lagoonal area of SRA during MIS 5. These materials appear as foreshore dipping beds some decimeters thick. At some points, a number of beds dip backshore. Scattered *Glycymeris* sp. shells are visible, as well as some rare *Persististrombus latus* (sin. *Strombus bubonius*) representatives. Today the ancient bar is occupied by a tourist center, but a section is still visible on each side of the road entering the resort.

2.2 Biological setting of the area around the Antas river

The area is placed in the “almeriense” sector of the “murciano-almeriense” chorological province (Rivas Martinez, 1988). The native plant species are resistant to drought and aridity, which occur during the summer and also in some winter-autumn months. Thus plant species have xerophytic characteristics, thus largely exposing soil and bed rock (Ferré Bueno, 1979).

The natural vegetation in the Almanzora valley falls in the climatic dominion of *Quercus ilicis* and Oleo-Ceratonion. The former holds *Quercus ilex*, *Q. coccifera*, *Juniperus phoenicea*, *Dhapne gnidium* and *Retama sphaenocarpa*, although *Q. ilex* is scarce. In many areas, *Pinus halepensis* has disappeared or remains as scattered individuals, and in wetter zones isolated individuals of *P. pinaster* are found. In general, only sub-serial bushes belonging to the *Q. ilex* dominion are present: *Rosamarinus officinalis*, *Cistus albidus*, *Rhamnus lycioides*, *Genista umbelata*, *Ulex* sp., and *R. spaenocarpa*. In more severely degraded areas, only species of *Thymus*, *Artesania glutinosa* and *Stipa tenacissima* are detected.

In the western-most sierras, a variety of the *Q. ilex* grade was observed: the sub-grade *Q. lusitanica-Acer monspesulamun granatensis* attesting a wetter and colder

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For the age calculation of SRA and ALF samples, aspartic acid and glutamic acid were chosen because they account for over ca. 50 % of the amino acid content in most ostracod valves (Kaufman, 2000; Bright and Kaufman, 2011). The numerical age of each bed was determined by introducing the aspartic acid and glutamic acid D/L values obtained in *C. torosa* individuals from the Alfaix terraces and the SRA borehole into the age calculation algorithms described by Ortiz et al. (2004b). The age of a single bed is the average of the numerical dates obtained for each amino acid D/L value measured in ostracods from that horizon. The age uncertainty is the standard deviation of all the numerical ages calculated from the amino acid D/L values of each level.

We used the age calculation algorithms obtained in ostracods collected in central and southern Spain by Ortiz et al. (2004b) for the dating these deposits because a similar thermal history can be inferred for these areas, as they are all located in the Mediterranean climatic zone of the Iberian Peninsula, with a similar mean annual temperature. Likewise, the age calculation algorithms were established for the ostracod species analyzed here (*C. torosa* and *H. reptans*), which show similar racemization rates for Middle and Lower Pleistocene localities (Ortiz et al., 2013). Thus DL ratios are directly comparable without any conversion factor.

3.2 Sedimentology

A total of 86 samples at 20–30 cm intervals were picked from the borehole core. These samples were analyzed for sedimentological, micropaleontological (especially, ostracoda and foraminifera) and palynological studies. The sampling interval was chosen on the basis of major changes in sediment color, grain size, and the presence of chemicals (gypsum crystals), and macrofossils.

Totally dried samples were gently crushed, soaked, passed through a 62 μm mesh sieve, and analyzed under microscope. Some samples were resin-embedded to study the sand fraction mineralogy in a thin section.

3.3 Palynology

A total of 37 samples were selected for palynological analysis. Pollen was extracted following standard protocols (Couteaux, 1977; Faegri et al., 1989; Girard and Renault-Miskovsky, 1969; Moore et al., 1991). The residuals were pollen-enriched through flotation in Thoulet's dense liquor (Goeury and Beaulieu, 1979) and later placed in Eppendorf tubes and preserved in glycerin.

Pollen grains were read and counted following Cambon (1981): 42 rows in each sample. The number of pollen grains in the samples oscillated between 110 and 716, although in some samples the number was so low (10–20) that they appeared in the pollen diagram as “presences”. In these cases the number of identifiable taxa was greatly reduced (5 or lower).

Statistical analysis was done using the package OLEA (M. Arroyo unpublished), which is based on philosophy of the TILIA[®] and TILIA-GRAPH[®] (Grimm, 1987, 1992) packages.

In the samples with a statistically significant number of pollen grains, we built AP/NAP logs and analyzed the rarefaction (Birks and Lyne, 1992) to determine vegetation diversity.

In order to facilitate the interpretation of the data, we built a synthetic pollinic diagram, which included the most significant taxa (*Pinus*, *Juniperus*) in the spectrum, as well as the largest taxa clusters (Mesophilous, Mediterranean, Xeric, Steppic).

Finally, all the sampled were subjected to a PCA (Principal Component Analysis) using the Biplot application from Excel-Microsoft.

3.4 Biomarker analyses

A total of 54 samples were collected for biomarker analysis, coarse-grained sediments being discarded. 5 g of sediment per sample was dried at 50 °C for 24 h and then ground with a mortar and pestle. Biomarkers were extracted in an accelerated solvent extractor (Dionex ASE 200) using dichloromethane/methanol 2 : 1 at 1500 psi and 175 °C (after

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explained as recent contamination associated with ground-water circulation in near-shore aquitards in arid areas under a marine intrusion regime. In fact, the analysis of total carbon in bulk sediments is inappropriate because significant differences between the radiocarbon content of the bulk sediment and true age can arise (Olsson, 1986; MacDonald et al., 1987; Lowe et al., 1988; Brown et al., 1989; Torres et al., 2010b). Pantaleón-Cano et al. (2003) dated a sample at 0.5 m of depth at 1390 ± 100 BP; however, although appearing reliable, it seems somewhat too young as implies that the deep river incision should occurred in historical times.

The possibility of ostracod reworking from older deposits is unlikely because the shells were not eroded, in some cases still being articulated. Furthermore, in some samples ostracods were found together with juvenile thin-shelled *Cerastoderma glaucum* representatives that were too fragile to be transported.

