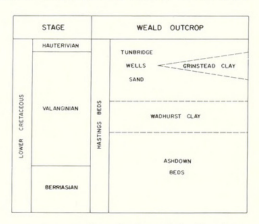
# PALYNOLOGY OF EARLY CRETACEOUS SOIL BEDS AND ASSOCIATED STRATA

by D. J. BATTEN

ABSTRACT. A distinctive palynologic assemblage is associated with *Equisetites* soil beds and fragment partings in the Wadhurst Clay of south-east England. Some of the components are probably the products of a local flora which grew in or near the *Equisetites* communities. Palynologic and palaeobotanic data suggest that the communities were isolated from the 'homogeneous Wealden delta flora', probably offshore from the delta complex. New taxa have been recorded from the assemblages. *Converrucosisporites venitus*, *Regresporites lophus*, *Retitriletes* sp., *Trilobosporites ivanovae*, and *Admolia amphidoxa* are described.

Equisetites ('horsetail') soil beds are common in the Wadhurst Clay (Lower Cretaceous; text-fig. 1) of south-east England. They constitute combinations of roots, rhizomes, and stems of the plant preserved in situ, generally in a matrix of interlaminated clays and fine-medium silts. They are the only truly autochthonous plant beds of botanic value in the Wealden. Some of the Equisetites fragment partings which occur above and occasionally within the soil beds are probably para-autochthonous. The majority of the rootlet beds in the succession have, however,

been truncated and there is nothing to indicate to which plants the roots were attached (although in the Wadhurst Clay they may frequently belong to Equisetites). It has proved difficult to determine both the environments of deposition in which Equisetites thrived and the factors which controlled its distribution. The Hastings Beds succession accumulated in an essentially non-marine basin (the Anglo-Paris Basin) with the result that similar aquatic faunas and floras lived both on the delta complex and offshore (Allen 1959, 1967). Although the properties of many of the sediment types are probably characteristic for special areas of formation, these areas are difficult to delimit. Four possible interpretations of the standard Hastings Beds megacyclothem, in which Equisetites soil beds play an important part, have been proposed by Allen (summarized 1967, text-fig. 1). The distribution of



TEXT-FIG. 1. Correlation of the Hastings Beds in south-east England (after Harland *et al.* 1967, and latest published information). Lithological boundaries are indicated by broken lines. Thickness of Wadhurst Clay varies from 30 to 70 m.

palynomorphs in samples from horizons within and adjacent to some of the Wadhurst Clay soil beds is considered here both in the light of these interpretations and with regard to the distribution of local flora.

### MATERIAL

The samples came from the Wadhurst Clay in the Cuckfield No. 1 Borehole, Sidnye Farm, Sussex (TQ 2962 2729, first redrilling, and TQ 2961 2731, second redrilling); Warlingham Borehole, Surrey

(TQ 3476 5719); Railway Brickyard, Sharpthorne, Sussex (TQ 3740 2729); and Freshfield Lane Brickworks, Danehill, Sussex (TQ 3813 2659). Most of the sampling horizons are figured in Batten (1968, text-figs. 1 and 2). Approximately 15 g of each sample from not more than 2 cm stratigraphic thickness were subjected to a standard preparation procedure (see Batten 1973) employing 50% hydrochloric acid, HF, nitric acid (< 30 minutes) or Schulze solution (< 10 minutes), ammonium hydroxide (5 minutes), and zinc bromide (sp. gr. c. 2·2). Strew slides were made with Clearcol; Euparal sealed the cover-slips to the slides.

# WADHURST CLAY EQUISETITES

Rootlets preserved as casts and infilled rhizomes are usually all that remain of Wadhurst *Equisetites* communities. Infilled stems in position of growth occur less often and rhizomes preserved as carbonaceous compressions are rare. The species preserved in the soil beds examined is *Equisetites lyellii* (Mantell) (1833). Since the species was erected, it has been consistently referred to as *lyelli*. Mantell's spelling is correct and is here reverted to (see Lanjouw *et al.* 1966, Recommendation 73C). *E. lyellii* is the only common plant megafossil in the Wadhurst Clay. *E. burchardti* Dunker and *E. yokoyamae* Seward, both tuber-bearing forms, were not encountered; they tend to occur in the more arenaceous facies of the succession (cf. Allen 1962).

The scarcity of other plants in the Wadhurst Clay suggests: (1) that most of the plant material derived from the lowland flora bordering the basin of deposition was deposited or comminuted before it reached the Wadhurst zone of deposition, (2) that the horsetail communities were isolated in some way from the bulk of the lowland flora. In addition, the preservation of *E. lyellii* may have been enhanced by two factors: (1) it was not dependent on chance deposition in an aqueous environment conducive to preservation because it probably grew in the shallow waters of the Wadhurst Clay, (2) although there is no direct evidence, the plants may have accumulated silica in their tissues as do species of extant *Equisetum* (e.g. *E. hyemale*; Lovering and Engel 1967).

Stem fragments of *E. lyellii* may be scattered through the clays and silts above or within the soil beds or concentrated in thin bands separated by more or less fragment-free horizons. The concentrations represent periods when deposition of waterlogged plant debris was more rapid than that of sediment deposition, and are probably the product of destruction of *E. lyellii* communities by water disturbance (Allen 1941; Batten 1968). The preservation state varies, but the accumulations which are well preserved may be nearly autochthonous. The absence of any large accumulations of *Equisetites* suggests that the communities were short lived (Allen 1941, 1960) and that *E. lyellii* was a pioneer coloniser (Hughes and Moody-Stuart 1967a); their existence was probably terminated by deepening water. Had the communities existed for many more than a few years, thin coal seams like those in the Yorkshire Jurassic (Harris 1953) might have been formed.

### SOIL BED AND FRAGMENT PARTING PALYNOLOGY

Soil beds and fragment partings often yield a distinctive palynologic assemblage. This assemblage is here designated as the SF (soil bed/fragment parting) type. During a routine examination of Wadhurst preparations most were assigned (at

the specific level) to assemblage-types 8 and/or 16 (Table 1 and Batten 1973). Important characters are: (1) the miospore *Pilasporites allenii* (described from a Wadhurst Clay soil bed preparation (CUC 792) and attributed by Batten 1968 to *Equisetites*) is abundant (V or F on Table 1), (2) there is usually a large amount of plant debris comprising brown 'wood' (probably mostly degraded *Equisetites* material) and *Equisetites* cuticle, (3) the general state of preservation of the miospores is usually fair to good.

Certain miospore taxa, viz. Retitriletes sp., Regresporites lophus gen. et sp. nov., Ischyosporites arkellii (Pocock 1970) comb. nov., and Admolia amphidoxa gen. et sp. nov. appear to be significantly associated. I. arkellii and A. amphidoxa occur more often and in greater abundances in SF assemblages than elsewhere, and Retitriletes sp. and Regresporites lophus have only been recorded from the SF type. Trilobosporites ivanovae sp. nov. and Patellasporites spp. occur more frequently associated with Equisetites than elsewhere in the Wadhurst Clay (although the former is more numerous and better preserved in the described Grinstead Clay sample). The frequency of occurrence of Pilosisporites spp., Converrucosisporites venitus sp. nov., Staplinisporites spp., Classopollis spp., and most of the other taxa recorded from soil beds and associated strata differs little from their frequency in the rest of the Wadhurst. However, megaspores have not been recovered, and the following are notably less common: Inapertisporites sp., Cerebropollenites mesozoicus (Couper 1958) Nilsson 1958, Couperisporites spp., Vitreisporites pallidus (Reissinger 1938) Nilsson 1958, Ceratosporites spp., and Celyphus rallus Batten 1973. Some soil bed and fragment parting samples do not yield an SF kind of assemblage (Table 1). Instead, the assemblages show little diversity and poor preservation and have (see below) a typical 'Wadhurst aspect' (assemblage-types 1, 4, 7, 9, 10, and 12 in Batten 1973).

### NUMERICAL COMPARISON OF ASSEMBLAGES

The relationships between forty-two selected assemblages recovered from soil beds and adjacent strata were determined by using both cluster analysis and nonmetric multidimensional scaling (see Kruskal 1964a, b). For cluster analysis both weighted and unweighted pair-group linkage methods were used (Sokal and Michener 1958; Sokal and Sneath 1963). The cluster program operated on matrices of coefficients of association derived by comparing data from counts of both 500 (Table 1) and 200 miospores, with and without other data (on preservation, abundance of wood, cuticle, etc.). The coefficients of association used were those of proportional similarity (Cos  $\theta$ ) of Imbrie and Purdy (1962), the Jaccard, and Sokal and Michener (see Cheetham and Hazel 1969). The multivariate data were converted to binary form to use the Sokal and Jaccard coefficients. The nonmetric multidimensional scaling program employed Cos  $\theta$  converted to a distance coefficient. Both procedures yielded similar results but those from the cluster analysis were more clear cut than the multidimensional scaling. Assemblages with an SF aspect formed one grouping and those identified as 'typical Wadhurst type' formed another. 'Borderline' assemblages which show only some of the characters of the SF type (i.e. those from samples CUC 792/1, CUC 792/4, CUC 792/6, DJB 329, and DJB 368;

Table 1) tended not to be well linked with either group. The records of Regresporites lophus, Ischyosporites arkellii, Admolia amphidoxa, Pilasporites allenii, Classopollis spp., Ceratosporites spp., and Gleicheniidites spp. were important in determining the grouping of the assemblages. The state of preservation and the abundance of brown wood and cuticle fragments were also important when all the data, not just the miospore content, were taken into account.

### PALAEOECOLOGY

Both onshore and offshore habitats have been postulated for the Equisetites communities (Allen 1959, 1967, text-fig. 1). On the delta top they could have thrived on levées and in backswamps and shallow ponds (Allen 1949, 1959, megacyclothem interpretations 1 and 2); offshore they could have become established in a band along much of the length of the delta shore-face and in other areas of shallow water (shoals) (Allen 1959, 1967, interpretations 3 and 4). Most Wadhurst Clay palynologic assemblages reflect the homogeneous delta flora in their composition. They are composed mainly of pteropsid and coniferopsid elements. Fern spores, especially the families Schizaeaceae and Gleicheniaceae, are prominent and, in the former, diverse. The miospores are, however, generally in a poor state of preservation, and when compared with the whole Wealden succession, show below average diversity. The general aspect of the assemblages, and the scarcity of both megaspores (Batten 1969) and determinable megascopic plant remains, apart from those of Equisetites, in the Wadhurst suggest that deposition took place far from source. The poor preservation of the palynomorphs can be accounted for by prolonged aerobic decay and reworking. All but the more resistant spores and plant fragments (fusain) would have been destroyed by the alternations of erosion and deposition on the delta complex.

The SF kind of assemblage has, however, a different aspect. The homogeneous delta flora forms only a background component. The better preservation suggests that some of the elements were not transported over long distances but were deposited in an environment where water transport was restricted to some extent, and where the duration of their exposure to aerobic decay depended on the rate at which they

TABLE 1. Plant megafossil and miospore content of 42 selected samples from Equisetites soil beds and associated strata. Sample numbers on left-hand side of table: prefix DJB = field sample, CUC = core sample from Cuckfield No. 1 Borehole, WM = core sample from Warlingham Borehole. On extreme right, record of assemblage-type assignments (see Batten 1973) noted during routine examination of Wadhurst Clay preparations (before counts of 500 miospores), and of SF kinds of assemblages (B = 'borderline' type).

