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Conducting tissues and phyletic relationships of bryophytes

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Internal specialized conducting tissues, if present, are restricted to the gametophytic generation in liverworts while they may occur in both generations in mosses. Conducting tissues are unknown in the anthocerotes. Water-conducting cells (WCCs) with walls perforated by plasmodesma-derived pores occur in the Calobryales and Pallaviciniaceae (Metzgeriales) among liverworts and in *Takakia* among mosses. Imperforate WCCs (hydroids) are present in bryoid mosses. A polarized cytoplasmic organization and a distinctive axial system of microtubules is present in the highly specialized food-conducting cells of polytrichaceous mosses (leptoids) and in less specialized parenchyma cells of the leafy stem and seta in other mosses including *Sphagnum*. A similar organization, suggested to reflect specialization in long-distance symplasmic transport of nutrients, also occurs in other parts of the plant in mosses, including rhizoids and caulonemata, and may be observed in thallus parenchyma cells of liverworts. Perforate WCCs in the Calobryales, Metzgeriales and *Takakia*, and hydroids in bryoid mosses, probably evolved independently. Because of fundamental differences in developmental design, homology of any of these cells with tracheids is highly unlikely. Likewise, putative food-conducting of bryophytes present highly distinctive characteristics and cannot be considered homologous with the sieve cells of tracheophytes.

Keywords: hydroids; sieve elements; food-conducting cells; leptoids; tracheary elements; water-conducting cells

1. INTRODUCTION

'he presence of an embryo, i.e. a stage in the life cycle uring which the sporophyte is associated with and epends on the gametophyte, is perhaps the most importnt unifying character of plants. For this reason the term mbryophytes' is receiving increasing favour as a more ppropriate name for 'plants', or 'land plants'. The bulk of norphological biochemical and molecular information ndicates that the embryophytes form a monophyletic roup together with charalean green algae (Garbary *et al.* 1993) and literature therein).

Further subdivision of embryophytes separates vascular lants from bryophytes, essentially on the basis of two lajor characters concerning sporophyte development and ascular tissues. In vascular plants, the embryo phase is elatively short and the sporophyte soon establishes direct ontact with the substrate, thus becoming independent rom the gametophyte. Moreover, the sporophyte evelops specialized vascular tissues, the xylem and hloem. In particular, the xylem contains wateronducting cells (WCCs), the tracheids and vessel lements, whose developmental pattern includes the eposition of a secondary lignified wall and final cytolasmic lysis.

The bryophytes traditionally include those embryohytes in which the sporophyte is permanently associated vith the gametophyte and never establishes direct contact vith the substrate. Traditionally, the bryophytes are set apart from vascular plants also on the basis of the lack of vascular tissues. Indeed, a number of bryophytes do contain vascular tissues including highly specialized WCCs which, like tracheids and vessel elements, undergo programmed cytoplasmic lysis. These cells, however, do not form lignified walls. For this reason the term 'tracheophytes' appears to be preferable to vascular plants when emphasis is put on vascular tissue as a diagnostic feature.

The bryophytes, as above defined, include three major groups: the anthocerotes (hornworts), mosses and liverworts. Phylogenetic links among these groups are largely a matter of speculation as there is no general agreement about whether the bryophytes sensu lato are a mono- or paraphyletic group (Garbary *et al.* 1993; Garbary & Renzaglia (1998) and literature therein). Their current taxonomical ranking spans from three classes in the same division (Pearson 1995) to three separate divisions (Bold *et al.* 1987). In this paper, we will avoid formal names as these might reflect a prejudicial assumption of relationships.

Relationships between the three bryophyte groups and tracheophytes are also far from resolved. Cladistic analyses setting the bryophytes as a paraphyletic group identify the anthocerotes (Sluiman 1985; Garbary & Renzaglia 1998) or the liverworts (Mishler & Churchill 1984, 1985; Bremer *et al.* 1987) as the sister group to the rest of embryophytes and either a clade of mosses plus liverworts (Garbary & Renzaglia 1998) or the mosses alone (Mishler & Churchill 1984, 1985) as the sister



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 \bigcirc igure 2. Transmission electron micrographs of moss hydroids. (a) *Mnium hornum*, sporophyte foot; hydroids with uniformly thin valls. (b) Polytrichum juniperinum, leafy shoot; hydroids with unevenly thickened walls. (c) Polytrichum formosum, leafy shoot; differntiating hydroid. Note the large pleomorphic nucleus (n); the longitudinal wall on the left is becoming thickened whilst the riginal transverse wall on the right is very thin. (d) Polytrichum formosum, leafy shoot; mature hydroids. Note the low electron pacity and absence of pores in the originally transverse walls (arrowed). Bar lines = $5 \,\mu m$.

