

# Climate and vegetation history of western Portugal inferred from Albian near-shore deposits (Galé Formation, Lusitanian Basin)

ULRICH HEIMHOFER\*†, PETER-A. HOCHULI‡, STEFAN BURLA§,  
FELIX OBERLI¶, THIERRY ADATTE||, JORGE L. DINIS# & HELMUT WEISSERT§

\*Institute of Geology, Leibniz University Hannover, 30167 Hannover, Germany

‡Palaeontological Institute, University of Zürich, CH-8006 Zürich, Switzerland

§Geological Institute, ETH Zürich, CH-8092 Zürich, Switzerland

¶Institute of Geochemistry and Petrology, ETH Zürich, CH-8092 Zürich, Switzerland

||Institute of Geology and Palaeontology, Université de Lausanne, CH-1015 Lausanne, Switzerland

#Departamento de Ciências da Terra, Universidade de Coimbra, 3000 Coimbra, Portugal

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**Abstract** – The late Early Cretaceous greenhouse climate has been studied intensively based on proxy data derived essentially from open marine archives. In contrast, information on continental climatic conditions and on the accompanying response of vegetation is relatively scarce, most notably owing to the stratigraphic uncertainties associated with many Lower Cretaceous terrestrial deposits. Here, we present a palynological record from Albian near-shore deposits of the Lusitanian Basin of W Portugal, which have been independently dated using Sr-isotope signals derived from low-Mg oyster shell calcite.  $^{87}\text{Sr}/^{86}\text{Sr}$  values fluctuate between  $0.707373 \pm 0.00002$  and  $0.707456 \pm 0.00003$ ; absolute values and the overall stratigraphic trend match well with the global open marine seawater signature during Albian times. Based on the new Sr-isotope data, existing biostratigraphic assignments of the succession are corroborated and partly revised. Spore-pollen data provide information on the vegetation community structure and are flanked by sedimentological and clay mineralogical data used to infer the overall climatic conditions prevailing on the adjacent continent. Variations in the distribution of climate-sensitive pollen and spores indicate distinct changes in moisture availability across the studied succession with a pronounced increase in hygrophilous spores in late Early Albian times. Comparison with time-equivalent palynofloras from the Algarve Basin of southern Portugal shows pronounced differences in the xerophyte/hygrophite ratio, interpreted to reflect the effect of a broad arid climate belt covering southern and southeastern Iberia during Early Albian times.

**Keywords:** Albian Stage, Sr-isotopes, palynology, clay mineralogy, palaeoclimate, Portugal.

## 1. Introduction

The Albian Age (112.0–99.6 Ma, Gradstein, Ogg & Smith, 2004) is generally viewed as a time characterized by a long-term increase of global temperatures, finally culminating in the Late Cretaceous thermal maximum (Huber, Hodell & Hamilton, 1995; Pucéat *et al.* 2003; Forster *et al.* 2007). Stratigraphically well-calibrated marine proxy records provide evidence for significant short-term fluctuations in tropical and boreal ocean water temperatures, which occur superimposed on the general warming trend during Albian times (Clarke & Jenkyns, 1999; Bice *et al.* 2006; Erbacher *et al.* 2011). Furthermore, deposition in the Tethys and the evolving North Atlantic Ocean was characterized by the repeated formation of organic-carbon-rich deposits, particularly during earliest and latest Albian times, known as Oceanic Anoxic Events 1b and d, respectively (e.g. Wilson & Norris, 2001; Herrle *et al.* 2003). These short-lasting events had a profound impact on marine biogeochemical cycles and were often associated with significant temperature

anomalies (Erbacher *et al.* 2001; Wagner *et al.* 2007; Hofmann *et al.* 2008). Major marine bio-events taking place in the course of the Albian affect planktonic foraminifera, radiolarians and calcareous nannofossils (Erbacher, Thurow & Littke, 1996; Tiraboschi, Erba & Jenkyns, 2009; Huber & Leckie, 2011). On land, this time interval was associated with the rising dominance of angiosperms in terrestrial ecosystems (Crane, Friis & Pedersen, 1995). The radiation and diversification of early angiosperms during Albian times had a strong impact on the coexisting plant communities, clearly visible in the concomitant decline of gymnosperm pollen and pteridophyte spores (Crane & Lidgard, 1989).

Compared to the marine realm, information on the climatic conditions prevailing on the continents during Albian times is relatively scarce. This results mainly from the difficulties associated with the stratigraphic assignment of Lower Cretaceous continental deposits, essentially caused by the lack of suitable age-diagnostic marker fossils. Furthermore, correlation with stratigraphically well-constrained marine strata is hampered by the frequent occurrence of hiatuses, highlighting the incompleteness of the continental sedimentary record.

†Author for correspondence: heimhofer@geowi.uni-hannover.de

However, some of these problems can be circumvented by studying near-shore marine deposits, which provide terrestrial climate signals in the form of spore-pollen assemblages and clay minerals.

The stratigraphic assignment of shallow marine palynomorph-bearing deposits can be significantly improved by the use of Sr-isotope stratigraphy (Burla *et al.* 2009). Sr-isotope values of low-Mg calcite fossils (e.g. brachiopods, belemnites, bivalves) reflect past seawater  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios (e.g. Jones *et al.* 1994; Veizer *et al.* 1999; Korte *et al.* 2003). Hence, temporal fluctuations in  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios can serve as an independent chronostratigraphic tool. In the last couple of years, the  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios of well-preserved rudist shells have been successfully utilized to date Cretaceous shallow-water carbonate platform deposits lacking accurate biostratigraphic constraints (Steuber *et al.* 2005; Huck *et al.* 2011). In more detritus-influenced depositional systems, the use of  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios from oyster shell low-Mg calcite represents a valuable alternative. A number of studies have demonstrated that oyster shells represent a reliable carrier of the Mesozoic seawater Sr-isotope signal (Jones *et al.* 1994; Denison *et al.* 2003; Burla *et al.* 2009; Schneider, Fürsich & Werner, 2009) and therefore hold a significant potential for the stratigraphic refinement of Mesozoic near-shore successions.

The Lusitanian Basin of western Portugal is a key area for the study of early angiosperm remains. Here, the discovery of a series of exceptionally preserved angiosperm mesofossil floras in the northern part has strongly influenced the current understanding of the evolutionary relationships and biology of early angiosperms (Friis, Pedersen & Crane, 2010) (Fig. 1). In contrast, the southern part of the basin bears near-shore marine deposits, which contain a well-preserved late Early Cretaceous palynoflora including a rich angiosperm pollen assemblage (Groot & Groot, 1962; Medus & Berthou, 1980; Medus, 1982; Trincão, 1990; Heimhofer *et al.* 2005, 2007). Together with the classical Potomac record (Doyle & Robbins, 1977; Hochuli, Heimhofer & Weissert, 2006), the Portuguese angiosperm pollen records from the Rodízio and Galé formations are regarded as a reference succession for the palynological changes observed in mid-latitude angiosperm palynofloras during Albian times (Field, Chatelet & Brodribb, 2009).

The aim of this study is twofold: (1) Establish a refined stratigraphic framework using Sr-isotope stratigraphy based on low-Mg calcite oyster shells for the Albian deposits of the Guincho section; (2) Provide information on the floral community structure and on the climatic history of the area based on total spore-pollen associations and clay mineral data. The integration of the new stratigraphic and palaeoclimatic findings with existing results on the angiosperm palynoflora will provide a better understanding of the temporal and palaeoclimatic context under which the Albian diversification of angiosperms took place.



Figure 1. Map of western Portugal showing the location of the coastal outcrop of the Guincho section. Asterisks correspond to sites bearing Early Cretaceous angiosperm mesofossils. Map modified after Heimhofer *et al.* (2007) and Friis, Pedersen & Crane (2010).

## 2. Geological setting

Lower Cretaceous sediments are well exposed in coastal outcrops of the Lusitanian Basin of Portugal. During Mesozoic times, the opening of the Atlantic Ocean resulted in the formation of a passive margin in western Iberia. In late Early Cretaceous times, thermally-induced basement uplift and isostatic adjustment led to the development of a basin-wide break-up unconformity in the Lusitanian Basin, followed by an episode of continental siliciclastic deposition directly overlying the unconformity (Dinis & Trincão, 1995). The subsequent long-term eustatic rise allowed the re-establishment of near-shore marine conditions with deposition of mixed carbonate-siliciclastic sediments that are the focus of this study.

The sedimentary succession investigated covers more than 125 m of Albian strata and is accessible along the cliffs north of Cabo Raso, about 5 km northwest of the village of Cascais (Fig. 1). Here, an essentially continuous section (referred to as Guincho section) crops out along the beach of Praia de Água Doce between the Forte de Cresmina and the Ponta da Galé (Rey, Bilotte & Peybernes, 1977; Rey, 1992).

