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CHEMICAL EVIDENCE FOR CELL WALL LIGNIFICATION AND THE EVOLUTION OF TRACHEIDS IN EARLY DEVONIAN PLANTS

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Anatomically preserved land plant fossils from the Lower Devonian Rhynie Chert contain conducting tissues with cells that range from dark-colored, elongated cells without secondary wall thickenings to tracheids similar to those of extant tracheophytes. A suite of tissue-specific microanalytical techniques was used to assess lignification in fossils of *Aglaophyton*, *Rhynia*, and *Asteroxylon*. Isotope ratio mass spectrometry provides millimeter-scale resolution of carbon isotopic abundances, whereas soft X-ray carbon (1s) spectromicroscopy provides micrometer-scale resolution of the preservation of organic molecular functionality. The isotopic and organic chemistry of Rhynie Chert plants indicates that the earliest vascular thickenings were probably unlignified and that cell wall lignification may have first appeared in the outer cortex. Only in more derived plants, it seems, was lignin deposited in conducting cells to produce the true tracheids seen today in vascular plants.

Keywords: xylem, tracheid, lignin, Rhynie Chert, paleobotany.

Introduction

Recently developed techniques of chemical microanalysis (Hemsley et al. 1996; Edwards et al. 1997; House et al. 2000; Boyce et al. 2001, 2002; Kudryavstev et al. 2001) make possible the paleontological study of ancient biochemistries at the cellular or tissue scale. Here we apply microanalytical methods to resolve the distribution of distinctive molecular and isotopic characteristics among tissues in three key early land plants from the Lower Devonian (Pragian) Rhynie Chert of Scotland. At Rhynie, siliceous spring deposits (Kidston and Lang 1921; Trewin 1994; Rice et al. 1995) have preserved-sometimes in life position and commonly with organic cell wall remnants and little compaction or deformation-a range of land plants that span the evolution of tracheids (Kidston and Lang 1917, 1920a, 1920b; Edwards 1993; Remy et al. 1993, 1994; Kenrick and Crane 1997). Our chemical analyses, thus, help to resolve the timing of the chemical and morphological innovations that collectively revolutionized water conduction in early land plants.

As defined in living plants, tracheids are elongated, water conducting cells of the vascular plant sporophyte that are dead at maturity and have cell walls strengthened with secondary thickenings and impregnated with the phenolic polymer lignin (essentially poly-bisphenol). Lignin occurs uniquely in vascular plants where it is embedded in the polysaccharide-based cell

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walls of some cell types. Detailed investigations of fossils from the Lower Devonian (fig. 1) have demonstrated that many of these cellular characteristics, all consistently associated with tracheids in living plants, accumulated sequentially through time. Protracheophyte fossils possess elongated cells that occupy the same central location as xylem in early tracheophyte axes, but the walls of these cells are smooth (Edwards 1986) and closely resemble the walls of water conducting cells in some mosses (Hébant 1977). Fossils of the rhyniophyte lineage, the sister group to the eutracheophyte clade that includes all living vascular plants (Kenrick and Crane 1997), possess tracheids with unusual S-type wall thickenings distinct from those seen in living plants (Kenrick et al. 1991a; Edwards 1993). Even the restriction of tracheids to the sporophyte generation is an independent characteristic that was one of the latest in the tracheid suite to evolve: the gametophyte fossil Sciadophyton contains S-type tracheids (Kenrick et al. 1991b). Lower Devonian eutracheophyte fossils have tracheids similar to those found in many plants today (Kidston and Lang 1920b).

The same range of fossil plants that show sequential structural innovations may preserve different stages in the physiological evolution of cell wall chemistry, including both lignin biosynthesis itself and the anatomical deployment of lignin within the plant. The presence of lignin in these fossils has been extensively debated, with reference made both to the taphonomy of cell ultrastructure and to the tracheids of basal extant lineages (Edwards 1993; Kenrick and Crane 1997; Friedman and Cook 2000 and accompanying W. G. Chaloner discussion).

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Because the isotopic and organic chemistries of lignin and



Fig. 1 Evolutionary relationships, reconstructions, stem cross sections, and conducting cell wall anatomy of (*A*) the protracheophyte *Aglaophyton major* (images modified from Gensel and Andrews 1984; Kenrick and Crane 1997), (*B*) the rhyniophyte *Rhynia gwynnevaughnii* (modified from Kenrick and Crane 1997), and (*C*) the eutracheophyte *Asteroxylon mackiei* (modified from Taylor and Taylor 1993; Kenrick and Crane 1997). Scale bars of reconstructions equal 1 cm; scale bars of cross sections equal 1 mm.