The AAR dating of the SRA record allowed us to determine that MIS11 (a single dating was obtained), MIS6 pro-parte, and MIS5 are represented. As the ages showed an uncertainty of about 25 %, the MIS5 sub-episode boundaries were refined on the basis of the palynological analysis and palaeoenvironmental information derived from sedimentology. This implies that MIS6-MIS5e was displaced 80 cm upwards, and we were unable to differentiate between MIS5b and MIS5a.

4.2 Sedimentology and fossil content

The SRA core comprised mainly material of detrital origin, predominantly clay, silt, and marls (Fig. 2). Six lithologies were observed:

- Gm: Gravel $C = 26\text{--}60$ mm (poorly recovered from the core), with clasts of quartz, dolostone, limestone and metamorphic rocks, with variable amounts of heterometric quartz sand, abundant mica flakes, and red-brown lutites (matrix?).
- Gf: fine-grained gravel ($C = 5$ mm) made of quartz and metamorphic rocks in a lutitic matrix with limonitized botroids (derived from pyrite), in which reworked marine microfossils were common.

chemistry of sediment records (Meyers and Ishiwatari 1993; Meyers 1997, 2003). However, the interpretation of organic geochemical signals is not always straightforward because in most cases organic matter is a mixture of components from many sources and with variable degrees of preservation.

n-Alkanes are among the biomarkers most commonly used. These compounds have the advantage that they are less susceptible to microbial degradation than other types of organic matter because they lack the functional groups that confer chemical reactivity, and also that they have low water solubility (Prahl and Carpenter, 1984; Meyers et al., 1995). The *n*-alkanes present in sediments reflect mainly the contribution from algae, bacteria, aquatic macrophytes, and land plants. Several *n*-alkane ratios can be used for paleoenvironmental reconstruction purposes.

The logs of the various indexes related to the *n*-alkane content, namely the *n*-alkane predominant chain, the average chain length (ACL), the aquatic macrophyte proxy (Paq), the Terrigenous/Aquatic Ratio of hydrocarbons (TAR_{HC}), and the relative percentages of C₂₇, C₂₉ and C₃₁ with respect to the sum C₂₇+ C₂₉+ C₃₁, are shown in Fig. 3.

The SRA cores samples showed *n*-alkanes with an odd-over-even carbon number predominance, with a chain length distribution ranging mainly from C₁₇ to C₃₁. We detected an overwhelming domination of C₃₁ *n*-alkane (44 over 54 samples), but in 10 of these samples the C₂₁ homolog was predominant.

The ACL (Poynter, 1989), calculated as $[(C_i \times i + C_{i+1} \times (i+1) + C_{i+2} \times (i+2) \dots + C_n \times n)] / (\sum C_i + C_{i+2} + \dots + C_n)$, with $i = 13$, $n = 33$, showed few oscillations (Fig. 3), with values varying between 21.9 (15.9 cm) and 28.0 (640 cm).

The Paq index, calculated as the $(C_{23} + C_{25}) / (C_{23} + C_{25} + C_{29} + C_{31})$ ratio (Ficken et al., 2000), allows discrimination between the relative contribution of aquatic macrophytes and terrestrial plants. Paq values registered several oscillations, but with a low range, falling between 0.05 (150 and 240 cm) and 0.25 (790 cm).

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Silliman et al. (1996) reported that the terrigenous / aquatic ratio, calculated as $(C_{31} + C_{29} + C_{27}) / (C_{15} + C_{17} + C_{19})$, distinguishes between terrestrial plant and algal inputs. The TAR_{HC} index varied between 39.9 (40 cm) and 1.4 (1660 cm).

The *n*-alkane composition as relative percentages of C_{27} , C_{29} and C_{31} with respect to the sum $C_{27} + C_{29} + C_{31}$ showed some oscillations (Fig. 3), but in all cases, the relative percentage of C_{31} accounted for more than 50 % of the *n*-alkanes present, with a maximum at 760 cm (70 %).

4.4 Palynology

Pollen was not present in the lowermost (2850–1830 cm) or uppermost (280–0 cm) part of the SRA record.

In the samples from the remaining record, we identified 39 taxa, of which 11 were trees (*Pinus*, *Juniperus*, *Corylus*, *Betula*, *Juglans*, *Quercus* deciduous, *Olea*, *Quercus* evergreen *Alnus*, *Fraxinus* and *Ulmus*), 6 bushes (*Calluna*, Ericaceae, *Myrtus*, Cistaceae, Pistacea, and Rosaceae), and 22 grasses (Asteraceae liguliflorae, Asteraceae tubuliflorae, Poaceae, *Sanguisorba*, *Artemisia*, Chenopodiaceae, *Ephedra*, *Plantago*, Polygonaceae, Rubiaceae, *Rumex*, *Urtica*, Apiaceae, Brassicaceae, Caryophyllaceae, Fabaceae, Boraginaceae, Scrophulariaceae, *Asphodellus*, *Potentilla*, Rhamnaceae and Violaceae). Also, we detected two aquatic species (Cyperaceae and *Alisma*). Monolete and trilete spores and 15 non-pollinic microfossils (NPMs) from various affinities were also found: 3b type (*Pleospora*) of dry character, 55A (*Sordaria*) 113 (*Sporomiella*), 368 (*Podospora*) and *Riccia* of coprophilic affinity, as well as the mesoeutrophic types 181 and 731 and the eutrophic 170 type (*Rivularia*). We also detected deforestation-linked type 207 (*Glomus* cf. *fasciolatum*) and soil erosion-linked *Pseudoeschizaeae circula*.

The pollen assemblage reveals a Mediterranean climatic scenario with fluctuations of moisture and, in some intervals, mainly in the upper half of the section, the presence of Nitrophilous taxa.

The pollen diagram (Fig. 4) reveals that tree cover was denser in the lower part of the record and more open at the top. Similarly, the shrubby stratum did not contribute greatly to the vegetation structure nor did NPM, aquatic taxa, or most of the elements identified in the trees or grasses groups.

