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Fungal remains in cordaite (Cordaiales) leaves from the Upper Pennsylvanian of central France

MICHAEL KRINGS, THOMAS N. TAYLOR, NORA DOTZLER & JEAN GALTIER



Little is known to date regarding the organisms that colonized the leaves of Carboniferous plants. This is due to a number of factors, including how the leaves were preserved, prepared for examination, and analyzed. Nevertheless, it is becoming increasingly clear that Carboniferous leaves were habitats for a wide variety of fungi. This paper describes fungal remains in cordaite leaves from the Upper Pennsylvanian Grand-Croix cherts in France. Within the intercellular system of the mesophyll occur intercalary and terminal swellings produced from an elaborate system of tenuous hyphae/filaments, as well as isolated putative (zoo-)sporangia and clusters of small unicells. Other fungal remains occur in bundle sheath cells. Although the systematic affinities of the fungi currently remain unknown, documenting their existence contributes to our understanding of the multiple levels of biological association and interaction that shaped the Carboniferous terrestrial ecosystems. • Key words: bundle sheath, Carboniferous, chert, fossil fungi, Grand-Croix, spongy parenchyma.

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Cordaiteae are one of the most conspicuous elements in Late Pennsylvanian and Early Permian tropical forest ecosystems (Barthel 1976, Šimůnek 2000, Martín-Closas & Galtier 2005). These plants, which are thought to represent a grade of evolution leading to modern conifers (e.g., Hilton & Bateman 2006, Gomankov 2009), were up to 45 m tall; the above-ground portion included massive, woody trunks and a crown of elongate, strap-shaped leaves, some of which were up to 70 cm in length (Šimůnek & Libertín 2006). Extending the length of the leaves are alternating bands of fibers and vascular bundles; their spacing and arrangement, together with leaf shape and epidermal anatomy, have been used to distinguish various leaf morphospecies (e.g., Harms & Leisman 1961, Šimůnek 2007). Recent research suggests that the cordaiteae were well-established in several different environments, including poorly to well drained mineral soils, as well as coastal and upland settings (Falcon-Lang 2003, 2005; Falcon-Lang & Bashforth 2004, 2005).

Although much is known about the biology and ecology of cordaiteae based on innumerable impression/compression specimens and permineralizations (Taylor *et al.* 2009), one

aspect that rarely is addressed concerns the various associations and interactions of these plants with other groups of organisms. For example, Cichan & Taylor (1982) described arthropod borings in the woody cordaitelean root *Premnoxylon*, and Strullu-Derrien *et al.* (2009) reported on arbuscular mycorrhizal fungi in *Radiculites*-type rootlets. While these associations have been demonstrated in secondary wood and the below-ground parts of the plants, we are unaware of any clearly defined evidence of associations of other organisms, especially arthropods and microorganisms, with the leaves of cordaiteae, with the possible exception of a few questionable structures interpreted as bite marks (Müller 1982) and leaf mines (Labandeira 1998). This is surprising since most cordaite leaves are quite large, and thus appear to have represented particularly suitable sites for arthropod herbivory and colonization by biotrophic and saprotrophic epiphyllous and endophytic microorganisms.

In this paper we report on the occurrence of fungi in structurally preserved cordaite leaves from the Upper Pennsylvanian (Carboniferous) Grand-Croix cherts of central France. The fungi occur within the intercellular system

of the mesophyll, as well as in the parenchymatous sheaths that surround the vascular bundles. This discovery provides the first evidence of cordaite leaves colonized by fungi. Moreover, the occurrence of some of the fungi is restricted to well-defined areas within the leaves, which may suggest that these organisms were biotrophs.

Material and methods

The thin sections containing the fragments of cordaite leaves come from the Grand-Croix cherts, which occur in the eastern part of the Saint-Étienne Basin in central France. They occur within the so-called “Poudingue mosaïque” at several localities in the vicinity of the city of La Grand-Croix (Galtier 2008). The stratigraphic provenance and age of the Grand-Croix cherts are difficult to resolve, but generally they are interpreted as belonging either to the terminal portion of the Rive de Gier Formation (dated as Stephanian A; see Vetter 1971), or to the sediments overlying the Rive de Gier Formation, which have been dated as earliest Stephanian B (Pruvost 1947). Regardless of the age, the cherts are probably reworked, and therefore may be slightly older, *i.e.* late Stephanian A (equivalent to the late Barruelian) in age (Doubinger *et al.* 1995).

The infected leaf fragments are contained in two different thin sections of Grand-Croix chert deposited in the paleobotanical collection of the Swedish Museum of Natural History (Naturhistoriska riksmuseet) in Stockholm (Sweden) under acquisition numbers S003419-05 (Fig. 1) and S003409-06 (Fig. 2). The fungi were examined in transmitted light, with digital images captured with a Leica DFC 480 digital camera and processed in Adobe Photoshop.