Miospore taxa: P = present (< 3% after count of 500), C = common (3-14·5%), V = very common (15-29·5%), F = frequent (30% or more). Taxa grouped within suprageneric categories except where indicated by solid line. Elements recorded subsequent to completion of count not included. Rootlets, rhizomes, and stems, X if recorded. Megascopic fragments of *Equisetites*: C = common, F = frequent. 1. *Cyclogranisporites* Pot. and Kr./*Granulatisporites* Ibr. emend. Pot. and Kr. 2. *Corrugatisporites* Th. and Pfl. ex Wey. and Greif./*Rugulatisporites* Th. and Pfl. 3. *Foveosporites* Balme/*Foveotriletes* v.d. Hammen ex Pot./*Sestrosporites* Dett. 4. *Lycopodiumsporites* Thierg. ex Del. and Spr./*Reticulisporites* Pot. and Kr. in Wey. and Krieg. 5. *Taurocusporites* Stover emend. Play. and Dett./*Polycingulatisporites* Simonc. and Kedv. emend. Play. and Dett.

8r0 8r0	860	Bro	Bro Dag	Bra	Bra	Bro	Bro	Bro	Bra	B G G	870	Bro	Bro	Bro	WW	cuc	cuc	CUC	CUC	cuc	cuc	CUC	cuc	cuc	cuc	cuc	cuc	CUC 788	CUC 78	cuc	cuc	S A A
366 367	365	364	360	359	358	357	356	330	346	347	349	327	324	320	1774/6	898	866/3	866	865/3	794/6	794/3	793	792/6	792/4	792/2	792/1	792	1	8/10-11A	788/6-9	788	SAMPLES TAXA
000			+	В		+		Р.		7	-		D		O			7	)	ס	-	0	$\Box$		+			τ		0	D	Auritulinosporites NILSSON LAEVIGAT
									-	U	P		7	O	О					ъ	-		$\Box$	D				+		1	D	Concavisporites jurienensis BALME Stereisporites PFL. in TH. and PFL.
0 0 0 0 0 0		P P	ס ס		D	ם ס ד	T .		D	7	סכ		+		Ъ		0	n	-				P		D T	ס		0	PP	יס	_	? Stereisporites sp. (undescribed) Undulatisporites PFL. in TH. and PFL.
< < <													< <	< <					0	F	T) ·	< T	1 <	< 0							2	TOTAL SMOOTH WALLED TRILETES
ס																										ס						Aconthotriletes NAUM. emend. POT. and KR. APICULA
ס	$\vdash$	-	סס	-	TO	+	Н	D	+	+	+	$\vdash$	+	+		Н	D	+	+	Н		סס	סי	-	O	P	+	+	ס	О	+	Apiculatisparis POT. and KR. Baculatisparites TH. and PFL.
	$\Box$	0	ס		Н				+				+		ъ	ס	ס		+			0 0	,			Н		+	$\vdash$	-	D	Ceratosporites COOK. and DETT.
0		ס	_	ס		τ			1				ר ס	O			D	Ţ		ס	1	0		ס		P	7	J		ס	4	Concavissimisporites DEL. and SPR. emend. DEL. et al.
0	$\vdash$	P	O		Φ.	D	סו		+	+	P	-	D	70	P	P	D	17	7	P	-	0	H	Ъ	+	Н	+	+	P	P P	+	Converrucosisporites POT. and KR. C. venitus SP. NOV.
		יס	ס	P			סי	О			TO		Ť		ס	-		7		-			סו			TO	ר ס	ס ד	-	_	ס	Cyclogranisporites / Granulatisporites 1.
++	Н	-	7		-	D		О	$\perp$	-			4		T	ס	_	1				0 0				ס				-	_	Foraminisporis KRUTZSCH Osmundacidites COUPER
TO	H	D	10				D		+	٦	0	1	D	+	-	D	_	+	+	P	_		+	ס	+	ס	_	0	P	D D	+	Pilosisporites DEL. and SPR.
		-	סס	ס		$\perp$							7	O	ס		0	U			ס ־	0						τ			ס	Verrucosisporites IBR emend. SMITH and B'WORTH
000	77		10	T	77 /	200		T)		7	100	70	D 3	D T			0	D 7	1 77		0				70	_	P	100	10	77		Verrucosisporites sp. (undescribed) Cicatricosisporites POT and GELL. MURORNA
110	1	1	1	0	0	1	1	0	1	1	1	1	0	0	-		1	1	1		1	1	1	-	1	1		0	P		1	Corrugatisporites/Rugulatisporites 2.
ס		ס				U								ס			ס	٦	)				Р		TO							Foveosporites/Foveotriletes/ Sestrosporites 3.
0 0		D .	סס	-		+	ס	-	ס	٦	1		+	-		P	О П	0	-		D -	0 0	_	D	+	-	1	D	P	ס	-	Klukisporites COUPER Lycopodiumsporites/Reticulisporites 4.
1		_	ס						1				+		0	-	9			P	9	-	ס	0		H		1		-	D	Lycopodiumsporites austroclavatidites (COOK) POCOCK
	П								1	1			1		ס	-	_	1		ס			ס					1				Microreticulatisporites KNOX emend. POT. and KR.
0	Н	ס	+	$\vdash$		ס ד	0		+	+	TO		+	+	PP	-	+	+	+	Н	י ס	0	ס	O	+	P	ס	+	P		+	Reticulisporites semireticulatus (BURGER) NORRIS Retitriletes v.d. HAMMEN ex PIERCE
					О	τ				$\pm$			$\pm$									$^{\dagger}$	$\Box$	-	ס	$\Box$	О	τ	,			Retitriletes sp.
ססס				C				ס	1	-		ס	1				_								ס ס	-	07		ס			Regresporites Iophus GEN. and SP. NOV.
0 0 0	-	ס	10	0	יס	0	О	-	+	+	ס פ	О	+	+	D		0	U	+	O	U	0 0	סו	O	+	P	-17	0	0	η.	D	Staplinisparites POCOCK/Coronatispora DETT.  Taurocusporites/Polycingulatisporites 5.
P					ס									+	Р								D			$\Box$	ס					Tripartina MALJ. ex POT.
77 -	ס	-	ס		0	7		ס	-		ס		1			_	_		-		ם י			P				-	-	Φ.	O	Ischyosporites BALME AURICULA
ססס		D D	(	Р		ס ס	H		י ס ס	0		P	+	+	Н	U	0	0 0	ס כ	0	-	0 0	סו	P	2 0	0	O	) (		P	+	I arkellii (POCOCK) comb. nov.  Malonisporites COUPER emend DETT.
		ם -	U	ס	_	στ			_	U		_	ס		ס		1			ס	-	0		ס								Trilobosporites PANT ex POT.
D G C		77	סס	7	D	200	P	-	D	DIT	10		2	00			0	0 7	77		0	7	D	70 0	D T	7	ד ס					T. ivanovae SP. NOV. Gleicheniidites ROSS ex DEL. & SPR. emend. DETT. TRICRASSA
, 000	+		0 0	U	0 0	, .		+	0	0 0			,,,	, ()		0			0	()	()	0 0	10	U	0 0	1	0 1		P		D	Gleicheniidites apilobalus BRENNER
													1							ס												Cingutriletes PIERCE emend. DETT CINGULA
++-	-	D	+	Н	-	+		+	+	+		H	+	ס	H	Н	+	+	ס	H	+	+	Н	+	+	H	+	+			+	Contignisporites DETT.  Densoisporites WEY, and KRIEG, emend, DETT.
		0	ס			O	$\Box$		+			ס	+	+	0	ס	ס			D		TO			T	Н	7	σ	,		1	Potellosporites GROOT and GROOT emend. KEMP. PATINA
סס	7	-	D		_	0		Р.	-		-	-	. 7	0 70	-	-	70	-						_	-	-						Patellasporites sp. (undescribed)
0 0	-	О	-	Н	+	ס ד	-	U	U	+	-	U	0 0	0 0	U	U	U	+	ס נ	-	ס-		ס	U	-	ס		0 0	ס	U	0	INDET. SCULPTURED TRILETES  Aequitrirodites DEL.and SPR. emend. COOK.and DETT. HILATE
						τ			1					I				İ							1							Cooksonites POCOCK emend DETT.
-		-	70		-				+	+			+	+			+	+	-		+	-		-	-	$\vdash$	D	+			-	Couperisporites complexus (COUPER) POCOCK  Couperisporites sp. (undescribed)
	H	-	U	Н	+	+	$\Box$		+	+	$\vdash$	Н	+	+			+	+			+	+	$\forall$	+	+	$\Box$	0	+			+	Rouseisporites POCOCK
									1				1					1			1		$\Box$				1				D	Marattisporites COUPER MONOLETE
ס ס	P	0 -	DC	T	D.	DT	P	77 -	דים	TO		ס	2 7	00	T		P -	0 7	)	0	D	0	0	P	DO	T	T) T	7 -	קו	0	0	Peromonolites ERDT. ex COUPER Aroucariacites COOK. ex COUPER ALETE
		-	ס			στ	T		7	nc			٦	ס ס	T	П				ס	(	0	T		T	D	ס	$\top$	TO			Inopertisporites vd. HAMMEN ex ROUSE
000	C	0	20	C	0	2	0	C	00	0	0	0	2	< 0	<	0	0	ד ס	0	0	< (	0	0	0	0	0	C	T	ס	0		Inaperturopollenites PFL.ex TH. and PFL. emend. POT.
) TI <		00	1	17	110	, (	U	11 4	+	110	()	<	0	-	0	0	+	1	1	0	U	0 0	0	< '	1	<	-1	1 11	n	0	<	Pilasporites allenii BATTEN Reticulatasporites IBR. emend. POT. and KR.
																												-			D	Toxodiaceaepollenites KR.
DE	T	T) -	n -	P		D T		ס ד			P		D		P				-		ם -		ס	1	-			T		-		Cerebropollenites mesozoicus (COUPER) NILSSON SACCITI
999														C < F													70					Tsugaepollenites POT. and VEN. emend. POT. BISACCATES excluding Vitreisporites
ס							ס								ס			τ	ס		ס				TO					ס		Vitreisporites LESCHIK emend. JANSONIUS
סס	$\overline{}$	_	U		$\rightarrow$	ס ד								ס ס	О	U	D	τ	ס	D	_	-	П	7	ס ס	-	1	T		Р.	O	Eucommidites ERDT. emend. HUGHES PLICATE
000	-	P	ס ס	D		ם סד	P			0 0		P .		< <	C	<	0	דוכ	סו	P		ס ס	P	17	ם ס	PC	דס	ם כ		P C	D	Cycadopites WODEH. ex WILS. and WEBST.  Classopollis PFL. emend. REYRE. POROSE
			70				ס									U				ס					ľ		1	Ĺ			1	Classopollis sp. (undescribed)
ס ס כ	0	ם	ס ס		-	ס ד		ם י	ט	T	0	יס-	OC	סכ	P		ם ס	דס	P		ם ס	סוס	ס	ס	-	ס	-	+	ס	Р Р	D D	Exesipollerites BALME
					-	ס									1		0	-	1		9	+	-	+	+	H	+	-		D.	-	Perinopollenites COUPER Spheripollenites COUPER INCERTAE SED
						U										ס				ס			ס							-	D	S. subgranulatus COUPER
		U	C	P	0	D D	H	0 -	_	U		-	O	+			T	τ	1	Р	O	P	P	ד ס	00		00		P		-	Admolia amphidoxa GEN. and SP. NOV.
ס ס ס	ם		+	0					9				-		ס	ס	ס	+	-	U	+	+	0	+	+	Р	0	1	P	U	+	Schizosporis COOK. and DETT. Celyphus rallus BATTEN
ס ס	0			12.2	)	<×					×			<×		×							×			×			×			ROOTLETS
×	יס		<×												120	VI	×			X	XX	IV	14		11	1×	$\times$	1	100	VI	×	BUITOMES (AMS SERVE)
(×		>	< ×		>	××	-	p -	D -	0	×		× >	<×	^	<u> </u>		n -	72			+						1				RHIZOMES (AND STEMS) MEGASCOPIC FRAGMENTS (Con F.)
0 0 0 0 0 ××	71 (	>	××		7	< ×				n C		F X	× >	<×	î		-		FX					(	) < ×	00 ;	××	×	TI ×			MEGASCOPIC FRAGMENTS (C or F) SF TYPE
(X (X	77 (X	0	××	× 00	F X	< ×		× >	× >	< 00		TI X		4 0		9	>	<×					8	B >	< ×	00 ;		×	TI ×		-	MEGASCOPIC FRAGMENTS (C or F)

For explanation of Table 1 see opposite.

were buried. The relative abundance of *Pilasporites allenii* and to a lesser extent of *Ischyosporites arkellii*, *Admolia amphidoxa*, and *Regresporites lophus* suggests both local production and some kind of ecological association between the parent plants. *R. lophus*, a relatively large thick-exined miospore, and *A. amphidoxa*, another thick-walled palynomorph, could not have been produced by plant species constituting part of the homogeneous delta flora. If they had been, they would have been recovered from sediments deposited closer to this source, particularly in medium and coarse siltstones of near-shore and back-delta facies. Thus *Equisetites* appears not to have been an important component of the onshore flora but probably thrived offshore in the pro-delta. This supports interpretations 3 and 4 of Allen (1959, 1967).

