

Re-examination of the palynological content of the Lower Cretaceous deposits of Angeac, Charente, south-west France: Age, palaeoenvironment and taxonomic determinations

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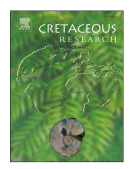
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| 1 | Re-examination of the palynological content of the Lower Cretaceous deposits of |
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| 2 | Angeac, Charente, south-west France: age, palaeoenvironment and taxonomic |
| 3 | determinations |
| 4 | |
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| 13 | |
| 14 | |
| 15 | Abstract |
| 16 | Further to the work published by Néraudeau et al. six years ago, palynological matter |
| 17 | recovered from the lignitic bone bed of Angeac in Charente (south-west France) has been re- |
| 18 | examined in order to provide more evidence of its age, the previous Hauterivian–Barremian |

¹⁹ interpretation being regarded as controversial. The samples come from four sedimentological

units, An2–An5, which together yielded an assemblage of 34 species, taxonomically

dominated by palaeoecologically significant lygodiaceous verrucate spores, including eight

species attributable to the genus *Concavissimisporites* and three species of *Trilobosporites*. A

²³ few bisaccate pollen grains, including *Vitreisporites pallidus*, have been recovered, but the

24 gymnospermous pollen spectrum is clearly dominated by the cheirolepidiaceous genus

25 *Classopollis*, its abundance diminishing progressively upwards from units An4 to An2. The

²⁶ great abundance and diversity of verrucate forms, along with the presence of other spores

| 27 | typical of Lower Cretaceous deposits, such as Aequitriradites verrucosus, and the scarcity of |
|----|---|
| 28 | specimens referable to Cicatricosisporites, render this assemblage most similar to those of the |
| 29 | Hastings Group of southern England, and the Bückeberg Formation in north-western |
| 30 | Germany. The time of deposition is, therefore, more likely to have been Berriasian- |
| 31 | Valanginian rather than Hauterivian–Barremian, as previously stated. The associated small |
| 32 | assemblage of megaspores is consistent with this determination. Except for Trilobosporites |
| 33 | and Concavissimisporites, the use of several other genera commonly applied to Mesozoic |
| 34 | verrucate spores, such as Converrucosisporites and Impardecispora, is considered |
| 35 | unnecessary. To support this assertion, a Principal Components Analysis has been carried out |
| 36 | on 120 verrucate spores from sedimentological units An2-4, taking into account ten |
| 37 | morphological variables. The results show that specimens attributable to Trilobosporites are |
| 38 | well be separated from the main cluster, which corresponds to Concavissimisporites, |
| 39 | underlining the futility of using more than two genera for the species concerned. |
| 40 | |
| 41 | Keywords: verrucate spores; Lygodiaceae; Schizaeaceae; megaspores; Wealden facies; |
| 42 | Berriasian; Valanginian; Charentes. |
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| 48 | Néraudeau) |
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| | |

- **1. Introduction**

| 53 | Comparatively little is known about the palaeontological content of French Lower |
|----|---|
| 54 | Cretaceous continental deposits, mainly because of the scarcity of outcrop. In this connection, |
| 55 | the lignitic bone-bed of Angeac in south-west France is remarkable in providing a rare |
| 56 | glimpse of the terrestrial palaeoenvironment in this part of the world. The flora and fauna |
| 57 | were described by Néraudeau et al. (2012). Since the discovery of the site in 2008, hundreds |
| 58 | of ornithomimosaur bones have been found (Allain et al., 2014), along with other vertebrate |
| 59 | remains, reminiscent of the fossil record from the Wealden succession of southern England |
| 60 | and Wealden-type facies in Spain. |
| 61 | Apart from the deposits of Angeac (Néraudeau et al., 2012), Cadeuil (Platel et al., 1976) |
| 62 | and Saint-Agnant (Vullo et al., 2012), no other outcrops of Wealden-type facies have been |
| 63 | described from south-western France. They have, however, been encountered in boreholes at |
| 64 | Saint-André de Lidon (Ternet and Berger, 1968), Soubran (Kromm et al., 1970), and Saint- |
| 65 | Romain-de-Benet (Deák and Combaz, 1967). The last of these yielded a rich palynological |
| 66 | assemblage that was considered by Deák and Combaz (1967) to be Valanginian–Hauterivian |
| 67 | in age. A few lowest Cretaceous outcrops, corresponding to "Purbeck facies", have also been |
| 68 | reported from the island of Oléron (Lantz, 1958) and from Cherves-Richemont (Bourgueil et |
| 69 | al., 1986; Colin et al., 2004; Benoit et al., 2017) (Fig. 1A). |

Until recently, the time of deposition of the Angeac sediments was considered to be Hauterivian–Barremian. Although it was noted in Néraudeau et al. (2012, p. 5) that verrucate spores tend to be especially common in deposits of Berriasian–Valanginian age, this younger determination was based on the basis of the recovery of a dinoflagellate cyst (dinocyst) identified as *Odontochitina imparilis* (Duxbury) Jain et Khowaja-Ateequzzaman. However, this age was questioned by Benoit et al. (2017), who dated the deposits as Berriasian owing to the recovery of the charophyte *Clavator grovesii* var. *grovesii* (Harris) Martin-Closas. The

52

- aim of our paper is, therefore, to re-assess the composition of the continental palynological
 assemblage recovered from Angeac in the light of this recent development. We also discuss
 the taxonomy of Early Cretaceous verrucate spores.
- 80

81 **2. Geological setting**

82

The Angeac site (45° 37' 59.08" N–0° 5' 7.27" W) is situated 700 m north of the village 83 of Angeac-Charente, between Angoulême and Cognac (Fig. 1A). The lignitic beds of 84 "Wealden-facies" underlie Pleistocene alluvium laid down by the Charente River. Seven 85 sedimentological units, An1–An7, were described by Néraudeau et al. (2012) from the top to 86 the base respectively. An 1 is composed of 0.10-0.20 m of fine white sand, with abundant fish 87 micro-remains: it was mainly exposed during the first excavation campaigns of 2010 and 88 2011. An2 comprises 0.20–0.40 m of yellow and grey calcareous and marl deposits, with 89 numerous dinosaur bones, other vertebrate remains, ostracods, charophytes, and pieces of 90 wood (Fig. 1B). The stratification of this bed is difficult to determine because it has been 91 affected by the trampling of dinosaurs (dinoturbation). An3 is a calcareous conglomerate 92 0.20–40 m thick, with abundant vertebrate remains, ostracods, charophytes, insect coprolites, 93 pieces of wood and other plant debris (Colin et al., 2011; Néraudeau et al., 2012; Benoit et al., 94 2017). And is about 0.5–1.20 m thick. It is composed of blue-grey lignitic clay, oxidized at 95 the top, and has yielded numerous plant and well-preserved vertebrate remains. The base of 96 An5 is not visible. It is more than 0.7 m thick and composed of green clay that is devoid of 97 macrofossils, and includes sub-angular limestone blocks that are likely to have been reworked 98 into this deposit (Néraudeau et al., 2012). An6 and An7 consist of grey limestone and red clay 99 respectively, but they were only seen in 2010, after the initial trench was extended laterally. 100 They did not yield any fossils. 101

102

103 **3. Material and methods**

104

105 *3.1. Material and laboratory processing methods*

Six samples were processed by E. Masure in order to provide the palynological results 106 presented in Néraudeau et al. (2012). Duplicates of these preparations (LEM1-LEM6) were 107 sent to one of us (DJB) in 2011 and these have now been (re)-examined for this paper. LEM1 108 comes from the An2 unit (Fig. 1B). LEM2 and LEM3 are from the An3 unit. LEM2 109 corresponds to the organic fraction in which the size of the particles ranges between 10 and 63 110 μm, whereas LEM3 includes particles ranging in size from 10 and 100 μm. LEM4 comes 111 from the boundary between units An3 and An4. LEM5 is from the boundary between An4 112 and An5, and LEM6 comes from within the An5 unit. There was no palynological preparation 113 from the An4 unit. However, a sample from this unit was processed in 2010 by DJB at 114 Aberystwyth University, Wales (preparation QPR 3669: QPR is an Aberystwyth University, 115 palynological preparation identifier), using a standard procedure involving immersion of the 116 sample in 10% HCl followed by digestion in 58-62% HF and brief oxidation of the organic 117 residue in fuming HNO₃, washing after each stage until the residue was neutral (for further 118 information, see Batten, 1999). 119

Four samples were also processed in Aberystwyth for their mesofossil content: preparations MFP422 from the boundary between An3 and An4, and MFP415, MFP423 and MFP512 from unit An4. All yielded very small assemblages of megaspores. Their extraction involved initial soaking of 100 g of rock in warm water on a hot plate followed by immersion of the sample in a solution of $Na_4P_2O_7$ in order to disaggregate it as much as possible before sieving over a 70 µm mesh to reduce its bulk prior to standard palynological processing using HCl and HF. The megaspores were picked out of the aqueous residues under a

stereomicroscope and housed dry in micropalaeontological slides. Some of the specimens
recovered were mounted on scanning electron microscope (SEM) stubs to which carbon tabs
had been fixed, coated with platinum–palladium, and examined under a Hitachi S47-100 SEM
in the Institute of Biological, Environmental and Rural Sciences (IBERS), Aberystwyth
University.

132

133 *3.2. Specimen counts*

For palynological preparations LEM1–LEM4, and QPR 3669, 250 specimens were counted from the slides under a light microscope. LEM5 yielded just 94 specimens, and only a few examples of *Classopollis* were recovered from LEM6.