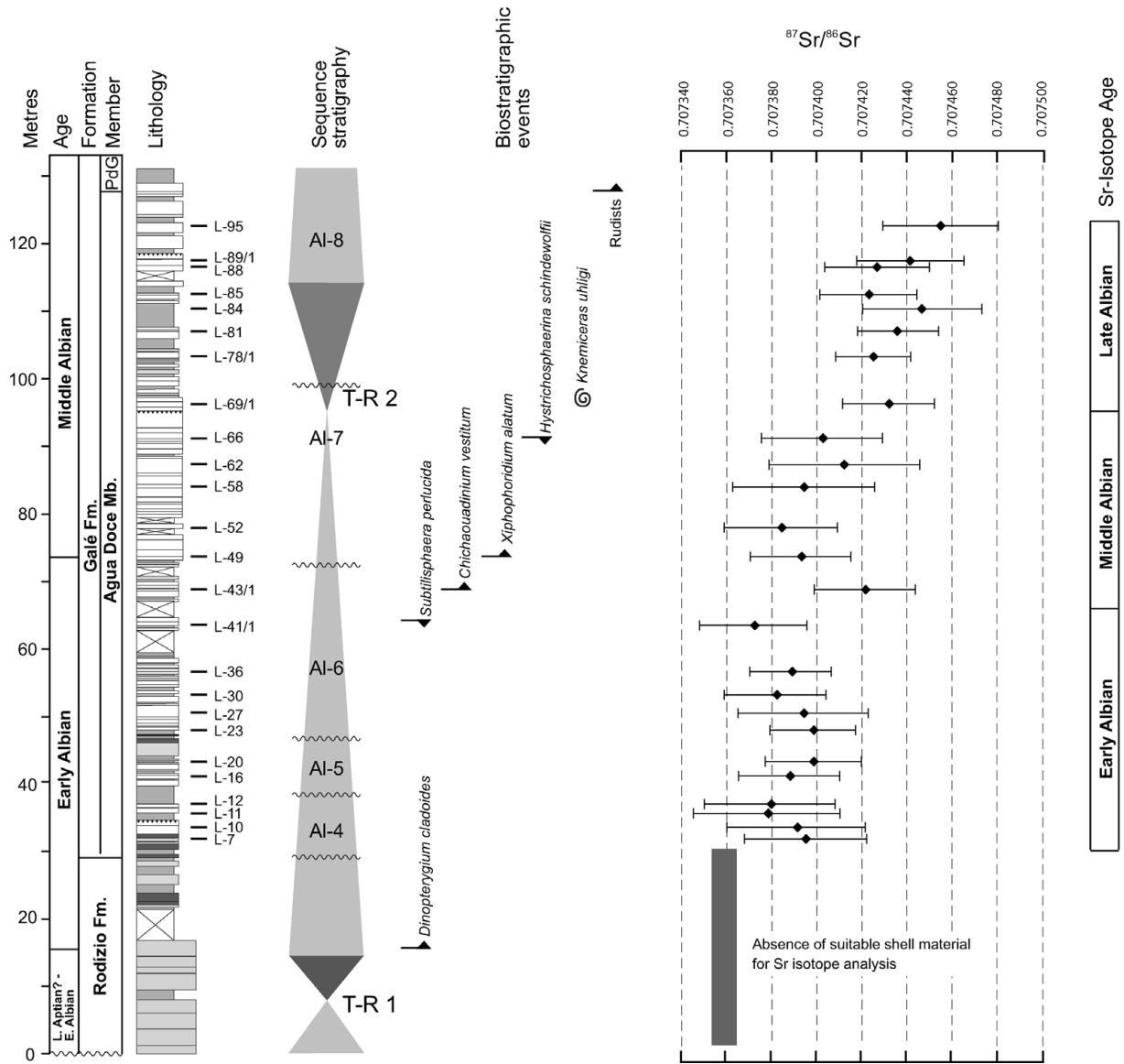


Figure 2. Simplified lithological log and lithostratigraphy of the Guincho section plotted against sequence-stratigraphy, first- and last occurrences of age-diagnostic dinoflagellate cysts and Sr-isotope ratios derived from oyster shell calcite. Both the previous (left-hand column) and the revised age assignment (right-hand column) are shown. Sequence-stratigraphy based on Dinis, Rey & de Graciansky (2002) and Rey (2006); biostratigraphy based on Heimhofer *et al.* (2007). PdG – Ponta da Galé Member.

Following Rey, Bilotte & Peybernes (1977) and Rey (1992, 2006) the section covers the Rodízio and the Galé formations with the latter being subdivided into two major lithological units, the Água Doce and Ponta da Galé members (Fig. 2). The Rodízio Formation covers the lowermost part of the section (0–29 m) and consists of thick-bedded and coarse-grained siliciclastic sediments with thin clayey and lignite-rich interbeds. At the studied locality, two fining-upward cycles can be differentiated, which consist of fluvial large-scale cross-bedded deposits, starting with conglomerates and passing into sandstones. The 18 to 22 m part of the section is currently not cropping out, being covered by beach sands.

The upper part of the Rodízio Formation, making the transition to the Água Doce Member of the Galé Formation, is characterized by a succession (22–29 m)

of reddish and grey sand- and siltstones alternating with purple to greenish claystones interpreted to reflect deposition in peritidal and/or coastal settings (Dinis, Rey & de Graciansky, 2002). This siliciclastic-dominated interval is overlain by the Água Doce Member, formed between 29 m and 71 m by fossiliferous inner shelf marls and sandy limestones, which alternate with varicoloured mudstones and sparse silt- or sandstone beds. The marls and limestones are strongly bioturbated and characterized by a rich macrofossil assemblage including oysters, gastropods, serpulids, thick-shelled bivalves and orbitolinids. Within this interval, a unit composed of thin-bedded ochre-coloured sand- and siltstones can be distinguished (43.5 m to 45.5 m). Abundant ripple structures and a fine lamination pattern as well as the occurrence of highly oxidized wood fragments point to deposition in a shallow, high-energy environment.

At ~ 71 m, the lithologically heterogeneous lower part of the Água Doce Member passes into inner shelf marls and limestones. The subsequent series (71–97.5 m) is dominated by thick-bedded and fossil-rich limestones, alternating with more marly or sand-rich interbeds. The limestones have a very distinct nodular appearance, most probably caused by intensive burrowing. This lithology is rich in large orbitolinids, gastropods and bivalves, whereas oysters, miliolinids and echinid remains are of subordinate importance. The limestone-dominated interval ends abruptly and is overlain by a succession of varicoloured peritidal mud- and siltstones with sandy and marly limestone intercalations (97.5–113 m). In this unit, the top of individual limestone beds is frequently marked by thick-shelled oyster colonies, often preserved in growth position. The greenish to dark-grey mudstones are characterized by millimetre-scale lamination, whereas the marly and sandy lithologies display intense bioturbation. This variegated interval passes again into thick-bedded limestones with thin clayey interbeds (113–131 m). The fossil content includes abundant oysters and other bivalves, echinoderm debris, gastropods, miliolinids and orbitolinids. The uppermost limestone bed (128 m) is characterized by abundant rudists, occurring reworked as well as in growth position, and serves as a distinct marker bed, namely as the lowermost bed of the Ponta da Galé Member.

The investigated succession has been studied in detail from a bio- and sequence-stratigraphic perspective by Rey, Bilotte & Peybernes (1977), Berthou & Schroeder (1979), Berthou & Leereveld (1990), Rey (1992, 2006) and Dinis, Rey & de Graciansky (2002). According to a revised biostratigraphic framework based on dinoflagellate cyst assemblages (Heimhofer *et al.* 2007), the upper part of the Rodízio Formation is characterized by an age not older than Early Albian, not excluding a Late Aptian age for its lower part. The overlying Água Doce Member is considered to be of Early and Middle Albian age. Biostratigraphic evidence derived from dinoflagellate cysts does not provide evidence for Upper Albian strata in the studied part of the succession.

Based on the sequence-stratigraphic interpretation of Dinis, Rey & de Graciansky (2002) and Rey, de Graciansky & Jacquin (2003), the studied section covers at least two second-order transgressive–regressive cycles (Fig. 2). A first progradational–retrogradational sequence is covered by the lower part of continental siliciclastic sediments of the Rodízio Formation. A second sequence is characterized by an extensive transgressive (~ 15–97.5 m) and a short regressive part and covers the upper Rodízio Formation and most of the Água Doce Member. The uppermost part of the studied succession corresponds to the transgressive part of a third second-order cycle, including the top of the Água Doce and the lowermost levels of the Ponta da Galé Member.

### 3. Palaeogeographic and palaeoclimatic framework

During late Early Cretaceous times, the Iberian Peninsula was situated at a palaeolatitude between 25° and 35°N (Sewall *et al.* 2007) and during most of this period formed part of an archipelago, which resulted from the expansion of epicontinental shallow seas during times of high sea-level (Chumakov *et al.* 1995; Hay *et al.* 1999). The Lusitanian Basin formed part of the eastern passive margin of the evolving North Atlantic and was strongly influenced by tectonic movements resulting in several regional unconformities and transgressive–regressive cycles (Dinis *et al.* 2008).

Generally, the late Early Cretaceous is considered a time characterized by exceptional global warmth as expressed in low equator–pole thermal gradients, high atmospheric CO<sub>2</sub> levels and the absence of large continental ice sheets (e.g. Huber, Hodell & Hamilton, 1995; Price, 1999; Haworth *et al.* 2005; Forster *et al.* 2007; Herman & Spicer, 2010). Following a phase of pronounced aridity during the Barremian to Aptian interval (Ruffell & Batten, 1990), also demonstrated to occur on the western Iberian margin (Berthou, Blanc & Chamley, 1982), the southward migration of the equatorial humid belt during Late Aptian to Albian times reflects a significant change in the overall palaeoclimatic patterns (Chumakov *et al.* 1995). According to the palaeoclimatic reconstructions of the latter authors, the southern part of Iberia was located within a broad zone of arid to semi-arid conditions (Northern Hot Arid belt, NHA), even with developed erg systems (Rodríguez-López *et al.* 2008, 2010), whereas the northern part was within the Northern Mid-latitude Warm humid belt (NMW) during Albian times.

Based on findings of plant macro-remains and on palynological assemblages from the Iberian Peninsula, different types of palaeovegetation and associated environments have been distinguished e.g. (Diéguez, Peyrot & Barrón, 2010). These include forest communities dominated by Taxodiaceae–Cupressaceae thriving in riparian or mangrove habitats as well as conifer forests composed of Cheirolepidaceae and tree ferns reflecting more stressed conditions (Peyrot *et al.* 2007). On the other hand, freshwater-related environments like back swamps or riverbanks were populated by hygrophilous plant communities including Bryophyta, Lycophyta and various types of ferns as well as plants assigned to the Cycadales/Ginkgoales/Bennettitales complex and early angiosperms (Coiffard, Gomez & Thevenard, 2007; Villanueva-Amadoz *et al.* 2010). In summary, the vegetation was characteristic of a winter-wet biome under a semi-arid subtropical climate (Diéguez, Peyrot & Barrón, 2010).