cellulose are distinct, chemical analyses may provide a more direct assessment of lignification in early fossil plants than inferences from preserved wall structure. For chemistry to be informative, however, comparative analyses must achieve a high degree of spatial resolution. The carbon isotopic values of lignin are 4‰–8‰ lighter than the cellulose of the same plant (Benner et al. 1987). However, because a host of environmental variables may also affect isotopic values (e.g., Buchmann et al. 1997; Sparks and Ehleringer 1997), lignified and unlignified tissues of the same age within the same fossil plant must be compared in order to distinguish the fractionation due to lignin biosynthesis from that due to environmental fluctuations. Because diagenetic alteration of fossil organic material results in the aromatization of originally nonaromatic material, including cellulose (Boyce et al. 2002), detection of aromatic chemistry in plant fossils is insufficient to indicate the original presence of lignin. One must compare the tissue chemistries of fossils having unambiguously lignified cells with those of fossils bearing tissues of unknown original composition and must do so using specimens that have experienced the same taphonomic history.

Recent taphonomic studies based on fine-scale tissue comparisons verify that the chemistry within fossils is preserved with sufficient spatial resolution to be physiologically informative (Boyce et al. 2001, 2002). Here we employ isotope ratio mass spectrometry on samples drilled manually from fossils (with millimeter-scale spatial resolution) and X-ray spectromicroscopy performed on peels of fossils (with micrometerscale spatial resolution).

Material and Methods

Organic and isotopic chemical analyses were performed on three genera from the Rhynie Chert representing lineages with distinct gradations of vascular organization: the eutracheophyte Asteroxylon mackiei, the rhyniophyte Rhynia gwynnevaughnii, and the protracheophyte Aglaophyton major. Additional isotopic controls were performed on silicified Araucarioxylon wood from the Lower Carboniferous Olentangy Shale of Ohio and extant, greenhouse-grown Psilotum nudum.

Isotope Ratio Mass Spectrometry

Samples for isotopic analysis were obtained by drilling the polished surfaces of hand samples containing silica-permineralized plant fossils. Diamond drill bits are preferable for softer cherts, including those preserving *Araucarioxylon*, but the substantially harder Rhynie Chert requires the use of carbon-free cubic boron nitride bits in order to avoid contamination of the sample with carbon eroded from the drill surface. Drill bits had a 1.2-mm head diameter and produced holes less than 1.5 mm wide. Drill speeds used were 15,000–18,000 rpm.

The powder resulting from drilling was collected frequently with a steel scalpel and stored in small aluminum foil packets until sufficient powder had been generated for analysis. The powder was then tapped into a tin sample boat. Drill bits, scalpels, and aluminum foil were cleaned before use by sonication for 15 min in hexane. The delicate sample boats were sonicated in hexane for 1 min, and the fossils were washed with hexane but not sonicated. The surfaces of fossils and all equipment were rinsed with ethanol and allowed to air dry after the collection of each sample. The drill and work area were wrapped in aluminum foil to provide a more dust-free work space that was easier to clean between samples. Sampling of a large quartz crystal following this procedure reveals that no more than 0.5 μ g of carbon contaminant is introduced during sample preparation (table 1).

Isotopic measurements were made with a Finnigan Delta Plus Excel isotope ratio mass spectrometer with a CE Instruments, NA 2500 series, elemental analyzer and a Conflo II interface. The gas chromatograph oven was set to 60°C for the fossil samples. Acetanilide standards were only included at the beginning of each set of analyses (followed by two or

Sample identity	Sample size (µg)	Voltage amplitude	Calculated % carbon	δ ¹³ C ‰
Controls to determine amount of external		-		
carbon contamination:				
Cleaned boat ^a	na	No peak	na	na
Quartz ^b	166	0.031	0.42	-24.47
Quartz	72	No peak	na	na
Araucarioxylon ring samples included in				
figures 2, 3:				
Ring 1A	1132	1.598	2.66	-21.00
Ring 1B	1381	2.100	2.87	-21.04
Ring 1C	2034	2.768	2.57	-21.01
Ring 1D	1805	2.434	2.55	-20.95
Ring 1 average				-21.00 ± 0.04
Ring 2A	1251	1.911	2.88	-21.13
Ring 2B	1536	2.365	2.92	-21.03
Ring 2C	1221	1.823	2.83	-21.04
Ring 2D	1710	2.569	2.84	-20.87
Ring 2 average				-21.02 ± 0.11
Ring 3A	824	1.209	2.77	-20.85
Ring 3B	852	1.226	2.72	-20.70
Ring 3C	1709	2.432	2.68	-20.51
Ring 3D	1174	1.645	2.64	-20.53
Ring 3E	1887	2.576	2.84	-20.65
Ring 3 average				-20.65 ± 0.14
Subsampling of a single <i>Araucarioxylon</i> sample included in figure 3:				
Subsample 1	83	0.111	3.07	-21.84
Subsample 2	219	0.262	2.78	-21.5
Subsample 3	36	0.079	5.08	-23.7
Subsample 4	138	0.181	3.00	-20.62
Subsample 6	24	0.187	17.99	-19.85
Subsample 7	44	0.076	3.95	-21.68
Subsample 8	19	0.039	4.71	-22.26
Subsample 9	288	0.333	2.66	-21.37
Subsample 10	109	0.144	3.08	-21.43
Subsample 11	157	0.186	2.75	-21.33
Subsample 12	187	0.213	2.65	-20.95
Subsample 13	619	0.652	2.47	-21.02
Subsample 14	344	0.400	2.72	-20.9
Subsample average				-21.42 ± 0.91
Additional Araucarioxylon samples included				
in figure 3:				
1	102	0.159	2.85	-21.32
2	634	0.922	2.67	-20.87
3	538	0.828	2.83	-20.24
4	243	0.353	2.69	-20.64
5	316	0.474	2.76	-20.27
6	676	0.954	2.59	-19.98
7	315	0.439	2.56	-20./1
8	336	0.428	2.33	-20.6/
י גער גער גער גער גער גער גער גער גער גער	328	0.560	3.13	-20./9
10	85	0.149	3.22	-20.82
11	232	0.340	2.68	-21.21
12	190	0.311	3.00	-20.46
13	338	0.535	2.90	-20.91
14	330	0.561	3.13	-20.53
15	442	0./4/	3.11	-20.35
16	1080	1./36	2.95	-20.87
Average				$-20.6/\pm0.35$

