

The Late Quaternary pollen sequence of Toll Cave, a palaeontological site with evidence of human activities in northeastern Spain

Juan Ochando^{a,*}, José S. Carrión^{a,b}, Ruth Blasco^c, Florent Rivals^{d,e,f}, Anna Rufà^g, Gabriela Amorós^a, Manuel Munuera^h, Santiago Fernández^a, Jordi Rosell^{d,e}

^a Department of Plant Biology (Botany Area), Faculty of Biology, University of Murcia, Campus de Espinardo, 30100 Murcia, Spain

^b Evolutionary Studies Institute, University of Witwatersrand, South Africa

^c Centro Nacional de Investigación Sobre la Evolución Humana (CENIEH), Paseo Sierra de Atapuerca 3, 09002 Burgos, Spain

^d Institut Català de Paleoecologia Humana i Evolució Social (IPHES), Campus Sescelades URV (Edifici W3), 43007 Tarragona, Spain

^e Universitat Rovira i Virgili (URV), Departament d'Història i Història de l'Art, Avinguda de Catalunya 35, 43002 Tarragona, Spain

^f ICREA, Pg. Lluís Companys 23, 08010 Barcelona, Spain

^g PACEA - UMR 5199 CNRS, Université de Bordeaux, Bâtiment B2. Allée Geoffroy Saint Hilaire CS 50023, 33615, Pessac, Cedex, France

^h Department of Agricultural Engineering, Polytechnic University of Cartagena, 30203, Cartagena, Spain

ARTICLE INFO

Keywords:

Palaeoecology
Palynology
Archaeobiology
Pleistocene
Holocene

ABSTRACT

Palynological investigations of Toll Cave, a carnivore and archaeological cave site in northeastern Spain, are presented. The inferred vegetation reveals the long-term permanence of mixed pine-oak forests through a long period of environmental changes within the interval MIS 4 to MIS 1, and probably before. A relatively high diversity of woody taxa is found, including conifers, mesophytic angiosperms, Mediterranean forest, and xerothermic scrub. The most outstanding findings include the abundance of *Pinus*, evergreen *Quercus*, and *Juniperus*; the continuous occurrences of deciduous *Quercus*, *Acer*, *Castanea*, *Betula*, *Fraxinus*, *Buxus*, *Olea*, *Salix*, and *Erica*, and the presence of *Abies*, *Taxus*, *Carpinus betulus*, *Tilia*, *Populus*, *Celtis*, *Juglans*, *Ulmus*, *Calicotome*, *Cistus*, *Ephedra fragilis*, *Myrtus*, *Pistacia*, *Rhamnus* and *Viburnum*. Together with the pollen record of the nearby Teixoneres Cave, this new data suggest the existence of woodland refugia during the coldest and most arid stages of the upper Pleistocene across this relatively high-latitude region within the Iberian Peninsula. This study also supports the occurrence of forest ecosystems within the Mediterranean-Eurosiberian ecotone of the Iberian Peninsula in the vicinity of *Homo* habitats, including Neanderthals.

1. Introduction

Bringing new palaeobotanical records for the Pleistocene of north-eastern Iberia, more specifically for the ecotonal territories between the present-day Mediterranean and Eurosiberian regions, is one of the aims of several ongoing research projects led by this research group (Carrión et al., 2010, 2011, 2013, 2015; González-Sampériz et al., 2010). Lacustrine, peat bog, and offshore pollen records provide valuable information on the changes in the vegetation and climate of the Pleistocene in the extreme south-west of the European continent (Allué et al., 2017a, 2018; Burjachs, 1994; Burjachs and Julià, 1994, 1996; Burjachs et al., 2012; Carrión et al., 2013; Pérez-Obiol, 1988, 1994; Pérez-Obiol and Julià, 1994; Sardella et al., 2019). However, for the study of the past habitats associated with human occupations, palynological records in

caves, when successful (sterility is common: Carrión et al., 2009), provide detailed information from the palaeobotanical point of view and a reconstruction closer to the probable local environment in which humans had to exercise their adaptive capacities. Some notorious examples in the region are Teixoneres Cave (López-García et al., 2012; Ochando et al., 2020a), Cova de L'Arbreda (Burjachs, 1987; Burjachs and Renault-Miskovsky, 1992), Abric Romaní (Allué et al., 2017b, Burjachs and Julià, 1994; Val-Peón et al., 2019), Cova del Parco (Bergadà et al., 1999), Balma del Gai (Allué et al., 2007), La Cativera (Allué et al., 2000) and Gabasa (González-Sampériz, 2004; González-Sampériz et al., 2003). In this context, it should be noted the concurrence of pollen on carnivore coprolites (Carrión et al., 1995a, b, 1999, 2004, 2006, 2007, 2008, 2018; Daura et al., 2017; De Porras et al., 2017; Gatta et al., 2016; González-Sampériz et al., 2003; Horwitz and

* Corresponding author.

E-mail address: juan.ochando@um.es (J. Ochando).

<https://doi.org/10.1016/j.quaint.2020.06.048>

Received 3 May 2020; Received in revised form 23 June 2020; Accepted 29 June 2020

Available online 24 July 2020

1040-6182/© 2020 Published by Elsevier Ltd.

Goldberg, 1989; Latorre et al., 2002; Marais et al., 2015; Ochando et al., 2020b, accepted; Scott, 1987, 1994; Yll et al., 2006). Coprolites often complete the palaeobotanical record as they incorporate pollen types strictly entomophilous which can be rarely seen in non-biogenic sediments (Carrión, 2002; Carrión et al., 2018, 2019a, c).

In this work, we present the results of the pollen analysis carried out in Toll Cave palaeontological and archaeological site, adding value and complementing it with a previous study in the adjacent Cova de Teixoneres in the province of Barcelona, northeastern Spain. The findings are substantial to the debate on the distribution of glacial refuges of temperate and Mediterranean woods in the northernmost territories of the northern Mediterranean peninsulas (Bhagwat and Willis, 2008; Giardini, 2007; Lawson et al., 2004; Magri et al., 2017; Margari et al., 2009; Pini et al., 2010; Sadori et al., 2008, 2016; Sinopoli et al., 2018; Tzedakis et al., 2002, 2003; Wagner et al., 2019), and to a theoretical framework for the adaptation of Neanderthals and *H. sapiens* to forested environments of the cold and dry end of the Pleistocene.

2. Physical setting, palaeontological findings and geochronology

Toll Cave (TC) (2° 09' 02" E, 41° 48' 25" N, 760 m a.s.l.) is located near Moià (Barcelona province, northeastern Spain). The cave is part of a karstic system with a number of galleries (Fig. 1) modelled through autochthonous limestone of Neogene origin, the Collsuspina Formation. The region is located between the Ter River to the north, which connect the inner area of northern Catalonia with the Mediterranean Sea, and the Llobregat River to the south (Rosell et al., 2017). TC, approximately 2 km long, is made up of three main galleries facing south, east and west (Fig. 1). The current entrance is located in the South Gallery gathering most archaeo-palaeontological records (Rosell et al., 2014).

During the 1950s, pioneering researchers highlighted the presence of a well stratified deposit of brown lutites, silt and sand over 9 m thick (Serra et al., 1957; Bergadà and Serrat, 2001). According to them, the sedimentary deposit was formed by a succession of different flooding episodes, which finished with the colmatation of the cave during the Holocene. The upper stratigraphic unit (0.5–1 m thick) contains Bronze Age and Neolithic materials, while the rest of units (up to 12) contain Middle Palaeolithic tools. The Upper Palaeolithic is not represented in the cave, probably due to the aridity of the period and the consequent lack of sedimentation. Regarding to Middle Palaeolithic times, TC is one of the most renowned sites for cave bear (*Ursus spelaeus*) (Serra et al., 1957; Fernández-García and López-García, 2013; Rosell et al., 2012, 2014, 2015).

Unit 1 (0.5–1 m thick) may be situated in the MIS 1, Unit 2 (thick sands and gravels, 25–30 cm thick) is most likely MIS 1, but perhaps covers part of MIS 2. Unit 3 (1 m thick, orange lutites rich in faunal remains, mainly cave bears) is MIS 3. Unit 4, formed by brown lutites and sands with significant lateral changes (Serra et al., 1957), is difficult to separate stratigraphically from the underlying deposits. Ongoing excavations are taken Unit 4 as a main goal.

The Pleistocene faunal record of TC is composed mainly by cave bears, suggesting a regular use of the cave for the hibernation of these animals. However, there are also other large predators such as spotted hyaenas (*Crocuta crocuta*), wolves (*Canis lupus*), and lions (*Panthera leo spelaea*), in addition to small carnivores such as badgers (*Meles meles*), lynxes (*Lynx pardina*), foxes (*Vulpes vulpes*) and wild cats (*Felis silvestris*), which shows other uses of the cave, probably carnivore dens. Among ungulates, it is worth mentioning the presence of rhinoceros (*Rhinocerotidae* indet.), horses (*Equus ferus*), European asses (*Equus hydruntinus*), red deer (*Cervus elaphus*), roe deer (*Capreolus capreolus*), aurochs (*Bos primigenius*), chamois (*Rupicapra rupicapra*), wild goat (*Capra*



Fig. 1. Toll Cave. a) Location in Mediterranean northeastern Spain near the Eurosiberian region, b) cave entrance, c) excavation area.

pyrenaica), wild boar (*Sus scrofa*) and rabbits (*Oryctolagus cuniculus*) (Serra et al., 1957). Their presence in the cave corresponds, in the majority of the cases, to the accumulations of carnivores. Ramírez-Pedraza et al. (2019) have identified this faunal assemblage as belonging to the Upper Pleistocene, probably associated to a hibernation lair in Unit 4. This assignation is based in the abundance and features of the remains of *Ursus spelaeus*, and the activity of other carnivores as well, such as spotted hyenas and wolves (Rosell et al., 2012, 2014, 2015). At the lowermost Unit 4, Serra et al. (1957) pointed out the presence of hippopotamus (*Hippopotamus* sp.), which would place the base of the sequence at any time in the late Middle Pleistocene.

The chronology of the site is still under scrutiny. Provisional results are based in ^{14}C AMS and amino acid racemization, dating on three teeth and four bone samples (Kromer et al., 2013; Ramírez-Pedraza et al., 2019) (Table 1). Unit 3 is dated in c. 43,130 cal BP. For Unit 4, two bear bone samples were dated outside the ^{14}C range (>49,000 cal BP). The bones of a large-sized ungulate are radiocarbon dated in 47,310 cal BP. Three bear tooth samples dated by amino acid racemization, two of them gave age results between 57,900 (LEB 12757) and 69,800 cal BP (LEB 12756). Another sample (LEB 12755) provided an age of c. 150,000 cal BP, which is considered unrealistic based on the low amino acid concentration (Table 1). In sum, the onset of the sequence is unknown. As we shall see later, it could start at the end of MIS 6 or any stage of MIS 5, but it cannot be categorically discarded that all the sequence is situated between MIS 4 and MIS 1, with clear evidences of MIS 3 accumulation of sediments and biotic remains into the cave. From an archaeological point of view, TC was interpreted in the 1950s as a hibernation lair for cave bears that alternated with dens of other carnivores. However, the location of a lithic industry piece of clear Mousterian manufacture at Unit 4, as well as some bear bones with cut marks, suggested that the cavity could be occasionally used by human groups (Rosell et al., 2014).

Research on the geo- and biochronological context of the TC sedimentary sequence must continue. Strengthening the existing age estimate via the use of complementary extended-range luminescence dating signals (particularly single-grain thermally transferred OSL; e.g., Arnold and Demuro, 2015; Demuro et al., 2019) could be of crucial importance.

3. Modern-day climate and vegetation cover

TC is located in the supramediterranean belt with subhumid ombroclimate and, biogeographically, it belongs to the Pyrenean Province of the Eastern Pyrenean Sector (Rivas-Martínez, 1987; Rivas-Martínez et al., 2007). The area is located in a temperate, oceanic bioclimate, within the upper limits of the Mediterranean Region (Sub-Mediterranean climate) and near the Eurosiberian Region of northern Iberia (Fig. 1). The nearby meteorological station in Moia shows annual rainfall of 749 mm and an annual average temperature of 12.3 °C.

The vegetation of the region is dominated by forests adapted to mild summers and cold winters, patchily altered by disturbances such as overgrazing, fires, and extreme weather events. *Buxus sempervirens*, *Pinus halepensis* and *Quercus pubescens/humilis* are placed in the surroundings of the cave, accompanied by other trees such *Acer campestre*/

monspessulanum, *Pinus nigra* subsp. *salzmannii*, *Quercus ilex* and *Quercus petraea* (Fig. 1). These forests also contain a dense woody stratum, with *Crataegus monogyna*, *Daphne laureola*, *Hedera helix*, *Ilex aquifolium*, *Juniperus communis*, *Laurus nobilis*, *Lonicera xylosteum*, *Pistacia terebinthus*, *Prunus spinosa*, *Quercus coccifera*, *Rhamnus saxatilis*, *Rubus ulmifolius*, *Ruscus aculeatus*, *Sorbus torminalis* and *Viburnum lantana*, among other species. It should also be mentioned the presence of a relic beech forest (*Fagus sylvatica*) a few kilometers east of the cave site.

4. Methodology and palynological potential

As recommended for archaeological sediments (Girard, 1975), sampling for pollen was conducted on several stratigraphical profiles within the southern gallery (Units 1–4) (Fig. 2). Preliminarily, one sample per unit was processed to evaluate the palyno-analytical potential of the cave infill. Once this prospective study was completed with success (e.g. good preservation, counts above 100, low number of indeterminable palynomorphs, relatively high palyno-diversity), a more exhaustive sampling was carried out, focusing in the same profiles, approximately 1–14 m from the main entrance: Unit 1, crust Units 1–2, Unit 2, crust Unit 2–3, and Unit 3, were sampled in square Q-14, and Unit 4 in squares P-08, M-01, M-02 and M-03 (Fig. 2). 27 samples were analysed and 25 were found polleniferous (Table 2). Two sterile samples came from Unit 4 (4.20, 4.30). Exposed surface layers of sediment were cleaned back and discarded to a depth of 5–10 cm in Units 1 and 2, and 10–15 cm in the rest of the units, in order to avoid potential sources of contamination by recent bioturbation (Fig. 2).