Given the above observations, we built a synthetic diagram taking in consideration various taxa (*Pinus*, *Juniperus*) and associations of trees (Mesophilous, Mediterranean), grasses (Xeric, Steppic and Nitrophilous), NPM type 3b, and Coprophilous taxa (Fig. 5).

5 Discussion

5.1 Sedimentary environments

We differentiated the following five lithological units in the SRA borehole on the basis of sedimentological characteristics and microfossil content (Fig. 2): Unit I (29.7–28.2 m: Pliocene yellow-grey marls and silts; Unit II (28.2–24.2 m loosely cemented gravels interbedded with sands of Pleistocene age); Unit III (24.2–18.2 m loosely cemented Pleistocene gravels and sands); Unit IV (18.2–3.0 m variegated lutites with bioclasts and gypsum); and Unit V (3.0–0.0 m laminated fine-grained sands). Because of some minor variations in lithology and biological content, Unit IV was sub-divided in ten (a–j) sub-episodes.

Although Pantaleón-Cano et al. (2003) identified these units in their borehole record, they observed peat deposits between 22.50 and ca. 20.60 m that were not present in the SRA core. Those authors did not provide the coordinates of their borehole, however, this discrepancy allows us to conclude that SRA and Pantaleón-Cano et al. (2003) boreholes were drilled in different areas of the paleo-lagoon.

Each sedimentary unit corresponded to a certain environment (Table 3, Fig. 2). With the exception of marine conditions that occurred during the Pliocene, the basin showed evidence of alluvial fan progradation alternating with lagoonal and palustrine episodes.

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5.2 Organic geochemistry (biomarkers)

A short description of the *n*-alkane indices used is given in Table 4. *n*-Alkane content provided reliable information on the local distribution of plants, thus completing the more regional data provided by pollen analysis.

The predominance of the C_{31} *n*-alkane in most of the samples indicated the main input of grasses along the whole record (Fig. 3). Levels maximizing in the C_{21} homolog can be linked to aquatic macrophyte dominance, usually associated with the presence of aquatic fauna remains (ostracods and/or pelecypoda), as occurred in sub-units IV-a (1690), IV-c (1390, 1360), IV-e (820, 790) and IV-i (550, 520). However, Unit II (2840), Unit III (2050) and sub-unit IV-b did not hold faunal remains (Fig. 2). In all cases ephemeral shallow water masses appear to have developed.

The variations in Average Chain Length (ACL) values were small, although some clarifications can be made. In this regard, the ACL values oscillated between 21.9 and 28.0, implying a reduced contribution of low molecular weight *n*-alkanes, usually linked to algae and bacteria. However, the ACL values were slightly lower in Units II, III and subunits IV-a and IV-b, indicating a growth of aquatic taxa.

The dominance of organic matter derived from terrestrial plants over algae input can also be interpreted from the TAR_{HC} index, which showed values higher than 1.5 in all cases, being higher than 3.0 in 80% of the samples.

The Paq values were originally defined and used to determine the relative contribution of macrophyte taxa and terrestrial plants to some African lakes (Ficken et al., 2000). According to Ficken et al. (2000), Paq values under 0.1 are linked to a dominant contribution from land plants, while values between 0.1 and 0.4 reflect significant inputs from emergent macrophytes. Paq values higher than 0.4 are typical in sediments with a major *n*-alkane input of submerged/floating macrophytes. In the SRA record, most of the samples showed Paq values between 0.05 and 0.25, indicating a major input of terrestrial plants and emergent macrophytes. The absence of submerged macrophyte dominance indicates that the water mass was limited in time and depth.

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In Phase I (1850–1700 cm), the pollen content was low, although *Pinus* and Poaceae were present. Towards the top, the presence of Mediterranean, Xeric, and Nitrophilous taxa increased and then decreased dramatically at the end of this phase, favoring the presence of *Juniperus*. We interpret these observations as an initial warm and dry climate, turning to a cold one, as attested by the development of *Juniperus*.

Phase II (1700–1550 cm) began with a sharp decrease in the presence of *Juniperus*, which favored the growth of Mesophilous taxa and later Mediterranean species, the frequencies of Xeric and Steppic taxa being low. Nitrophilous taxa increased and scattered samples of Aquatic taxa appeared. We interpret these climatic conditions as temperate-warm and humid.

In Phase III (1550–1330 cm), there was a clear dominance of *Pinus*, followed by Xeric and Steppic taxa and, to a lesser extent, Nitrophilous and NPM-3B taxa. We interpret these data as indicative of a dry and colder period, as attested by the low development of Mesophilous and Mediterranean taxa.

In Phase IV (1330–1120 cm), there was a clear expansion of Mediterranean taxa, with some oscillations, together with an increasing presence of Xeric and Steppic taxa, as well as *Juniperus* towards the top, marking a temperature increase under dry conditions.

In Phase V (1130–840 cm), Mediterranean and Mesophilous taxa clearly dominated, though with some oscillations. There was a clear decline in the presence of *Pinus*, and *Juniperus* disappeared. At the beginning of this phase the presence of Aquatic and Nitrophilous taxa increased and in the upper part they were replaced by Xeric taxa and NPM-3b, ending with a short period in which the pollen content was practically absent. We interpret these observations as indicating a warm period that was initially relatively humid and that evolved towards dry conditions; this phase can be correlated with MIS5e.

Phase VI (840–700 cm) started with an increase in *Juniperus* and Steppic taxa, which were later substituted by Mediterranean and Mesophilous ones. The presence

of Aquatic and Nitrophilous taxa showed a clear increase. These data indicate that temperate and humid conditions dominated during this phase.

In Phase VII (700–290 cm), the presence of *Pinus* and *Juniperus* fluctuated and alternated, and both genera tended to diminish towards the top of the record. There was a significant increase in Steppic and Xeric taxa. In spite of this general trend, Mediterranean taxa showed a progressive growth accompanied by a slight development of Mesophilous ones. These data reflect warm and dry conditions, with some periods with extremely water shortage.

