SYMМЕТРИЧЕСКАЯ БИФУРКАЦИЯ ЛИСТЬЯ АЛЕТОПТЕРИС УРОФИЛЛЯ

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ABSTRACT

A symmetrical bifurcation of an Alethopteris urophylla frond from the Pennsylvanian (Upper Carboniferous) of the Northern France coalfield shows some still-attached remains of foliage. Such rare specimens contribute, step by step, to a better understanding of the frond architecture of the genus Alethopteris, which is of the bifurcate-pinnate type. A few complementary general comments are given.

Key words: Palaeobotany, Carboniferous, Medullosales, Alethopteris, frond architecture.

INTRODUCTION

Zeiller (1888: 226-227) suggested that the frond of Alethopteris lonchitica (Schlotheim, 1804, 1820) probably exhibited a symmetrical dichotomy. His argument was based on a specimen (Zeiller, 1886: pl. 31) showing a correlation between the variation in size of the attached inner secondary pinnae and their location inside the presumed bifurcation. Unfortunately, owing to its large dimensions, the specimen could only be partly illustrated, and the bifurcation itself was not represented. The lack of formal proof for the connection of the primary rachides probably entailed that little attention was paid to Zeiller’s suggestion. Accordingly, most reconstructions of the Alethopteris Sternberg, 1825 frond were essentially proposing a strictly pinnate, non-dichotomizing frond (e.g. Buisine, 1961: figs 2a, 6; Laveine, 1967: fig. 2; Wnuk & Pfefferkorn, 1984: fig. 13).

Fortunately, the record of a specimen of Neuralethopteris Cremer, 1893, exhibiting a frond main bifurcation with still-attached remains of foliage (Laveine et al., 1993: pl. 1), together with the record (Goganova et al., 1993)
of remarkable remains of Cardioneuropteris Goganova, Laveine, Lemoigne & Durante, 1993, allowed Laveine et al. (1993) to reconsider formally the frond architecture of the taxa of the whole group. The latter authors provided a fairly extensive discussion on the matter, and proposed new frond reconstructions exhibiting naked petioles and a main symmetrical bifurcation (Laveine et al., 1993: fig. 3; Goganova et al., 1993: fig. 6). Laveine (1997: fig. 2) proposed the denomination ‘bifurcate-pinnate’ for such fronds. However, the assignment of Alethopteris to that type of frond was proposed by Laveine et al. (1993) only by analogy. They could only present (Laveine et al., 1993: pl. 3, fig. 1) a specimen showing two fragments of large frond bifurcations devoid of attached foliage, and ascribed them to the genus Alethopteris solely on the basis of association. The relationship therefore remained uncertain. Accordingly, Zodrow (2002: 177) noted that ‘a bifurcate alethopterid frond, …, remains to be demonstrated’. This remark emphasizes that decisive specimens are hard to come by. Consequently, any specimen that brings decisive information on frond organization is worth recording.

**MATERIAL AND METHODS**

The specimen described herein (accession number: MBL 630905) belonged initially to the collections of the Laboratory of Palaeobotany of Lille University. It was rediscovered during the transfer of the collections to the Lille Coal Museum in 2008. Apparently, this specimen was collected a long time ago from the coalfield of Northern France, probably from a spoil heap. No precise localisation is available.

The specimen required only slight mechanical degaging and was photographed using plain-light illumination. Terminology for frond architecture follows that used in Laveine (1997).

**DESCRIPTION**

The slab is 24 cm long and 25 cm wide. It bears the bifurcate fragment of an Alethopteris frond. The petiole, at least 22 mm wide, is only preserved in its top part, for 25 mm from the inside angle of bifurcation (Figs 1, 2a, 2b). The two primary rachides are at ca 50° angle. The right primary rachis is preserved straight in 180 mm length, after which it is bent slightly and broken off 20 mm further. Its width changes from 15 mm proximally to 13 mm distally. The left primary rachis is preserved for a length of 140 mm; its distal part is not preserved because of preferential splitting of the rock along the plane of a root remain of Radicites H. Potonié, 1893, located below the level of the Alethopteris foliage. The width of the left primary rachis changes from 14 mm proximally to 8 mm distally, but the original width was larger because the proximal part of the rachis is broken along its left border. Furthermore, it appears to have been more markedly crushed than the right primary rachis, as attested by the presence of prominent longitudinal compression ribs (Figs 2a, 3a).