Description

The infected leaf fragments are represented by mm-sized pieces of leaf tissue. One fragment appears in a longitudinal (radial) section (Fig. 1), the other in a paradermal (sagittal) plane (Fig. 2). The fragments are identified as cordaite leaves based on internal structure (*e.g.*, compare Figs 1A, 2A with Harms & Leisman 1961, pl. 126, figs 1, 3), as well as the morphology and arrangement of stomata (Fig. 2B). Be-

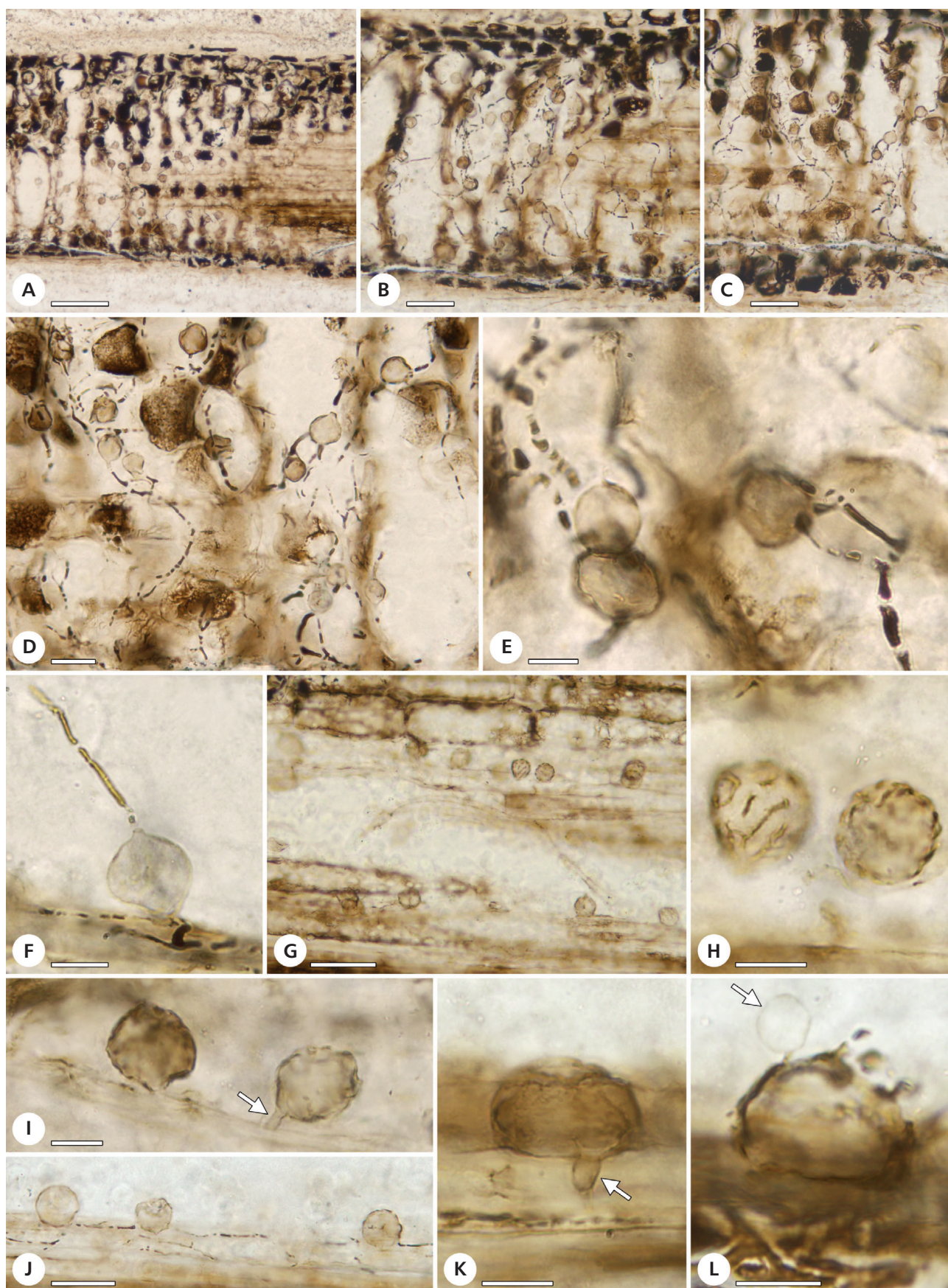
cause we are dealing with isolated leaf fragments, and thus cannot determine whether both fragments come from the same type (*i.e.* morphospecies) of cordaite leaf, we will describe the fungal remains from each leaf piece separately under the designation *leaf fragment 1* and *leaf fragment 2*.