To interpret the pollen assemblages, we performed a principal components analysis (Fig. 6) on the data used for the synthetic diagram. Component 2 (explaining 56.55 % of the variability) grouped samples on the basis of the development of the local Mesophilous and Mediterranean taxa (positive values) versus open landscapes and regional *Pinus* forest (negative values). In contrast, Component 3 (explaining 72.15 % of the variability) clustered taxa marking warmer conditions (positive values) and those indicating temperate or cool conditions (negative values).

According to these results, cold-to-temperate and humid conditions developed during MIS6, coinciding with the paleoclimatological interpretation of the Fuentillejo core (Ruiz Zapata et al., 2012), located 300 km north-west of the SRA borehole in the same climatic zone (Fig. 1), as well as with the growth of tufa deposits in the Iberian Peninsula (Torres et al., 2005; Ortiz et al., 2009). In contrast, warm and arid conditions occurred at the end of MIS6 (140–133 ka) both in the Fuentillejo (Ruiz Zapata et al., 2012) and SRA cores.

At the beginning of MIS5 climatic conditions were warmer but less dry, with the recovery of the Mediterranean forest, Mesophilous taxa and *Pinus* in Phase 5, which corresponds to MIS5e, and coinciding with pollen data of the Fuentillejo core (Ruiz Zapata et al., 2012). Fuentillejo maar lake developed under a continental-mediterranean climate.

During the rest of MIS5, the alternation of *Juniperus*, Xeric and Mesomediterranean taxa indicate short cold-dry episodes within this interglacial period.

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Thus the palynological data revealed that paleoenvironmental fluctuations were not as extreme as in other areas of the Mediterranean realm and Central Europe.

6 Conclusions

The abundance of the ostracod *C. torosa* in almost all the beds of the SRA core allowed AAR dating of the Middle-Upper Pleistocene record of the Vera Basin in the vicinity of Garrucha village (between the mouths of the rivers Antas and Aguas), with a sedimentary hiatus between MIS11 and MIS6 and 5 deposits. It is noticeable that MIS5 was extensively represented in the SRA core, covering almost 11.5 m (14.5–3.0 m). The accuracy of AAR has been tested through the coincidence of U/Th and AAR ages of the neighbor zone of Alfaix.

It was proved a clear relationship between the *P. latus* bearing MIS5 bar along the Garrucha coastline and the fine grained lagoon record of the same age that allowed the paleoenvironmental reconstruction. In fact, MIS5 coastal bars played a decisive role in the present-day coastline morphology allowing the permanence of fossil or still-active lagoons. There are other examples along the Spanish Mediterranean coast of active lagoons where the active bar results to be anchored on fossil bars (MIS5): Calblanque (Lillo, 1988) and Mar Menor in Murcia (Lillo, 1988; Colodrón et al., 1977), Elche (Blázquez, 2005), La Mata (Somoza et al., 1986; Zazo et al., 1993; Santisteban et al., 2004), and Pego (Torres et al., 2013) in Alicante. In any case it seems that a rich paleo-environmental record seems to be waiting for future research.

On the basis of the lithology and sedimentological characteristics, we distinguished six associations of lithofacies, corresponding to lagoon deposits and distal alluvial fan influence.

Organic geochemistry, namely *n*-alkane content, which is extensively used in paleoenvironmental reconstruction, proved highly suitable for the analysis of the SRA record. Our results reveal that there was a constant and considerable input of terres-

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trial plant-derived biomarkers to a shallow wetland, where emergent macrophytes were also present in varying amounts.

Sedimentation during MIS11 was partially represented by alluvial fan sediments (Unit II) that retracted towards its end (Unit III). Unfortunately, we were unable to interpret the paleoclimatic conditions because the record was devoid of pollen content.

The end of MIS6 was confidently dated and represented by playa-lake marginal deposits (subunit IV-a). Towards the end of MIS6, the alluvial system was reactivated, as reflected in fine gravel and sand deposition (subunit VI-b). Frequent root tubuli indicate aerial exposure. The pollen content analysis revealed that this stage showed milder climatic conditions (warm-temperate) than the beginning of MIS5e (cool), although varying from dry to humid.

MIS5 was confidently dated and presumably all the sub-stages were represented, although AAR ages showed some uncertainties. During MIS5 a series of three lagoon-alluvial couplets were identified through sedimentological characteristics and the micropaleontological content. The lagoonal episodes were marked by the presence of autochthonous foraminifers and brackish-water mollusks and ostracods, the presence of which indicates larger marine incursions during the MIS5e substage. Although these marine conditions never prevailed, as the lower presence of algae biomarkers suggest, the environment was characterized by intermittent sea-water entrance and emergent macrophyte colonization but with dominant periods of dryness in which interstitial gypsum lenses developed, the dried surface of which was grass-colonized, as attested by biomarker analysis.

Alternation of episodes of lagoonal and alluvial fan progradation were also observed in the Pleistocene record of the nearby Elche and Pego basins (Fig. 1; cf. Blázquez, 2005; Blázquez and Usera, 2010; Torres et al., 2013).

Our analysis of pollen content revealed that nuanced Mediterranean biomes were ever-present, showing a small degree of decoupling with sedimentary environments.

- MIS5e is well represented by lagoonal sediments with ostracods, foraminifera and brackish water pelecipoda (subunits IV c–e). According to the pollen record, cli-

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mate conditions changed from warm-dry to warm and more humid, ending with more temperate conditions.

- MIS5d can be correlated with the sedimentary subunit IV-f, marking some progradation of the alluvial systems, and temperate and humid conditions.
- MIS5c correlates with subunit IV-g in which lagoonal conditions prevailed. According to the pollen record, warm and dry conditions prevailed.
- MIS5b corresponds to subunit IV-h, which marks a new alluvial progradation with azoic sediments, although the pollen record indicated warm and dry climatic conditions.
- MIS5a (subunits IV-i and IV-j) began with the re-establishment of lagoonal conditions (ostracods, brackish water pelecipoda, Helicidae) under warm and dry conditions, although the end of this period was marked by alluvial fan progradation.

The paleogeographical reconstruction suggests that the present-day Antas river channel incised an ancient lagoon at the end of Upper Pleistocene after a sudden shift southwards.