137

138 *3.3. Analysis of verrucate spores*

Owing to their morphological resemblance to the products of the extant fern Lygodium, 139 Cretaceous verrucate spores are considered to belong to the Schizaeales (Bolchovitina, 1961; 140 Ivanova in Samoilovitch and Mtchedlishvili, 1961; Fensome, 1987: see Discussion, Section 141 5). From the 1950s to the late 1980s, tens of species of eight genera (*Concavissimisporites*, 142 Converrucosisporites, Impardecispora, Maculatisporites, Rubinella, Trilobosporites, 143 *Tuberositriletes* and *Varirugosisporites*) were erected to accommodate upper Mesozoic 144 verrucate spores, principally on the basis of the outline of their amb, and the size and 145 arrangement of the vertucae on a specimen. This led to considerable overlap between generic 146 and specific diagnoses (Potonié and Kremp, 1954; Delcourt and Sprumont, 1955; Döring, 147 1964; Venkatachala et al., 1969; Dörhöfer, 1977), as previously underlined and discussed by 148 Batten (1986). 149

The history of classification of these spores is well summarized by Fensome (1987, pp.
19, 20), who broadened the diagnosis of *Concavissimisporites*, and considered

Tuberositriletes and Impardecispora to be junior synonyms. Tuberositriletes was erected by 152 Döring (1964) to accommodate Mesozoic forms similar to Convertucosisporites Potonié et 153 Kremp, 1954, because he considered the Carboniferous holotype of the type species of the 154 latter to be an atypical developmental stage. Venkatachala et al. (1969) designated 155 Impardecispora apiverrucata (Couper, 1958) as the type of Impardecispora. This species had 156 previously been accommodated in *Trilobosporites*, a genus that is distinguishable from 157 *Concavissimisporites* by the fact that it possesses one or several valvae at each apex, and/or 158 displays a more or less complete network of muri in the apical region (Fensome, 1987). 159 Srivastava (1977) considered Maculatisporites Döring, 1964 to be a junior synonym of 160 Concavissimisporites, but Fensome (1987, p. 19) noted that the morphological features of the 161 type species, *M. undulatus* Döring, 1964 do not clearly demonstrate this. The genera 162 Rubinella and Varirugosisporites were erected by Maljavkina (1949, emended by Potonié, 163 1960) and Döring (1965) respectively to accommodate Jurassic vertucate forms, but have 164 rarely been used by other authors. The diagnoses of both are very broad and can easily be 165 accommodated within Fensome's (1987) emended diagnosis of Concavissimisporites. 166 A few years later, Waksmundzka (1992) adopted an even more reductive approach and 167 placed Convertucosisporites, Concavissimisporites, Convertucitriletes, Gemmatriletes, 168 Tuberositriletes, Maculatisporites, Rubinella and Varirugosisporites in synonymy with 169 *Verrucosisporites*. She considered the basis for separation of these genera was insufficient, 170 this being mainly the general outline of the spores. However, Verrucosisporites was 171 established by Ibrahim (1933), and emended later by Potonié and Kremp (1954) and Krutzsch 172 (1959), among others, to accommodate Palaeozoic spores covered with warts or small 173 verrucae, which more closely resemble those produced by the Osmundaceae than the 174 Lygodiaceae. Also, placing in synonymy too many taxa can defeat the purpose of achieving a 175 sensible, stratigraphically and biologically relevant taxonomy. Indeed, Waksmundzka (1992, 176

p. 47) noted that the spores belonging to her emended version of *Verrucosisporites* are 177 imprecisely related to the Pteridophyta. Hence, we do not accept her solution to the problem 178 of too many genera for verrucate miospores. We prefer to base our identifications on 179 Fensome's major revision (1987) which, despite his synthesis and practical approach, has 180 never been discussed, most of the genera noted above still being encountered in the literature 181 (e.g., Burden and Hills, 1989; Song et al., 2000; Dejax et al., 2007). This not only renders 182 comparisons between assemblages challenging, but also diminishes the stratigraphic 183 significance of the species, because too many names are available for the same morphotype 184 and closely similar forms. 185

As a result, it was decided to carry out a Principal Components Analysis (PCA) of the 186 verrucate spores in the Angeac palynomorph assemblages, one of the purposes being to 187 determine whether *Concavissimisporites* can truly be regarded as encompassing a 188 morphological group that is well separated from *Trilobosporites* (see Section 4.2.). Indeed, 189 since numerous verrucate spores were recovered from the samples corresponding to 190 sedimentological units An2-4, we decided to identify them *a priori* following Fensome's 191 emended diagnoses as species of either Concavissimisporites or Trilobosporites. The PCA 192 was then carried out using 120 specimens from the three units (50, 20, and 50 for An2, An3 193 and An4, respectively). The purposes of this statistical analysis were to see whether it is 194 possible to (1) obtain several morphological groups that would cluster according to the 195 diagnoses of already established species, (2) determine those groups that can be divided into 196 two or more genera, and (3) ascertain whether there are any significant morphological 197 differences between the specimens recovered from the different lithological units. 198 Ten variables were taken into account (Fig. 2): 199

200 S: size of specimen (= maximum diameter).

- SA: smallest value of the reflex, acute or obtuse angle formed by the two tangents of the
 starting and arrival points of the arc or curve formed by the outline of the interradial region of
 a spore.
- LA: largest value of the reflex, acute or obtuse angle formed by the two tangents of the
- starting and arrival points of the arc or curve formed by the outline of the interradial region of

a spore.

- ²⁰⁷ MiD/MaD: ratio between the minimum and maximum distances from the centre to the edge of
- the spore (excluding the verrucae).
- mDV: mean diameter of verrucae on one spore; 32 verrucae were counted on each specimen,
- when possible. Most of the means have been calculated based on approximately 20 values.
- ²¹¹ mHV: mean height of verrucae on one spore.
- 212 STDV: standard deviation of the diameter of verrucae on one spore.
- STHV: standard deviation of the height of verrucae on one spore.

ET: exine thickness.

- PC: per cent cover of the verrucae on the proximal face of a spore. 0.15: 0–30%; 0.45: 31–
- 216 60%; 0.75: 61–94%; 0.95: 95–100%.
- All of the values have been divided by the size of the specimen, and scaled. They were measured on clear images of the specimens under a light microscope. The angles were measured using the free java application *OnScreenProtractor*. The various lengths and diameters were measured using the software *Inkscape*. The PCA was carried out using the software *Past*.

222

3.4. Repository

All of the palynological preparations and the megaspores illustrated are housed in the Geological Institute of the University of Rennes 1, under collection numbers IGR-PAL-5735–

- IGR-PAL-5753 for the palynological slides, and IGR-PAL-5780–IGR-PAL-5784 for the
 megaspores illustrated.
- 228

229 **4. Results**

230

231 4.1. Palynological assemblages

Altogether 34 taxa (excluding those placed in comparison apart from

Concavissimisporites sp. cf. C. crassatus) have been identified at genus or species level in the 233 seven samples examined (Fig. 3). The most common of these are the cheirolepidiaceous 234 pollen grain Classopollis (Fig. 4C, D), verrucate spores such as Concavissimisporites 235 montuosus (Döring) Fensome (Fig. 5J, K, M), Concavissimisporites apiverrucatus (Couper) 236 Döring (Fig. 5A, B), Concavissimisporites exquisitus (Singh) Fensome (Fig. 5C, D) and 237 Trilobosporites canadensis Pocock (Fig. 6H), the papillate spore Pilosisporites 238 trichopapillosus (Thiergart) Delcourt et Sprumont (Fig. 4P), the hilate spore Aequitriradites 239 verrucosus (Cookson et Dettmann) Cookson et Dettmann (Fig. 4G), and various smooth 240 walled spores belonging to the genera Cyathidites, Deltoidospora, and Gleicheniidites (Fig. 241 4H, J). Freshwater algae such as Ovoidites spriggi (Cookson et Dettmann) Zippi (Fig. 4N) are 242 relatively common, but no more dinoflagellate cysts were encountered during the re-243 examination. 244

An2 yielded an assemblage of 16 species (Fig. 3), numerically and taxonomically dominated by verrucate spores (Fig. 7). It is the only assemblage in which there are no *Classopollis* pollen, the gymnosperm component being represented by only two inaperturate pollen grains referable to *Araucariacites australis* Cookson. Eighteen species were recovered from the An3 unit. Verrucate spores are less diverse than in the An2 unit with eight species, but the gymnosperms are better represented, with numerous specimens referable to

| 251 | Classopollis torosus (Reissinger) Couper, and bisaccates such as Abietineaepollenites sp. |
|-----|--|
| 252 | (Fig. 4A). This is the only assemblage in which a specimen of <i>Cicatricosisporites hallei</i> |
| 253 | Delcourt et Sprumont was found (Fig. 4M). The sample taken from the boundary between |
| 254 | An3 and An4 yielded 16 species, five of which are exclusive to this sample: Callialasporites |
| 255 | dampieri (Balme) Dev (Fig. 4B), Densoisporites microrugulatus Brenner (Fig. 4K), cf. |
| 256 | Foraminisporis wonthaggiensis (Cookson et Dettmann) Dettmann (Fig. 4L), |
| 257 | Microreticulatisporites sp. cf. M. diatretus Norris (Fig. 4I), and Podocarpidites ellipticus |
| 258 | Cookson (Fig. 4F). The assemblage from the lignitic clays of An4 is the most diverse, having |
| 259 | yielded 21 species, three of which were recorded only from this level: Retitriletes sp. (Fig. |
| 260 | 4O), Cerebropollenites mesozoicus Couper, and Vitreisporites pallidus (Reissinger) Nilsson |
| 261 | (Fig. 4E). This sample yielded proportionally the greatest abundance of <i>Classopollis</i> (Fig. 7), |
| 262 | but also, and in common with An2, the most diverse verrucate spore assemblage (nine |
| 263 | species). The An4/An5 sample yielded a moderately diverse, albeit small assemblage of 15 |
| 264 | species, including six verrucate taxa. Overall, the relative abundance of <i>Classopollis</i> tends to |
| 265 | decrease upwards in the succession from An4/An5 to An2, whereas the abundance of the |
| 266 | verrucate and smooth walled spores increases greatly in this direction. |
| 267 | Also, significant in this context is the fact that specimens identified as Trilobosporites |
| | |

Also, significant in this context is the fact that specimens identified as *Trilobosporites* are more abundant in the palynomorph assemblage recovered from An2 than in the older
 deposits, and that overall, verrucate spores in the An2 assemblage are generally larger and
 more sparsely sculptured than those recovered from the other units (see PCA plots, Figs 8, 9).