Leaf fragment 1 (Fig. 1)

This fragment of leaf (Fig. 1A) is approximately 20 mm long and 0.5 mm wide. Within the leaf are several different types of fungal remains, with the most conspicuous being a mycelium of tenuous hyphae or filaments (each of which up to 2 µm in diameter) producing both intercalary and terminal swellings. This mycelium occurs exclusively within the intercellular system of the mesophyll (Fig. 1B, C), most frequently in the large intercellular spaces of the spongy parenchyma (Fig. 1D). The swellings range from spherical to slightly spindle-shaped and are up to 18 µm in diameter. The wall is thin, unornamented, and lacks any observable exit pore or tube (Fig. 1E). What is unusual about this fossil fungus is that the swellings and hyphae/filaments appear to have been preserved in a different state of development. In general, the swellings show little evidence of decay. The hyphae/filaments are preserved in place and in a configuration that indicates their exact relationship with the swellings during life (*e.g.*, Fig. 1D, E). However, they appear to have been in an advanced state of disintegration at the time of preservation. This is evidenced by the opaqueness of most of the hyphae/filaments (Fig. 1E). Moreover, almost all hyphae/filaments are not represented by continuous threads, but rather consist of irregular segments that, although dissociated, are still aligned to reflect hyphae/filaments. While most of the swellings occur freely within the intercellular spaces, some are attached to the outer surfaces of parenchyma cells (Fig. 1F).

In addition to this conspicuous fungus, there are several other types of fungal remains within the intercellular system. One of these consists of more or less spherical structures up to 18 µm in diameter that are attached to outer host cell wall surfaces, preferably to outer surfaces of bundle sheath cells (Fig. 1G). These spherical structures are relatively thin-walled, and some distinctly ornamented by discontinuous and irregularly distributed ridges (Fig. 1H). While most of the spheres are sessile, some possess a short,

Figure 1. *Leaf fragment 1.* • A – longitudinal (radial) cross section of cordaite leaf, showing mesophyll and part of a bundle sheath (to the right of image); bar = 100 µm. • B, C – mycelium within intercellular system of mesophyll; bar = 50 µm. • D – detail of Fig. 1C, showing mycelium with intercalary and terminal spherical structures; bar = 20 µm. • E – spherical structures; note irregular segments that although disassociated are still aligned to reflect hyphae/filaments; bar = 10 µm. • F – spherical structure with dissociated hyphae/filaments at both ends; bar = 10 µm. • G – spherical structures attached to outer surfaces of bundle sheath cells; bar = 50 µm. • H – detail of spherical structure ornamentation; bar = 10 µm. • I – stalked spherical structures; bar = 10 µm. • J – spheres on outer host cell surface; note basal mycelium extending out on host cell surface; bar = 30 µm. • K – spherical structure with what appears to be an infection peg (arrow); bar = 10 µm. • L – putative mycoparasite (arrow) attached to spherical structure; bar = 10 µm.



narrow stalk (arrow in Fig. 1I). Both unornamented and ornamented spheres arise from, or give rise to, delicate mycelia extending out on the outer surface of the host cells to which the spheres are attached (Fig. 1J). These mycelia are identical in appearance to that seen in the intercellular system; the individual hyphae/filaments, however, are slightly narrower, *i.e.* up to 1 µm in diameter. We are unsure as to whether this type of fungus is simply a different form of the type that occurs freely in the intercellular system, or represents some other fungus, or is perhaps a different developmental stage of the form described above.

Still other spherical structures attached to outer host cell wall surfaces are distinctly larger (up to 25 µm in diameter) and have more opaque walls (Fig. 1K, L). Some of these latter spheres show what appears to be an infection peg penetrating into the lumen of the host cell (arrow in Fig. 1K). In one example there is a small, thin-walled spherule attached to one of the larger spheres (arrow in Fig. 1L).

Leaf fragment 2 (Fig. 2)

This fragment of leaf (Fig. 2A) measures 1.3 × 7 mm, and shows several tissues of the leaf that are not well recognizable in *leaf fragment 1* [*e.g.*, the stomatiferous epidermis (Fig. 2B) and subepidermal tissue (Fig. 2C)]. Fungal remains within this leaf (Fig. 2D, E) consist of spherical to pyriform or elliptical structures up to 17 µm in diameter that may be free within the intercellular spaces of the mesophyll or attached to the outer wall surfaces of mesophyll cells (Fig. 2F, G, J–N). The wall of some of these spheres appears double (*e.g.*, Fig. 2K), suggesting that they contain a slightly smaller inner body, which may represent a spore or simply a preservational artifact caused by shrinkage of the lumen and separation of the wall layers. It is also possible, however, that the wall of the spheres is composed of different wall components giving the impression of a double wall. Some of the spherical structures have 1–2 hyphal/filament fragments attached, indicating both intercalary and terminal development; an elaborate mycelium like that seen in the intercellular system of *leaf fragment 1*, however, is not recognizable. It is interesting to note that spheres are more frequent toward the margin of the intercostal fields (Fig. 2F, G), and in these regions the fungi usually occupy the corners of the intercellular spaces that are directly adja-

