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New insights into the Barremian–lower Aptian calcareous nannofossils of the Mediterranean Tethys: chronostratigraphic and paleobiogeographic implications Roque Aguado^{a,*} raguado@ujaen.es, Miguel Company^b, Luis O'Dogherty^c, José Sandoval^b, Mathieu Martinez^d

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Abstract

A detailed study of calcareous nan rofossil assemblages from twelve uppermost lower Barremian—lower Aptian sertions in the Subbetic domain of the Betic Cordillera was performed. Seven nev na. nofossil species (*Crucibiscutum bastetanum*, *Crucibiscutum gracile*, *Chiastozygus i mprostauros*, *Cyclagelosphaera platyaspis*, *Lithraphidites aichmoides*, *Lithraphidites pugio*, and *Rhagodiscus sicutclipeus*) are described, one species is emended (*Lithraphidites magnus*) and the taxonomic concept of the marker species *Hayesites irregularis* is discussed and clarified. The detailed stratigraphic ranges of the new species, together with those of other relevant taxa, are determined and correlated to standard ammonite biostratigraphy. Age estimates of biostratigraphically relevant calcareous nannofossil biohorizons are calculated using astrochronologically tuned cyclostratigraphic data. Five new calcareous nannofossil subzones are proposed

which enhance upper Barremian biostratigraphic resolution at a regional scale and are directly correlated with respect to the standard Tethyan ammonite zonation. Two of the new species described here are used as biostratigraphic markers for the newly proposed subzones. The duration of each subzone is provided through astrochronological calibration. This study allows the refinement of the calcareous nannofossil zonation for the Mediterranean–Atlantic province of the Tethyan domain. The implications of these new results are discussed regarding the extant definition and use of the Barremian/Aptian boundary. The morpho-evolutionary trends of selected nannofossil groups are reviewed in relation to the latest Barremian–Aptian paleogeographic changes, showing these were a prominent factor controlling calcareous nannoplankton evolution and biogeographical distribution in the vest European-Atlantic region.

Keywords: Taxonomy; Bio rec graphy; Bio-chronostratigraphy; Barremian—Aptian boundary; Betic Cordillera

1. Introduction

The Barremian Stage (~125 °8 to ~121.40 Ma; Martinez et al., 2020) records a particular interval marked by notable environmental and biotic changes. These include two episodes of accelerated environmental change (e.g., Föllmi, 2012): the Mid-Barremian Event (MBE; Coccioni et al., 2003; Sprovieri et al., 2006; Yilmaz et al., 2012; Huck et al., 2013; Aguado et al., 2014a) and the Taxy Episode (TE; Masse and Machhour, 1998; Moullade et al., 1998a; Föllmi, 2012; Wissler et al., 2002; Godet et al., 2006; Stein et al., 2011; Martinez et al., 2020). Both episodes were linked to carbon-cycle perturbations locally related to enhanced preservation of organic-rich marine sediments.

One of the most convenient ways to identify these episodes is through their C-isotope imprints, as their lithological expression in different basins may vary depending on the local depositional and environmental conditions (e.g., Jenkyns, 2010). The calibration of the resulting C-isotope curves to biostratigraphy, chronostratigraphy, magnetostratigraphy and to geochronology will greatly impact global correlations (e.g., Ogg et al., 2016; Olierook et al., 2019; Gradstein et al., 2020; Martinez et al., 2020; Castro et al., 2021; Zhang et al, 2021).

Tethyan ammonite biostratigraphy (e.g., Reboulet et al., 2018) represents, in the Barremian, a high-precision reference scale to which visiotope data could be tied. In the absence of ammonites, biostratigraphic scales in each on calcareous nannofossils, planktonic foraminifera and/or radiolarians) may be used (e.g., Thierstein, 1971, 1973, 1976; Sissingh, 1977; Roth, 1978; Perch-Nielsen, 1985; Coccioni and Premoli-Silva, 1994; Jud, 1994; O'Dogherty, 1994; Taralower et al., 1995; Premoli-Silva et al., 2018). However, the biostratigraphioneous function of these microfossil groups is currently much lower compared to that observed using ammonites. In addition, the calcareous nannofossil zonation, have a low resolution with regard to counterpart schemes developed for the West European Province of the Boreal Realm (e.g., Jakubowski, 1987; Bown et al., 1998; Jeremiah, 2001). This may be due, in part, to the spread of sedimentary facies unfavorable for a good preservation of the taxa (i.e., limestones) in the Tethyan Realm, leading to a poor investigation in land sections where nannofossils can be correlated to other reference groups such as ammonites.

The Barremian successions of argillaceous limestones and marlstones of the Subbetic Domain in the Betic Cordillera are well dated by ammonites (e.g., Company et al., 1992, 1995, 2003; Aguado et al., 1992, 1997), and proved to contain abundant and moderately to well-preserved calcareous nannofossil assemblages (e.g., Aguado et al., 1997, 2014a). Some of these successions were also studied for C-isotope stratigraphy, both MBE and TE were identified in them (e.g., Aguado et al., 2014a; Martinez et al., 2020), and they have been recently calibrated by astrochronology (Martinez et al., 2020). These successions represent good candidates to improve Tethyan calcareous nannofossil biostratigraphy by providing adequate chronos, attgraphical and geochronological frameworks to tie all the biostratigraphic events.

The aim of this paper is to improve the resolution of the Barremian calcareous nannofossil biozonation for the Meditarranean so it can be used as an alternative to ammonite zones. The stratigraphic range of the calcareous nannofossil species with biostratigraphic potential, together with that of other relevant taxa, will be directly correlated to standard ammonite biostratigraphy and to the geochronologic scale. Finally, as the Barremian Annian boundary is included within the interval studied, it will be discussed in the harbor of new findings.

2. Geological setting

All sections studied belong to the Subbetic Domain (Fig. 1), a complex tectonostratigraphic unit that paleogeographically corresponds to the pelagic areas of the southern passive margin of the Iberian Plate during the Alpine tectonic cycle (Triassic to early Miocene). During the Jurassic and Early Cretaceous this region underwent extensional tectonics associated with the seafloor spreading of the North Atlantic in a

transform continental margin setting (Vera, 2001, 2004; Martín-Chivelet et al., 2019). A paleolatitude of ~26–28° N (Fig. 2) has been inferred for the Barremian–early Aptian position of the Subbetic Domain (e.g., Barron et al., 1981; Stampfli and Kozur, 2006; Cao et al., 2017; Barrier et al., 2018), being located near the boundary between the arid-tropical and humid-temperate climatic belts (e.g., Masse et al., 1993; http://www.scotese.com/ecretcli.htm; Fig. 2).

Except for X.Cp₂ section (see Chapter 3), all sections studied belong to the widely extended Lower Cretaceous Carretero Formation (Figs. 5, -1). Roughly, the lithologic succession of this formation (e.g., de Gea, 2004) consists of a rhythmic alternation of yellowish to gray argillaceous limestone beds (5–°9 cm thick) and gray marlstone interbeds (3–760 cm thick). The lime fraction is partially made up of calcareous nannofossil remains and carbonate particles of micritic size (micarbs) which probably originated from the adjacent platforms. Clay minerals are, by far, the main components of the detrital fraction (e.g., Agradese al. 2008). Overlying the Carretero Formation, some sections include parts of the Argos, Fardes, or Carbonero formations (Figs. 3, 4; see also Supplementary Neterial 1 (SM1) for details on lithostratigraphy and biostratigraphy).

3. Material and methods

A total of 575 samples, spread across 12 sections were studied. These include, from West to East (Fig. 1):

- Caprés (A in Fig. 1. Section X.Cp₂; 38.233078°N, 1.134206°W; 16 samples; Fig. 4).
- Arroyo de Gilico (B in Fig. 1. Section X.V₁; 38.159931°N, 1.678208°W; 77 samples; Fig. 3).

- Barranco de Cavila (C in Fig. 1. Section X.Kv: 38.051514°N, 1.885713°W; 57 samples; Fig. 3. Section X.Kv₂: 38.052469°N, 1.888964°W; 222 samples; Fig. 3).
- Río Argos (D in Fig. 1. Section X.Ag6: 38.051514°N, 1.885713°W; 15 samples; Fig. 4).
- Rambla Seca (E in Fig. 1. Section RA03: 37.794622°N, 2.030255°W; 32 samples; Fig. 3).
- Cerro Trompeta y Cortijo del Hielo (F in Fig. 1. Section X.CT: 37.866290°N, 2.575907°W; 8 samples; Fig. 4. Section X.HA: 37.851783°N 2.588921°W; 29 samples; Fig. 4).
- Barranco de las Azadillas (G in Fig. 1. Section X.7: 57 779300°N, 2.708924°W; 12 samples; Fig. 4).
- La Frontera (H in Fig. 1. Section X.F: 37.591)35°N, 3.602561°W; 56 samples; Fig. 4. Section X.F1: 37.593226°N, 3.603831°W, 39 samples; Fig. 4).
- La Coronilla (I in Fig. 1. Section X.CO₂: 37.546093°N, 3.937009°W; 13 samples; Fig. 4).

Although sections X.Cp₂, PA.33, X.CT, X.HA, X.F, X.F1 and X.CO₂ have been studied previously (see SM₁, new smear-slides were prepared and studied for all samples. Samples from X.V₁, X.Kv, X.Kv₂, X.Ag6 and X.Z sections (370) are new. The number of samples studied in each section and their respective positions are stated in figures 3, 4, and in Supplementary Material 2 (SM2). The sampling intervals were highly variable (Figs. 3 and 4), from 6–7 cm (e.g., sections X.V₁ and X.Kv₂) to several meters (e.g., lower part of X.F section). Except for samples from the RA03 section and those from the uppermost Barremian to lower Aptian parts of the X.F and X.F1 sections, the remainder of them are directly correlated to ammonite zones/subzones (Aguado, 1994;

Aguado et al., 1992, 1997, 2017; Company et al., 1995, and this paper). In the present work, the standard ammonite zonation for the Mediterranean Province (Reboulet et al., 2018) is followed, and used as a reference to calibrate the calcareous nannofossil bioevents. In the absence of magnetostratigraphical data, the base of the *Deshayesites oglanlensis* ammonite Zone (AZ) is used here to determine the base of the Aptian (Reboulet et al., 2011, 2018).

Simple permanent smear slides (Bown and Young, 1998) were counted with coverslips for routine biostratigraphic analysis and examined for nanctossil content using a polarizing light microscope Olympus BHSP at 1200 × agnification. At least a complete longitudinal traverse (= 200 fields of vie v; y: reface area of one field = 2.37 \times 10⁻² mm²) was studied, but in some slides s xx (a) (up to 4) longitudinal traverses were studied. Special care was taken to prepare the smear slides as uniformly as possible, so that particle density on the slide surpose was kept between 30–50% (Baccelle and Bosellini, 1965). To determine the tratigraphic position of the horizon from which the wide-canal nannoconids are nore abundant than narrow-canal nannoconids (wc>nc Event) in some sections (\$\lambda 4/\dagger 3, X.F, X.HA, X.Kv and X.Kv2), nannoconids were counted in 50 fields of view on samples selected according to their biostratigraphic position. The percentages of wide canal (including Nannoconus bucheri, N. circularis, N. truittii; N. sp. cf. N. truittii, N. vocontiensis and N. wassallii) and narrow canal (including Nannoconus bermudezii and N. steinmannii) forms were calculated (see SM2). The used taxonomic framework is based on Perch-Nielsen (1985), Bown (1998, 2005), Aguado et al. (1997, 2014b, c), and the nannotax website (http://www.mikrotax.org/Nannotax3/index.php?dir=Mesozoic).

Preservation in the assemblages was evaluated in each sample (see SM2) based on the visual criteria established by Roth and Thierstein (1972) and Roth (1983). Limestone beds were avoided for sampling, as they contain impoverished (poorly to moderately preserved) assemblages with moderate to heavily overgrown taxa. Overall abundance per sample was estimated by counting the total number of nannofossils in 20 random fields of view.