271

4.2. Megaspores

The four samples processed for their mesofossil content yielded only a few megaspores and fragments thereof, most of which are referable to the genus *Erlansonisporites* (Figs. 10, 11A, B), and faecal pellets, probably attributable to the activities of termites (Colin et al.,

2011). The sample from the boundary between An3 and An4 (preparation MFP422) also 276 proved to contain a few specimens of a form tentatively recorded here as *Minerisporites* sp. 277 (Fig. 11C-F). Apart from a couple specimens that are currently unidentifiable without 278 recourse to examination under an SEM, all other megaspore remains are too damaged or 279 fragmentary to be positively identified: among these are possible representatives of 280Paxillitriletes. Associated with the megaspores in MFP422 are isolated, comparatively large 281 specimens of the freshwater algae Schizosporis reticulatus Cookson et Dettmann emend 282 Pierce and Ovoidites spriggii (Cookson et Dettmann) Zippi. 283 Most of the specimens of *Erlansonisporites* (Figs. 10E, F, 11A, B) are closely similar to 284 Erlansonisporites sp. sensu Batten, 1975, pl. 13, fig. 6, a selaginellalean megaspore from the 285 upper Berriasian-Valanginian Ashdown Formation in Sussex, south-east England (e.g., 286 Batten, 2011): this is consistent with the suggested revised age determination of the Angeac 287 deposits herein. Numerous microspores, all exhibiting the same morphology, are lodged 288 within the muri of its reticulate sculpture, especially on the proximal surface close to the 289 triradiate flange. In the dispersed state, these are probably attributable to the small spore 290 genus Patellasporites, or perhaps Uvaesporites. 291 The specimens of *Minerisporites* sp. are atypical for this genus in that, although zonate, 292

they have a virtually smooth surface and a triradiate flange that is of uneven elevation, indeed it can be partly spinose (Fig. 11C). The outer edge of the zona is also uneven to weakly spinose, features that might equally suggest a representative of *Henrisporites*, albeit again atypical. More specimens are required before this form can be satisfactorily identified.

297

4.3. Systematic palaeontology of the vertucate spores

The PCA2 plot (Fig. 9) supports Fensome's (1987) taxonomic approach to the
 identification of vertucate spores, the specimens from Angeac only being attributable to two

| 301 | genera, Concavissimisporites and Trilobosporites. They are morphologically well separated, |
|-----|---|
| 302 | principally on the account of their size, the relative size of their verrucae, and the variation in |
| 303 | the dimensions of verrucae on a specimen. Three species of Trilobosporites, and eight species |
| 304 | of Concavissimisporites have been identified, along with several intermediate forms. Except |
| 305 | for Concavissimisporites apiverrucatus, specimens assigned to a particular species are |
| 306 | clustered together, although the boundaries between the clusters are not always clearly |
| 307 | defined. It was thought that the semi-quantitative variable corresponding to the per cent cover |
| 308 | of verrucae (PC) might be biasing the results of the PCA in assuming too much importance |
| 309 | compared to the other variables. Thus, another PCA was run without the per cent cover |
| 310 | variable, but the results were very similar, indicating that this did not significantly affect the |
| 311 | plot. |
| 312 | |
| 313 | Genus Concavissimisporites Delcourt et Sprumont, 1955, emend. Fensome, 1987. |
| 314 | |
| 315 | Type species: Concavissimisporites verrucosus Delcourt et Sprumont, 1955 emend. Delcourt |
| 316 | et al., 1963. |
| 317 | |
| 318 | Concavissimisporites apiverrucatus (Couper, 1958) Döring, 1965 (Fig. 5A, B) |
| 319 | |
| 320 | Material. Angeac units An2, An3/4, An4, An4/5: 11 specimens, 57 (72) 105 µm in diameter. |
| 321 | Remarks. Concavissimisporites apiverrucatus is differentiated from the other species of this |
| 322 | genus in that the sculpture of the apices is slightly to distinctly coarser than over the poles and |
| 323 | in interradial regions. In the case of the Angeac specimens, the polar and interradial regions |
| 324 | are almost smooth (Fig. 5B), granulate (Fig. 5A), or sculptured with verrucae. Specimens |
| 325 | attributed to this species are the only ones that do not form a united cluster on PCA2 (Fig. 9), |

- principally because the main diagnostic characters mentioned above have not been considered
- 327 as a variable, these being very difficult to calculate.
- 328 Occurrence. This species occurs widely in rocks ranging in age from latest Jurassic to Albian
- (e.g., Fensome, 1987; Burden and Hills, 1989). In France, it has been recorded, as
- 330 Trilobosporites apiverrucatus Couper, from "Wealden" (Delcourt and Sprumont, 1959),
- ³³¹ lower Valanginian (Vakhrameev and Kotova, 1980), upper Barremian–lower Aptian
- (Herngreen, 1971) and upper Albian (Bardet et al., 1991) deposits.

333

Concavissimisporites sp. cf. *C. crassatus* (Delcourt et Sprumont, 1955) Delcourt et al., 1963

335 (Fig. 5L, N)

336

- 337 *Material*. Angeac unit An2: three specimens 61 (63) 67 µm in diameter.
- *Remarks.* Specimens Q35.4 and B28.4 (Fig. 5L, N) fit the description given by Delcourt et al.
- (1963, p. 285), bearing in mind the changes made by Fensome to the generic diagnosis (1987,
- $_{340}$ $\,$ p. 19). No formal size range is given by Delcourt et al., but their illustrated specimen is 78 μm
- in diameter. Delcourt and Sprumont (1955, p. 26) stated that the "radius" of their form *crassatus* is 40–55 µm. As a result, we only compare our specimens to this species because
 they are smaller.
- On the PCA2 (Fig. 9), the squares pertaining to *C*. sp. cf. *crassatus* are situated at the edge of the cluster that corresponds to *C. montuosus*. It could be argued that these specimens are also identifiable as *C*. sp. cf. *C. montuosus*, the limiting character being the small size of the verrucae.
- Occurrence. Concavissimisporites crassatus was originally described from Wealden-type
 strata in Belgium (Delcourt and Sprumont, 1955; Delcourt et al., 1963). It has not been
 encountered previously in France, but has been found in Albian strata of the Bay of Biscay

(Batten, 1979), Berriasian–mid-Valanginian strata in the Netherlands (Burger, 1966), upper 351 Berriasian, and upper Barremian-lower Aptian rocks in Germany (Dörhöfer, 1977; Lister and 352 Batten, 1995), and within Cenomanian deposits of Spain (Solé de Porta, 1978). It was also 353 recorded by Fensome (1987) from Upper Jurassic to lower Valanginian strata in western 354 Canada. 355 356 Concavissimisporites exquisitus (Singh, 1971) Fensome, 1987 (Fig. 5C, D) 357 358 Material. Angeac units An2, An3, An3/4, An4, An4/5: 19 specimens, 47 (59) 75 µm in 359 diameter. 360 *Remarks.* This species is defined by its triangular to convex outline, coupled with bulbous 361 verrucae 6–10 µm wide. In his original diagnosis, Singh (1971, p. 116) stated that the 362 verrucae are spaced 1-4 µm apart. The verrucae on the specimens from Angeac are more 363 closely spaced overall, being separated by no more than 1 µm. The size-range given by Singh 364 is 68 (71) 84 µm, but this is based on only four specimens, and the only specimen of this 365 species recovered by Fensome is 59 µm in diameter (1987, pl. 4, fig. 11). Some relatively 366 small, rounded specimens, such as R40.2 (Fig. 5E), are referred to Concavissimisporites sp. 367 cf. C. exquisitus because they show proportionally smaller verrucae. They are more similar to 368 Verrucosisporites major (Couper) Burden et Hills, a form that occurs widely in Jurassic and 369 Cretaceous strata. 370 The cluster corresponding to this species on the PCA2 plot is very close to that 371 representing Concavissimisporites ferniensis (Fig. 9), which is, however, distinguished from 372 C. exquisitus in having significantly lower vertucae separated by narrow canals that form a 373

more or less distinct negative reticulum.

| 375 | Occurrence. This species was originally recovered from Albian strata in Alberta, Canada |
|------------|--|
| 376 | (Singh, 1971), but Fensome (1987) found it in upper Jurassic deposits. It has not been |
| 377 | encountered previously in France, but Mohr (1989) mentioned its presence in upper |
| 378 | Tithonian–lower Berriasian strata in Portugal. |
| 379 | |
| 380 381 | Concavissimisporites ferniensis (Pocock, 1970) Fensome, 1987 (Fig. 5F–H) |
| 382 | Material. Angeac units An2, An3, An3/4, An4: seven specimens 45 (56) 78 µm in diameter. |
| 383 | Remarks. For differences between Concavissimisporites ferniensis and C. exquisitus, see |
| 384 | remarks above under the latter species. In common with C. exquisitus, the specimens assigned |
| 385 | to C. ferniensis are very similar to Verrucosisporites major, which however differs in usually |
| 386 | being smaller and in lacking a negative reticulum (see Couper, 1958). |
| 387 | Occurrence. This species has only been recovered from Jurassic sedimentary rocks in western |
| 388 | Canada (Fensome, 1987), but very similar forms, identified as Leptolepidites major Couper, |
| 389 | have been found in various French Jurassic, Albian and Cenomanian strata (Fauconnier, 1979; |
| 390 | Fechner and Dargel, 1989; Bignot et al., 1994). |
| 391 | |
| 392 | Concavissimisporites montuosus (Döring, 1964) Fensome, 1987 (Fig. 5J, K, M) |
| 393 | |
| 394 | Material. Angeac units An2, An3, An3/4, An4, An4/5: 26 specimens 52 (72) 105 µm in |
| 395 | diameter. |
| 396 | Remarks. This species is characterized by being sculptured with typically well-spaced |
| 397 | verrucae $2-5 \ \mu m$ in diameter. It is the most common verrucate spore in the Angeac |
| 398 | succession. A few specimens have been referred to Concavissimisporites sp. cf. C. montuosus |
| 399 | (e.g., N26.1, not illustrated), because they have more closely spaced and smaller verrucae. |

| 400 | Occurrence. The stratigraphic range of Concavissimisporites montuosus extends from upper |
|-------------------|--|
| 401 | Oxfordian to Albian. It is a distinctive element in many Volgian–Valanginian palynofloras |
| 402 | (Fensome, 1987). It has not been found in France previously, but has been recovered, as |
| 403 | Converrucosisporites montuosus Döring, from lowest Cretaceous strata in north-western |
| 404 | Germany (Dörhöfer and Norris, 1977). According to these authors, an abundance of this form |
| 405 | is characteristic of their upper Berriasian palynofloral suite "Hils 1". |
| 406 407 408 | Concavissimisporites robustus Dörhöfer, 1977 (Fig. 5I) |
| 409 | <i>Material</i> . Angeac units An2, An3: two specimens 59 and 66 μ m in diameter. |
| 410 | Remarks. Concavissimisporites robustus differs from the other species encountered in having |
| 411 | a more or less uniformly thick exine coupled with poorly delineated verrucae that are either |
| 412 | sparsely or densely distributed. The two specimens recovered from Angeac are weakly |
| 413 | sculptured. The PCA2 plot shows that they cluster closest to the specimens identified as C . |
| 414 | uralensis, mainly because of the scarcity of the verrucae on the proximal face. |
| 415 | Occurrence. This species was originally described from the upper Berriasian palynofloral |
| 416 | suites "Hils 1" and "Hils 2" in north-western Germany (Dörhöfer, 1977). It has also been |
| 417 | recorded from uppermost Jurassic strata in western Canada (Fensome, 1987) and middle |
| 418 | Berriasian deposits in eastern Maryland, USA (Doyle, 1983). |
| 419 | |
| 420 | Concavissimisporites uralensis (Bolchovitina, 1961) Fensome, 1987 (Fig. 6D) |
| 421 | |
| 422 | Material. Angeac units An2, An4: three specimens 56 (68) 90 µm in diameter. |
| 423 | Remarks. This species can appear very similar to the specimens identified here as |
| 424 | Concavissimisporites apiverrucatus: they differ in being more sparsely sculptured. They also |