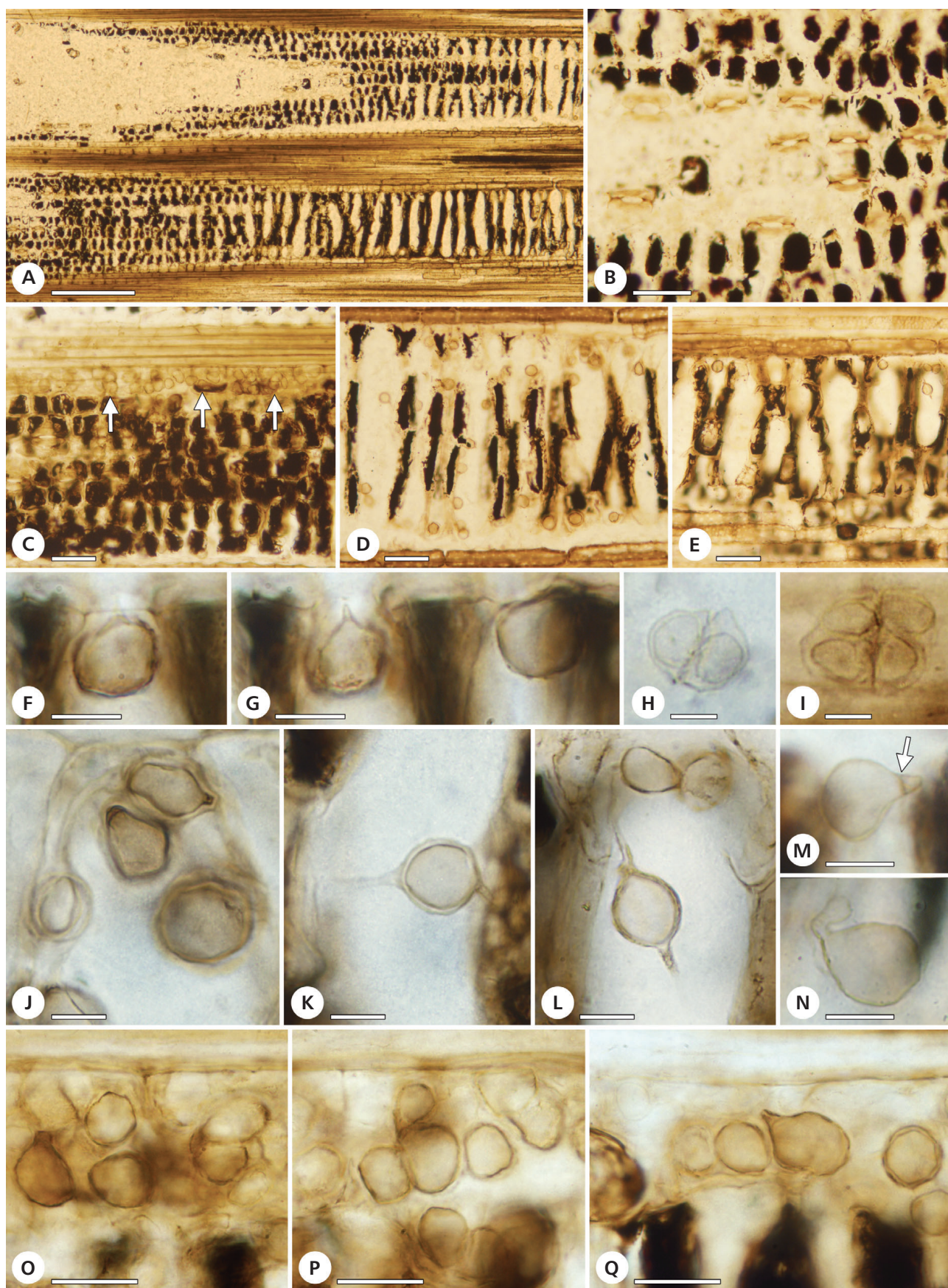
cent to the bundle sheaths. In some instances (*e.g.*, Fig. 2J) the spheres in the corners appear to be surrounded by a diffuse, translucent investment. Many of the spheres contain slightly pointed or irregularly elongate protrusions (Fig. 2G, M, N) that may be more opaque; some of the protrusions appear to be separated from the main sphere by a septum (arrow in Fig. 2M).

