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- 1 The classic Lower Devonian plant-bearing deposits of northern New Brunswick, eastern
- 2 Canada: dispersed spore taxonomy and biostratigraphy

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- 7

# 8 ABSTRACT

This paper describes dispersed spore assemblages recovered from the Lower Devonian Val 9 10 d'Amour and Campbellton formations exposed along the Restigouche River in northern New Brunswick, eastern Canada. The dispersed spore assemblages were recovered from seven of 11 12 the nine sampled sections and biostratigraphical analysis suggests that these can be assigned to the polygonalis-emsiensis, annulatus-sextantii and douglastownense-eurypterota Spore 13 Assemblage Biozones. This indicates that these strata range in age from Pragian to late 14 15 Emsian [or possibly earliest Eifelian]. The new biostratigraphical control enables accurate stratigraphical correlation of the nine sections and provides important age constraints for the 16 17 famous early land plant megafossil and non-marine invertebrate/vertebrate assemblages recovered from these strata. Two new dispersed spore species are described: Emphanisporites 18 genselae sp. nov. and Emphanisporites morrisae sp. nov. 19

20

21 *Key words:* Pragian; Emsian; Lower Old Red Sandstone.

# **1. Introduction**

25	Among the earliest reports of early land plant fossils are those from the Lower			
26	Devonian deposits exposed along the Restigouche River in northern New Brunswick, eastern			
27	Canada (Dawson, 1882). Subsequent work on these fossiliferous horizons has been critical in			
28	developing our understanding of Lower Devonian plants (Gensel and Andrews, 1984).			
29	However, placing the plants in a secure stratigraphical context has been hampered because			
30	the stratigraphy of the deposits was unclear and they were poorly age-constrained. Recent			
31	work on the stratigraphy, sedimentology and palaeoecology of the deposits (Kennedy and			
32	Gibling, 2011; Kennedy et al., 2012a,b, 2013), coupled with analysis of their dispersed			
33	spore/pollen assemblages (this work), now enables these plant communities to be considered			
34	within a rigid stratigraphical/biostratigraphical framework.			
35				
36	2. Geological setting			
	2. Geological setting			
36	2. Geological setting A narrow belt of Lower Devonian strata belonging to the Campbellton Formation			
36 37				
36 37 38	A narrow belt of Lower Devonian strata belonging to the Campbellton Formation			
36 37 38 39	A narrow belt of Lower Devonian strata belonging to the Campbellton Formation crops out along the southern shore of the Restigouche River in northern New Brunswick (Fig.			
36 37 38 39 40	A narrow belt of Lower Devonian strata belonging to the Campbellton Formation crops out along the southern shore of the Restigouche River in northern New Brunswick (Fig. 1). To the south of it lies a thicker development of volcaniclastic deposits belonging to the			
36 37 38 39 40 41	A narrow belt of Lower Devonian strata belonging to the Campbellton Formation crops out along the southern shore of the Restigouche River in northern New Brunswick (Fig. 1). To the south of it lies a thicker development of volcaniclastic deposits belonging to the Val d'Amour Formation (Fig. 1). The stratigraphy of the Campbellton Formation, and its			

45 famous due to the rich terrestrial-freshwater biotas they yield, including terrestrial plants

46 (Gensel and Andrews, 1984) and arthropods (Shear et al. 1996), and freshwater aquatic
47 invertebrates (e.g. Miller, 2007) and fish (e.g. Miller et al., 2003). Recently, Kennedy and
48 Gibling (2011) undertook extensive logging/mapping of the formations that has led to a much
49 better understanding of their sedimentology and stratigraphical relationships (Figs 2-3).

Regionally these deposits form part of the Gaspé Belt whose genesis and history are 50 intimately associated with the Acadian Orogeny. They form the final terrestrial phase of basin 51 52 infilling following significant uplift. The Val d'Amour Formation comprises volcaniclastic deposits deposited in a shallow marine environment to the east and subaerially to the west. 53 Following a period of uplift, and possible unconformity, the terrestrial fluviatile deposits of 54 55 the Campbellton Formation were deposited in an intermontane basin surrounded by a rugged mountainous landscape (Wilson et al., 2004, 2005). Deposition took place at approximately 56 35°S on the southeast margin of Euramerica near the margin between the arid and warm 57 58 temperate climate belts.

Radiometric dating of rhyolites near the top of the Val d'Amour Formation give an age of 407.4 +/- 0.8 Ma which indicates a late Pragian age (Wilson et al., 2005). Colin McGregor studied dispersed spore assemblages from the Campbellton Formation but his findings are included in unpublished Geological Survey of Canada reports and published only as personal communications in various papers reporting on the biotas of the deposits. A discussion of this work is included within the descriptions of the various sections examined during the study reported herein (see below).

The boundary between the Val d'Amour and Campbellton formations is difficult to
interpret. This is because sedimentary horizons are present in the Val d'Amour Formation
volcanics, but towards the boundary with the Campbellton Formation these become more
common and sedimentary and volcanic units are intercalated. In places there is clear evidence

of sediments overlying volcanic rocks with angular unconformity. However, this may be very
local. In other places there is clear interdigitation with sediments and lavas/ash falls.

Kennedy and Gibling (2011) undertook detailed bed-by-bed logging of all of the 72 73 available coastal exposure of the Campbellton Formation and attempted to produce a composite log from six logged sections (Fig. 3). The eastern outcrop includes a 4 km long 74 coastal exposure that follows strike. Kennedy and Gibling logged 938 m of strata (sections III 75 76 to VI). Two logs (sections I and II) were undertaken in the western outcrop encompassing 82 m of strata. These strata belong to the basal part of the formation and were considered 77 equivalent to the lower strata logged in section III of the eastern outcrop. Kennedy and 78 79 Gibling subdivided the strata into six facies associations: (1) restricted lacustrine; (2) marginal lacustrine; (3) near-shore lacustrine; (4) coastal-deltaic; (5) sandy to gravelly 80 alluvial plain; (6) gravelly proximal alluvial environments. Building on this stratigraphical 81 82 and sedimentological framework Kennedy et al. (2012a,b) undertook a detailed analysis of the palaeoenvironments and biota of the deposits. 83

84

# 85 **3. Materials and methods**

86

Samples for palynological analysis were collected from nine sections (A-I) from
throughout the sequence during fieldwork in 1998, 2007, 2013 and 2016. Additional samples
from specific plant beds were supplied by Pat Gensel. Details of all analysed samples are
provided in Appendix 1 and their geographical and stratigraphical position illustrated in Figs
1 and 3, respectively. Twenty grams of each rock samples was processed using standard
HCl/HF/HCl palynological acid maceration techniques followed by heavy liquid separation

93	using $ZnCl_2$ . Sieving was undertaken using a 20 $\mu$ m mesh. Most of the residues are of low			
94	thermal maturity and did not require oxidation. However, some samples from adjacent to			
95	igneous intrusions or extrusions were coalified and oxidised for between 10 and 20 minutes			
96	using Schultz solution. Residues were strew mounted and attached to slides using Petropox			
97	resin. All materials (rock, residue, slides) are housed in the collections of the Centre for			
98	Palynology of the University of Sheffield. Slides of each sample were scanned using a light			
99	microscope and all spore taxa present identified (largely using the spore taxonomy			
100	established by McGregor (1973, 1977) in his classic monograph on the spores from coeva			
101	correlative strata at nearby Gaspé). A semi-quantitative count of 200 fossil spores was			
102	undertaken on each productive sample.			
103				
104	4. Systematic palaeontology			
104	4. Systematic paraeontology			
105				
106	4.1. Introduction			
107	All of the taxa encountered in this study are listed in Table 1 and figured (Plates I-			
108	VIII). Most of these taxa are well known and described in detail in the monograph of			
109	dispersed spores from coeval strata from nearby Gaspé (McGregor 1973, 1977). Two new			
110	spore taxa were discovered during the course of this investigation and are described in the			
111	brief taxonomic section that follows.			
112				
113	4.2. Description of new taxa			