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igure 3. (a-c) Details of polytrichalean hydroids. (a) Polytrichum formosum, sporophyte foot. Note the thick longitudinal walls and he thin, originally transverse walls (arrowed). (b,c) Polytrichum formosum, leafy shoots. (b) Obliterated plasmodesmata between a ydroid (h) and an adjacent cortical cell. (c) Remnants of plasmodesmata in an originally transverse wall of a differentiating ydroid. (d-f) Takakia, water-conducting cells. (d) Light micrograph, longitudinal 1 µm section. The central water-conducting ells (arrowed) are similar in size and shape to the adjacent cortical cells. (e) In longitudinal sections, the water-conducting cells ave transverse septa containing numerous plasmodesmata-derived pores. (f) Details of the pores in an end wall. (g) Detail of he wall between a water-conducting cell (w) and an adjacent cortical cell. Bar lines: $a, e = 5 \mu m; b, c = 1 \mu m; d = 100 \mu m;$ $= 0.5 \mu m; g = 0.25 \mu m.$

 Table 1. Taxonomic distribution and main features of internal pater-conducting cells in bryophytes

	-	
ICES	nthocerotes	unknown
BIOLC	Calobryales	slightly elongate cells with thin walls and large pores produced by lysis of primary wall associated with modified plasmodesmata
OYAL [ETY]	Metzgeriales (Pallaviciniaceae)	elongate cells with thickened walls and large pits produced by dissolution of secondary wall material associated with modified plasmodesmata; polyphenols in cell walls
HE R OCH	Metzgeriales (Moerckia)	slightly elongate cells with swollen walls. Remnants of plasmodesmata visible occasionally
	Marchantiales s.l.	unknown ^a
	Jungermanniales	unknown
ANSACTI	nosses Takakia	slightly elongate cells with thin walls and small pores derived from plasmodesmata
T T T	Andreaeidae	unknown
	Sphagnidae	unknown
	Bryidae	imperforate and very highly elongate cells with thick and/or thin walls; thin walls loosely textured; polyphenols in cell walls

See text for discussion of possible water-conducting elements in *'onocephalum* (Kobiyama & Crandall-Stotler 1999).

roup to tracheophytes. Among the most relevant torphological characters considered are stomata, present the sporophyte in some anthocerotes and mosses, and iternal conducting tissues, occurring in the majority of tosses and few liverwort taxa.

The present paper analyses ultrastructural and developental features of water- and food-conducting tissues in ryophytes, with emphasis on their possible significance in the context of phyletic interrelationships of embryohytes. Much of the present knowledge on this issue is use to the seminal work by the late Charles Hébant, to /hom this paper is dedicated.

2. WATER-CONDUCTING TISSUES

(a) Mosses

Three major moss groups are currently recognized, e. the Andreaeidae, Sphagnidae and Bryidae, the last peristomate mosses) being considered advanced relative b the first two (Crosby 1980). An internal strand of VCCs is lacking in both the Andreaeidae, including the ewly introduced genus *Andreaeobryum* (Murray 1988) figure 1*a*), and Sphagnidae (Ligrone & Duckett 1998*a*). 'he latter are characterized by specialized dead cells, the hyalocysts, forming an effective external waterconducting system (Mozingo *et al.* 1969).

An internal water-conducting system occurs in the Bryidae (figures 1, 2 and 3a-c) and is considered as a distinctive character of this group. This consists of highly elongate cells lacking cytoplasmic contents at maturity. Traditionally these cells are called hydroids, a term introduced by Potonié (1883) and first used for mosses in Tansley & Chick's (1901) classical paper on conducting tissues of bryophytes. The hydroids, alone or associated with stereids (thick-walled living cells), form a central strand of varying size (figure 1) in the leafy stem of the gametophyte and/or in the sporophyte seta, which is sometimes referred to as hydrom. In many instances, hydroids also occur in the leaf nerve, where they are usually associated with stereids and specialized parenchyma cells referred to as deuters (Hébant 1977). Absence of a central strand in certain genera (figure 1b) and species of Bryidae is frequent (Hébant 1977, 1979) and is interpreted as a result of reduction (Schofield 1985). In line with this notion is the fact that the more primitive members of the Bryidae, i.e. the nematodontous order Polytrichales (including the Dawsoniaceae), have highly developed conducting strands. Conversely, a tendency towards a reduction of the water-conducting system is observed in the more advanced groups. Some moss species are known where a strand of conducting cells occurs in the seta but is lacking in the gametophyte (Hébant 1977). The converse is true in Grimmia pulvinata, where the developing sporophyte remains almost entirely ensheathed by the upper leaves of the parental shoot.