### 4. Materials and methods

#### 4.a. Strontium isotope stratigraphy

For the analysis of the strontium isotope composition, a total of 25 visually well-preserved oysters were



collected from the Água Doce Member of the Galé Formation. Oysters are relatively abundant in the detritus-rich shoal-water deposits and inner shelf limestones of the Guincho section, often occurring in the form of regular pavements covering entire bedding planes. Oysters collected from the succession belong to the genus *Exogyra* and *Ostrea* (Rey & Cugny, 1978) and occur throughout the entire Galé Formation providing an essentially continuous coverage of the studied strata. Only the lowermost part of the Água Doce Member and the siliciclastic sediments of the Rodízio Formation were devoid of suitable shell material.

Oysters precipitate a shell consisting of low-Mg calcite and composed of individual foliate and prismatic layers (Carter, 1990). The specific mineralogy makes their calcitic shells relatively robust against subsequent diagenetic alteration processes which may affect the primary chemical composition of the shell material. Therefore, oyster shell calcite is considered a reliable carrier of the seawater Sr-isotope signal as demonstrated by previous studies (Jones *et al.* 1994; Veizer *et al.* 1999; Denison *et al.* 2003; Burla *et al.* 2009; Schneider, Fürsich & Werner, 2009). However, conventional sampling of thin-shelled oysters can result in contamination by adjacent substrate and/or altered shell material or in insufficient sampling amounts. To circumvent this problem, a system combining laser ablation (LA) with multi-collector inductively coupled plasma mass spectrometry (MC-ICP-MS) was used for this study, which yields precise and accurate Sr-isotope data at a high spatial resolution (e.g. Christensen *et al.* 1995; Woodhead *et al.* 2005; Burla *et al.* 2009).

Following visual inspection, the selected oyster shells were carefully cleaned to remove matrix material. Cleaned shells were cut using a diamond saw, placed on a specimen slide and polished. Following conventional petrographic inspection, all shell samples were investigated using a cold cathode luminescence (CL) optical system (Technosyn CL8200 MK II) at ETH Zurich, Switzerland. Only non-luminescent shell portions of the inner, foliate shell layer were used for subsequent laser ablation sampling.

Sr-isotope compositions were measured on a Nu Plasma 1700 MC-ICP-MS coupled to an Excimer 193 nm ArF laser (see Burla *et al.* 2009 for details). Guided by prior CL mapping, typically 15 spots were located on each sample and ablated for 50 seconds each (120  $\mu\text{m}$  spot size). The measurements were internally corrected for mass bias and interference by Rb, Kr and Ca argides, using equations given in Burla *et al.* (2009 and online supporting information). Groups of one to five sample measurements were bracketed by measurements on recent oyster shell material, the latter yielding a mean  $^{87}\text{Sr}/^{86}\text{Sr}$  ratio of  $0.709161 \pm 0.000016$  (external reproducibility at 1 standard deviation,  $n = 16$ ). With the use of these bracketing standards, a renormalization to a value of 0.709175 was carried out on the samples in order to minimize ratio drift and to render the data compatible for comparison with the LOWESS model for seawater Sr evolution

(Howarth & McArthur, 1997; McArthur, Howarth & Bailey, 2001). Precision for runs on pristine samples ranges from  $\pm 0.000008$  to  $\pm 0.000016$  ( $1\sigma$ ; Table 1) and includes an error component from the renormalization procedure. Average external reproducibility calculated from repeated measurements on nine different samples from a companion study (Burla *et al.* 2009) was  $\pm 0.000009$  ( $1\sigma$ ).

#### 4.b. Palynology

A total of 36 rock samples from the Guincho section were prepared for palynological analysis. Despite the selection of apparently well-suited samples (including fresh medium- to dark-grey silt- and claystones), 16 samples were barren of palynomorphs. Cleaned, crushed and weighed samples (20 to 80 g) were treated with HCl and HF following standard palynological preparation techniques (e.g. Traverse, 2007). All 20 productive samples were studied for their sporomorph content (spores and pollen). A minimum of 200 (average of 240) sporomorphs were determined and counted per slide. Light photomicrographs were taken using an Olympus BX 51 light microscope (LM) equipped with an Olympus DP 12 digital camera. The preservation of the studied palynomorphs is fairly good to excellent. Individual grains exhibit no obvious signs of post-depositional degradation. Thermally unaltered preservation of organic matter is indicated by the virtually unchanged colouring of the palynomorphs ( $\text{TAI} < 2$  according to the thermal alteration index of Staplin, 1982).

#### 4.c. Clay mineralogy

A total of 25 samples from the Guincho section were analysed for their clay mineralogy. Clay mineral analyses were based on methods described by Adatte, Stinnesbeck & Keller (1996). Ground chips were mixed with deionized water (pH 7–8) and agitated. The carbonate fraction was removed by addition of 10% HCl at room temperature for 20 minutes or longer until all the carbonate was dissolved. Ultrasonic disaggregation was accomplished during 3 minute intervals. The insoluble residue was washed and centrifuged (5–6 times) until a neutral suspension was obtained (pH 7–8). Separation of two different grain-size fractions ( $< 2 \mu\text{m}$  and 2–16  $\mu\text{m}$ ) was obtained by the timed settling method based on Stokes law. The selected fraction was then pipetted onto a glass plate and air-dried at room temperature. X-ray diffraction (XRD) analysis of oriented clay samples were carried out after air drying at room temperature and under ethylene-glycol solvated conditions. The intensities of selected XRD peaks characterizing each clay mineral present in the size fraction (e.g. chlorite, mica, kaolinite and illite/smectite mixed-layers) were measured for a semi-quantitative estimate of the proportion of clay minerals present in two size fractions ( $< 2 \mu\text{m}$  and 2–16  $\mu\text{m}$ ). Determination of the illite/smectite (I/S)

Table 1. Strontium isotopic results for the Água Doce Member of the Galé Formation

Sample	Fm. member	Height [m]	$^{87}\text{Sr}/^{86}\text{Sr}$	$2\sigma$	LOWESS ages [Ma]			Stage
					Age	Lower limit	Upper limit	
L-7	Água Doce	30.7	0.707396	0.000027	108.6	93.5	110.1	Early Albian
L-10	Água Doce	32.4	0.707392	0.000030	108.8	93.2	110.3	Early Albian
L-11	Água Doce	34.4	0.707379	0.000032	109.4	104.7	110.8	Early Albian
L-12	Água Doce	35.8	0.707381	0.000028	109.3	105.1	110.6	Early Albian
L-16	Água Doce	40.0	0.707389	0.000022	109.0	104.7	110.2	Early Albian
L-20 top	Água Doce	42.1	0.707400	0.000021	108.3	102.7	109.8	Early Albian
L-23	Água Doce	46.6	0.707400	0.000019	108.3	103.4	109.7	Early Albian
L-27 top	Água Doce	49.3	0.707395	0.000028	108.6	93.4	110.2	Early Albian
L-30	Água Doce	52.0	0.707383	0.000022	109.2	105.6	110.3	Early Albian
L-36	Água Doce	55.4	0.707390	0.000018	108.9	105.3	110.0	Early Albian
L-41/1	Água Doce	62.2	0.707373	0.000023	109.6	107.7	110.7	Early Albian
L-43/1	Água Doce	67.6	0.707422	0.000022	104.7	94.6	108.9	Middle Albian
L-49	Água Doce	72.4	0.707394	0.000022	108.7	103.7	108.0	Middle Albian
L-52	Água Doce	76.7	0.707386	0.000025	109.1	104.9	110.3	Middle Albian
L-58	Água Doce	82.7	0.707396	0.000031	108.6	93.3	110.2	Middle Albian
L-62	Água Doce	86.0	0.707413	0.000033	107.0	93.8	109.7	Middle Albian
L-66	Água Doce	89.9	0.707404	0.000026	108.0	93.7	109.8	Middle Albian
L-69/1	Água Doce	95.0	0.707433	0.000020	NA	95.5	108.2	NA
L-78/1	Água Doce	102.0	0.707426	0.000016	103.8	95.3	108.4	Late Albian
L-81	Água Doce	105.8	0.707437	0.000018	NA	95.9	107.7	NA
L-84	Água Doce	109.0	0.707447	0.000026	NA	96.1	107.6	NA
L-85	Água Doce	111.2	0.707424	0.000021	104.2	94.8	108.8	Late Albian
L-88	Água Doce	115.2	0.707428	0.000023	103.4	94.9	108.7	Late Albian
L-89/1	Água Doce	116.2	0.707442	0.000024	NA	95.9	107.7	NA
L-95	Água Doce	121.3	0.707456	0.000025	NA	97.2	105.3	NA

After internal correction for instrumental mass fractionation using  $^{88}\text{Sr}/^{86}\text{Sr}$ , the data have been normalized to a value of 0.709175 based on standard-sample bracketing, using modern oyster shells as standard (for details see Burla *et al.* 2009 and online supporting information). The LOWESS ages (Howarth & McArthur, 1997; McArthur, Howarth & Bailey, 2001) and their bracketing values are derived from Look-Up Table Version 4: 08 / 04, the latter by combining  $2\sigma$  measurement uncertainties with error envelopes of the LOWESS curve. NA – no match of  $^{87}\text{Sr} / ^{86}\text{Sr}$  with the LOWESS curve.

mixed-layers and their content in smectitic expandable layers is based on the method of Moore & Reynolds (1997).