For the extraction of palynomorphs in the laboratory, we followed the conventional chemical analysis described by Dumbleby (1985) with the modifications proposed by Girard and Renault-Miskovsky (1969). In order to evaluate the quality of the laboratory preparation procedures and identify palynological sterility, we added to each sample three tablets of *Lycopodium* spores of a known concentration. After being treated at the laboratory, the samples were mounted on slides with liquid paraffin. The palynological identification and counting was made by conventional microscopy (400x and 1000x) using a transmitted light optical microscope, and the pollen reference collection of the Department of Plant Biology, University of Murcia. The pollen counts were treated with Tilia Graph 1.7.16 software in order to construct pollen diagrams (Figs. 3–8). A total of 8703 palynomorphs were identified; 6825 pollen grains and 1878 spores. Spores, non-pollen microfossils, and the pollen grains of Asteraceae (Asteroideae, Cichorioideae, *Anthemis* t. and *Centaurea montana* t.) were excluded from the pollen sum assuming over-representation, as a consequence of local over-deposition. The number of pollen taxa is between 5 and 39 per sample, with a total of 66 taxa being recognized. The percentage of indeterminable is lower than 10% (Table 2), and the preservation was generally good. The lowest pollen concentrations were recorded in the lowermost samples of Unit 4, and the maxima were found in the upper sample of Unit 1, perhaps due to compaction of the Holocene sediments. The highest number of taxa are recorded in Unit 1.

Table 1

Geochronological data for Toll Cave. Further details of the ^{14}C AMS can be found in Ramírez-Pedraza et al. (2019).

	Lab. Ref.	Unit	Field ref.	Depth (Z)	Mat.	Taxon	Technique	Dating (cal BP)	Error 1σ (Ka)	Observ.
^b MPI	S-EVA 27843	3	T'11/3/P17/21	−190	Bone	<i>Cervus elaphus</i>	^{14}C AMS	43,130	0,34	
^a LEB	LEB 12755	4	T'11/4/Q12/19	−174	Tooth	<i>Ursus spelaeus</i>	Amino acid racemization	150,000		Low quantity of amino acid
^b MPI	S-EVA 27845	4	T'11/4/Q12/60	−191	Bone	<i>Ursus spelaeus</i>	^{14}C AMS	>49,000		
^b MPI	S-EVA 27851	4	T'11/3/Q11/44	−199	Bone	<i>Ursus spelaeus</i>	^{14}C AMS	>49,000		
^a LEB	LEB 12757	4	T'11/4/P16/66	−199	Tooth	<i>Ursus spelaeus</i>	Amino acid racemization	57,900		
^b MPI	S-EVA 27850	4	T'11/3/P16/71	−200	Bone	Large sized ung.	^{14}C AMS	47,310	0,54	Cut marks
^a LEB	LEB 12756	4	T'11/4/P13/29	−203	Tooth	<i>Ursus spelaeus</i>	Amino acid racemization	69,800		

^a LEB: Laboratorio de Estratigrafía Biomolecular (Madrid, Spain).

^b MPI: Max Planck Institute (Leipzig, Germany).

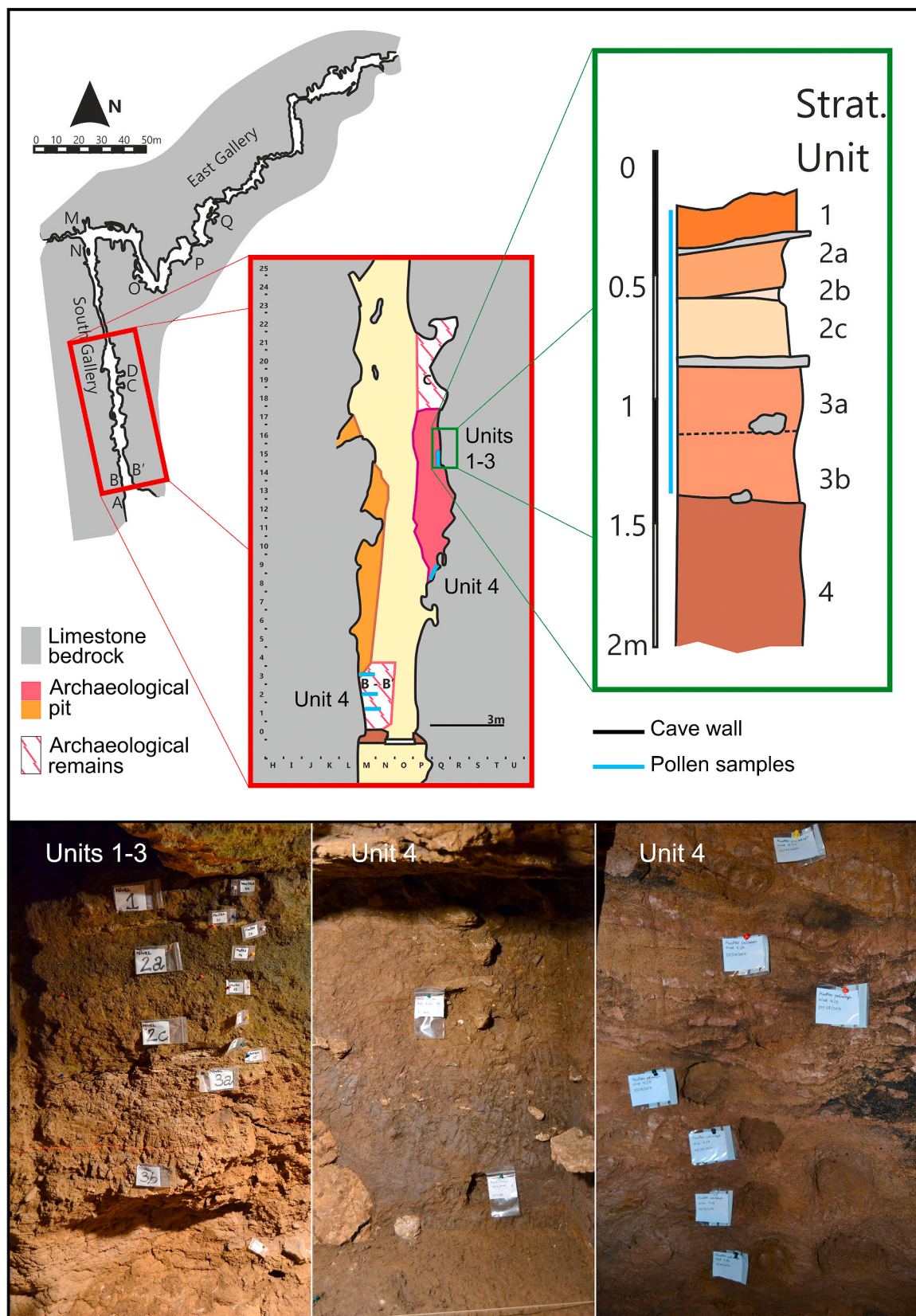


Fig. 2. Stratigraphical units sampled for pollen analysis in Toll Cave.

Table 2
Summary of palynological features of the Toll samples.

Sample	Unit	N°	Field data	Depth (cm)	Gross Weight (g)	Net Weight (g)	Concentration (grains/g)	Indeterminable (%)	*Pollen sum	Number of taxa (Pollen)	Spores sum
1.1	1	1	Q14/ x=1637;Y=1400	27	81.2	67.8	4638,29	3,97	302	36	58
1.2	1	2	Q14/ x=1637;Y=1400	30	73.8	56,6	745,05	0,22	456	39	202
Crust 1.2-2a.1											
2a.1	2	4	Q14/ x=1637;Y=1400	34	43.8	43,8	144,55	0,88	341	31	59
2a.2	2	5	Q14/ x=1637;Y=1395	39	80.8	55,0	3566,62	4,71	318	35	26
2c.1	2	6	Q14/ x=1637;Y=1400	42	76.3	31,3	1795,46	0,21	469	31	59
2c.2	2	7	Q14/ x=1637;Y=1390	55	76.8	55,1	851,42	3,43	349	34	23
Crust 2c.2-3a											
3a	3	8	Q14/ x=1635;Y=1327	63	74.7	55,5	744,28	0,23	426	27	52
3b	3	9	Q14/ x=1637;Y=1400	70	74.7	74,7	109,45	0,72	416	26	48
4.1	4	10	Q14/ x=1660;Y=1333	100	77,3	61,4	195,13	2,46	203	22	10
4.2	4	11	P08/ x=1557;Y=642	125	76,5	70,3	474,74	2,36	212	17	10
4.3	4	12	P08/ x=1557;Y=634	125	75,3	75,3	3731,31	0,81	491	24	54
4.4	4	13	P08/ x=1557;Y=682	150	72,4	66,5	2391,97	0,41	487	18	17
4.6	4	14	P08/ x=1545;Y=694	175	73,2	72,0	2833,61	0,20	495	19	27
4.8	4	15	P08/ x=1545;Y=687	203	76,8	66,6	470,75	3,79	211	16	30
4.10	4	16	M01/ x=1230;Y=60	222	78,1	68,7	223,97	4,45	202	16	88
4.12	4	17	M02/ x=1280;Y=108	244	76,5	49,5	65,62	8,78	205	17	146
4.14	4	18	M02/ x=1248;Y=107	305	76,4	54,8	158,00	6,47	201	20	197
4.16	4	19	M02/ x=1248;Y=105	333	78,0	56,5	31,47	8,33	144	26	239
4.18	4	20	M02/ x=1248;Y=105	361	77,2	75,0	13,46	5,13	39	14	220
4.20	4	21	M02/ x=1270;Y=106	390	75,7	70,8	24,49	9,52	21	6	0
4.22	4	22	M02/ x=1239;Y=108	401	76,5	68,1	27,28	6,81	44	10	43
4.24	4	23	M02/ x=1239;Y=108	426	75,9	41,7	0	0	0	0	0
4.26	4	24	M03/ x=1210;Y=202	446	76,2	64,6	75,51	9,52	21	5	0
4.28	4	25	M03/ x=1209;Y=202	486	75,1	47,6	48,73	1,75	57	9	27
4.30	4	26	M03/ x=1217;Y=203	517	75,1	40,3	44,41	6,38	47	12	234
	4	27	M03/ x=1219;Y=203	542	77,0	44,4	151,41	5,13	39	7	9
	4			569	72,2	64,2	0	0	0	0	0
TOTAL 6196											1878

^a Pollen sum column without Asterioideae, Cichorioideae, Anthemis type and Centaurea montana type.

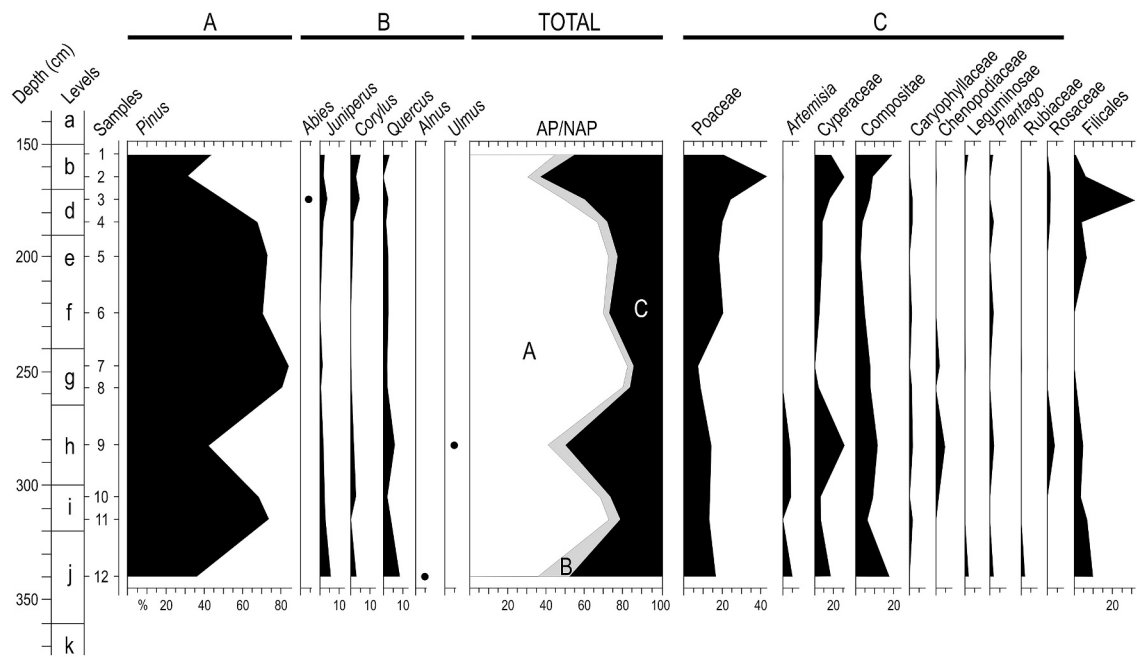


Fig. 3. Pollen diagram from Toll Cave redrawn from [Donner and Kurtén \(1958\)](#).

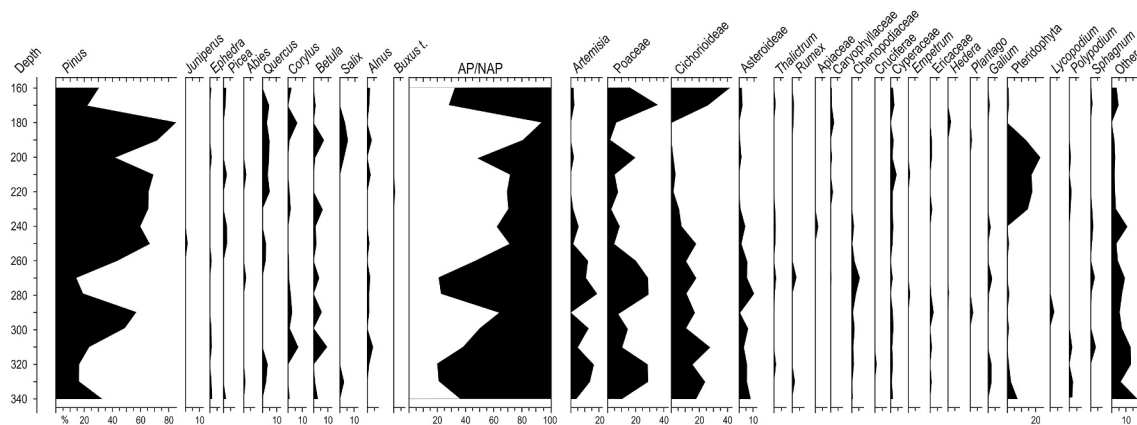


Fig. 4. Pollen diagram from Toll Cave redrawn from [Menéndez-Amor and Florschütz \(1962\)](#).