Photomicrographs of calcareous nannofossils were acquired with an Olympus Camedia C5050 camera attached to the Olympus BHSP microscope, using cross-polarized light (XPL). A wide variability of size was observed for sont taxa recorded throughout the studied interval. To test whether size of specific tata fuctuates throughout the studied interval, specimens of *Lithraphidites* spp. (120) *Mayesites irregularis* group (140) and *Flabellites oblongus* (120) were photographed and measured. In addition, some specimens (6–22) of the new species described were also photographed and measured. The results are documented in the £ M2. Measurements were taken using ImageJ software, with an accuracy of ± 0.03 µm. Regarding precision, in a set of 10 repetitive measurements of 10 specimes, the error was lower than ±0.14 µm with a 95% confidence level. The lengths (L) and widths (W) of *Lithraphidites* spp., *H. irregularis* gr. and *F. oblongus* specimens were later used to test whether these three groups consist of several taxa.

To test the supposed multimodality in the populations of the measured taxa, mixture analyses were applied to the whole data sets, (i.e., all specimens of each group pooled together) using the free PAST v4.05 software package (Hammer et al., 2001). This statistical analysis is a maximum-likelihood method for estimating the descriptive

parameters (mean, standard deviation and proportion) of two or more distinct distributions, based on an initially pooled univariate sample (Hammer and Harper, 2006). The minimum values of the Akaike Information Criterion (AIC, Akaike, 1974) helped to identify the groups obtained by mixture analysis with the lowest overfitting (Hammer et al., 2001). The number of bins for histograms in mixture analyses follows the Sturges' rule ($k = (\log_2 n) + 1$), where k is the number of bins and n the number of observations).

4. Results

4.1 Nannofossil abundance and preservation

Calcareous nannofossil abundance usually fluctuates cetween > 5 and 30 specimens per field of view (SFOV), with only discrete samples showing abundances below 5 SFOV, while 22 samples were barren (see SN.2). Nannofossil assemblages are rich and moderately to well preserved, showing slight to moderate overgrowth except for those from the organic-rich beds of X + Ag = And + X + Z = And + Ag = And + And + Ag = And + Ag = And + Ag = And + And +

4.2 Mixture analyses

Fluctuations observed in length (L) and width (W) in the pool of the measured specimens of *Lithraphidites* spp. (Fig. 5A–D) suggest that several species can be differentiated. The results of a mixture analysis performed on the W values (Fig. 5E) reveal three different-sized groups. The first group corresponds to *L. carniolensis* (Fig. 5A), while the second and third group correspond to *L. aichmoides* (Fig. 5B; Chapter 6) and to the *L. pugio/L. magnus* plexus respectively (Fig. 5C, D). The same statistical analysis was applied to the length (L) of all specimens with W greater than 4.5 µm (*L*.

pugio/L. magnus plexus). The frequency histogram (Fig. 5G) reveals the existence of two different-sized groups, which represent the new species *L. pugio* and *L. magnus* (Fig. 5C, D; Chapter 6). Figure 5H shows the descriptive parameters resulting from the mixture analysis performed on W in the whole data set and on L in those specimens having a W greater than 5 μm. However, total length (L) proved not to be an outright criterion to separate the latter two species, being more effective distinguishing both taxa using the L/W ratio (Fig. 6; see also Chapter 6).

Hayesites irregularis is an important marker species in the Tethyan Realm (e.g., Thierstein, 1973; Perch-Nielsen, 1985; Bralower et al., 1995). Its taxonomic clarification can increase the biostratigraphic precision of stratigraphic correlations. Variations observed in the length (L) and wide (CV) of the specimens measured in this study (Fig. 7A, B) suggest the presence of two morphotypes in the *H. irregularis* group (gr.). This was corroborated by the results of a mixture analysis on the L/W ratio of the complete pool of measurements (Fig. 7C). Forms with a nearly circular outline, (L/W smaller or equal to 1.12), being younger in time, were assigned to *H. irregularis* sensu stricto (s.str.). Specimens having a more elongated outline (L/W>1.12) and poorly defined elements, being slightly older, were assigned to *H. irregularis* morphotype E (*H. irregularis* E; Chapter 6).

Mixture analyses performed on size measurements (L and W) of specimens of Flabellites oblongus sensu lato (s.l.) spread across the uppermost lower Barremian lower Aptian of the study sections (see SM2 and Fig. 8) provided non-conclusive results. These showed similar AIC values for unimodal/bimodal solutions suggesting a

rather continuous increase in size through the complete data pool (Fig. 8) thereby hindering a clear separation of different populations (see Chapter 6).

4.3. Sequence of nannofossil bioevents

One hundred and four taxa were identified through the complete interval studied, including 7 new species (see SM2 and Chapter 6). The following sequence of bioevents (from bottom to top) has been observed:

- Presence of *Braarudosphaera hockwoldensis* (*Holcod^{*}scu^{*} fallax* ammonite subzone [ASz]).
- Lowest occurrence (LO) of Flabellites oblongus (h. lcodiscus caillaudianus ASz).
- LO of Lithraphidites aichmoides (Mouton'ce 'as moutonianum ammonite zone [AZ]).
- Successive LOs of *Phosterolithus prossir*, Crucibiscutum gracile and Lithraphidites pugio (Gassendiceras alpinum ASz).
- LO of Rhagodiscus sicutclipeus (Avermost part of the Gerhardtia sartousiana ASz).
- Successive LOs of *Chiasto vgus lamprostauros*, *Micrantholithus stellatus* and *Rhagodiscus gallagheri* (Car ardtia provincialis ASz).
- LO of *Hayesites irrequaris* E, highest occurrences (HOs) of *Lithraphidites pugio* and *Lithraphidites aichmoides*, followed by the successive LOs of *Lithraphidites magnus*, *Crucibiscutum bastetanum* and *Nannoconus* sp. cf. *N. truittii* (*Hemihoplites feraudianus* ASz).
- LO of *Hayesites irregularis* s. str. (*Imerites giraudi* AZ)
- LO of *Stoverius acutus*, followed by the wc>nc Event and the HO of *Crucibiscutum* bastetanum (lower part of the *Martelites sarasini* AZ).
- LOs of Nannoconus truittii and Rhagodiscus angustus (Deshayesites oglanlensis AZ).

- Presence of *Conusphaera rothii* and *Micrantholithus stellatus* (lower part of the *Deshayesites forbesi* AZ).

A summary of the main bioevents recorded and their stratigraphic context is shown in Figure 9.

5. Discussion

5.1. Biostratigraphic remarks

Although most of the taxa shown in Figure 9 have stratigran in value, several of them (e.g., Chiastozygus lamprostauros, Crucibiscutum gracile, Lithraphidites magnus, Nannoconus sp. cf. N. truittii, Phosterolithus prosiii, Phagodiscus sicutclipeus and Stoverius acutus) are rare and/or have a spotty and discontinuous record (drawn in dashed line in Fig. 9) throughout a sui stantial part of their respective stratigraphic ranges.

Other taxa have unclear LO. due to the presence of transitional morphologies in the stratigraphic record. Such in the case of the LOs of *Rhagodiscus gallagheri* (obscured by the presence of transmonal forms from small *Rhagodiscus asper*) and *Lithraphidites magnus* (difficult to differentiate from transitional specimens from *Lithraphidites pugio* in the upper part of the *Hemihoplites feraudianus* ASz). The LO of *Nannoconus truittii* (upper part of *Deshayesites oglanlensis* AZ) is masked by the record of forms similar to this species reported here as *Nannoconus* sp. cf. *N. truittii* (see Chapter 6: Figures 3, 4, and range charts of SM2). Finally, specimens of *Rhagodiscus gallagheri* transitional to *Rhagodiscus angustus* were observed from the upper part of the *Martelites sarasini* AZ. The oldest forms assignable to *R. angustus* were recorded from the uppermost part of

the *Deshayesites oglanlensis*/lowermost *Deshayesites forbesi* AZs upwards, which agrees with the observations of Rutledge and Bown (1996).

Other bioevents, such as the FOs of *Flabellites oblongus* and *Hayesites irregularis* have been used as biostratigraphic markers (e.g., Thierstein, 1973, 1976; Bralower et al.1995), but they were never adequately correlated to ammonite biostratigraphy. The LO of *F. oblongus* was recently identified in the lower part of the *Moutoniceras moutonianum* AZ (Aguado et al., 2014a; Martinez et al., 2020). Here we report the LO of *F. oblongus* from the *Holcodiscus caillaudianus* ASz in the X.Kv and X.V₁ sections. This species is very rare and shows a spotty record throughout the *H. caillaudianus* ASz and lowermost part of the *M. moutonianum* AZ. It accesistent record (Figs. 3, 9) correlates with the level where Aguado et al. (2014a) and Martinez et al. (2020) reported the LO of this species. *Hayesites irregularis* E is rather rare near its LO and is more abundant from the upper part of the *Hemihoplites feraudianus* ASz. The LO of *H. irregularis* s.str. (Chapter 4.3 and Fig. 9) was not recorded until the mid part of the *Imerites giraudi* AZ (Figs. 2, 4; SM2).

Finally, a small grou₁ or taxa have shown fairly continuous and consistent records. These include *Lithraphidites pugio*, whose LO was already successfully used in regional correlation (as *L.* sp. cf. *L. magnus*) by Martinez et al. (2020), *Micrantholithus stellatus*, and the distinctive *Crucibiscutum bastetanum* (Fig. 9 and SM2). In addition, the stratigraphic horizon from which the proportion of wide-canal nannoconids outnumber the narrow-canal nannoconids (wc>nc Event, Figs. 3, 4, 9; sections X.HA, X.Kv, X.Kv₂ and SM2), has also proven to be a good correlation event. In some sections (RA03, X.F, X.F1), this horizon is masked by a stratigraphic break surface (see

Figs. 3, 4, and SM2). The onset of the 'nannoconid decline' event (e.g., Erba et al.,

1999, 2019; Channell et al., 2000; Erba 2004; Tremolada et al., 2006) was not

determined in this study.

5.2. Proposed biozones

Here we document a detailed zonation of the upper Barremian-lower Aptian interval of

the Mediterranean area. The zonation uses the scheme of Bralower et al. (1995) by

adding new subzones as digits. The proposed subzones are proposed on a regional scale

(e.g., Martinez et al., 2020), but they could be applied to the Mediterranean–Atlantic

area, as some of the marker species were also recorded from outside of the Betic

Cordillera (chapters 5.4, 6).

NC5E1 Subzone. (New subzone)

Base: LO of Flabellites oblongus. Calibrated here to 124.72 Ma, slightly earlier than in

Martinez et al. (2020). This bioever also corresponds to the base of the NC5E Subzone

(Bralower et al., 1995).

Top: LO of *Lithraphidites prizio* (calibrated to 123.57 Ma).

Age and stratigraphic range: Latest early Barremian—earliest late Barremian (upper part

of the *Holcodiscus caillaudianus* ASz–lower part of the *Gassendiceras alpinum* ASz).

This subzone has a duration of 1.15 myr following the chronostratigraphy of Martinez

et al. (2020).

NC5E2 Subzone. (New subzone)

Base: LO of *L. pugio*.

Top: LO of *Micrantholithus stellatus* (calibrated to 122.64 Ma).

Age and stratigraphic range: Mid-part of the late Barremian (lower part of the *G. alpinum* ASz–upper part of the *Gerhardtia provincialis* ASz). Duration of 0.93 myr according to Martinez et al. (2020).

NC5E3 Subzone. (New subzone)

Base: LO of M. stellatus.

Top: LO of *Hayesites irregularis* E (calibrated here to 122.45 Ma), slightly earlier than in Martinez et al., (2020).

Age and stratigraphic range: Mid-part of the late Barrer nan. (upper part of the *G. provincialis* ASz–lower part of the *Hemihoplites feraucianus* ASz). Duration of 0.19 myr following Martinez et al. (2020).

NC6A1 Subzone. (New subzone)

Base: LO of *H. irregularis* E.

Top: HO of *Crucibiscutum bastota.* " n (calibrated here to 121.97 Ma).

Age and stratigraphic range. Upper part of the late Barremian (lower part of *H*. feraudianus ASz–lower part of Martelites sarasini AZ). Duration of 0.48 myr following Martinez et al. (2020)

NC6A2 Subzone. (New subzone)

Base: HO of C. bastetanum.

Top: HO of Conusphaera rothii.

Age and stratigraphic range: Latest Barremian–early Aptian (lower part of the *M. sarasini* AZ–*Hedbergella excelsa*). This subzone has a duration much greater than 0.97 myr following Martinez et al. (2020).