Two fragmentary secondary rachides are preserved on the outer side of the right primary rachis. The proximal one, 28 mm long and 6 mm wide at the base, is located at 35 mm from the bifurcation. It bears a few fragmentary last order pinnae. The most proximal one, on the acroscopic side, is the better preserved (Figs 2a, 3e). It is 45 mm long and comprises at both sides seven lateral...
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pinnules below a rather short terminal pinnule. The next secondary rachis departure is located at 95 mm from the first one (Figs 2a, 3b: right-hand side). Unfortunately, it is only preserved for a very short distance: 4 mm. It is 5 mm wide, with no preserved foliage. These departures were attached adaxially to the primary rachis (Fig. 2c), on the face opposite to the viewer, which explains that some proximal parts of the foliage are covered by the crushed primary rachis (Fig. 3b).

Only one secondary rachis, 20 mm long and more or less 5 mm wide, is partly preserved at the external side of the left primary rachis (Figs 2a, 3c, arrow). It bears at its acroscopic side an incomplete ultimate pinnula. That secondary departure is slightly plunging into the rock matrix. It is located at 70 mm from the bifurcation and is, therefore, not quite symmetric to the corresponding right-hand-side secondary rachis. Assuming that the spacing of the outer secondary rachides was the same for both primary rachides, that original asymmetry, together with the shorter preservation of the left primary rachis explains why there is not another external secondary rachis departure visible at the left side of the specimen.

Inside the bifurcation, four secondary pinnulae are more or less well preserved, two attached to the left primary rachis (A and C on Fig. 1, moving proximo-distally), and two at the right primary rachis (B and D on Fig. 1). These secondary pinnulae are inserted almost symmetrically, A and C respectively at 30 mm and 104 mm, and B and D at 35 mm and 110 mm from the inside of the bifurcation. Because of space constraints, secondary pinnulae A and B are shorter and slightly less differentiated than secondary pinnulae C and D.

Secondary pinnula B, ca 50 mm long, is rather nicely preserved, slightly bent downwards in its distal part, with its very apex missing. It bears at both basiscopic and acroscopic sides a single ultimate pinnula, followed by several elongate pinnulae, the more proximal ones being 18 mm long and 5 mm wide (Fig. 3d). The basiscopic ultimate pinnula (Figs 1, 2a) of secondary pinnula B (with six and ? seven lateral pinnulae) is slightly more developed than the acroscopic one (with four lateral pinnulae). Unfortunately, the symmetrical secondary pinnula A is not so well preserved, especially at its basiscopic side. However it shows a similar degree of differentiation for its acroscopic side, i.e. a basal ultimate pinnula (with four lateral pinnulae) followed by several elongate pinnulae, evidently diminishing in size where approaching the partly preserved terminal pinnula (Figs 2a, 3d). Secondary pinnula A is also ca 50 mm long. Both secondary pinnulae A and B were inserted at 80° on their respective primary rachides. They are markedly overlapping (Figs 1, 2a, 3d).

Secondary pinnula D, ca 85 mm long, is also rather well preserved, although markedly distorted upwards distally. At both basiscopic and acroscopic sides, respectively, it bears four and three ultimate pinnulae, followed on either side by a lobate pinnula and several elongate pinnulae, gradually diminishing in size where approaching the partly preserved terminal pinnula. The acroscopic ultimate pinnulae are 30 mm long, and bear four or five lateral pinnulae each. The basiscopic pinnulae appear more developed than those at the acroscopic side (Figs 2a, 3b). Only the distal part of the first basiscopic pinnula is visible, its proximal part being hidden by the superimposed primary rachis. It was ca 60 mm long. The next basiscopic ultimate pinnula, also partly covered by the primary rachis, is ca 45 mm long and bears seven lateral pinnulae at its anadromic side. The following basiscopic foliage elements decrease in size and differentiation as usual. The symmetrical secondary pinnula C is not well preserved, especially at its acroscopic side because the rock split in favour of foliage debris of Sphenopteris sp., present on a plane located slightly below that of the Alethopteris specimen (Fig. 2a). Fortunately, the acroscopic side, although not perfectly preserved, is seen to show an approximately similar degree of differentiation as the acroscopic side of secondary pinnula D, i.e. with three ultimate pinnulae followed by a lobate pinnula (Fig. 3a). More distally, the preservation is defective where approaching the poorly preserved terminal pinnula. Secondary pinnula C is ca 83 mm long. Both secondary pinnulae C and D were also inserted at 80° on the respective primary rachides. When straightening the distal part of the rachis of secondary pinnula D, taphonomically distorted upwards, it appears that the tips of pinnulae C and D were originally slightly overlapping (Figs 1, 2a).