115	Genus <i>Emphanisporites</i> McGregor 1961	
116	Type species: Emphanisporites rotatus McGregor 1961	
117		
118	Emphanisporites genselae sp. nov. (Plate V, 5-8)	
119		
120	Holotype: Plate V, fig. 5, Sample BNB29/1, E.F.No. (K41/3).	
121		
122	Derivation of name: after Pat Gensel who provided samples used in this study and invaluable	
123	guidance during field work on the Gaspé and New Brunswick Lower Devonian sequences.	
124		
125	Diagnosis: An Emphanisporites characterised by a prominent, wide equatorial crassitude and	
126	proximal face ornamented by radial ribs, which are highly variable in width and often divide	
127	and cover most of the interradial area.	
128		
129	Description: Amb subcircular. Trilete mark distinct. Suturae straight and simple extending to	

129 near the spore equator. Equatorial crassitude up to 5 µm wide and very distinct, apparently 130 131 with a groove on the distal surface demarcating the inner margin of the crassitude. Contact areas ornamented with highly variable, more-or-less radially arranged, muri. The muri are of 132 highly variable width and frequently coalesce in a general direction towards the proximal 133 pole. This leaves irregular, sinuous gaps between muri of  $<1 \,\mu$ m. The muri finish short of the 134 trilete mark leaving the sutures in a distinct topographic low (indentation). Due to their 135

136	irregular nature they end near the sutures along their entire length (i.e. they do not all		
137	converge on the proximal pole). Distal surface entirely laevigate.		
138			
139	Dimensions: 42(45)46 µm (5 specimens measured).		
140			
141	Occurrence: Present in two samples from the Val d'Amour Formation of New Brunswick,		
142	Canada (Pragian-?earliest Emsian polygonalis-emsiensis Spore assemblage Biozone) (Table		
143	1).		
144			
145	Comparison and remarks: The highly irregular nature of the radially arranged proximal muri		
146	distinguish this taxon from previously described species of Emphanisporites (reviewed in		
147	Taylor et al. 2011).		
148			
149	Emphanisporites morrisae sp. nov. (Plate V, 12-14)		
150			
151	Holotype: Plate V, fig. 12, Sample BNB33/1, E.F.No. (N43).		
152			
153	Derivation of name: after Jenny Morris who provided invaluable assistance in the field		
154	working on the Gaspé and New Brunswick Lower Devonian sequences.		
155			

Diagnosis: An Emphanisporites characterised by its relatively large size and large number of
robust radially-arranged proximal muri.

159	Description: Amb subcircular to subtriangular. Trilete mark indistinct. In some specimens		
160	straight suturae, accompanied by lips ca. 1 $\mu$ m wide, extend from the proximal pole to the		
161	equator. In other specimens the suturae appear to be gaping resulting in a triangular section of		
162	the proximal surface, centred on the proximal pole, that is missing. Equatorial crassitude		
163	distinct and up to 2 $\mu$ m wide. Contact areas ornamented with a large number (up to 40) of		
164	crowded, radially-arranged muri. Muri robust, thinning from the equatorial margin towards		
165	the proximal pole, from 4 to 1 $\mu$ m in width. Muri occasionally bifurcating. Distal surface		
166	entirely laevigate.		
167			
168	Dimensions: 59(73)107 µm (21 specimens measured).		

170 Occurrence: Present in samples from the Val d'Amour and Campbellton formations of New
171 Brunswick, Canada (Pragian-?earliest Emsian *polygonalis-emsiensis* Spore assemblage
172 Biozone) (Table 1).

*Comparison and remarks:* The relatively large size and large number of prominent radially175 arranged proximal muri distinguish this taxon from previously described species of
176 *Emphanisporites* (reviewed in Taylor et al. 2011).

#### 180 *4.3.1. Laevigate retusoid trilete spores*

181 McGregor (1973, 1977) subdivided the laevigate retusoid spores he encountered in Gaspé into several genera (Calamospora, Retusotriletes, Deltoidospora) and numerous 182 species. The criteria used to distinguish these taxa were rather subtle, and although we could 183 identify most of them, it was decided to group them together as Retusotriletes spp., with 184 positive identification of only the most distinctive taxa (R. triangulatus, R. cf. rotundus, R. 185 186 eslae). This is because these taxa are morphologically similar and also because they are not biostratigraphically significant. They are, however, extremely common and comprise up to 187 90% of spores in the assemblages studied. Plate I figures a selection of laevigate retusoid 188 189 spores including some of the more distinctive and common taxa encountered.

190

#### 191 *4.3.2. Ornamented retusoid trilete spores*

Apiculiretusispora spp. is abundant in all of the samples studied and is highly
distinctive as it is bilayered with the outer ornamented layer often in the process of separating
from the inner body and sloughing off. McGregor (1973, 1977) described six species of this
genus from Gaspé (as well as similar spores he placed with *Apiculatisporis microconus*Richardson 1965, *Apiculatasporites perpusillus* (Naumova) McGregor 1973 and *Anapiculatisporites* sp.). In this study only four of these taxa were recognised (see Table 1
and Plate I).

Of the five *Dibolisporites* species recognised by McGregor (1973, 1977) from Gaspé
all are present in the New Brunswick sequence except for *D. variegatus* McGregor 1973 (see

Table 1 and Plate II). *Dibolisporites* spp. occurs in all samples and is a persistent but lowabundance component.

All five *Dictyotriletes* species recognised by McGregor (1973, 1977) in Gaspé are
 present in the New Brunswick sequence (see Table 1 and Plate II).

205

## 206 *4.3.3. Laevigate crassitate trilete spores*

Simple laevigate crassitate spores are present in most samples and are referred to as *Ambitisporites* spp. (Plate III, fig. 1). They become less common in increasingly younger
spore zones. Similar forms from Gaspé were described by McGregor (1973, 1977) as species
of *Punctatisporites*.

211

#### 212 *4.3.4. Ornamented crassitate trilete spores*

213 Various ornamented crassitate trilete spores were encountered (Table 1). These include forms assigned to Synorisporites (Plate III, figs 4-5,9) those with an ornament of 214 elements with a spatulate tip (Raistrickia sp.) (Plate III, figs 6-7) and rare but distinctive taxa 215 216 with a distal ornament of anastomosing muri (Plate III, figs 2-3) or an annulus (Amicosporites jonkeri) (Plate III, fig. 8). These all occurred sporadically and in low numbers (Table 1). 217 218 Forms with a more prominent equatorial crassitude, verging on patinate structure, were included in Lycopodiacidites ogygius, Verruciretusispora dubia, Verruciretusispora 219 multituburculata, Verrucosisporites devonicus, Verrucosisporites polygonalis? and 220 Verrucosisporites sp. A. (see Table 1 and Plates III-IV). 221

Other ornamented crassitate spores include five species of *Brochotriletes* (Plate IV), seven species of *Emphanisporites* (Plate V) and *Camarozonotriletes sextantii* (Plate IV, figs 7-8). The latter also occurs within a large laevigate body with a trilete mark (Plate IV, fig.14) as previously reported among *in situ* spores recovered from the plant *Chaleuria cirrosa* (Andrews et al. 1974).