The local flora may be poorly represented in the soil beds and fragment partings. Possible reasons for the fluctuations in abundances include depositional environments unfavourable for preservation, winnowing, low rate of spore production, seasonal production, and variable rate of sedimentation (Batten 1968). Fragment partings yield *P. allenii* abundances more consistently than soil beds, probably because they contain a greater number of dispersed *E. lyellii* sporangia. Allen has suggested (*Pal. Ass. Circ.* 64, 1971, unpublished) that the larger Hastings Beds *Equisetites* (i.e. *E. lyellii*) may have flourished during periods of low salinity (cf. Anderson *et al.* 1967), that low salinity transgressions might have been followed by offshore *Equisetites*, and that brackish transgressions might not. The lack of dinoflagellates and acritarchs in soil beds and adjacent strata suggests that *Equisetites* flourished in fresh or only weakly saline waters. However, disagreement over the salinity ranges of several of the fossils of the Wadhurst Clay confuses the picture.

The Classopollis group of species is an important component of many Wadhurst assemblages. Hughes and Moody-Stuart (1967a) proposed that the group was not available for deposition in back-delta environments but that the parent plants were favourably placed for their pollen to be preserved in offshore deposits; they suggested a coastal swamp habitat comparable with the Tertiary and Recent mangroves. There is, however, no direct (megafossil) evidence for this and by contrast, Vakhrameev (1970) has suggested that the parent plants of Classopollis were psammophilic, preferring upland slopes, and able to tolerate drought conditions. An attempt to determine the relationship between Classopollis producers and Equisetites using palynology has merely shown that the Classopollis parent plants are not significantly associated with Equisetites, nor do they show an antipathetic relationship. There are some roots of unknown origin, which are larger than those of Equisetites, preserved in the coarse-grained Ashdown and Tunbridge Wells Sand facies (e.g. at Hastings, East Grinstead, and Pembury; White 1928, Allen 1959); Hughes and Moody-Stuart (1967a) suggested that Classopollis plants might have borne them, but attempts to prove this by palynologic association have been unsuccessful.

The affinities of the palynomorph taxa associated with *Pilasporites allenii* are uncertain because it is hazardous to compare them either with the spores of modern plants or with modern plant associations. It is probable that *Ischyosporites arkellii* is a filicalean spore (produced by a member of the Schizaeaceae?). *Regresporites lophus* is probably a fern spore, but *Retriletes* sp. could have been produced by a member of the Lycopodiaceae or of the Bryophyta, and the affinities of *Admolia* 

amphidoxa can only be guessed at (Thallophyta (Algae)?, Bryophyta?). However, the palynologic record does suggest that other plants grew, if not amongst the *Equisetites*, at least near by. That remains of the parent plants have not been recorded may be due either to non-preservation or lack of recognition. The only other record of plants suggested as being ecologically associated with *Equisetites* is that of 'doubtful leafy liverworts' (Allen 1959, p. 295).

TABLE 2A. Records of miospores tentatively attributed to the Equisetaceae (*Equisetites/Equisetum*) or compared with the spores of *Equisetum* (see *Pilasporites alleni* discussion). The spores that Rogalska (1962) attributed to cf. *Equisetum* differ from her earlier assignments (1954, 1956) in lacking a 'perisporium'. She noted that the earlier records may represent another genus.

Author	Record	Age
Rogalska 1956	Equisetum sp.	Lower Jurassic
Rogalska 1962	Cf. Equisetum	Jurassic
Góczán 1956	Calamospora cf. 'cf. C. pallida Schemel'	Lower Jurassic
Nagy 1958	Cf. Equisetum	Pliocene
Kara-Murza 1960	Leiotriletes spp.	Mesozoic
Chun'-Bin Chzhan 1962	Equisetum sp.	Lower Cretaceous
Verbitskaja 1962	Leiotriletes spp.	Cretaceous
Simonesics 1964	Perinosporites sphaericus Simonesics 1964	Miocene
Voevodova (in Pokrovskaja and Stel'mak) 1964	Equisetum chassynense Voevodova 1964	Lower Cretaceous
Piel 1971	?Equisetum sp.	Oligocene

TABLE 2B. Dispersed spores which bear some resemblance to Pilasporites allenii.

Author	Record	Age
Nakoman 1964	Inaperturopollenites problematicus Nakoman 1964	Oligocene
Wall 1965	Concentrisporites hallei (Nilsson 1958) Wall 1965	Lower Jurassic
Goubin 1965	Inaperturopollenites cf. orbicularis Nilsson 1958	Middle Jurassic
Goubin 1965	Laricoidites desquamatus Goubin 1965	Upper Triassic
Volkheimer 1968	Inaperturopollenites velatus Volkheimer 1968	Jurassic

### SYSTEMATIC PALAEONTOLOGY

Ischyosporites arkellii (Pocock 1970) comb. nov., recorded in graded comparison form (see Hughes and Moody-Stuart 1967b, p. 355), Retitriletes sp., and the four new taxa referred to in the text are here described and compared. Pilasporites allenii Batten 1968 is also discussed. Assemblage-type identifications (see Batten 1972, 1973) are given for the assemblages from which the taxa are described. The age/stage from which a compared species was described is given if it came from outside the Cretaceous period.

Miospores examined with the scanning electron microscope were individually mounted on 'Durofix' or double-sided 'Sellotape' on 12-mm diameter stubs, or strew mounted on stubs without adhesive. Most mounts were coated with carbon but gold-palladium was also used. The assemblage slides are labelled with the preparation number prefixed by 'T'. A holotype specimen is present with topotypes on a strew slide of a single preparation. The slides containing the figured specimens are in the Sedgwick Museum, Cambridge. Representative specimens of the new species are in the Institute of Geological Sciences, Leeds. Stage co-ordinates given refer to Leitz Laborlux (Ll) microscope, number 557187, Sedgwick Museum.

### Turma TRILETES Reinsch emend. Dettmann 1963

# Infraturma APICULATI Bennie and Kidston emend. Potonié 1956 Genus CONVERRUCOSISPORITES Potonié and Kremp 1954 Converrucosisporites venitus sp. nov.

Plate 41, figs. 1-9; Plate 42, figs. 1-5

Type sample. CUC 924, Cuckfield No. 1 Borehole, Sidnye Farm, Sussex (TQ 2961 2731), depth 924 ft (281.6 m); upper Ashdown Sand, Valanginian? Grey (N5-N7) laminated fine siltstone, sedimentary mica, bedding disturbed, rootlets in situ. Preparation T206; ten minutes Schulze solution, cleared in dilute NH<sub>4</sub>OH, mineral separation, strew slides with Clearcol. AT1.

Holotype. Slide T206/3, Ll 48·4 125·2; Pl. 41, fig. 3.

Diagnosis. Miospore, trilete, mean maximum diameter 41.5  $\mu$ m, standard deviation 4.1  $\mu$ m (100 specimens). Amb triangular; concave, straight or slightly convex sides, rounded angles. Proximal face slightly convex, may be raised up along margins of laesurae; distal surface convex. Laesurae distinct,  $> \frac{3}{4}$  radius. Exine two-layered, total thickness 1–2  $\mu$ m. Inner layer smooth 0.75–1.5  $\mu$ m thick. Outer layer 0.5–1  $\mu$ m thick, carries sculpture on both proximal and distal surfaces. Sculpture, which may be only weakly developed on proximal face, generally consists of grana and verrucae of low elevations ( $< 1-2 \mu$ m high) and variable maximum diameter (up to 6  $\mu$ m). Elements spaced  $< 1-2 \mu$ m (sometimes up to 4  $\mu$ m) apart; bases rounded to irregular in outline, distinct or (more usually) partially fused, leading to the formation of irregular rugulae.

Description. The observed limits of the maximum diameter of the spores are  $33-53 \,\mu m$  (coefficient of variation 9.8%). Fifty per cent of the specimens are in polar aspect; none are in equatorial view. Proximal arcuate folds resulting from compression of the raised up area of the proximal face may be present (Pl. 41, fig. 5). The sculptural elements are sparsely distributed on four specimens (Pl. 41, fig. 6), and predominantly dome shaped on seven (Pl. 41, fig. 3). Four show isolated clavae or dome shaped verrucae up to 6  $\mu m$  high (Pl. 41, fig. 1) and two have almost smooth proximal

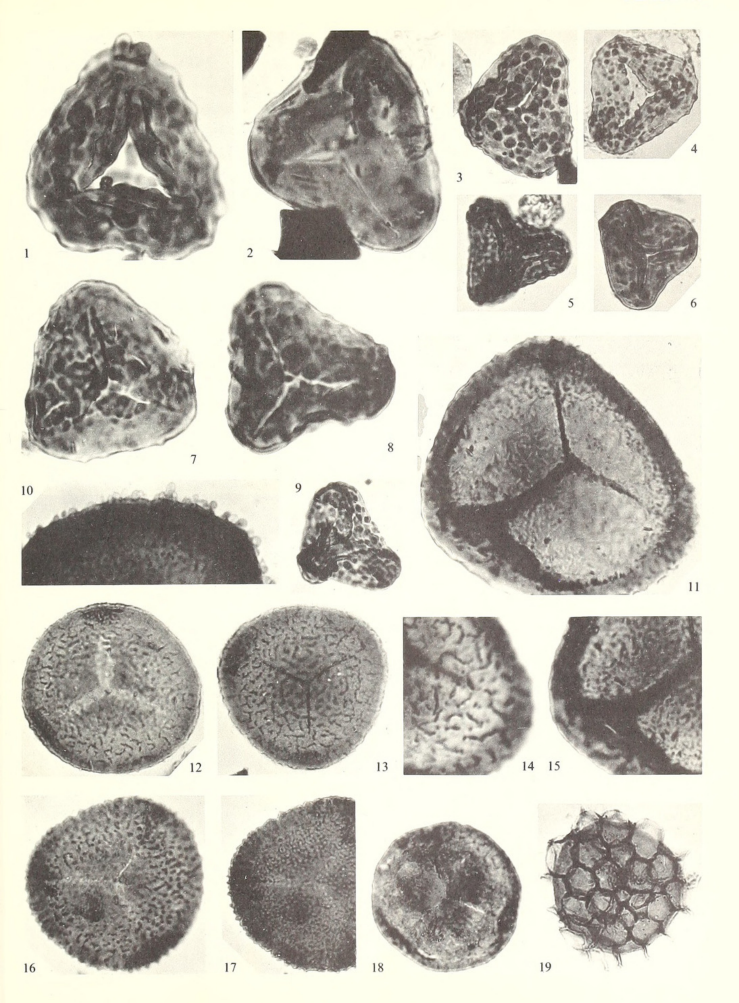
### EXPLANATION OF PLATE 41

Figs. 1–9. *Converrucosisporites venitus* sp. nov. Preparation T206. 1, Showing variable shape and distribution of sculptural elements and two layered exine, T206/1, Ll 54·1 123·2, ×1000. 2, Proximal surface almost smooth, T206/1, Ll 58·7 116·6, ×1000. 3, Holotype, T206/3, Ll 48·4 125·2, ×500. 4, T206/3, Ll 42·9 118·2, ×500. 5, Proximal kyrtomes, T206/1, Ll 38·7 120·2, ×500. 6, Sparsely distributed sculptural elements, T206/2, Ll 40·3 118·7, ×500. 7, T206/1, Ll 53·3 118·2, ×1000. 8, T206/1, Ll 43·3 118·1, ×1000. 9, T206/3, Ll 41·6 118·8, ×500.