In one area of the leaf fragment, structures similar to those that are present in the intercellular system occur densely clustered within the parenchymatous cells of a bundle sheath (arrows in Fig. 2C; Fig. 2O–Q), in the outer cell layer directly adjacent to the subepidermal tissues and spongy parenchyma. Attachment to subtending hyphae in these clusters was not observed, and thus also infection pathways could not be determined. Fig. 2H, I shows clusters of 2–8 minute, thin-walled spherical unicells up to 5 µm in diameter, each of which is surrounded by what appears to be a translucent sheath. The arrangement of the components in several-cell units is always regular and consistent.

Discussion

Despite the fact that there is an increasing number of records of well preserved Carboniferous fungi (*e.g.*, Taylor *et al.* 1994; Krings *et al.* 2009a–c, 2010a–c, 2011; Strullu-Derrien *et al.* 2009, 2011), very few of these are reports on fungi associated with leaves of vascular plants. Moreover, these few occurrences are usually based on single reports (*e.g.*, Oliver 1903, Barthel 1961, Krings 2001, Krings *et al.* 2009b, Hübers *et al.* 2011), rather than including several infected specimens of the same leaf type that could provide a more inclusive understanding of the fungi (Krings *et al.* 2010c). This is puzzling since leaf endophytes represent a significant proportion of today's total fungal diversity (Petrini 1991, Rodriguez *et al.* 2009), and all major groups of fungi were in existence by the Carboniferous (Blair 2009). The fungal remains in cordaite leaves from Grand-Croix are special among the leaf-inhabiting fungi known from the Carboniferous. Firstly, they do not represent single occurrences, but rather entire assemblages of morphologically and in part spatially distinct fungal remains. Secondly, the assemblages represent the first documented evidence of fungi colonizing cordaite leaves, which are distinct among the leaf types occurring in Carboniferous coal swamp

Figure 2. *Leaf fragment 2.* • A – slightly oblique paradermal (sagittal) section of cordaite leaf, showing vascular bundles and mesophyll; bar = 500 µm. • B – detail of epidermis (with stomata) and subepidermal tissue; bar = 100 µm. • C – detail of subepidermal tissue; note clusters of fungal remains in bundle sheath cells (arrows); bar = 50 µm. • D, E – fungal remains in intercellular system of spongy parenchyma; bars = 50 µm. • F, G – spherical structures in intercellular spaces adjacent to bundle sheath; bars = 10 µm. • H, I – pair and small cluster of thin-walled unicells, each surrounded by what appears to be a translucent sheath; bars = 5 µm. • J–N – spherical structures in intercellular system of spongy parenchyma; arrow in Fig. 2M indicates septum at the base of protrusion; bars = 10 µm. • O–Q – clusters of spherical structures in bundle sheath cells (see arrows in Fig. 2C); bars = 20 µm.



plants (*i.e.* microphylls and compound fronds), and thus may have provided some special features for colonization.

We are uncertain as to what major groups the fungi represented by the remains in these two cordate leaf fragments may belong to. Moreover, it is difficult to identify biological species because the different types of fungal remains within the leaves may not all represent separate species, but rather different life history stages of the same organism. It is also possible that the individual types of remains each represent several morphologically similar species. Unfortunately, this lack of preciseness is an inherent problem in paleomycology that has been detailed previously (Sherwood-Pike 1991, Krings *et al.* 2010c), and thus need not to be addressed here again. The fungal remains in *leaf fragment 1* based on their overall morphology might represent rhizomycelial systems and zoosporangia of monocentric (*e.g.*, Fig. 1K, L) and polycentric (*e.g.*, Fig. 1D, J) chytrids (Chytridiomycota) (see Karling 1977). However, affinities with other groups of Fungi or fungus-like microorganisms (*e.g.*, Hyphochytridiomycota, Peronosporomycetes) cannot be ruled out. The same is true of the intercalary and terminal spherical structures occurring in the intercellular system of *leaf fragment 2* (*e.g.*, Fig. 2F, G, J–N). The clusters of intracellular spheres illustrated in Fig. 2O–Q resemble chytrid-like organisms described previously from Grand-Croix (compare Fig. 2C, O–Q with Krings *et al.* 2009c, pl. I, fig. 5a–c). The latter occur in cortical cells of a specimen of the zygopteridalean reproductive structure *Biscalitheca* cf. *musata*. The affinities of the pairs and clusters of small unicells illustrated in Fig. 2H, I remain elusive, and could include algae, cyanobacteria, and some fungal structures. It is interesting to note that similar structures have been reported from the Lower Devonian Rhynie chert where they occur in the rhizome cortex of the early land plant *Horneophyton lignieri* (Taylor *et al.* 1992, figs 1–12) and from Viséan (Middle Mississippian) cherts of France where they occur freely in the chert matrix (Krings *et al.* 2009a, pl. III, figs 4–6). The Rhynie and Viséan unicells, however, are larger (*i.e.* >10 µm in diameter) and distinguished from the Grand-Croix fossils by the presence of a short discharge tube extending from the unicells. The Viséan specimens have been interpreted as zoosporangia of an endobiotic chytrid that formed holocarpic thalli in host cells (Krings *et al.* 2009a).