227

### 228 *4.3.5. Patinate trilete spores*

Laevigate patinate spores are represented by the taxa *Archaeozonotriletes chulus*(Plate VI, fig. 2) and ?*Archaeozonotriletes* sp. of McGregor 1973 (Plate VI, fig. 1).
Ornamented forms are notably rare with only very rare representatives of the genera *Cymbosporites* and *Chelinospora* (see Table 1 and Plate VI).

233

### 234 *4.3.6. Trilete spores with multiple wall layers*

Structurally more complex spores, with distinct inner bodies +/- zona, include 235 236 Acinosporites lindlarensis (Plate VI, figs 4,7,11), Ancyrospora loganii (Plate VIII, figs 5-7), Camptozonotriletes caperatus (Plate VII, figs 5,8), Grandispora douglastownensis (Plate 237 VIII, figs 1-4), Zonotriletes brevivelatus (Plate VII, figs 1,4,7,10), and other rare forms (Plate 238 VII, figs 2-3,6). A. lindlarensis is highly variable in terms of size and ornament (See 239 Richardson et al. 1993) as has previously been noted based on studies of in situ spores 240 (Richardson et al. 1993; Gensel and Kasper 2005; Gensel and Albright 2006; Wellman et al. 241 2009). Grandispora douglastownensis is highly variable regarding ornament (Plate VIII, figs 242 1-4) but this is considered to represent intraspecific variation within a single taxon (see 243 discussion in Wellman and Gensel 2004). 244

# 246 4.3.7. Cryptospores

247	Cryptospores are very rare in the assemblages and include the pseudodyad	
248	Pseudosyadospora petasus (Plate VII, fig. 9) and permanent tetrad Tetrahedraletes	
249	medinensis. No true dyads (or hilate cryptospores dissociated from them) were reported.	
250		
251	5. Biostratigraphical analysis	
252		
253	5.1. Introduction	
254	Nine sections (A - I) were sampled (Figs 1-3). Most of the samples yielded	
255	assemblages containing abundant and well preserved spores that were either of low thermal	
256	maturity or cleared easily using Schultz solution. Details of all samples, including those that	
257	were barren, are provided in Appendix 1. The best preserved spore assemblages from each of	
258	the nine sampled sections $(A - I)$ were logged and the distribution of recorded taxa is	
259	reported in Table 2. Fig. 4 is a summary diagram indicating the spore zonation, ages and	
260	correlation of the studied sections.	

Previous palynological investigation of these deposits has been reported on by Colin McGregor and Elliot Burden. McGregor's work consists of unpublished Geological Survey of Canada reports and personal communications. These are quoted in various papers concerning descriptions of the palaeontology (e.g. Gensel et al. 1991) and stratigraphy of these beds (e.g. Kennedy and Gibling 2011) and are outlined in the discussion below. The spore assemblages are correlated using the spore zonation scheme of McGregor and Camfield

267	(1976) and McGregor (1977). It should be noted that this differs somewhat from the later
268	spore zonation of Richardson and McGregor (1986) and cross correlation is difficult (see
269	Figure 1 (insert) of Richardson and McGregor 1986). Burden's spore-based biostratigraphical
270	interpretations are discussed in Wilson et al. (2004, 2005).

## 272 5.2. Section A (Val d'Amour Formation)

Wilson et al. (2004, 2005) mapped an outcrop of the Val d'Amour Formation around 273 Dalhousie. It is exposed in coastal sections south and west of Dalhousie, and in the latter it is 274 275 unconformably overlain by the Campbellton Formation (section III of Kennedy & Gibling 2011). Two samples yielding palynomorphs were recovered in the section south of Dalhousie 276 (Walker Brook Member in Wilson et al. 2005). Sample BNB30 is from a small pocket of 277 278 sediments intercalated between lavas, with a notable accumulation of pillow lavas stratigraphically above. Sample BNB29 occurs higher in the sequence and is from an 279 exposure of highly fossiliferous sediments rich in marine invertebrate remains. Both samples 280 contain abundant marine and non-marine palynomorphs. The marine palynomorphs consist of 281 diverse assemblages of acritarchs, chitinozoans and scolecodonts, in addition to large 282 283 fragments of arthropod cuticle. The non-marine palynomorphs consist of abundant, but not diverse, spore assemblages in addition to abundant large fragments of plant debris (including 284 285 banded tubes and cuticles). The dispersed spore assemblages are virtually identical in both 286 samples (Table 2) and are placed with the PE Spore Assemblage Biozone of Richardson & 287 McGregor (1986), based on the presence of the nominal species of the zones and the overall characteristics of the spore assemblages, indicating a Pragian-?earliest Emsian age. This zone 288 289 equates to the PoW Oppel Zone of Streel et al. (1987). The presence of D. wetteldorfensis may further constrain the assemblages to the W Interval Zone of Streel et al. (1987). The 290

presence of *Acinosporites lindlarensis* in sample BNB30 is interesting as this occurrenceappears to be earlier than expected.

293

294 5.3. Section B (Val d'Amour Formation)

There is an excellent section of the Val d'Amour Formation exposed in the road cut 295 either side of Route 11 directly west of Sugarloaf Mountain (logged and described by 296 297 Kennedy et al. 2013). The strata are more-or-less perpendicular and strike parallel to the road. Moving west along the road, from its junction with Route 270, in the cut on the south side of 298 299 the road a volcaniclastic sequence is dominated by tuffs. The first significant development of sediments is at N47°59.177 / W066°042.027 and consists of an approximately 5m thick 300 sequence from which 4 samples were collected (BNB34 - coal; BNB33 - siltstone with plant 301 302 fragments; BNB 32 – coaly horizon; BNB 31 – siltstone). Further west, on the north side of the road cut at N47°59.142 / W066°42.096, further sediments are exposed from which 5 303 samples were collected (BNB16 - coal; BNB17 - siltstone; BNB18 - siltstone with 304 fragments of the plant Taeniocrada (see also Kennedy et al. 2013 for a discussion of plant 305 fragments recovered from this section); BNB 19 directly above BNB18; BNB20 directly 306 307 below BNB18). Slightly further west Sample BNB21 (dark organic rich shale) was collected from the sediments that unconformably overlie a hummocky volcanic surface well exposed in 308 309 the road cut. A similar horizon was collected (BNB22) on the south side of the road cut.

The samples all yield essentially the same spore assemblage (Table 2). It is dominated by simple *Ambitisporites*, *Retusotriletes*, *Apiculiretusispora* and *Dibolisporites* but with rare more elaborate spores. The spore assemblage can be equated with the PE Spore Assemblage Biozone of Richardson and McGregor (1986) indicating an early (but not earliest) Pragian -?earliest Emsian age. It contains both of the eponymous species and numerous other species 315 characteristic of this zone (e.g. *Brochotriletes foveolatus*?, *Dictyotriletes favosus*,

316 *Clivosispora verrucata*). This zone equates to the PoW Oppel Zone of Streel et al. (1987).