 $\label{eq:Table 1} Table \ 1 \\ \delta^{13}C \ Values \ for \ Control \ Samples \ of \ Quartz \ and \ Araucarioxylon \ Wood$

^a No peak was ever detected with the cleaned blank boats run with all subsequent analyses.

^b Quartz sampled with cubic boron nitride and Araucarioxylon with diamond drill bits.

	Sample size	Voltage	Calculated %	
Sample identity	(μg)	amplitude	carbon	δ ¹³ C ‰
Samples of extant Psilotum:				
Psilotum 1-V ^a	301	3.757	48.02	-25.51
Psilotum 1-I	330	3.22	37.5	-24.92
Psilotum 1-O	334	4.349	50.14	-25.19
Psilotum 2-V	381	4.559	46.06	-25.34
Psilotum 2-I	310	3.199	39.68	-24.96
Psilotum 2-O	386	4.869	48.53	-25.22
Psilotum 3-V	436	5.586	49.68	-25.75
Psilotum 3-I	304	3.323	42.16	-25.03
Psilotum 3-O	317	5.992	73.05	-25.26
Average for vasculature minus inner corr	tex per stem: -0.56 ± 0.17			
Largest samples of Aglaophyton fossils inc	luded			
in figure 5:				
Aglaophyton 1-V ^b	886	0.215	0.56	-25.42
Aglaophyton 1-IC	1046	0.145	0.32	-25.61
Aglaophyton 1-OC	931	0.278	0.67	-26.2
Aglaophyton 2-V	1254	0.199	0.37	-24.79
Aglaophyton 2-IC	ca. 1000	0.066	0.11	-24.89
Aglaophyton 2-OC	1080	0.073	0.17	-25.69
Aglaophyton 3-V	682	0.065	0.24	-22.77
Aglaophyton 3-IC	906	0.147	0.42	-22.75
Aglaophyton 3-OC	921	0.101	0.28	-23.36
Average for vasculature minus inner cor	tex per stem: 0.09 ± 0.11			
Average for outer cortex minus vasculat	ure per stem: -0.76 ± 0.16			
Average for outer cortex minus inner co	rtex per stem: -0.67 ± 0.12			
Additional, smaller Aglaophyton samples	-			
included in figure 5:				
Aglaophyton 4-V	751	0.104	0.39	-24.15
Aglaophyton 4-IC	1216	0.085	0.18	-24.67
Aglaophyton 4-OC	697	0.066	0.25	-23.11
Aglaophyton 5-V	840	0.093	0.28	-23.44
Aglaophyton 5-IC	705	0.06	0.26	-27.33
Aglaophyton 5-OC	865	0.067	0.19	-24.18
Aglaophyton 6-V	572	0.06	0.26	-25.46
Aglaophyton 6-IC	720	0.055	0.19	-24.49
Aglaophyton 6-OC	933	0.1	0.27	-24.1
Aglaophyton 7-V	255	0.072	0.56	-20.37
Aglaophyton 7-IC	334	0.047	0.31	-19.64
Aglaophyton 7-OC	283	0.057	0.42	-20.14
Aglaophyton 8-V	?	0.053	0.13	-24
Aglaophyton 8-IC	357	0.07	0.49	-24.15
Aglaophyton 8-OC	148	0.036	0.61	-25.81
Average for vasculature minus inner corr	tex per stem: 0.57 ± 1.95			
Average for outer cortex minus vasculat	ure per stem: 0.02 ± 1.30			
Average for outer cortex minus inner co	rtex per stem: 0.59 ± 1.86			
Samples of <i>Rhynia</i> fossils included in figur	e 6:			
Rhynia 1-V ^c	886	0.153	0.43	-24.3
Rhynia 1-C	1004	0.137	0.34	-24.05
Rhynia 2-V	684	0.158	0.58	-24.13
Rhynia 2-C	672	0.105	0.39	-23.37
Rhynia 3-V	1380	0.083	0.15	-23.74
Rhynia 3-C	1410	0.167	0.29	-24.34
Rhynia 4-V	1110	0.105	0.23	-29.7
Rhynia 4-C	891	0.103	0.28	-25.09
Rhvnia 5-V	895	0.073	0.21	-24.23
Rhvnia 5-C	770	0,105	0.34	-25.95
Rhynia 6-V	632	0.081	0.32	-23.14
Rhynia 6-C	730	0.085	0.29	-21.89
Rhvnia 7-V	888	0.077	0.22	-23.39
Rhvnia 7-C	934	0.083	0.22	-22.62
	201	0.000	·	22.02