## 5. Results

### 5.a. Sr-isotope ratios of oyster shell calcite

The Sr-isotope data are graphically displayed in Figure 2 and summarized in Table 1. Measured Sr-isotope ratios of oyster calcite from the Água Doce Member of the Galé Formation at Guincho vary between  $0.707373 \pm 0.000023$  and  $0.707456 \pm 0.000025$  ( $2\sigma$ ). The Sr-isotope record shows a distinct trend towards higher Sr ratios as a function of stratigraphic height. The lower part (22 m to 71 m) is characterized by  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios fluctuating between  $0.707379 \pm 0.000032$  and  $0.707400 \pm 0.000021$  with the exception of two outliers in its upper part (samples L-41/1 and L-43/1). On average the middle part of the Água Doce Member (72–90 m) shows slightly higher ratios ranging between  $0.707386 \pm 0.000025$  and  $0.707413 \pm 0.000033$ . The strata of the upper Água Doce Member (91 m to 128 m) display the highest ratios with values between  $0.707424 \pm 0.000021$  and  $0.707456 \pm 0.000025$  with the highest Sr-isotope ratio occurring in the uppermost sample measured (L-95, 121.2 m).

Up to now, a number of similar late Early Cretaceous  $^{87}\text{Sr}/^{86}\text{Sr}$  curves, both in terms of the trend and absolute values, have been published (Bralower *et al.* 1994; Jones *et al.* 1994; McArthur, Howarth & Bailey, 2001). These curves show a distinct minimum in the Late Aptian followed by a subsequent rise towards higher ratios during Albian times. Following a steep increase during latest Aptian and Early Albian times, the curve starts to flatten out during the Middle Albian with the Late Albian being characterized by comparatively stable values reflecting a plateau phase of the seawater strontium trend. Our results obtained on oyster calcite appear to conform to such a pattern both in terms of trend and absolute values.

Converted to numerical ages according to the LOWESS best-fit curve (SIS Look-Up Table Version 4: 08 / 04; Howarth & McArthur, 1997; McArthur, Howarth & Bailey, 2001),  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios range from  $109.4 +1.4/-4.7$  Ma (L-11) to  $108.3 +1.4/-4.7$  Ma (L-23), disregarding samples L41–1 and L43–1. This would indicate an Early to Middle Albian age based on the timescale of Gradstein, Ogg & Smith (2004) for the lower part of the Água Doce Member. The middle part of this member is assigned to the Middle Albian ( $109.1 +1.2/-4.2$  Ma (L-32) to  $107.0 +2.7/-13.2$  Ma (L-62)), whereas its uppermost interval would be Late Albian (104.2 to 103.4 Ma with an error envelope ranging from *c.* 108.8 to 94.8 Ma) in age (Fig. 3).

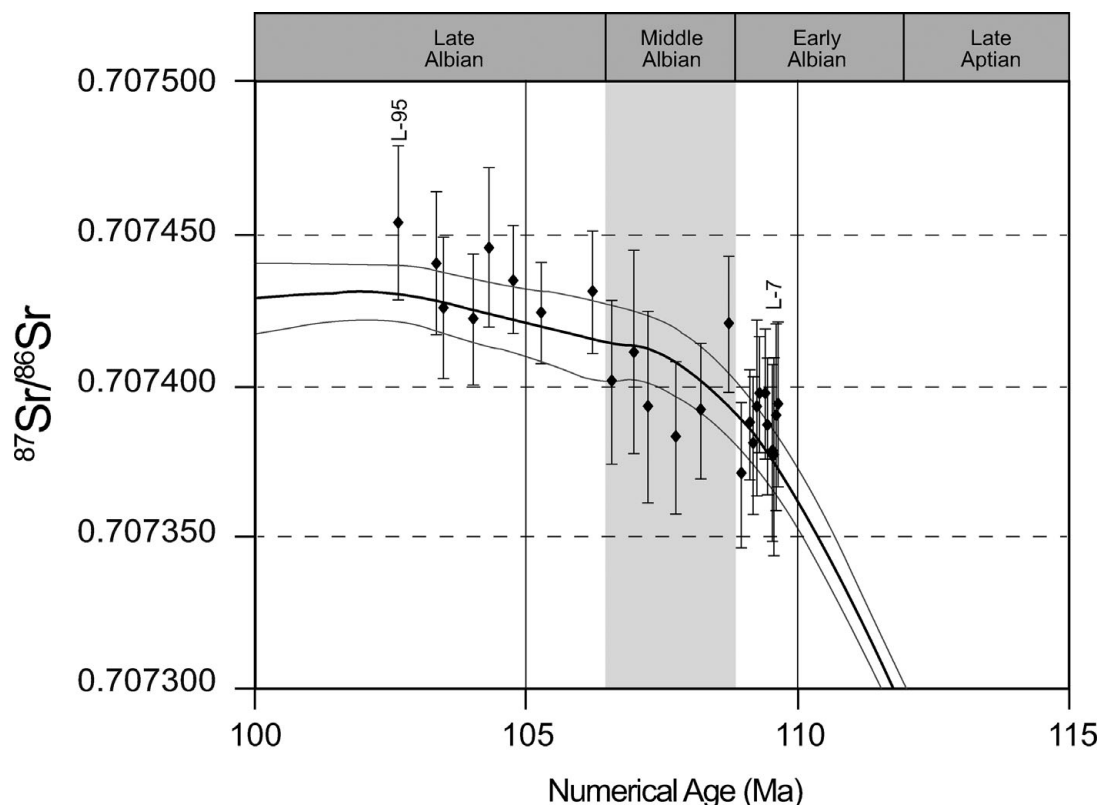


Figure 3. Comparison of LOWESS best-fit curve (black line) and 95 % confidence intervals (thin grey lines) versus Sr-isotope ratios derived from the Guincho section. To reduce the effect of changes in sedimentation rate, the Guincho Sr-isotope record has been subdivided into three stratigraphic intervals (following the revised age assignment to Early, Middle and Late Albian in Fig. 2). The corresponding sets of data points derived from those three stratigraphic intervals have been plotted against numerical age.

However, owing to the flat gradient of the seawater Sr-isotope curve from the Middle Albian onwards, the error derived from the combined uncertainties of the measurements and of the SIS look-up table is considerable and results in significant uncertainties regarding the chronostratigraphic position of this part of the section (Table 1).

### 5.b. Spore-pollen assemblage

The sediments of the Guincho section contain a rich and diverse terrestrial palynoflora composed of various gymnosperm pollen, pteridophyte spores and angiosperm pollen taxa. Numerous spores and gymnosperm pollen were determined on the generic level; other forms were treated as groups (e.g. bisaccate pollen). In the studied record we differentiated 14 genera of gymnosperm pollen (including 7 species) and 20 genera of spores (including 14 species). Angiosperm pollen show a high diversity comprising 50 different forms, which have been previously described in detail by Heimhofer *et al.* (2007). To obtain information on changes in humidity, a xerophyte/hygrophite ratio has been calculated following the concept of earlier studies (Visscher & van der Zwan, 1981; Hochuli *et al.* 2010). In this study, all spores are regarded as hygrophytic plants whereas the xerophyte group is represented solely by *Classopollis* spp.

Based on quantitative distribution of the major pollen groups, four different local pollen zones (LPZ) are distinguished (Fig. 4) from the base to the top of the studied succession. The lowermost LPZ I (0–38.0 m) is characterized by an increase in *Classopollis* spp. increasing from less than 5 % to up to 44 %, high abundances of *Inaperturopollenites* spp. (up to 34 %) and a decline in *Perinopollenites* spp. (from 28 % to less than 10 %) (Fig. 5k, m–q, s, t, x). Other gymnosperm pollen (including bisaccate pollen, *Araucariacites* spp., *Exesipollenites* spp.; Fig. 5a, b, d, j, v, w, y) occur in low numbers, whereas trilete spore abundances vary between 4 % and 17 %. In this interval, the majority of the spores belong to the spore taxa *Cicatricosisporites* spp., *Deltoidospora* spp. and *Gleicheniidites* spp. (Fig. 6a–c, e, f, r, w) and show a mean diversity of seven genera. Pollen assigned to *Afropollis* cf. *jardinus* are a rare but regular component of LPZ I where they account for up to 2 % (Fig. 5aa, bb). Similarly, *Sciadopityspollenites* spp., *Eucommiidites* spp. and *Callialasporites* spp. occur as subordinate floral elements (Fig. 5c, e, f, h, i, r, z). In LPZ I, angiosperm pollen are dominated by monocolpate forms and account for up to 8 % (average 4.1 %). Tricolpate forms are restricted to rare occurrences of *Tricolpites* spp., *Phimopollenites* spp. and *Retitricolpites* spp.

LPZ II includes the interval between 38.0 m and 76.6 m and is characterized by a prominent peak