5. Pollen sequence

5.1. Previous palynological research

Several previous pollen studies have been performed at TC. [Donner and Kurtén \(1958\)](#) analysed 12 pollen samples, at irregular intervals but at specified depth from –155 to –340 cm, collected from the witness section “Sondage B” ([Fig. 3](#)) ([Serra et al., 1957](#)), situated at the cave entrance. Several years later, [Menéndez-Amor and Florschütz \(1962\)](#), published a second study, allegedly from 19 samples, at 10 cm intervals from –160 to –340 cm, of the same profile ([Fig. 4](#)). As mentioned by [Butzer and Freeman \(1968\)](#), it is highly likely that “the two sets of pollen data refer to the same beds of the same section”. Allegedly, both studies were aimed at studying the Neolithic vegetational landscapes, and interpreting the potential changes from the Pleistocene to the Holocene as well. Although the AP and NAP curves in both diagrams may be somewhat comparable, there are enough details that allow us to make different palaeobotanical and palaeoecological interpretations. [Menéndez-Amor and Florschütz \(1962\)](#) interpreted the presence of pollen of deciduous arboreal species such as *Quercus*, *Corylus*, *Alnus*, *Betula*, and *Salix* as by long distance transport. In addition, the pollen diagram

shows two distinct kinds of alternating pattern: one with high pine values (60–90%) accompanied by grasses, and another one with low pine values (15–45%) together with abundant grasses, *Artemisia* and other Asteraceae ([Fig. 4](#)). In contrast, [Donner and Kurtén \(1958\)](#) contend the existence of local broad-leaf trees while their pollen diagram shows less fluctuation in the main pollen contributors ([Fig. 3](#)). All these differences could be reflecting on such factors as identification of pollen in poor condition, selective destruction of pollen types through different techniques of sample preparation, and size sample ([Butzer and Freeman, 1968](#)). It bears emphasis that both papers ([Donner and Kurtén, 1958](#); [Menéndez-Amor and Florschütz, 1962](#)) fail to reckon that most of the cave had been extensively anthropogenically modified by Neolithic groups. Therefore, it means that the first meter below the Neolithic unit, where these authors took the samples, could be mixed with different materials, possibly older. In fact, in more recent excavations, remains of cave bear bones have been found along with ceramics and beads. Some of the criticism made later by [Butzer and Freeman \(1968\)](#), on the work of [Donner and Kurtén \(1958\)](#), and [Menéndez-Amor and Florschütz \(1962\)](#) may be justified by these factors.

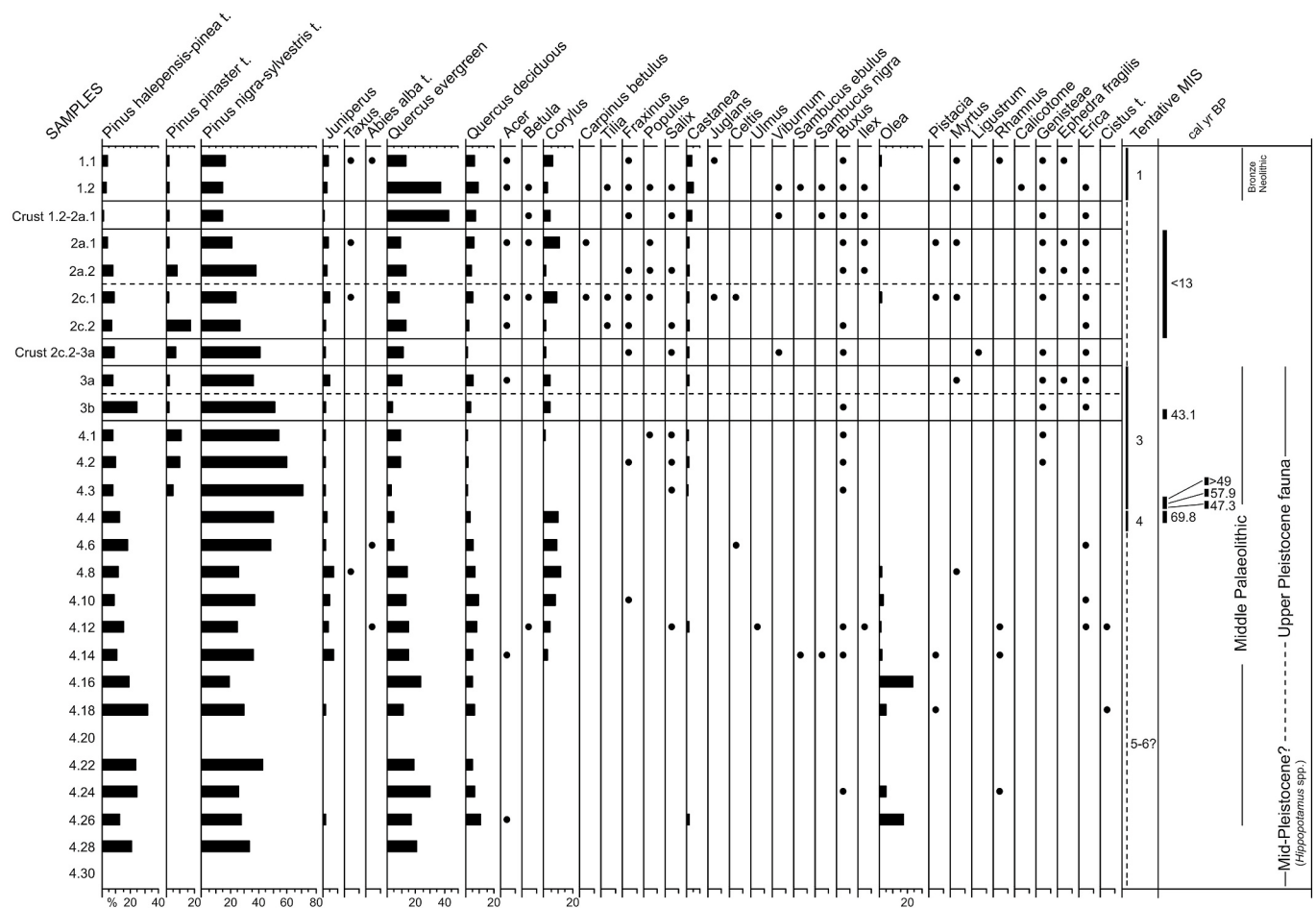


Fig. 5. Pollen diagram of Toll Cave (this paper) including principally the woody component. Asteroideae, Cichorioideae, *Anthemis* type, and *Centaurea montana* type are out from the total pollen sum. Black dots for percentages below 3%. Dashed lines for subdivisions within archaeological units.

5.2. General features of the TC pollen record

Overall, the TC pollen record is characterized by the prevalence and continuity of a *Pinus-Quercus* forest, which is in some stages accompanied by *Corylus*, cf *Juniperus* and to a lesser extent, *Castanea* t (type), *Buxus*, *Olea*, *Salix* and *Erica* (Figs. 5–7). The woody component exhibits high cover and noticeable diversity, and includes a combination of mesophytes, such as deciduous trees that feature Eurosiberian temperate forests (*Acer*, *Betula*, *Celtis*, *Fraxinus*, *Juglans*, *Populus*, *Carpinus betulus*, *Tilia*, *Ulmus*), and Mediterranean taxa in addition to evergreen oaks and pines (*Viburnum*, *Pistacia*, *Myrtus*, *Calicotome*, *Cistus*, *Ephedra fragilis*, *Rhamnus*). The xero-heliophytic component (Poaceae, *Artemisia*, *Amaranthaceae*, *Ephedra*) is relatively unimportant, with the exception of samples 1.1, 2a.1, 2c.2, crust 2c.2-3a, and 4.28 (Figs. 5–7).

A broad spectrum of non-pollen palynomorphs (NPPs) have been found in TC, including fungal, fern, algal, and moss spores (Fig. 8; Table 3). A majority of the NPPs identified in TC also occur in the vicinity at the Pleistocene sediments of Teixoneres Cave such as *Glomus*, *Diporisorites*, *Hypoxylonites*, *Inapertisorites*, *Microsporionites* and *Monoporisorites* (Fig. 8; Table 3). However, the number of NPPs in Teixoneres is higher, with exclusive taxa such as *Asplenium*, *Dicellaesporites*, *Dyadosporites*, *Fusiformisporites*, *Uncinulites*, *Striadiporites*, *Quilonia*, among others. The taphonomical processes involved in these similarities and differences are unknown to us.

5.3. TC pollen stratigraphy

Unit 4 includes samples from 4.30 to 4.1 (Figs. 5–7) and depicts a

palynozone dominated by arboreal pollen (AP), accounting 100% of the total counts in several samples. The frequencies of *Pinus nigra-sylvestris* are c. 19–71%, *Pinus halepensis-pinea* varies between 8 and 32%, with evergreen *Quercus* ranging between 3 and 30%, *Olea* exceeds 23%, deciduous *Quercus* reaches 11%, *Juniperus* is below 8%, and *Corylus* at 12%. Other lesser contributors to AP include *Pinus pinaster*, *Salix*, *Buxus*, *Castanea*, *Pistacia*, *Myrtus*, *Rhamnus* and *Erica*. In the non-arboreal pollen (NAP) category, Poaceae attains relative abundances close to 21% in sample 4.28, *Artemisia* reaches 8% and *Amaranthaceae*, Cichorioideae, Asteroideae, Fabaceae, *Lotus*, Brassicaceae, Urticaceae and *Plantago lanceolata*, are frequent. Fungal spores abound, especially *Glomus* (1–83%), *Glomus* chlamydospores, *Ctenosporites*, *Hypoxylonites*, and *Monoporisorites*. *Algae* shows maxima above 82% (Fig. 8).

Unit 3 includes pollen samples 3a and 3b (Figs. 5–7) and is also clearly dominated by trees, with AP higher than 81%. The abundance of pines is noteworthy, with a decreasing trend for *Pinus halepensis-pinea* throughout the zone. *Pinus nigra-sylvestris* is consistently high, fluctuating between 36 and 51%, while *Pinus pinaster* does not exceed 1%. *Quercus* shows an increasing trend from 7% (3b) up to 16% (3a). Evergreen *Quercus* ranges from 4 to 10% and deciduous *Quercus* between 3 and 5%. *Corylus* (4–5%), *Juniperus* (2–4%) and *Erica* (1–4%) are noteworthy, as are the occurrence of *Acer*, *Castanea*, *Buxus*, *Myrtus*, *Ephedra fragilis* and *Genisteae*. *Artemisia* (4–5%) and Poaceae (2–3%) are not abundant, neither are Fabaceae, *Lotus*, Asteroideae, Cichorioideae and *Centaurea montana*. The abundance of *Glomus* is outstanding (50–70%) (Fig. 8). *Dicellaesporites*, *Inapertisorites*, *Multicellites*, *Scleroderma* and Type 209 are frequent, while bryophyte spores abound in 3b (Fig. 8).

In the crust sample 2c.2-3a, AP continues to be dominant, surpassing

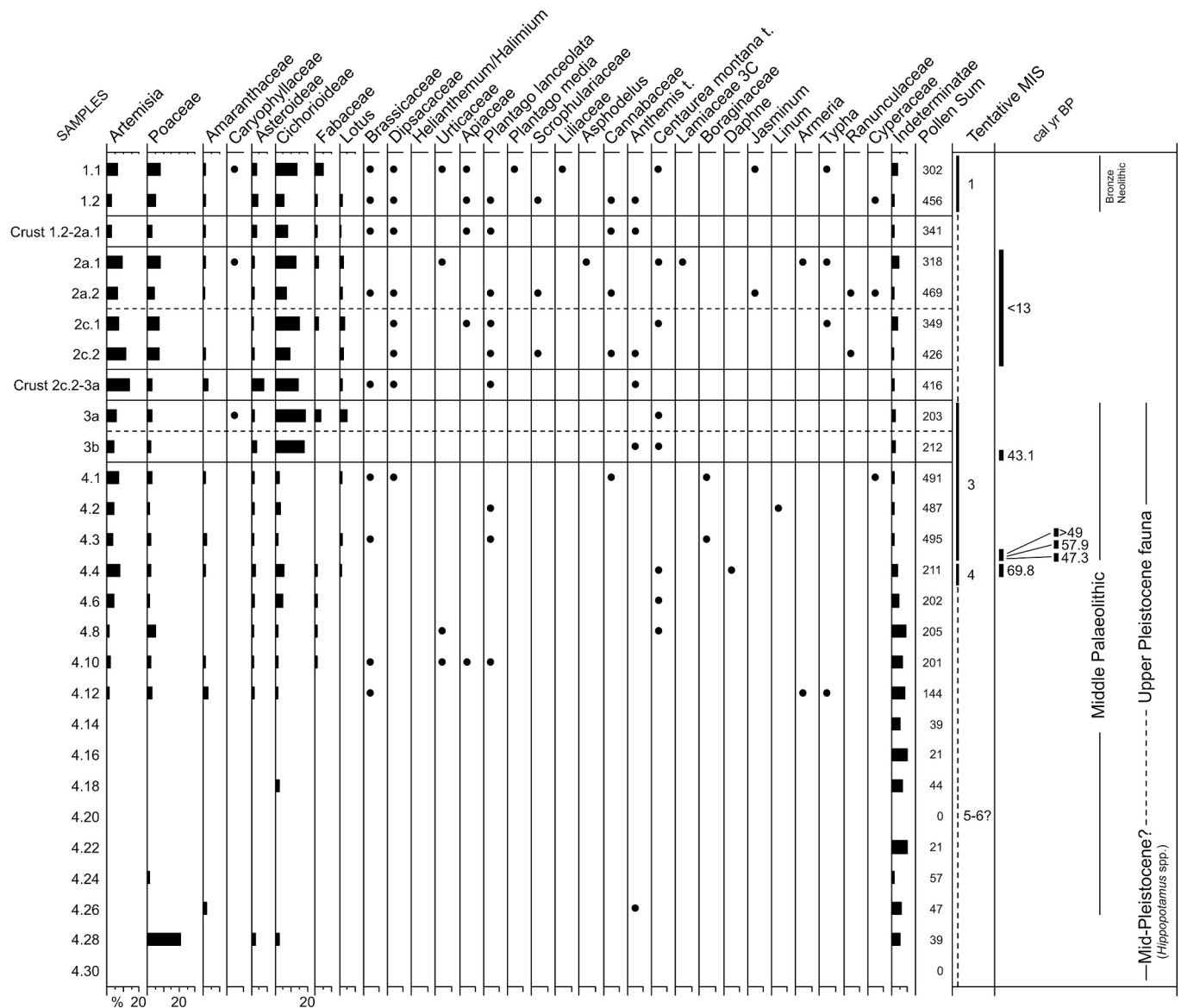


Fig. 6. Pollen diagram of Toll Cave (This paper) including mainly non arboreal elements. Asteroideae, Cichorioideae, *Anthemis* type, and *Centaurea montana* type are out from the total pollen sum. Black dots for percentages below 3%. Dashed lines for subdivisions within archaeological units.