| 425 | differ from specimens of C. robustus in having a thinner exine. It could be argued that they |
|-----|---|
| 426 | are intermediate between C. apiverrucatus and C. robustus. However, the PCA clustered all |
| 427 | three specimens together, quite far away from the other clusters, which is why we decided to |
| 428 | regard them as belonging to a distinct species (see Section 6.2 for the biological relevance of |
| 429 | this decision). |
| 430 | Occurrence. Concavissimisporites uralensis was originally described from Barremian-Aptian |
| 431 | strata in western Kazakhstan (Bolchovitina, 1961). It has also been recorded from Ryazanian |
| 432 | (lowest Berriasian) strata in western Canada (Fensome, 1987). |
| 433 | |
| 434 | Concavissimisporites verrucosus Delcourt et Sprumont, 1955 emend. Delcourt et al., 1963 |
| 435 | (Fig. 6A, B) |
| 436 | |
| 437 | Material. Angeac units An3, An4, An4/5: six specimens 73 (81) 100 µm in diameter. |
| 438 | Remarks. This species differs from the other verrucate spores from Angeac by its large size, |
| 439 | and in having relatively small, closely spaced verrucae. The specimen D20.4 (Fig. 6C) is |
| 440 | considered to be an intermediate form between C. verrucosus and C. montuosus, because it is |
| 441 | sculptured with relatively large, closely spaced verrucae. |
| 442 | Occurrence. The stratigraphic range of C. verrucosus extends from Middle Jurassic to Lower |
| 443 | Cretaceous. In western Europe, it occurs principally within lowest Cretaceous and |
| 444 | "Wealden"-type strata (Burger, 1966; Bolchovitina, 1971; Dörhöfer and Norris, 1977). |
| 445 | |
| 446 | Genus Trilobosporites Pant, 1954 ex Potonié, 1956 emend. Fensome, 1987 |
| 447 | |
| 448 | Type species: Trilobosporites hannonicus (Delcourt et Sprumont, 1955) Potonié, 1956 |
| 449 | |

450 Trilobosporites aequiverrucosus Dörhöfer, 1977 (Fig. 6I)

451

452 *Material*. Angeac unit An4: one specimen, 100 µm in diameter.

Remarks. This species differs from *Trilobosporites canadensis* in having proportionally 453 smaller unfused, closely spaced vertucae in polar and interradial regions. As stated by 454 Dörhöfer (1977, pp. 53, 54), this species is considered to include many specimens previously 455 attributed to Trilobosporites bernissartensis (Delcout et Sprumont) Potonié, for which the 456 type and size of the sculptural elements were not specified. The re-illustrations of the holotype 457 and paratypes in Delcourt et al. (1963, pl. 43, figs. 11–14) are considered inadequate, because 458 several morphotypes are displayed. As a result, Dörhöfer considered that a precise 459 interpretation of *T. bernissartensis* is not possible. 460 Occurrence. The species was originally described from upper Berriasian-lower Valanginian 461 beds in western Germany (Dörhöfer, 1977). As Trilobosporites bernissartensis, it has been 462 found in Wealden-type strata in Belgium (Delcourt and Sprumont, 1955; Delcourt et al., 463 1963), and in Purbeck and Wealden beds in southern England and stratigraphically equivalent 464 deposits in western France (Couper, 1958; Lantz, 1958; Deák and Combaz, 1967). It has also 465 been reported from lowest Aptian strata in south-eastern France (de Reneville and Raynaud, 466 1981), and within upper Barremian–lower Aptian strata in northern France (Herngreen, 1971). 467 However, the specimen illustrated by Herngreen (1971, pl. 3, fig. 1) is in fact attributable to 468 Trilobosporites hannonicus (Delcourt et Sprumont) Potonié, so we do not accept this record 469 of occurrence. 470

471

472 Trilobosporites aornatus Döring, 1965 (Fig. 6E, F)

473

474 *Material*. Angeac units An2, An3, An4: three specimens 68 (78) 96 μm in diameter.

| 475 | <i>Remarks</i> . The morphology of the specimens recorded as <i>Trilobosporites aornatus</i> have a |
|-----|---|
| 476 | maculate sculpture and a large verruca or rounded valva in equatorial radial regions. K28.3 |
| 477 | (Fig. 6G) is referred to Trilobosporites sp. cf. T. aornatus, because of the presence of a few |
| 478 | verrucae close to each of the angles, in common with the specimen illustrated by Norris as |
| 479 | Trilobosporites obsitus Norris (1969, pl. 106, fig. 7) (see Section 5.2. for the biological |
| 480 | relevance of this species). |
| 481 | Occurrence. Döring (1965) described this species from Upper Jurassic-lowermost Cretaceous |
| 482 | strata in eastern Germany. In southern England, it is typical of the upper Berriasian-lower |
| 483 | Valanginian palynofloral suite C of Norris (1969), and as discussed by Dörhöfer and Norris |
| 484 | (1977). In North America, it ranges into the lower Barremian (Burden and Hills, 1989). |
| 485 | |
| 486 | Trilobosporites canadensis Pocock, 1962 (Fig. 6H) |
| 487 | |
| 488 | Material. Angeac units An2, An3, An3/4, An4, An4/5: 11 specimens 69 (88) 105 µm in |
| 489 | diameter. |
| 490 | Remarks. Trilobosporites canadensis is characterized by its coarse interradial and polar |
| 491 | verrucae, which sometimes merge to form elongate elements, along with one or two large, |
| 492 | pronounced, strongly projecting verrucae or valvae in equatorial radial regions. This is the |
| 493 | most common form of Trilobosporites at Angeac, especially in the An2 unit. |
| 494 | Occurrence. This species was originally described from "Neocomian" deposits in western |
| 495 | Canada (Pocock, 1962). In Western Europe, it has been reported from upper Berriasian-lower |
| 496 | Valanginian beds in western Germany (Dörhöfer, 1977), probable Berriasian Purbeck beds in |
| 497 | southern England (as T. cf. canadensis; Hunt, 1985), Tithonian–Berriasian deposits in |
| 498 | Portugal (Mohr, 1989) and within upper Barremian-lower Aptian strata in northern France |
| 499 | (Herngreen, 1971). However, the morphology of the specimens illustrated in Herngreen's |

paper (pl. 3, figs. 2–4) is closer to that of *Concavissimisporites montuosus* than to *T*. *canadensis*.

502

503 5. Discussion

504

505 5.1. Age determination and comparisons

We discuss the previous identification in Néraudeau et al. (2012) of the dinoflagellate 506 cyst Odontochitina imparilis (Duxbury) Jain et Khowaja-Ateequzzaman in the Angeac 507 assemblage. In his description of Muderongia imparilis, Duxbury (1980, p. 128) indicated the 508 presence of three well-developed horns as a diagnostic feature, with the lateral horn being 509 greatly reduced. A lateral horn seems to be present on the Angeac specimen, but it is folded 510 behind the periblast. There are two antapical horns, one shorter than the other. The longer of 511 the two is very corroded. A vestigial apical horn is present. The original description also states 512 that the endoblast and periblast are joined together in the epicystal region, which does not 513 seem to be the case of the Angeac specimen, the endoblast being proportionally much smaller 514 than the periblast. 515

The species has usually been found within Hauterivian–Barremian deposits (Duxbury, 516 1980; Jain and Khowaja-Ateequzzaman, 1984; Leereveld, 1997), hence the previous 517 attribution of a Hauterivian-Barremian age to the Angeac deposits. However, the species has 518 also been recorded from upper Valanginian strata (Burger, 1996). Although the poor 519 preservation of the Angeac specimen renders the identification questionable, the short range 520 of the species may well be related to the general scarcity of this form and linked to specific 521 environmental conditions (Masure, pers. comm. 2017), so a find of O. imparilis in older 522 deposits is not impossible. 523

The Angeac specimen is also similar to *Muderongia simplex* Alberti, which has one apical, two lateral, and two antapical horns. The holotype of this species is 151 μ m long, which is almost the same as the Angeac specimen at 152 μ m. However, they differ in that the latter possesses a proportionally small endoblast, and does not display a second lateral horn, although it is possible that has been destroyed or is obscured by folding. The stratigraphic range of *Muderongia simplex* extends from upper Tithonian to upper Valanginian (Riding et al., 2000).

Dating Lower Cretaceous deposits using miospores can be challenging, the forms 531 encountered often being long ranging. For instance, Aequitriradites verrucosus and 532 *Pilosisporites trichopapillosus* are present through all of the Lower Cretaceous in western 533 Europe. In France, A. verrucosus has only been reported recently within Cenomanian deposits 534 (Fleury et al., 2017). Batten (1979) recorded it from Aptian–Albian strata in the Bay of 535 Biscay. It occurs rarely elsewhere in Europe but has nonetheless been documented from 536 Aptian and Wealden deposits in England (Ruffell and Batten, 1994 and Batten unpublished, 537 respectively) and in Wealden-type facies in Germany (Döring, 1964). 538 Pilosisporites trichopapillosus ranges from uppermost Jurassic (Couper and Hughes,

Pilosisporites trichopapillosus ranges from uppermost Jurassic (Couper and Hughes,
1963) to Upper Cretaceous (Robaszynski et al., 1985), but is more frequently recovered from
Lower Cretaceous deposits (Bolchovitina, 1971; Dörhöfer, 1977; Batten, 1979; Doubinger
and Mas, 1981; Batten and Li, 1987). In France, it has only been documented from Tithonian
strata (Couper and Hughes, 1963).

Nevertheless, dating can be more precise if the composition of entire assemblages is
taken into account. A few attempts at stratigraphic correlations using Lower Cretaceous
miospore assemblages were made during the late 1960s and early 1970s. A method
introduced by Hughes and Moody-Stuart (1969) consisted of using several types of *Cicatricosisporites* to correlate between different beds in a Wealden borehole and an outcrop

in south-east England. Their abandonment of Linnaean nomenclature in favour of biorecords
(Hughes and Moody-Stuart, 1969; Hughes, 1970) meant that their method did not gain
widespread support among palynologists (Batten, 1986). Regardless, we could not apply a
similar approach to the Angeac assemblages using species of *Cicatricosisporites* because only
a few specimens of this taxon were recovered.