5.3. Calcareous nannofossil bioevents, C-isotope stratigraphy and the Barremian/Aptian boundary: an update

Currently, the boundary between the Barremian and Aptian Stages is provisionally located at the base of the CM0r magnetochron, and the Gorgo a Cerbara section is its possible Global Stratotype Section and Point (Erba et al., 1996). This designation was made based on the supposed proximity of the lower boundary of CM0r to the LO of deshayesitid ammonites (=base of the Aptian used by ammonite *pecialists; e.g., Reboulet et al., 2018). However, magnetostratigraphic record is very poorly preserved in hemipelagic sections where ammonite zonations are established and, to date, no direct calibration of magnetic polarity and ammonite zones is so far available for the upper Barremian–lowermost Aptian interval.

The LO of *Hayesites irregularis* has been reported in Gorgo a Cerbara from ~6 m below the base of CM0r (Patruno et al. 2014). New findings (Martinez et al., 2020, and present paper) indicate that be LO of *H. irregularis* gr. occurs in the lower part of the *Hemihoplites feraudianus* ASz (see chapters 4.3 and 5.1), below that reported by Aguado et al. (1997). This is consistent with the stratigraphy of Gorgo a Cerbara proposed by Frau et al. (2018), and indicates that the LO of *H. irregularis* is far below the base of the Aptian as defined by the first appearance of deshayesitids.

The wc>nc Event (e.g., Erba et al., 1999; Larson and Erba, 1999; Channell et al., 2000; Bellanca et al., 2002; Tremolada et al., 2006; Erba et al., 2019) was recorded in the Gorgo a Cerbara section within CM0r (Channell et al., 2000; Patruno et al., 2015). This horizon has been here correlated to the lower part of the *Martelites sarasini* AZ (Figs. 3,

4, 9 and SM2). This change in the proportions of narrow/wide canal nannoconids requires time-consuming analyses to be determined, and probably is not a good alternative as a primary marker to be used in the definition of the base of the Aptian, but it could be used as a secondary indicator at least in the Tethyan Realm.

The HO of the new coccolith species *Crucibiscutum bastetanum* follows the wc>nc Event and slightly predates the onset of the C-isotope Intra-Sarasini Negative Excursion (ISNE) linked to the Taxy Episode (Föllmi, 2012; Frau, 2020: Nartinez et al., 2020). According to the chronology of Martinez et al. (2020), this bioevent predates the Barremian/Aptian boundary by 0.57 myr, being a moderately good approximation to it. However, this new species has been identified only in the Subbetic Domain as yet.

The LOs of *Nannoconus truittii* and *Kras odiscus angustus* (upper part of *Deshayesites oglanlensis*—lower part of *Deshayesur's forbesi* AZs) do not provide reliable events because they are masked by the presence of specimens with transitional morphologies (see chapters 5.1 and 6). No other reliable calcareous nannofossil event has been recorded from the Betic Cordillera, until the 'nannoconid crisis', which has been correlated to the upper part of the *D. forbesi* AZ (Aguado et al., 1999; Moreno-Bedmar et al., 2009, 2012).

The C-isotope ISNE, has been identified in several sections within the mid–upper part of the *Martelites sarasini* AZ (Kuhnt et al., 1998; Moullade et al., 1998a, b; Godet et al., 2006; Frau, 2020; Martinez et al., 2020). Its top is close to the base of the Aptian as defined by ammonite biostratigraphy. In sections from SE France and Spain (see also Sanchez-Hernandez and Maurrasse, 2016), the ISNE is made of two spikes with lower

C-isotope values separated by a short excursion with slightly higher values (Fig. 10). In the Gorgo a Cerbara section the ISNE consists of only one spike in the δ^{13} C curve, which is located at the top of CM0r and correlates with the change from the 'Maiolica' to the 'Marne a Fucoidi' formations (e.g., Godet et al., 2006; Stein et al., 2011; Frau et al., 2018).

Figure 10 summarizes the findings of the current study around the Barremian–Aptian interval and correlates the X.Kv₂ section and some relevant French and Italian sections by using bio- and chemostratigraphy. Given the proximity of the ISNE (especially its top) to the base of *D. oglanlensis* AZ, we think this negative excursion of the δ^{13} C_{carb} is an alternative for defining the base of the Aptian.

5.4. Barremian—Aptian calcareous natinofossil morpho-evolutionary trends and paleobiogeography

Although the driving factors for siz hanges of calcareous nannofossils have been discussed in the last decades, both for living assemblages and for fossil species, these remain still unclear. In living assemblages, some link was found within coccolith morphology and environmental factors such as sea-water temperature and salinity (e.g., Bollmann and Klaas, 2008; Bollmann et al., 2009; Triantaphyllou et al., 2010). In deeper time, studies are more problematic due to difficulties in the reconstruction of the paleoenvironmental conditions and the effects of preservation in the fossil assemblages. Despite this, factors such as sea-water temperature (e.g., Bornemann and Mutterlose, 2006; Linnert and Mutterlose, 2013; Wulff et al., 2020), nutrient (e.g., Erba et al., 1995; Giraud et al., 2006; Linnert and Mutterlose, 2013; Lübke et al., 2015; Wulff et al., 2020) and light availability (e.g., Lübke et al., 2015; Lübke and Mutterlose, 2016),

acidification (e.g., Erba et al., 2010), or trace metal- and CO₂ water concentrations (e.g., Faucher et al., 2017) are thought to have influenced coccolith growth and calcification. Coccolith size change in the fossil record has been also attributed to evolutionary processes (López-Otálvaro et al., 2012; Gollain et al., 2019). Changes toward decreasing size are usually claimed to represent a response to stressful and unstable environmental conditions (eutrophication, lower temperature and light availability, increased CO₂ concentrations and acidification) in some fossil coccolith species, while other taxa do not show size alterations. Long-term size increase, are however claimed to be related to evolutionary processes. Since most of the cosc ved biometric changes in this study (e.g., in *Flabellites oblongus*, in the new *I itn. aphidites* morphospecies and in the *Hayesites irregularis* gr.) represent mainly lon 3-1c m size increases, they are interpreted as related to evolutionary processe.

The biometric studies cited above were all performed on coccolith species. However, most of the biometric data presented in this study correspond to nannoliths whose possible relationship with corconals is uncertain and whose ecological preferences are mostly unknown.

However, our data suggest that some link existed between morphologic change/evolutionary patterns and paleogeography. Here we review the morphoevolutionary trends of selected nannofossil groups in relation to the latest Barremian–Aptian paleogeographic changes.

The genus *Lithraphidites* first appeared in the Berriasian (e.g., Perch-Nielsen, 1985) and consist of bladed rod-shaped nannoliths that taper towards both ends, have a cruciform

cross-section and may have a minute basal murolith coccolith. Janin (1988) suggested affinities of these nannoliths with Actinozoa that most authors do not find compelling, and it is currently thought that they may represent disarticulated spines of a heterococcolith

(https://www.mikrotax.org/system/index.php?taxon=Lithraphidites&module=ntax_mes ozoic).

A common evolutionary trend within the genus *Lithraphidires* Consists of a widening of the longitudinal blades in a symmetrical or asymmetrical way. This trend episodically originated in several species across the Cretaceous (e.g., *L. houghtonii* and *L. moray-firthensis* in the Aptian; *L. alatus*, *L. acutus* and *L. eccontricus* in the Albian-Cenomanian; *L. praequadratus*, *L. quadratus*, and *L. kennethii* in the Campanian-Maastrichtian). The current work shows this trend also operated on the Tethyan-Atlantic *Lithraphidites* across the lawst early Barremian-late Barremian (Figs. 11, 12, 13), leading to the differentiation of three species (*L. aichmoides*, *L. pugio* and *L. magnus*; Chapter 6). These construte the earlier known record of this evolutionary trend within the genus *Lithraphialites*.

Besides the Subbetic Domain, *L. magnus* was recorded in the North American Basin (Covington and Wise, 1987), West Iberian Continental Margin (Bralower et al., 1994) and northern Gargano (Cobianchi et al., 1997). This suggests a westernmost Tethys–North Atlantic paleogeographic record for this species (Fig. 2). The records of *L. aichmoides* and *L. pugio* are so far limited to the Subbetic as both are new species described in the current work and remain undifferentiated from *L. magnus* in the previous papers.

Lithraphidites houghtonii is a species with a very similar morphology to L. pugio/L. magnus, but being noticeably smaller (length 9–12 μm). It was recorded (Jeremiah, 2001; Bottini and Mutterlose, 2012) throughout the lower Aptian (uppermost part of Prodeshayesites fissicostatus AZ) to lower upper Aptian of the North Sea area and Lower Saxony Basin (Fig. 2). As the evolutionary lineage defined by the Tethyan-Atlantic L. aichmoides-L. pugio-L. magnus is earlier, we speculate these taxa (especially L. magnus) could be the ancestors of the Boreal species. In relation to this, the latest Barremian–early Aptian paleogeography (Fig. 2), together with sea level rise (e.g., Hallam, 1992; Mutterlose, 1992b; 1996; Hardenbol et al. 1998; Mutterlose and Böckel, 1998) would have allowed the migration of some roperations of L. magnus into the North Sea Basin, where they evolved into *L. n. m. htonii* (Fig. 11). Several authors record a homogenization followed by `Tthyan influx of marine invertebrate faunas (Mutterlose, 1987, 1998; Mutterlose and Böckel, 1998) and calcareous nannofloras (Mutterlose, 1987, 1989, 1992a 1936; Jeremiah, 2001) into the West European Province (WEP) of the Bore Realm throughout the early Aptian, which gives additional support to the suggested migration.

The marker species *Flabellites oblongus* and *Hayesites irregularis* gr. also underwent morphologic changes across the late Barremian–early Aptian here interpreted as the result of long-term evolutionary processes. The early coccoliths of *F. oblongus* are small, progressively increasing to medium size from the *Gerhardtia provincialis* ASz upwards (Figs. 8, 9; see Aguado et al., 1997; Bown, 2005). Regarding the *H. irregularis* gr., aside from the two morphotypes identified by mixture analyses (Chapter 4.2, 6; Fig. 7), a gradual increase in size has been observed in the specimens of *H. irregularis* s.str.

across the *Martelites sarasini* AZ (SM2). The LOs of *F. oblongus* and *H. irregularis* are often recorded, as rare species, from the lower Aptian interval (*Prodeshayesites fissicostatus/tenuicostatus—Deshayesites forbesi* AZs) in the WEP of the Boreal Realm (e.g., Erba et al., 1996; Bown et al., 1998; Mutterlose and Böckel, 1998; Jeremiah, 2001). However, these records correspond to the entry of the species in the Boreal Realm, which is concomitant with the homogenization of floras and faunas and Tethyan influx described above, and do not correspond to the true LOs of these species.

During the late Barremian, the WEP of the Boreal Real n and the Tethyan Realm were nearly isolated from each other (e.g., Ziegler, 1990; Mytterlose, 1992b; Barrier et al., 2018). However, Jeremiah (2001) recorded short influxes of Micrantholithus stellatus in rocks equivalent to the uppermost Barremia. Parancyloceras bidentatum AZ in the North Sea Basin (Fig. 2). The LO of M. **ellatus* in the Subbetic Domain has been recorded in the upper part of the Ce. ha. dtia provincialis ASz (see chapters 4.3 and 5.1, Fig. 9), that is, much earlier that no occurrence in the North Sea Basin. Based on its morphological similarities viti, the extant species Braarudosphaera bigelowii, the genus Micrantholithu is commonly interpreted as a marginal/neritic taxon (e.g., Roth, 1994; Applegate et al., 1989; Street and Bown, 2000; Bown, 2005; Bottini and Mutterlose, 2012; Quijano et al., 2012; Aguado et al., 2014a, b). The short influxes of M. stellatus in the North Sea Basin suggest that shallow seaways intermittently connected the WEP of the Boreal Realm and the Tethyan Realm already during the late Barremian, favoring the expansion of Tethyan neritic nannofloras into the Boreal Realm. As no Boreal taxa were recorded in our sections during this interval, we suggest this influx of neritic Tethyan taxa (M. stellatus) into the Boreal Realm was probably favored by increasing temperatures related to the Taxy Episode.

Late Barremian episodes of sea isolation, coincident with regressive periods, favored the allopatric speciation of calcareous nannoplankton and the apparition of endemic taxa under restricted conditions, (e.g., in the WEP of the Boreal Realm; Jakubowski, 1987; Crux, 1989; Bown et al., 1998; Jeremiah, 2001). During moderate to extensive high stands, seaways opened improving the communication between the Tethyan Realm and the WEP of the Boreal Realm, and nannofloral exchange (e.g., Mutterlose, 1992a, b, 1996). All the described biogeographic data suggest that paleography played a fundamental role in the evolution and spatial distribution of calcareous nannoplankton in the west European-Atlantic region during the late Barremian—early Aptian interval.