(Continued)					
Sample identity	Sample size (µg)	Voltage amplitude	Calculated % carbon	δ ¹³ C ‰	
Rhynia 8-V	161	0.063	0.78	-24.83	
Rhynia 8-C	295	0.051	0.36	-20.82	
Average for vasculature minus cortex per stem: -1.	17 ± 2.16				
Largest samples of <i>Asteroxylon</i> fossils included in figure 6:					
Asteroxylon 1-V ^c	1415	0.584	0.98	-24.39	
Asteroxylon 1-C	1129	0.072	0.17	-25.22	
Asteroxylon 2-V	794	0.15	0.47	-24.49	
Asteroxylon 2-C	1180	0.113	0.24	-25.36	
Average for vasculature minus cortex per stem: 0.85	± 0.03				
Additional samples of <i>Asteroxylon</i> fossils included in figure 6:					
Asteroxylon 3-V	201	0.107	1.01	-21.65	
Asteroxylon 3-C	201	0.047	0.51	-20.36	

Table 2

Note. For each taxon, stems are arranged in order of descending average sample size.

^a For *Psilotum*, tissues sampled for each stem are (V) central vasculature, (I) inner cortical tissues, and (O) outer tissues including the stem surface.

^b For Aglaophyton, tissues sampled for each stem are (V) central vasculature, (IC) inner cortex, and (OC) outer cortex avoiding the stem surface.

^c For Rhynia and Asteroxylon, tissues sampled for each stem are (V) central vasculature and (C) cortex.

three blank sample boats) and at the end after all fossil samples had been run in order to eliminate the possibility that trace residue from the carbon-rich acetanilide standards might contaminate fossil samples.

Controls for this analytical protocol included repeated sampling along successive Araucarioxylon tree rings; measurements within rings were up to 15 mm apart, but each was no more than 1.5 mm away from a measurement in an adjacent ring. In living plants, carbon isotopic values are stable within individual tree rings but can vary between adjacent rings (Leavitt and Long 1982; Loader et al. 1995). The pattern observed in living plants is preserved in Araucarioxylon, demonstrating that small spatial variations in carbon isotopic ratios can be retained in fossils and measured with $\pm 0.1\%$ resolution (fig. 2; table 2). However, a further control study, in which a single large Araucarioxylon drill sample was subdivided into 14 aliquots of varying size, demonstrated that this resolution is strongly dependent on sample size and is progressively lost as the carbon content of samples decreases below ca. $3-5 \mu g$ (fig. 3; table 2). This control is key; it demonstrates that carbon isotopic analysis will only be informative when sufficient carbon is drilled from permineralized fossils.

Rhynie Chert samples prepared for isotopic analyses were drilled from cross sections of fossil stems (fig. 4). Tissues sampled in *Aglaophyton* were the central conducting strand, inner cortex, and outer cortex (avoiding the actual stem surface) (fig. 5). Only a central conducting strand and general cortical sample could be collected in *Rhynia* because of its small size (fig. 6). Since the steles of *Aglaophyton* and *Rhynia* are each less than 1 mm wide, the vascular samples from these species always included a relatively constant area of the adjacent inner cortex. Central vasculature and cortex were also sampled in *Asteroxylon* (fig. 6). Though generally aligned, the sampled stems were all at slightly different angles within the rocks. Because of this variation, sample drilling was limited to within

a millimeter of the surface to insure that a different tissue than intended was not sampled at greater depth.

Carbon isotopic values were also measured in *Psilotum*. Tissues sampled were the stele, the inner cortex, and the stem periphery, which included the outer cortex and epidermis. Samples were excised with a scalpel, freeze-dried overnight, and ground with a mortar and pestle before analysis.