DISCUSSION

Morphological comments

The asymmetry in the insertion of the first outer secondary pinnulae, combined with the poorer preservation of the specimen on its left-hand side, might mean that the specimen in hand does not correspond to the main bifurcation of the Alethopteris frond. However, it is not uncommon to find occasionally such kind of asymmetrical specimens for bipartite fronds known to exhibit usually a rather more symmetrical architecture. Good examples are for instance the type specimen of Neuropteris heterophylla (Bromgniart, 1831: pl. 71), the type specimen of Laveineopteris guardinis (Grand’Eury, 1890: pl. 22, fig. 2), the specimens of Eusphenopteris scribanii Van Amerom, 1975 and Eusphe-
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nopteris sp. as illustrated in Laveine (1993: pl. 1, fig. 2, and in pl. 3, fig. 1 respectively). Conversely, the almost symmetrical location of the insertions of the inner secondary pinnae, the similar degree of differentiation of the preserved elements of foliage, and their overlapping along the bisector of the internal angle are clearly in favour of a fundamentally symmetric organisation for the present specimen. It is only unfortunate that the petiole is so incompletely preserved. However, the few petiolar striae visible at the lower part of the specimen (Figs 1, 2b, arrow) are oriented also more or less in the direction of the bisector. Though not wholly conclusive, this reinforces the impression that the specimen corresponds to the main (and only, in the case of Alethopteris bifurcation of the frond. The general organisation of the present specimen fits the frond reconstruction as proposed in Laveine et al. (1993: text-fig. 3). However, the basal 15 mm-width of the right primary rachis (the best preserved), as compared to the 20 mm-width of the Zeiller (1886: pl. 31) specimen (at a more distal location in the frond), and the 45 mm-width of the primary rachides of the specimen illustrated in Laveine et al. (1993: pl. 3, fig. 1) does suggest that the specimen in hand corresponds to a rather small size frond. The shorter length of the ultimate pinnæ of the present specimen, as compared to the length of the ultimate pinnæ of the specimen illustrated in Laveine (1986: pl. 1, fig. 2), i.e. 60 mm as against 150 mm, also attests to a rather small size frond. This would explain the slightly lesser differentiation of the foliage especially on the outer side (Fig. 3e), as compared to the reconstruction proposed in Laveine et al. (1993: text-fig. 3).

Taxonomical comments
The general outline and the rather poorly preserved venation of the pinnules of the present specimen suggest an assignment to Alethopteris lonchitica (Schlotheim, 1804, 1820), as understood by Zeiller (1886-1888) for specimens from northern France. However, Wagner & Álvarez-Vázquez (2008) reappraised the characteristics of the taxon as figured by Zeiller, and concluded that it corresponded to Alethopteris urophylla (Brongniart, 1834) Goeppter, 1836.

GENERAL CONSIDERATIONS

General morphological considerations
Zodrow (2002: fig. 18, and 2007: fig. 16) presented sketch drawings of an alethopterid tree carrying fronds exhibiting a main symmetrical bifurcation, and also some large foliage segments inserted well down on the petiole, below the main bifurcation. That reconstruction is based on rachis remains (Zodrow, 2002: fig. 9) showing an “Y axis” with several short rachial departures. Conversely, another specimen (Zodrow, 2002: figs 3, 4) shows, in organic connection to the stem, several naked petioles, i.e. with no rachial departures for a distance of at least 60 cm from the base, as is emphasized by Zodrow (2002: 192). Consequently, there is some discrepancy in the information provided by these two specimens. Unfortunately, all the rachides of large dimensions recorded from the site by Zodrow (2002) are devoid of attached foliage remains, thus leaving the usual uncertainty as to the correct taxonomic assignment of the rachides. It is known, for instance for the genus Eusphenopteris Simson-Scharold, 1934, that some bipartite fronds may (e.g. Zeiller, 1886: pl. 4, fig. 1; pl. 5, figs 1-2) or may not (e.g. Laveine, 1993: pl. 3, fig. 2; pl. 5, fig. 1) bear some lateral secondary pinnæ on the petiole, but such a possibility remains to be determined for the genus Alethopteris. The lack of a preserved petiole for the specimen in hand implies a lack of information in this respect. The few known specimens of fronds belonging to genera of the same natural group (e.g. Neuralethopteris, Cardioneuropteris), with a partly preserved petiole and with some remains of foliage attached, all show naked petioles.