6. Systematic paleontology (by R. Aguado)

The taxonomic descriptions below follows he terminology guidelines of Young et al. (1997) and the higher taxonomy follows Bown and Young (1997). Only taxonomic references that do not appear in Bo vr. (1998) or cannot be found in the Nannotax website (http://www.mikrot.x.org/Nannotax3/) are provided in the reference list. In the following descriptions, L = ½ ngth, W = width, XPL = cross-polarized light, CNZ /CNSz = calcareous annofossil Zone/Subzone.

Images and smear slides of type material are stored in the Department of Geology of the University of Jaén (DG image numbers bracketed in the descriptions below). All species names are registered at the Plant Fossil Names Registry (PFNR; https://plantfossilnames.org) of the National Museum Prague. The corresponding PFNR nomenclatural act number is indicated for each new species. The reference calcareous nannofossil biozonation used is that of Bralower et al. (1995), with the modifications

introduced in this paper. The reference planktonic foraminifera biozonation is that of Coccioni et al. (2007).

Order Podorhabdales Rood et al., 1971, emend. Bown, 1987

Family Biscutaceae Black, 1971

Genus Crucibiscutum Jakubowski, 1986

Crucibiscutum bastetanum sp. nov.

PFNR: PFN001007.

(Figs. 14A–N)

1997 Crucibiscutum salebrosum Aguado et al., Figs 8.19, 8.20, 8.22 non Fig. 8.21.

2014a Crucibiscutum hayi Aguado et al., Figs. 4.8 -4.10.

2014b Crucibiscutum hayi Aguado et al., Fig. (13.

Derivation of name: After the Basterni, an old Iberian tribe that populated the southeastern part of the Iberian Peninsula, the type area of this species.

Diagnosis: Small to medium. sized (3.4–5.5 μm) broadly elliptical to normally elliptical Crucibiscutum coccoliths with a relatively narrow central area spanned by off-axial (slightly rotated) crow bars. The rim is bicyclic, with the bright inner cycle being wider (~1 μm width) than the outer dark cycle.

Description: Under XPL, the bright inner cycle of the rim has equal or greater width than the outer dark cycle. The central area is nearly filled by a relatively thick cross slightly rotated with respect to the axes of the ellipse defined by the rim. The cross is bright (although not as bright as the inner cycle of the rim) under XPL when oriented at 45°, and dark when oriented to 0°. The long and short arms of the central cross usually have slightly different rotation.

Dimensions: L = 3.4–5.5 (average 4.6) μ m; W = 2.9–4.5 (average 3.8) μ m; specimens measured: 22.

Holotype: Figs. 14A, B [DG2802, DG2803] (are same specimen). Holotype dimensions: $L=4.9~\mu m,\,W=3.9~\mu m.$

Paratypes: Figs. 14C, D [DG2804, DG2805] (are same specimen); Figs. 14E, F [DG2806, DG2807] (are same specimen).

Type level: Uppermost Barremian, sample X.HA-2 [DGF001] (NC6A1) nannofossil Subzone; upper part of *Imerites giraudi* AZ).

Type locality: X.HA section, Province of Granada, southeastern Spain.

Occurrence: This species has a short range. Its LO was found near the top of the Gerhardtia sartousiana AZ, (upper part of the Her uncolites feraudianus ASz), which is equivalent to the lowermost part of the NC6A! CNSz, and is shortly preceded by the LO of the primitive forms (elongated) of Jayesites irregularis E. Its HO was recorded in the lower part of the Martelites sacrasini AZ (upper boundary of the NC6A1 CNSz), shortly following the nc>wc Event Chapter 4.3, Fig. 9). Usually this species is a rare component of the assemblages, being more abundant from the upper part of the Imerites giraudi AZ to its HO, but the ws a rather consistent record. It was recorded in sections X.Ag₆, X.F, X.F1, A.Ya, X.Kv, X.Kv₂, and RAO3 from the Subbetic Domain in the Betic Cordillera (southern Spain). Its HO is used here to define the upper boundary of the NC6A1 CNSz.

Remarks: Crucibiscutum bastetanum differs from Crucibiscutum salebrosum (Berriasian–Hauterivian), Crucibiscutum ryazanicum (Berriasian–Valanginian) and Crucibiscutum pinnatus (Hauterivian–Barremian) by having a rim with a wider (~1 μm) bright inner cycle when viewed in XPL and a slightly larger size. Crucibiscutum salebrosum has, in addition, a cross aligned with the axes of the ellipse instead of a

slightly rotated one as in C. bastetanum. Crucibiscutum bastetanum differs from

Crucibiscutum neuquenensis (lower Hauterivian) and Crucibiscutum trilensis (upper

Hauterivian) by having a narrower central area, which is nearly filled by a relatively

thick cross, and a wider (~1 µm) bright cycle of the rim when viewed in XPL.

Crucibiscutum hayi (Upper Albian) and Crucibiscutum gracile (below) have wider

(more open) central areas and narrower axial cross bars. Palaeopontosphaera giraudii

(de Kaenel, 2020) has a greater size, with a bright inner cycle of the rim of similar width

to the outer dark cycle, and an axial cross. Following de Kaener et al. (2020), the range

of P. giraudii is early Hauterivian–late Barremian (HO in T vandenheckii AZ), being

earlier in time than C. bastetanum and there is a stratigiophical displacement between

these two species.

Crucibiscutum gracile sp. nov.

PFNR: PFN001008.

(Figs. 14O-AB)

1987 Cruciplacolithus hayi Tovington and Wise, Pl. 1, Fig. 6.

1994 Corollithion cf. C ach losum Bralower et al., Figs. 19.31, 19.32.

1997 Crucibiscutum 'ayı Aguado et al., Figs. 8.23, 8.24.

Derivation of name: From the Latin word gracilis meaning 'slim', 'slender', 'delicate',

referring to the narrow rim and slim axial cross of this species.

Diagnosis: Small (3.4–4.7 μ m) sub-circular to broadly elliptical (axial ratio ~1.1)

Crucibiscutum coccoliths with a wide central area spanned by slim axial cross bars. The

rim is narrow, bicyclic, with the bright inner cycle being usually slightly narrower (~0.5

µm width) than the outer dark cycle.

Description: The central area is wide (around twice as wide as the rim width) and partially covered by a slim bright cross with arms aligned with respect to the axes of the ellipse defined by the rim. The central cross is slightly brighter under XPL when oriented at 45° than when oriented to 0° .

Dimensions: L = 3.4-4.7 (average 4.1) μ m; W = 3.2-4.4 (average 3.8) μ m; specimens measured: 20.

Holotype: Figs. 14O, P [DG2808, DG2809] (are same specimen). Holotype dimensions: $L = 4.4 \mu m$, $W = 4.2 \mu m$.

Paratypes: Figs. 14Q [DG2810]; Figs. 14R, S [DG2811, L 72812] (are same specimen). Type level: Uppermost Barremian, sample X.HA-2 [DCF001] (NC6A1) nannofossil Subzone; upper part of the *Imerites giraudi* AZ.

Type locality: X.HA section, Province of Gran. de, southeastern Spain.

Occurrence: Found throughout the upper 3 arremian to lower Aptian (uppermost part of the NC5E1 to NC6A2 CNSzs; lower part of the Gassendiceras alpinum ASz to Hedbergella excelsa Zone of plank oric foraminifera) in sections X.Cp₂, X.F, XF1, X.HA, X.Kv, X.Kv₂, and R. 93 from the Subbetic Domain in the Betic Cordillera (southern Spain). This species was also recorded, as Cruciplacolithus hayi, in DSDP Site 603 off Cape Halteras by Covington and Wise (1987) and as Corollithion cf. C. achylosum across the upper Barremian-lowermost Aptian of the ODP Site 641 by Bralower et al (1994).

Remarks: Crucibiscutum gracile differs from all previously described Neocomian Crucibiscutum species (C. giraudii, C. neuquenensis, C. pinnatus, C. ryazanicum, C. salebrosum, C. trilensis) including C. bastetanum (above) by having a wider central area and a narrower rim with a lower axial ratio (~ 1.1), usually showing a sub-circular outline. Crucibiscutum gracile also differs from C. bastetanum, C. trilensis, and C.

ryazanicum in having an axial slim cross spanning the central area. Crucibiscutum

gracile has lower axial ratio and a wider and open central area than C. pinnatus.

Crucibiscutum hayi, from the Upper Albian, has a greater axial ratio, a wider rim and a

central area slightly narrow than that of C. gracile.

Family Cretarhabdaceae Thierstein, 1973

Genus Flabellites Thierstein, 1973

Flabellites oblongus (Buky, 1969) Crux in Crux et al., 1982

Remarks: A known morphological trend for the specimens of Flabellites oblongus along the upper Barremian–Albian (e.g., Aguado et al., 195.: Bown, 2005) is the increase in coccolith size. Recently, de Kaenel et al. (2020) e.ected a new species (Flabellites eclepensensis) for those small (3 to < ur.) specimens previously assigned to F. oblongus (e.g., Aguado et al., 2014a, However, the mixture analysis performed in the present study (Chapter 4.1 and Fig. 8) suggest a rather continuous increase in size of the measured population. The selected size of 5 µm by de Kaenel et al. (2020) to separate the specimens assigned to F. eclepensensis within the complete pool of F. oblongus s.l. seems arbitrary. In the absence of any other distinctive biometric (e.g., ellipticity) or morphological character that could help in a net separation of both species, we opted here to retain the wide concept (F. oblongus s.l., or simply F. oblongus) of the species.

Order Eiffellithales Rood et al, 1971

Family Chiastozygaceae Rood et al., 1973

Genus Chiastozygus Gartner, 1968

Chiastozygus lamprostauros sp. nov.

PFNR: PFN001011.

(Figs. 14AC–AP)

1997 Chiastozygus sp. cf. C. litterarius Aguado et al., Figs. 15, 16.

2014a Chiastozygus litterarius Aguado et al., Figs. 3.1–3.3.

Derivation of name: From the Greek words lampros meaning 'bright', and stauros meaning 'cross', 'saltire', referring to the XPL image of this species.

Diagnosis: Small to medium-sized (4.3–7.2 μ m) normally elliptical Chiastozygus coccoliths with a narrow (~1.2 μ m) rim. The relatively via central area is covered by a diagonal cross whose arms are seen simple and brighter than the rim when the longitudinal direction is oriented at 45° under XPI.

Description: This species has an axial ratio 11 and a relatively narrow rim, which is dark and diffusely bicyclic under XPL. The diagonal cross spanning the central area have narrow (~0.5 μm) arms that show median lines and remain dark when the longitudinal direction is oriented at 0°. These arms are seen simple (without median lines) and bright under XPL when the longitudinal direction is oriented at 45°. The angle between the arms 01° hy cross is greater (~105°) in the longitudinal direction than in the transverse direction (~75°). The arms of the diagonal cross are asymmetrical (slightly rotated) with respect to the axes of the ellipse. In some specimens, the diagonal cross supports the base of a distal stem.

Dimensions: L = 4.3-7.2 (average 5.6) μm ; W = 3.4-5.5 (average 4.2) μm ; specimens measured: 20.

Holotype: Figs. 14AC, AD [DG2813, DG2814] (are same specimen). Holotype dimensions: $L = 6.2 \mu m$, $W = 5.0 \mu m$.

Paratypes: Figs. 14AE, AF [DG2815, DG2816] (are same specimen); Figs. 14AL, AM [DG2817, DG2818] (are same specimen).

Type level: Uppermost Barremian, sample X.HA-7 [DGF002] (NC6A2 CNSz; Martelites sarasini AZ).

Type locality: X.HA section, Province of Granada, southeastern Spain.

Occurrence: Found, as a rare species, from the upper Barremian (upper part of the NC5E2 CNSz, Gerhardtia provincialis ASz) to uppermost lower Aptian (NC7A2 CNSz, Schackoina cabri Zone of planktonic foraminifera). Peccaded in sections X.Cp₂, X.F, X.F1, X.HA, X.Kv, X.Kv₂, and RA03 from the Subjection Domain in the Betic Cordillera (southern Spain).