Another interesting but difficult question concerns the timing of when the fungi entered and colonized the host leaves. In general, this question relates to whether the leaves were alive and fully intact when colonization by the fungi occurred, or had become moribund but were still attached to the tree, or had already been abscised and were lying on the forest floor. Based on the conditions of the interior tissues, especially the spongy parenchyma (Figs 1A, B, 2A, D, E), one may suggest that the leaves were alive at the time of colonization. Moreover, two of the

fungi in *leaf fragment 1* (Fig. 1D, J) appear in a life history stage in which sporangia/propagules have been produced, but the producing hyphal/filament system has not yet completely undergone disintegration. This sequence of events suggests that the time of infection predates the condition at the time of preservation by some time. Arguing in favor of colonization during the lifetime of the leaves is also the occurrence of intracellular clusters of fungal remains that are restricted to parenchymatous bundle sheath cells (Fig. 2C, O–Q). On the other hand, the infected fragments represent minute portions of usually rather large leaves, and it is reasonable to assume that this fragmentation was the result of considerable mechanical forces that occurred prior to fossilization. If this is accurate then one might challenge the assumption that infection occurred during the lifetime of the leaf, because the delicate hyphal/filament systems present in *leaf fragment 1* (Fig. 1B–D, J) do not show evidence of mechanical distortion, which would certainly have occurred if the fungi were present when the leaf became fragmented. This line of reasoning rather suggests that colonization occurred after the leaf abscised from the parent plant and became fragmented, and thus that the fungi represent saprotrophs. Compounding the determination of the timing of when a fungus actually entered its host plant with regard to the Grand-Croix fossils is a general lack of information on the sequence of events and timing of precisely when silicification occurred.

In spite of the inherent problems, the discovery of fungi in cordate leaves has scientific merit. Firstly, it provides a general framework with which to consider the utilization of microhabitats by fungi during the Carboniferous. In this context it will be interesting to see if morphologically similar types of fungi were generalists and colonized the leaves of different plants, and/or whether these particular types of fungi were general in overall preference for plant organs (*e.g.*, roots, shoot axes, leaves, reproductive structures). Secondly, despite the lack of information on the biological affinities of the fungi, it is possible to trace certain features of how the fungi functioned and interacted with their host. For example, the hyphal/filament network of the fungal morphotype shown in Fig. 1B–E is in the process of disintegration, while the spherical structures formed by this network are consistently intact. This suggests to us that in this particular fungus the mycelium had completed its function at the time of fossilization. Moreover, the spatial arrangement of the fungal remains within the leaves suggests that the spongy parenchyma may have represented a particularly suitable site for fungal colonization, perhaps related to the presence of large intercellular spaces. As a result, the fungal remains reported in cordate leaf fragments from Grand-Croix expand our knowledge of both the fungi and what kinds of plant tissues they inhabited in the Carboniferous. Reports such as this represent initial steps in better understanding of the Carboniferous swamp forest

communities and the multiple levels of biological association and interaction that shaped these complex ecosystems some 300 Ma ago.