The presence of *Dictyotriletes subgranifer* may restrict the assemblage to the Su Interval Zone of Streel et al. (1987) constraining the age to late Pragian - ?earliest Emsian. All of the samples yield only spores and plant fragments with no marine elements suggesting deposition in a subaerial environment.

321

## 322 5.4. Section C (Val d'Amour Formation / Campbellton Formation contact)

There is an excellent section purportedly representing the contact between the Val 323 D'Amour and Campbellton formations exposed in the roadcut of Route 134 Beauvista Street 324 that heads northwest from Sugarloaf Mountain towards the Restigouche River. The roadcut 325 326 exposes mainly tuffs, but contains a thin sedimentary sequence that Miller (2007, p.983) considers to "...sit at the base of the Campbellton Formation in contact with the Val d'Amour 327 Formation Rhyolite.". It is from this section that Doran (1980) described specimens of the 328 trimerophyte plant *Psilophyton crenulatum* exceptionally preserved by a fall of volcanic ash 329 (plant locality Q of Gensel & Andrews (1984) and Gensel et al. (1991)). McGregor (in Doran 330 331 1980) reported on a spore assemblage recovered from the sediments. He equated the spore assemblages with the CE and AL(S) zones of McGregor (1977). This was based on the 332 333 presence of Brochotriletes hudsonii, Camptozonotriletes caperatus, Dictyotriletes canadensis, D. emsiensis, Emphanisporites erraticus, E. schultzii and ?Enigmophytospora 334 335 simplex. A sample was collected during this study (Appendix 1) and based on the spores present (Table 2) it is assigned to the AS Spore Assemblage Biozone of Richardson and 336 337 McGregor (1986) which is equivalent to the AB Oppel Zone of Streel et al. (1987). These designations were based on the presence of all of the nominal species of the zones and also 338

339 general consideration of the composition of the assemblage. This suggests an early Emsian

340 age. Only terrestrial palynomorphs were recovered suggesting that the sediments

accumulated as terrestrial fluvial-lacustrine deposits between ash falls directly onto the landsurface.

343

344 5.5. Section D (Campbellton Formation Western Belt sections I & II sensu Kennedy &
345 Gibling 2011)

The Western Belt of the Campbellton Formation includes Sections I and II of Kennedy & 346 347 Gibling (2011). It contains a basal contact between Val d'Amour Formation rhyolites and basal sediments of the Campbellton Formation as described in Wilson et al. (2005) and 348 Kennedy and Gibling (2011, p. 1574). A radiometric age of 407.4 (+/-0.8) Ma has been 349 350 obtained from the rhyolites (Wilson et al. 2004, 2005). Gensel and Albright (2006) note that spore assemblages recovered from this sequence belong to the AS Spore Assemblage 351 Biozone of Richardson & McGregor (1986) and "The presence of Calyptosporites 352 heisdorfensis indicates an age no older than late Emsian (McGregor, written communication 353 to McCutcheon 1996)". 354

A rich invertebrate and fish fauna occurs in these beds (Whiteaves 1881; Jones 1889; 355 Woodward 1892; Traquair 1893; Miller et al. 2003; Miller 2007). Kennedy and Gibling 356 (2011) interpreted Section I as a coastal deltaic sequence containing both aquatic fauna and 357 plant remains but noted that it is difficult to decide if the delta was feeding into a lacustrine, 358 359 brackish or marine environment. Arguments have previously been put forward for the nonmarine affinities of various fish (Kennedy and Gibling 2011) and invertebrates (Morris 1985; 360 Gray 1988) recovered from these beds. However, both Blieck and Cloutier (2000) and 361 McGregor (in Gamba 1990) report the presence of marine acritarchs and prasinophcean cysts. 362

Section II is interpreted as fluvial based on sedimentological grounds (Kennedy and Gibling
2011). Plant locality 3 of Grierson and Hueber (1968) (GH3) occurs low in section I (Figs 34).

Samples BNB36-41 are from Section I. BNB36b is the lowest in the sequence and is 366 at the unconformity where a mudstone fills a fissure in the Val d'Amour Formation rhyolites. 367 BNB36a is from indurated sediments directly above this and represents a fish bed. This 368 locality was termed the "Athoville Section" by Miller (2007). Higher in the sequence samples 369 BNB37 and BNB38 are mudstones containing ostracods and sample BNB39 is a mudstone 370 slightly higher in the sequence. Samples BNB40-41 are from another fish/eurypterid bed 371 372 (Miller 2007). BNB40 is from the actual fish/eurypterid bed and is extremely hard. Sample BNB41 is from directly below this and is a softer plant hash bed. Sample BNB35 is from east 373 and belongs to Section II and probably represents the highest beds exposed in the sequence. 374

The samples all yield essentially the same spore assemblage (Table 2). It is dominated 375 by simple Retusotriletes, Apiculiretusispora and Dibolisporites but with rare more elaborate 376 377 spores. The spore assemblage can be equated with the AS Spore Assemblage Biozone of Richardson and McGregor (1986) indicating an early - mid Emsian age. It contains both of 378 379 the eponymous species and numerous other species characteristic of this zone (e.g. 380 Acinosporites lindlarensis, Dictyotriletes canadensis, Emphanisporites schultzii). The assemblage can be correlated with the AB Oppel Zone of Streel et al. (1987) suggesting a 381 more restricted early Emsian age. Marine palynomorphs, in the form of acritarchs, are 382 relatively common in samples BNB40, BNB41 and BNB35 suggesting some marine 383 384 influence higher in the sequence.

5.6. Section E (Campbellton Formation Eastern Belt Section III sensu Kennedy & Gibling
2011)

Kennedy and Gibbling (2011)'s section III is a 250m+ section with sediment 388 intermittently exposed that covers the eastern end of the Eastern Belt of the Campbellton 389 Formation. Productive samples in this study are currently restricted to strata in the lower part 390 of the sequence around plant localities N and M of Gensel and Andrews (1984) and Gensel et 391 392 al. (1991) that include the type localities for *Sawdonia acanthotheca* (Gensel et al. 1975) and Zosterophyllum divaricatum (Gensel 1982a). A total of 6 samples were collected, from a 393 variety of lithologies, but all yield essentially the same dispersed spore assemblage (Table 2). 394 395 Previously, for plant locality M Gensel (1982a) reported McGregor pers. com. as suggesting that palynological analysis indicates an Emsian age and Gensel (1991) noted that D. C. 396 McGregor, unpublished Geological Survey of Canada Report #F1-6-1979 suggested an early 397 398 Emsian age. Gensel and Andrews (1984) and Gensel et al. (1991) indicated that the assemblage belonged to the PE Spore Assemblage Biozone of Richardson and McGregor 399 400 (1986). We confirm this designation that suggests an early (but not earliest) Pragian -401 ?earliest Emsian age. The assemblage is dominated by simple Ambitisporites, Retusotriletes, 402 Apiculi retusispora and Dibolisporites but with rare more elaborate spores. It contains both of 403 the eponymous species of the zone and numerous other species characteristic of this zone (e.g. Brochotriletes foveolatus?, Camptozonotriletes caperatus, Dictyotriletes favosus, 404 *Clivosispora verrucata*). The zone equates to the PoW Oppel Zone of Streel et al. (1987). The 405 406 presence of *Dictyotriletes subgranifer* may restrict the assemblage to the Su Interval Zone constraining the age to late Pragian - ?earliest Emsian. All of the samples yield only spores 407 and plant fragments with no marine elements. 408

409 Plant localities higher in the sequence (K and L in Andrews and Gensel 1984 and410 Gensel et al. 1991) are probably younger as they yield spore assemblages considered to

411 belong to the AL(S) zone of McGregor (1977) (Andrews & Gensel 1984; Gensel et al. 1991;

412 Kennedy et al. 2013). Li et al. (2000) reported that spore assemblages belonging to the AS

413 Spore Assemblage Biozone were recovered from this higher sequence in the section.

414 Unfortunately this study failed to recover productive palynological samples from this part of415 the section.

416

417 5.7. Section F (Campbellton Formation Eastern Belt Section IV lower part sensu Kennedy &
418 Gibling 2011)

Kennedy and Gibbling (2011)'s section IV covers the western end of the Eastern Belt
of the Campbellton Formation and essentially consists of two separated sections: a lower 28
m section and an upper18 m section separated by ca. 126 m of section. We subdivided these
into Section F (lower) and Section G (upper).