(Santonian–Maastrichtian) in rim construction, and optical behavior from which is differentiated by having the arms of the diagonal cross slightly rotated with respect to the axes of the ellipse (asymmetric). It differs from Chiastozygus bifarius (Albian–Maastrichtian) and Chiastozygus bifarius (Aptian–Coniacian) by having a narrower, simple (with no n. dian lines when oriented at 45°) diagonal cross with slightly asymmetrical (rotalea) arms with respect to the axes of the ellipse. Chiastozygus litterarius (Aptian–Nastrichtian) is a rather poorly documented taxon originally described from the upper Maastrichtian sedimentary successions of Poland, and is characterized by having a weakly birefringent diagonal cross instead of the bright cross when oriented at 45° which is present in C. lamprostauros. Chiastozygus tenuis (another poorly documented species) has a slim diagonal cross symmetrical with respect of the axes of the ellipse instead of a broader, slightly rotated one as present in C. lamprostauros. Chiastozygus lamprostauros differs from other species of this genus (C. antiquus, C. garrisoni, C. stylesii and C. trabalis) by having a dark diffusely bicyclic

rim. Probably most of the late Barremian–early Aptian *C. litterarius* specimens from the literature should be assigned to *C. lamprostauros*.

Family Rhagodiscaceae Hay, 1977

Genus Rhagodiscus Reinhardt, 1967

Rhagodiscus sicutclipeus sp. nov.

PFNR: PFN001009.

(Figs. 14AQ–BD)

2014a Rhagodiscus sp. cf. R. achlyostaurion Aguado et al., 1-1gs. 3.48-50.

2014b Rhagodiscus sp. cf. R. achlyostaurion Aguado e. al., Fig. 3.3.

Derivation of name: From the Latin words cic. *r_leaning 'as', 'such as', and clipeus, the elliptical war shield used by the ancie. *Creeks and Romans which resembles the XPL image of this species.

Diagnosis: Small to medium-sized 4.6–5.9 μm) normally elliptical *Rhagodiscus* coccoliths with a relatively vide central area bearing a small spine base, which is bright in XPL. The rim and central area images are distinctly dark in XPL.

Description: This species has an elliptic rim (axial ratio ~1.4) and central area that remain relatively dark under XPL. The central area bears a small spine base, which is seen as a bright, solid (not hollow) circle traversed by four sharp extinction gyres. The spine base is relatively small (~1.3 μm in diameter), remains isolated in the middle of the central area and does not reach the internal margin of the rim.

Dimensions: L = 4.6-5.9 (average 5.1) μ m; W = 3.3-4.6 (average 3.7) μ m; specimens measured: 20.

Holotype: Fig. 14AQ [DG2819]. Holotype dimensions: $L = 5.3 \mu m$, $W = 3.7 \mu m$.

Paratypes: Figs. 14AR, AS [DG2820, DG2821].

Type level: Upper Barremian, sample X.HA-4 [DGF003] (NC6A1 CNSz; Martelites sarasini AZ).

Type locality: X.HA section, Province of Granada, southeastern Spain.

Occurrence: Found, as a rare species, from the upper Barremian (upper part of the NC5E2 CNSz, Gerhardtia sartousiana ASz) to uppermost lower Aptian (NC7A2 CNSz, Deshayesites forbesi AZ). Recorded in sections X.Ag₆, X.Cp₂, X.F, X.F1, X.HA, X.Kv, X.Kv₂, and RA03 from the Subbetic Domain in the Betic Cordillera (southern Spain).

Remarks: Rhagodiscus sicutclipeus is rather similar to Phagodiscus achlyostaurion (upper Aptian–Coniacian) from which differs by it's insticeably smaller central spine base, which does not seem hollow and does not reach the internal margin of the rim. Rhagodiscus pancostii (Lower Turonian) is smaller in size (holotype length = 3.96 µm) and lacks the distinctly optical features present in the rim and central area of R. sicutclipeus. Rhagodiscus hampton is upper Aptian–Upper Albian) is greater in size and has a less clearly defined (usually absent) bright spine base. Rhagodiscus buisensis lacks the solid birefringent or me base (de Kaenel et al., 2020) present in R. sicutclipeus. Probably most of the Barremian–lower Aptian occurrences of R. achlyostaurion cited in the literature correspond to R. sicutclipeus.

Order Watznaueriales Bown, 1987

Family Watznaueriaceae Rood et al., 1971

Genus Cyclagelosphaera Noël, 1965

Cyclagelosphaera platyaspis sp. nov.

PFNR: PFN001010.

(Figs. 12N–Q)

Derivation of name: From the Greek words platys, meaning 'wide' and aspis meaning 'shield', alluding to the wider distal shield of this species.

Diagnosis: Medium to large-sized (7.0–8.4 μm) Cyclagelosphaera coccoliths having a wide distal shield and a very small (1.1–1.4 μm) central area.

Description: Circular placoliths with proximal and distal shield elements composed by R-units. In distal view, the V-unit forms a narrow distal cycle around the central area and is seen as a thin, dark line under XPL. The central area is very small (1.1–1.4 μm) compared to the distal shield (7.0–8.4 μm), and is completely closed.

Dimensions: W = 7.0-8.4 (average 7.5) µm; specimens measured: 6.

Holotype: Fig. 12N [DG2827]. Holotype dime wions: $W = 8.4 \mu m$, central area = 1.1 μm .

Paratypes: Figs. 12O, P, Q [DG2820, DG2829, DG2847].

Type level: Uppermost Barremian, ample X.Kv₂-700 [DGF004] (NC6A2 CNSz; lower part of the *Martelites sarasi.* i Az.).

Type locality: X.Kv₂ section Province of Murcia, southeastern Spain.

Occurrence: Sporacing any recorded throughout the upper Barremian to lowermost Aptian (uppermost part of the *Gassendiceras alpinum* ASz to the *Deshayesites oglanlensis* AZ). This interval is equivalent to the uppermost part of the NC5E1–NC6A2 CNSzs. This species is a very rare component of the assemblages. It has been recorded from sections X.Kv, X.Kv₂ and X.Ag₆ in the Subbetic Domain of the Betic Cordillera (southern Spain).

Remarks: Cyclagelosphaera platyaspis differs from Cyclagelosphaera margerelii by having a noticeably larger size (7.0–8.4 μm instead of 4–6 μm), although both species

have a very small central area closed by calcite elements. *Cyclagelosphaera platyaspis* differs from *Cyclagelosphaera brezae* by having a distinct V-unit cycle around the central area, and from *C. argoensis*, *C. jiangii*, *C. lacuna* and *C. wiedmannii* by the lack of central opening or open central area. *Cyclagelosphaera rotaclypeata* and *C. deflandrei* both have central areas wider than that of *C. platyaspis*, which also lacks the raised central plugs present in *C. reinhardtii* and *C. shenleyensis*.

Nannoliths Incertae sedis

Family Microrhabdulaceae Deflandre, 1963

Genus Lithraphidites Deflandre, 1963

Lithraphidites aichmoides sp. nov.

PFNR: PFN001012.

(Figs. 12A–H)

Derivation of name: From the Gree words aichme meaning 'spearhead', and the suffix -oides meaning 'resembling 'o'.

Diagnosis: Very large (15.3–25.3 μ m) Lithraphidites nannolith having an outline similar to a bladed of ger. In the direction opposite to the 'handle', the outline is slightly convex and decreases its width slowly before tapering toward the point. The maximum width oscillates between 2.7 and 4.1 μ m.

Description: Calcareous rods with cruciform cross-section built of long blades of identical optical orientation. The outline is similar to a bladed dagger in which the 'handle' (\sim 1.1 μm width and frequently missing by breakage) is followed by a wider expansion (2.7–4.1 μm width) which slowly decreases its width before tapering toward the point.

Dimensions: L = 15.3–25.3 (average 19.6) μ m; W = 2.7–4.1 (average 3.6) μ m; specimens measured: 30.

Holotype: Fig. 12A [DG2822]. Holotype dimensions: $L=20.1~\mu m,~W=3.2~\mu m.$

Paratypes: Figs. 12B-D [DG2823-DG2825], H [DG2826].

Type level: Uppermost Barremian, sample X.Kv₂-235 [DGF005] (NC5E2 CNSz; lowermost part of the *Gerhardtia sartousiana* ASz).

Type locality: X.Kv₂ section, Province of Murcia, southeastern Spain.

Occurrence: Recorded from the uppermost lower Barremian (upper part of the Moutoniceras moutonianum AZ) to the upper Barremian (uppermost part of the Hemihoplites feraudianus ASz), within NC5E1–NC6A! CNSzs. This species is a rare component of the assemblages, but has a consistent and continuous record until the uppermost part of the Gerhardtia provincia'is ASz. It has been recorded from sections X.CO₂, X.CT, X.F, X.Kv, X.Kv₂, and X.V₁ from the Subbetic Domain in the Betic Cordillera (southern Spain). The LO of L. aichmoides has been recorded 467 cm above that of Flabellites oblongus (base of NC5E1 CNSz) in section X.V₁.

Remarks: Lithraphidites aic. moules differs from L. magnus (emend.) and L. pugio by having a more stylized outline, with a maximum width smaller than 4.5 µm. This width appears as the more appropriate to separate both species according to the measurements made in the present study (Fig. 5). Lithraphidites aichmoides differs from Lithraphidites alatus (Upper Albian–Lower Cenomanian) by having an outline similar to a bladed dagger instead of a 'closed umbrella', the latter decreasing quickly in width from the end close to the 'handle' towards the point. It differs from Lithraphidites houghtonii (Boreal lower–upper Aptian) by having a greater length and a smaller width.

Lithraphidites magnus (Covington and Wise, 1987) comb. & stat. nov.

PFNR: PFN001015.

(Figs. 13F–J)

1987 Lithraphidites alatus ssp. magnus Covington and Wise, Pl. 10, Figs. 3, 4, 8?.

1994 Lithraphidites alatus ssp. magnus Bralower et al., Figs. 20.7, 20.8.

2020 Lithraphidites magnus Martinez et al., Fig. 4M.

Emended diagnosis: Very large (usually >21 μm) Lithraphidites nannolith having an outline similar to a bladed dagger. The 'handle' (~1.1 μm width and sometimes missing by breakage) is followed by a wider expansion (4.5–7.0 μm, width) which nearly maintains its width a long distance (around three fifths of the nannolith length) before tapering toward the point. The maximum width is greater than 4.5 μm and the L/W ratio is greater than 3.5 (average 4.1).

Remarks: Covington and Wise (1987, Σ. €31) originally described this taxon, but as a subspecies of Lithraphidites alatus (L. alatus ssp. magnus). Lithraphidites magnus is here erected to the category of spec Δ, because the lack of stratigraphic connection with L. alatus and the existence C morphologic differences between both taxa. While L. magnus is restricted to the permost Barremian–lowermost Aptian, L. alatus ranges from Upper Albian to Lower Cenomanian. The original description of L. magnus was based on two SEM micrographs in which specimens are foreshortened due to tilting or partially covered. The estimated holotype dimensions were 5.7 μm width and ~20 μm long. However, the holotype appears to be a broken specimen, its true length probably being ~25 μm. The length in the population measured in the present paper oscillates between 21.2 and 26.2 μm (average 23.8 μm). After a biometric study (Fig. 6), all specimens with L/W ratio smaller than 3.7 (usually with L<21 μm) were here assigned to Lithraphidites pugio n. sp., which is characterized by a wider and shorter outline (see

below). It seems a width greater than 4.5 μ m is also a good value to characterize these forms differentiating them from *Lithraphidites aichmoides* n. sp. (Fig. 5).

Occurrence: Covington and Wise (1987) recorded this taxon in sediments from core 44 of DSDP Hole 603B (North American Basin of North Atlantic). These were dated as lower Aptian, although no convincing evidence supporting this age was provided from calcareous nannofossil assemblages. Bralower et al. (1994) reported this species from upper Barremian-lower Aptian sediments of DSDP Site 398 (West Iberian continental margin). Cobianchi et al. (1997) recorded this species throughout upper Barremian lower Aptian (M. hoschulzii to H. irregularis CNZs) in nor hern Gargano (Apulia, Italy). However, the species concept of these authors of bably includes L. magnus and L. pugio as differentiated here, which originates ar expanded range. In the Subbetic Domain of the Betic Cordillera, L. magnus 'sa, been recorded from the uppermost Barremian (uppermost part of the Heiriho plites feraudianus ASz) to the lower Aptian (lower part of the Deshayesites forberi AZ), thoroughout most of the lowermost part of NC6A CNSz. This species is a rare component of the assemblages, but has a consistent and continuous record until the upper part of the Martelites sarasini AZ, from which it becomes rare. It has been regarded from sections X.Cp₂, X.F, X.F1, X.HA, X.Kv, X.Kv₂, and RA03, in the Subbetic Domain of the Betic Cordillera (southern Spain).