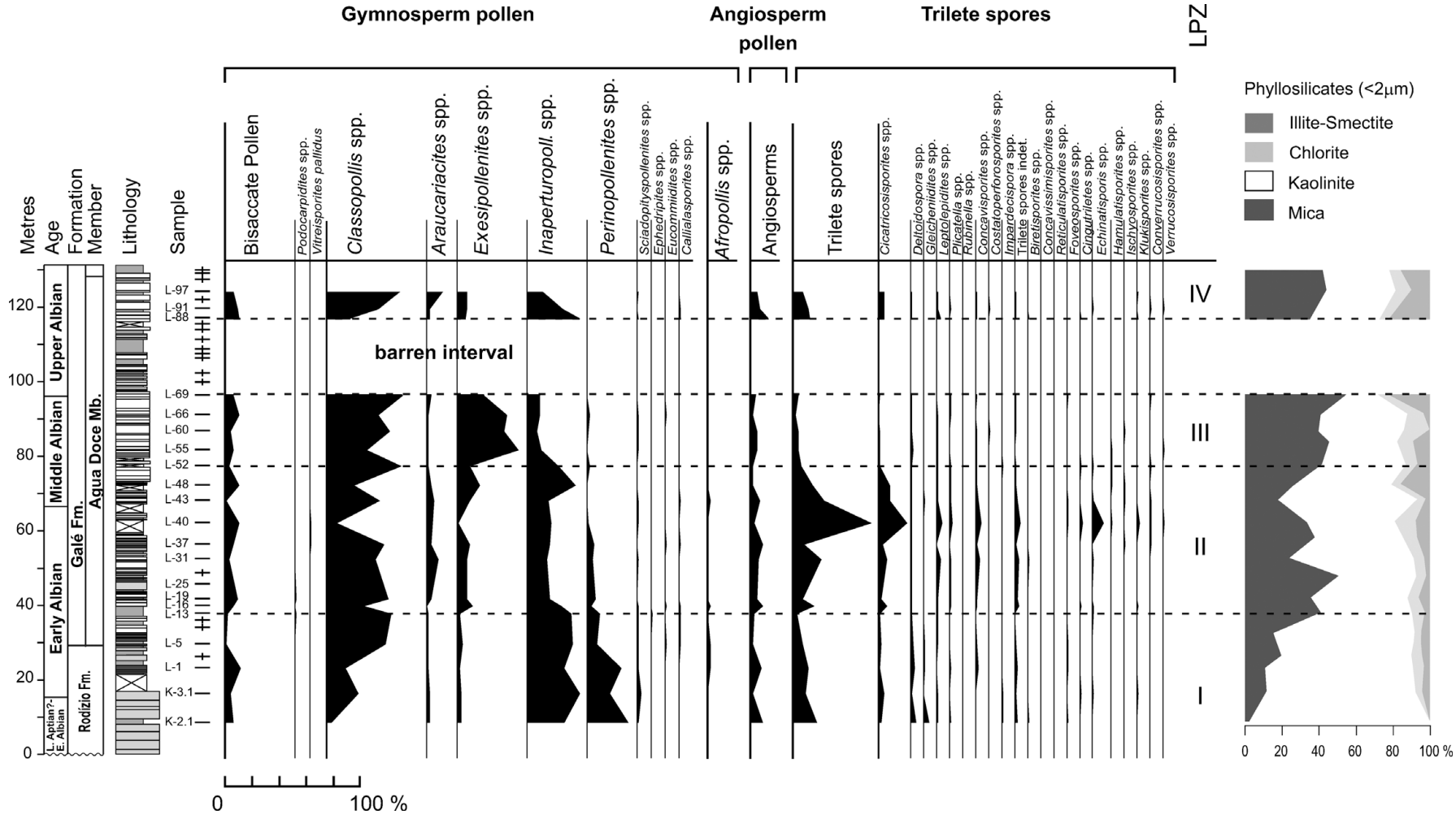


Figure 4. Simplified lithological log, lithostratigraphy, relative abundance [%] of spores and pollen and clay mineral distribution through the Guincho section, Lusitanian Basin, Portugal. LPZ – local pollen zone.



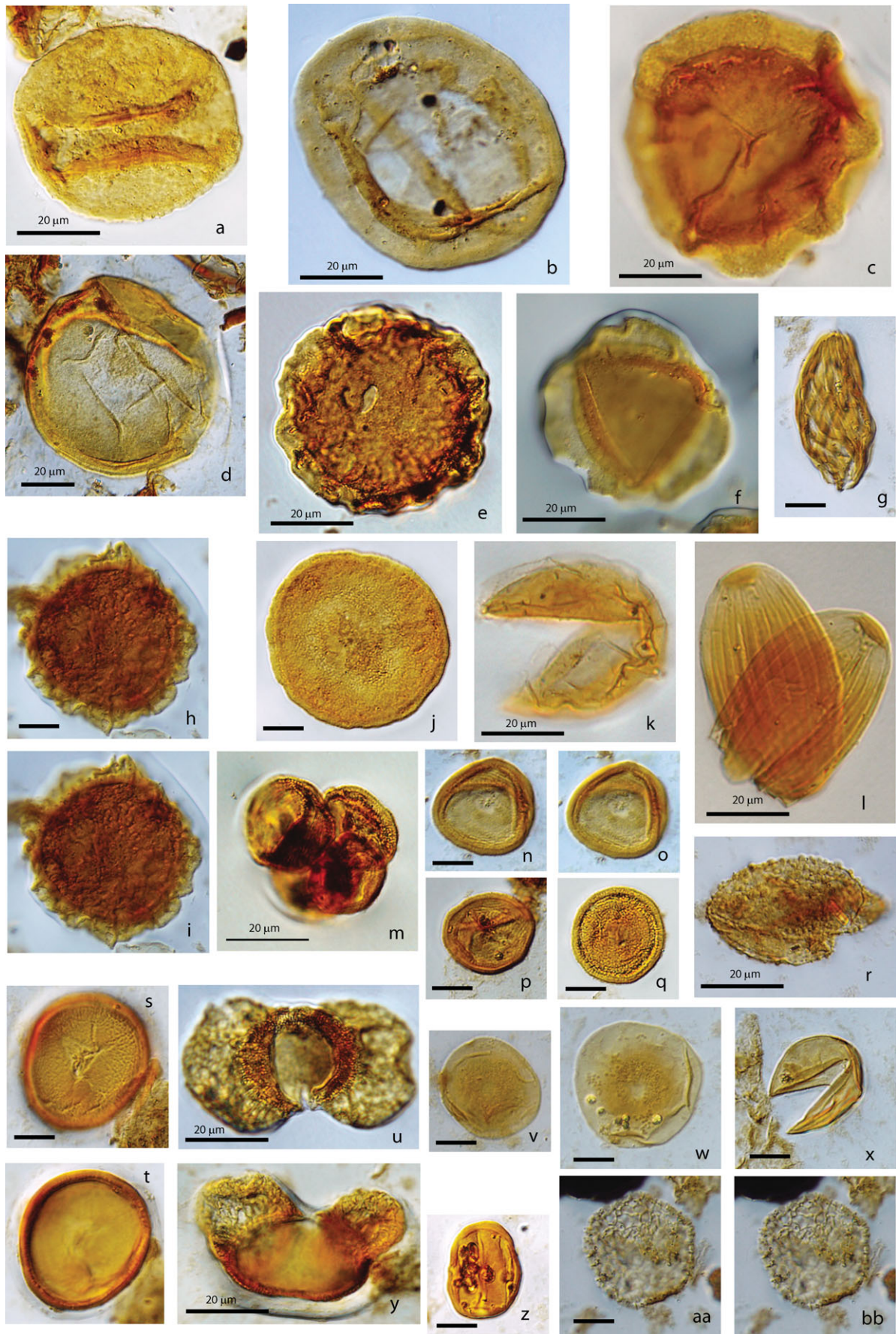


Figure 5. (Colour online) Photomicrographs of selected gymnosperm pollen grains from the Guincho section. Scale bar is 10  $\mu\text{m}$  if not indicated differently. (a) *Araucariacites* sp. (granulate form); L-72-5; 9.0/134.1; (b) *Araucariacites* cf. *australis* Cookson, 1947; L-72-5; 6.7/143.8; (c) *Callialasporites* sp.; L-91.5; 9.2/134.2; (d) *Araucariacites australis* Cookson, 1947; L-66-4; 12.1/145.5;

in trilete spores reaching up to 54%. This increase in spore abundance is most pronounced in the *Cicatricosisporites* group, but can also be observed in spores assigned to *Plicatella* spp., *Leptolepidites* spp., *Concavisorites* spp., *Cingutritetes* spp. and *Echinatisporites* spp. (Fig. 6d, g, h, m, z). Peak abundance of trilete spores is paralleled by the highest spore diversity with 13 different genera. The increase in spore abundance is accompanied by a significant decline in *Classopollis* spp., *Exesipollenites* spp., *Inaperturopollenites* spp. and a further decrease in *Perinopollenites* spp., whereas *Araucariacites* spp. displays a slight increase, reaching up to 8% of the sporomorph count. *Exesipollenites* spp. shows an increase at the beginning of LPZ II to values of up to 11%, but decreases to less than 2% during peak spore abundance. Compared to LPZ I, bisaccate pollen show no significant variations in abundance and fluctuate between 2% and 11%. Noteworthy is the occurrence of *Podocarpidites* spp. (Fig. 5u) and *Vitreisporites pallidus* within LPZ II. Angiosperm pollen represent a relatively common element within this interval (up to 8%; average 4.9%), and the first occurrence of several new tricolpate taxa is observed (including *Artiopollis* spp., *Hammenia* spp. and *Striatopollis* spp.)

The palynoflora of LPZ III (76.6–95.0 m) is characterized by low abundances of *Inaperturopollenites* spp., *Araucariacites* spp. and trilete spores, with the latter group showing a reduced mean diversity (average of 5.8 genera). In contrast, *Classopollis* spp. and *Exesipollenites* spp. show comparatively high abundances, the count of the latter reaching up to 42%. Together with *Sciadopityspollenites* spp. and *Eucommiidites* spp., pollen assigned to *Perinopollenites* spp. are of subordinate importance and account for less than 2% of the assemblage. *Ephedripites* spp. (Fig. 5g) and *Callialasporites* spp. are virtually absent. Compared to LPZ II the relative contribution of angiosperm pollen is slightly reduced within LPZ III and accounts for less than 5% (average 3.4%).

Following a palynologically barren interval between 95.0 m and 115.2 m, the assemblage characterizing the uppermost LPZ IV (115.2–127.8 m) shows an increase in *Classopollis* spp. which is paralleled by a decrease in *Inaperturopollenites* spp. In contrast to the previous LPZ, *Exesipollenites* spp. occurs in low abundances of less than 8%, whereas trilete spores show a return to values of up to 12% (with *Cicatricosisporites* spp. accounting for 5%). *Perinopollenites* spp. is virtually

absent in this zone. Angiosperm pollen shows the highest relative abundance of the entire record reaching up to 12% (average 7.9%).