423 Section G is very rich in plant beds including plant localities F, E, A, G and H of Gensel and Andrews (1984) and Gensel et al. (1991). Gensel (1982a) reported that dispersed 424 spore assemblages belonging to the lower *Grandispora* subzone of McGregor (1977) 425 indicated a late Emsian age for plant locality A (D. C. McGregor, unpublished Geological 426 Survey of Canada Report #F1-6-1979). Gensel (1982b) reported that dispersed spore 427 assemblages belonging to the Grandispora subzone of McGregor (1973) indicated a late but 428 not latest Emsian age for plant locality E (D. C. McGregor, unpublished Geological Survey 429 of Canada Report F1-11-1981). Gensel and Albright (2006) suggested that plant locality F 430 431 belonged to the AS Spore Assemblage Biozone of Richardson and McGregor (1986). Jensen and Gensel (2013) noted that McGregor, in a 1979 pers. comm. to Gensel, suggested that the 432 433 three plant localities A, G and H all belonged to the lower Grandispora subzone of McGregor (1977). 434

435 Samples were collected from throughout section F including plant beds E and A (Appendix 1). All of the samples yield essentially the same dispersed spore assemblage 436 (Table 2). This can be equated with the AS Spore Assemblage Biozone of Richardson and 437 438 McGregor (1986) indicating an early - mid Emsian age. It contains both of the eponymous species and other species characteristic of this zone (e.g. Acinosporites lindlarensis and 439 Emphanisporites schultzii). Convincing examples of Grandispora douglastownense are not 440 441 present in the samples studied precluding assignment to the overlying DE Spore Assemblage Biozone of Richardson & McGregor (1986). The spore assemblage can be equated to the AB 442 443 Oppel Zone of Streel et al. (1987) suggesting a more restricted early Emsian age (Fig. 4). Marine palynomorphs, in the form of acritarchs, occur rarely in sample BNB12 high in the 444 section. 445

446

5.8. Section G (Campbellton Formation Eastern Belt Section IV upper part sensu Kennedy &
Gibling 2011)

Section G is from the upper18 m of Kennedy and Gibbling (2011)'s section IV (see 449 above). It contains the important plant localities B and I of Gensel and Andrews (1984) and 450 451 Gensel et al. (1991). Andrews et al. (1975) reported that McGregor pers. com. suggested plant locality B was late Emsian (uppermost Lower Devonian) based on spore studies. 452 453 Subsequently, Gensel (1982a) noted that McGregor, unpublished Geological Survey of Canada Report #F1-6-1979, suggested a late Emsian age. Gensel and Andrews (1984) and 454 455 Gensel et al. (1991) placed plant locality B as the youngest in the Campbellton Formation and belonging to the Grandispora subzone of the annulatus-lindlarensis zone of McGregor 456 457 (1977). Gensel and Kasper (2005) reported McGregor (pers. comm. 1975) as stating they belong to the Grandispora subzone of the annulatus-lindlarensis zone of McGregor (1977) 458

which is equivalent to high in the AS Spore Assemblage Zone of Richardson & McGregor(1986).

Samples were collected from throughout section G including plant beds I and B 461 (Appendix 1). All of the samples yield essentially the same dispersed spore assemblage 462 (Table 2). They belong to the DE Spore Assemblage Biozone of Richardson and McGregor 463 (1986) suggesting a late Emsian-?earliest Eifelian age. The assemblage is dominated by 464 465 simple Retusotriletes, Apiculiretusispora and Dibolisporites but with rare more elaborate spores. It contains one of the eponymous species of the zone (*Grandispora douglastownense*) 466 and other species characteristic of this zone (e.g. Ancyrospora loganii). The zone equates to 467 the Cor and Pro (but not Vel) Interval zones of the AP Oppel Zone of Streel et al. (1987) 468 (Fig. 4). Unfortunately none of the taxa that designate Interval Zones are present in the 469 Canadian spore assemblages so further subdivision is not possible. Marine palynomorphs, in 470 471 the form of acritarchs, occur rarely in sample BNB48, BNB15, BNB47 suggesting possible marine influence (or reworking). 472

473

474 5.9. Section H (Campbellton Formation Eastern Belt Section V sensu Kennedy & Gibling
475 2011)

476 No productive samples were recovered from these strata (Appendix 1).

477

5.10. Section I (Campbellton Formation Eastern Belt Section VI sensu Kennedy & Gibling
2011)

480 No productive samples were recovered from these strata (Appendix 1).

#### 482 6. Discussion

483

484 Biostratigraphical analysis of dispersed spore assemblages from the best exposed sections of the Val d'Amour and Campbellton formations of northern New Brunswick enable 485 their age dating and correlation (Fig. 4). It is evident that there is a complex relationship 486 regarding the boundary between the two formations. The Val d'Amour Formation is 487 predominantly volcanic consisting of lavas and ash falls. However, sedimentary intercalations 488 489 are present throughout. In the east (Section A) they are marine and older (PE SAB/PoW OZ[?W IZ]). In the west (Section B) they are terrestrial and slightly younger (PE SAB/PoW 490 OZ[Su IZ]). Situating the actual boundary between the two formations is problematic as it 491 492 would appear that sedimentary intercalations increase in frequency but volcanic activity persists. Thus it is impossible to draw a boundary between a solely/predominantly volcanic 493 and a solely/predominantly sedimentary formation. In the west we have Section C, with 494 sediments of (AS SAB/AB OZ) age intercalated among volcanic rocks, in very close vicinity 495 to basal sediments of Section D, apparently unconformably overlying a volcanic sequence of 496 497 exactly the same spore zone and age. In contrast, however, in the east the thick unit of basal sediments of Section E, that overlies a predominantly volcanic sequence, is older and of (PE 498 499 SAB/PoW OZ[Su IZ]) age. This sedimentary sequence is relatively thick and possibly 500 extends up into the succeeding spore zone (AS SAB/AB OZ) but without the high proportion 501 of associated volcanic rocks as seen in the west in Section C.

Higher sediments in the sequence, which clearly belong to the Campbellton
Formation rather than Val d'Amour Formation, are demonstrably younger. However, their
relationships are not as clear cut as previously considered. To the west Sections F and G,

previously considered to represent a continuous sequence, are shown to differ in age. Section
F is older and of (AS SAB/AB OZ) age. Section G is young and of (DE SAB/AP OZ pars
[Cor and Pro IZ]) age. This suggests a significant gap in the sequence between Sections F and
G (although the magnitude of this gap may appear exaggerated because of the absence of
critical taxa used to identify Oppel and Interval zones in the scheme of Streel et al. 1987).
Unfortunately palynomorph assemblages have not been recovered from Section H and I to
the east.

