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The origin and early evolution of tracheids in vascular plants: integration of palaeobotanical and neobotanical data

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Although there is clear evidence for the establishment of terrestrial plant life by the end of the Ordovician, the fossil record indicates that land plants remained extremely small and structurally simple until the Late Silurian. Among the events associated with this first major radiation of land plants is the evolution of tracheids, complex water-conducting cells defined by the presence of lignified secondary cell wall thickenings. Recent palaeobotanical analyses indicate that Early Devonian tracheids appear to possess secondary cell wall thickenings composed of two distinct layers: a degradation-prone layer adjacent to the primary cell wall and a degradation-resistant (possibly lignified) layer next to the cell lumen. In order to understand better the early evolution of tracheids, developmental and comparative studies of key basal (and potentially plesiomorphic) extant vascular plants have been initiated. Ultrastructural analysis and enzyme degradation studies of wall structure (to approximate diagenetic alterations of fossil tracheid structure) have been conducted on basal members of each of the two major clades of extant vascular plants: Huperzia (Lycophytina) and Equisetum (Euphyllophytina). This research demonstrates that secondary cell walls of extant basal vascular plants include a degradation-prone layer ('template layer') and a degradation-resistant layer ('resistant layer'). This pattern of secondary cell wall formation in the water-conducting cells of extant vascular plants matches the pattern of wall thickenings in the tracheids of early fossil vascular plants and provides a key evolutionary link between tracheids of living vascular plants and those of their earliest fossil ancestors. Further studies of tracheid development and structure among basal extant vascular plants will lead to a more precise reconstruction of the early evolution of water-conducting tissues in land plants, and will add to the current limited knowledge of spatial, temporal and cytochemical aspects of cell wall formation in tracheary elements of vascular plants.

Keywords: tracheid; xylem; Huperzia; Equisetum; cell wall; developmental evolution

1. INTRODUCTION

Due of the most significant sets of evolutionary events in he history of life on Earth was the migration of complex fe forms from water-based environments to land in the harly Palaeozoic. These events began with the migration f aquatic photosynthetic organisms on to land some 75 Myr ago and resulted in a veritable explosion of volutionary innovation and consequent diversification of errestrial ecosystems.

Prior to the input of significant energy into terrestrial cosystems by land plants (embryophytes), terrestrial nvironments were occupied by various heterotrophic and utotrophic bacteria, protists, fungi, lichens and some mple algae (Gray & Shear 1992; Gray 1993; Knoll 1994; Cenrick & Crane 1997*a*,*b*). Within 75 Myr of the origin of and plants, terrestrial photosynthetic organisms (embryohytes) had undergone a major evolutionary radiation and stablished conditions for the colonization and subsequent iversification of various metazoan lineages on land. Some 50 Myr after the origin of land plants, the surface of the

Earth was dominated by the highly diverse Carboniferous forest ecosystems—a radical change from the pre-Ordovician bacterial, fungal and algal biotic crusts.

Photosynthesis in an aerial environment requires high levels of gas exchange (uptake of CO_2) and is facilitated in the sporophytes of all land plants, with the exception of liverworts, by the formation of stomatal openings in the outer surface of the plant body. These pores, however, also result in significant losses of gas-phase water from the internal tissues of the plant body to the external environment. As a consequence, to remain hydrated terrestrial plants larger than a few centimetres require specialized tissues to transport water from the plant-soil interface to the aerial portions of the plant body (Raven 1993). Biophysical models clearly indicate (Raven 1984, 1993) that any significant increase in the stature of land plants was predicated on the evolution of highly specialized hollow and dead water-conducting cells, whose rates of conductance have been calculated to be 1×10^7 times greater than equivalent living cells with cytoplasmic contents (Raven 1984, 1993).

Euphyllophytina



igure 1. Phylogenetic relationships of land plants. Evidence ased on intron distribution indicates that liverworts are the ster group to all other land plants (Qiu et al. 1998). The terrelationship of hornworts and mosses is unresolved. glaophyton does not have tracheids with secondary hickenings, and is hypothesized to be a close (extinct) relative f tracheophytes or vascular plants. Rhyniopsida (all extinct) re hypothesized to be the sister group to the Lycophytina nd Euphyllophytina, which together comprise the utracheophytes.

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It has long been known that certain members of the verworts and mosses possess elongate, non-lignified cells hat are dead at maturity and may function in water onduction (Hébant 1977; Scheirer 1980; Mishler & hurchill 1984, 1985; Ligrone & Duckett 1996; Renzaglia t al. 1997; Ligrone et al., this issue»). However, much emains to be learned of the structure, development, funcion and evolutionary homologies of these cells. What is bundantly clear is that the fossil record documents that and plants remained extremely small and structurally imple until the Late Silurian, ca. 425 Myr ago (Gensel & undrews 1987; Gray & Shear 1992). This first burst of ructural diversification among land plants was assoiated with the evolution of tracheids (Knoll & Rothwell 🕖 981; Knoll & Niklas 1987; Raven 1993; Kenrick & Crane 997a), developmentally complex water-conducting cells efined by the presence of secondary cell walls, lignificaion and programmed cell death.