Figs. 10–17. *Regresporites lophus* gen. et sp. nov. Preparation T209. 10, Equatorial sculpture, T209/1, L1 57·0 129·1, ×1000. 11, 14, 15, Same specimen. 11, Raised lips; 14, distal sculpture; 15, proximal sculpture; T209/1, L1 61·7 129·1, ×1000. 12, T209/1, L1 63·3 125·7, ×500. 13, T209/3, L1 58·4 126·0, ×1000. 16, 17, Same specimen, holotype. 16, Distal focus; 17, equatorial focus; T209/1, L1 41·5 126·4, ×500.

Fig. 18. cfA. Regresporites lophus gen. et sp. nov. Specimen corroded, preparation T210/13, Ll 58·7 123·2, × 500.

Fig. 19. Retitriletes sp. Preparation T210/13, Ll 54·3 115·7, × 500.



BATTEN, Cretaceous spores

surfaces (Pl. 41, fig. 2). Predepositional and diagenetic corrosion is shown by tearing and pitting of the exine of some specimens.

Discussion. The sculptural elements of Converrucosisporites cameroni (de Jersey 1962) Playford and Dettmann 1965 (Triassic) rarely coalesce, and although they consist mostly of verrucae, they may also comprise some bacula, spinae, coni, and grana. The closely spaced verrucae on the exine of C. geniculatus Deák and Combaz 1967 are more rounded and more strongly developed. 'Trilitisporites' rariverrucatus Danzé-Corsin and Laveine 1963 (lower Lias) has more or less rounded verrucae 1–3 µm high scattered widely over its surface. Concavissimisporites southeyensis Pocock 1970 (lower Bajocian) is larger and has a sexine which is thicker in equatorial radial regions than interradially. Concavisporites variverrucatus Couper 1958 is somewhat larger and has a thicker exine. Converrucosisporites venitus resembles some Carboniferous species of the genera Pustulatisporites Potonié and Kremp 1954 and Converrucosisporites. The specimens ascribed by Lantz (1958, pl. 1, fig. 12) to Leptolepidites cf. verrucatus Couper 1953 are similar. Forms recorded as Converrucosisporites venitus have been recovered from the Ashdown Beds, Wadhurst Clay, Lower Tunbridge Wells Sand, and Grinstead Clay (Berriasian-Valanginian?).

# Infraturma MURORNATI Potonié and Kremp 1954

# REGRESPORITES gen. nov.

Type species. R. lophus sp. nov.

Diagnosis. Miospore, trilete, rounded triangular to subcircular in equatorial outline. Extremities of laesurae usually merge with equatorial or proximal subequatorial exine thickening. Distal sculpture of irregular, occasionally anastomosing rugulae  $\pm$  scattered grana, verrucae, clavae, or bacula. Proximal sculpture similar but reduced, may consist only of grana, rarely scabrate.

Discussion. Spores which structurally resemble Regresporites include Retusotriletes Naumova 1953 emend. Streel 1964 and related genera (e.g. Apiculiretusispora Streel 1964) described from the Palaeozoic. Rugulatisporites Thomson and Pflug 1953 is used for rugulate Mesozoic spores but the diagnosis of the genus does not allow for the inclusion of forms which show modifications of the exine in equatorial

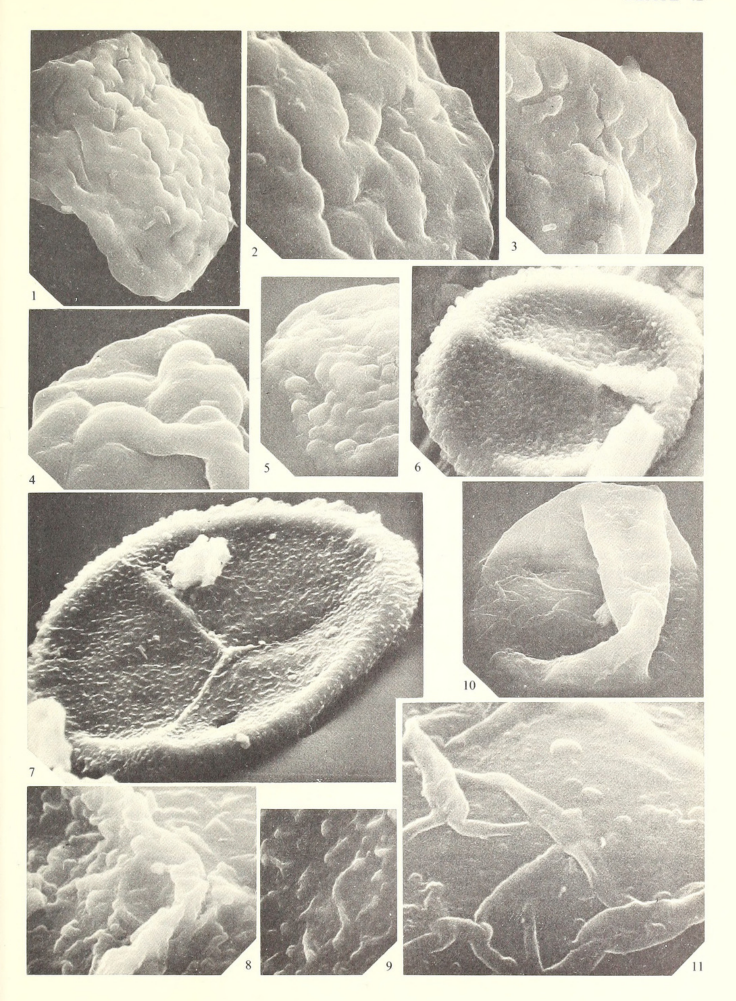
#### EXPLANATION OF PLATE 42

All scanning electron micrographs.

Figs. 1-5. Convertucosisporites venitus sp. nov. Preparation T206. 1, 2, Specimen on stub (SH) DB 44. 1, Specimen obliquely compressed, ×1500. 2, Detail of distal sculpture, ×3500. 3, Second specimen, distal view, ×1500, (SH) DB 50. 4, Third specimen, detail of proximal radial region, ×3500, (SH) DB 44. 5, Part of distal surface of fourth specimen, ×1500, (SH) DB 44.

Figs. 6-9. Regresporites lophus gen. et sp. nov. Preparation T209. 6, Slightly oblique compression, × 1000, (SH) DB 36. 7, Second specimen showing raised lips and thickened proximal equatorial regions, × 1000, (SH) DB 1a. 8, Third specimen, detail of proximal pole, × 4000, (SH) DB 34. 9, Distal sculpture detail of fourth specimen, × 2000, (SH) DB 36.

Figs. 10–11. *Pilasporites allenii* Batten 1968. Preparation T210. Specimen on (SH) DB 47. 10, ×1500. 11, Detail of loosened ?ektexine showing crinkling and attached globules, ×7500.



BATTEN, Cretaceous spores

or subequatorial regions. *Lycopodiacidites* Couper 1953 emend. Potonié 1956 is generally used for azonate miospores which are verrucate to rugulate on their distal surface. *Hamulatisporis* Krutzsch 1959, erected from the Tertiary, is a subtriangular spore with hamulate sculpture.

# Regresporites lophus sp. nov.

Plate 41, figs. 10-17; Plate 42, figs. 6-9; Plate 44, figs. 1-6

Type sample. CUC 791/11, Cuckfield No. 1 Borehole, Sidnye Farm, Sussex (TQ 2962 2729), depth 791 ft 11 in (241·4 m), Wadhurst Clay, Valanginian? Medium grey (N5) clay, thin fine silt laminations, Equisetites in situ and fragments of same. Preparation T209; 10 minutes Schulze solution, cleared in dilute NH<sub>4</sub>OH, mineral separation, strew slides with Clearcol. AT8. Holotype slide T209/1, Ll 41·5 126·4; Pl. 41, figs. 16, 17.

Diagnosis. Miospore, trilete, mean maximum diameter  $62.6 \mu m$ , standard deviation  $5.5 \mu m$  (100 specimens). Amb rounded triangular to subcircular. Distal surface (when uncompressed) more strongly convex than proximal. Laesurae length  $\frac{3}{4}$  spore radius or more, straight or (rarely) slightly sinuous. Lips may be raised,  $0.5-2 \mu m$  wide at base on each side of laesurae and up to  $5 \mu m$  high (measured on thirty-eight specimens). Extremities of laesurae usually merge with equatorial or subequatorial exinal thickening which appears as a dark zone in transmitted light. Width of this zone usually  $5-7 \mu m$  interradially and  $6-11 \mu m$  radially; inner margin usually indistinct.

Exine 2-3  $\mu$ m thick except near equator where it may be up to 4.5  $\mu$ m thick in interradial regions and up to 5  $\mu$ m thick at angles, probably two-layered; outer layer thin (c. 1  $\mu$ m?), carries sculpture. A subcircular area (outline indistinct) at proximal pole may be slightly darkened in transmitted light (= slight thickening of exine); observed limits of maximum diameter 19-36  $\mu$ m, mean 26.3  $\mu$ m (sixty-six specimens).

Proximal sculpture usually of small grana and/or occasionally anastomosing irregular rugulae (up to 1  $\mu$ m high, < 4  $\mu$ m in length, up to 3  $\mu$ m apart). Distal and equatorial sculpture of rugulae  $\pm$  scattered grana, verrucae, clavae, and baculae; rugulae irregular, occasionally anastomosing, up to 4  $\mu$ m apart, 4–10  $\mu$ m long, usually 0·5–2  $\mu$ m (rarely up to 3·5  $\mu$ m) wide, and < 2  $\mu$ m (rarely up to 4  $\mu$ m) high.

Description. The observed limits of the maximum diameter of these spores are  $51-80 \mu m$  (coefficient of variation 8.7%). Seventy-five of the specimens are in polar aspect but none are in equatorial view. The laesurae are sometimes only weakly developed (Pl. 42, fig. 6 in part; Pl. 44, fig. 4). The dimensions of the sculptural elements on the proximal face tend to increase towards the equator. Eighty-two specimens have both grana and rugulae proximally (Pl. 41, fig. 15; Pl. 42, fig. 7). Most specimens are very compressed, usually in polar orientation or slightly obliquely. The proximal face is often concave (Pl. 42, figs. 6, 7; Pl. 44, fig. 4). The sculptured layer of the exine has been loosened on, and partially removed from, some specimens. Twenty-four specimens show a single fold which in most cases is developed subparallel to the equator.