77%. *Pinus nigra-sylvestris* (41%), evergreen *Quercus* (12%), and *Pinus halepensis-pinea* (8%) are noticeable. *Pinus pinaster* is 6% and *Erica* less than 2%. Other AP contributors are *Juniperus*, deciduous *Quercus*, *Corylus*, *Fraxinus*, *Salix*, *Castanea*, and Genisteae. Poaceae does not exceed 3%, while *Artemisia* is more frequent than in Unit 3, exceeding 13%. Among the fungal spores, *Monoporosporites* stands out as being above 48%, and *Glomus*, *Hypoxylonites* and *Diporisorites* are frequent. The ferns are represented by *Polypodium* (4%) (Fig. 8).

Unit 2 includes samples 2a.1, 2a.2, 2c.1 and 2c.2 (Figs. 3–5). AP is above 73%, with *Pinus nigra-sylvestris* (21–39%), *Pinus pinaster* (1–17%), evergreen *Quercus* (8–13%), *Corylus* (1–11%), *Pinus halepensis-pinea* (3–9%), deciduous *Quercus* (2–6%), *Juniperus* (2–4%), and *Erica* (1–3%) as principal taxa, accompanied by *Taxus*, *Acer*, *Betula*, *Carpinus betulus*, *Fraxinus*, *Salix*, *Buxus*, *Myrtus* and Genisteae. *Artemisia* (6–11%) and Poaceae (5–8%) are dominant within NAP. Amaranthaceae, Asteroideae, Cichorioideae, *Lotus*, Dipsacaceae, *Plantago lanceolata*, Cannabaceae and *Centaurea montana*, are frequent. *Glomus* (39–83%) is abundant, and *Diporisorites*, *Gelasinospora*, *Hypoxylonites*, *Inapertisporites*, *Monoporosporites* and *Scleroderma*. and fern spores, including *Polypodium*, occur in this zone. The ferns are represented by *Triletes* and

Polypodium. Bryophyte spores exceed 9% (Fig. 8).

The pollen sample Crust 1.2-2a.1 shows AP dominant, surpassing 90%. Evergreen *Quercus* (43%), *Pinus nigra-sylvestris* (15%), and deciduous *Quercus* (7%) are noticeable. *Castanea* is 4% and *Salix* less than 4%. Other AP contributors are *Pinus halepensis-pinea*, *Pinus pinaster*, *Juniperus*, *Fraxinus*, *Buxus*, *Erica*, and Genisteae. Poaceae does not exceed 4%, while *Artemisia* is less frequent than in Unit 2, below 3%. Among the fungal spores, *Glomus* stands out as being above 44%, and *Hypoxylonites*, *Ctenosporites*, *Fractisporonites*, and *Reduviasporonites* are frequent.

Unit 1 includes samples 1.1 and 1.2 (Figs. 3–5). The predominant pollen type continues to be AP, with total values higher than 71%. The abundance of *Quercus* (19–47%) is remarkable, with evergreen *Quercus* ranging between 13 and 38%, and deciduous *Quercus* between 6 and 9%. *Pinus halepensis-pinea* (3–4%) and *Pinus nigra-sylvestris* (14–17%) decrease with respect to the previous units. The relatively high frequencies of *Corylus* (2–7%), *Castanea* (3–5%), *Juniperus* (3–4%), and Genisteae (2–4%) are noticeable. Other lesser contributors to AP include *Pinus pinaster*, *Taxus*, *Acer*, *Fraxinus*, *Viburnum*, *Buxus*, *Myrtus*, *Rhamnus* and *Erica* are well represented. In the non-arboreous pollen (NAP) category, Poaceae attains relative abundances close to 9% in sample 1.1,

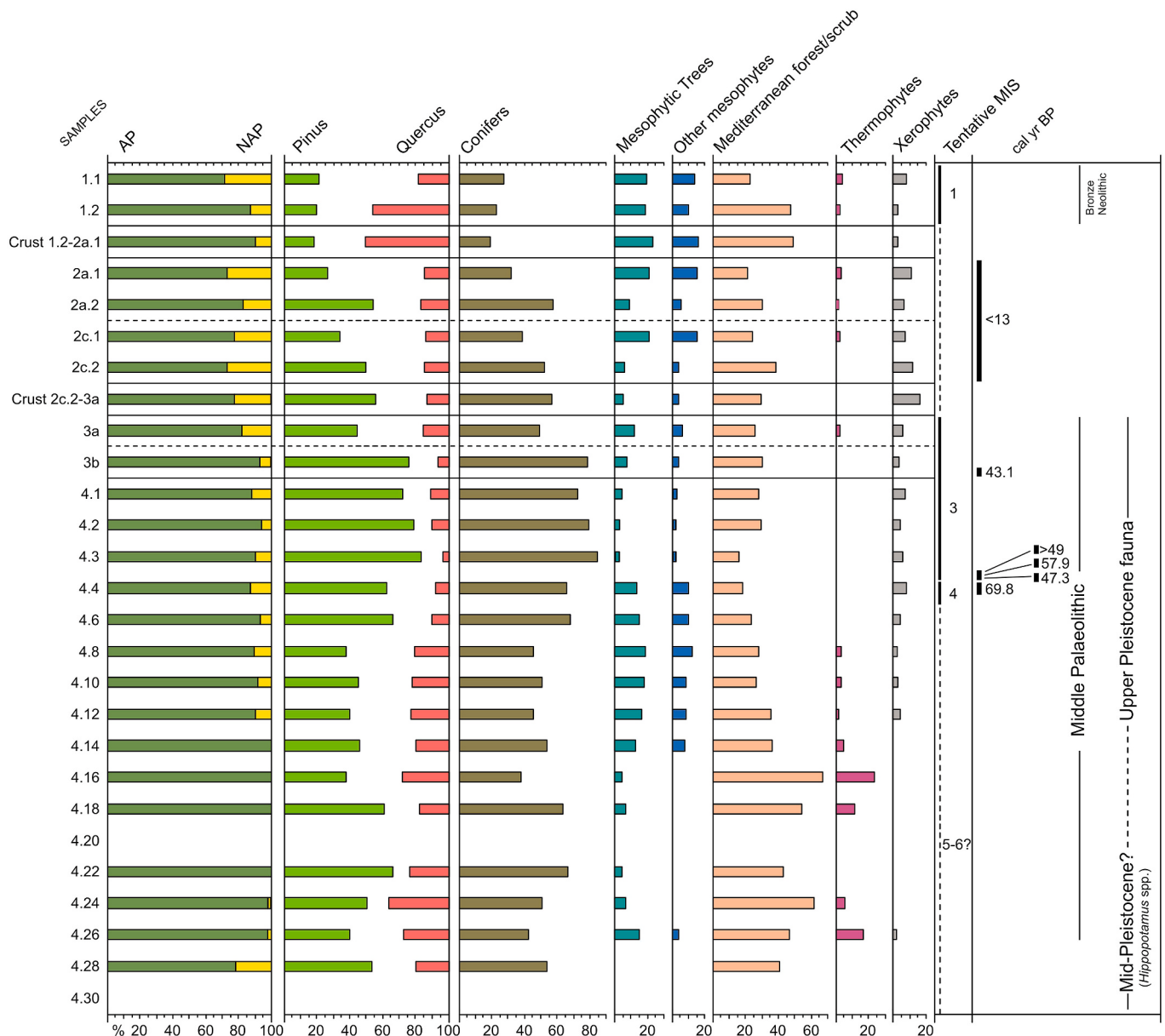


Fig. 7. Synthetic pollen diagram of Toll Cave (this paper) including ecological groups and the main pollen contributors. Conifers include *Pinus*, *Juniperus*, *Taxus* and *Abies*. Mesophytic trees include deciduous *Quercus*, *Acer*, *Betula*, *Corylus*, *Carpinus betulus*, *Tilia*, *Fraxinus*, *Populus*, *Salix*, *Castanea*, *Juglans*, *Celtis*, *Ulmus*, *Sambucus nigra* and *Ilex*. Other Mesophytes comprise mesophytes trees minus deciduous *Quercus*. Mediterranean forest/scrub include evergreen *Quercus*, *Pinus halepensis*, *Pinus pinaster*, *Viburnum*, *Sambucus ebulus*, *Buxus*, *Olea*, *Pistacia*, *Myrtus*, *Calicotome*, *Ephedra fragilis*, *Erica* and *Cistus*. Thermophytes include *Olea*, *Pistacia*, *Myrtus*, *Calicotome*, *Ephedra fragilis*, *Cistus* and *Asphodelus*. Xerophytes include *Artemisia*, *Amaranthaceae*, *Asphodelus*, *Lamiaceae* and *Ephedra fragilis*. The dashed lines indicate subdivisions within an archaeological unit.

Artemisia reaches 7% and *Amaranthaceae*, *Cichorioideae*, *Asterioideae*, *Fabaceae*, *Lotus*, *Dipsacaceae* and *Brassicaceae*, are frequent. Fungal spores are abundant, especially *Glomus* (17–62%), *Diporisorites* (10–28%), *Monoporisorites* (3–12%), *Dictyosporites*, and *Inapertisporites*, while bryophyte spores abound in 1.1.

6. Palaeoecological and palaeobotanical remarks

In view of the findings described above, here we postulate in the study area a long-term permanence of forest ecosystems dominated by pines and oaks with an important contribution of *Corylus*, *Juniperus* and *Castanea*, which were continuously accompanied by other trees such as *Abies*, *Taxus*, *Acer*, *Betula*, *Carpinus betulus*, *Tilia*, *Celtis*, *Fraxinus*, *Juglans*, *Buxus*, *Ilex*, *Populus*, *Salix*, and *Ulmus*, as well as Mediterranean elements such as *Pistacia*, *Myrtus*, *Calicotome*, *Cistus*, *Ephedra fragilis*, *Ligustrum*,

Rhamnus and *Viburnum* (Figs. 5 and 7). The xero-heliophytic component (*Artemisia*, *Poaceae*, *Amaranthaceae*, *Erica*, *Ephedra*) would be relatively unimportant with the exception of some phases (1.1, 2a.1, 2c.2, crust 2c.2-3a, and 4.28) where the landscapes opened somewhat probably as consequence of cold dry spells (Fig. 6). However, the woodland structure and composition would have remained unaltered and both the forest density and the thermophytic component (e.g. *Olea*, *Buxus*, *Pistacia*, *Myrtus*, *Fraxinus*, *Populus*, *Salix*, *Castanea*) preserved with minor variations (Fig. 7).

Palaeoenvironmental analyses and studies of small mammals (including rodents) in the site also suggests the predominance of local woodlands (Allué et al., 2013; Fernández-García and López-García, 2013; López-García et al., 2012). This inference is supported by isotopic carbon studied by Ramírez-Pedraza et al. (2019) who point out that the average value of $\delta^{13}\text{C}$ in *Ursus spelaeus* (only adults) corresponds to

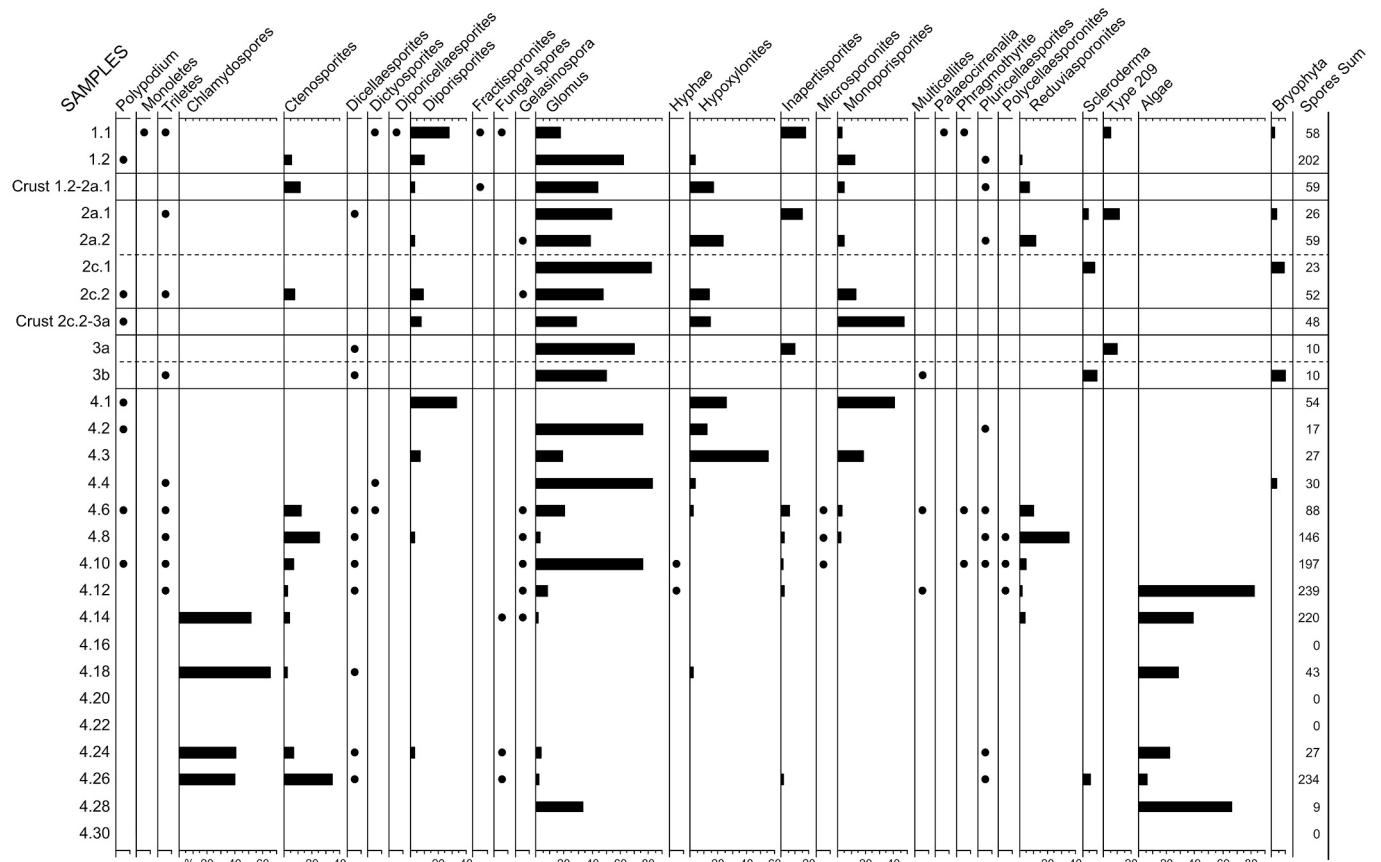


Fig. 8. Palynogram of spore and non-pollen palynomorphs of Toll Cave (this paper). Black dots indicate percentages below 3%. All taxa percentages calculated out from the total pollen sum.