2. TRACHEIDS OF THE EARLIEST VASCULAR PLANTS

Recent phylogenetic analyses indicate that tracheidearing plants (tracheophytes) are monophyletic (figure 1) nd defined by the presence of secondary cell wall thickenhgs in water-conducting cells (Kenrick & Crane 1997a, b). volutionary biologists (Banks 1975; Gensel & Andrews 987; Taylor & Taylor 1993; Kenrick & Crane 1997*a*,*b*)

have hypothesized that a rapid diversification among early vascular plants (tracheophytes) produced three major clades (figure 1), each of which is characterized by a particular tracheid type among its earliest members (Kenrick & Crane 1997*a*,*b*).

Rhyniopsida is hypothesized to be the sister group to a monophyletic eutracheophyte clade that includes all extant vascular plants, as well as many of their extinct relatives (figure 1; Kenrick & Crane (1997a,b) and references therein). Members of the Rhyniopsida, all of which are extinct, are characterized by the presence of S-type tracheids (after the genus Sennicaulis). Two monophyletic lineages have been recognized within the eutracheophyte clade (figure 1): the Lycophytina (lycophytes and their extinct ancestors, the zosterophylls), whose earliest members have G-type tracheids (after the zosterophyll genus Gosslingia); and the Euphyllophytina (the extinct trimerophytes, eusporangiate ferns, leptosporangiate ferns, sphenopsids, Psilotaceae, progymnosperms and seed plants), whose earliest (trimerophyte) members have P-type tracheids (after the genus Psilophyton).

S-type tracheids of rhyniopsids have annular or helical thickenings and lateral walls that appear to be made of a spongy or reticulate material (figure 2a-c). It has been suggested that the 'pockets' in the spongy wall material may represent unlignified portions that were preferentially degraded during fossilization (Kenrick & Crane 1991; Kenrick et al. 1991). A very thin degradationresistant layer of cell wall material with micropores covers the entire spongy layer of wall material adjacent to the cell lumen (figure 2; Kenrick et al. 1991; Kenrick & Crane 1991, 1997a, b). It is unclear whether the micropores are a preservational artefact (i.e. small pockets of unlignified wall material that were preferentially degraded during fossilization) or were a real structural component of this innermost wall layer.

G-type tracheids of early zosterophylls have annular or helical secondary cell wall thickenings that exhibit two layers: a carbonaceous dark layer closest to the cell lumen, and a light layer that appears to represent a mineralized hollow core of each gyre (figure 2d-f) (Kenrick & Edwards 1988; Kenrick & Crane 1991; Kenrick et al. 1991). The hollow core in wall thickenings of these fossil cells may represent an unlignified portion of the thickening that was preferentially degraded during fossilization (Kenrick & Edwards 1988; Kenrick & Crane 1991). Between gyres of secondary thickenings, lateral walls in G-type tracheids exhibit holes ranging in size from less than 1 µm to 4 µm (Kenrick & Edwards 1988). These holes have been interpreted as perforations in the primary wall (Hartman 1981; Hueber 1983; Rayner 1984) or as small pits in a layer of lignified secondary wall that is continuous with the degradationresistant dark layer of the annular or helical thickenings (Brauer 1980; Taylor 1986; Kenrick & Edwards 1988; Edwards 1993; Kenrick et al. 1991; Kenrick & Crane 1991, 1997a). Alternatively, these holes could represent small pockets of non-lignified cell wall material that were degraded and lost during the process of fossilization.

P-type tracheids (Kenrick & Crane 1997a, b) are found in early euphyllophytes, and have bordered pits (figure 2g-i). Strands of secondary wall material traverse the pit



igure 2. The three major types of early tracheids. (a) Reconstruction of Rhynia, a member of the Rhyniopsida that produced -type water-conducting cells. Reproduced from Kenrick & Crane (1997a). Copyright permission of the Smithsonian Institution 🖳 ress. (b) Drawing of the wall of an S-type water-conducting cell showing the wall thickenings with a spongy (alveolate) structure nd a thin microporate surface on the lumen face of the tracheid. Reproduced from Kenrick & Crane (1997a). Copyright ermission of the Smithsonian Institution Press. (c) SEM of a single gyre from an S-type cell of Sennicaulis. Note the spongy wall 🔿 onstruction within the gyre at left, and the microporate nature of the inner surface of the tracheid. Reproduced from Kenrick et 🟹!. (1991). Copyright permission of the Palaeontological Association. (d) Reconstruction of Zosterophyllum, a member of the ycophytina that produced G-type water-conducting cells. Reproduced from Kenrick & Crane (1997a). Copyright permission of ne Smithsonian Institution Press. (e) Drawing of the wall of a G-type water-conducting cell showing the wall thickenings that tck material in the core and holes in the surface of the lumen face of the tracheid. Reproduced from Kenrick & Crane (1997a). topyright permission of the Smithsonian Institution Press. (f) SEM of a portion of a G-type cell of Gosslingia. Note the spongy all construction within the gyre at left, and the microporate nature of the inner surface of the tracheid. Reproduced from Cenrick & Crane (1997a). Copyright permission of the Smithsonian Institution Press. (g) Reconstruction of Psilophyton, a nember of the Euphyllophytina that produced P-type water-conducting cells. Reproduced from Kenrick & Crane (1997a). opyright permission of the Smithsonian Institution Press. (h) Drawing of the wall of a P-type water-conducting cell showing the all thickenings that lack material in the core and continuity of the wall material that overlies the pits and has holes. Reproduced from Kenrick & Crane (1997a). Copyright permission of the Smithsonian Institution Press. (i) SEM of a portion of a P-type cell f Psilophyton. Note the secondary cell wall thickenings appear hollow (arrow). Reproduced from Hartman & Banks (1980). opyright permission of the Botanical Society of America.