Another method involved the use of palynological assemblage-types for correlating 554 Wealden deposits in southern England (Batten, 1973). Seventeen assemblages were described, 555 taking into account preservation state, diversity of miospores, average size of trilete spores, 556 abundance of brown wood, and the abundance of certain taxa. The total Angeac assemblage 557 (i.e. the overall composition of the assemblages recovered from An2-5) is very similar to 558 Batten's Assemblage Type 15 (AT 15). This was characterized by the common presence of 559 *Concavissimisporites*, coupled with a "Trilete spore content dominated by average-sized or 560 large forms; miospore diversity average or large; general state of preservation of miospores 561 good or fair; brown wood and cuticle P, C or F [present, common, or frequent]; Pilosisporites 562 P or C [present or common], *Cicatricosisporites* C or V [common or very 563 common].....Trilobosporites occurs in more than 70% of the assemblages referable to this 564 AT" (Batten, 1973, pp. 13, 14). The only difference from this description is the scarcity of 565 *Cicatricosisporites.* Following the same study, and using current stratigraphic terminology 566 (e.g., Batten, 2011), this assemblage-type has mostly been recovered from the Ashdown 567 Formation, including the Fairlight Clays facies, and the Grinstead Clay Formation, all of 568 which are part of the Hastings Group, which was deposited between the mid Berriasian and 569 the late Valanginian. 570

The Angeac assemblage overall is also very similar to those recovered from the Bückeberg Formation, especially the upper Berriasian palynofloral suites "Hils 1" and "Hils 2", which are characterized by abundant *Concavissimisporites montuosus*, large

Trilobosporites, and the relative scarcity of Cicatricosisporites (Dörhöfer, 1977; Dörhöfer and 574 Norris, 1977). In France, Berriasian palynological assemblages have been recovered from 575 Jura (eastern France) and potentially Oléron island (Fig. 1). The Angeac assemblage is 576 somewhat similar to that recovered by Médus and Mojon (1991) from upper Berriasian strata 577 of southern Jura in that the latter is characterized by an abundance of *Densoisporites* 578 microrugulatus and Trilobosporites. It was compared to palynofloral suite "Hils 4" (Médus 579 and Mojon, 1991, p. 57), but only a few illustrations were provided to support the 580 identifications, and D. microrugulatus is not abundant in the Angeac assemblage. The 581 uppermost Berriasian assemblage from Jura recovered earlier by Taugourdeau-Lantz and 582 Donze (1971) is less similar, with few vertucate spores and several dinoflagellate cyst species 583 in association. The palynoflora recovered from the "Purbeck beds" of Oléron island (Lantz, 584 1958) includes similar forms of Trilobosporites, but the types of Concavissimisporites are less 585 diverse, and the specimens illustrated seem comparatively small. 586 The Angeac assemblage is, however, less clearly correlated with younger palynofloras, 587 such as that recovered from the Wealden-facies, dinosaur-bearing pit of Bernissart (Belgium, 588 Dejax et al., 2007), considered to be Hauterivian in age. The samples from this pit yielded 589 specimens of *Trilobosporites* that are similar to the species *T. hannonicus* (Delcourt et 590 Sprumont) Potonié, which has not been recorded from Angeac, more species of 591 *Cicatricosisporites*, and angiospermid pollen grains. Correlations with the assemblage 592 recovered from Saint-Romain-de-Benet, regarded as Valanginian-Hauterivian in age, are 593 quite difficult to make because of a lack of precision in the description of the assemblage as 594 indicated by such statements as "on n'observe pratiquement pas de pollens 595 d'Angiospermes" in Deák and Combaz (1967, p. 72), meaning that they "almost did not 596

⁵⁹⁷ observe any angiosperm pollen grains". Their stratigraphic range chart (p. 71) is rather

⁵⁹⁸ inaccurate because most of the species they note as restricted to the Valanginian–Hauterivian

have, in fact, wider ranges. In any case, they do not seem to have encountered many vertucate
spores, and their species of *Trilobosporites* are quite different from those recovered from
Angeac.

Although most of the palynomorphs identified herein indicate a stratigraphic
distribution that is not confined to the lowest Cretaceous, previous observations in Néraudeau
et al. (2012, p. 5) and our new data suggest that the Angeac assemblage is more likely to have
been deposited during the Berriasian–early Valanginian rather than in the Hauterivian–
Barremian, as previously stated.

607

5.2. Biological relevance of the verrucate spores

The family Schizaeaceae originally included five extant genera: *Schizaea* Sm., *Actinostachys* Wall., *Mohria* Sw., *Anemia* Sw., and *Lygodium* Sw., which are widely distributed in predominantly tropical or southern warm-temperate parts of the world (Tryon and Lugardon, 1991). However, Smith et al. (2006) considered that these genera in fact belong to three families of the order Schizaeales: Schizaeaceae (for *Schizaea* and *Actinostachys*), Anemiaceae (for *Anemia*, including *Mohria*), and Lygodiaceae (for *Lygodium*).

Unequivocal members of this order first appeared in the Early or Middle Jurassic, but 616 they did not become widespread until the Early Cretaceous, as indicated by occurrences of 617 their spores in numerous deposits throughout Eurasia (Bolchovitina, 1971; Van 618 Konijnenburg-Van Cittert, 2002). The Schizaeaceae produce monolete, bean-shaped spores, 619 with diverse ornamentation. The Anemiaceae have trilete spores with coarse, compact ridges. 620 These have abundant Lower Cretaceous analogues, such as Cicatricosisporites, and 621 specimens found *in situ* within fertile plant organs associated with the ferns *Ruffordia*, 622 Pelletieria, and Schizaeopsis (e.g., Hughes and Moody Stuart, 1966). Lygodium spores are 623

| 624 | trilete, and can be tuberculate, verrucate or reticulate (Tryon and Lugardon, 1991). Thus, |
|------|---|
| 625 | Early Cretaceous verrucate spores have often been compared to the genus Lygodium (e.g., in |
| 626 | Delcourt and Sprumont, 1955; Bolchovitina, 1961; Ivanova in Samoilovitch and |
| 627 | Mtchedlishvili, 1961; Fensome, 1987). |
| 628 | Mesozoic spores comparable to Lygodium have only been found in situ in Jurassic rocks |
| 629 | in northern England (Van Konijnenburg-Van Cittert, 1981), but they have a reticulate |
| 630 | sculpture. They have been encountered in sorophores from Turonian deposits in New Jersey, |
| 631 | USA (Gandolfo et al., 2000), and compared to the extant species Lygodium palmatum |
| 632 | (Bernh.) Sw. on account of their trilete, psilate spores, among other characters. Dispersed |
| 633 | spores are much more abundant. Bolchovitina (1961) produced a comprehensive monograph |
| 634 | on fossil and extant spores of the former family Schizaeaceae (now Schizaeales), in which she |
| 635 | listed 62 species of extant Lygodium (Smith et al., 2006 considered that the Lygodiaceae |
| 636 | comprise only about 25 species), remarked on and illustrated 27 of these, and listed, |
| 637 | commented, described or re-described, and illustrated 46 species of dispersed fossil spores |
| 638 | referable to Lygodium. |
| 639 | The spores of the extant species are usually described according to their shape, |
| 640 | thickness of exine, colour and sculpture. The last of these may vary within a species, such as |
| 641 | in L. flexuosum (L.) Sw. and L. japonicum (Thunb.) Sw., which produce smooth to slightly |
| 642 | tuberculate spores. However, when a species also displays a coarser sculpture, several authors |
| 643 | (e.g., Couper, 1958; Dettmann and Clifford, 1991) have considered associated smooth spores |
| 644 | to be immature forms. Spore polymorphism has been recognized in a few anemiaceous |
| 645 | species (Dettmann and Clifford, 1991), but it would seem that in general mature spores of |
| 646 | extant species of Lygodium can be well separated on the basis of their sculpture (Bolchovitina, |
| 647 | 1961; Tryon and Lugardon, 1991; see also SEM micrographs of Lygodium spores in the |
| - 10 | Correll University website) |

648 Cornell University website).

| 649 | Hence, most of the morphological clusters of the verrucate spore species displayed by |
|-----|--|
| 650 | PCA2 (Fig. 9) could well also have biological significance. Specimens assigned to |
| 651 | Concavissimisporites uralensis, which are defined by their weak sculpture, could in fact be |
| 652 | immature forms of other species of Concavissimisporites. The same applies to Trilobosporites |
| 653 | aornatus, which could comprise immature specimens of T. aequiverrucosus or T. canadensis. |
| 654 | Intermediate forms such as specimen K28.3 (Fig. 6G) would support this suggestion. The |
| 655 | most abundant verrucate spore in the Angeac palynomorph assemblages, |
| 656 | Concavissimisporites montuosus, is similar to the spores of the extant species Lygodium |
| 657 | circinatum (Burm. f.) Sw., in that both display quite well spaced, massive verrucae. |
| 658 | Specimens assignable to the genus Trilobosporites are somewhat similar to the spores of |
| 659 | Lygodium merrillii Copel., which display coarse distal muri that appear as a thickening of the |
| 660 | exine in apical regions on the proximal face. However, no spores of extant species of |
| 661 | Lygodium seem to develop valvae, which is all the more reason to maintain Trilobosporites as |
| 662 | a separate genus for fossil spores with this character. Some species of Concavissimisporites, |
| 663 | such as Concavissimisporites exquisitus, also do not seem to have any modern analogues. |
| 664 | |

- 665 5.3. Palaeoenvironmental implications
- 666

The great abundance of verrucate spores in the assemblages suggests the presence of ferns showing a strong affinity with species of the extant genus *Lygodium*, a pantropical climbing schizaealean, concentrated especially in Malaysia and Central America (Kramer in Kramer and Green, 1990). These ferns are typical of warm, humid environments. Their fossil analogues may well have grown along riverbanks or in understories of forests (Van Konijnenburg-Van Cittert, 2002). The composition of the Angeac palynological assemblages is consistent with the previous interpretation of the depositional environment at this site as a

swamp, but the absence of marine dinoflagellate cysts apart from the single specimen
identified as *Odontochitina imparilis*, suggests very limited connection to the sea (Néraudeau
et al., 2012). Nevertheless, this is attested by the relative abundance in unit An2 of brackish
ostracods such as *Fabanella boloniensis* (Schudack and Schudack, 2011) and the presence at
different levels (An3–4, An1) of a few benthic foraminifers (*Ammocycloloculina* sp., *Trocholina odukpaniensis*) and echinoid or bryozoan debris (Néraudeau et al., 2012 and
unpublished data).