Discussion. Forms identified as Regresporites lophus have only been recorded from the Wadhurst and Grinstead Clays (Valanginian?).

# Genus RETITRILETES van der Hammen *ex* Pierce 1961 *Retitriletes* sp.

Plate 41, fig. 19; Plate 43, fig. 1

Description. The observed limits of the maximum diameter of this miospore, excluding sculpture, are 40-75  $\mu$ m (twenty specimens). The amb is circular or subcircular in outline. An irregular weakly developed (discontinuous) triradiate ridge may be present on the proximal face extending to the equator of the spore body. The exine is 2-3  $\mu$ m thick and sculptured with a reticulum both on the distal face and equatorially. On the proximal face the sculpture is reduced, sometimes to a low reticulum, but more often to irregular grana, and muri < 3  $\mu$ m high. Near the trilete mark the exine may be scabrate. The distal and equatorial muri are 0.5-1.5  $\mu$ m wide at their bases but rapidly become membraneous above and are up to 12  $\mu$ m high. The lumina are 4-21  $\mu$ m in maximum diameter and circular to polygonal in outline.

Discussion. Species of Januasporites Pocock 1962 emend. Singh 1964 are alete miospores possessing a two-layered exine. The outer layer is absent over a more or less circular area on the distal face. The muri of Reticulatisporites arcuatus Brenner 1963 are thicker and more irregular in width. The proximal sculpture of Hymenoreticulisporites altimuronatus Döring 1964 (Jurassic; Upper Malm) is not reduced. The laesurae of R. castellatus Pocock 1962 are not visible and the proximal sculpture is not reduced. Woodsia reticulata Bolkhovitina 1953 appears to possess a clearly defined triradiate mark. R. incomptus Manum 1962 (Tertiary) has muri of lower elevations and is clearly triradiate. Zlivisporis blanensis Pacltová 1961 is clearly trilete, lacks sculpture on the proximal face, and the arrangement of the muri is different. Lycopodiumsporites marginatus Singh 1964 differs chiefly in having a stronger trilete mark. Retitriletes sp. bears some resemblance to palynomorphs which have been referred to species of the acritarch genus Cymatiosphaera (cf. Wall 1965, pl. 9, fig. 8). Retitriletes sp. has only been recorded from the Wadhurst Clay (Valanginian?).

Infraturma AURICULATI Schopf emend. Dettmann 1963

Ischyosporites arkellii (Pocock 1970) comb. nov.

1970 Dictyotriletes arkelli Pocock, p. 52, pl. 9, fig. 16A; D. arkellii, p. 119.

Pocock's species is included here in *Ischyosporites* because it has a valvate exine.

Cf. B. Ischyosporites arkellii

Plate 43, figs. 2-14; Plate 44, figs. 7-11; Plate 46, fig. 2

Sample. CUC 792, Cuckfield No. 1 Borehole, Sidnye Farm, Sussex (TQ 2962 2729), depth 792 ft (241·4 m); Wadhurst clay, Valanginian? Medium dark grey (N4) clay, thin fine silt laminations. Equisetites in situ and fragments of same. Preparation T210; 10 minutes Schulze solution, cleared in dilute NH<sub>4</sub>OH, mineral separation, strew slides with Clearcol. AT8/16.

Description. A trilete miospore with a mean maximum equatorial diameter of  $43.6 \mu m$ 

(standard deviation  $5\cdot 1~\mu m$ , observed limits  $32-57~\mu m$ , coefficient of variation  $11\cdot 7\%$  (100 specimens)) and a mean maximum polar diameter (including membraneous lips of trilete mark) of  $45~\mu m$  (twenty-two specimens). The amb has straight to slightly convex sides and rounded angles. The proximal face is slightly, and the distal face markedly, convex. The laesurae are straight or slightly sinuous and are longer than three-quarters of the spore radius. The membraneous lips of the trilete mark are up to  $5~\mu m$  high and have a basal width of  $1\cdot 5-2~\mu m$ . The (perfect or imperfect) distal reticulum encroaches on to the otherwise smooth proximal face at the angles. The muri are rounded or (sometimes) flat-topped,  $0\cdot 5-3\cdot 5~\mu m$  (usually  $1-2\cdot 5~\mu m$ ) high, highest where they join and at the angles. They are  $1-2\cdot 5~\mu m$  wide except at intersections where the width may increase to  $4~\mu m$ . The lumina are generally subcircular to polygonal and  $2-7~\mu m$  in diameter but may be larger and irregular if the reticulum is imperfect. They tend to be smallest in equatorial radial regions.

Thirty-two specimens are in polar and twenty-two in equatorial aspect. One weakly sculptured specimen has unusually convex sides (Pl. 43, fig. 9) but this is probably because the triradiate lips have parted. The compression of the specimens varies from relatively uncompressed to very compressed. A few show signs of corrosion in the form of pitting of the exine.

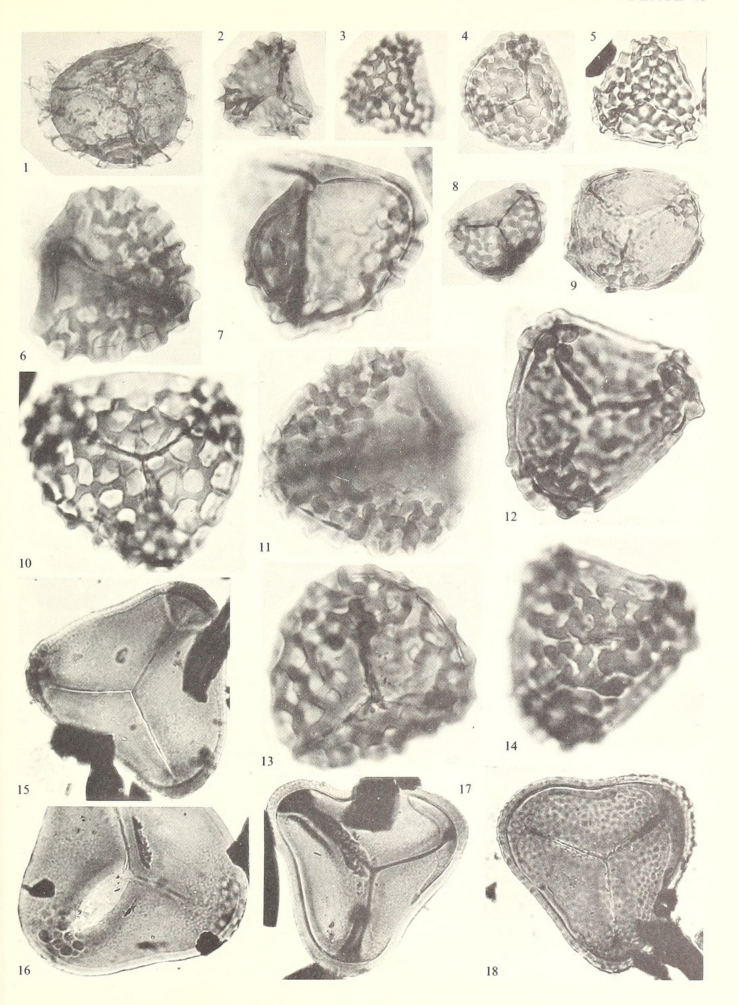
Discussion. The lumina of Ischyosporites crateris Balme 1957 are larger and the exine is thicker. I. punctatus Cookson and Dettmann 1958 has a thicker exine, foveoreticulate sculpture and wider muri. I. marburgensis de Jersey 1963 (Jurassic) has a different sculpture and a thicker exine. 'Lycopodiacidisporites' cerniidites (Ross 1949; Delcourt and Sprumont 1955) Danzé-Corsin and Laveine 1963 (in Briche et al. 1963; Lower Lias), has cristate sculpture disposed more or less concentrically and not forming a true reticulum. Lycopodiumsporites crassimacerius Hedlund 1966 has a thicker exine, is sculptured with a polygonal reticulum, and has higher muri. The distal sculpture of 'Ischyisporites' puzzlei Levet-Carette 1966 consists of little cavities, rounded or more often vermiform in shape, separated by muri 3–5 μm wide. Ischyosporites disjunctus Singh 1971 is larger, and the muri are wider and the exine thicker. The nature of the murornate sculpture and/or other features distinguishes I. arkellii from the following: L. dentimuratus Brenner 1963, 'Ischyisporites' surangulus Levet-Carette 1964 (Bajocian), Foveasporis microfovearis Krutzsch 1959 (Tertiary), Ischyosporites asolidus (Krutzsch 1959) Krutzsch 1967 (Tertiary), I. foveasolidus

# EXPLANATION OF PLATE 43

Fig. 1. Retitriletes sp. Preparation T210/14, Ll 47-9 123-0, ×500.

Figs. 2–14. cfB. *Ischyosporites arkellii* (Pocock 1970) comb. nov. Preparation T210. 2, 3, Same specimen. 2, Proximal focus; 3, distal focus; T210/14, L1 30·3 121·6, ×500. 4, T210/11, L1 35·6 121·2, ×500. 5, T210/2, L1 42·7 118·0, ×500. 6, Lateral view, T210/2, L1 28·7 118·7, ×1000. 7, Oblique view, T210/1, L1 48·3 115·8, ×1000. 8, T210/14, L1 25·2 119·7, ×500. 9, Specimen weakly sculptured, unusually convex sides, T210/2, L1 46·0 122·8, ×500. 10, Distal focus, T210/13, L1 60·7 123·7, ×1000. 11, Lateral view, T210/14, L1 48·6 123·0, ×1000. 12, 14, Same specimen. 12, Proximal focus; 14, distal focus; T210/3, L1 48·0 122·2, ×1000. 13, Oblique view, T210/14, L1 47·5 122·0, ×1000.

Figs. 15–18. *Trilobosporites ivanovae* sp. nov. Preparation T061. 15, Holotype T061/1, L1 43·6 109·8, ×500. 16, T061/1, L1 37·3 111·8, ×500. 17, Margins of laesurae thickened, T061/1 L1 52·7 113·0, ×500. 18, T061/1, L1 27·0 113·8, ×500.



BATTEN, Cretaceous spores

Krutzsch 1967 (Tertiary), I. tuberosus Döring 1965, and F. triangulus Stanley 1965 (Paleocene).

The spores *Dictyotriletes arkelli* Pocock 1970 from the lower Bajocian of western Canada (not recorded from younger strata) are similar to those described here. Some characters differ slightly, but there are not sufficient criteria for the erection of a new species for the Wealden spores. The illustrations of *Klukisporites pseudoreticulatus* Couper 1958 in Agasie (1969) resemble some of those of *I. arkellii* shown here. Forms comparable with *I. arkellii* have been recorded from the Ashdown Beds, Wadhurst Clay, Lower Tunbridge Wells Sand, and Grinstead Clay (Berriasian-Valanginian?).

# Trilobosporites ivanovae sp. nov.

Plate 43, figs. 15-18; Plate 45, figs. 1-8; Plate 46, figs. 1, 5-9

Type sample. DJB 18, Philpots Quarry, West Hoathly, Sussex (TQ 3592 3221), 5·4 m above base of Grinstead Clay, Valanginian? Light brown (5YR 6/4) medium silt, plant fragments. Preparation T061; 20 minutes HNO<sub>3</sub>, cleared in dilute NH<sub>4</sub>OH, mineral separation, strew slides with Clearcol. AT13/14; S4, 5. Holotype slide T061/1, Ll. 43·6 109·8; Pl. 43, fig. 15.