Zodrow (2002: 187) suggested that ‘intercalated pinnate structures’ may possibly be present in the Alethopteris frond, but this suggestion is also based on association, the remains of ramifications being devoid of attached foliage. The taxonomic assignment is, therefore, disputable. In some cases, because of a marked adaxial insertion of secondary rachides (e.g. Buisine, 1961: pl. 13, fig. 1, re-illustrated in Wagner & Álvarez-Vázquez, 2008: figs 8-9), it may appear, when the fossil remain is seen from the abaxial side, that some ultimate pinnæ are directly attached to the corresponding primary rachis (e.g. Figs 1, 2a, 3b). However, this corresponds clearly to a taphonomic effect, with the primary rachis partly covering the most proximal part of the secondary rachides and, consequently, hiding the basiscopic part of the proximal ultimate pinnæ. This point has been made before (e.g. Buisine, 1961: pl. 34, explanation of figure 1). Of course, it is not excluded that a fossil remain of Alethopteris might be recorded some day showing a rachial intercalated pinnæ, such as for the specimen of the recent fern Pteridium aquilinum (Linné, 1753) Kuhn, 1879, illustrated here (Figs 4a, b, arrow), but this is theoretical. Such cases are very rare teratological cases,
with normally only a minute chance to being preserved as fossils. Another example is given here (Figs 4c, arrow, 4d) of a plant of Osmunda regalis Linné, 1753, with a frond showing secondary pinnae inserted alternately, whereas the other fronds of the same plant show the usual habit, with secondary pinnae oppositely attached (Fig. 4c). Another exceptional teratological case of development is presented in Josten et al. (2001: pls 1-6) for a calamitalean remain. Such teratological modifications never justify discarding the general diagnostic characteristics of the corresponding taxa. Obviously, all the large, rather numerous remains now available of true Alethopteris foliage (excluding, of course, those belonging to Callipteridium Weiss, 1870) attest clearly that the lack of rachial intercalated elements on the primary and secondary rachides is the normal, usual condition for Alethopteris.

A last morphological characteristic worthwhile being discussed is the degree of foliage differentiation of the Alethopteris frond. The proposed reconstruction in Laveine et al. (1993: text-fig. 3) shows a quadripinnate stage of differentiation in the proximal part of the primary pinnae, principally on the outer side. Unfortunately, the present specimen corresponds to a frond of rather small size. Even if better preserved, it would be unlikely to provide additional information of substance in this matter. As mentioned earlier, it is far from easy to record accurate specimens, and it is practically impossible to find such specimens for all the Alethopteris species. Despite the large amount of large specimens of Alethopteris stored in the collections of the Lille Museum of Geology there is only one specimen (MBL 19116) that brings slight information on that point (Fig. 5). It belongs to the Alethopteris missouriensis D. White, 1899, complex (see Wagner & Álvarez-Vázquez, 2008, for recent taxonomical considerations on this matter). The specimen shows three parallel fragments of large secondary pinnae. The parallel course of the two secondary rachides visible suggests that they were inserted on a common primary rachis, unfortunately not preserved (Fig. 5a). As a matter of consequence, that fossil remain appears at first sight tripinnate. However, a close examination of the long ‘ultimate’ pinnae of the specimen reveals that some basal ‘pinnules’ show a beginning of lobation (Figs 5b, 5c, arrows), thus attesting indubitably to a beginning of quadripinnatifid stage of foliage differentiation for the corresponding primary pinna (according to the terminology for frond architecture used in Laveine, 1997). Otherwise, for all the other specimens available showing a gradual differentiation of pinnules into ultimate pinnae (e.g. Wagner & Álvarez-Vázquez, 2008: figs 2, 9) it is impossible to decide on the true location of these specimens. They may either be close to the apex of the corresponding primary pinna (which is the more likely), or close to the apex of a proximal outer secondary pinna of a huge frond. It is therefore impossible to conclude safely on the degree of segmentation of the corresponding fronds.