Table 3

Non-pollen palynomorphs from Toll and Teixoneres sequences.

Caves	Toll-Teixoneres	Toll	Teixoneres
Non-pollen palynomorphs	<i>Polypodium</i>	<i>Chlamydozporites</i>	<i>Asplenium</i>
	<i>Monoletes</i>	<i>Palaeocirrenalia</i>	<i>Dicellaesporites</i>
	<i>Triletes</i>	<i>Phragmothyrites</i>	<i>Dyadosporites</i>
	<i>Ctenosporites</i>		<i>Fusiformisporites</i>
	<i>Dicellaesporites</i>		<i>Palambages</i>
	<i>Dictyosporites</i>		<i>Quilonia</i>
	<i>Diporicellaesporites</i>		<i>Striadiporites</i>
	<i>Diporisporites</i>		<i>Trichothyrites</i>
	<i>Fractisporites</i>		<i>Uncinulites</i>
	<i>Gelasinospora</i>		Type 224
	<i>Glomus</i>		Type 232
	<i>Hyphae</i>		Type 339
	<i>Hypoxylonites</i>		
	<i>Inapertisporites</i>		
	<i>Microsporites</i>		
	<i>Monoporisporites</i>		
	<i>Multicellites</i>		
	<i>Pluricellaesporites</i>		
	<i>Polycellaesporites</i>		
	<i>Reduviasporites</i>		
	<i>Scleroderma</i>		
	Type 209		
	Algae		
	Bryophyta		

animals with a dominant consumption of C_3 plants. $\delta^{13}C$ in bear caves are more negative than in the rest of the herbivores and carnivores analysed, and may certainly indicate the presence of a closed forest habitat for this species.

This palaeoenvironmental scenario involves a glacial refugium for

woodland ecosystems with relatively dense cover and high species diversity. A pollen study in the nearby Teixoneres Cave confirms this pattern during MIS 3 and MIS 2 (Ochando et al., 2020a). Indeed, the Toll-Teixoneres ecosystems identify a so far unknown, reservoir of angiosperm forest during the Pleistocene in northern Iberia. Similar findings can only be found far southwards within the Iberian Pleistocene such as during the Mid-Pleistocene Bolomor (Ochando et al., 2019), and the Upper Pleistocene Navarrés (Carrión and Dupré, 1996; Carrión and van Geel, 1999; Carrión et al., 1999), littoral of Murcia at Perneras and Sima de las Palomas (Carrión and Dupré, 1994; Carrión et al., 1995a, 2003), and Gibraltar caves (Carrión et al., 2008, 2018), but doubtless, mixed deciduous-evergreen forests are not so manifest in the south. The TC new pollen record in relatively high altitude within Spain must therefore be incorporated to the discussion on glacial refugia for temperate trees in the Mediterranean Peninsulas (Bhagwat and Willis, 2008; Carrión et al., 1999, 2003; Giardini, 2007; González-Sampériz et al., 2010; Lawson et al., 2004; Magri and Sadori, 1999; Magri et al., 2017; Manzano et al., 2017; Margari et al., 2009; Pini et al., 2010; Sadori et al., 2008, 2016; Sinopoli et al., 2018; Tzedakis, 1994, 1999; Tzedakis et al., 2002, 2003; Wagner et al., 2009, 2014; Willis, 1994).

The evidence of evergreen *Quercus* (mainly *Quercus ilex*) during the Quaternary glacial stages indicated limited abundance in northern Iberia (Uzquiano et al., 2016). The presence of deciduous oaks has been nevertheless shown in the ecotonal territories from Mediterranean to Eurosiberian regions (Aranbarri et al., 2016; Blanco-Castro et al., 1997; García-Mijangos et al., 2015; Gil-Romera et al., 2014; González-Sampériz, 2004; González-Sampériz et al., 2004, 2017; Loidi, 2017; Morales-Molino and García-Antón, 2014; Salomón et al., 2016). Pine forests (*Pinus halepensis*, *Pinus nigra* and *Pinus sylvestris*) may have been, however, widespread in the region during the study period, even in cold stages, as shown by a number of palaeobotanical studies (Allué

et al., 2007, 2018; Bergadà et al., 1999; Burjachs, 1994, 2009; Burjachs and Julià, 1994; Burjachs and Renault-Miskovsky, 1992; González-Sampériz, 2004; González-Sampériz et al., 2003; Pérez-Obiol and Julià, 1994; Val-Peón et al., 2019; Yll, 1995). *Pinus pinaster* was rather a component of mixed forests with deciduous and evergreen oaks. A critical study at the local scale is by López-García et al. (2012) who analysed charcoals remains in hearths within units II, IIb and III from Teixoneres showing the abundance of *Pinus pinea/pinaster* and *Pinus sylvestris*.

Carpinus betulus is worth commenting because its current distribution in the Iberian Peninsula is restricted (Postigo-Mijarra et al., 2010). *Carpinus* has existed in the Peninsula since the Oligocene (Postigo-Mijarra et al., 2008, 2009), and there are palynological evidences during the Middle Pleistocene (Desprat et al., 2005, 2007; García Antón and Sainz Ollero, 1991). Fossil remains for *Carpinus* are still usual in the late Pleistocene (Postigo-Mijarra et al., 2008, 2009, 2010; Carrión et al., 2013) demonstrating its well-known resistance to cold conditions. Several localities pertinent to this case are Abric Romaní (Burjachs et al., 2012) during MIS 3, Pla de l'Estany-Garrotxa (Burjachs, 1990, 1994) during the Eemian (substage 5e), and Cañizar de Villarquemado record (González-Sampériz et al., 2013; García-Prieto, 2015) during MIS 5. Thereafter during the Holocene, *Carpinus* populations were depleted (Postigo-Mijarra et al., 2008; Magri et al., 2017) leading to its almost complete disappearance. However, its native presence has been confirmed in a few inhabited areas between the Cantabrian Mountains and the extreme west of the Pyrenees (Aizpuru and Catalán, 1984). The increased vulnerability of *Carpinus* communities through glacial-interglacial cycles, together with ecological factors, may have caused the disappearance of this taxon during the Holocene, whose extirpation from the Iberian Peninsula is still poorly understood (Magri et al., 2017). The presence of hornbeam (*Carpinus betulus*) at TC reinforces previous palynological data (Burjachs, 1990, 1994; Burjachs et al., 2012; González-Sampériz et al., 2013; Postigo-Mijarra et al., 2010) that supports the naturalness of the current populations in southwestern Europe.

7. Neanderthals in forest ecosystems

Although the archaeological and palaeontological studies carried out in TC (Ramírez-Pedraza et al., 2019; Rosell et al., 2012, 2014, 2015; Serra et al., 1957) point towards a prolonged use over time of the cave as a hibernation lair for carnivores, the occurrence some of lithic tools of clear Mousterian manufacture at Unit 4, and bear bones with cut marks, suggests that the cavity was occasionally used by Middle Palaeolithic human groups (Rosell et al., 2014). Therefore, the studied pollen sequence site can be directly compared with human and cave bears occupation histories at TC, and the relevant connections between these inhabitants and their ecosystem. Archaeological excavations in TC have shown a broad faunal diversity with species belonging to large and small carnivores, ungulates and small vertebrates (Rosell et al., 2012, 2014; 2015; Serra et al., 1957). At synchronous phases, it is also noteworthy in the nearby Teixoneres site the presence of a wide spectrum of edible plants accessible in the proximity, such as holly oak (*Quercus ilex*), olive (*Olea europaea*), Mediterranean hackberry (*Celtis australis*), elderberry (*Sambucus nigra*), hazelnut (*Corylus avellana*), walnut (*Juglans regia*) and chestnut (*Castanea sativa*).

The idea that Neanderthals were significant inhabitants of woodlands as well has been suggested by several authors (Carrión and Walker, 2019; Carrión et al., 2018, 2019b; Finlayson and Carrión, 2006, 2007; Rosas, 2016; Stewart, 2005; Stewart et al., 2019). Similarly, Nabais and Zilhão (2019) consider that Neanderthals were highly knowledgeable of their environments, integrating small prey in their diet, such as rabbits, turtles and birds since very early (e.g., Blasco, 2008; Blasco and Fernández Peris, 2009; Blasco et al., 2013; Blasco et al., 2014; Blasco et al., 2016; Fiore et al., 2016; Morin et al., 2019). Clearly, Neanderthals adapted to a wide range of environments and were able to develop

complex, diverse and successful subsistence strategies in each territory (Spagnolo et al., 2019). Thus, fuel acquisition strategies have been seen on the part of Neanderthals, as well as occupation patterns in the Northwest of the Iberian Peninsula through taxonomic and taphonomic studies with charcoal (Allué et al., 2017a). Nevertheless, the pressures from hunting and predation in difficult terrain will also have led to risks of injury and mortality in even the most favourable environments, as shown by Spikins et al. (2019). Interestingly, Stewart et al. (2019) contend that North European Neanderthals were particularly adapted to the conditions of temperate episodes during which a wooded landscape and rich faunal diversity existed. In this context, the regional palaeovegetation picture shown here has important implications for existing arguments about the habitats and long survival of Neanderthals in the Iberian Peninsula (Carrión, 2004; Finlayson and Carrión, 2007; Higham, 2014; Jiménez-Espejo et al., 2007; Stewart, 2005; Wood et al., 2013; Zilhão et al., 2017).

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

The development of this work was supported by Project (CGL-BOS2015-68604-P), funded by: FEDER/Ministry of Science and Innovation - Agencia Estatal de Investigación Project (PID2019-1049449BB-I00), funded by: FEDER/Ministry of Science and Innovation - Agencia Estatal de Investigación and Fundación Séneca (grant number 20788/PI/18). The research at Teixoneres Cave is supported by projects CLT009-18-00055 and 2017 SGR 836 from the Generalitat de Catalunya, and projects HAR2016-76760-C3-1-P and PID2019-103987GB-C31 from Spanish AEI. J Rosell develops his work within the Spanish AEI/FEDER projects PGC2018-093925-BC32 and CGL2016-80000-P, and R Blasco within PID2019-104949GB-I00. A Rufà is a beneficiary of a postdoctoral research contract funded by the IdEx University of Bordeaux Investments for the Future program (ANR No.-10-IDEX-03-02) and also collaborates in the Generalitat de Catalunya projects CLT009-18-00053 and CLT009-18-00054. Special thanks to the excavation team for their very useful help during the fieldwork seasons.