Diagnosis. Miospore, trilete, mean maximum diameter 92  $\mu$ m, standard deviation 8·3  $\mu$ m (100 specimens). Amb triangular with slightly concave, straight or slightly convex sides and rounded angles. Distal surface more convex than proximal. Laesurae about  $\frac{3}{4}$  spore radius, may divide near extremities, bordered by simple membraneous lips  $1-2\cdot5$   $\mu$ m high. Exine may be slightly thickened for 4-7  $\mu$ m on each side of laesurae (forty-four specimens). Width of thickened zone may increase slightly near equatorial radial regions and merge with equatorial radial thickenings.

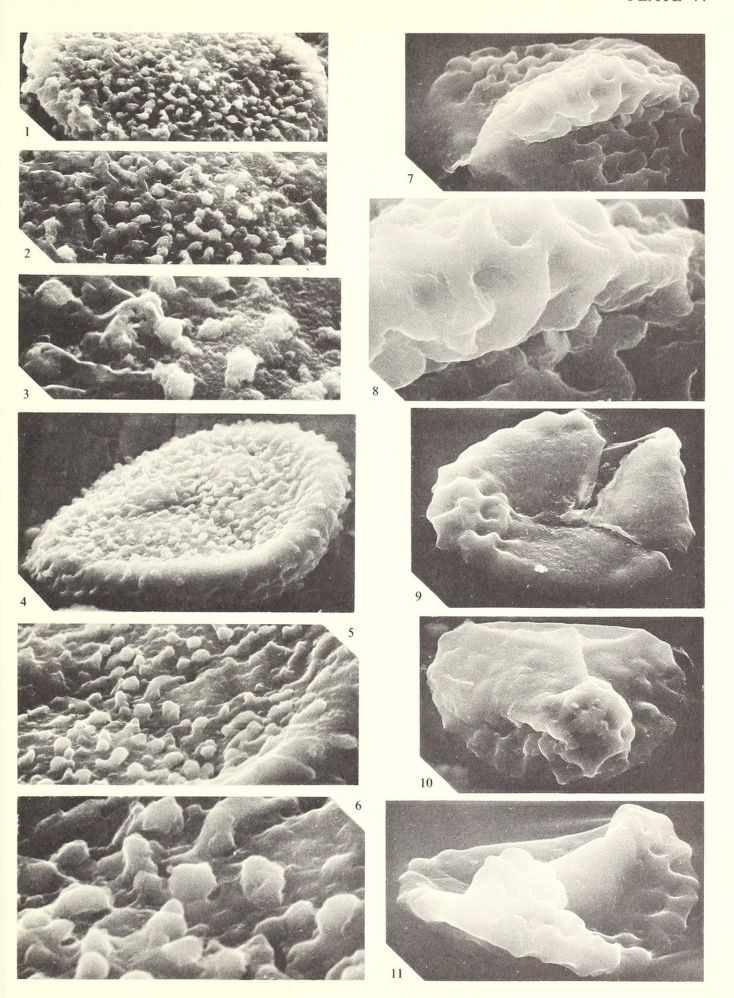
Exine mainly 3–5  $\mu$ m thick but generally thicker (up to 8  $\mu$ m) in equatorial radial regions; increase in thickness varies from 0·5 to 4  $\mu$ m. Thickened areas extend from 14 to 25  $\mu$ m over proximal and distal faces; boundaries usually indistinct. Exine scabrate, maculate, or sculptured with closely spaced (0·25  $\mu$ m or less apart) grana and/or verrucae. Verrucae subcircular, polygonal, or somewhat irregular in shape, usually < 1  $\mu$ m high and with a diameter of < 3  $\mu$ m interradially and < 5  $\mu$ m at the angles, may form a negative reticulum. Sculpture often more strongly developed on distal face and/or near equatorial radial regions. Forms with verrucate distal exines are usually maculate or scabrate in vicinity of proximal pole and in proximal interradial regions.

## EXPLANATION OF PLATE 44

All scanning electron micrographs.

Figs. 1-6. Regresporites lophus gen. et sp. nov. 1, 2, 3, Fifth specimen, on stub (SH) DB 36. 1, Part of distal surface, ×1000. 2, Sculpture detail, ×1500. 3, Sculpture detail, ×4000. 4, 5, 6, Sixth specimen, (SH) DB 36. 4, Proximal surface, trilete mark weakly developed, ×1000. 5, Sculpture detail, ×2000. 6, Sculpture detail, ×5000.

Figs. 7-11. cfB. *Ischyosporites arkellii* (Pocock 1970) comb. nov. Preparation T210. 7, 8, Specimen laterally compressed, (SH) DB 46. 7, ×1500. 8, Sculpture detail, ×3500. 9, Second specimen, proximal surface, ×1500, (SH) DB 44. 10, Third specimen, lateral view, ×1500, (SH) DB 46. 11, Fourth specimen, ×1500, (SH) DB 44.



BATTEN, Cretaceous spores

Description. The observed limits of the maximum equatorial diameter of this miospore are 75-126  $\mu$ m (coefficient of variation 9%) and of the maximum polar diameter are 62-80  $\mu$ m (seven specimens). Although the increase in thickness of the exine from interradial to radial regions is usually about the same at each angle, differences of up to 3.5  $\mu$ m have been noted. There is no increase in thickness at one or two of the angles of three specimens (Pl. 45, fig. 6). Three specimens which are sculptured with a few verrucae up to 7  $\mu$ m in diameter and 3  $\mu$ m in height (Pl. 45, figs. 10-12) are considered to be extreme variants.

Discussion. Trilobosporites (Trilobosporites) tenuiparietalis Döring 1965 has a thinner exine and more strongly concave sides. Maculatisporites microverrucatus Döring 1964 and M. undulatus Döring 1964, both from the Wealden A of Westmecklenburg, lack thickening of the exine in equatorial radial regions and are somewhat smaller; the exine of M. microverrucatus is also thinner. Rare specimens of T. ivanovae resemble Lygodium (Tuberculata) triangulatum E. Ivanova 1961 but the latter has a thinner exine.

The descriptions and illustrations of both Lygodium (Cavernosotriangulata) cavernosum E. Ivanova 1961 and L. (C.) cavernosum var. tuberculatum E. Ivanova 1961 (in Samoilovitch et al. 1961; Cenomanian) suggest that these taxa may fall within T. ivanovae, but in Ivanova's descriptions are insufficient to confirm this. Some of the weakly sculptured specimens of T. ivanovae resemble both T. (Trilobosporites) aornatus Döring 1965 (Upper Malm) and T. (T.) crassiangularis Döring 1965 (German Wealden A).

Forms comparable with *T. ivanovae* have been recorded from the Wadhurst Clay, Lower Tunbridge Wells Sand, and Grinstead Clay (Berriasian-Valanginian?).

### Turma ALETES Ibrahim 1933

Subturma AZONALETES Luber emend. Potonié and Kremp 1954 Genus PILASPORITES Balme and Hennelly 1956 Pilasporites allenii Batten 1968

Plate 42, figs. 10, 11

A specimen of *Pilasporites allenii* from the type sample (CUC 792) was illustrated in Batten (1968, pl. 123, fig. 11) and described as having granules and small verrucate

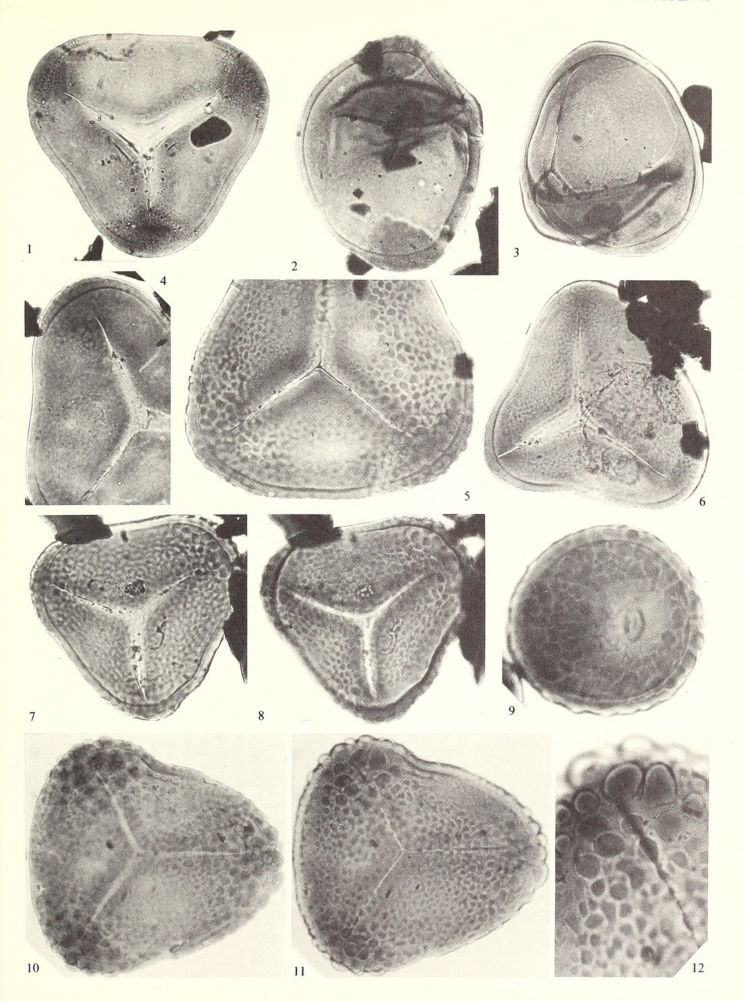
### EXPLANATION OF PLATE 45

All figures  $\times$  500 unless otherwise specified.

Figs. 1-8. Trilobosporites ivanovae sp. nov. Preparation T061. 1, Specimen with scabrate exine and scattered corrosion pits, T061/1, L1 35·1 112·2. 2, Lateral view showing equatorial radial thickenings, T061/1, L1 57·1 111·1. 3, Oblique orientation, T061/1, L1 35·7 114·3. 4, Small verrucae on distal face near equatorial radial thickening, T061/1, L1 43·0 111·1. 5, Unusually large specimen, T061/1, L1 44·9 114·2. 6, Exine weakly thickened in equatorial radial regions, T061/1, L1 35·3 113·7. 7, 8, Same specimen, strongly sculptured. 7, Equatorial focus; 8, distal focus; T061/1, L1 41·9 121·0.

Fig. 9. Admolia amphidoxa gen. et sp. nov. Preparation T209/2, Ll 39·9 124·2, ×1000.

Figs. 10–12. cfA. *Trilobosporites ivanovae* sp. nov., showing strongly developed sculpture in equatorial radial regions. 10, Equatorial focus; 11, proximal focus; 12, detail, ×1000; preparation T053/2, Ll 36·0 114·3.