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pertures and connect the pit borders that overlie the pit avity (Gensel 1979; Hartman & Banks 1980). Holes ccur in this area of additional secondary cell wall ornanentation (Gensel 1979; Hartman & Banks 1980; Cenrick & Crane 1997*a*,*b*). In the fossil record, P-type racheids are similar to G-type cells in possessing econdary wall thickenings (between the pits) that appear ollow (figure 2).

Although various aspects of secondary cell wall atterning differ between fossil S-, G- and P-type traceids (figure 2), all of these early water-conducting cells ossess secondary cell wall thickenings composed of two istinct layers: a degradation-resistant (possibly lignified) →ayer next to the cell lumen and a degradation-prone 🗳 ayer closest to the primary cell wall (Kenrick & dwards 1988; Kenrick & Crane 1991; Kenrick et al. 1991; Udwards 1993). S-, G- and P-type tracheids may all be Ovolutionarily homologous and represent developmental ransformations of a rudimentary tracheid type from a ommon ancestor of all vascular plants (Kenrick & Irane 1991 1997a,b; Cook & Friedman 1998). Alternaively, there could have been two or three separate origins f water-conducting cells with secondary wall thickenings i.e. 'tracheids' are homoplasious) (Kenrick & Crane 991).

3. TRACHEARY ELEMENTS OF EXTANT VASCULAR PLANTS

Until quite recently (Cook and Friedman 1998), all tudies of tracheary element (tracheid and vessel element) ifferentiation and fine structure had been conducted on ighly derived vascular plants, namely conifers and flowring plants (Esau *et al.* 1963, 1966*a,b*; Wooding & Northote 1964; Cronshaw & Bouck 1965; O'Brien & Thimann 967; Hepler & Fosket 1970; Esau 1978; Daniel & Nilsson 984; Uehara & Hogetsu 1993; Fineran 1997). As a result, urrent biochemical and cell biological models of racheary element development (Boudet *et al.* 1995; Jarceló 1997) are based solely on seed plants (Cook & riedman 1998).

Electron micrographs of tracheary elements in conifers nd angiosperms depict secondary wall thickenings that re essentially homogeneous. Although seed plant traceids typically have a three-layered secondary cell wall S1, S2, and S3 layers), these layers of cell wall are all eavily lignified and differ mostly in the orientation angle) of microfibril deposition (Esau 1977; Boudet et al. 995). In seed plants that have been studied, lignification Uegins at the cell periphery and gradually progresses owards the cell lumen as centripetal wall deposition rogresses (Hepler & Fosket 1970; Liu et al. 1994; Boudet t al. 1995; Barceló 1997). Most significantly, tracheary lements of extant seed plants are not known to exhibit a egradation-prone (possibly unlignified) layer of cell wall naterial between an outer wall and an innermost egradation-resistant layer of cell wall, as is characteristic f tracheids in early tracheophytes. Thus, despite major rogress in reconstructing the structural diversity of early ascular plant tracheids, interpretation of the develophental and evolutionary relationships between these early ossil tracheids and those of extant vascular plants has emained uncertain (Cook & Friedman 1998).

Patterns of tracheid differentiation and mature fine structure are virtually unknown in basal vascular plants (i.e. the pteridophytes). Most basic information on tracheid secondary wall patterning in pteridophytes can be traced to the studies of Bierhorst (1958, 1960), Wilder (1970), Morrow & Dute (1997), Carlquist & Schneider (1997a,b, 1998a,b, 1999), Schneider & Carlquist (1997, 1998a, b, 1999a, b) and Carlquist et al. (1999). Interestingly, Bierhorst reported an 'unlignified or very faintly lignified' core at the base of secondary cell wall thickenings in Lycopodium, Equisetum, Psilotum and various ferns (Bierhorst 1958, 1960). Unfortunately, photomicrographs of this supposedly unlignified core in the tracheid walls of primitive vascular plants were never published, so it is difficult to evaluate these observations. Nevertheless, Bierhorst's reports remained intriguing and several palaeobotanists suggested a relationship between the purported 'unlignified core' and the missing layers of wall material in early fossil tracheids (Brauer 1980; Taylor 1986; Kenrick & Crane 1991; Kenrick et al. 1991).

Comparative studies of basal and potentially plesiomorphic vascular plants are essential if we are to understand better the early evolution of tracheids. Knowledge of the relationship between development and mature tracheid wall structure in primitive extant vascular plants can be used to aid in the interpretation of the diverse tracheid wall structures of early fossil vascular plants. Indeed, recent (Cook & Friedman 1998) and ongoing developmental studies of tracheid wall structure in basal extant vascular plants have begun to yield information on critical intermediate character states that link the structures of water-conducting cells of extant vascular plants to those of their 400 Myr-old ancestors. In addition, this research is beginning to provide the requisite information for an explicit model for the evolution of cellular differentiation patterns that could have produced the tracheids of the earliest vascular plants and their extant evolutionary descendants.