Carbon Absorption X-Ray Spectroscopy

Organic materials preserved in the central conducting strands of the three Rhynie taxa were compared using carbon (1s) X-ray Absorption Near Edge Spectroscopy (C-XANES) in the 280–300 eV frequency range, just below the carbon 1s absorption edge. X-ray absorbance in this frequency range results from various electron core level (1s) transitions to stable bound states (e.g., π^* states) that correlate with specific functional groups (Stöhr 1992). C-XANES data were acquired using the scanning transmission X-ray microscope and microspectrometer located at the X1A beam line at the National Synchrotron Light Source at Brookhaven National Laboratory. Energy resolution is 0.3 eV, and the spatial resolution is ca. 100 nm.

Each sample spectrum has been corrected for background absorption by subtraction of a control spectrum (without a sample) and is presented in absorption mode as the $-\log (T/T_o)$, where T and T_o are the transmittance of the sample and background, respectively. Quantitative comparison of the different samples requires a self-consistent fitting of the C-XANES spectra with Gaussian bands (at 285.2 eV [Aryl-C, H], 287.1 eV [Aryl-O], 287.9 eV [C_{Al}-H], 288.5 eV [carboxylate], 289.5 eV [C_{Al}-O], and 291 eV [absorption associated with the ionization edge]) in order to extract relative absorption band intensities. After normalization of band intensities to account for differences among organic functional groups (Ishii and Hitch-



Fig. 2 Petrified *Araucarioxylon* wood from the Olentangy Shale. *A*, Carbon elemental map with dark indicating low and light indicating high carbon abundance, $50-\mu$ scale (modified from Boyce et al. 2001). *B*, Sampling and averages of carbon isotopic measurements from successive wood rings. Measurement details found in table 1. Scale bar is 1 mm.

cock 1988; Francis and Hitchcock 1992; Hitchcock et al. 1992), compositional estimates were made using scaled intensities based on published spectra of known compounds (Ishii and Hitchcock 1988; Hitchcock et al. 1992; Ade 1998).

Fossil samples for X-ray spectroscopy must be demineralized and thin enough (70-200 nm) for X-ray transmission. This thickness was achieved with a minimal loss of spatial resolution by the production of cellulose acetate peels of Rhynie fossil blocks using hydrofluoric acid and following standard procedures (Joy et al. 1956). Peels were embedded in epoxy and sectioned to 100-200 nm perpendicular to the peel surface. This preparation obscures the ultrastructural continuity of individual walls because visualization is in the same plane as the deformation caused by the acetate sheet on organic matter exposed by etching. Despite this deformation, however, the cell type under analysis is still tightly controlled (fig. 7; Boyce et al. 2002). Only the conducting cells of the fossils investigated here have a sufficient density of organic wall material for this technique to be practical. Detailed discussions of sample preparation (Cody 1996; Boyce et al. 2002) and equipment specifications (Jacobsen et. al. 1991) are available elsewhere.

Results

Isotopic Analyses

Overall, the carbon isotopic values of the fossils were similar to those of *Psilotum*. The average of δ^{13} C values for *Psilotum* was $-25.2\% \pm 0.3\%$ and for the larger Rhynie Chert samples

was $-24.5\% \pm 1.0\%$. (δ^{13} C values of atmospheric and surface ocean CO₂ fluctuated during the Early Devonian, but in general they are estimated to have been ca. 1‰ more positive than today; Veizer et al. 1999.) The increased variance for the Rhynie specimens reflects the inclusion of multiple individuals from three different taxa (as opposed to the repeated sampling of a single *Psilotum* plant). *Araucarioxylon* samples were also within the range of extant plants that use C₃ photosynthesis: mean δ^{13} C was $-20.8\% \pm 0.3\%$ for samples larger than 100 µg.

Within *Psilotum*, carbon isotopic values of the lignified central vasculature were $0.6\% \pm 0.2\%$ more negative than those of the contemporaneously produced unlignified inner cortex (table 2). As with the fossil *Araucarioxylon* controls (figs. 2, 3), the isotopic measurements from Rhynie fossils are extremely variable in smaller samples but show a decrease in variability with increasing sample size (figs. 5, 6).

Rhynia samples contain little organic matter, and so, despite repeated measurements, isotopic results remained variable and, therefore, physiologically inconclusive across all sample sizes (fig. 6; table 2). In contrast, the larger samples from *Aglaophyton* produced replicable isotopic ratios. The central conducting strand and inner cortex have approximately equal isotopic values when tissues are compared within cross sections of individual stems. These inner tissues, however, can be differentiated isotopically from the anatomically distinct outer cortex: the outer cortex is $0.7\% \pm 0.1\%$ more negative (fig. 5; table 2). *Asteroxylon* shows a different pattern of isotopic heterogeneity; the larger of our few samples show a negative isotopic shift of 0.8‰ in the cortex relative to the central xylem (fig. 6B; table 2).