References

- Aizpuru, I., Catalán, P., 1984. The presence of the bombean in the Iberian Peninsula. *An. J. Bot. Madrid* 41, 143–146.
- Allué, E., Angelucci, D., Cáceres, L., Flocchi, C., Fontanals, M., García, M., Huguet, R., Ollé, A., Saladié, P., Vergès, J.M., Zaragoza, J., 2000. El registro paleoecológico y arqueológico de La Catiuera (El Catllar, Tarragona): datos preliminares sobre el límite Pleistoceno-Holoceno en el sur de Cataluña. *Actas do 3º Congresso de Arqueologia Peninsular. IX Contributos das ciências e das tecnologias para a arqueologia da Península Ibérica*, pp. 81–96. Porto.
- Allué, E., Nadal, J., Estrada, A., García-Argüelles, P., 2007. Los datos antracológicos de la Balma del Gai (Bages, Barcelona): una aportación al conocimiento de la vegetación y la explotación de los recursos forestales durante el Tardiglacial en el NE peninsular. *Trab. Prehist.* 64, 87–98.
- Allué, E., Burjachs, F., Vernet, J.-L., Morales, J.I., Rodríguez-Hidalgo, A., Cebrià, A., Rosell, J., 2013. Cova del Toll (Moià, Bages): perspectiva paleoambiental I arqueobotànica del Plistocè I Holocè. *Quad. Preh. Cat.* 21–38.
- Allué, E., Picornell-Gelabert, L., Daura, J., Sanz, M., 2017a. Reconstruction of the palaeoenvironment and anthropogenic activity from the upper Pleistocene/Holocene anthracological records of the NE Iberian peninsula (Barcelona, Spain). *Quat. Int.* 457, 172–189.
- Allué, E., Solé, A., Burguet-Coca, A., 2017b. Fuel exploitation among Neanderthals based on the anthracological record from Abric Romaní (Capellades, NE Spain). *Quat. Int.* 431, 6–15.
- Allué, E., Martínez-Moreno, J., Roy, M., Benito-Calvo, A., Mora, R., 2018. Montane pine forests in ne Iberia during MIS 3 and MIS 2. A study based on new anthracological evidence from Cova gran (santa linya, iberian pre-pyrenees). *Rev. Palaeobot. Palynol.* 258, 62–72.
- Aranbarri, J., Bartolomé, M., Alcolea, M., Sancho, C., Celant, A., González-Sampériz, P., Arenas, C., Magri, D., Rodríguez-Lázaro, J., 2016. Palaeobotanical insights from

- early-mid Holocene fluvial in the Moncayo natural park (Iberian range, NE Spain): regional correlations and biogeographic implications. *Rev. Palaeobot. Palynol.* 234, 31–43.
- Arnold, L.J., Demuro, M., 2015. Insights into TT-OSL signal stability from single-grain analyses of known-age deposits at Atapuerca, Spain. *Quat. Geochronol.* 30, 472–478.
- Bergadà, M.M., Serrat, D., 2001. Seqüència sedimentària i paleoambiental de la Cova del Toll (Moia): darretes aportacions. *Modilium* 24, 7–22.
- Bergadà, M., Burjachs, F., Fullola, J.M., 1999. Évolution paléoenvironnemental du 14.500 au 10.000 BP dans le Prépyrénées catalans: La Grotte du Parco (Alòs de Balaguer, Lleida, Espagne). *L'Anthropologie* 103, 249–264.
- Bhagwat, S.A., Willis, K.J., 2008. Species persistence in Northern glacial refugia of Europe: a matter of chance or biogeographical traits? *J. Biogeogr.* 35, 464–482.
- Blanco-Castro, E., Casado-González, M.A., Costa-Tenorio, M., Escribano-Bombín, R., García-Antón, M., Génova-Fuster, M., Gómez-Manzanque, F., Moreno-Saiz, J.C., Morla-Juaristi, C., Regato-Pajares, P., Sainz-Ollero, H., 1997. Los Bosques Ibéricos. Una Interpretación Geobotánica. *Planeta*, S.A., p. 572.
- Blasco, R., 2008. Human consumption of tortoises at level IV of bolomor cave (valencia, Spain). *J. Archaeol. Sci.* 35, 2839–2848.
- Blasco, R., Fernández Peris, J., 2009. Middle Pleistocene bird consumption at level XI of bolomor cave (valencia, Spain). *J. Archaeol. Sci.* 36, 2213–2223.
- Blasco, R., Finlayson, C., Rosell, J., Sánchez Marco, A., Finlayson, S., Finlayson, G., Negro, J.J., Giles Pacheco, F., Rodríguez-Vidal, J., 2014. The earliest pigeon fanciers. *Sci. Rep.* 4, 5971.
- Blasco, R., Rosell, J., Fernández Peris, J., Arsuaga, J.L., Bermúdez de Castro, J.M., Carbonell, E., 2013. Environmental availability, behavioural diversity and diet: a zooarchaeological approach from the TD10-1 sublevel of Gran Dolina (Sierra de Atapuerca, Burgos, Spain) and Bolomor Cave (Valencia, Spain). *Quaternary Science Reviews* 70, 124–144. <https://doi.org/10.1016/j.quascirev.2013.03.008>.
- Blasco, R., Rosell, J., Rufà, A., Sánchez Marco, A., Finlayson, C., 2016. Pigeons and choughs, a usual resource for the Neanderthals in Gibraltar. *Quaternary International* 421, 62–77. <https://doi.org/10.1016/j.quaint.2015.10.040>.
- Burjachs, F., 1987. Palinología de los niveles Gravetienses, Solutrense y Postsolutrense de la Cova de l'Arbreda (Serinyà, Girona). *Actas de la VII Reunión sobre Cuaternario. Universidad de Santander, AEQUA, Santander, Cantabria*, pp. 19–21.
- Burjachs, F., 1990. Palinología dels doïmens de l'Alt Empordà i dels diposits quaternaris de la cova de l'Arbreda (Serinyà, Pla de l'Estany) i del Pla de l'Estany (Olot, Garrotxa). *Evolució del paisatge vegetal i del clima des de fa mes de 140.000 anys al NE de la Península Ibérica*. Ph.D. Thesis. Universitat Autònoma de Barcelona, Bellaterra, p. 324.
- Burjachs, F., 1994. The palynology of the upper Pleistocene and Holocene of the North East Iberian Peninsula: Pla de l'Estany (Catalonia). *Hist. Biol.* 9, 17–33.
- Burjachs, F., 2009. Paleoambient del Tardiglacial al sud dels Pirineus vist a través de la Palinologia. In: Mercadal, O., Coord (Eds.), *Els Pirineus i les àrees circumdants durant el Tardiglacial. Mutacions i filiacions tecnoculturals, evolució paleoambiental (16.000-10.000 BP)*. Museu Cerdà, Puigcerdà, pp. 151–162.
- Burjachs, F., Renault-Miskovsky, J., 1992. Paléoenvironnement et paléoclimatologie de la Catalogne durant près de 30,000 ans (du Würmien ancien au début de l'Holocène) d'après la Palynologie du site de l'Arbreda (Gérone, Catalogne). *Quaternaire* 3, 75–85.
- Burjachs, F., Julià, R., 1994. Abrupt climatic changes during the last glaciation based on pollen analysis of the abric Romaní, Catalonia, Spain. *Quat. Res.* 42, 308–315.
- Burjachs, F., Julià, R., 1996. Palaeoenvironmental evolution during the middle-upper palaeolithic transition in the NE of the Iberian peninsula. In: Carbonell, E., Coord (Eds.), *The Last Neanderthals, the First Anatomically Modern Humans*. URV, Tarragona, pp. 377–383.
- Burjachs, F., López-García, J.M., Allué, E., Blain, H.-A., Rivals, F., Bennàsar, M., Expósito, I., 2012. Palaeoecology of Neanderthals during dansgaard-oeschger cycles in northeastern Iberia (abric Romaní): from regional to global scale. *Quat. Int.* 247, 26–37.
- Butzer, K.W., Freeman, L.G., 1968. Pollen analysis at the Cueva del Toll, Catalonia: a critical re-appraisal. *Geol. Mijnbouw* 47, 116–120.
- Carrión, J.S., 2002. A taphonomic study of modern pollen assemblages from dung and surface sediments in arid environments of Spain. *Rev. Palaeobot. Palynol.* 120, 217–232.
- Carrión, J.S., 2004. The use of two pollen records from deep sea cores to frame adaptive evolutionary change for humans: a comment on "Neanderthal extinction and the millennial scale variability of OIS3" by F. d'Errico and M.F. Sánchez-Goni. *Quat. Sci. Rev.* 23, 1217–1219.
- Carrión, J.S., Dupré, M., 1994. Pollen data from mousterian sites in southeastern Spain. *AASP Contrib. Ser.* 29, 17–26.
- Carrión, J.S., Dupré, M., 1996. Late Quaternary vegetational history at Navarrés, Eastern Spain. A two core approach. *New Phytol.* 134, 177–191.
- Carrión, J.S., van Geel, B., 1999. Fine-resolution upper weichselian and Holocene palynological record from Navarrés (valencia, Spain) and a discussion about factors of mediterranean forest succession. *Rev. Palaeobot. Palynol.* 106, 209–236.
- Carrión, J.S., Walker, M.J., 2019. Background to neanderthal presence in Western mediterranean Europe. *Quat. Sci. Rev.* 217, 7–44.
- Carrión, J.S., Dupré, M., Fumanal, M.P., Montes, R., 1995a. A palaeoenvironmental study in the semi-arid south-eastern Spain: the palynological and sedimentological sequence at Perneras cave (Lorca, Murcia). *J. Archaeol. Sci.* 22, 355–367.
- Carrión, J.S., Munuera, M., Dupré, M., 1995b. Estudios de Palinología arqueológica en el sureste ibérico semiárido. *Cuaternario Geomorfol.* 9, 17–31.
- Carrión, J.S., Munuera, M., Navarro, C., Burjachs, F., Dupré, M., Walker, M., 1999. Palaeoecological potential of pollen records in caves: the case of Mediterranean Spain. *Quat. Sci. Rev.* 18, 67–78.
- Carrión, J.S., Yll, E.I., Walker, M.J., Legaz, A.J., Chain, C., 2003. Glacial refugia of temperate, Mediterranean and Ibero-North African flora in South-Eastern Spain: new evidence from cave pollen at two Neanderthal man sites. *Global Ecol. Biogeogr.* 12, 119–129.
- Carrión, J.S., Yll, R., Riquelme, J.A., González, P., 2004. Perspectivas del análisis polínico de coprolitos y otros depósitos biogénicos útiles en la inferencia paleoambiental. In: *Miscelanea en Homenaje a Emiliano Aguirre. Volumen II: Paleontología*. Museo Arqueológico Regional, Alcalá de Henares, pp. 129–140.
- Carrión, J.S., Scott, L., Marais, E., 2006. Environmental implications of pollen spectra in bat droppings from south-eastern Spain and potential for palaeoenvironmental reconstructions. *Rev. Palaeobot. Palynol.* 140, 175–186.
- Carrión, J.S., Scott, L., Arribas, A., Fuentes, N., Gil-Romera, G., Montoya, E., 2007. Pleistocene landscapes in central Iberia inferred from pollen analysis of hyena coprolites. *J. Quat. Sci.* 22, 191–202.
- Carrión, J.S., Finlayson, C., Fernández, S., Finlayson, G., Allué, E., López-Sáez, A., López-García, P., Fuentes, N., Gil, G., González-Sampériz, P., 2008. A coastal reservoir of biodiversity for Upper Pleistocene human populations: palaeoecological investigations in Gorham's Cave (Gibraltar) in the context of the Iberian Peninsula. *Quat. Sci. Rev.* 27, 2118–2135.
- Carrión, J.S., Fernández, S., González-Sampériz, P., Leroy, S.A.G., López-Sáez, J.A., Burjachs, F., Gil-Romera, G., Rodríguez-Sánchez, E., García-Antón, M., Gil-García, M.J., Parra, I., Santos, L., López-García, P., Yll, E.I., Dupré, M., 2009. Sterility cases and causes in Quaternary pollen analysis in the Iberian Peninsula: the advantages of reporting bad luck. *Internet Archaeol.* <http://intarch.ac.uk/journal/issue25/5/toc.html>.
- Carrión, J.S., Fernández, S., González-Sampériz, P., Gil-Romera, G., Badal, E., Carrión-Marco, Y., López-Merino, L., López-Sáez, J.A., Fierro, E., Burjachs, F., 2010. Expected trends and surprises in the lateglacial and Holocene vegetation history of the Iberian peninsula and balearic islands. *Rev. Palaeobot. Palynol.* vol. 162, 458–475. Special issue: Carrión, J.S., Leroy, S. (Eds.), *Iberian Floras through Time: Land of Diversity and Survival*.
- Carrión, J.S., Rose, J., Stringer, C., 2011. Early human evolution in the western Palaearctic: ecological scenarios. *Quat. Sci. Rev.* 30, 1281–1295.
- Carrión, J.S., Fernández, S., González-Sampériz, P., López-Merino, L., Peña, L., Burjachs, F., López-Sáez, J.A., García-Antón, M., Carrión-Marco, Y., Uzquiano, P., Postigo, J.M., Barrón, E., Allué, E., Badal, E., Dupré, M., Fierro, E., Munuera, M., Rubiales, J.M., García-Amorena, I., Jiménez-Moreno, G., Gil-Romera, G., Leroy, S., García-Martínez, M.S., Montoya, E., Fletcher, W., Yll, E., Vieira, M., Rodríguez-Ariza, M.O., Anderson, S., Peñalba, C., Gil-García, M.J., Pérez-Sanz, A., Albert, R.M., Díez, M.J., Morales, C., Gómez-Manzanque, F., Parra, I., Ruiz-Zapata, B., Riera, S., Zapata, L., Ejarque, A., Vegas, T., Rull, V., Scott, L., Andrade, A., Pérez-Díaz, S., Abelschaad, D., Moreno, E., Hernández-Mateo, L., Ochando, J., Pérez Navarro, M.A., Sánchez-Baena, J.J., Riquelme, J.A., Iglesias, R., Franco, F., Chaín, C., Figueiral, I., Grau, E., Matos, M., Jiménez-Espejo, F., Arribas, A., Garrido, G., Finlayson, G., Finlayson, C., Ruiz, M., Pérez-Jordá, G., Miras, Y., 2013. *Paleoflora Ibérica: Plioceno-Cuaternario*, 2 vols. Ministerio de Economía y Competitividad. Universidad de Murcia y Fundación Séneca, Madrid, 84-616-6797-2.
- Carrión, J.S., Fernández, S., González-Sampériz, P., López-Merino, L., Peña, L., Burjachs, F., López-Sáez, J.A., García-Antón, M., Carrión-Marco, Y., Uzquiano, P., Postigo, J.M., Barrón, E., Allué, E., Badal, E., Dupré, M., Fierro, E., Munuera, M., Rubiales, J.M., García-Amorena, I., Jiménez-Moreno, G., Gil-Romera, G., Leroy, S., García-Martínez, M.S., Montoya, E., Fletcher, W., Yll, E., Vieira, M., Rodríguez-Ariza, M.O., Anderson, S., Peñalba, C., Gil-García, M.J., Pérez-Sanz, A., Albert, R.M., Díez, M.J., Morales, C., Gómez-Manzanque, F., Parra, I., Ruiz-Zapata, B., Riera, S., Zapata, L., Ejarque, A., Vegas, T., Rull, V., Scott, L., Andrade, A., Pérez-Díaz, S., Abelschaad, D., Moreno, E., Hernández-Mateo, L., Ochando, J., Pérez Navarro, M.A., Sánchez-Baena, J.J., Riquelme, J.A., Iglesias, R., Franco, F., Chaín, C., Figueiral, I., Grau, E., Matos, M., Jiménez-Espejo, F., Arribas, A., Garrido, G., Finlayson, G., Finlayson, C., Ruiz, M., Pérez-Jordá, G., Miras, Y., 2015. Cinco millones de años de cambio florístico y vegetal en la Península Ibérica e Islas Baleares. Ministerio de Economía y Competitividad, Madrid. <https://play.google.com/store/books/details/JOS%C3%89CARRI%C3%93N.coordinador.CINCO.MILLONES.DEA%C3%91OS.DE?id=JEh1CQAAQBAJ>.
- Carrión, J.S., Ochando, J., Fernández, S., Munuera, M., Amorós, G., Blasco, R., Rosell, J., Finlayson, S., Giles, F., Jennings, R., Finlayson, G., Giles-Pacheco, F., Rodríguez-Vidal, J., Finlayson, C., 2018. Last Neanderthals in the warmest refugium of Europe: palynological data from vanguard cave. Review of palaeobotany and palynology, special issue. In: Carrión, J.S., deMenocal, P., Scott, L. (Eds.), *Human Evolution and Palaeofloras: The Contribution and Potential of Palaeobotany in the Environmental Reconstruction of Hominin-Bearing Sites*, vol. 259, pp. 63–80. *Rev. Palaeobot. Palynol.*
- Carrión, J.S., Fernández, S., Jiménez, J., Munuera Giner, M., Ochando, J., Amorós, G., Ponce de León, M., Zollikofer, Ch., Martín-Lerma, I., Toro-Moyano, I., Hajdas, I., Walker, M.J., 2019a. The sequence at Carrihueta Cave and its potential for research into Neanderthal ecology and the Mousterian in southern Spain. *Quat. Sci. Rev.* 217, 194–216.
- Neanderthals: ecology and evolution. In: Carrión, J.S., Lalueza, C., Stewart, J. (Eds.), *Quat. Sci. Rev. Special Issue*. 217, 1–6.
- Paleofloras, paleovegetation and human evolution. In: Carrión, J.S., Scott, L., deMenocal, P. (Eds.), *Rev. Palaeobot. Palynol. Special Issue*. 267, 32–38.
- Daura, J., Sanz, M., Allué, E., Vaquero, M., López-García, J.M., Julià, R., Ortiz, E., Sánchez Marco, A., Skinner, A.R., Domenech, R., Martinell, J., Arnold, L.J., Carrión, J.S., 2017. Cova del Coll Verdager (Cervelló, Barcelona): the palaeoenvironmental reconstruction of the MIS 3 based on a terrestrial archive and the implications for the last Neanderthals's landscape in the Mediterranean coast of the Iberian Peninsula. *Quat. Sci. Rev.* 177, 34–56.