The new age dates and stratigraphical correlations have important implications regarding the ages of the important terrestrial plant/animal and freshwater aquatic invertebrate/vertebrate fossil assemblages described from these sequences. Details of the plant assemblages is summarised in Tables 3-4 with their stratigraphical relationships illustrated in Fig. 4. It is noteworthy that the numerous plant localities belong to three different spore zones.

The oldest plant localities (N and M=?H1) belong to the PE SAB (Fig. 4). *Sawdonia acanthotheca* and *Zosterophyllum divaricatum* are exclusive this zone (Table 4). The spore assemblages are dominated by *Retusotriletes* spp. (Table 2) that are produced by these zosterophylls (Table 3).

At least 9 plant assemblages belonging to the AS SAB occur in four of the sections (C,D,E,F) (Fig. 4). They yield rich plant assemblages containing rhyniophytes, zosterophylls, lycopsids, trimerophytes and various plants of uncertain affinity (*Bitelaria, Chaleuria, Loganophyton*). The spores *Retusotriletes* spp. and *Apiculiretusisporites* spp., known to be produced by zosterophylls and trimerophytes (Table 3), dominate these assemblages. The spores of *Chaleuria* (*Camarozonotriletes sextantii*) and *Leclercqia* (*Acinosporites lindlarensis*) are locally abundant.

529	Plant assemblages I and B, from section G, belong to the DE SAB (Fig. 4). They yield
530	rich plant assemblages containing rhyniophytes, zosterophylls, lycopsids, trimerophytes and
531	the more complex/advanced plant of uncertain affinity Oocampsa (Andrews et al. 1975)
532	(Table 4). The spores Retusotriletes spp. and Apiculiretusisporites spp., known to be
533	produced by zosterophylls and trimerophytes (Table 3), still dominate these assemblages. The
534	spores of Oocampsa (Grandispora douglastownense) appear for the first time and are
535	relatively abundant. Other more complex spores, such as Ancyrospora loganii with its
536	grapnel-tipped processes, appear for the first time. Spores bearing grapnel-tipped processes
537	are most likely produced by lycopsids (Wellman 2002).

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547

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704	
705	FIGURE CAPTIONS
706	
707	Fig. 1. Location map of the study area showing the distribution of the geological formations
708	and the location of the studied sections (A-I).
709	
710	Fig. 2. Stratigraphical sequence in the region illustrating the relationship of the sections
711	studied (A-I). The location of a radiometric age date is indicated with an asterisk.
712	
713	Fig. 3. Details of the stratigraphic sequences (modified from Kennedy and Gibling 2012).
714	
715	Fig. 4. Summary diagram indicating the spore zonation, ages and correlation of the studied
716	sections (A-I).
717	
718	
719	PLATE CAPTIONS
720	

721	Plate I. Dispersed spores from the Lower Devonian Val d'Amour and Campbellton			
722	formations of New Brunswick (eastern Canada). All spores at magnification x1000. Taxon			
723	name followed by sample/slide number and England Finder co-ordinates.			
724	1. Retusotriletes sp. [BNB11/1 (Q44)].			
725	2. Retusotriletes triangulatus (Streel) Streel 1967 [BNB45/1 (G46/4)].			
726	3. Retusotriletes cf. rotundus (Streel) Streel 1967 [BNB47/1 (T27)].			
727	4. Retusotriletes eslae Cramer and Diez 1975 [BNB44/1 (O31/1)].			
728	5. Retusotriletes eslae Cramer and Diez 1975 [PG6/1 (N25/3)].			
729	6. Apiculiretusispora plicata (Allen) Streel 1967 [BNB45/1 (Y30/3)].			
730	7. Apiculiretusispora plicata (Allen) Streel 1967 [BNB45/1 (C37/4)].			
731	8. Apiculiretusispora minor McGregor 1973 [BNB11.1 (E50/2)].			
732	9. Apiculiretusispora minor McGregor 1973 [BNB45/1 (D37/2)].			
733	10. Apiculiretusispora minor McGregor 1973 [BNB45/1 (O37)].			
734	11. Apiculiretusispora arenorugosa McGregor 1973 [BNB11/1 (R23)].			
735	12. Apiculiretusispora brandtii Streel 1964 [BNB45/1 (U55)].			
736				
737	Plate II. Dispersed spores from the Lower Devonian Val d'Amour and Campbellton			
738	formations of New Brunswick (eastern Canada). All spores at magnification x1000 unless			
739	otherwise stated. Taxon name followed by sample/slide number and England Finder co-			
740	ordinates.			

- 1. Dibolisporites echinaceus (Eisenack) Richardson 1965 [BNB42/1 (W55/3)].
- 742 2-3. Dibolisporites quebecensis McGregor 1973 [BNB35/1 (B37/3)].
- 4. *Dibolisporites quebecensis* McGregor 1973 [BNB30/1 (C37/2)].
- 5. Dibolisporites eifeliensis (Lanninger) McGregor 1973 [BNB30/1 (E29/2)].
- 6. *Dibolisporites eifeliensis* (Lanninger) McGregor 1973[BNB39/1 (O24)].
- 746 7. Dibolisporites wetteldorfensis Lanninger 1968 [BNB29/1 (W29/3)].
- 747 8-9. *Dibolisporites wetteldorfensis* Lanninger 1968 [BNB23 (G37/4)].
- 10. Dictyotriletes emsiensis (Allen) McGregor 1973 [BNB40/1 (M28/3)].
- 11. Dictyotriletes subgranifer McGregor 1973 [BNB23/1 (X32)].
- 12. Dictyotriletes favosus McGregor & Camfield 1976 [BNB42/1 (J28/1)].
- 13. Dictyotriletes emsiensis (Allen) McGregor 1973 [BNB33/1 (H46/3)].
- 14. Dictyotriletes ?gorgoneus Cramer 1966 [BNB34/1 (P34)].
- 15. Dictyotriletes canadensis McGregor 1973 [BNB40/1 (G29)] Magnification x500.
- 754
- 755 Plate III. Dispersed spores from the Lower Devonian Val d'Amour and Campbellton
- formations of New Brunswick (eastern Canada). All spores at magnification x1000. Taxon
- name followed by sample/slide number and England Finder co-ordinates.
- 758 1. Ambitisporites sp. [BNB33/1 (D38/3)].
- 2. Crassitate trilete spore with a distal ornament of anastomosing muri [BNB45/1 (J49/3)].