Lithraphidites pugio sp. nov.

PFNR: PFN001013.

(Figs. 12I–M; Figs. 13A–E)

2017 Lithraphidites magnus Aguado et al., Fig. 2.

2020 Lithraphidites sp. cf. L. magnus Martinez et al., Fig. 4N.

Derivation of name: From the Latin word *pugio*, the double-edged, straight short dagger used by the ancient Romans, whose outline resembles that of this species.

Diagnosis: Very large (13.4–23.5 μ m) Lithraphidites nannolith having an outline similar to a short bladed dagger. In the direction opposite to the 'handle', the outline is convex and quickly decreases its width before tapering toward the point. The maximum width oscillates between 4.5 and 8.1 μ m.

Description: Calcareous rods with cruciform cross-section built of long blades of identical optical orientation. The outline is similar to a bladed αωgger in which the 'handle' (~1.1 μm width and frequently missing by breakage) is followed by a wider expansion (4.5–8.1 μm width) which nearly maintains ως width a short distance (around a half of the nannolith length) before quickly tapering toward the point. The L/W ratio is smaller than 3.5 (average 2.9).

Dimensions: L = 13.4–23.5 (average 18.1), μ m; W = 4.5–8.1 (average 6.2) μ m; specimens measured: 48.

Holotype: Fig. 12I [DG2842]. Holotype dimensions: $L = 19.6 \mu m$, $W = 7.8 \mu m$. *Paratypes*: Figs. 12K, L, [D 32843, DG2844]; *Lithraphidites* sp. cf. *L. magnus* in Martinez et al., 2020; Fig. 4N, which is reproduced here as Fig. 13A, [DG2845]; Fig. 13B, [DG2846].

Type level: Uppermost Barremian, sample X.Kv₂-228 [DGF006] (NC5E2 CNSz; lowermost part of the *Gerhardtia sartousiana* ASz).

Type locality: X.Kv₂ section, Province of Murcia, southeastern Spain.

Occurrence: In the Subbetic Domain of the Betic Cordillera, *L. pugio* has been recorded throughout part of the upper Barremian (from the middle part of the *Gassendiceras* alpinum ASz to the uppermost part of the *Hemihoplites feraudianus* ASz; base of NC5E2 to basal part of NC6A1 CNSzs). This species is scarce, but has a consistent

record in the assemblages from the X.F, X.HA, X.Kv, X.Kv₂ and X.V₁ sections.

Martinez et al. (2020) used the LO of this species to correlate the uppermost part of the X.V₁ and the lowermost part of the X.Kv₂ sections. This bioevent has shown biostratigraphic potential to be used in correlation, at least at a regional scale (chapters

Remarks: Lithraphidites pugio differs from L. magnus (emend.) by having a shorter (usually < 21 μm) and wider outline, with an average L/W ratio ~2.9. The expansion of the blades decreases more quickly in L. pugio than in L. magnus. It differs from L. aichmoides by having a greater width (>4.5 μm) and ne ray parallel sides (by around the half of its length) in side view, before tapering at the end Lithraphidites pugio differs from Lithraphidites alatus (Upper Albian–Lower Cenemanian) by having an outline similar to a short bladed dagger instead of a 'c' and umbrella', the latter having a straight to slightly concave outline quarkly decreasing in width from the end close to the 'handle' towards the point. Jeremian (2001) described Lithraphidites houghtonii from the lower–upper Aptian transition (2 shayesites forbesi to Epicheloniceras martinoides AZs) of the Central North S. a. This later species is morphologically rather similar to L. pugio, from which it different having a smaller size (length = 9–12 μm, width ~5.6 μm).

Radiate multielement group

Genus Hayesites Manivit, 1971

Hayesites irregularis E

(Figs. 15A–R)

5.1 and 5.2).

2020 Hayesites irregularis E Martinez et al., Figs. 4I–L.

Diagnosis: Specimens of *H. irregularis* (Thierstein in Roth and Thierstein, 1972)

Applegate et al. in Covington and Wise, 1987 with an elongated outline (L/W>1.12)

usually showing poorly defined radial elements under cross-polarized light microscopy

(XPL). See section 4.2 for details.

Description: A mixture analysis performed on the L/W ratio of 140 specimens of the *H. irregularis* gr. (Fig. 7C and SM2) indicates that two populations are present within this pool. Those forms with nearly circular outline, (L/W smaller or equal to 1.12), being younger in time, were assigned to *Hayesites irregularis* s.str These specimens having a more elongated outline (L/W>1.12), generally showing poorly defined radial elements and being slightly older, were assigned to *H. irregularis* morphotype E (*H. irregularis* E).

Dimensions: L = 2.8–5.3 (average 3.8) μ m; W = 2.2–4.3 (average 3.0) μ m; specimens measured: 51.

Occurrence: Hayesites irregularis E was recorded from the lower part of the Hemihoplites feraudianus ASz in the Subbetic Domain of the Betic Cordillera (sections RA03, X.Ag₆, X.F, X.HA, Y. Kv, X.Kv₂) and extends throughout the Imerites giraudi—Deshayesites oglanlensis A.7, of the upper Barremian—lowermost Aptian. The LO of this species has been used here to define the base of the NC6A1 CNSz. The LO of H. irregularis s. str. was recorded from the middle part of I. giraudi AZ.

Nannoliths

Order Braarudosphaerales Aubry, 2013 emend. Lees and Bown, 2016

Family Nannoconaceae Deflandre, 1959

Genus Nannoconus Kamptner, 1931

Nannoconus sp. cf. N. truittii

(Figs. 15AK–AM)

Diagnosis: Specimens of Nannoconus resembling N. truittii ssp. rectangularis Dères and Achéritéguy, 1980 in being wider than tall in side view and having a central canal of similar width than the wall. They differ from N. truittii ssp. rectangularis by showing a slightly conical, instead of rectangular, outline.

Dimensions: L = 5.6-9.9 (average 7.4) µm; W = 8.1-11.9 (average 10.2) µm; specimens measured: 28.

Occurrence: These forms were recorded (Figs. 3, 4; section: RA03, X.Ag6, X.CP₂, X.F, X.F1, X.HA, X.Kv, X.Kv₂) from the upper part of the Momihophites feraudianus ASz to the Deshayesites forbesi AZ (NC6A1–NC6A2 CNSz.)

Remarks: Nannoconus truittii ssp. frequens an N. truittii ssp. truittii are respectively taller than wide and of similar height par width in side view (Brönnimann, 1955; Dères and Achéritéguy, 1980), differing from N. sp. cf. N. truittii. Nannoconus truittii ssp. rectangularis was described from the Albian (Dères and Achéritéguy, 1980), while N. sp. cf. truittii has been recorded form the uppermost Barremian–lower Aptian.

7. Conclusions

A detailed study of the calcareous nannofossil assemblages from twelve uppermost lower Barremian–lowermost Aptian sections in the Subbetic Domain of the Betic Cordillera, well dated by ammonite biostratigraphy, has allowed to the identification of seven new species. The new taxa *Crucibiscutum bastetanum*, *Crucibiscutum gracile*, *Chiastozygus lamprostauros*, *Cyclagelosphaera platyaspis*, *Lithraphidites aichmoides*, *Lithraphidites pugio*, and *Rhagodiscus sicutclipeus* are described, one species is

emended (*Lithraphidites magnus*) and the taxonomic concept of the marker species *Hayesites irregularis* is discussed and clarified.

The stratigraphic ranges of each new species, together with those of other relevant markers, are directly correlated to standard ammonite biostratigraphy and tied to the geochronological scale using a previous astrochronological calibration of the complete interval studied.

Five new calcareous nannofossil subzones are proposed (NC5E1, NC5E2, NC5E3, NC6A1, and NC6A2) which allow the refinement of the zonation for the Mediterranean—Atlantic Province of the Tethyan Pearm and are directly correlated with respect to the standard Tethyan ammonite zonation. Two of the new species described here are used as biostratigraphic mark are or the new proposed subzones. The duration of each one of these subzones is also provided through a previous astrochronological calibration.

The implications of these results on the extant definition of the Barremian/Aptian boundary are discussed. In the absence of magnetostratigraphic data, we chose the definition of the Barremian/Aptian boundary in coincidence with the LO of the ammonite species $Deshayesites\ oglanlensis$. Regarding calcareous nannofossils, none of the biostratigraphic reliable markers coincides with this boundary. The recorded bioevent closer to the Barremian/Aptian boundary was the HO of the new species $Crucibiscutum\ bastetanum$, which correlates to the lower part of the $Martelites\ sarasini$ AZ. Based on its proximity to the base of the $D.\ oglanlensis\ AZ$, the ISNE of the $\delta^{13}C$

curve may be an alternative for the definition of the base of the Aptian, if it is identified in additional locations.

The morpho-evolutionary trends and paleogeographic distribution of some selected nannofossil taxa (mainly from genera *Flabellites*, *Lithraphidites* and *Micrantholithus*) across the interval studied were analyzed. This analysis suggests that paleogeography played a fundamental role as a factor controlling the evolution of the calcareous nannoplankton in the west European-Atlantic region during the Interval.

The following are the supplementary data related to use article.

Supplementary Material 1 (SM1): Detailed 'oction, lithostratigraphy and previous data on biostratigraphy of the sections studied

Supplementary Material 2 (SM2): a) stratigraphic ranges of individual calcareous nannofossil species in the 12 sections studied; b) size measurements of the new species described and the additional haxa included in the biometric study.

Supplementary Material 5. A phabetic list of all taxa cited in the text, with author attributions and date.

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Author statement

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Delaration of competing interest

None.

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Figure 1. Geologic sketch of the central and eastern sectors of the Betic Cordillera, showing the present-day location of the sections included in this study. A: X.Cp₂ section; B: X.V₁ section; C: X.Kv and X.Kv₂ sections; D: X Ag₆ section; E: RA03 section; F: X.CT and X.HA sections; G: X.Z section; H^{*} A.^{T^{*}} and X.F1 sections; I: X.CO₂ section.

Figure 2. Paleogeographic map of the Western. Fethys—Atlantic connection for the late Barremian (~123 Ma) with indication of the paleolatitude (modified from Ziegler, 1990 and Barrier et al., 2018). Location of reveral areas and sections referenced throughout the text. Sb: Subbetic Basin. 398: I SDP Site 398 (Bralower et al., 1994); 603: DSDP Site 603 (Covington and Wi. e., 1>87); 641: ODP Site 641 (Bralower et al., 1994); Cg: UK North Sea Central Grabol (Well 22/26a-2); Mf: Moray-Firth area (Well 21/2-6); Nj: North Jens-1 Weil Sp: Speeton cliffs (see Jeremiah, 2001). Gp: Gargano Promontory (Cobianchi et al., 1997); Ls: Lower Saxony Basin (Malkoč et al., 2010).

Figure 3. Lithologic sketches of sections X.V₁, X.Kv, X.Kv₂ and RA03, with indication of formations, stage, ammonite/calcareous nannofossil biostratigraphy, thickness, position and number (in parentheses) of samples studied and main calcareous nannofossil bioevents recorded. Marker species names are in bold typeface.

Figure 4. Lithologic sketches of sections X.Ag₆, X.CT, X.Z, X.HA, X.F1, X.F, X.CO₂ and X.Cp₂, with indication of formations, stage, ammonite/calcareous nannofossil biostratigraphy, thickness, position and number of samples studied and main calcareous nannofossil bioevents recorded. Marker species names are in bold typeface.

Figure 5. Cross-plot of width (W) vs. length (L) for 120 upper Barremian *Lithraphidites* spp. specimens. A–D: Cross-plots of width vs. length for the different taxa and sketches of the side and top views of each one (upper right), with indication of measured parameters. E, F: Frequency histograms of the width (E) and length (F) of the complete data set. The distribution curves in E result from a mixture analysis applied to the whole data set using the widths of all measured *Lithraphiduce* spp. specimens. G: Frequency histogram showing the distribution curves resulting from a mixture analysis applied to the length of all measured specimens. Aving a width greater than 5 μm. H: Table showing the descriptive statistical parameters (proportion, mean, and standard deviation) of the mixture analyses μer formed on the width and length of the measured specimens. Dashed vertical ines on the cross-plot are the widths suggested to separate *Lithraphidites carniolensu*. From the new species *L. aichmoides* (3 μm) and *L. aichmoides* from the magnus/L. pugio plexus (5 μm). Circles with cross on the graph indicate the calculated mean values for each species/group of taxa. W was measured in the area with the maximum expansion of the blades of *Lithaphidites* spp.