- Demuro, M., Arnold, L.J., Aranburu, A., Sala, N., Arsuaga, J.-L., 2019. New bracketing luminescence ages constrain the Sima de los Huesos hominin fossils (Atapuerca, Spain) to MIS 12. *J. Hum. Evol.* 131, 76–95.
- De Porras, M.E., Maldonado, A., Latorre, C., Betancourt, J.L., 2017. Late Quaternary environmental dynamics in the Atacama Desert reconstructed from rodent midden pollen records. *J. Quat. Sci.* 32, 665–684.
- Desprat, S., Sánchez Goñi, M.F., Turon, J.-L., McManus, J.F., Loutre, M.F., Duprat, J., Malaize, B., Peyron, O., Peypouquet, J.-P., 2005. Is vegetation responsible for glacial inception during periods of muted insolation changes? *Quat. Sci. Rev.* 24, 1361–1374.
- Desprat, S., Sánchez Goñi, M.F., Naughton, F., Turon, J.-L., Duprat, J., Malaize, B., Cortijo, E., Peypouquet, J.-P., 2007. Climate variability of the last five isotopic interglacials: direct land-sea-ice correlation from the multiproxy analysis of North-Western Iberian margin deep-sea cores. *Dev. Quat. Sci.* 7, 375–386.
- Dimbleby, G.W., 1985. *The Palynology of Archaeological Sites*. Academic Press, London.
- Donner, J.J., Kurtén, B., 1958. The floral and faunal succession of “Cueva del Toll”, Spain. *Eiszeitalt. Ggw.* 9, 72–82.
- Fernández-García, M., López-García, J.M., 2013. Palaeoecology and biochronology based on the rodents analysis from the late pleistocene/holocene of Toll cave (Moia, Barcelona). *Spanish J. Palaeontol.* 28, 227–238.
- Finlayson, C., Carrión, J.S., 2006. Neandertales y humanos modernos en ecosistemas mediterráneos. In: Carrión, J.S., Fernández, S., Fuentes, N. (Eds.), *Paleoambientes y Cambio Climático*. Quaderna. Fundación Séneca, Agencia Regional de Ciencia y Tecnología (Murcia).
- Finlayson, C., Carrión, J.S., 2007. Rapid ecological turnover and its impact on Neanderthal and other human populations. *Trends Ecol. Evol.* 22, 213–222.
- Fiore, L., Gala, M., Romandini, M., Cocca, E., Tagliacozzo, A., Peresani, M., 2016. From feathers to food: Reconstructing the complete exploitation of avifaunal resources by Neanderthals at Fumane cave, unit A9. *Quaternary International* 421, 134–153. <https://doi.org/10.1016/j.quaint.2015.11.142>.
- García-Antón, M., Sainz-Ollero, H., 1991. Pollen records from the Middle Pleistocene atapuerca site (burgos, Spain). *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 85, 199–206.
- García-Mijangos, I., Campos, J.A., Biurrun, I., Herrera, M., Loidi, J., 2015. Marcescent Forests of the Iberian Peninsula: Floristic and Climatic Characterization. Springer, pp. 119–138.
- García-Prieto, E., 2015. *Dinámica paleoambiental durante los últimos 135.000 años en el Alto Jiloca: el registro lacustre de El Cañizar* (PhD Thesis). Universidad de Zaragoza, Spain.
- Gatta, M., Sinopoli, G., Giardini, M., Giaccio, B., Hajdas, I., Pandolfi, L., Bailey, G., Spikins, P., Rolf, M.F., Sadori, L., 2016. Pollen from Late Pleistocene hyena (*Crocota crocata spelaea*) coprolites: an interdisciplinary approach from two Italian sites. *Rev. Palaeobot. Palynol.* 233, 56–66.
- Giardini, M., 2007. Late quaternary vegetation history at stracciaccappa (rome, central Italy). *Veg. Hist. Archaeobotany* 16, 301–316.
- Gil-Romera, G., González-Sampériz, P., Lasheras-Álvarez, L., Sevilla-Callejo, M., Moreno, A., Valero-Garcés, B., López-Merino, L., Carrión, J.S., Pérez Sanz, A., Aranbarri, J., García-Prieto Fonce, E., 2014. Biomass-modulated fire dynamics during the last glacial-interglacial transition at the central Pyrenees (Spain). *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 402, 113–124.
- Girard, M., 1975. Prélèvement d'échantillons en grotte et station de terrain sec en vue de l'analyse pollinique. *Bull. Soc. Préh. Fr.* 72, 158–160.
- Girard, M., Renault-Miskovsky, J., 1969. Nouvelles techniques de préparation en palynologie appliquées à trois sédiments du Quaternaire final de l'Abri Corneille (Istres-Bouches-du-Rhône). *Bull. Assoc. Fr. Et. Quat.* 4, 275–284.
- González-Sampériz, P., 2004. Evolución paleoambiental del sector central de la cuenca del Ebro durante el Pleistoceno superior y Holoceno. Instituto Pirenaico de Ecología-CSIC, Zaragoza, p. 210.
- González-Sampériz, P., Montes, L., Utrilla, P., 2003. Pollen in hyena coprolites from Gabasa cave (northern Spain). *Rev. Palaeobot. Palynol.* 126, 7–15.
- González-Sampériz, P., Valero-Garcés, B., Carrión, J.S., 2004. Was the Ebro valley a glacial refugium for temperate trees? *An. Biol.* 26, 13–20.
- González-Sampériz, P., Leroy, S., Carrión, J.S., García-Antón, M., Gil-García, M.J., Figueiral, I., 2010. Steppes, savannahs and botanic gardens during the Pleistocene. *Rev. Palaeobot. Palynol.* 162, 427–457.
- González-Sampériz, P., García-Prieto, E., Aranbarri, J., Valero Garcés, B.L., Moreno, A., Gil-Romera, G., Sevilla Callejo, M., Santos, L., Morellón, M., Mata, P., Andrade, A., Carrión, J., 2013. Reconstrucción paleoambiental del último ciclo glacial-interglacial en la Iberia continental: la secuencia del Cañizar de Villarquemado (Teruel). *Cuadernos de investigación geográfica*, pp. 49–76.
- González-Sampériz, P., Aranbarri, J., Pérez-Sanz, A., Gil-Romera, G., Moreno, A., Leunda, M., Sevilla-Callejo, M., Corella, J.P., Morellón, M., Oliva, B., Valero-Garcés, B., 2017. Environmental and climate change in the southern central Pyrenees since the last glacial maximum: a view from the lake records. *Catena* 149, 668–688.
- Higham, T., 2014. The timing and spatiotemporal patterning of Neanderthal disappearance. *Nature* 512, 306–309.
- Horwitz, L.K., Goldberg, P., 1989. A study of Pleistocene and Holocene hyaena coprolites. *J. Archaeol. Sci.* 16, 71–94.
- Jiménez-Espejo, F., Martínez-Ruiz, F., Finlayson, C., Paytan, A., Sakamoto, T., Ortega, M., Finlayson, G., Iijima, K., Gallego, D., Fa, D., 2007. Climate forcing and Neanderthal extinction in southern Iberia: insights from a multiproxy marine record. *Quat. Sci. Rev.* 26, 836–852.
- Kromer, B., Lindauer, S., Synal, H.A., Wacker, L., 2013. MAMS-A new AMS facility at the curt-engelhorn-centre for achemetry, mannheim, Germany. *Nucl. Instrum. Methods Phys. Res. Sect. B Beam Interact. Mater. Atoms* 294, 11–13.
- Latorre, C., Betancourt, J.L., Rylander, K.A., Quade, J., 2002. Vegetation invasions into absolute desert: a 45,000 year rodent midden record from the Calama-Salar de Atacama basins, northern Chile (lat 22°/24°S). *Bull. Geol. Soc. Am.* 114, 349–366.
- Lawson, I.T., Frogley, M., Bryant, C., Preece, R., Tzedakis, P., 2004. Lateglacial and Holocene environmental history of the ioannina basin, north-west Greece. *Quat. Sci. Rev.* 23, 1599–1625.
- Loidi, J., 2017. *The Vegetation of the Iberian Peninsula*. Springer. <https://doi.org/10.1007/978-3-319-54784-8>.
- López-García, J.M., Blain, H.-A., Burjachs, F., Ballesteros, A., Allué, E., Cuevas-Ruiz, G.E., Rivals, F., Blasco, R., Morales, J.L., Hidalgo, A.R., Carbonell, E., Serrat, D., Rosell, J., 2012. A multidisciplinary approach to reconstructing the chronology and environment of Southwestern European Neanderthals: the contribution of Teixoneres cave (Moia, Barcelona, Spain). *Quat. Sci. Rev.* 43, 33–44.
- Magri, D., Sadori, L., 1999. Late Pleistocene and Holocene pollen stratigraphy at lago di Vico, central Italy. *Veg. Hist. Archaeobotany* 8, 247–260.
- Magri, D., Di Rita, F., Aranbarri, J., Fletcher, W., González-Sampériz, P., 2017. Quaternary disappearance of tree taxa from Southern Europe: timing and trends. *Quat. Sci. Rev.* 163, 23–55.
- Manzano, S., Carrión, J.S., López-Merino, L., González-Sampériz, P., Munuera, M., Fernández, S., Martín-Lerma, I., Gómez Ferreras, M.C., 2017. Mountain strongholds for woody angiosperms during the late Pleistocene in SE Iberia. *Catena* 149, 701–712.
- Marais, E., Scott, L., Gil-Romera, G., Carrión, J., 2015. The potential of palynology in fossil bat-dung from Arnhem Cave, Namibia. *Trans. Roy. Soc. S. Afr.* 70, 1–7.
- Margari, V., Gibbard, P.L., Bryant, C.L., Tzedakis, P.C., 2009. Character of vegetational and environmental changes in Southern Europe during the last glacial period; evidence from Lesbos Island, Greece. *Quat. Sci. Rev.* 28, 1317–1339.
- Menéndez-Amor, J., Florschütz, F., 1962. Análisis polínico de sedimentos tardi-glaciarios en la Cueva del Toll (Moia, Barcelona). *Estud. Geol.* 18, 93–95.
- Morales-Molino, C., García-Antón, M., 2014. Vegetation and fire history since the last glacial maximum in an inland area of the Western Mediterranean Basin (Northern Iberian Plateau, NW Spain). *Quat. Res.* 81, 63–77.
- Morin, E., Meier, J., El Guennouni, K., Moigne, A.-M., Lebreton, L., Rusch, L., Valensi, P., Conolly, J., Cochard, D., 2019. New evidence of broader diets for archaic Homo populations in the northwestern Mediterranean. *Science Advances* 5, eaav9106. <https://doi.org/10.1126/sciadv.aav9106>.
- Nabais, M., Zilhao, J., 2019. The consumption of tortoise among last interglacial Iberian Neanderthals. *Quat. Sci. Rev.* 217, 225–246.
- Ochando, J., Carrión, J.S., Blasco, R., Fernández, S., Amorós, G., Munuera, M., Sanudo, P., Fernández Peris, J., 2019. Silvicolous Neanderthals in the far west: the mid-pleistocene palaeoecological sequence of bolomor cave (valencia, Spain). *Quat. Sci. Rev.* 217, 247–267.
- Ochando, J., Carrión, J.S., Blasco, R., Rivals, F., Rufá, A., Demuro, M., Arnold, L.J., Amorós, G., Munuera, M., Fernández, S., Rosell, J., 2020a. Neanderthals in a highly diverse, Mediterranean-Eurosiberian forest ecotone: the Pleistocene pollen record of Teixoneres Cave, Northeastern Spain. *Quat. Sci. Rev.* Accepted.
- Ochando, J., Carrión, J.S., Rodríguez-Vidal, J., Jiménez-Arenas, J.M., Fernández, S., Amorós, G., Munuera, M., Scott, L., Stewart, J.R., Knul, M.V., Toro-Moyano, M., Ponce de León, M., Zollikofer, C., 2020b. Palynology and chronology of hyaena coprolites from the piñar karstic caves las ventanas and carihuela, southern Spain. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 552, 109771. <https://doi.org/10.1016/j.palaeo.2020.109771>.
- Pérez-Obiol, R., 1988. Histoire Tardiglaciaire et Holocène de la végétation de la région volcanique d'Olot (N.E. Péninsule Ibérique). *Pollen Spores* 30, 189–202.
- Pérez-Obiol, R., 1994. Análisis polínicos de sedimentos lacustres y de suelos de ocupación de La Draga (Banyoles, Pla de l'Estant). In: Mateu, I., Dupré, M., Güemes, J., Burgaz, M.E. (Eds.), *Trabajos de Palinología Básica y Aplicada*. Universitat de València, València, pp. 277–284.
- Pérez-Obiol, R., Julià, R., 1994. Climatic change on the iberian peninsula recorded in a 30,000 Yr pollen record from lake banyoles. *Quat. Res.* 41, 91–98.
- Pini, R., Ravazzi, C., Reimer, P.J., 2010. The vegetation and climate history of the last glacial cycle in a new pollen record from Lake Fimon (Southern Alpine foreland, N-Italy). *Quat. Sci. Rev.* 29, 3115–3137.
- Postigo-Mijarra, J.M., Gómez-Manzanque, F., Morla, C., 2008. Survival and long-term maintenance of tertiary trees in the Iberian Peninsula during the Pleistocene: first record of *Aesculus* L. (Hippocastanaceae) in Spain. *Veg. Hist. Archaeobotany* 17, 351–364.
- Postigo-Mijarra, J.M., Barrón, E., Gómez-Manzanque, F., Morla, C., 2009. Floristic changes in the iberian peninsula and balearic islands (south-west Europe) during the cenozoic. *J. Biogeogr.* 36, 2025–2043.
- Postigo-Mijarra, J.M., Morla, C., Barrón, E., Morales-Molino, C., García, S., 2010. Patterns of extinction and persistence of arctotertiary flora in Iberia during the quaternary. *Rev. Palaeobot. Palynol.* 162, 416–426.
- Ramírez-Pedraza, I., Tornero, C., Pappa, S., Talamo, S., Salazar-García, D.C., Blasco, R., Rosell, J., Rivals, F., 2019. Microwear and isotopic analyses on cave bear remains from Toll Cave reveal both short-term and long-term dietary habits. *Sci. Rep.* 9, 5716.
- Rivas-Martínez, S., 1987. Memoria del Mapa de Series de Vegetación de España. ICONA, Serie Técnica, Madrid.
- Rivas-Martínez, S., Asensi, A., Díez-Garretas, B., Molero, J., Valle, F., Cano, E., Costa, M., López, M.L., Díaz, T.E., Prieto, J.A.F., Llorens, L., del Arco, M., Fernández, F., Sánchez-Mata, D., Penas, A., Masalles, R., Ladero, M., Amor, A., Izco, J., Amigo, J., Loidi, J., Molina, J.A., Navarro, C., Cantó, P., Alcaraz, F., Bascónes, J.C., Soriano, P., 2007. Mapa de series, geoseries y geopermaseries de vegetación de España. Memoria del Mapa de Series de Vegetación de España. Parte I. It. Geobot, pp. 5–436.
- Rosas, A., 2016. La evolución del género ‘Homo’. Los Libros de la Catarata-CSIC, Madrid.