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References

- BARTHEL, M. 1961. Ein Pilzrest aus dem Saarkarbon. *Geologie* 10, 856–857.
- BARTHEL, M. 1976. Die Rotliegendflora Sachsens. *Abhandlungen des Staatlichen Museums für Mineralogie und Geologie zu Dresden* 24, 1–190.
- BLAIR, J.E. 2009. Fungi, 215–219. In HEDGES, S.B. & KUMAR, S. (eds) *The Timetree of Life*. Oxford University Press, New York, NY.
- CICHAN, M.A. & TAYLOR, T.N. 1982. Wood-borings in *Premnoxylon*: plant-animal interactions in the Carboniferous. *Palaeogeography, Palaeoclimatology, Palaeoecology* 39, 123–127. DOI 10.1016/0031-0182(82)90075-X
- DOUBINGER, J., VETTER, P., LANGIAUX, J., GALTIER, J. & BROU-TIN, J. 1995. La flore fossile du bassin houiller de Saint-Etienne. *Mémoires du Museum national d'Histoire naturelle, Paris* 164, 1–357.
- FALCON-LANG, H.J. 2003. Anatomically-preserved cordaitalean trees from a dryland alluvial plain setting, Joggins, Nova Scotia. *Atlantic Geology* 39, 259–265.
- FALCON-LANG, H.J. 2005. Small cordaitalean trees in a marine-influenced coastal habitat in the Pennsylvanian Joggins Formation, Nova Scotia. *Journal of the Geological Society of London* 162, 485–500. DOI 10.1144/0016-764904-080
- FALCON-LANG, H.J. & BASHFORTH, A.R. 2004. Pennsylvanian uplands were forested by giant cordaitalean trees. *Geology* 32, 417–420. DOI 10.1130/G20371.1
- FALCON-LANG, H.J. & BASHFORTH, A.R. 2005. Morphology, anatomy, and upland ecology of large cordaitalean trees from the Middle Pennsylvanian of Newfoundland. *Review of Palaeobotany and Palynology* 135, 223–243. DOI 10.1016/j.revpalbo.2005.04.001
- GALTIER, J. 2008. A new look at the permineralized flora of Grand-Croix (Late Pennsylvanian, Saint-Etienne basin, France). *Review of Palaeobotany and Palynology* 152, 129–140. DOI 10.1016/j.revpalbo.2008.04.007
- GOMANKOV, A.V. 2009. Pollen evolution in cordaites and early conifers. *Paleontological Journal* 43, 1245–1252. DOI 10.1134/S0031030109100062
- HARMS, V.L. & LEISMAN, G.A. 1961. The anatomy and morphology of certain *Cordaite* leaves. *Journal of Paleontology* 35, 1041–1064.
- HILTON, J. & BATEMAN, R.M. 2006. Pteridosperms are the backbone of seed-plant phylogeny. *Journal of the Torrey Botanical Society* 133, 119–168. DOI 10.3159/1095-5674(2006)133[119:PATBOS]2.0.CO;2
- HÜBERS, M., BOMFLEUR, B., KRINGS, M. & KERP, H. 2011. An Early Carboniferous leaf-colonizing fungus. *Neues Jahrbuch für Geologie und Paläontologie, Abhandlungen* 261, 77–82. DOI 10.1127/0077-7749/2011/0150
- KARLING, J.S. 1977. *Chytridiomycetorum Iconographia. An Illustrated and Brief Descriptive Guide to the Chytridiomycetous Genera with a Supplement of the Hyphochytridiomycetes*. viii + 414 pp. J. Cramer, Vaduz.
- KRINGS, M. 2001. Pilzreste auf und in den Fiedern zweier Pteridospermen aus dem Stefan von Blanzky-Montceau (Zentralfrankreich). *Abhandlungen des Staatlichen Museums für Mineralogie und Geologie zu Dresden* 46/47, 189–196.
- KRINGS, M., DOTZLER, N., GALTIER, J. & TAYLOR, T.N. 2009a. Microfungi from the upper Visean (Mississippian) of central France: Chytridiomycota and chytrid-like remains of uncertain affinity. *Review of Palaeobotany and Palynology* 156, 319–328. DOI 10.1016/j.revpalbo.2009.03.011
- KRINGS, M., DOTZLER, N., TAYLOR, T.N. & GALTIER, J. 2009b. A Late Pennsylvanian fungal leaf endophyte from Grand-Croix, France. *Review of Palaeobotany and Palynology* 156, 449–453. DOI 10.1016/j.revpalbo.2009.04.010
- KRINGS, M., GALTIER, J., TAYLOR, T.N. & DOTZLER, N. 2009c. Chytrid-like microfungi in *Biscaliitheca* cf. *musata* (Zygopteridales) from the Upper Pennsylvanian Grand-Croix cherts (Saint-Etienne Basin, France). *Review of Palaeobotany and Palynology* 157, 309–316. DOI 10.1016/j.revpalbo.2009.06.001
- KRINGS, M., DOTZLER, N., TAYLOR, T.N. & GALTIER, J. 2010a. A fungal community in plant tissue from the Lower Coal Measures (Langsettian, Lower Pennsylvanian) of Great Britain. *Bulletin of Geosciences* 85, 679–690. DOI 10.3140/bull.geosci.1209
- KRINGS, M., DOTZLER, N., TAYLOR, T.N. & GALTIER, J. 2010b. Microfungi from the upper Visean (Mississippian) of central France: Structure and development of the sporocarp *Mycocarpon cinctum* nov. sp. *Zitteliana A* 50, 127–135.
- KRINGS, M., TAYLOR, T.N., GALTIER, J. & DOTZLER, N. 2010c. Microproblematic endophytes and epiphytes of fern pinnules from the Upper Pennsylvanian of France. *Geobios* 43, 503–510. DOI 10.1016/j.geobios.2010.02.002
- KRINGS, M., DOTZLER, N., GALTIER, J. & TAYLOR, T.N. 2011. Oldest fossil basidiomycete clamp connections. *Mycoscience* 52, 18–23. DOI 10.1007/s10267-010-0065-4
- LABANDEIRA, C.C. 1998. Early history of arthropod and vascular plant associations. *Annual Review of Earth and Planetary Sciences* 26, 329–377. DOI 10.1146/annurev.earth.26.1.329
- MARTÍN-CLOSAS, C. & GALTIER, J. 2005. Plant taphonomy and paleoecology of Late Pennsylvanian intramontane wetlands in