4. BACKGROUND PHYLOGENETIC INFORMATION

Phylogenetic analyses indicate that extant vascular plants are distributed in two major sister clades (Raubeson & Jansen 1992), the Lycophytina and the Euphyllophytina (figure 3). Among extant Lycophytina (Lycopodiaceae, Selaginellaceae, Isoetaceae), homosporous Lycopodiaceae are sister to Selaginellaceae plus Isoetaceae (figure 3) (Therrien & Haufler 1997); while within the Lycopodiaceae, terrestrial (non-epiphytic) species of Huperzia are basal (Wagner & Beitel 1992; Wikström & Kenrick 1997). Among members of the Euphyllophytina, the phylogenetic positions of sphenopsids (Equisetum and its extinct relatives), Psilotaceae (Psilotum and Tmesipteris), Marattiales, Ophioglossales and Filicales (leptosporangiate ferns) remain unresolved (figure 3), although Psilotaceae may be closely related to Ophioglossales, a result found in several recent phylogenetic analyses (Manhart 1994, 1995; Hasebe et al. 1995; Pryer et al. 1995; Wolf 1997).

To reconstruct the characters associated with the common ancestors of extant vascular plants, it is important to study basal representatives of both the Lycophytina and the Euphyllophytina. Homosporous lycopods

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igure 3. Current hypothesis of phylogenetic relationships mong extant vascular plants. See text for additional iformation.

Ich as *Huperzia* may well provide critical information bout the plesiomorphic structural features of the ommon ancestors of extant lycophytes. Given the uncerainties of euphyllophyte interrelationships, information n tracheid structure and development from *Equisetum*, *Silotum* and *Ophioglossum* (or *Botrychium*) will be critical to econstructing those features of tracheids that are likely b have characterized the common ancestors of extant uphyllophytes.

5. TRACHEID DEVELOPMENT AND STRUCTURE IN HUPERZIA

Huperzia lucidula, a member of the most basal extant lade within the Lycopodiaceae, is highly plesiomorphic mong vascular plants and can be considered a true ving fossil. Morphologically, it is almost indistinguishble from the Lower Devonian fossil lycopod Drepanohycus. It has recently been discovered that secondary cell all structure in tracheids of *Huperzia* is significantly ifferent from what is known of 'model systems' among onifers and angiosperms (Cook & Friedman 1998). fore importantly, there is compelling evidence (see elow) that tracheid secondary cell wall structure in *Iuperzia* shows a high degree of structural correspondence () the secondary wall thickenings of early fossil tracheids. In Huperzia, like many lycophytes, the vascular ylinder of the stem takes the form of an actinostele, with rotoxylem at the ends of xylem arms and phloem and arenchyma located between the arms of xylem. Protoylem elements are few and small in diameter, in comparon with metaxylem elements. At maturity, protoxylem cacheids of stems exhibit annular secondary wall thick- \bigcirc nings that are occasionally interconnected via vertical or anted thickenings, forming a loose reticulum (figure 4a). hort, helical thickenings are sometimes seen in cells with nnular thickenings. Longitudinal sections of metaxylem acheids demonstrate that bordered pits of various sizes, hapes and patterns appear in secondary walls. Early

metaxylem elements are small in diameter and possess circular or oval pits with borders. Oval or elongate pits are found in later-formed tracheids of larger diameter (figure 4b).

Differentiation of lateral walls in protoxylem and metaxylem tracheids of *Huperzia* involves three discrete stages of cell wall deposition, each of which produces a cell wall layer with distinct properties (Cook & Friedman 1998). As is true for all plants, cell wall formation is centripetal. The first layer to mature is the primary cell wall, which is smooth, homogeneous, and assumes a light grey appearance under the transmission electron microscope (TEM). Deposition of the primary cell wall is completed before synthesis of the two layers that compose the secondary cell wall is initiated.

Secondary cell wall deposition begins at the lumen surface of the primary cell wall. A first-formed layer of secondary cell wall is deposited over the surface of the primary cell wall (figure 4c, e), except in areas that will develop into pit membranes (metaxylem) or that lie between gyres of secondary wall material (protoxylem). In Huperzia, the first-formed secondary cell wall layer determines the pattern of further secondary cell wall deposition and has been named the 'template layer' (Cook & Friedman 1998). The template layer exhibits dark, mottled staining under the TEM (figure 4), which matches the contents of dictyosome-derived vesicles (figure 4c) that fuse with the plasmalemma and contribute to the synthesis of this layer of secondary cell wall. Each vesicle contains a single, electron-opaque particle, and it is likely that this structure is directly related to the electron-opaque particles found within the template layer (figure 4d, f).

After deposition of the template layer of secondary cell wall is completed in *Huperzia*, an additional and structurally distinct layer of secondary cell wall (the 'resistant layer') is deposited on the lumen surface of the template layer. This later-formed layer is first discernible next to the cell lumen as a thin, very lightly stained layer of newly synthesized cell wall material (figure 4g). The TEM shows that dictyosome-derived vesicles (figure 4d) that appear to be contributing to this second phase of secondary wall formation differ markedly from those associated with the synthesis of the template layer. The contents of these vesicles lack the electron-opaque particles associated with formation of the template layer. As the later-formed resistant wall layer continues to increase in thickness (figure 4g-i) and to mature, it stains more darkly.

Cell autolysis completes tracheid differentiation in Huperzia (figure 4j). At maturity, three distinct layers of cell wall can be discerned in Huperzia lucidula: a homogeneous primary cell wall; a mottled, heterogeneous template layer that covers much of the primary cell wall; and a homogeneous layer of secondary cell wall that overlies the template layer. The structural distinctness of the two layers of secondary thickenings in Huperzia is also apparent in longitudinal sections of water-conducting cells that have been prepared for scanning electron microscopy (SEM) (figure 5). Often, as a result of mechanical stress associated with microtomy, the inner resistant layer of the secondary cell wall becomes physically detached from the template layer, which remains attached to the underlying primary cell wall (figure 5).