- Rosell, J., Blasco, R., Morales, J.I., Rivals, F., Rodríguez-Hidalgo, A., Cebrià, A., Camarós, E., 2012. Compartint l'espai: la interacció entre homínids i carnívors al nord-est peninsular (Cova del Toll i Cova de les Teixoneres, Moia, Bages). In: Argemí, M., et al. (Eds.), *Actes I Jornades d'Arqueologia de La Catalunya Central* 2010. Generalitat de Catalunya. Departament de Cultura, pp. 47–51.
- Rosell, J., Blasco, R., Rivals, F., Chacón, M.G., Blain, H.-A., López, J.M., Picin, A., Camarós, E., Rufà, A., Sánchez, C., Gómez, G., Arilla, M., Gómez de Soler, B., Bustos, G., Iriarte, E., Cebrià, A., 2014. Cova del Toll y Cova de les Teixoneres, Moia, Barcelona. In: Sala Ramos, R. (Ed.), *Los cazadores recolectores del Pleistoceno y del Holoceno en Iberia y el Estrecho de Gibraltar: Estado actual del conocimiento del registro arqueológico*. Universidad de Burgos, pp. 302–307.
- Rosell, J., Blasco, R., Rivals, F., Chacón, M.G., Picin, A., Iriarte, E., López-García, J.M., Blain, H.-A., Camarós, E., Arilla, M., Fábregas, J., Rufà, A., Sánchez-Hernández, C., Gómez, G., Gómez de Soler, B., Bustos, G., Cebrià, A., 2015. Els ossos hibernen, els neandertals passen: noves dades sobre els desplaçaments per la Catalunya Central dels grups humans del Paleolític mig a partir dels resultats de les coves del Toll i de les Teixoneres (Moia, Bages). In: Blasco, M.A. (Ed.), *Actes III Jornades d'Arqueologia de la Catalunya Central* 2014. Generalitat de Catalunya. Departament de Cultura, pp. 95–99.
- Rosell, J., Blasco, R., Rivals, F., Chacón, M.G., Arilla, M., Camarós, E., Rufà, A., Sánchez-Hernández, C., Picin, A., Andrés, M., Blain, H.A., López-García, J.M., Iriarte, E., Cebrià, A., 2017. A resilient landscape at Teixoneres cave (MIS 3; Moia, Barcelona, Spain): the Neanderthals as disrupting agent. *Quat. Int.* 435, 195–210.
- Sadori, L., Zanchetta, G., Giardini, M., 2008. Last Glacial to Holocene palaeoenvironmental evolution at Lago di Pergusa (Sicily, Southern Italy) as inferred by pollen, microcharcoal, and stable isotopes. *Quat. Int.* 181, 4–14.
- Sadori, L., Koutsodendris, A., Panagiotopoulos, K., Masi, A., Bertini, A., Combourieu-Nebout, N., Francke, A., Kouli, K., Joannin, S., Mercuri, A.M., Peyron, O., Torri, P., Wagner, B., Zanchetta, G., Sinopoli, G., Donders, T.H., 2016. Pollen-based paleoenvironmental and paleoclimatic change at Lake Ohrid (South-Eastern Europe) during the past 500 ka. *Biogeosciences* 13, 1423–1437.
- Salomón, R., Rodríguez-Calcerrada, J., Zafra, E., Morales-Molino, C., Rodríguez-García, A., González-Doncel, I., Oleksyn, J., Zytowski, R., López, R., Miranda, J.C., Gil, L., Valbuena-Carabana, M., 2016. Unearthing the roots of degradation of *Quercus prenaica* coppices: a root-to-shoot imbalance caused by historical management? *For. Ecol. Manage.* 363, 200–211.
- Sardella, R., Iurino, D.A., Mecozzi, B., Sigari, D., Bona, F., Bellucci, L., Coltorti, M., Conti, J., Lembo, G., Muttillio, B., Mazzini, I., 2019. Grotta Romanelli (Lecce, Southern Italy) between past and future: new studies and perspectives for an archaeo-geosite symbol of the Palaeolithic in Europe. *Geohistory* 11, 1413–1432.
- Scott, L., 1987. Pollen analysis of hyena coprolites and sediments from Equus cave, Taung, southern Kalahari (South Africa). *Quat. Res.* 28, 144–156.
- Scott, L., 1994. Palynology of Late Pleistocene hyrax middens, southwestern Cape Province, South Africa: a preliminary report. *Hist. Biol.* 9, 71–81.
- Serra, R.J., Villalba, J.F., Thomas, J.M., 1957. *Alentours de Barcelone et Moia*. Livret guide des excursions B2-B3. INQUA, V Congrès international Madrid-Barcelona, p. 35.
- Sinopoli, G., Masi, A., Regattieri, E., Wagner, B., Francke, A., Peyron, O., Sadori, L., 2018. Palynology of the last interglacial complex at Lake Ohrid: palaeoenvironmental and palaeoclimatic inferences. *Quat. Sci. Rev.* 180, 177–192.
- Spagnolo, V., Marciari, G., Aureli, D., Berna, F., Toniello, G., Astudillo, F., Boschin, F., Boscato, P., Ronchitelli, A., 2019. Neanderthal activity and resting areas from stratigraphic unit 13 at the Middle Palaeolithic site of Oscurusciuto (ginosa - taranto, southern Italy). *Quat. Sci. Rev.* 217, 169–193.
- Spikins, P., Needham, A., Wright, B., Dytham, C., Gatta, M., Hitchens, G., 2019. Living to fight another day: the ecological and evolutionary significance of Neanderthal healthcare. *Quat. Sci. Rev.* 217, 98–118.
- Stewart, J.R., 2005. The ecology and adaptation of Neanderthals during the non-analogue environment of oxygen isotope stage 3: armageddon or entente? The demise of the European Neanderthals in isotope stage 3. *Quat. Int.* 137, 35–46.
- Stewart, J.R., García-Rodríguez, O., Knul, M.V., Sewell, L., Montgomery, H., Thomas, M. G., Diekmann, Y., 2019. Palaeoecological and genetic evidence for Neanderthal power locomotion as an adaptation to a Woodland environment. *Quat. Sci. Rev.* 217, 310–315.
- Tzedakis, P.C., 1994. Vegetation change through glacial-interglacial cycles: a long pollen sequence perspective. *Philos. Trans. R. Soc. London, Ser. A B* 345, 403–432.
- Tzedakis, P.C., 1999. The last climatic cycle at Kopais, central Greece. *J. Geol. Soc.* 156, 425–434.
- Tzedakis, P.C., Lawson, I.T., Frogley, M.R., Hewitt, G.M., Preece, R.C., 2002. Buffered vegetation changes in a Quaternary refugium: evolutionary implications. *Science* 297, 2044–2047.
- Tzedakis, P.C., McManus, J.F., Hooghiemstra, H., Oppo, D.W., Wijnstra, T.A., 2003. Comparison of changes in vegetation in Northeast Greece with records of climate variability on orbital and suborbital frequencies over the last 450,000 years. *Earth Planet Sci. Lett.* 212, 197–212.
- Uzquiano, P., Ruiz-Zapata, MaB., Gil-García, MaJ., Fernández, S., Carrión, J.S., 2016. Late quaternary developments of mediterranean oaks in the atlantic domain of the iberian peninsula: the case of the cantabrian region (N Spain). *Quat. Sci. Rev.* 153, 63–77.
- Val-Peón, C., Expósito, I., Soto, M., Burjachs, F., 2019. A taphonomic approach to the pollen assemblage from layer M of the Abric Romaní archaeological site (NE Iberian Peninsula). *Rev. Palaeobot. Palynol.* 270, 19–39.
- Wagner, B., Lotter, A.F., Nowaczyk, N., Reed, J.M., Schwalb, A., Sulpizio, R., Valsecchi, V., Wessels, M., Zanchetta, G., 2009. A 40,000-year record of environmental change from ancient Lake Ohrid (Albania and Macedonia). *J. Paleolimnol.* 41, 407–430.
- Wagner, B., Wilke, T., Krastel, S., Zanchetta, G., Sulpizio, R., Reicherter, K., Leng, M.J., Grazhdani, A., Trajanovski, S., Francke, A., Lindhorst, K., Levkov, Z., Cvetkoska, A., Reed, J.M., Zhang, X., Lacey, J.H., Wonik, T., Baumgarten, H., Vogel, H., 2014. The SCOPSCO drilling project recovers more than 1.2 million years of history from Lake Ohrid. *Sci. Drill.* 17, 19–29.
- Wagner, B., Vogel, H., Francke, A., Friedrich, T., Donders, T., Lacey, J.H., Leng, M.J., Regattieri, E., Sadori, L., Wilke, T., Zanchetta, G., Albrecht, C., Bertini, A., Combourieu-Nebout, N., Cvetkoska, A., Giaccio, B., Grahzdani, A., Haufe, T., Holtvoeth, J., Joannin, S., Jovanovska, E., Just, J., Kouli, K., Kousis, I., Koutsodendris, A., Krastel, S., Lagos, M., Leicher, N., Levkov, Z., Lindhorst, K., Masi, A., Melles, M., Mercuri, A.M., Nomade, S., Nowaczyk, N., Panagiotopoulos, K., Peyron, O., Reed, J.M., Sagnotti, L., Sinopoli, G., Stelbrink, B., Sulpizio, R., Timmermann, A., Tofilovska, S., Torri, P., Wagner-Cremer, F., Wonik, T., Zhang, X., 2019. Mediterranean winter rainfall in phase with African monsoons during the past 1.36 million years. *Nature* 573, 256–260.
- Willis, K.J., 1994. The vegetational history of the Balkans. *Quat. Sci. Rev.* 13, 769–788.
- Wood, R.E., Barroso-Ruiz, C., Caparrós, M., Jordá Pardo, J.F., Galván Santos, B., Higham, T.F.G., 2013. Radiocarbon dating casts doubt on the late chronology of the Middle to Upper Palaeolithic transition in southern Iberia. *Proc. Natl. Acad. Sci. Unit. States Am.* 110, 2781–2786.
- Yll, E.I., 1995. Estudi de l'evolució de la vegetació i el clima durant el Tardiglacial i el Postglacial a partir d'anàlisis pollíniques del Delta de l'Ebre i de Menorca. *Publicacions de la Universitat Autònoma de Barcelona. Edició Microfotogràfica. ETD Micropublicacions, SL.*
- Yll, R., Carrión, J.S., Marra, A.C., Bonfiglio, L., 2006. Pollen in late Pleistocene hyena coprolites from san teodoro cave (sicily, Italy). *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 237, 32–39.
- Zilhão, J., Anesin, D., Aubry, T., Badal, E., Cabanes, D., Kehl, M., Klasen, N., Lucena, A., Martín-Lerma, I., Martínez, S., Matias, H., Susini, D., Steier, P., Wild, E.M., Angelucci, D.E., Villaverde, V., Zapata, J., 2017. Precise dating of the middle-to-upper paleolithic transition in Murcia (Spain) supports late neandertal persistence in Iberia. *Heliyon* 3, e00435.