### 5.c. Clay mineral assemblage

The clay mineral assemblage of the Guincho section (fraction < 2 µm) is dominated to varying degrees by mica and kaolinite whereas chlorite and I/S mixed-layer clay minerals represent only a subordinate fraction (Fig. 4). The relative proportion of the different clay minerals shows distinct variations within the stratigraphic succession. Mica accounts for 30.3% on average and shows a continuous increase throughout the succession ranging from less than 2.4% in the coarse-grained siliciclastic sediments of the Rodízio Formation to reach > 40% at the base of the Ponta da Galé Member. The increasing trend in mica abundance is punctuated by two peaks of 49.5% (at 47.7 m) and 54.0% (at 95.0 m), respectively. Compared to mica, kaolinite shows an opposite pattern with high kaolinite abundances (> 70%) prevailing within the Rodízio Formation and lower part of the Água Doce Member and reduced abundances (~ 40%) above, in most of the Água Doce Member (mean of 56.2%). A distinct minimum of 18.0% is paralleled by peak mica abundances at 95.0 m. Chlorite shows a relatively stable abundance pattern, fluctuating between 2.6% and 14.0% (mean of 6.3%) in most samples (exception of 0% in the lowermost sample). In contrast, the I/S mixed-layers show a slow but distinct increase in abundance from 0.0% at the very base to 24.5% in the uppermost interval (mean of 7.6%). Following the criteria of Moore & Reynolds (1997), identified I/S are of R1–R3 type with 15–20% smectitic expandable layers.

The obtained results agree well with other studies on the clay assemblages of the Upper Aptian and Albian deposits of the region, showing a change from kaolinite to a predominance of illite approximately in Middle Albian times (Berthou, Blanc & Chamley, 1982). Around 100 to 150 km further north in the basin, kaolinite is much more important all over the Figueira da Foz Formation (Upper Aptian to lower Cenomanian) (e.g. Grade & Moura, 1992), with an increase in smectite and I/S mixed-layers in the presumably Upper Albian to lower Cenomanian levels (Rocha & Gomes, 1995).

(e) *Callialasporites dampieri* (Balme) Sukh-Dev, 1961; L-16-3; 8.4/141.0; (f) *Callialasporites trilobatus* (Balme) Sukh Dev, 1961; L-72-5; 11.3/132.5; (g) *Ephedripites* cf. *barghoornii-staplinii* formgroup (Pocock) sensu Herngreen, 1973; L-16-4; 5.0/133.1; (h, i) *Callialasporites* aff. *dampieri* (Balme) Sukh-Dev, 1961; L-16-4; 9.2/121.7; (j) *Araucariacites* sp. (finely echinate form); L-55-5; 12.2/127.2; (k) *Perinopollenites* sp.; L-88-4; 8.0/135.0; (l) *Steevesipollenites* sp.; K-3.1.4; 19.7/142.0; (m) *Classopollis* sp. tetrad; L-55-5; 19.2/138.8; (n, o) *Classopollis* sp.; L-66-4; 7.0/134.3; (p) *Classopollis* sp.; L-55-5; 12.2/134.4; (q) *Classopollis* sp. (finely echinate form); L-55-5; 7.8/141.7; (r) *Sciadopityspollenites* sp.; L-16-4; 9.0/135.7; (s, t) *Classopollis* sp. (smooth form); L-66-4; 9.2/144.8; (u) *Podocarpidites* sp.; L-60-5; 10.3/127.4; (v) *Exesipollenites* sp.; L-66-4; 18.7/130.1; (w) *Exesipollenites tumulus* Balme, 1957; L-66-4; 7.2/144.2; (x) *Inaperturopollenites* sp.; L-66-4; 7.0/137.0; (y) *Pinuspollenites* sp.; L-66-4; 18.7/140.0; (z) *Eucommiidites* sp.; L-66-4; 13.2/130.3; (aa, bb) *Afropollis* cf. *jardinus*; (Brenner) Doyle, Jardiné & Doerenkamp, 1982; L-16-4; 8.0/136.3.



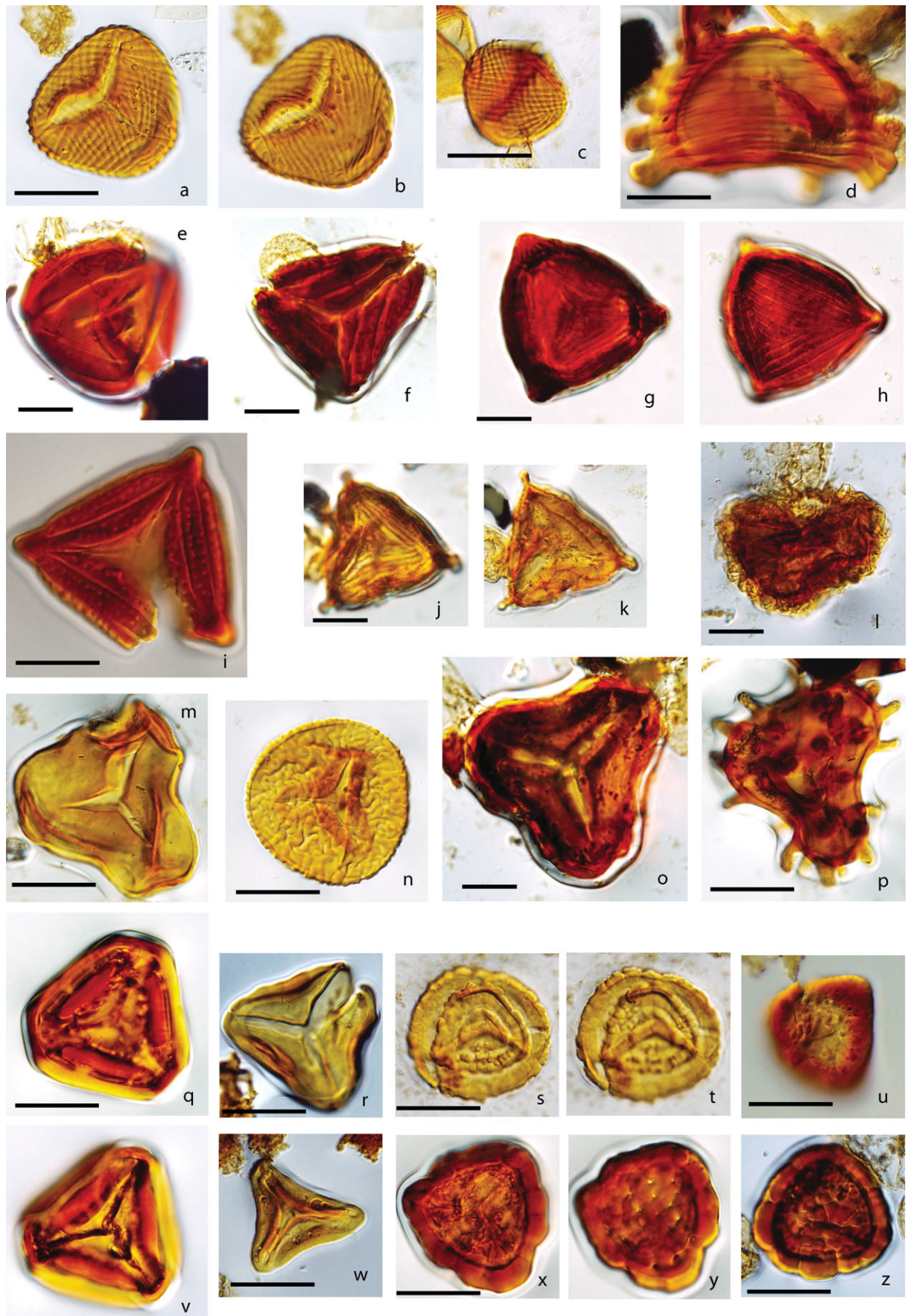


Figure 6. (Colour online) Photomicrographs of selected spores from the Guincho section. Scale bar is 20  $\mu\text{m}$  on all photomicrographs. (a, b) *Cicatricosporites* cf. *hallei* Delcourt & Sprumont, 1955; L-55-5; 16.1/134.7; (c) *Cicatricosporites* cf. *venustus* Deák, 1963; L-66-4; 14.8/146.2; (d) *Plicatella* cf. *crystata* (Markova) Davies, 1985; L-52.6; 10.4/129.0; (e) *Cicatricosporites* cf. *patapscoensis* Brenner, 1963; L-16-4; 12.3/122.2; (f) *Cicatricosporites* sp.; L-16-3; 16.7/124.8; (g, h) *Plicatella* cf. *concentrica* (Kemp) Davies,

## 6. Discussion

### 6.a. Stratigraphic assignment

Generally the  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios derived from low-Mg oyster calcite of the studied Guincho section match the global marine Sr-isotope trend for the Albian Stage, both in terms of the general rise in the ratio and the absolute values (McArthur, Howarth & Bailey, 2001). Correlation with existing Sr-isotope curves from open marine environments (Bralower *et al.* 1997) allows a tentative distinction between Lower, Middle and Upper Albian strata. Comparison with oyster-derived Sr-isotope results from the Portuguese Algarve region generated by the same analytical technique (Burla *et al.* 2009) shows a good match between the two records for the Early to Middle Albian interval, enabling correlation between the two successions. On the other hand, certain discrepancies are observed between the Guincho results and an oyster-derived Sr-isotope record from the Albian of northern Texas, USA (Denison *et al.* 2003). Both records show a distinct increase in  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios for the Early to Middle Albian interval, but both amplitudes and absolute values differ significantly. The steeper gradient and the significantly higher  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios (reaching up to 0.70750) of the Texan oyster record may reflect freshwater influx, which could have shifted the original seawater Sr-isotope composition to more radiogenic ratios (Ingram & Sloan, 1992; Korte *et al.* 2003). A similar mechanism may be invoked to explain the small-scale shifts observed in the Guincho Sr-isotope record, in particular in its lower part. Here, the observed gradual low-amplitude fluctuations may reflect variations in the riverine input of radiogenic Sr derived from continental silicate weathering.