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igure 4. (a) SEM of a longitudinal section of protoxylem elements of a stem of *Huperzia*. (b) SEM of a longitudinal section of netaxylem elements of a stem of *Huperzia*. Early metaxylem is to the left (reticulate secondary cell walls). (c) Earliest stage of econdary cell wall deposition on to the primary cell wall (P) in *Huperzia*. The budding face of a Golgi body (G) can be seen, s well as vesicles apparently fusing with the cell wall to form the template layer (T). These vesicles each contain a single lectron-dense particle that appears to be incorporated into the template layer. (d) Deposition of the resistant layer (R) of econdary cell wall material in a tracheid of *Huperzia*. Note that the vesicles derived from Golgi bodies are now electronranslucent. (e-j) Development of secondary wall thickenings in *Huperzia* (longitudinal views). A first-formed layer of secondary vall, the template layer, is deposited on the primary cell wall (e, f). Subsequently, a later-formed layer of secondary wall, the esistant layer, is deposited on the surface of the template layer (g-i). The resistant layer appears unstained or very lightly tained when first deposited, but later appears grey. In mature, dead tracheids (j), the template layer is distinct from the esistant layer of secondary cell wall, and can be recognized by the inclusion of electron-dense particles.

Huperzia remains the sole extant basal vascular lant for which there is ultrastructural developmental nformation on cell wall formation in tracheids. urther developmental studies of other lycophytes and arious basal euphyllophytes will be central to deterning how general the pattern of secondary cell wall prmation in *Huperzia* may be among extant vascular lants.

6. EXPERIMENTAL CELL WALL DEGRADATION IN TRACHEIDS IN HUPERZIA AND EQUISETUM

S-, G- and P-type tracheids preserved in the Silurian and Early Devonian were invariably subject to diagenetic alterations. During the time between plant death and permanent incorporation in the fossil record, plant parts are often attacked by a variety of cell-wall-degrading

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igure 5. (a,b) SEM images of the secondary thickenings of racheids in *Huperzia* in which the resistant layer (**R**) of the econdary cell wall has become physically separated from the emplate layer (arrows). The primary cell wall (**P**) can be een in both cells.

nzymes of fungi and bacteria. To simulate degradation rocesses and patterns experienced by fossilized cells, *Iuperzia* stem segments have been experimentally treated rith cell-wall-degrading enzymes (Cook & Friedman 998). Although it is not possible to replicate exactly the onditions of preservation in the fossil record, enzyme egradation studies can help determine if there is correpondence between the different layers of cell walls of racheids in extant *Huperzia* and the reported patterns of ifferential tracheid wall preservation in early vascular lants.

Stem segments 2 mm long were immersed for two to our weeks in an aqueous solution of 2% pectinase (from he fungus *Rhizopus*) and 2% cellulase (from the fungus enicillium), rinsed in water and treated for 2 h in aqueous % osmium tetroxide, before usual preparation for elecon microscopy. These experiments demonstrated that he two layers of secondary cell wall material in tracheids f Huperzia are distinct chemically, as well as structurally nd developmentally. The primary cell wall and firstormed layer of secondary cell wall (at the cell periphery) re significantly degraded by a mixture of pectinase and ellulase. After enzyme treatment, the primary cell wall nd the first-formed layer of secondary cell wall (template yer) were either entirely missing or were represented by delicate reticulum of remaining cell wall material $\prod_{i=1}^{n}$ figure 6b). The later-formed secondary cell wall layer next to the cell lumen) was not degraded by enzyme Creatment. This portion of the secondary wall in Huperzia as been named the 'resistant layer' (Cook & Friedman 998).

Experimental evidence for a degradation-prone emplate layer in *Huperzia* (Cook and Friedman 1998), is ne first conclusive report for this type of secondary cell 'all organization in an extant vascular plant. Cell wall egradation experiments have now been extended to the racheids of *Equisetum*. When subjected to a solution of ectinase and cellulase, secondary thickenings in *Equi*tum tracheids show clear evidence of degradation of the rimary cell wall and the base of the secondary cell wall figure 7). An overlying cap of secondary cell wall shaped much like a mushroom in sectional view)



Figure 6. Comparison of wall layers in tracheids of Huperzia. (a) Traditional TEM view (control) of a single secondary cell wall thickening with a template layer (T) and a resistant layer (R). The primary cell wall (P) is at the right. (b) In cells subjected to enzyme treatment (see text), the template layer has been degraded and appears reticulate, while the resistant layer remains intact. The primary cell wall is largely degraded and is not apparent. (c) Secondary cell wall of tracheid which has been post-stained with potassium permanganate, an indicator of the presence of lignin. The primary cell wall appears to lack significant amounts of lignin, while the template layer is partially stained, suggesting the presence of some lignin. The resistant layer is heavily stained, indicating the presence of large amounts of lignin. (d) Secondary cell wall of tracheid which has been subjected to enzyme treatment and then analysed cytochemically for the presence of lignin. The resistant layer is intact and heavily lignified. The template layer is largely degraded, but shows signs of a reticulate distribution of lignin.

adjacent to the lumen does not show evidence of degradation by a combination of cellulase and pectinase. These experiments provide, for the first time, compelling evidence that a degradation-prone (template) layer and a degradation-resistant layer are present in the secondary thickenings of tracheids in a member of the euphyllophyte clade.