Acknowledgements. Funding was obtained through the projects “Paleoclimatological revision of climate evolution in Western Mediterranean region” (European Union, CE-FI2W-CT91-0075) and “Paleohidrogeological Data Analysis and Model Testing” (European Union, FIKW-CT-2001-00129). The Biomolecular Stratigraphy Laboratory has been partially funded by ENRESA.

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Table 1. Amino acid racemization ratios measured in the ostracod shells from the different horizons of Alfaix fluvial terraces and SRA borehole.

Locality	Species	N	D/L Asp	D/L GLu	Age (ka)
AFX-1	<i>C. torosa</i>	3	0.426	0.177	194.7 ± 17.1
AFX-3	<i>C. torosa</i>	6	0.325 ± 0.008	0.202 ± 0.030	155.1 ± 68.7
AFX-4	<i>C. torosa</i>	7	0.402 ± 0.051	0.214 ± 0.058	205.6 ± 40.1
SRA2-300	<i>C. torosa</i>	2	0.268 ± 0.008	0.108 ± 0.009	72.3 ± 25.6
SRA2-870	<i>C. torosa</i>	3	0.353 ± 0.033	0.128 ± 0.049	117.9 ± 47.8
SRA2-900	<i>C. torosa</i>	3	0.363 ± 0.035	0.112 ± 0.037	156.5 ± 4.8
SRA2-1050	<i>C. torosa</i>	5	0.342 ± 0.044	0.137 ± 0.003	136.6 ± 45.3
SRA2-1200	<i>C. torosa</i>	7	0.344 ± 0.023	0.162 ± 0.024	139.5 ± 35.9
SRA2-1380	<i>C. torosa</i>	7	0.381 ± 0.011	0.179 ± 0.024	168.0 ± 28.6
SRA2-1500	<i>C. torosa</i>	14	0.349 ± 0.021	0.185 ± 0.032	156.7 ± 48.4
	<i>H. reptans</i>				
SRA2-1740	<i>C. torosa</i>	5	0.370 ± 0.018	0.133 ± 0.036	143.9 ± 58.0
SRA2-1830	<i>C. torosa</i>	1	0.541	0.393	421.8 ± 77.7
SRA2-2270	<i>C. torosa</i>	12	0.405 ± 0.018	0.275 ± 0.062	308.7 ± 42.3
SRA2-2540	<i>C. torosa</i>	4	0.514 ± 0.032	0.235 ± 0.035	289.4 ± 54.8

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Table 2. Lithology observed in thin sections of resin-impregnated samples previously water-sieved ($63\ \mu$).

Depth (cm)	Lithology
400	carbonates, tourmaline, clorite, zoisite, epidote, quartz.
900	zoisite, epidote, garnet, fragments of saussuritized rock, limestone, calcite, quartz (sometimes with undulating extinction), opaque minerals and muscovite.
1500	quartz, zoisite, micaschist, quartzite, muscovite, biotite, saussuritized plagioclase and opaque minerals (probably magnetite or pyrite)
1620	quartz, muscovite, biotite, clorite, limonitized rock fragments, tourmaline, zoisite, quartzite and sandstone.
1960	quartz, limestone, micaschist, plagioclases, garnet, muscovite, micaschist, opaque minerals, garnet, anfibol, anfibol and micaceous quartzite.
2050	quartz, limestone, micaschist, plagioclases, garnet, muscovite, micaschist, opaque minerals, garnet, anfibol, anfibol and micaceous quartzite.
2360	quartz, calcite, plagioclase, zoisite-epidote, garnet, muscovite, biotite, clorite, cloritized biotite, sericite, tourmaline and fragments of micaschist, quartzite and calcite.

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Table 3. Lithology, fossil content and paleoenvironmental interpretation of Units and Sub-Units identified in SRA borehole core (Au F: Autochthonous foraminifers; Al. F.: Allochthonous foraminifers).

Units	Interval (cm)	Sub-units	Lithology	Fossil content	Paleoenvironmental interpretation
V	300-0		brown fine-medium grained homometrical laminated sands (Fl), with charcoal, plant debris, seeds and pollen grains, some gypsum, gravel at the top	Al F, Au F at the top AF	Alluvial with marine input at its top
IV	1830–300	IVj 520–280	Brown-beige sandy/silty carbonate lutites Fm-Fs, angulous and homometrical quartz grains, mica flakes abundant	Al F, land snail, <i>C. torosa</i> rare	alluvial with small ponds
		IVi 580–520	Light gray carbonate lutites	Al F, rare Au F, brackish water pelecipoda, land snails, ostracods	lagoon
		IVh 640–580	Brown lutites with fine sand, lenticular gypsum	Al F	distal deposits (lobe) prograding alluvial lagoon
		IVg 700–640	Silty clay (Fm) with lenticular gypsum. Fe patches	Au F	distal deposits (lobe) prograding alluvial lagoon
		IVf 760–700	Interbeds of scarcely sandy lutites (Fs) and lutitic sands (Sf), lenticular gypsum from absent to very abundant, fine grained quartz and mica flakes.	Al F	distal deposits (lobe) prograding alluvial lagoon
		IVe 850–760	(Myc). White carbonate lutites, gypsum very abundant (tabular and lenticular), tubuli, quartz sand (rare).	Al F, Au F, ostracods	lagoon
		IVd 1240–850	white-light gray carbonate lutites (Myc; lutitic fraction accounts > 80%). Some tabular gypsum (1180 and 970), charcoal (1090), mica flakes, and limonitized tubuli (940). At the top medium-fine grained quartz sand interbeds (Sf).	Azoic, only a single level with brackish water ostracods	frequently dried-up lagoon with two alluvial incursions
		IVc 1420–1240	beige, brown and gray lutites-Fm (lutitic fraction accounts > 80%). Iron oxides, botroidal oxidized pyrite noduli.	Al F, Au F, brackish water pelecipoda, ostracods	lagoon-brackish waters
IVb 1540–1420	1830–1540	IVb 1540–1420	fine gravel and sand (Gfs), botroidal limonite rare, rhizotubules rare	Rare Al F	alluvial fan toe
		IVa 1830–1540	Green and grey marls-Myc facies (lutitic fraction accounts > 90%), angulous fine grained quartz and mica flakes, limonitized root casts. Tabular gypsum rare	Rare Al F, rare Au F (1740)	mud flat playa/margin playa lake
III	2360–1830		Lutitic sands. Fs (lutitic fraction > 60% in all samples). Sands are medium-fine grained and made of quartz, mica flakes abundant. Two Mc interbeds (2300–2270 and 1830–1820) contained well preserved autochthonous ostracods.	Abundant Al F	fan toe/mud flat deposits. Two short palustrine events
II	2840–2360		Gm though poorly recovered. Polymictic gravel (metamorphic, dolostone, quartz), C: 20–60 mm, with coarse quartz sand, limonitized root casts, mica flakes abundant.	Al F	Alluvial fan (channel?) deposits
I	2960–2840		Carbonate lutites, with very fine grained quartz grains and mica flakes, Fe stains, botroidal limonite (former pyrite), bioclasts (forams) very abundant	Al F	deep marine environment