BATTEN, Cretaceous spores

elements attached to a closely adhering perine. These elements (Pl. 42, fig. 11) are similar to the 'resin droplets' observed by Smith and Butterworth (1967, p. 145) on the Carboniferous miospore *Cadiospora magna* Kosanke 1950, and to the 'globules of ?tapetal material' (Evans 1970, p. 65) characteristically attached to the 'exoexine' of species of *Diaphanospora* Balme and Hassell 1962 emend. Evans 1970. They may also be comparable with Ubisch bodies (cf. Rowley 1963; Pettitt 1966). Lugardon (1969) showed similar bodies adhering to the perispore of spores of *Equisetum maximum* Lamk. Abnormal spores (not Ubisch bodies) attached to *P. allenii* exines occasionally occur in Wealden dispersed spore preparations. They are comparable with those of Gould (1968, pl. 2, fig. 11). Specimens of *P. allenii* may pseudomorph *Cycadopites* or *Taxodiaceaepollenites* and resemble probable algal bodies, e.g. *Halosphaeropsis liassica* Mädler 1963 (Lias) and *Leiosphaeridia pusilla* Mädler 1963 (Lias). Species of *Inaperturopollenites* generally have thinner exines containing many more secondary folds than is usual in *P. allenii*.

P. allenii was compared by Batten (1968) with other miospore species, some attributed to the Equisetaceae (Equisetites/Equisetum). Additional records of dispersed miospores which have been (tentatively) attributed to the family or genera or compared with the spores of Equisetum are listed on Table 2. Other dispersed spore species resembling P. allenii are also listed. Perhaps most closely comparable with P. allenii is P. marcidus Balme 1957 (Batten 1968, p. 641), from the Lower Jurassic of Western Australia, and recorded from both the Jurassic and Cretaceous. It may have affinities with the Equisetaceae (Balme 1957, p. 28) but other species of Pilasporites (including the type species P. calculus Balme and Hennelly 1956, P. plurigenus Balme and Hennelly 1956, and others) are unlikely to be so related.

# INCERTAE SEDIS ADMOLIA gen. nov.

Type species. A. amphidoxa sp. nov.

*Diagnosis*. Spherical or subspherical body in uncompressed state. Exine differentially thickened, smooth or sculptured, may show a circular or subcircular perforation (rarely a pore) on thinnest part.

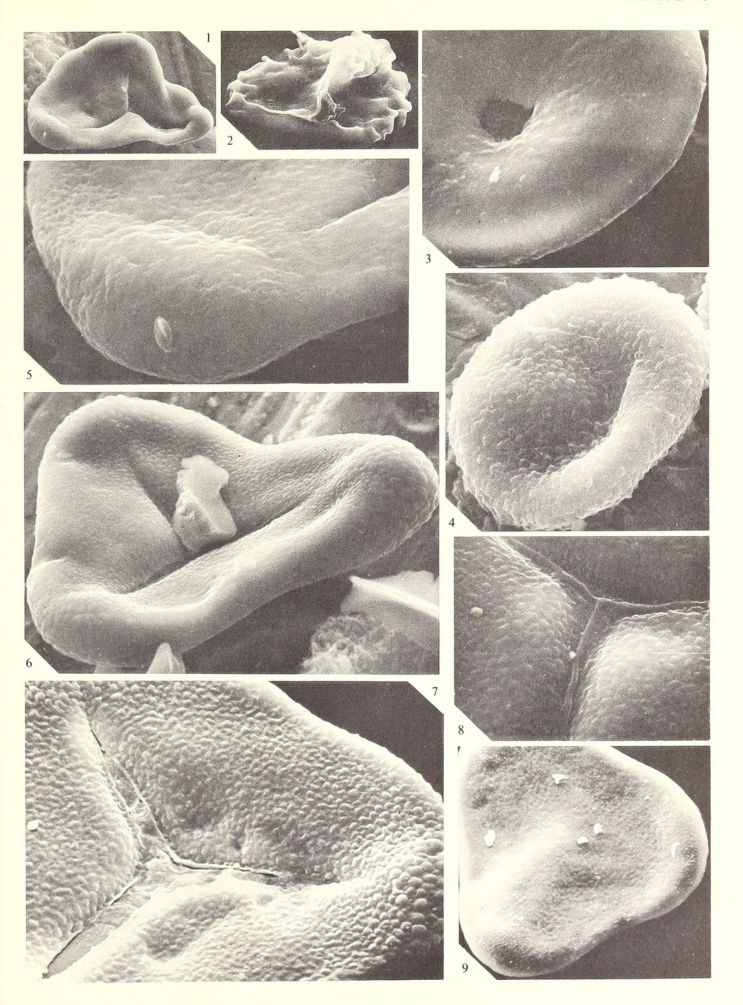
Discussion. Forms referable to Exesipollenites have a circular depression, which probably represents a pore, surrounded by exine thickening. The wall of Discoperculina Phillips 1971 is thickest by the operculum; this palynomorph also has a thin-

### EXPLANATION OF PLATE 46

All scanning electron micrographs.

Figs. 1, 5–9. *Trilobosporites ivanovae* sp. nov. Preparation T061. 1, 5, Specimen on stub (SH) DB 33. 1, Proximal view, ×400. 5, Detail of radial equatorial region, ×2000. 6, Second specimen, ×1000, (SH) DB 33. 7, Third specimen, part of proximal exine, ×1000, (SH) DB 8. 8, Fourth specimen, detail of proximal pole, ×2000, (SH) DB 33. 9, Fifth specimen, distal surface, ×500, (SH) DB 8. Fig. 2. cfB. *Ischyosporites arkellii* (Pocock 1970) comb. nov. Fifth specimen, ×750, (SH) DB 46.

Figs. 3-4. Admolia amphidoxa gen. et sp. nov. Preparation T209. 3, Specimen with pore, ×2000, (SH) DB 34. 4, Second specimen, ×1500, (SH) DB 36.



BATTEN, Cretaceous spores

walled inner body. O. carbonis, of the monotypic genus Operculites Newman 1965 is smaller than A. amphidoxa, bears a rounded or helicoidal operculum of width about  $\frac{1}{3}-\frac{2}{3}$  grain diameter attached at one side when in place, and the exine is thinner and not differentially thickened.

Admolia amphidoxa sp. nov.

Plate 45, fig. 9; Plate 46, figs. 3, 4; Plate 47, figs. 1-14

Type sample. CUC 791/11, preparation T209; details under Regresporites lophus sp. nov. Holotype slide T209/2, Ll 27·4 115·1; Pl. 47, fig. 5.

Diagnosis. Mean maximum diameter of grain  $42.5~\mu m$ , standard deviation  $4.7~\mu m$  (200 specimens). Amb circular or subcircular. A circular or subcircular perforation of exine (rarely a pore or tear) 5–8  $\mu m$  in diameter, may be present on thinnest part of differentially thickened exine. Perforation frequently only partially separates (sub)circular area of exine (operculum) from surrounding exine, but separation may be as much as 3  $\mu m$ . Thickness of exine mainly between 2 and  $4.5~\mu m$ ; difference between thinnest and thickest parts varies considerably (from  $< 0.5~to 3~\mu m$ ), thickest on face opposite perforation. Exine smooth, gently undulating, or scabrate, or sculptured with micrograna, grana, or dome-shaped verrucae or combinations of these. Sculpture often more weakly developed on thinnest part of wall. Height of verrucae usually 1  $\mu m$  or less, but may be as much as 3  $\mu m$ . Elements sometimes separated by distances greater than their diameter but usually closely spaced and partially coalescent, may form a negative reticulum.

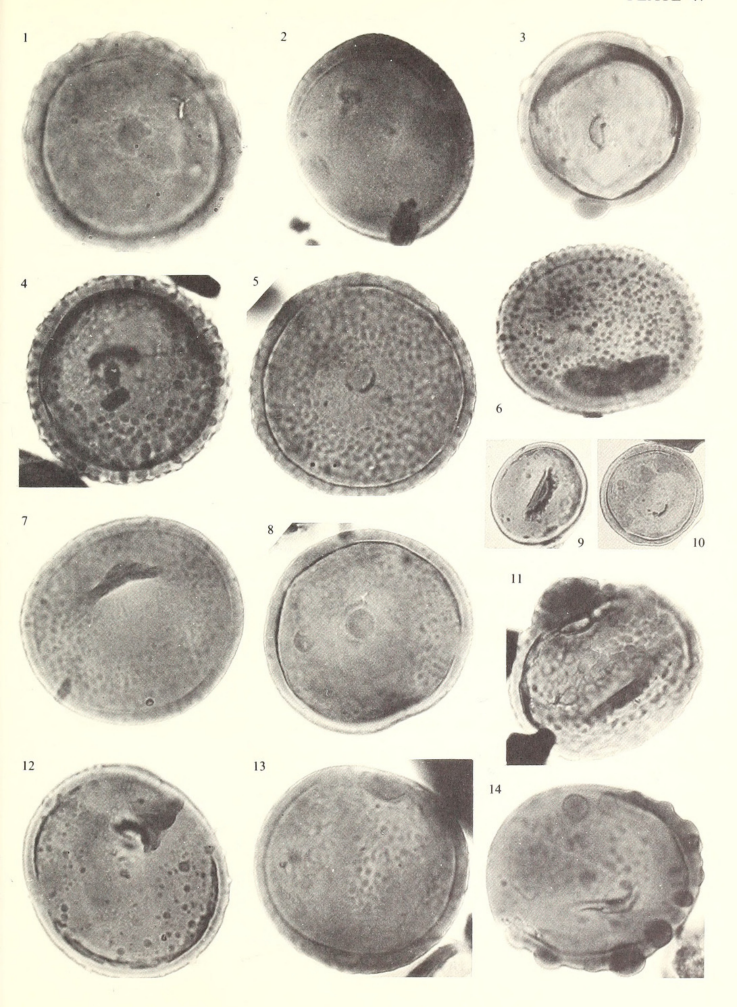
Description. The observed limits of the maximum diameter of the body of this palynomorph are  $33-57~\mu m$  (coefficient of variation  $11\cdot1\%$ ). The exine of seven specimens is c.  $1\cdot5~\mu m$  around the perforation. One-hundred-and-two specimens show a small circular or subcircular perforation in the exine (Pl. 47, fig. 8), eight show a pore or tear (Pl. 46, fig. 3). The perforation may be obscured by a small fold developed on the thin area, and it is invisible in some compression states or orientations. Discrete grana, bacula, or verrucae may be superimposed on low verrucae. The specimens are compressed but generally well preserved.

Discussion. Foveoinaperturites paucipunctatus Pierce 1961 is larger and has an infrapunctate, sparsely foveolate exine. Some phytoplankton show a naturally constituted exit hole. Forms probably comparable with A. amphidoxa have been seen in the Tertiary (A. E. González Guzmán, pers. comm.). The relatively small size variation

### EXPLANATION OF PLATE 47

All figures  $\times 1000$  unless otherwise specified.

Figs. 1–14. *Admolia amphidoxa* gen. et sp. nov. Preparation T209. 1, T209/2, Ll 37·7 109·3. 2, Showing perforation of thinnest part of differentially thickened exine, T209/2, Ll 52·1 113·0. 3, T209/2, Ll 25·3 117·8. 4, T209/1, Ll 54·6 116·7. 5, Holotype, T209/2, Ll 27·4 115·1. 6, T209/2, Ll 29·0 123·8. 7, Perforated area of exine on fold, T209/2, Ll 38·3 119·6. 8, T209/2, Ll 42·2 124·2. 9, Smooth exine, T209/2, Ll 28·8 114·1, × 500. 10, Partial separation of circular area from surrounding exine, T209/2, Ll 25·8 118·9, × 500. 11, Large verrucae, T209/2, Ll 33·6 115·3. 12, T209/2, Ll 34·7 124·7. 13, T209/2, Ll 51·1 118·0. 14, Large dome-shaped verrucae, T209/1, Ll 47·2 129·6.