No palaeoenvironmental variation has previously been noted for the period of
deposition of the sediments comprising units An5–An2. However, the relative composition of
Angeac palynological assemblages changes from An4/5 up to An2, the abundance of
verrucate spores clearly increasing at the expense of *Classopollis*, which diminishes greatly
(Fig. 7). The verrucate spores recovered from unit An2 appear to be generally larger than the
specimens recovered from the older units An3 and An4 (Fig. 8).

From a sedimentological perspective, the succession of green clay (An5), dark clay (An4) and calcareous conglomerate (An3), and then of silty limestone (An2) and fine sand (An1) implies two successive sequences of increasing energy in the depositional environment, which suggests progressive flooding of the swamp by a river nearby, ending with the establishment of a floodplain. This environmental change could explain the increase of the abundance of verrucate spores up-section, ferns being generally more abundant than gymnosperms in a floodplain (Coiffard et al., 2006, 2007).

694

695 **6. Conclusion**

696

Our taxonomic study of the Angeac palynological assemblages provides a rare insight
 into French palynofloras of Berriasian–Valanginian age. They have previously only been

| 699 | documented from deposits in the Jura mountains, the Vocontian Basin, and on Oléron island |
|--|--|
| 700 | (Lantz, 1958; Taugourdeau-Lantz and Donze, 1971; Médus and Mojon, 1991; Kujau et al., |
| 701 | 2013), but the Angeac assemblages seem more continental not only because of the abundance |
| 702 | of verrucate spores but also owing to the scarcity of marine indicators in the preparations. |
| 703 | All of the verrucate spores are attributable to either <i>Concavissimisporites</i> or |
| 704 | Trilobosporites, which in turn are believed to have been produced by ferns showing strong |
| 705 | affinities with extant Lygodium, a fern genus that typically inhabits warm, humid |
| 706 | environments. These ferns could well have grown on the margins of the swamp envisaged by |
| 707 | Néraudeau et al. (2012) and been widely dispersed on the developing floodplain. |
| 708 | Despite the important work of Bolchovitina (1961, 1971), in the light of recent research |
| 709 | on modern representatives of the Lygodiaceae a taxonomic revision of Lower Cretaceous |
| 710 | lygodiaceous spores in the Northern Hemisphere is now needed in which both their biological |
| | |
| 711 | affinities and stratigraphic distribution are taken into account. |
| 711 712 | affinities and stratigraphic distribution are taken into account. |
| | affinities and stratigraphic distribution are taken into account. Acknowledgements |
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| 712713714715 | Acknowledgements We are very grateful to E. Masure for making available the palynological preparations |
| 712 713 714 715 716 | Acknowledgements We are very grateful to E. Masure for making available the palynological preparations for analysis and for her useful insight concerning dinoflagellate identification. We also thank |
| 712 713 714 715 716 717 | Acknowledgements We are very grateful to E. Masure for making available the palynological preparations for analysis and for her useful insight concerning dinoflagellate identification. We also thank L. Londeix, who processed an additional sample from Angeac unit An2 at Bordeaux |
| 712 713 714 715 716 717 718 | Acknowledgements We are very grateful to E. Masure for making available the palynological preparations for analysis and for her useful insight concerning dinoflagellate identification. We also thank L. Londeix, who processed an additional sample from Angeac unit An2 at Bordeaux University. We are grateful to the Audoin Society for access to the quarry, to R. Allain, who |
| 712 713 714 715 716 717 718 719 | Acknowledgements We are very grateful to E. Masure for making available the palynological preparations for analysis and for her useful insight concerning dinoflagellate identification. We also thank L. Londeix, who processed an additional sample from Angeac unit An2 at Bordeaux University. We are grateful to the Audoin Society for access to the quarry, to R. Allain, who took time to show the Angeac excavation site to FP and discuss the latest discoveries there, |

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| 940 | |
| 941 | |
| 942 | Figure captions |
| 943 | |
| 944 | Fig. 1. A. Map showing the geographical and geological location of Angeac, along with other |
| 945 | outcrop localities and boreholes in which "Purbeck" and Wealden-type facies are preserved. |
| 946 | Soubran is mentioned in the introduction (Section 1), but is located about 50 km south-east of |

 $_{947}$ Royan, and is not, therefore, indicated on the map. GPS coordinates of the site: latitude: 45°

| 948 | 37' 59.08" N; longitude: 0° 5' 7.27" W. B . Synthetic stratigraphic section of Angeac |
|-----|---|
| 949 | sedimentological units. The solid circles indicate parts of the section from which the |
| 950 | palynological samples were taken (modified after Néraudeau et al., 2012, fig. 1). |
| 951 | |
| 952 | Fig. 2. Schematic representation of a verrucate spore showing seven out of ten variables |
| 953 | measured to run a PCA. S: maximum diameter; SA, LA: smallest and largest value of the |
| 954 | reflex, acute or obtuse angle formed by the two tangents of the starting and arrival points of |
| 955 | the arc or curve formed by the outline of the interradial region of a spore; miD: minimum |
| 956 | distance from the centre to the edge of the spore (excluding the verrucae); maD: maximum |
| 957 | distance from the centre to the edge of the spore (excluding the vertucae); DV: diameter of |
| 958 | verrucae on one spore; HV: height of verrucae on one spore; ET: exine thickness. |
| 959 | |
| 960 | Fig. 3. Species list of palynomorphs recovered from sedimentological units An2–An4 at |
| 961 | Angeac. |
| 962 | |
| 963 | Fig. 4. Palynomorphs recovered from Angeac sedimentological units An2–An4. |
| 964 | Accompanying data are palynological preparation and slide numbers followed by England |
| | |

Finder coordinates. Authors of taxa not otherwise cited in the text are not listed in the 965

references. Scale bar represents 10 µm. A. Abietineaepollenites sp.; LEM4-1, E23.4. B. 966

Callialasporites dampieri (Balme, 1957) Dev, 1961; LEM4-2, R53.0. C, D. Classopollis 967

torosus (Reissinger, 1950) Couper, 1958; QPR3669-2; V20.4, K47.3. E. Vitreisporites 968

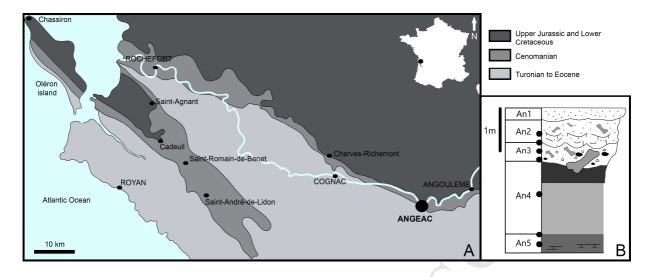
- pallidus (Reissinger) Nilsson, 1958; QPR3669-3, S35.3. F. Podocarpidites ellipticus 969
- Cookson, 1947; LEM4-2, U31.0. G. Aequitriradites verrucosus (Cookson et Dettmann, 1958) 970
- Cookson et Dettmann, 1961; QPR3669-3, W38.1. H. Cyathidites australis Couper, 1953; 971
- LEM4-1, G52.2. I. Microreticulatisporites sp. cf. M. diatretus Norris, 1969; LEM4-1, X45.0. 972

| 973 | J. Gleicheniidites apilobatus Brenner, 1963; QPR3669-4, Q20.2. K. Densoisporites |
|-----|---|
| 974 | microrugulatus Brenner, 1963; LEM4-1, H30.4. L. cf. Foraminisporis wonthaggiensis |
| 975 | (Cookson et Dettmann, 1958) Dettmann, 1963; LEM4-1, E30.2. M. Cicatricosisporites hallei |
| 976 | Delcourt et Sprumont, 1955 sensu Burger, 1966; LEM3-2, G39.4. N. Ovoidites spriggi |
| 977 | (Cookson et Dettmann, 1959) Zippi, 1998; LEM3-2, L34.4. O. Retitriletes sp.; LEM3-2, |
| 978 | D33.2. P. Pilosisporites trichopapillosus (Thiergart, 1949) Delcourt et Sprumont, 1955; |
| 979 | QPR3669-3, V31.2. |
| 980 | |
| 981 | Fig. 5. Species of <i>Concavissimisporites</i> recovered from Angeac sedimentological units An2– |
| 982 | An4. Accompanying data are palynological preparation and slide numbers followed by |
| 983 | England Finder coordinates. Authors of taxa not otherwise cited in the text are not listed in the |
| 984 | references. Scale bar represents 10 µm. A, B. Concavissimisporites apiverrucatus (Couper, |
| 985 | 1958) Döring, 1965; LEM4-1, D33.1; QPR3669-2, E24.1. C, D. Concavissimisporites |
| 986 | exquisitus Singh, 1971; QPR3669-3, O27.4, G36.3. E. Concavissimisporites sp. cf. C. |
| 987 | exquisitus Singh, 1971; LEM2-1, R40.2. F-H. Concavissimisporites ferniensis (Pocock, |
| 988 | 1970) Fensome, 1987; QPR3669-2, M53.4, N20.4; LEM2-3, C33.0. I. Concavissimisporites |
| 989 | robustus Dörhöfer, 1977; LEM2-3, R34.0. J, K, M. Concavissimisporites montuosus (Döring, |
| 990 | 1964) Fensome, 1987; QPR3669-2, W31.0; LEM1-1, Q49.0; QPR3669-2, T28.1. L, N. |
| 991 | Concavissimisporites sp. cf. C. crassatus (Delcourt et Sprumont, 1955) Delcourt, Dettmann et |
| 992 | Hughes, 1963; LEM1-1, Q35.4, B28.4. |
| 993 | |