The clear implication of these results from *Equisetum*, coupled with the recent findings in *Huperzia* (Cook & Friedman 1998), is that the common ancestor of the Lycophytina and Euphyllophytina is likely to have produced tracheids with secondary cell wall thickenings, comprised of an underlying degradation-prone template layer and a subsequently formed degradation-resistant layer.

7. CYTOCHEMICAL LOCALIZATION OF LIGNIN IN THE TRACHEIDS OF *HUPERZIA*

Lignin is one of the most degradation-resistant biopolymers on Earth (Graham 1993). This suggests that those

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igure 7. Secondary cell wall thickenings in the tracheids of *quisetum.* (a) Traditional TEM view (control) of a single econdary cell wall thickening. The primary cell wall (P) is on he right. (b) In cells subjected to enzyme treatment (see ext), a template layer (T) is apparent and has been partially egraded. A resistant layer (R) is also quite clearly seen. The primary cell wall is partially degraded. These data idicate that at least in one member of the euphyllophyte lade, an organization of secondary cell walls may involve

first-formed partially lignified template layer and a ater-formed heavily lignified resistant layer.

ortions of the cell walls of early fossil tracheids that were ot degraded during preservation in the Silurian and Devonian may have been heavily lignified. We sought to etermine whether there is a correspondence between the esults of our laboratory-based enzyme degradation tudies and patterns of lignin distribution within the cell valls of tracheids of extant basal vascular plants.

Potassium permanganate has been widely used as a xative or a post-fixation stain to distinguish lignified cell valls from those that are unlignified (Hepler & Fosket 970; Barceló 1997; Fineran 1997). Thin sections of nzyme-degraded *Huperzia* tracheids were post-stained vith permanganate and examined under the TEM for ytochemical analysis of lignin distribution. The resistant ayer of secondary wall stained darkly and homoge-eously, while the template layer stained in a reticulate nanner (figure 6d). What remains of the primary wall ppears to contain little, if any, lignin.

To cross-correlate cytochemical data for enzymeegraded cells with untreated tracheids, TEM-level cytohemical localization of lignin in *Huperzia* was performed based on a modification of the 'Coppick and Fowler' nethod, see Fineran (1997) for methods) on intact traceids (figure 6c). These data indicate that the resistant ayer is heavily lignified, that the template layer is artially lignified in a reticulate manner, and that the rimary cell wall of *Huperzia* tracheids appears to lack ignificant quantities of lignin.

8. RELATIONSHIPS BETWEEN THE TRACHEIDS OF EXTANT BASAL VASCULAR PLANTS AND THE EARLIEST FOSSIL VASCULAR PLANTS

Extant lycopods are more closely related to zosterohylls and fossil lycopods than they are to other vascular lants (figure 1). Therefore, one might expect tracheids of *luperzia* to be more similar to the G-type cells typical of osterophylls and early fossil lycopods than to fossil Snd P-type cells or the tracheids of other extant plants. A rominent feature of G-type cells is the presence of holes a resistant lateral walls between annular or helical secondary thickenings (figure 8). These holes have been interpreted as perforations in the primary wall (Hartman 1981; Hueber 1983; Rayner 1984) or as small pits in a layer of lignified secondary wall that is continuous with the degradation-resistant dark layer of the annular or helical thickenings (Brauer 1980; Taylor 1986; Kenrick & Edwards 1988; Edwards 1993; Kenrick *et al.* 1991; Kenrick & Crane 1991, 1997*a*). Tracheids of *Huperzia* do not possess holes in lateral secondary walls comparable with those between thickenings in G-type tracheids (figure 8).

The biochemical nature of the template layer in Huperzia, its susceptibility to degradation and its position at the base of secondary cell wall thickenings strongly suggest developmental and structural homology with the degradation-prone wall layer in the secondary cell wall thickenings of the G-type tracheids characteristic of early members of the Lycophytina. For many years, palaeobotanists suggested that the absent layer of wall material in the secondary cell walls of early fossil tracheids represents an unlignified or poorly lignified wall layer that did not survive the fossilization process (Kenrick & Crane 1991, 1997a,b; Kenrick & Edwards 1988; Brauer 1980). Documentation of a degradation-prone, partially lignified template layer and a heavily lignified resistant layer in the secondary cell wall thickenings of Huperzia (Cook & Friedman 1998) provides a critical link between the water-conducting cells of Late Silurian and Early Devonian lycophytes and those of their extant descendants (figure 8).

Structural correspondence of secondary cell wall layers of basal extant vascular plant tracheids can be extended beyond the ancestral G-type cells of the early Lycophytina to the P-type cells representative of the earliest members of the Euphyllophytina (the sister group to lycophytes). Secondary cell wall thickenings of fossil P-type tracheids also have an absent core layer (figure 8; Hartman & Banks 1980; Edwards 1993). Results of experimental degradation studies of tracheids of extant Equisetum show a degradation-prone layer is present in the secondary thickenings that positionally matches the template layer of Huperzia and underlies a resistant layer (figure 8). The clear implication of these findings is that the common ancestor of lycophytes and euphyllophytes (together the eutracheophyte clade) produced tracheids with secondary thickenings composed of a first-formed poorly lignified template layer that was susceptible to degradation during fossilization and a second-formed heavily lignified resistant layer that is still present in 390 Myr-old tracheids.