- the Graissessac-Lodève basin (Languedoc, France). *Palaios* 20, 249–265.
- MÜLLER, A.H. 1982. Über Hyponome fossiler und rezenter Insekten, erster Beitrag. *Freiberger Forschungsheft C* 366, 7–27.
- OLIVER, F.O. 1903. Notes on fossil fungi. *New Phytologist* 2, 49–53. DOI 10.1111/j.1469-8137.1903.tb05809.x
- PETRINI, O. 1991. Fungal endophytes of tree leaves, 179–197. In ANDREWS, J.H. & HIRANO, S.S. (eds) *Microbial Ecology of Leaves*. Springer-Verlag, New York.
- PRUVOST, P. 1947. Le Stéphaniens du bassin houiller de la Loire. *Annales de la Société Géologique du Nord* 67, 279–295.
- RODRIGUEZ, R.J., WHITE, J.F. JR., ARNOLD, E.E. & REDMAN, R.S. 2009. Fungal endophytes: diversity and functional roles. *New Phytologist* 182, 314–330. DOI 10.1111/j.1469-8137.2009.02773.x
- SHERWOOD-PIKE, M. 1991. Fossils as keys to evolution in fungi. *BioSystems* 25, 121–129. DOI 10.1016/0303-2647(91)90018-G
- ŠIMŮNEK, Z. 2000. Cuticles of *Cordaites* from the Westphalian, Stephanian and Autunian of the Bohemian Massif (Czech Republic) (a preliminary study). *Acta Palaeobotanica* 40, 25–34.
- ŠIMŮNEK, Z. 2007. New classification of the genus *Cordaites* from the Carboniferous and Permian of the Bohemian Massif, based on cuticle micromorphology. *Acta Musei nationalis Pragae, Series B – historia naturalis* 62, 97–210.
- ŠIMŮNEK, Z. & LIBERTIN, M. 2006. *Cordaites schatzlarensis* sp. nov. and *Samaropsis newberryi* (Andrews) Seward from the Westphalian (Carboniferous) of the Žacléř area (Czech Republic). *Review of Palaeobotany and Palynology* 138, 43–62. DOI 10.1016/j.revpalbo.2005.10.001
- STRULLU-DERRIEN, C., RIOULT, J.P. & STRULLU, D.G. 2009. Mycorrhizas in Upper Carboniferous *Radiculites*-type cordaitalean rootlets. *New Phytologist* 182, 561–564. DOI 10.1111/j.1469-8137.2009.02805.x
- STRULLU-DERRIEN, C., KENRICK, P., RIOULT, J.P. & STRULLU, D.G. 2011. Evidence of parasitic Oomycetes (Peronosporomycetes) infecting the stem cortex of the Carboniferous seed fern *Lyginopteris oldhamia*. *Proceedings of the Royal Society, Series B* 278, 675–680. DOI 10.1098/rspb.2010.1603
- TAYLOR, T.N., GALTIER, J. & AXSMITH, B.J. 1994. Fungi from the Lower Carboniferous of central France. *Review of Palaeobotany and Palynology* 83, 253–260. DOI 10.1016/0034-6667(94)90073-6
- TAYLOR, T.N., REMY, W. & HASS, H. 1992. Fungi from the Lower Devonian Rhynie chert: Chytridiomycetes. *American Journal of Botany* 79, 1233–1241. DOI 10.2307/2445050
- TAYLOR, T.N., TAYLOR, E.L. & KRINGS, M. 2009. *Paleobotany. The Biology and Evolution of Fossil Plants*. xxi + 1230 pp. Elsevier/Academic Press Inc., Burlington MA, London, San Diego CA, New York NY.
- VETTER, P. 1971. Le Carbonifère supérieur et le Permien du Massif Central, 169–213. In JUNG, J. (ed.) *Symposium*. Plein-Air Service, Clermont-Ferrand.