Figure 6. Cross-plot of length/width (L/W) ratio vs. length (L) for 69 *Lithraphidites* specimens of the *L. pugio/L. magnus* plexus. Note as all specimens assigned to *L. magnus* have L >24 μ m and L/W ratio >3.5, being located within the upper right quadrant of the graph.

Figure 7. A. Parameters measured in 140 specimens of the *Hayesites irregularis* group. B) Cross-plot of L vs. W. Shaded dots indicate initially assumed *H. irregularis* E. Regression lines considering all specimens (solid) and the separated populations of *H. irregularis* s.str. and *H. irregularis* E (dashed). C) Frequency histogram showing the distribution curves resulting from a mixture analysis applied to the L/W ratio of all measured specimens. The small table shows the descriptive statistical parameters (proportion, mean, and standard deviation) resulting from the nileture analysis. Two populations are revealed based on the L/W ratio. The forms with L/W > 1.12 were assigned to *H. irregularis* E. See text for details.

Figure 8. X.- Cross-plot of coccolith length (L) vs. width (W) for 120 specimens of Flabellites oblongus s.l., referred to a monite zones (different symbols for data points), with frequency histograms of the convolete data set. The distribution curves on the histograms result from applying mixture analyses. Dashed-line curves (A) result from the discrimination of two populations within the whole data set. Solid continuum line curves (B) result from the discrimination of only one population in the complete data set. Y.- Table shown, the descriptive statistical parameters resulting from the mixture analyses, where the two options (A and B) are considered. Mout = Holcodiscus caillaudianus ASz + Moutoniceras moutonianum AZ; Vand = Toxancyloceras vandenheckii AZ; Sart = Gerhardtia sartousiana ASz; Prov = Gerhardtia provincialis ASz; Fera = Hemihoplites feraudianus ASz; Gira=Imerites giraudi AZ; Sara=Martelites sarasini AZ; Ogla=Deshayesites oglanlensis AZ; Forb=Deshayesites forbesi AZ.

Numbers in parentheses represent the specimens measured in each interval. Note the

nearly continuous distribution of the coccolith size, where two populations can not be clearly differentiated and the close AIC values for cases A and B.

Figure 9. Geochronostratigraphic sketch showing a compilation of the main biostratigraphic events recorded in this study. Geochronology and ammonite stratigraphy after Martinez et al. (2020). Positions of the MBE (Mid-Barremian Episode) TE (Taxy Episode) and C-isotope Intra-Sarasini Negative Excursion (ISNE) are based on data from Martinez et al. (2020). Calcareous name fossil subzones are those proposed here. The stratigraphic ranges of all new species described in present study and the newly described calcareous nannofossil subzones are calibrated to ammonite zones and to geochronology. Ranges with a tack lines correspond to marker species. Intervals of the species ranges with very are/spotty record are marked with dashed line. Numbers in parentheses a feet to the numerical ages (in Ma) of the bioevents associated to zonal marker, using the chronology of Martinez et al. (2020). The web-ne horizon indicates the strategraphic position where the wide-canal nannoconids outnumber the varrow canal nannoconids (web-ne Event).

Figure 10. Correlatio. Of the uppermost Barremian—lowermost Aptian interval of the sections of Barranco de Cavila (X.Kv₂), Angles, Casis-La Bédoule and Gorgo a Cerbara using ammonite and C-isotope data. Light green shaded band correspond to the ISNE associated to the Taxy Episode. Light blue shaded band correspond to the OAE 1a. Lines of correlation: dashed lines based on ammonite biostratigraphy, dotted-dashed lines are based on calcareous nannofossils, and dotted line is based on C-isotope curves. Data for X.Kv₂ section after Martinez et al. (2020) and this paper; data for Cassis-La Bédoule after Kuhnt et al (1998) and Moullade et al. (1998a); data for Angles after

Delanoy (1995), Wissler et al. (2002) and Godet et al. (2006); data for Gorgo a Cerbara after Stein et al. (2011, 2012) and Frau et al. (2018). In Angles section, the boundary between *Martelites sarasini* and *Deshayesites oglanlensis* AZs marked with a star is after Godet et al. (2006), the boundary marked with two stars (bed 197b) is after Delanoy (1995), and reinterpreted here following Reboulet et al. (2011).

Figure 11. Proposed morpho-evolutionary sketch for the genus *Lithraphidites* across the late Barremian–early late Aptian. Chronostratigraphic scale folic wing Martinez et al. (2020). Calcareous nannofossil zones/subzones follow Frail wer et al. (1995) and modifications proposed in this work. Tethyan ammonic zones after Reboulet et al. (2018). NW Europe ammonite zones follow Mutterloca (1992b), Casey et al. (1998), Mutterlose and Böckel (1998), Malkoč et al. (2013), and Bottini and Mutterlose (2012). Stratigraphic range of *Lithraphidites in wintonii* and *Lithraphidites moray-firthensis* after Jeremiah (2001) and Bottini and Mutterlose (2012). Sea-level curves (dashed line = long-term; continuous line = short-term) after Hardenbol et al. (1998). MBE = Mid-Barremian Episode, TE = T. xy Episode, ISNE = C-isotope Intra-Sarasini Negative Excursion. Duration of the MBE, TE and ISNE after Martinez et al. (2020). Duration of the OAE 1a after Li Call (2008), Malinverno et al. (2010) and Giraud et al. (2018).

Figure 12. Cross-polarized light micrographs of several specimens of some of the new species described here. Holotypes/paratypes are indicated. Species name and section - sample of provenance are shown for each one.

Figure 13. Cross-polarized light micrographs of several specimens of some of the new species described here. Holotypes/paratypes are indicated. Species name and section - sample of provenance are shown for each one.

Figure 14. Cross-polarized light micrographs of several specimens of some of the new/emended species described here. Holotypes/paratypes are indicated. Species name and section - sample of provenance are shown for each one.

Figure 15. Cross-polarized light micrographs of several specimens of the *Hayesites irregularis* group, with differentiation of the two morphotypes described in the text and *Nannoconus* sp. cf. *N. truittii*. Species name and section - sample of provenance are shown for each one.

Highlights

- Seven new upper Barremian na ur) fossil species are described.
- Tethyan upper Barremian alca eous nannofossil biostratigraphy is improved.
- The latest Barremian δ^{12} C excursion should be considered as a potential marker for the base of the Aptian
- Calcareous nannofoss. morpho-evolutionary trends were controlled by paleogeography.

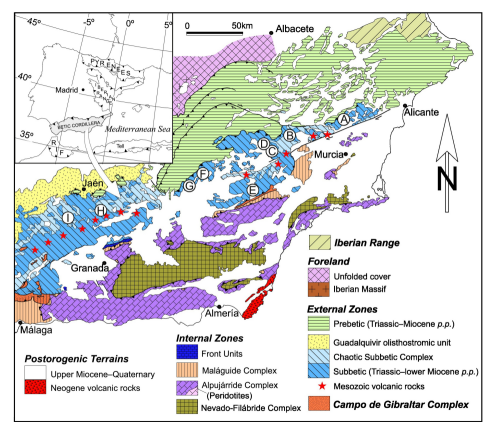


Figure 1

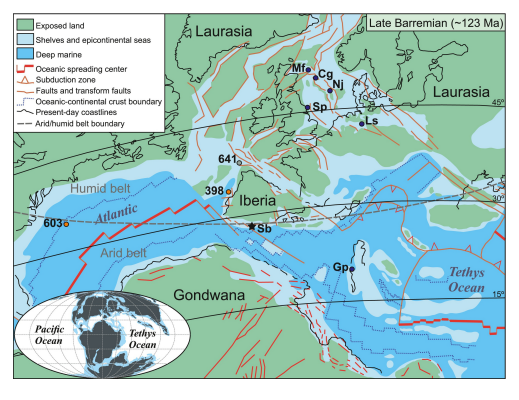


Figure 2

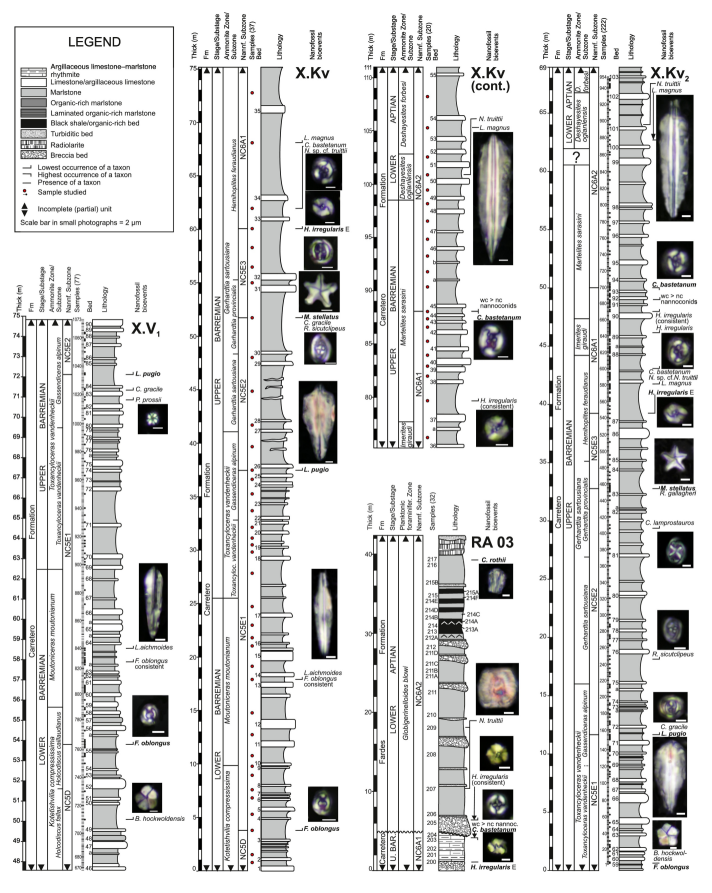


Figure 3

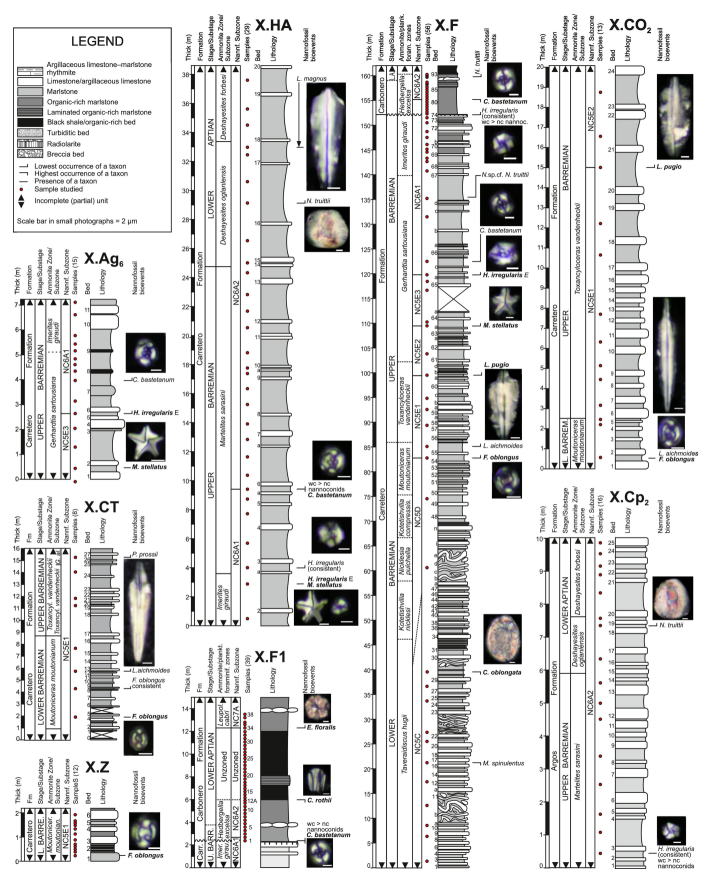


Figure 4

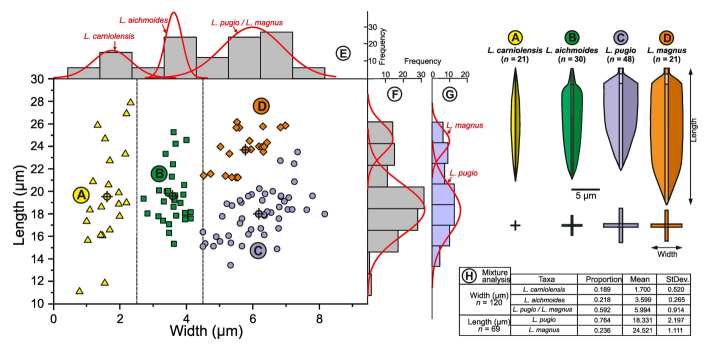


Figure 5

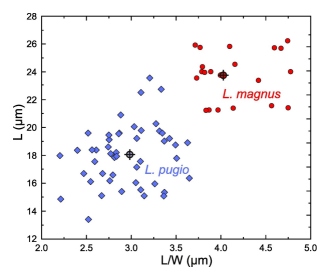
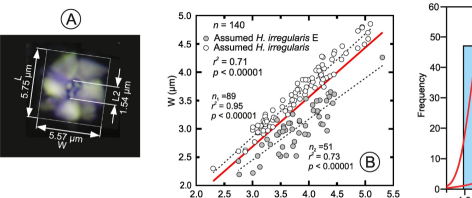