- 3. Crassitate trilete spore with a distal ornament of anastomosing muri [BNB34/1 (N40/1)].
- 761 4-5. Synorisporites cf. verrucatus Richardson & Lister 1969 [BNB33/1 (G30)].
- 762 6. *Raistrickia* sp. [BNB36A/1 (N45/1)].
- 763 7. *Raistrickia* sp. [BNB33/1 (L45/4)].
- 764 8. Amicosporites jonkeri Riegel 1973 [BNB33/1 (L29/4)].
- 765 9. Synorisporites papillensis McGregor 1973 [BNB33/1 (U40/2)].
- 766 10. Verrucosisporites sp. A [PG3/1 (N40/1)].
- 11. Crassitate trilete spore with a distal ornament of verrucae [BNB40/1 (O46/2)].
- 12. Lycopodiacidites ogygius McGregor 1973 [BNB23/1 (M40)].
- 13. Verruciretusispora multituburculata (Lanninger) McGregor 1973 [BNB39/1(Q34/4)].
- 770 14. Verrucosisporites sp. A [BNB39/1 (M30/3)].
- 15. Verruciretusispora dubia (Eisenack) Richardson & Rasul 1978 [BNB47/1 (F50)].
- 16. Verrucosisporites sp. A [BNB37/1 (Q31)].
- 17. Verrucosisporites devonicus McGregor 1973 [BNB49/1(U40/4)].
- 774
- 775 Plate IV. Dispersed spores from the Lower Devonian Val d'Amour and Campbellton
- formations of New Brunswick (eastern Canada). All spores at magnification x1000. Taxon
- name followed by sample/slide number and England Finder co-ordinates.
- 1. Verrucosisporites polygonalis? Lanninger 1968 [BNB35/1 (J30/4)].

- 2. Verrucosisporites polygonalis? Lanninger 1968 [BNB45/1 (D37)].
- 780 3. *Brochotriletes* sp. B [BNB40/1 (T54)].
- 781 4. *Brochotriletes* [PG2/1 (S32)].
- 782 5. *Clivosispora verrucata* McGregor 1973 [PG6/1 (G41.4)].
- 6. *Clivosispora verrucata* McGregor 1973 [BNB42/1 (T25)].
- 784 7. *Camarozonotriletes sextantii* McGregor & Camfield 1976 [PG3/1(F31/2)].
- 8. *Camarozonotriletes sextantii* McGregor & Camfield 1976 [BNB11/1 (B46.3)].
- 9. *Brochotriletes foveolatus*? Naumova 1953 [BNB23/1 (M40)]. Singleton.
- 10. Brochotriletes bellatulus Steemans 1989 (=Brochotriletes sp. B McGregor 1973)
  [BNB42/1 (G49/3)].
- 789 11. Brochotriletes sp. A McGregor 1973 [BNB48/1 (M52)].
- 12. Tetrad of *Brochotriletes robustus* (Scott & Rouse) McGregor 1973 [BNB23/1 (N44/3)].
- 13. Brochotriletes robustus (Scott & Rouse) McGregor 1973 [PG2/1 (V43)].
- 14. Camarozonotriletes sextantii McGregor & Camfield 1976 [BNB05/1 (Q34)]. Spore
- enclosed within a large laevigate body with trilete spore as also reported by Andrews *et al.*(1974).

- 796 Plate V. Dispersed spores from the Lower Devonian Val d'Amour and Campbellton
- 797 formations of New Brunswick (eastern Canada). All spores at magnification x1000 unless

798	otherwise stated. Taxon name followed by sample/slide number and England Finder co-
799	ordinates.

- 800 1. *Emphanisporites annulatus* McGregor 1961 [BNB47/1(M36/4)].
- 2. Emphanisporites erraticus (Eisenack) McGregor 1961 [BNB35/1 (L37)].
- 802 3. *Emphanisporites schultzii* McGregor 1973 [PG2/1 (T38/1)].
- 4. *Emphanisporites rotatus* (McGregor) McGregor 1973 [PG2/1 (T30/2)].
- 5. *Emphanisporites genselae* sp. nov. Holotype. [BNB29/1(K41/3)].
- 805 6. *Emphanisporites genselae* sp. nov. [BNB29/2 (R37/2)].
- 806 7. *Emphanisporites genselae* sp. nov. [BNB29/1 (W38)].
- 807 8. *Emphanisporites genselae* sp. nov. [BNB30/1 (Y37/3)].
- 808 9. *Emphanisporites* spp. Singleton. [BNB36A/1(Q37/2)].
- 809 10. *Emphanisporites* spp. Singleton. [PG2/1 (W30)].
- 810 11. *Emphanisporites micrornatus* Richardson & Lister 1969 [BNB29/1 (C30/1)].
- 811 12. *Emphanisporites morrisae* sp. nov. Holotype. [BNB43/1 (Y40/1)]. Magnification x500.
- 812 13. *Emphanisporites morrisae* sp. nov. [BNB20/1 (G36/4)].
- 813 14. *Emphanisporites morrisae* sp. nov. [BNB33/1(N43)].

- 815 Plate VI. Dispersed spores from the Lower Devonian Val d'Amour and Campbellton
- 816 formations of New Brunswick (eastern Canada). All spores at magnification x1000. Taxon
- name followed by sample/slide number and England Finder co-ordinates.
- 818 1. ?*Archaeozonotriletes* sp. McGregor 1973 [PG2/1 (U40/4)].
- 819 2. Archaeozonotriletes chulus (Cramer) Richardson & Lister 1969 [BNB30/1 (P40)].
- 820 3. *Cymbosporites yorkensis* McGregor 1973 [BNB23/1 (P25/1)].
- 4. Acinosporites lindlarensis Riegel 1968 [PG2/1 (044/1)].
- 822 5-6. *Chelinospora* spp. McGregor 1973 [BNB36A/1 (R31/1)].
- 823 7. Acinosporites lindlarensis Riegel 1968 [BNB38/1 (E43/1)].
- 824 8-9. *Chelinospora* spp. [BNB45/1 (Y41/1)].
- 825 10. *Chelinospora* spp. [PG6/1 (X41)].
- 826 11. Acinosporites lindlarensis Riegel 1968 [BNB11/1 (E33/3)].

828 Plate VII. Dispersed spores from the Lower Devonian Val d'Amour and Campbellton

829 formations of New Brunswick (eastern Canada). All spores at magnification x1000 unless

otherwise stated. Taxon name followed by sample/slide number and England Finder co-

831 ordinates.

- 1. Zonotriletes brevivelatus Breuer and Steemans 2013 [BNB45/1 (M25/3)].
- 2-3.*Camptozonotriletes* sp. in McGregor 1973. Singleton. [BNB36A/1 (G46)].
- 4. *Zonotriletes brevivelatus* Breuer and Steemans 2013 [BNB45/1 (V28/2)].

835	5. Camptozonotriletes	caperatus McGregor	1973 [BNB45/1 (D24)].

836	6. Zonate-pseudosaccate spore with conate ornament	. Singleton.	[BNB32/1	(V42)].
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- 837 7. *Zonotriletes brevivelatus* Breuer and Steemans 2013 [BNB45/1 (R31/2)].
- 838 8. *Camptozonotriletes caperatus* McGregor 1973 [BNB38/1 (L43/3)].
- 839 9. *Pseudodyadospora petasus* Wellman & Richardson 1993 [PG6/1 (X31/3)].
- 10. *Zonotriletes brevivelatus* Breuer and Steemans 2013 [BNB43/1 (S48/3)].
- 841 11. *Camptozonotriletes caperatus* McGregor 1973 [PG3/1 (U35/1)]. Magnification x500.
- 842

Plate VIII. Dispersed spores from the Lower Devonian Val d'Amour and Campbellton
formations of New Brunswick (eastern Canada). All spores at magnification x500 unless
otherwise stated. Taxon name followed by sample/slide number and England Finder coordinates.

1. Grandispora douglastownense McGregor 1973 [PG3TOP/2 (R41/1)].

2. *Grandispora douglastownense* McGregor 1973 [PG3TOP/14 (U32)].

849 3. *Grandispora douglastownense* McGregor 1973 [PG3TOP/14 (V32/2)].