Fig. 3 Carbon isotopic measurements of petrified *Araucarioxylon* wood from the Olentangy Shale plotted against voltage peak amplitude, which is a measure of sample carbon content as detected by the mass spectrometer. Measurements from a single large homogenous sample being subdivided into 14 samples of varying size (circles). The smallest sample that was attempted was not recognized by the mass spectrometer. Measurement details found in table 1. Subsample measurements plotted against background of all other wood measurements (squares); details found in table 1.



Fig. 4 Example of *Aglaophyton* stem used for isotopic analysis. Letters indicate drill holes for samples of (*V*) vasculature, (*I*) inner cortex, and (*O*) outer cortex. Only a single type of cortical measurement was made in both *Rhynia* and *Asteroxylon*. Scale bar is 1 mm.

Organic Analyses

As previously documented, Asteroxylon tracheid walls possess two distinct organic zones, similar to the biochemical zonation seen in the tracheids of other fossils and extant plants (Boyce et al. 2002). In contrast, X-ray spectroscopy indicates that the walls of conducting cells in Aglaophyton and Rhynia are more homogenous. Moreover, C-XANES spectra acquired for quantitative organic analysis of the cell walls in the three taxa show that conducting cell walls in Aglaophyton and Rhynia are extremely similar to each other but different from those of Asteroxylon (fig. 8; table 3). Aglaophyton and Rhynia conducting cells retain as much oxygen-bonded aromatic carbon as the tracheid wall layers of Asteroxylon but contain less aromatic carbon overall. Aglaophyton and Rhynia also show greater proportional representation of aliphatic carbon as well as ether and alcohol groups. One of the two wall layers observed in Asteroxylon tracheids reveals a chemical structure reminiscent of those seen in Aglaophyton and Rhynia; it has a similar proportion of single-bonded oxygen groups as well as abundances of aliphatic and aromatic carbon that are intermediate between those of the other Asteroxylon tracheid layer and the conducting cell walls of the other two taxa.

Discussion

Evidence for Lignification in Rhynie Chert Fossils

Isotopic comparisons of the lignified xylem protostele and unlignified inner cortex of *Psilotum* show that lignified tissues are 0.7‰ or 0.8‰ more negative than unlignified tissues (table 2). The full range of isotopic differentiation of 4‰–8‰ between pure lignin and cellulose (Benner et al. 1987) cannot be expected from tissue-scale analyses because lignin exists *in vivo* only as a lignocellulosic complex. The isotopic analyses of rings in the Lower Carboniferous *Araucarioxylon* wood indicate that isotopic differences as small as those distinguishing lignified and unlignified tissues can be preserved in fossil plants and can be accurately measured within $\pm 0.1\%$ (fig. 2). However, control studies also indicate that this high degree of isotopic resolution is lost as sample size decreases to below 3–5 μ g of carbon (fig. 3). Analyses must, therefore, always be evaluated in terms of sample size, a requirement of particular importance for Rhynie, because preserved fossils are small and have low carbon contents.

Despite the technical limitations on isotopic analyses of Rhynie fossils, *Aglaophyton* presents a clear and consistent isotopic pattern (table 2). The conducting strand and inner cortex are isotopically indistinguishable. Since the thin-walled parenchyma cells of the inner cortex are probably the least likely in the plant to be lignified, this result suggests that the conducting strand was also unlignified. However, a negative isotopic shift



Fig. 5 Comparisons of tissue-specific carbon isotopic values of individual *Aglaophyton* stems. Each datum represents the carbon isotopic difference between two of the three sampled tissues (vasculature, inner cortex, and outer cortex) from a single stem plotted against the average voltage peak amplitude of the two samples. Details of larger sample measurements included in table 2, those of smaller samples in table 2.



Fig. 6 Comparisons of carbon isotopic values from the vasculature and cortex of individual stems of *Rbynia* and *Asteroxylon*. Each datum represents the carbon isotopic difference between the two tissues from a single stem plotted against the average voltage peak amplitude of the two samples. Details of *Rbynia* measurements included in table 2, those of *Asteroxylon* samples in table 2.

of the same magnitude as that seen between lignified and unlignified tissues of *Psilotum* characterizes peripheral tissues of *Aglaophyton*. This result suggests that *Aglaophyton*'s outer cortex may have been lignified even if the conducting cells were not. Other than lignin, cuticle is the only prominent plant constituent that could both produce this cortical isotopic shift and survive diagenesis. Nuclear magnetic resonance spectroscopy of bulk Rhynie macerates, however, indicates that these plants contained no more aliphatic carbon than would be expected from surficial cuticle (G. D. Cody and C. K. Boyce, unpublished data), which was avoided during isotopic sampling.

The inconclusive isotopic results for *Rhynia* and *Asteroxylon* point out an important limitation: tissues can be complexly heterogeneous even at the millimeter scale of sampling. *Rhynia* possesses a distinct outer cortex similar to that of *Aglaophyton*, but because *Rhynia* stems are no more than 3–4 mm in diameter, only a single cortical measurement was possible. Incorporation of a varying ratio of inner and outer cortex in this single cortical measurement may account for some or all of the isotopic variability that persists in the largest *Rhynia* samples (fig. 6). If the isotopic variability is indeed caused by imprecise tissue sampling in *Rhynia*, this would suggest that chemical differentiation between tissues was present.