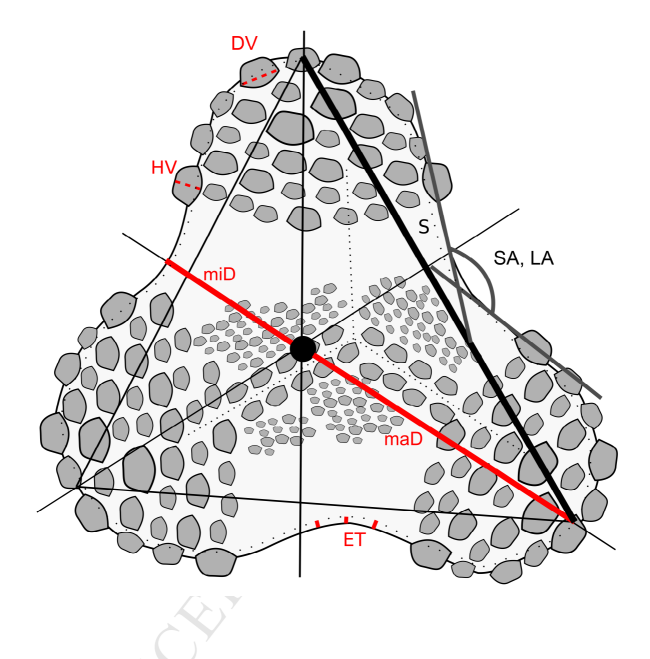
Fig. 6. Species of *Concavissimisporites* and *Trilobosporites* recovered from Angeac
sedimentological units An2–An4. Accompanying data are palynological preparation and slide
numbers followed by England Finder coordinates. Authors of taxa not otherwise cited in the
text are not listed in the references. Scale bar represents 10 µm. A, B. *Concavissimisporites*

| 998 | verrucosus (Delcourt et Sprumont, 1955) Delcourt, Dettmann et Hughes, 1963; QPR3669-3, |
|------|--|
| 999 | E20.1, V23.0. C. Concavissimisporites sp. cf. C. verrucosus (Delcourt et Sprumont, 1955) |
| 1000 | Delcourt, Dettmann et Hughes, 1963; QPR3669-3, D20.4. D. Concavissimisporites uralensis |
| 1001 | (Bolkhovitina, 1961) Fensome, 1987; LEM1-1, Y42.3. E-G. Trilobosporites aornatus |
| 1002 | Döring, 1965; LEM2-1, S32.1; LEM3-2, B32.0; LEM1-1, K28.3. H. Trilobosporites |
| 1003 | canadensis Pocock, 1962; LEM1-1, A28.2. I. Trilobosporites aequiverrucosus Dörhöfer, |
| 1004 | 1977; QPR3669-3, S30.0. |
| 1005 | |
| 1006 | Fig. 7. Palynological composition of Lower Cretaceous sedimentological units An2–An4 at |
| 1007 | Angeac: 250 specimens were counted for each sample except for that from the boundary |
| 1008 | between An4 and An5, which yielded only 94 specimens. |
| 1009 | |
| 1010 | Fig. 8. PCA1 plot of PC1 vs. PC2, with 76.2 % of the total variance extracted, showing the |
| 1011 | distribution of specimens according to their occurrence in the sedimentological units. For |
| 1012 | explanation of abbreviations, see caption to Fig. 2. |
| 1013 | |
| 1014 | Fig. 9. PCA2 plot of PC1 vs. PC2, with 76.2 % of the total variance extracted, showing |
| 1015 | clusters corresponding to species of Concavissimisporites and Trilobosporites. For |
| 1016 | explanation of abbreviations, see caption to Fig. 2. |
| 1017 | |
| 1018 | Fig. 10. Megaspores from Angeac. A–D. Erlansonisporites sp. from unit An4, mesofossil |
| 1019 | preparation MFP512, SEM stub DJB2014/18, specimen 23, IGR-PAL-5780. A. Whole |
| 1020 | specimen. B. Close-up of part of triradiate flange and adjacent reticulate wall. C. Detail of |
| 1021 | part of reticulate sculpture showing the structure of the outermost part of the protective wall: |
| 1022 | the surface consists of a network of sporopollenin threads, beneath which is an irregular, |
| | |

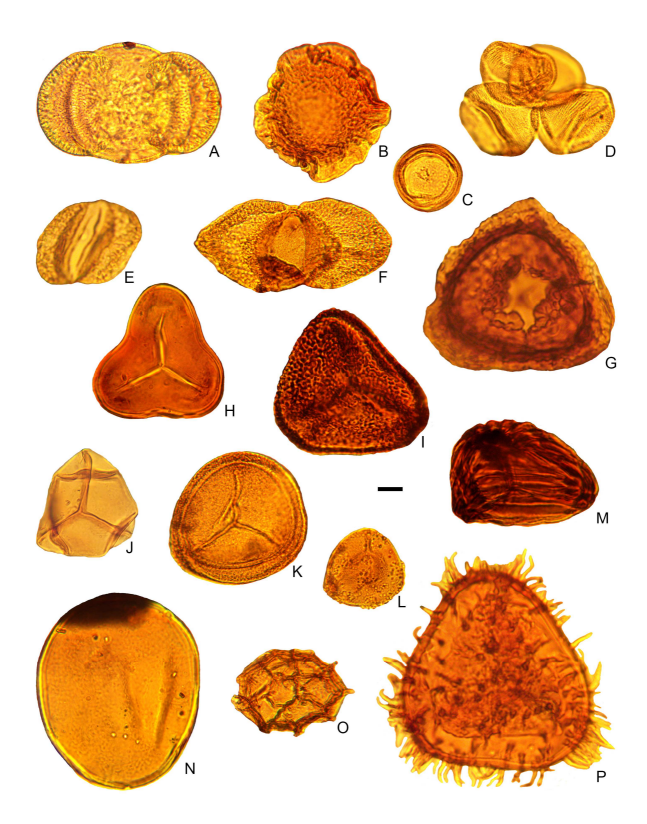
perforated layer. D. The same at higher magnification. E, F. Erlansonisporites sp., cf. 1023 Erlansonisporites sp. in Batten (1975, pl. 13, fig. 6). Mesofossil preparation MFP422, from 1024 the boundary between An3 and An4, SEM stub DJB2014/20, specimen 1, IGR-PAL-5781. E. 1025 whole specimen. **F**. Close-up of reticulate sculpture. Scale bars: A, E: 200 µm; B: 50 µm; C: 1026 10 μm; D: 5 μm; F: 20 μm. 1027 1028 Fig. 11. Megaspores from Angeac, all from mesofossil preparation MFP422, from the 1029 boundary between An3 and An4, SEM stub DJB2014/20, specimens 4 (IGR-PAL-5784), 2 1030 (IGR-PAL-5782) and 3 (IGR-PAL-5783), respectively. A, B. Erlansonisporites sp., cf. 1031 Erlansonisporites sp. in Batten (1975, pl. 13, fig. 6). A. Whole specimen in lateral view. B. 1032 Close-up of a small part of the reticulate sculpture showing microspores lodged within the 1033 muri of the reticulum. C, D. Specimen tentatively identified as *Minerisporites* sp., an atypical 1034 representative of this genus (see discussion in text). C. Whole specimen in polar view. D. 1035 Close-up of perforated surface. E, F. Another example. E. Specimen in lateral view. F. Close-1036 up of undulating distal surface just below equatorial flange (zona). Scale bars: A, C, E: 200 1037 μm; B, D, F: 20 μm. 1038

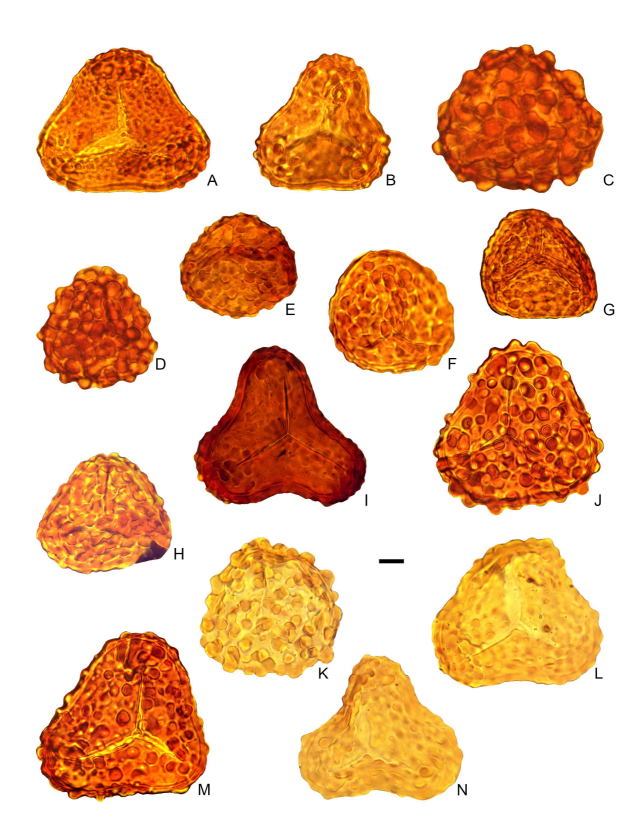


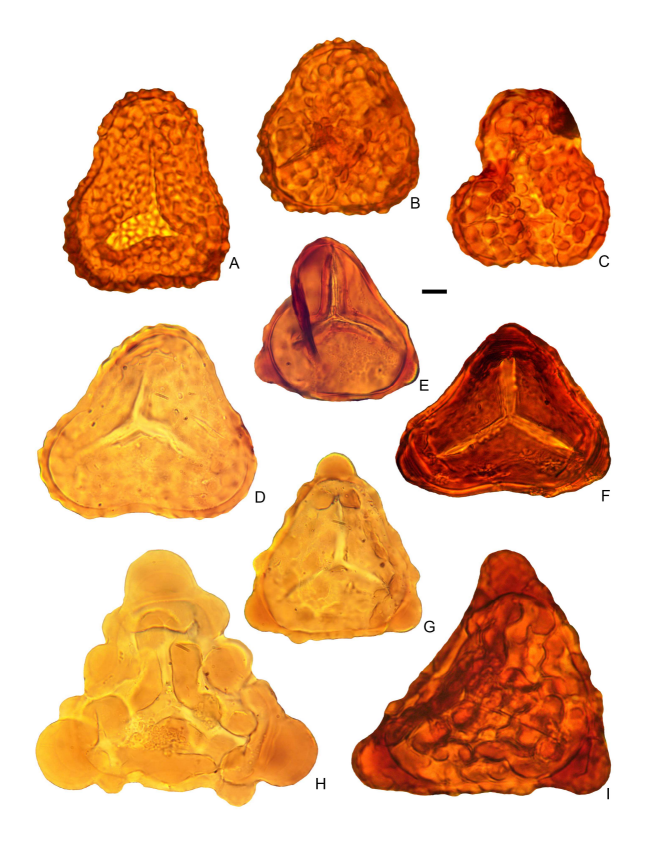
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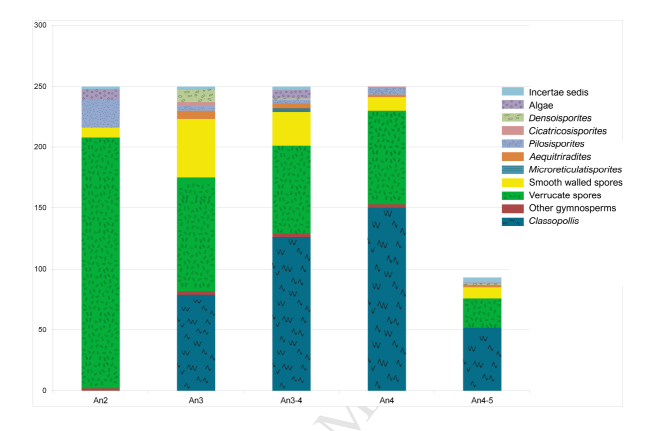