Figure 6



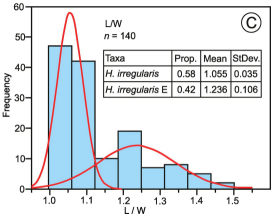


Figure 7

L (µm)

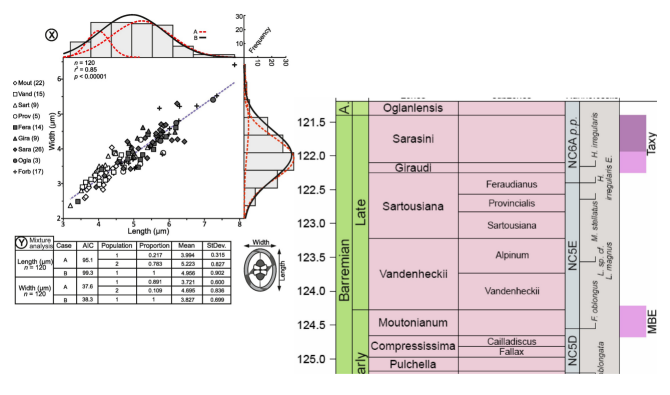


Figure 8

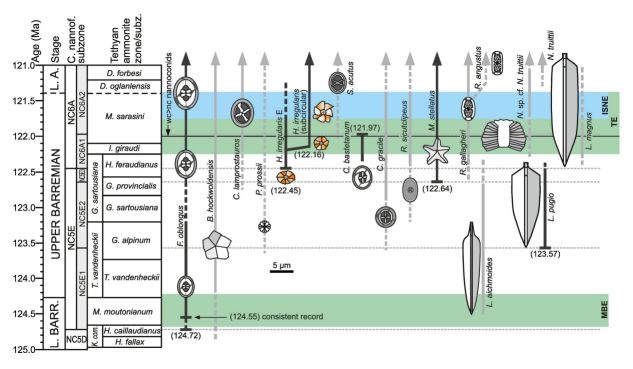


Figure 9

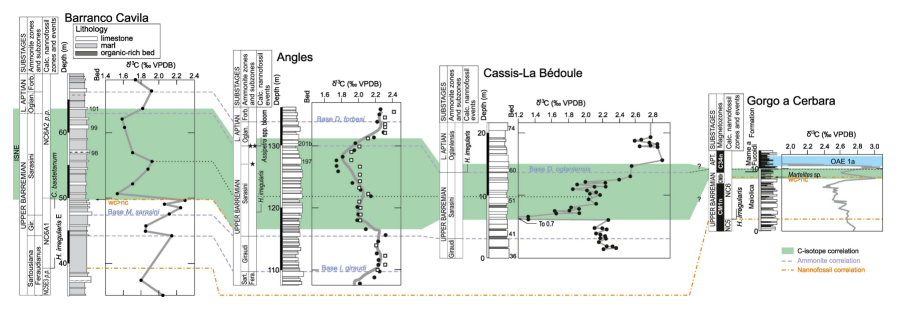


Figure 10

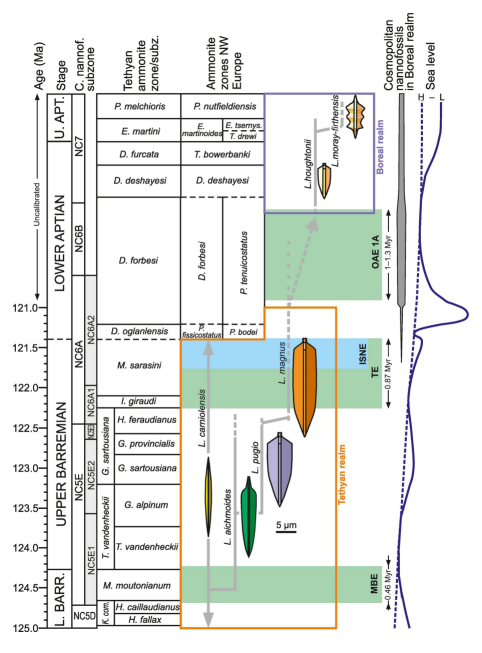


Figure 11

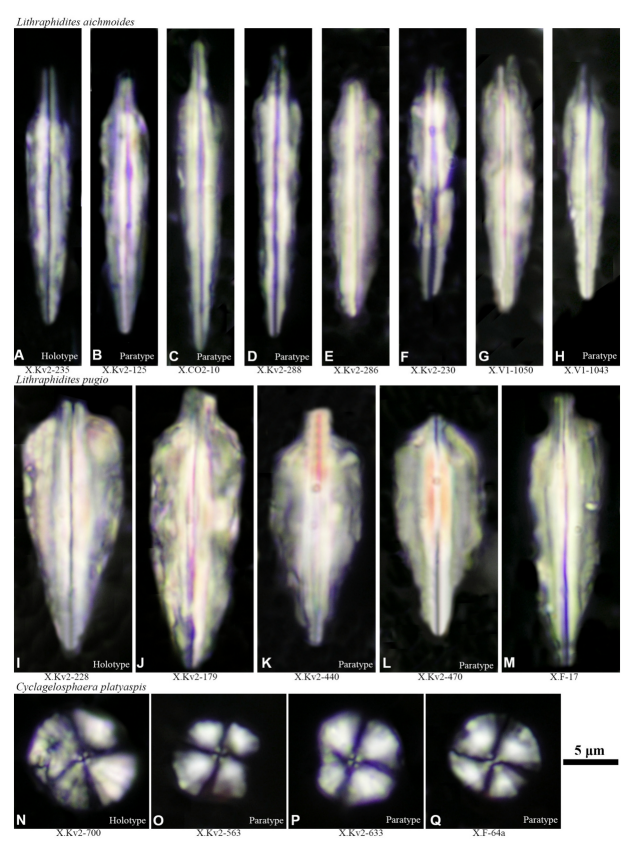


Figure 12

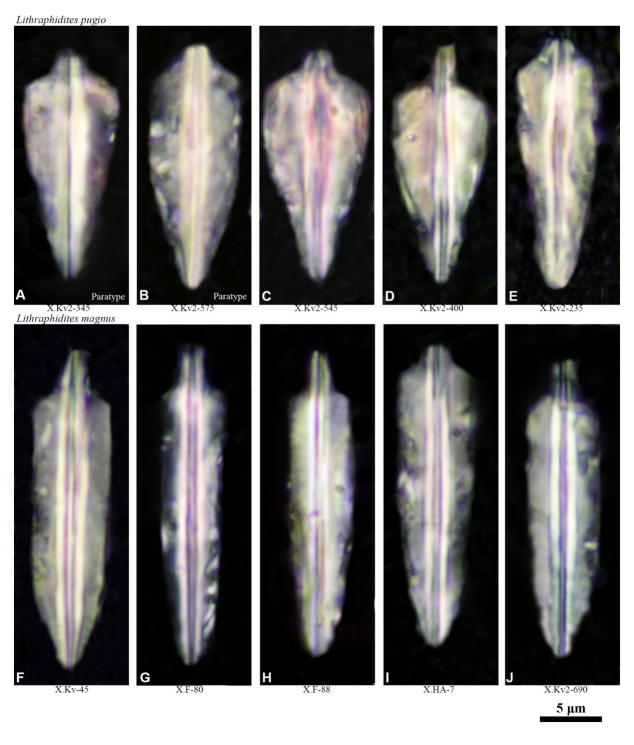


Figure 13

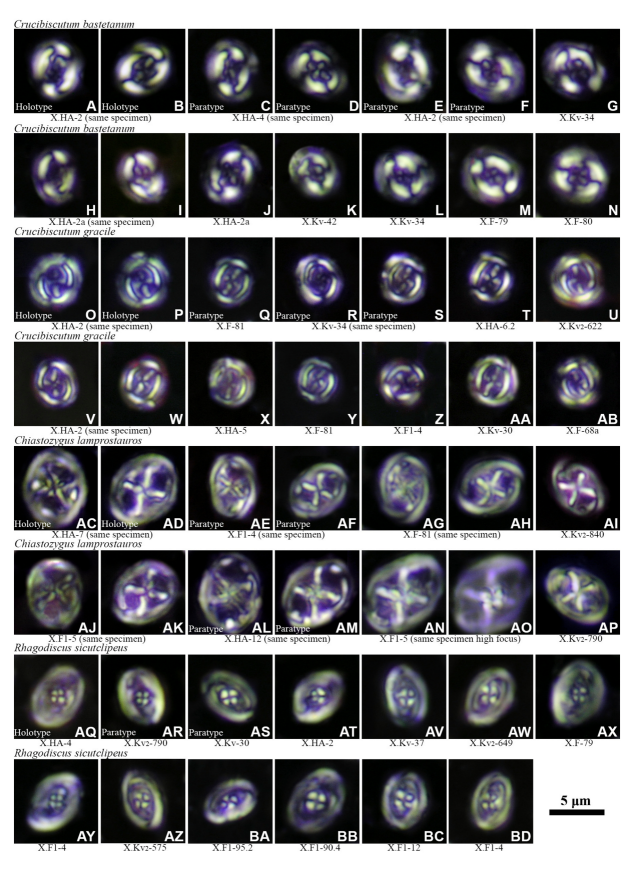


Figure 14

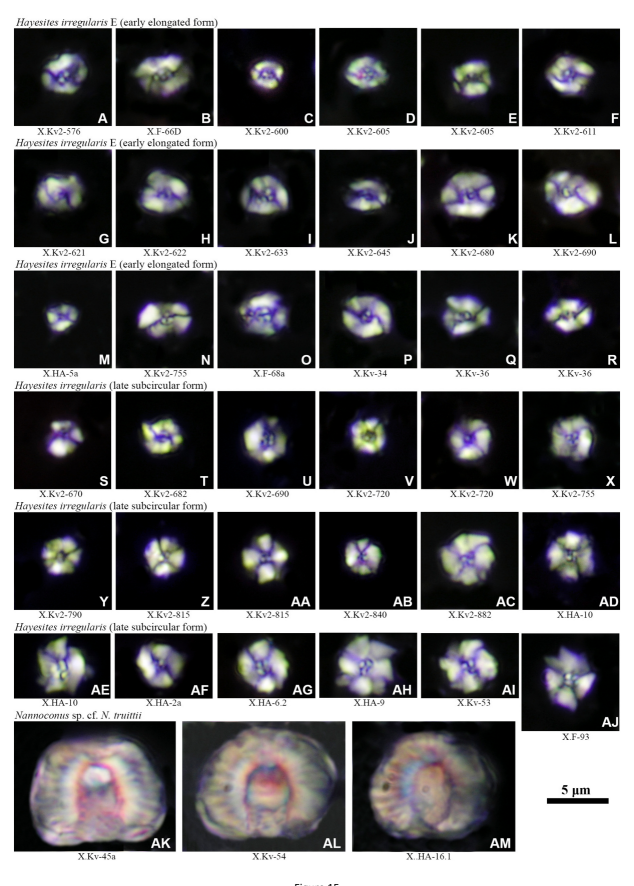


Figure 15