The cortex of Asteroxylon is even more problematic: lig-

nified leaf traces run throughout the cortex, so at this sampling scale, completely unlignified tissues are not available for comparison to the central stele. The aerenchymatous nature of *Asteroxylon* cortex (Edwards 1993) further complicates microsampling. The few samples available for *Asteroxylon* suggest that the cortex might be isotopically more negative than the lignified central xylem, perhaps due to isotopic depletions associated with the degradation of the aerenchymatous tissues. Without a finer scale of sampling (e.g., House 2000), however, this difference cannot be resolved.

X-ray spectromicroscopy demonstrates that two distinct organic components contribute to the tracheid cell walls of *Asteroxylon* but only a single component is found in the conducting cells of *Aglaophyton* and *Rhynia* (fig. 8). The relative



Fig. 7 Xylem analyzed with X-ray spectromicroscopy for (*A*) *Aglaophyton*, (*B*) *Rhynia*, and (*C*) *Asteroxylon*. Images in the left column are of the central vasculature of the fossil stems, and those on the right are of a peel of each fossil embedded in epoxy for X-ray analysis. The microtomed face from which the epoxy-embedded peels were sampled is on the right hand side of each peel image. In each case, only the xylem had sufficient organic matter to be distinguishable in the peel and to permit chemical analysis. Anatomical correlations between fossil and peel images are not exact due to subsequent polishing and sampling of the fossils. Scale bar is 1 mm.



Fig. 8 Stacked plots of carbon near edge X-ray absorption spectra of the vascular cell walls of Rhynie Chert taxa. Two distinct chemistries are found in *Asteroxylon*, but *Rhynia* and *Aglaophyton* share a single chemistry. Columns *a*–*d* correspond to absorption frequencies for different types of carbon bonds: *a*, aromatic C; *b*, aromatic C-OH; *c*, COOH; *d*, COH.

broadness of the aromatic 1s to π^* absorption bands in these spectra is indicative of a broad distribution of variously substituted aromatic molecules, consistent with a high degree of diagenetic alteration. However, the overall similarity of the spectra suggests similar original biochemistries subjected to the same taphonomic history. Because of this diagenetic alteration, X-ray microscopy alone cannot indicate whether the two chemically distinct components correspond to lignified and unlignified walls or lignified walls with different ratios of lignin and cellulose (Boyce et al. 2002). However, since isotopic evidence suggests that lignin was absent from conducting cells of *Aglaophyton*, observed differences between *Asteroxylon* and *Aglaophyton* organic chemistry may indicate that the two vascular organic signals of *Asteroxylon* represent a wall layer that is relatively heavily lignified and one that is unlignified or at most lightly lignified. Likewise, the close similarities of organic constituents in *Aglaophyton* and *Rhynia*, in conjunction with *Aglaophyton* isotopic data, suggest that the conducting cells of *Rhynia* were poorly lignified or perhaps even unlignified, despite the presence of tracheids with secondary thickenings. This evidence suggesting that S-type thickenings were poorly lignified provides support for the previous claims (Kenrick and Crane 1991; Kenrick et al. 1991*a*; Friedman and Cook 2000) that the unusual microstructure of rhyniophyte S-type tracheids results from more extensive degradation than is characteristic of eutracheophyte tracheids.

That said, the chemical evidence presented here is at odds with some taphonomic arguments for the distribution of lignin in Rhynie plants. The dark coloring of protracheophyte and rhyniophyte conducting cells has been taken as prima facie evidence for lignification (Edwards 1993; Kenrick and Crane 1997; Friedman and Cook 2000). The absence of corroborating chemical evidence, however, forces us to entertain the alternative explanation that dark color reflects the abundance rather than the chemical composition of preserved organic matter: dark conducting cells could have had relatively thick, but unlignified, primary walls. However, we cannot rule out the possibility that the differential preservation of conducting cell walls in rhyniophytes and protracheophytes was imparted by phenolic compounds present in amounts too small to influence isotopic composition.

As noted above, microchemical analyses of Rhynie fossils provide evidence that lignification may have originated in the peripheral tissues of protracheophytes and only later been coopted for the strengthening of tracheids in eutracheophytes. This perspective is also supported by anatomical and taphonomic arguments. The presence of partially decayed stems that maintain their cylindrical shape despite the complete loss of the inner tissues (Trewin 1994) is consistent with lignification of the outer cortex (although this feature perhaps does not rule out a lignified conducting strand; degradation of the surrounding inner cortex might be sufficient to disrupt the central strand regardless of wall chemistry). Positive evidence that these conducting cells were not lignified comes from anatomical studies of bryophytes: one of the characteristics of the unlignified, bryophyte water conducting cells is that, when on the periphery of a conducting strand, these dead cells have concave cell walls that respond to the pressure imposed on them by the turgor of adjacent living cells (Hébant 1977). The conducting cells of Aglaophyton and Rhynia show a similar response to the surrounding cells, suggesting their walls were not rigid (e.g., cf. figs. 340, 341 in Hébant 1977).