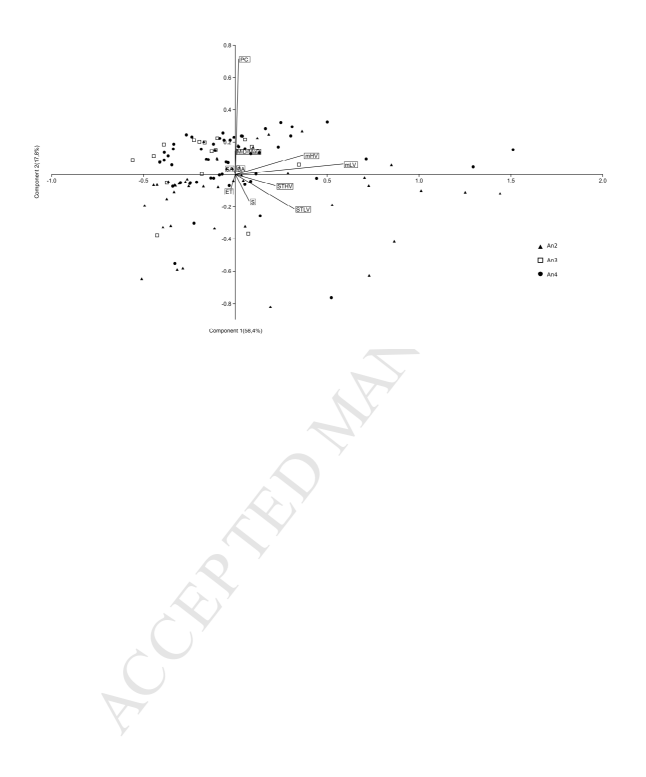
| | An2 | An3 | An3/4 | An4 | An4/5 | An5 |
|---|-----|-----|-------|-----|-------|-----|
| Abietineaepollenites sp. | | o | | | | |
| Aequitriradites verrucosus | | o | o | 0 | o | |
| Araucariacites australis | o | | | | | |
| Callialasporites dampieri | | | o | | | |
| Cerebropollenites mesozoicus | | | | 0 | | |
| <i>Cicatricosisporites hallei sensu</i> Burger, 1966 | | ٥ | | | | |
| Cicatricosisporites sp. cf. C. sternum sensu Burger, 1966 | | o | | o | | |
| Classopollis torosus | | 0 | 0 | 0 | 0 | o |
| Concavissimisporites apiverrucatus | 0 | | 0 | 0 | • | |
| Concavissimisporites exquisitus | o | o | o | 0 | o | |
| Concavissimisporites ferniensis | o | o | o | o | | |
| Concavissimisporites montuosus | o | ٥ | 0 | 0 | o | |
| Concavissimisporites robustus | o | 0 | | | | |
| Concavissimisporites sp. cf. C. crassatus | 0 | | | | | |
| Concavissimisporites sp. cf. C. exquisitus | 0 | ٥ | | o | | |
| Concavissimisporites sp. cf. C. montuosus | 0 | o | | | | |
| Concavissimisporites sp. cf. C. robustus | 0 | | | | | |
| Concavissimisporites uralensis | o | | | o | | |
| Concavissimisporites verrucosus | | 0 | | o | 0 | |
| Cyathidites australis | | | o | 0 | | |
| Cyathidites sp. | 0 | ٥ | 0 | o | 0 | |
| Deltoidospora sp. | o | 0 | 0 | o | 0 | |
| Densoisporites microrugulatus | | | 0 | 0 | o | |
| cf. Foraminisporis wonthaggiensis | | | 0 | | | |
| Gleicheniidites apilobatus | o | | | o | o | |
| Microreticulatisporites sp. cf. M. diatretus | | | o | | | |
| Ovoidites spriggi | | o | | | | |
| Ovoidites sp. | 0 | o | | | o | |
| Pilosisporites trichopapillosus | o | o | o | 0 | o | |
| Podocarpidites ellipticus | | | 0 | | | |
| Retitriletes austroclavatidites | | o | | | o | |
| Retitriletes sp. | | | | o | | |
| Trilobosporites aequiverrucosus | | | | o | | |
| Trilobosporites aornatus | o | o | | o | 0 | |
| Trilobosporites canadensis | 0 | o | o | o | o | |
| Trilobosporites sp. cf. T. aornatus | o | | | | | |
| Vitreisporites pallidus | | | | 0 | | |

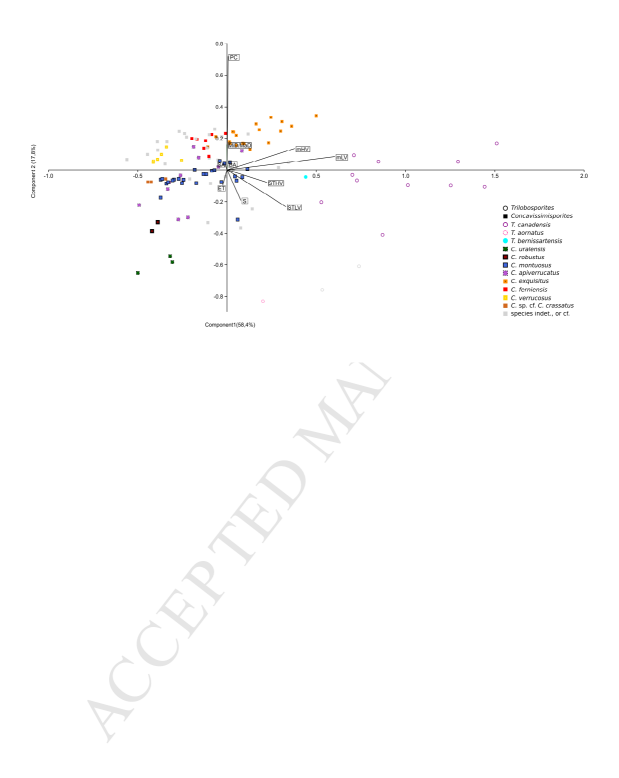


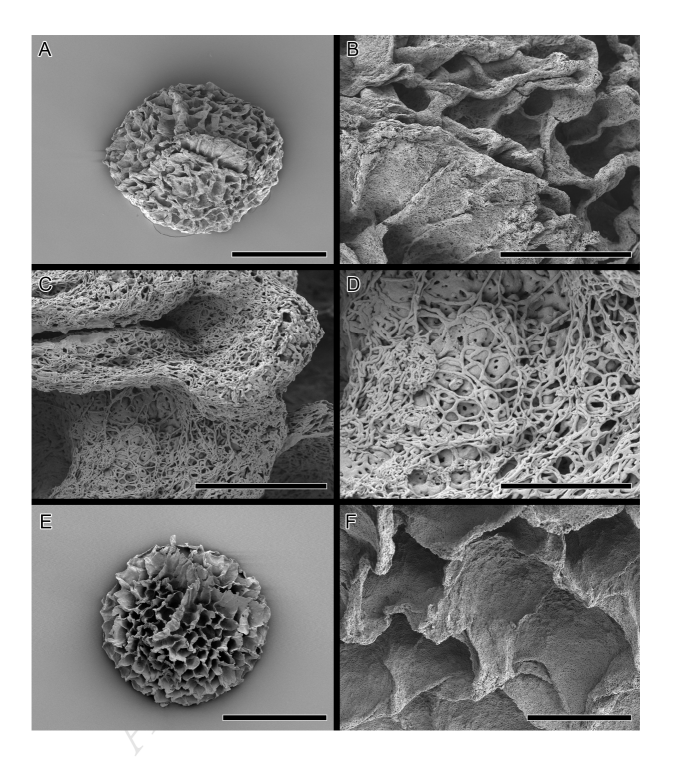


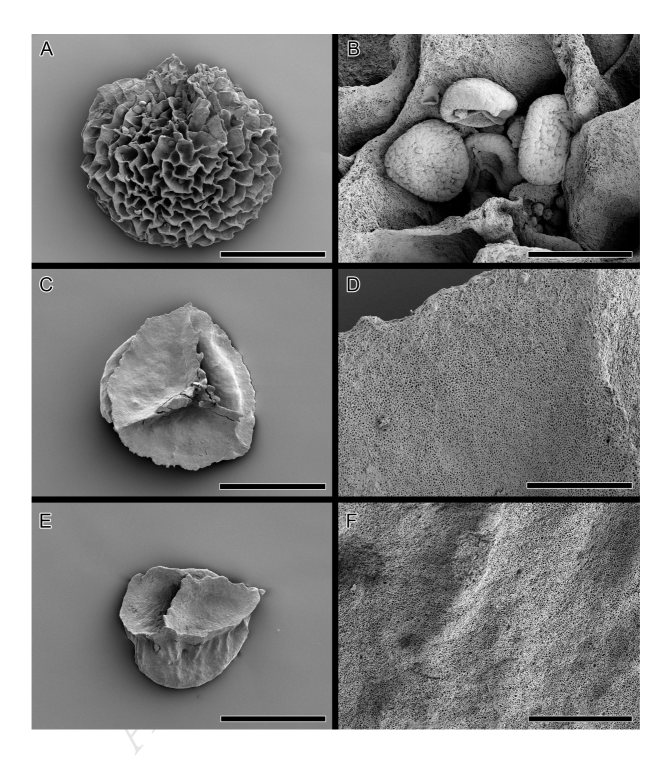












Highlights

Revision of the palynology of Lower Cretaceous strata at Angeac, western France

Deposition is considered to have occurred during the Berriasian-Valanginian

Eight species of verrucate spores were recovered from the non-marine deposits

All are referred to either Concavissimisporites or Trilobosporites

The taxonomy and biological relevance of the verrucate spores are discussed