Generally the age assignment based on the isotope record of the Guincho section is in good agreement with existing age interpretations, thus confirming the accuracy of previous stratigraphic studies based on macrofauna and orbitolinids (Rey, Bilotte & Peybernes, 1977; Berthou & Schroeder, 1979), dinoflagellate cysts (Berthou & Leereveld, 1990; Heimhofer *et al.* 2007) and sequence-stratigraphic considerations (Rey, 1992; Dinis, Rey & de Graciansky, 2002; Rey, de Graciansky & Jacquin, 2003). A late Early Albian age for the lower part of the Água Doce Member is corroborated by the consistent occurrence of *Dinopterygium cladoides*.

A distinct offset in the Sr-isotope record in the middle part of the Água Doce Member at  $\sim 95$  m (between samples L-66 and L-69/1) may reflect a sedimentary hiatus. Evidence is provided by the occurrence of a distinct hardground (*discontinuités sédimentaires* in

Rey, 1992), which is considered to reflect the maximum flooding surface of a second-order cycle (Dinis, Rey & de Graciansky, 2002; Rey, de Graciansky & Jacquin, 2003).

Stratigraphic discrepancies exist with regard to the position of the upper part of the Água Doce Member and the lowermost part of the Ponta da Galé Member ( $\sim 100$ – $131$  m; Fig. 2). According to the numerical age assignment derived from the Sr-isotope data, this interval belongs to the early Late Albian (*c.* 104.2 to 103.4 Ma nominal; however, note the error envelope allows for a range of *c.* 108.8 to 94.8 Ma). Similarly, Rey, Bilotte & Peybernes (1977) considered the interval above the bed including the ammonite *Knemiceras uhligi* ( $\sim 95$  m) to be Late Albian in age, and Berthou & Schroeder (1979) indicate an Early to Middle Albian age for the carbonates below  $\sim 85$  m based on the occurrence of the orbitolinid *Simplorbitolina manasiconus*. In contrast, a Middle Albian age has been proposed for the interval above  $\sim 92$  m based on the absence of a characteristic Late Albian dinoflagellate cyst assemblage (Heimhofer *et al.* 2007). However, given the near-shore character of the deposits in this part of the succession, the lack of typical Late Albian dinoflagellate taxa may result from the proximity and restricted character of the depositional setting.

In summary, considering the biostratigraphic constraints and the  $^{87}\text{Sr}/^{86}\text{Sr}$  chronostratigraphic best-fit, the boundary between the Middle and Late Albian is tentatively located in the studied section between 89.9 m (sample L-66) and 95.0 m (sample L-69). The Sr-isotope results of the Guincho section further support the suitability of oyster-derived low-Mg calcite for stratigraphic applications and corroborate earlier biostratigraphic studies.

### 6.b. Significance of near-shore marine palynomorph records

When dealing with sporomorph assemblages obtained from near-shore marine sedimentary deposits, potential effects of hydrodynamic sorting during transportation and deposition have to be considered. Differential sorting due to varying buoyancy or sporomorph size and density can result in significant bias of the studied association (Tyson, 1995; Traverse, 2007). This applies in particular to bisaccate conifer pollen grains, which can remain afloat for weeks explaining the observed relative off-shore increase in bisaccate pollen in modern continental margin sediments (Heusser & Balsam, 1977). In the studied succession the relative abundance

1985; L-16-3; 6.4/140.3; (i) *Costatoperforosporites triangulatus* Deák, 1962; L-55-5; 13.5/128.0; (j, k) *Appendicisporites potomacensis* Brenner, 1963; L-16-3; 12.2/123.7; (l) *Crybelosporites cf. pannuceus* (Brenner) Srivastava, 1977; L-16-3; 18.4/131.2; (m) *Concavisporites* sp.; L-16-4; 8.2/124.9; (n) *Hamulatisporites* sp.; L-55-5; 18.9/132.6; (o) *Impardecispora* sp.; L-16-4; 8.3/126.3; (p) *Trilobosporites apibaculatus* Brenner, 1963; L-16-3; 9.6/131.8; (q, v) *Matonisporites excavatus* Brenner, 1963; L-55-5; 7.9/141.5; (r) *Gleicheniidites cf. senonicus* Ross, 1949; L-60-5; 17.8/136.3; (s, t) *Taurocusporites cf. segmentatus* Stover, 1962; L-66-4; 16.2/146.3; (u) *Cingitriteles cf. congruens* Pierce, 1961; L-88.4; 8.8/131.8; (w) *Gleicheniidites cf. circinidites* (Cookson) Brenner, 1963; L-55-5; 14.8/137.0; (x, y) *Klukisporites* sp.; L-55-5; 9.0/133.0; (z) *Leptolepidites* sp.; L-55-5; 12.2/129.2.



of bisaccate pollen varies between 2 % and 12 %, but does not show significant fluctuations with stratigraphic height. In fact, during Albian times sedimentation in the Lusitanian Basin was affected by distinct sea-level variations (Dinis, Rey & de Graciansky, 2002; Rey, de Graciansky & Jacquin, 2003) as indicated by changes from peritidal (estuary to upper delta) to inner and shallow platform depositional settings (Rey & Cugny, 1978). However, the relatively stable distribution of bisaccates does not show a clear sporomorph sorting due to varying distance from the shoreline. Bias of the quantitative composition of sporomorph assemblages due to differential sorting has also been observed between thin-walled, non-saccate pollen grains and dense, ornamented trilete spores, with the latter being preferentially deposited more proximally, often in the vicinity of a river mouth (Heusser & Balsam, 1977; Tyson, 1995). Again, the Guincho record does not display a strong correspondence between trilete spore abundance and sea-level variations since the exceptionally high spore abundances in LPZ II occur during a long-lasting transgressive phase, even if the minimum in LPZ III coincides with the second-order highest sea-level. The deposits in LPZ II show no sign of enhanced riverine detrital input or sea-level lowering, which could explain the pronounced peak in spore percentages. Hence the effects of hydrodynamic sorting processes are considered to be of minor importance for most of the observed sporomorph distribution patterns. This assumption is in agreement with the view that sporomorph assemblages obtained from marginal marine settings reflect changes in vegetation patterns on the adjacent hinterland rather than sorting during transport and deposition (Tyson, 1995; Abbink, 1998; Traverse, 2007).

#### 6.c. Evolution of vegetation patterns and climate

The composition of the sporomorph assemblages obtained from the Albian deposits of the Guincho section is in good accordance with existing sporomorph records from other parts of Iberia (Groot & Groot, 1962; Medus & Berthou, 1980; Medus, 1982; Taugourdeau-Lantz *et al.* 1982; Solé da Porta *et al.* 1994; Peyrot *et al.* 2007; Diéguez, Peyrot & Barrón, 2010; Villanueva-Amadoz *et al.* 2010). The composition fits well into the palaeophytogeographic scheme of Brenner (1976) in which the occurrence of various types of conifer-derived gymnosperm pollen (e.g. *Classopollis* spp., *Araucariacites* spp., *Inaperturopollenites* spp. and of bisaccate pollen grains of pinacean affinity) and the occurrence of diverse pteridophyte spore associations (including numerous representatives of the Schizaeaceae and Gleicheniaceae) place the Guincho record into the Southern Laurasian floral province. However, minor influence of the Northern Gondwana province adjacent to the south is indicated by the continuous occurrence of typical Gondwanan floral elements including *Afropollis* spp. as well as by the regular occurrence of pollen grains of the *Ephedripites* group.

A position at the southernmost margin of the Southern Laurasian floral province is in good agreement with the concept of a transitional zone between the two major floral provinces, which incorporates floral elements from both provinces to varying extent (Hochuli, 1981).

Based on the presumed ecological preferences and habitat requirements of the parent plants of the observed sporomorph groups, the distribution patterns can be interpreted in terms of changes in palaeovegetation and corresponding environmental conditions. Abbink (1998) and Abbink, Van Konijnenburg-Van Cittert & Visscher (2004) provide a comprehensive compilation of presumed ecologies of the parent plants of Late Jurassic–earliest Cretaceous sporomorph groups, which serves as a basis for the interpretation of the Guincho record. A distinct shift from forest communities dominated by cupressacean and taxodiacean conifers to the dominance of *Classopollis*–(Cheirolepidaceae) is observed in LPZ I (late Early Albian). Conifers of the family Cheirolepidaceae are considered thermophilous and drought-resistant and have thrived in a wide range of habitats, ranging from well-drained upland to coastal lowlands to even saline habitats (Vakhrameyev, 1982; Watson, 1988; Heimhofer *et al.* 2008; Mendes *et al.* 2010). In contrast, Taxodiacean conifers (as indicated by *Perinopollenites*) are interpreted to form part of lush vegetation thriving predominantly in wet lowland environments (Pelzer, Riegel & Wilde, 1992). Hence, the marked increase in cheirolepidacean at the expense of taxodiacean conifers during LPZ I most probably reflects a shift towards more arid conditions in the hinterland of the Lusitanian Basin. This climatic trend is supported by the clay mineral assemblage, where a decline in kaolinite abundance can be observed. Formation of kaolinite is usually associated with intense chemical weathering under humid tropical to subtropical conditions (Gaucher, 1981; Chamley, 1989). An inverse trend back towards a more humid climate characterizes the late Early Albian to early Middle Albian interval of LPZ II. Here, a pronounced peak in pteridophytes, particularly of ferns of the families Schizaeaceae and Matoniaceae as well as lycopsids, is accompanied by a marked decrease in cheirolepidacean conifers and points to the establishment of more humid conditions. High abundances of pteridophytes are characteristic of relatively moist and lush vegetation occurring, for example, along riversides and/or in coastal lowlands (Van Konijnenburg-Van Cittert & Van der Burgh, 1989; Pelzer, Riegel & Wilde, 1992; Abbink, 1998). Peak abundances in pteridophytes are accompanied by *Vitreisporites*-producing pteridosperms, which are typically associated with warm and rather wet habitats (Pelzer, 1984; Van Konijnenburg-Van Cittert & Van der Burgh, 1989). A trend towards a more humid climate may be interpreted from a subtle increase in kaolinite content and decreasing mica abundances, reflecting a shift from physical to predominantly chemical weathering processes. Following this period of comparatively humid conditions, a reverse trend

towards a more arid climate is indicated in LPZ III by a pronounced increase of *Exesipollenites* spp., most probably produced by bennettitalean or cupressacean plants. According to Watson & Sincock (1992) and Hotton & Baghai-Riding (2010) the parent plants of *Exesipollenites* preferred a dry microclimate and were able to withstand a certain saline influence. This climatic interpretation is supported by sedimentological evidence. Reduced riverine siliciclastic input to the basin is indicated by the predominance of carbonate-rich deposits during LPZ III. Similarly, decreasing percentages of kaolinite point to a reduction in chemical weathering rates, most probably owing to limited water availability. A slightly modified picture emerges from the Late Albian interval of LPZ IV, where moderate abundances of pteridophytes occur in association with cupressacean, araucariacean and taxodiacean conifers and angiosperms. A rise in *Classopollis* pollen may indicate increasing dominance of drought-adapted cheirolepidaceans. Increased abundances of I/S mixed-layer minerals point to the erosion of poorly drained soils in the hinterland (Deconinck & Chamley, 1995). The climatic condition prevailing during LPZ IV are probably best described as semi-arid, reflecting conditions in between the enhanced aridity of LPZ III and the humid phase of LPZ II.