Note: Au F: Autochthonous foraminifers; Al. F.: Allochthonous foraminifers.

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Table 4. Information provided by the *n*-alkane ratios.

<i>n</i> -Alkane ratio	Information provided
Predominant <i>n</i> -alkane chain	Provides information of the origin of the biomolecules: phytoplankton and algae are dominated by low molecular weight compounds, maximizing at C ₁₇ (Gelpi et al., 1970; Blumer et al., 1971; Cranwell et al., 1987); submerged and floating macrophytes maximize at C _{21–25} (Cranwell, 1984; Ogura et al. 1990; Viso et al. 1993), high molecular weight <i>n</i> -alkanes (>C ₂₅) are typical of terrestrial vascular plants (Cranwell et al., 1987).
Average chain length	Indicates the origin of biomarkers: phytoplankton and algae have lower number of carbon atoms in the <i>n</i> -alkane molecules (Gelpi et al., 1970; Cranwell et al., 1987); submerged/floating macrophytes maximize at C ₂₁ , C ₂₃ and C ₂₅ ; terrestrial vascular plants maximize in >C ₂₅ (Cranwell et al., 1987).
Aquatic proxy (P _{aq}) (C ₂₃ +C ₂₅) / (C ₂₃ +C ₂₅ + C ₂₉ +C ₃₁)	Reflects the input of submerged/floating aquatic macrophytes relative to emergent and terrestrial plant input (Ficken et al., 2000). P _{aq} < 0.1 is linked to a dominant contribution of terrestrial plants; P _{aq} = 0.1–0.4 marks a marked development of emergent macrophytes; P _{aq} > 0.4 indicates a dominant input of submerged and floating macrophytes.
Terrigenous/Aquatic Ratio of hydrocarbons (TAR _{HC}) = C ₂₇ +C ₂₉ +C ₃₁ / (C ₁₅ +C ₁₇ +C ₁₉)	Reflects the input of terrestrial plants relative to algae input (Silliman et al., 1996).
Relative percentage of C ₃₁ <i>n</i> -alkane with respect to the sum C ₂₇ +C ₂₉ +C ₃₁	Large amounts of C ₃₁ in sediments are associated with dry phases and extensive grass cover and pines (Schwarz et al., 2002; Ortiz et al., 2010; Torres et al., 2013)

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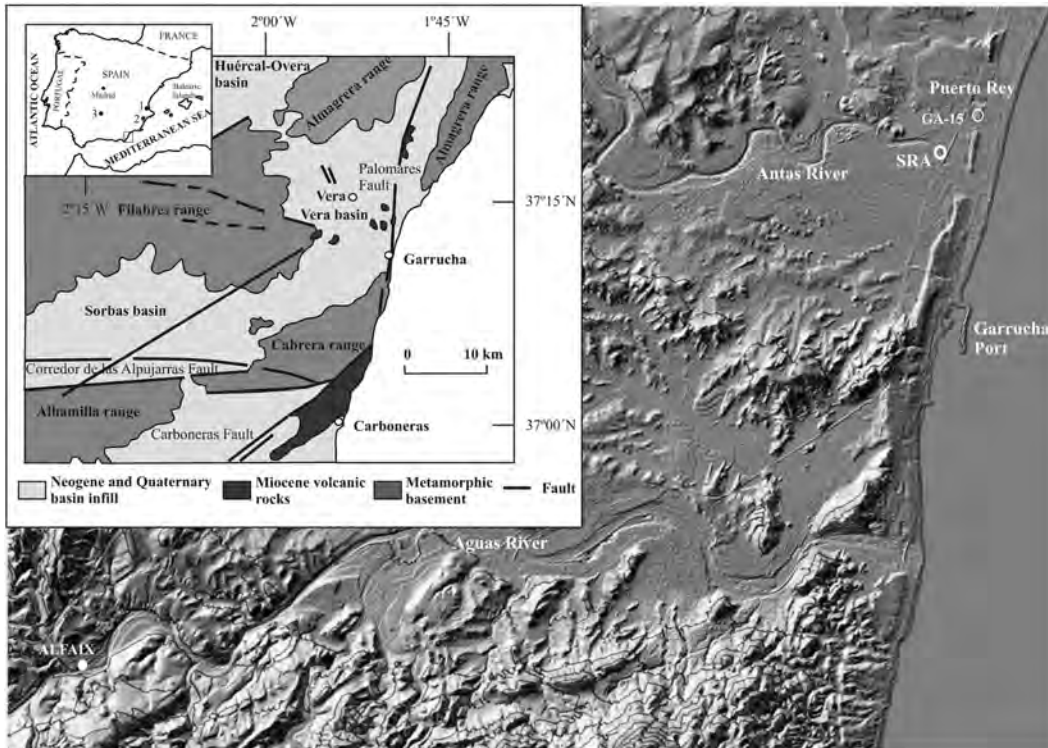


Figure 1. Geographical and Geological setting with the position of the localities. Other localities cited in the paper are also shown (1-Pego Basin; 2-Elche Basin; 3-Fuenteillejo).