**General taxonomical and palaeogeographical considerations**

According to the general outline of the pinnules, with a marked decurrent deltoid base, and to the aspect of their venation, the present bifurcate specimen must be assigned to Alethopteris urophylla, as re-described by Wagner & Álvarez-Vázquez (2008). Incidentally, it brings the opportunity to reappraise both a general taxonomical and palaeogeographical point dealt with by Wagner & Álvarez-Vázquez (2008). These authors (2008: 170, bottom of the left column) mentioned in the specimens *excludenda* the specimen illustrated by Laveine et al. (2003: pl. 10, fig. 1) as *Alethopteris cf. lonchitica*, and representing the distal part of an ultimate pinna. The specific assignment of the single specimen recorded being proposed with reservation in Laveine et al. (2003), it would be endless to discuss on the full accuracy of that specific assignment. However, Wagner & Álvarez-Vázquez (2008) suggesting a possible relationship of the specimen with *Neopteris obliqua* (Brongniart, 1833) Zeiller, 1888 *sensu lato*, it implies that the generic assignment is also disputed. The suggested relationship with *Neopteris obliqua* is not satisfactory. The marked catadromatic attachment of the lamina of the few lateral pinnules preserved is indisputably of *Alethopteris* type, as well as the marked midvein with its attachment slightly shifted toward the anadromic side of the pinnule. It is never the case for *Neopteris obliqua*, for which the midvein is always only faintly marked, and always strongly decurrent for the pinnules at such a distal location. Definitely, if the specific assignment can be questioned, this is not the case for the generic assignment. This point would not have been worthy of discussion here, were it not linked with another point discussed in Wagner & Álvarez-Vázquez (2008: 166) contesting, albeit implicitly, the presence of the genus *Alethopteris* in China. Therefore, it is necessary to emphasize that another specimen of *Alethopteris* illustrated in Laveine et al. (2003: pl. 18, fig. 1), also questionable as to its specific assignment.

**Figure 5.** a, large frond fragment of *Alethopteris missouriensis* D. White, 1899, showing three large parallel fragments of secondary pinnae more or less well preserved, and very likely inserted on a common primary rachis, not preserved. Scale bar represents 5 cm. Origin: Nord/Pas-de-Calais coalfield, Fouquières-les-Lens, seam Marcel, Faisceau d’Ernestine, Upper Bolsovian. Repository: Musée de Géologie, Ville de Lille, accession n° MBL 19116. b, enlargement of the right-hand part of the lowermost secondary pinna of the same specimen, the arrows point to the elements enlarged on figure 5c. Scale bar represents 1 cm. c, enlargement of the right part of the lowermost secondary pinna in the vicinity of the secondary rachis, the two arrows points to some lobate ‘pinnules’, attesting to a beginning of quadripinnatifid stage of differentiation. Scale bar represents 5 mm.
exhibits characteristics rendering totally indisputable the generic assignment. Therefore, it can be safely concluded that the genus *Alethopteris*, as well as the genus *Lonchopteris* Brongniart, 1828 (Zhang et al., 1993: pls 19-20), although rarely recorded, are definitely present in the Pennsylvanian (Upper Carboniferous) deposits (and also in the lowermost Permian deposits for *Alethopteris*) of Northwest China. The paucity of records is clearly linked with the fact that it concerns remote areas difficult to access. There is little doubt that future investigations will extend the general palaeogeographical distribution of these genera, as ultimately concluded also by Wagner & Álvarez-Vázquez (2008: 181).

**SUMMARY AND CONCLUSION**

The specimen subject of the present paper brings a definite argument attesting to the bipartite architecture of the frond for the genus *Alethopteris*, a characteristic which remained until now to be formally established. Despite the fact that *Alethopteris* is abundantly recorded from the Pennsylvanian deposits in the Euramerican area, experience shows that it is far from easy to find significant specimens recording its organization. Now a bipartite architecture has been proved for one of the most common species of *Alethopteris*, it means that the large specimen of *Alethopteris* dealt with in Laveine (1986) is only part of one of the two symmetrical primary pinnae of the frond. Accordingly, it can now be firmly concluded that the large fronds of *Alethopteris* may have reached a width of about four metres.

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