#### 6.d. Palaeoenvironmental implications

Evaluation of the palynological data indicates that during Early to Middle Albian times the hinterland of the southern Lusitanian Basin was covered by evergreen vegetation, adapted to semi-humid conditions. Mixed conifer forests, essentially composed of Cheirolepidaceae, Taxodiaceae–Cupressaceae and Araucariaceae, sometimes accompanied by different types of Podocarpaceae and Pinaceae, prevailed. The forest understorey was composed of ferns of the groups Schizaeaceae, Gleicheniaceae and Cyatheaceae/Dicksoniaceae/Dipteridaceae and as well as of lycopsids and early angiosperms. Similar floral assemblages have been described from the Albian of Iberia, from the Basque-Cantabrian basin (Barrón, Comas-Rengifo & Elorza, 2001) and from the Oliete Sub-basin (Peyrot *et al.* 2007).

However, the stratigraphic succession of the palynofloras reflect some marked changes in the plant communities, including the demise of taxodiacean conifers (LPZ I), a pronounced increase in pteridophyte spores, particularly of Schizaeaceae (LPZ II) and a rise in Cupressaceae (LPZ III) as well as an overall increase in angiosperm pollen throughout the studied section. Whereas the increase in the abundance of angiosperm pollen is interpreted to reflect the incipient radiation of this group during Albian times (Medus & Berthou, 1980; Heimhofer *et al.* 2005, 2007) observed variations in the distribution of gymnosperm pollen and pteridophyte spores are best explained as a response of vegetation to changes in regional climatic conditions, in particular to moisture availability. Changes in

humidity during the studied time interval are well expressed in the xerophyte/hygrophite ratio (Fig. 7). The dataset indicates significant variations in the temporal distribution of moisture availability, with an increase in regional humidity during late Early Albian times. Coeval deposits from the Algarve Basin, located ~ 200 km towards the south, show no evidence for increased humidity, neither in the palynoflora nor in the clay mineral composition (Heimhofer *et al.* 2008). Here, the palynological assemblages are strongly dominated by *Classopollis* pollen, resulting in a constantly high xerophyte/hygrophite ratio throughout the Early Albian interval (Fig. 7).

Late Aptian to early Cenomanian deposits in the northern part of the western Iberian margin include very high proportions of kaolinite and are interpreted as a braided fluvial system (Dinis *et al.* 2008), pointing to a climate that was, at least seasonally, wet. In contrast, sedimentological evidence for relatively arid conditions prevailing in eastern Iberia during Albian times is derived from the presence of a subtropical erg-margin system (Rodríguez-López *et al.* 2008, 2010). Sand sea development resulted from a northward shift of the northern subtropical high-pressure system, probably linked to the opening of the equatorial Atlantic gateway (Poulsen, Gendaszek & Jacob, 2003). Arid palaeoclimatic conditions prevailing in southern and eastern Iberia during latest Early Cretaceous times are in line with the palaeogeographic reconstructions of Chumakov *et al.* (1995). According to these authors, the boundary between the so-called Northern Hot Arid belt and the Northern Mid-latitude Warm humid belt divides the Iberian Peninsula in an E–W direction. The position of the studied Guincho section close to the boundary of the two belts made this area sensitive to climatic changes resulting in expansion or contraction of the latitudinally arranged climatic zones. Hence, the occurrence of a more hygrophilous vegetation during late Early Albian times may reflect a transient expansion of the Northern Mid-latitude Warm humid belt towards the south, probably driven by the rearrangement of atmospheric or oceanic heat transport.

#### 7. Conclusions

(1) The near-shore marine deposits of the Guincho section of the Lusitanian Basin, W Portugal were deposited in Early to early Late Albian times as indicated by Sr-isotope data derived from low-Mg oyster calcite. The new Sr-isotope record corroborates the existing biostratigraphic assignment of the lower Água Doce Member as being Early to Middle Albian in age. However, high  $^{87}\text{Sr}/^{86}\text{Sr}$  values between  $0.707424 \pm 0.00002$  and  $0.707456 \pm 0.00003$  characterizing the uppermost Água Doce Member, point to an age not older than early Late Albian. The Sr-isotope data obtained using laser-ablation multi-collector ICP-MS highlight the suitability of low-Mg oyster shell calcite

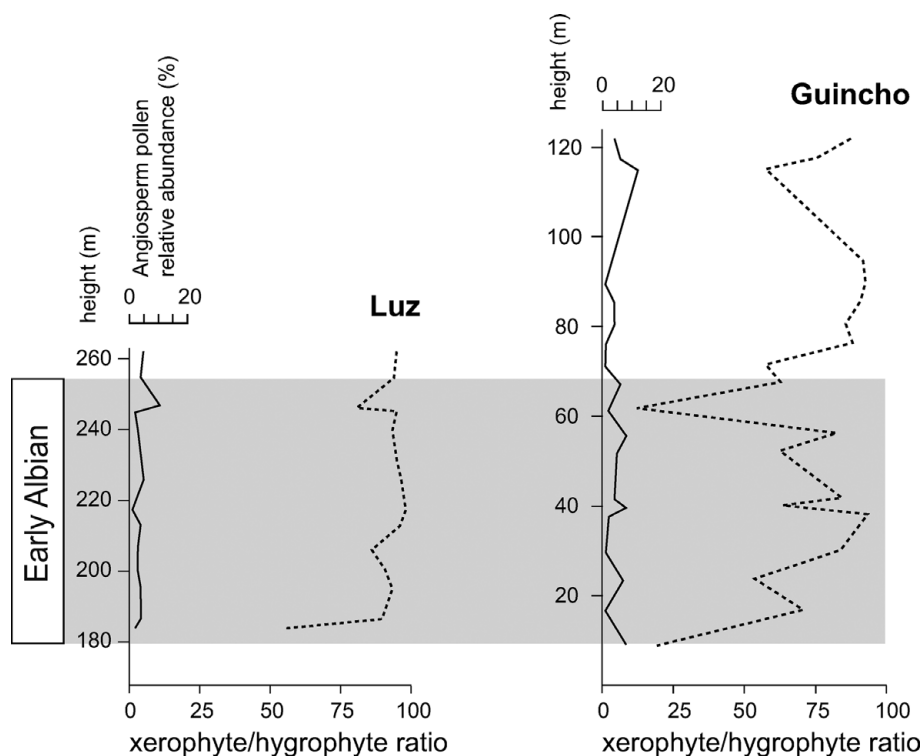


Figure 7. Comparison of the xerophyte/hygrophyte ratios (dashed line) and angiosperm pollen relative abundances (solid line) between the Early Albian records of Guincho (Lusitanian Basin, W Portugal) and the Luz (Algarve Basin, S Portugal). Xerophyte/hygrophyte ratio of the Luz record is based on Heimhofer *et al.* (2008), angiosperm pollen abundance is based on Heimhofer *et al.* (2007), and stratigraphic assignment follows Burla *et al.* (2009).

for the age assignment of Cretaceous near-shore marine deposits.

(2) The sedimentary deposits are characterized by a rich and diverse palynoflora essentially composed of conifer-derived gymnosperm pollen, trilete pteridophyte spores and angiosperm pollen. According to the spore-pollen data, the hinterland of the Lusitanian Basin was covered by an evergreen vegetation composed of mixed conifer forests (including Cheirolepidaceae, Taxodiaceae–Cupressaceae, Araucariaceae and subordinately Podocarpaceae, Pinaceae) with the understorey being composed of various ferns (Schizaeaceae, Gleicheniaceae, Cyatheaceae, Dicksoniaceae, Dipteridaceae), and a relatively diverse angiosperm assemblage. This vegetation is interpreted to correspond to semi-humid conditions.

(3) Across the stratigraphic succession, the spore-pollen record shows some distinct fluctuations in assemblage composition. Similarly, the clay mineral assemblages vary with stratigraphic height. Important changes in the plant community structure include the demise of taxodiacean conifers (LPZ I), a pronounced increase in pteridophyte spores (LPZ II) and a rise in bennettitalean or cupressacean-derived pollen (*Exesipollenites* spp.; LPZ III). In concert with the clay mineral data, these changes in the distribution patterns of gymnosperm pollen and pteridophyte spores are interpreted as reflecting changes in regional climate patterns, particularly in moisture availability. Distinct differences in the xerophyte/hygrophyte ratio between the Guincho record and data from the southern Algarve

Basin are interpreted as reflecting the position of gross climatic belts, with southern and southeastern Iberia being under the influence of the northern subtropical high-pressure system during Early Albian times.

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