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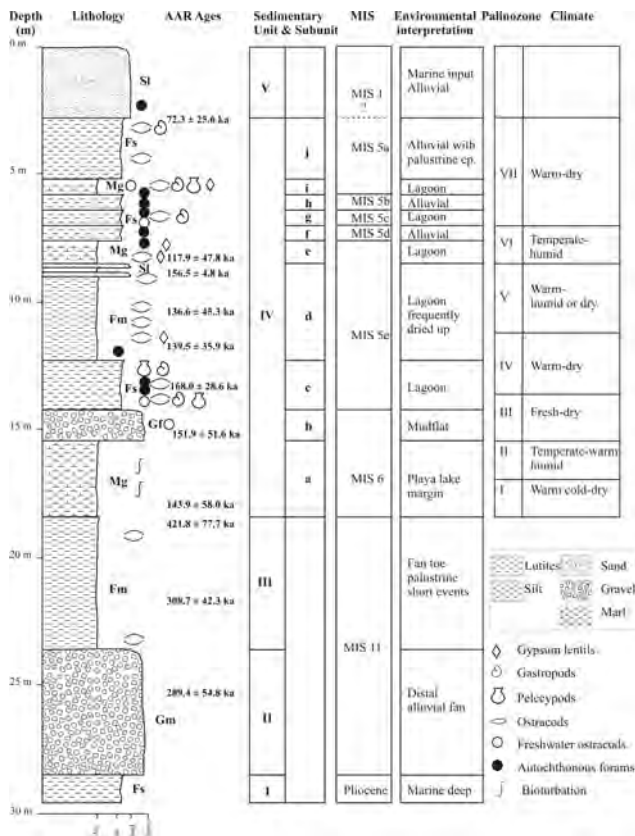


Figure 2. Stratigraphy and chronology of SRA core with the different sedimentological, micropaleontological and palynological units defined together with the palaeoenvironmental and palaeoclimatological interpretation.

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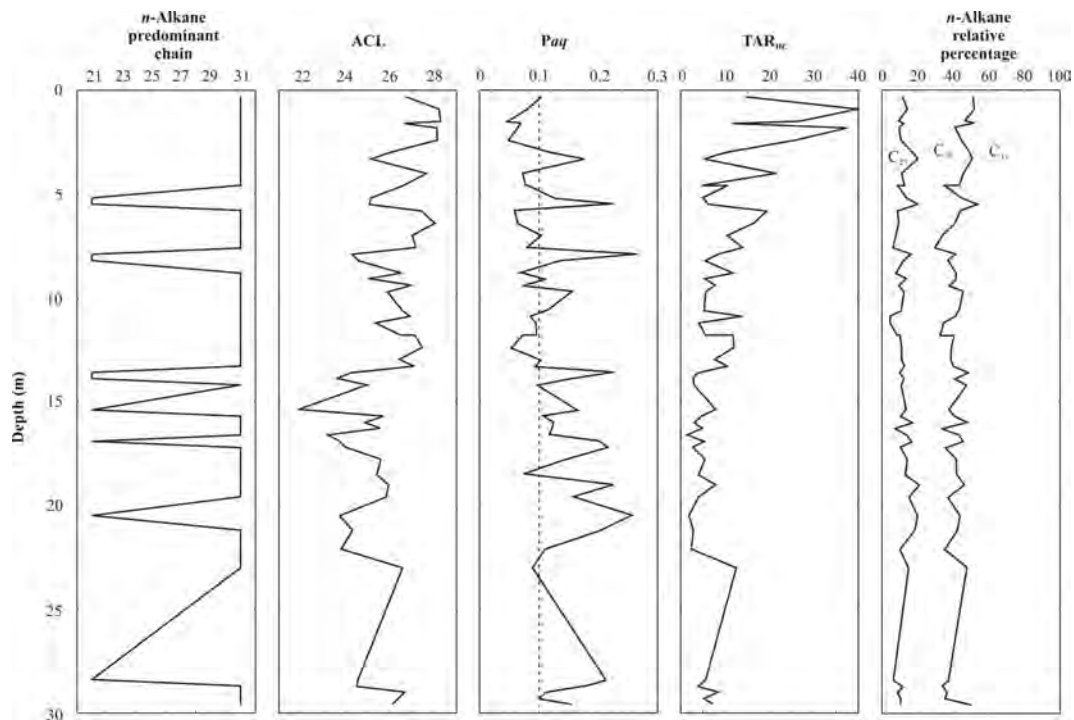


Figure 3. Profiles of the *n*-alkane predominant chain, ACL, Paq index, TAR_{HC} and the relative percentages of C₂₇, C₂₉ and C₃₁ with respect to the sum C₂₇ + C₂₉ + C₃₁ in SRA core.

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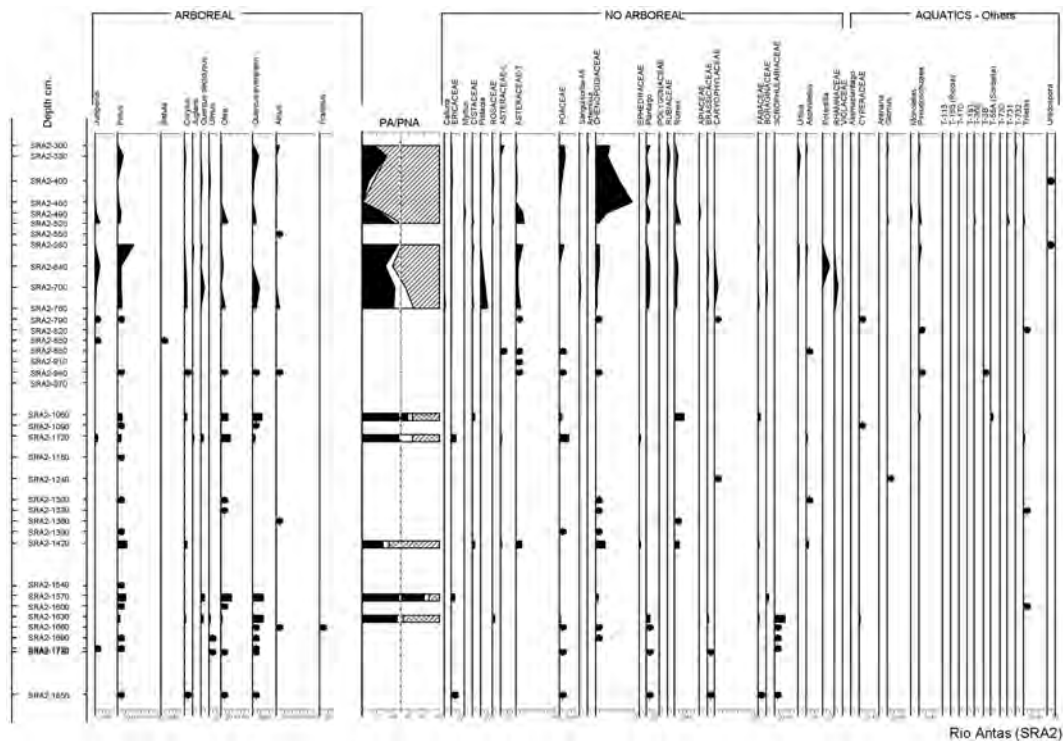