- 4. *Grandispora douglastownense* McGregor 1973 [PG3TOP/14 (V32/4)]. Specimen laterally
  compression.
- 5-6. *Ancyrospora loganii* McGregor 1973 [BNB47/1 (K47/4)]. (6) at magnification x1500.
- 853 Specimen laterally compressed. Close up of ornament illustrates bifurcate-tipped processes.
- 7. Ancyrospora loganii McGregor 1973 [BNB15/1 (X37)]. Magnification x1000.

855				
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857	TABLES			
858				
859	Table 1. Dispersed spore taxa identified in this study.			
	Retusotriletes eslae Cramer & Diez 1975 [Plate 1, figs 4-5]			
	Retusotriletes cf. rotundus (Streel) Streel 1967 [Plate 1, fig. 3]			
	Retusotriletes triangulatus (Streel) Streel 1967 [Plate 1, fig. 2]			
	Retusotriletes spp. [Plate 1 fig. 1]			
	Aniculiratusisnora aranorugosa McGregor 1073 [Plate 1 fig 1			

Apiculiretusispora arenorugosa McGregor 1973 [Plate 1, fig. 11] Apiculirestusispora brandtii Streel 1964 [Plate 1, fig. 12] Apiculiretusispora minor McGregor 1973 [Plate 1, figs 8-10] Apiculiretusispora plicata (Allen) Streel 1967 [Plate 1, figs 6-7] Apiculiretusispora spp. Dibolisporites echinaceus (Eisenack) Richardson 1965 [Plate 2, fig. 1] Dibolisporites eifeliensis (Lanninger) McGregor 1973 [Plate 2, figs 5-6] Dibolisporites quebecensis McGregor 1973 [Plate 2, figs 2-4] Dibolisporites wetteldorfensis Lanninger 1968 [Plate 2, figs 7-9] Dibolisporites spp. Dictyotriletes canadensis McGregor 1973 [Plate 2, fig. 15] Dictyotriletes emsiensis (Allen) McGregor 1973 [Plate 2, figs 10,13] Dictyotriletes favosus McGregor & Camfield 1976 [Plate 2, fig. 12] Dictyotriletes ?gorgoneus Cramer 1966 [Plate 2, fig. 14] Dictyotriletes subgranifer McGregor 1973 [Plate 2, fig. 11] Ambitisporites sp. [Plate 3, fig. 1] Amicosporites jonkeri Riegel 1973 [Plate 3, fig. 8] *Raistrickia* sp. [Plate 3, figs 6-7] Synorisporites papillensis McGregor 1973 [Plate 3, fig. 9] Synorisporites cf. verrucatus Richardson & Lister 1969 [Plate 3, figs 4-5] Crassitate trilete spore with a distal ornament of anastomosing muri [Plate 3, figs 6-7] Crassitate trilete spore with a distal ornament of vertucae [Plate 3, fig. 9] Lycopodiacidites ogygius McGregor 1973 [Plate 3, fig. 12] Verruciretusispora dubia (Eisenack) Richardson & Rasul 1978 [Plate 3, fig. 15] Verruciretusispora multituburculata (Lanninger) McGregor 1973 [Plate 3, fig. 13] Verrucosisporites devonicus McGregor 1973 [Plate 3, fig. 17] Verrucosisporites polygonalis? Lanninger 1968 [Plate 4, figs 1-2] Verrucosisporites sp. A [Plate 3, figs 10,14,16] Brochotriletes bellatulus Steemans 1989 (=Brochotriletes sp. B McGregor 1973) [Plate 4, fig. 10] Brochotriletes foveolatus? Naumova 1953 [Plate 4, fig. 9] Brochotriletes robustus (Scott & Rouse) McGregor 1973 [Plate 4, figs 12-13] Brochotriletes sp. A McGregor 1973 [Plate 4, fig. 11] Brochotriletes sp. B [Plate 4, figs 3-4] Emphanisporites annulatus McGregor 1961 [Plate 5, fig. 1]

*Emphanisporites erraticus* (Eisenack) McGregor 1961 [Plate 5, fig. 2] Emphanisporites genselae sp. nov. [Plate 5, figs 5-8] Emphanisporites micrornatus Richardson & Lister 1969 [Plate 5, fig. 11] *Emphanisporites morrisii* sp. nov. [Plate 5, figs 12-14] Emphanisporites rotatus (McGregor) McGregor 1973 [Plate 5, fig. 4] Emphanisporites schultzii McGregor 1973 [Plate 5, fig. 3] Camarazonotriletes sextantii McGregor & Camfield 1976 [Plate 4, figs7-8,14] Clivosispora verrucata McGregor 1973 [Plate 4, figs 5-6] Archaeozonotriletes chulus (Cramer) Richardson & Lister 1969 [Plate 6, fig. 2] ?Archaeozonotriletes sp. McGregor 1973 [Plate 6, fig. 1] Cymbosporites yorkensis McGregor 1973 [Plate 6, fig. 3] Chelinospora spp. [Plate 6, figs 5-6,8-9,10] Acinosporites lindlarensis Riegel 1968 [Plate 6, figs 4,7,11] Ancyrospora loganii McGregor 1973 [Plate 8, figs 5-7] Zonotriletes brevivelatus Breuer & Steemans 2013 [Plate 7, figs 1,4,7,10] Camptozonotriletes caperatus McGregor 1973 [Plate 7, figs 5,8,11] *Camptozonotriletes* sp. in McGregor 1973 [Plate 7, figs 2-3] Grandispora douglastownense McGregor 1973 [Plate 8, figs 1-4] Zonate spp. [Plate 7, figs 2-3,6] Pseudodyadospora petasus Wellman & Richardson 1993 [Plate 7, fig. 9] Tetrahedraletes medinensis (Strother & Traverse) Wellman & Richardson 1993

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**Table 2.** Distribution of spore taxa from the sections/samples analysed.

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Table 3. Plant taxa reported from the Val d'Amour and Campbellton formations and details
of their in situ spores. In terms of localities: SI – SVI refer to the sections of Kennedy &
Gibling (2011); A-Q refer to the plant-bearing localities of Gensel & Andrews (1984) and
Gensel et al. (1991); GH1-GH3 refer to plant-bearing localities of Grierson & Hueber (1968);
H1-H3 refer to plant-bearing localities of Hueber (1968). Facies Associations (FA) are from
Kennedy et al. (2012b). The stratigraphical relationship of these localities, some of which
represent the same locality, is summarised in Fig. 4.

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Table 4. Plant taxa reported from individual localities. Data is from references in Table 3
supplemented by data in Kennedy et al. (2012). In terms of localities: Sections A-H refer to
those used in this paper; SI – SVI refer to the sections of Kennedy & Gibling (2011); A-Q
refer to plant-bearing localities of Gensel & Andrews (1984) and Gensel et al. (1991); GH1GH3 refer to plant-bearing localities of Grierson & Hueber (1968); H1-H3 refer to plant-

bearing localities of Hueber (1968). Facies Associations (FA) are from Kennedy et al.

878 (2012b). The stratigraphical relationship of these localities, some of which represent the

- same locality, is summarised in Fig. 4. \* refers to spore age constraints reported by McGregor
- 880 (see Li et al. 2000).

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883	APPENDIX
884	
885 886 887	<b>Appendix 1.</b> Details of samples. Sections A-I refer to sections collected. The number in Parenthesis (I-VI refers to the equivalent section of Kennedy and Gibling 2012). + indicates productive samples that were logged and counted.
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