Many features of development of tracheary element walls at the cellular level appear to be the same in *Huperzia* and seed plants: the primary wall reaches maximum thickness before secondary wall layers are laid down; microtubules, endoplasmic reticulum and dictyosomes line up along the forming secondary wall; dictyosome-derived vesicles fuse with the plasma membrane near the developing secondary thickenings; the cell undergoes programmed cell death (Baucher *et al.* 1998). Developmental differences do exist, however, between tracheary elements of seed plants and those of *Huperzia*. Most significant is that in those seed plants studied, secondary cell wall thickenings of mature tracheary elements appear homogeneous (figure 8) when Downloaded from rstb.royalsocietypublishing.org The origin and evolution of tracheids in vascular plants W. E. Friedman and M. E. Cook 865



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igure 8. Schematic of longitudinal view of cell wall thickenings in fossil S-, G- and P-type tracheids and in tracheids of *Huperzia*, *Quisetum* and extant angiosperms (and conifers). The initial diversification of vascular plants is hypothesized to have produced n early divergent Rhyniopsida (all extinct) with S-type tracheids; the Lycophytina (lycopsids and zosterophylls) with G-type acheids (in Early Devonian members); and the Euphyllophytina (all other vascular plants), with P-type tracheids (in Early Devonian members). The cell lumen is to the right in each diagram. Cell wall thickenings in S-type cells have a thick, partially esistant spongy layer that is covered by a thin microporate resistant layer adjacent to the cell lumen. Cell wall thickenings in i - and P-type cells appear to have a core of degradation-prone wall material that is covered by a thick, resistant layer adjacent o the cell lumen. The degradation-prone layer of G- and P-type cells is absent from tracheids. In P-type tracheids holes are parent in a layer of degradation-resistant cell wall that overlies pits. In *Huperzia* and *Equisetum*, secondary cell wall thickenings of extant seed plant tracheary elements appear to be homogeneous and lack any quivalent to a degradation-prone layer.

iewed with light microscopy and TEM. (This also ppears to be true of secondary cell wall thickenings in *Quisetum*, even though degradation studies indicated that ne distribution of lignin is not homogeneous.) Seed lants typically have secondary cell walls with three ayers (S1, S2 and S3) that differ in the orientation of ellulose microfibril deposition, but are all heavily lignied (Boudet *et al.* 1995). Lignification begins at the cell eriphery (middle lamella and primary cell wall) and rogresses towards the cell lumen as centripetal wall eposition continues (Boudet *et al.* 1995; Barceló 1997). In ontrast, in *Huperzia* and *Equisetum*, lignification appears reak or absent in the primary cell wall, heterogeneous nd somewhat more prominent in the template layer, and heaviest in the layer closest to the cell lumen.

Ultimately it will be essential to study the developent, biochemical nature and degradation properties of ther extant euphyllophyte clades, such as Psilotaceae, phioglossales, Marattiales, leptosporangiate ferns, *Ginkgo* nd cycads, to determine how widespread the pattern of racheid wall structure seen in *Huperzia* and *Equisetum* is mong extant vascular plants. Given that conifers and ngiosperms appear to lack a template layer similar to nat which has been found in basal vascular plants, phyloenetically based comparative analyses of tracheid ructure among diverse euphyllophyte clades will be eccessary to address the question of whether the template and rest of secondary thickenings in tracheary elements was pst prior to the origin of seed plants or within seed lants.

Developmental analysis and enzyme degradation experiments also suggest a possible connection between the tracheids of Huperzia and Equisetum, and the more primitive fossil S-type cells. It has been suggested that the spongy layer of S-type cell walls survived in the fossil record because it was partially resistant to degradation, perhaps due to thin veins of resistant biopolymers running through a mass of less resistant material (figure 8; Kenrick et al. 1991). Similarly, the template layers of Huperzia and Equisetum do not completely disappear under conditions of experimental degradation, but instead yield a reticulate condition that bears considerable resemblance (figures 6b and 7b) to the spongy wall layer of S-type cells (figure 2c). Thus, the template layer of Huperzia and Equisetum, as well as the missing core of secondary cell wall in P-type and G-type tracheids, may be structurally and evolutionarily homologous to the spongy layer of S-type cells.

9. WORKING HYPOTHESIS FOR THE EARLY EVOLUTION OF TRACHEIDS

Documentation of a degradation-prone template layer and a degradation-resistant layer in the secondary cell wall thickenings of the primitive vascular plants *Huperzia* and *Equisetum* provides the critical link between the water-conducting cells of Early Devonian tracheophytes and those of their extant descendants. While the secondary cell wall thickenings of tracheids of previously studied extant vascular plants (Esau *et al.* 1963, 1966*a,b*; PHILOSOPHICAL TRANSACTIONS

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Vooding & Northcote 1964; Cronshaw & Bouck 1965;)'Brien & Thimann 1967; Hepler & Fosket 1970; Esau 978; Daniel & Nilsson 1984; Uehara & Hogetsu 1993; 'ineran 1997) appear to differ significantly from those of arly fossil vascular plants, developmental data from *Iuperzia*, and the results of enzyme degradation studies in *Iuperzia* and *Equisetum*, demonstrate a clear structural orrespondence between the tracheids of extinct and xtant primitive vascular plants. Moreover, tracheid evelopment in *Huperzia* provides the essential informaion needed to propose an explicit model for the evolution f secondary wall thickenings in vascular plants.