Figure 4. Pollen diagram of SRA core.

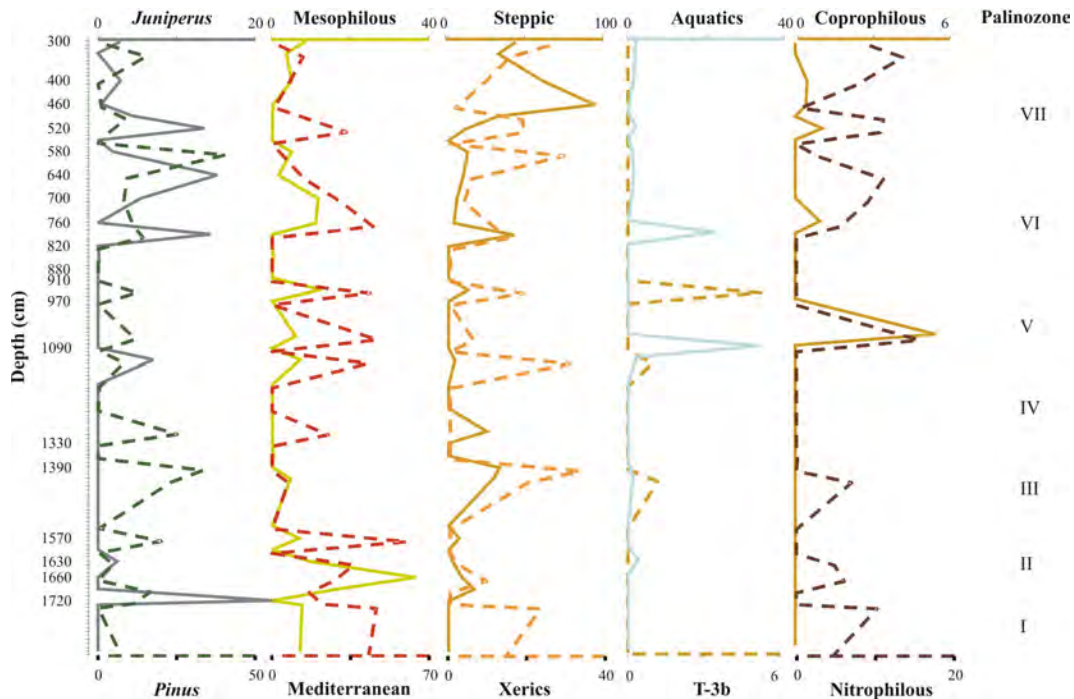


Figure 5. Synthetic pollen diagram of SRA core.

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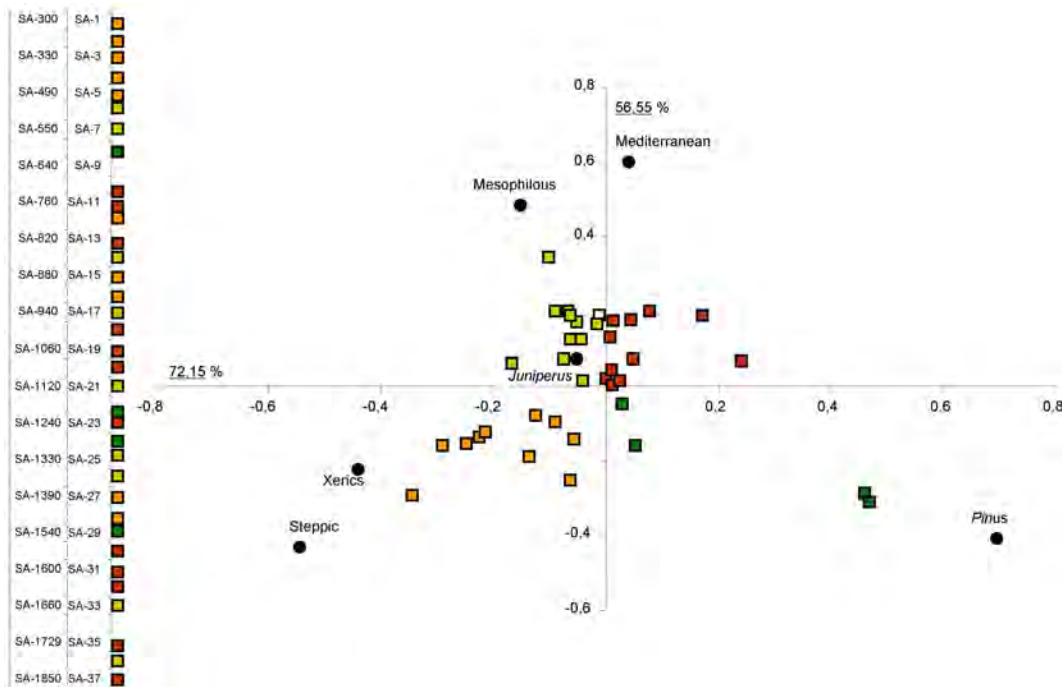


Figure 6. Principal component analysis of the pollen content in SRA core.

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