BATTEN, Cretaceous spores

and the variety of sculpture observed in this population is usual. DJB 294, a sample from the Wadhurst Clay of the High Brooms Brick and Tile Company's pit at Southborough, Kent (TQ 5948 4189), however, yielded an assemblage composed almost entirely of smooth forms. The High Brooms assemblages are mostly smaller and darker in colour than the population on which this species is based. Specimens identified as *A. amphidoxa* have so far only been recorded from the Wadhurst and Grinstead Clays (Valanginian?).

Acknowledgements. The work for this paper was carried out whilst in receipt of a N.E.R.C. grant. I thank the directors of Robertson Research International for facilities and for a generous contribution towards the cost of publication. I am grateful to Mr. John Odell for help with the computer programming, and to Professor P. Allen for commenting on the manuscript. Material from the Cuckfield No. 1 and Warlingham Boreholes and access to the log of the former was made available by the Institute of Geological Sciences.

### REFERENCES

- AGASIE, J. M. 1969. Late Cretaceous palynomorphs from northeastern Arizona. *Micropaleontology*, **15**, 13–30, 4 pls.
- ALLEN, P. 1941. A Wealden soil bed with *Equisetites lyelli* (Mantell). *Proc. Geol. Ass.* 52, 362-374, 1 pl. —— 1949. Wealden petrology: the Top Ashdown Pebble Bed and the Top Ashdown Sandstone. *Q. J. geol. Soc. Lond.* 104, 257-321, 1 pl.
- —— 1959. The Wealden environment: Anglo-Paris Basin. Phil. Trans. R. Soc. 242B, 283-346.
- —— 1960. Geology of the Central Weald: a study of the Hastings Beds. Geol. Ass. Guide, 24, 28 pp.
- —— 1962. The Hastings Beds deltas: recent progress and Easter field meeting report. *Proc. Geol. Ass.* 73, 219–243, 3 pls.
- —— 1967. Strand-line pebbles in the mid-Hastings Beds and the geology of the London Uplands. Old Red Sandstone, New Red Sandstone and other pebbles. Conclusion. Ibid. **78**, 241–276, 3 pls.
- ANDERSON, F. W., BAZLEY, R. A. B. and SHEPHARD-THORN, E. R. 1967. The sedimentary and faunal sequence of the Wadhurst Clay (Wealden) in boreholes at Wadhurst Park, Sussex. *Bull. geol. Surv. Gt. Br.* 27, 171–235, 7 pls.
- BALME, B. E. 1957. Spores and pollen grains from the Mesozoic of Western Australia. Rep. Fuel Res. Sect. C.S.I.R.O. Aust. 48 pp., 11 pls.
- BATTEN, D. J. 1968. Probable dispersed spores of Cretaceous *Equisetites*. *Palaeontology*, **11**, 633-642, 1 pl.
- —— 1969. Some British Wealden megaspores and their facies distribution. Ibid. 12, 333–350, 6 pls.
- —— 1972. Recognition of the facies of palynologic assemblages as a basis for improved stratigraphic correlation. *Proc. 24th Int. Geol. Cong.* 7, 367–374.
- —— 1973. Use of palynologic assemblage-types in Wealden correlation. *Palaeontology*, **16**, 1–40, 2 pls. BRICHE, P., DANZÉ-CORSIN, P. and LAVEINE, J. P. 1963. Flore infraliasique du Boulonnais. *Mém. Soc. Géol. N.* **13**, 1–143, 11 pls.
- CHEETHAM, A. H. and HAZEL, J. E. 1969. Binary (presence-absence) similarity coefficients. *J. Paleont.* 43, 1130-1136.
- CHZHAN, CHUN'-BIN 1962. A spore-pollen complex from Lower Cretaceous strata in the Djuijui district, province of Tseiansu. *Acta palaeont. sin.* **10,** 246–286, 6 pls. [In Chinese, Russian summary.]
- EVANS, P. R. 1970. Revision of the miospore genera *Perotrilites* Erdtm. ex Couper 1953 and *Diaphanospora* Balme and Hassell 1962. *Bull. Commonw. Aust. Dept. Nat. Dev. Bur. Min. Res.* 116, 65–74, 3 pls.
- GÓCZÁN, F. 1956. Pollenanalytische (palynologische) Untersuchungen zur Identifizierung der liassischen Schwarzkohlenflöze von Komló. Földt. Evk. 45, 167–212, 20 pls.
- GOUBIN, N. 1965. Description et répartition des principaux pollenites Permiens, Triasiques et Jurassiques des Sondages du bassin de Morondava (Madagascar). Rev. Inst. Fr. Petr. 20, 1415-1461, 8 pls.
- GOULD, R. E. 1968. Morphology of *Equisetum laterale* Phillips 1829, and *E. bryanii* sp. nov. from the Mesozoic of south-eastern Queensland. *Aust. J. Bot.* 16, 153–176, 3 pls.
- HARLAND, W. B. et al. (eds.) 1967. The fossil record. London (Geological Society), xii+828 pp.

HARRIS, T. M. 1953. The geology of the Yorkshire Jurassic flora. Proc. Yorks. geol. Soc. 29, 63-71.

HUGHES, N. F. and MOODY-STUART, J. C. 1967a. Palynological facies and correlation in the English Wealden. *Rev. Palaeobotan. Palynol.* 1, 259–268.

IMBRIE, J. and PURDY, E. G. 1962. Classification of modern Bahamian carbonate sediments. *Mem. Am. Ass. Petrol. Geol.* 1, 253-272.

KARA-MURZA, E. N. 1960. Palynological evidence for the stratigraphical subdivision of Mesozoic deposits of the Khatanga Depression. *Trudy nauchno-issled. Inst. Geol. Arkt.* 109, 134 pp., 22 pls. [In Russian.]

KRUSKAL, J. B. 1964a. Multidimensional scaling by optimizing goodness of fit to a nonmetric hypothesis. *Psychometrika*, **29**, 1–27.

—— 1964b. Nonmetric multidimensional scaling: a numerical method. Ibid. **29**, 115–129.

LANJOUW, J. et al. (eds.) 1966. International Code of Botanical Nomenclature. Utrecht, The Netherlands. LANTZ, J. 1958. Étude palynologique de quelques échantillons Mésozoiques du Dorset (Grande-Bretagne). Rev. Inst. Fr. Petr. 13, 917-942, 7 pls.

LOVERING, T. S. and ENGEL, C. 1967. Translocation of silica and other elements from rock into *Equisetum* and three grasses. *Prof. Pap. U.S. geol. Surv.* **594-B**, 16 pp.

LUGARDON, B. 1969. Sur la structure fine des parois sporales d'*Equisetum maximum* Lamk. *Pollen Spores*, 11, 449-474, 8 pls.

MANTELL, G. A. 1833. The geology of the south-east of England. London. 415 pp.

NAGY, E. 1958. Palynologische Untersuchung der am Füsse des Mâtra-Gebirges gelagerten oberpannonischen Braunkohle. Földt. Evk. 47, 1–352, 29 pls.

NAKOMAN, E. 1964. Étude palynologique de quelques échantillons de lignite provenant du Bassin de Thrace (Turquie). *Annls Soc. géol. N.* **84**, 289–302, 1 pl.

PETTITT, J. M. 1966. Exine structure in some fossil and recent spores and pollen as revealed by light and electron microscopy. *Bull. Br. Mus.* (*Nat. Hist.*), *Geol.* 13, 221–257, 21 pls.

PIEL, K. M. 1971. Palynology of Oligocene sediments from central British Columbia. Can. J. Bot. 49, 1885–1920, 17 pls.

POCOCK, S. A. J. 1970. Palynology of the Jurassic sediments of western Canada, Part 1. Terrestrial species. *Palaeontographica*, **130B**, 12–136, 22 pls.

POKROVSKAJA, I. M. and STEL'MAK, N. K. (eds.) 1964. Atlas of the Lower Cretaceous spore-pollen complexes of certain regions of the U.S.S.R. *Trudy VSEGEI*, N.S. 124, 552 pp., 69 pls. [In Russian.]

ROGALSKA, M. 1954. Spore and pollen analysis of the Liassic coal of Blanowice in Upper Silesia. *Biul. Inst. Geol.* 89, 5-46, 12 pls. [In Polish, English summary.]

—— 1956. Spore and pollen analysis of the Liassic deposits of the Mroczków-Rozwady area in the Opoczno District. Ibid. 104, 5–89, 34 pls. [In Polish, English summary.]

—— 1962. Spore and pollen grain analysis of Jurassic sediments in the northern part of the Cracow-Wieluń Cuesta. *Pr. Inst. Geol.* 30, 495–524, 2 pls. [In Polish, English summary.]

ROWLEY, J. R. 1963. Ubisch body development in *Poa annua*. Grana palynol. 4, 25–36.

SAMOILOVICH, S. R. *et al.* 1961. Pollen and spores of western Siberia; Jurassic to Paleocene. *Trudy VNIGRI*, 177, 659 pp., 84+65 pls. [In Russian.]

SIMONCSICS, P. 1964. Einige neue Sporen aus dem Salgótarjáner Kohlengebiet in Ungarn. Fortschr. Geol. Rheinld Westf. 12, 97–104, 3 pls.

SMITH, A. H. V. and BUTTERWORTH, M. A. 1967. Miospores in the coal seams of the Carboniferous of Great Britain. *Spec. Paper*, *Palaeont*. 1, 324 pp., 27 pls.

SOKAL, R. R. and MICHENER, C. D. 1958. A statistical method for evaluating systematic relationships. *Kans. Univ. Sci. Bull.* **38**, 1409–1438.

— and SNEATH, P. H. A. 1963. *Principles of numerical taxonomy*. San Francisco, Freeman, 359 pp.

VAKHRAMEEV, V. A. 1970. Range and palaeoecology of Mesozoic conifers, the Cheirolepidiaceae. *Paleont. zhur.* 1, 19–34. [In Russian.]

VERBITSKAJA, Z. I. 1962. Palynologic evidence and stratigraphic subdivision of Cretaceous deposits of the Suchansky coal basin. *Trudy Lab. Geol. Uglja*, **15**, 166 pp., 24 pls. [In Russian.]

VOLKHEIMER, W. 1968. Esporas y granos de polen del Jurasico de Neuquen (Republica Argentina). 1. Descripciones sistematicas. *Ameghiniana*, **5**, 333–370, 9 pls.

WALL, D. 1965. Microplankton, pollen, and spores from the Lower Jurassic in Britain. *Micropaleontology*, 11, 151-190, 9 pls.

WHITE, H. J. O. 1928. Geology of the country near Hastings and Dungeness. Mem. geol. Surv. U.K. iv+104 pp., 6 pls.

Typescript received 31 July 1972.



Batten, D. J. 1973. "Palynology of early Cretaceous soil beds and associated strata." *Palaeontology* 16, 399–424.

View This Item Online: <a href="https://www.biodiversitylibrary.org/item/197406">https://www.biodiversitylibrary.org/item/197406</a>

Permalink: <a href="https://www.biodiversitylibrary.org/partpdf/173202">https://www.biodiversitylibrary.org/partpdf/173202</a>

# **Holding Institution**

Smithsonian Libraries and Archives

# Sponsored by

**Biodiversity Heritage Library** 

## **Copyright & Reuse**

Copyright Status: In Copyright. Digitized with the permission of the rights holder.

License: <a href="http://creativecommons.org/licenses/by-nc/3.0/">http://creativecommons.org/licenses/by-nc/3.0/</a></a><a href="https://www.biodiversitylibrary.org/permissions/">https://www.biodiversitylibrary.org/permissions/</a>

This document was created from content at the **Biodiversity Heritage Library**, the world's largest open access digital library for biodiversity literature and archives. Visit BHL at <a href="https://www.biodiversitylibrary.org">https://www.biodiversitylibrary.org</a>.