The evolution of complex water-conducting cells is nlikely to have occurred in a single step. Rather, racheids almost certainly evolved through a series of novations and modifications of development. Although recapitulation' explanations for the evolution of organsmal development are frequently problematic (Alberch & Glanco 1996 and references therein), the developmental

equence of secondary cell wall patterning associated with racheid development in the basal lycopod *Huperzia* (and lmost certainly in *Equisetum*, a basal euphyllophyte) may ndeed recapitulate aspects of the developmental evoluion of tracheids in early vascular plants.

Our developmental data are congruent with the ypothesis that evolution of water-conducting cells in and plants commenced with the programmed autolysis of ells. We hypothesize that the origin of secondary cell vall thickenings in tracheids began with deposition of a vartially degradation-resistant template layer of cell wall n to the inner surface of the primary cell wall prior to ell autolysis. A further innovation, a more heavily lignied resistant layer, was added to the process of secondary ell wall deposition, and this secondary cell wall organzation can be found as a thin, innermost (next to the umen) layer in mature S-type conducting cells of thyniopsida.

At some point in the Late Silurian or Early Devonian, nodification of the process of tracheid cell wall eposition led to a decrease in the proportion of the egradation-prone template layer and an increase in the roportion of the resistant layer within secondary cell valls, as found in G- and P-type fossil tracheids. The esistant layer represents 2% of the thickness of econdary wall thickenings in S-type cells, 30% in Gnd P-type cells (Cook & Friedman 1998), and well over 0% in extant Huperzia and Equisetum. This trend towards increasing amounts of degradation-resistant (lignified) econdary cell wall material has previously been noted UKenrick & Edwards 1988; Kenrick & Crane 1997a). continuation of the trend towards reduction of the Semplate layer and augmentation of the resistant layer has roduced the characteristic secondary thickenings of traceids in extant seed plants, in which no equivalent of a emplate layer has been reported and the entire econdary wall is highly lignified and resistant to degraation.

10. SUMMARY

The origin of vascular plants occurred during the ilurian, well over 400 Myr ago. This evolutionary transion in structural and physiological complexity is believed

to have been one of the most significant evolutionary events during the entire history of land plants. The development of lignified water-conducting tissues played a major role in the remarkable morphological and anatomical radiation of land plants during the Silurian and the Devonian (Raven 1993). Following the evolution of lignified vascular tissues, greater stature of the primary plant body was achieved, secondary growth evolved and a host of physiological modifications were likely to have been possible as a consequence of the improved efficiencies of lignified water-conducting tissues. Through integration of phylogenetically based palaeobotanical and neobotanical studies of tracheid structure we are beginning to address explicitly how, in a historical and mechanistic sense, lignified vascular tissues evolved. What is clearly needed is far more information about the structure, development and biochemical nature of cell walls in tracheids among extant basal vascular plants.

Beyond the evolutionary significance of studying tracheid structure in plesiomorphic vascular plants, additional developmental information on the course of lignification in basal vascular plant tracheids will be central to the formulation of more inclusive 'models' of tracheid structure and differentiation. Our current knowledge of spatial and temporal aspects of lignin deposition during cell wall differentiation for all vascular plants has been limited to seed plants (Boudet *et al.* 1995; Barceló 1997). If we are to ever fully understand the molecular and cellular basis for differentiation of tracheids, more phylogenetically global models of development will be essential, and these models will most certainly depend on a thorough understanding of the tracheids of basal vascular plants.

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Discussion

V. G. Chaloner (Department of Geology, Royal Holloway Iniversity of London, UK). It is now widely accepted that he conducting elements at the core of the axis of the Devonian plant Aglaophyton are not technically tracheids, ince they show no evidence of gyres of (annular or elical) wall thickenings. On these grounds, most authors ave excluded that genus from the tracheophytes (see, for xample, the 'protracheophytes' of Kenrick & Crane 997).

Nonetheless, the walls of these presumed waterconducting cells of Aglaophyton have the same distinctive black coloration that characterizes the presumably lignified xylem elements of the other true tracheophytes (e.g. Rhynia, Asteroxylon) in the Rhynie Chert. Inevitably this suggests similarity of the original wall chemistry of all these tissues. Do you think that it is possible that these walls in Aglaophyton were indeed lignified, but perhaps rather late in their ontogeny, after axial elongation was completed? If this was the case, no provision for wall stretching-no gyres of secondary wall thickeningwould have developed. If these cells were lignified, would they then qualify as tracheids? Has the difference between Aglaophyton and Rhynia perhaps been overrated?

W. E. Friedman. It is entirely possible that the walls of water-conducting elements in Aglaophyton were lignified. As you suggest, these smooth-walled metaxylem cells that lack evidence of gyres could have been stretched during axis elongation, and lignified after elongation of the immediately proximal tissues of the axis had ceased. If this is the case, these water-conducting cells of Aglaophyton would differ from the similar (and potentially homologous) hydroids of some mosses by virtue of the presence of lignified walls. Although never explicitly discussed in the literature, it is possible that the water-conducting elements of Aglaophyton could represent a character reversal from lignified tracheids with gyres to tracheids that lack gyres, but retain lignin. If this were the case, the distinction (structurally and phylogenetically) between Rhynia and Aglaophyton might be significantly less than presently assumed, and both taxa might be members of the same rhyniophyte clade. A possibly analogous case can be found in Nothia, which Kenrick and Crane hypothesize to be a lycophyte. The water-conducting cells of Nothia lack gyres and if the phylogenetic placement is correct, this would represent a secondary reversion in this character.

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