Evolution of the Tracheid Cell Type

An initial evolutionary independence of lignification and conducting cell morphology is reasonable on several grounds. First, although many living tracheophytes require lignified xylem to support plant mass, the thin central strand of conducting tissue in rhyniophytes would have contributed far less than turgor pressure to axis support (Speck and Vogellehner 1988). Moreover, while the discontinuous rings of secondary

Chemical Structures of Tracheid Cell Wall Derived from C-XANES Spectra						
Peak		eV	Asteroxylon			
	Transition		Layer A	Layer B	Rhynia	Aglaophyton
Aromatic carbon C _{AR} -C,H	$1s-\pi^{a}$	285.1	5.1ª (54%)	5.2 (49%)	5.0 (44%)	4.9 (42%)
Aromatic carbon C _{AR} -O	$1s-\pi^{a}$	287.2	0.9 (9%)	0.8 (7%)	1.0 (9%)	1.1 (9%)
Aliphatic carbon CH ₁₋₃	$1s - \sigma^{ab}$	287.8	0.3 (3%)	0.7 (7%)	1.1 (10%)	1.5 (13%)
Carboxylate COOR,H	$1s-\pi^{a}$	288.5	0.5 (5%)	0.5 (5%)	0.6 (5%)	0.5 (4%)
Aliphatic ether or alcohol						
Č _{AL} -O	$1s$ - σ^{ab}	289.5	2.7 (28%)	3.5 (33%)	3.7 (32%)	3.8 (32%)
No. oxygen atoms per						
aromatic ring			0.9	0.8	1.0	1.1

Table 3

^a Abundances cast in terms of number of carbons normalized to six aromatic carbons (i.e., a single aromatic ring) and in terms of percent of total carbon in parentheses.

^b The low energy $1s - \sigma^*$ transitions involve C-H bonding.

wall material in rhyniophytes could have helped prevent cell collapse under the negative pressure of water transport, they could not have supported the axis (reviewed in Bateman et al. 1998). If lignification was involved in the structural support of these fossil axes, it would optimally lie in the stem periphery (Niklas 1997), as is suggested for *Aglaophyton*.

Raven (1984) proposed that lignin's original function was antimicrobial and that only later did it come to play a structural role in xylem cells. Consistent with this hypothesis, phenolic compounds have been identified in charophyte algae (Delwiche et al. 1988) and bryophytes (Kroken et al. 1996). Raven's proposal implies that lignin in early land plants might have been concentrated along the periphery of axes, and the differential preservation of outer cortex in the Devonian fossil *Gosslingia* has been adduced as evidence for such a distribution (Kenrick and Edwards 1988). Thus, the evolution of lignified tracheids might have only involved the expression in a different cell type of a preexisting polyphenolic pathway and its targeted cell wall deposition.

The evolution of tracheid wall thickening independent of lignification is also consistent with xylem development in living plants. In the *Zinnia* model system of tracheid differentiation, patterned lignin deposition is dependent on the prior initiation of cellulose secondary thickenings and is disrupted by inhibitors of cellulose synthesis (reviewed in McCann 1997). This developmental pattern has been confirmed in pteridophytes with tracheid ultrastructure more similar to early fossils (Friedman and Cook 2000).

Mosiac evolution is not uncommon for plants (Knoll et al. 1984). Convergent evolution of form is also widespread (e.g., Boyce and Knoll 2002), and virtually all cell characteristics associated with tracheids have evolved repeatedly among the bryophytes. The prevalence of phenolic wall chemistry in peripheral tissues was mentioned above. Phenolic chemistry is also present in some bryophyte water conducting cells but only within derived polytrichalean mosses (Ligrone et al. 2000).

Cell elongation and anatomical variations in cell wall thickness are found in nearly all taxa. Helical, cellulosic wall thickenings similar to those of Rhynia tracheids occur in the elaters of liverworts and occasionally in hornwort columellae (Hébant 1977). Cell wall pitting is found in a variety of cell types, including conducting cells, in many taxa (Hébant 1977; Ligrone et al. 2000). Programmed cell death as part and parcel of cell function occurs in liverwort elaters, in water conducting cells, and in cortical cells of a variety of groups (Hébant 1977). Auxin gradients established by polar auxin transport are used for the patterning of vascular cell differentiation in tracheophytes (Sachs 1991), and similar gradients have been documented in liverworts (Maravalo 1980). What makes tracheids unique is that this entire suite of chemical, developmental, and morphological traits co-occur in a single cell type shaped by sequential evolutionary events in early land plants.

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