The hydroids in the leafy stems of mosses arise from the inner cell produced by the first division of the apical derivatives (Berthier 1972). The initials are shortly rectangular cells with numerous plasmodesmata in their oblique terminal walls and fewer in the longitudinal walls. Hydroid differentiation is very fast and involves pronounced cellular elongation (figure lf,g) during which the cellular ends become so strongly tapered that the demarcation between longitudinal and transverse walls is solely indicated by the presence of Y-shaped junctions (figure 2d). The nucleus becomes elongate and pleomorphic (figure 2c) and may undergo endopolyploidization (Hallet 1972). Very early in development the plastids lose their starch content and become relatively small and irregular in shape. In some members of the Polytrichales (figures 2b,c and 3a) new wall material is deposited along longitudinal walls, but it is not clear if this is to be considered as a secondary wall (Hébant 1974). The newly deposited material causes the obliteration of the plasmodesmata between the hydroids and adjoining cortical cells (figure 3b). Hydroids in bryalean mosses have uniformly thin walls (figure 2a). Differentiation concludes with cytoplasmic degeneration and the alteration of walls, including the disappearance of all the plasmodesmata.

The modified walls have amorphous appearance with low electron opacity and may contain remnants of disrupted plasmodesmata (figure 3c) but always lack pores. Because of their loosely fibrillar texture (figure 2d) in transmission electron micrographs, it has been assumed that similar to tracheary elements (O'Brien 1974), maturational changes in the walls of hydroids involve



igure 4. Conducting cell differentiation in *Haplomitrium hookeri*. (a) Transverse section of an underground axis. Note the avestitive of mucilage (m) and the small central strand of water-conducting cells (arrowed). (b) At higher magnification the maller size and absence of contents distinguishes the water-conducting cells (arrowed) from the adjacent food-conducting lements. (c) Longitudinal section showing the transverse end walls of the water-conducting cells. (d) Plasmodesmata in the end vall between two differentiating water-conducting cells. (e-g) Details of the plasmodesmata-derived perforations in mature vater-conducting cells. (e, f) Longitudinal and transverse sections through end walls. (g) Pits ending blindly against adjacent bod-conducting cells (f). Bar lines: $a, c = 100 \,\mu\text{m}; b = 25 \,\mu\text{m}; d, f, g = 1 \,\mu\text{m}; e = 0.25 \,\mu\text{m}.$

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igure 5. Water-conducting cells in *Symphyogyna*. (a,b) Light micrographs. (a) Two strands of narrow water-conducting cells are isible in a recently bifurcated thallus. (b) Higher magnification. Note the thick walls of the conducting cells. (c,d) Transmission lectron micrographs showing details of wall differentiation. (c) At the onset of cell maturation the thin walls contain normal lasmodesmata and numerous cortical microtubules (arrowed) are present. (d) Thickening wall. Collars of electron-transparent naterial are associated with the plasmodesmata. (e, f) Scanning electron micrographs of mature water-conducting cells showing e elongated pits in the thickened walls. (Micrographs kindly supplied by Dr W. Frey). (g,h) Transmission electron micrographs f mature conducting cells showing the large pits in the now highly electron-opaque walls. Bar lines: $a = 100 \,\mu\text{m}$; $b = 10 \,\mu\text{m}$; $= 0.2 \,\mu\text{m}$; $d, e = 0.1 \,\mu\text{m}$; $f-h = 20 \,\mu\text{m}$.

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onsequence of cell extension. Only unthickened walls or vall areas undergo these structural alterations in polytrihaceous mosses, while in *Timmiella* (Ligrone et al. 1980) nd other bryalean mosses the walls are modified hroughout except at the cellular corners. Where the ydroids abut parenchyma cells the wall alteration stops t the middle lamella region of these cells (Ligrone et al. 980). In no case has lignin been detected in hydroids Hébant 1974, 1977), although histochemical evidence 🔀 -oints to the presence of polyphenolic compounds 🖳 Scheirer 1980). Hydroids also lack secondary wall atterns such as spirals, bands or pitting. A chemical Utudy of whole plants confirms that mosses do not contain gnin as defined in angiosperms and gymnosperms but nay produce other types of polyphenolic compounds Miksche & Yasuda 1978).

nzymatic removal of non-cellulosic polysaccharides

Hébant 1977). However, cytochemical evidence for such

vall 'hydrolysis' has never been produced. The apparently

hydrolysed' appearance of the walls may be simply a

Mature hydroids are very variable in size, ranging rom 200 to $1500 \,\mu\text{m}$ in length and 10 to $25 \,\mu\text{m}$ in width, ccording to the species and organs. The overlap contact rea of adjacent hydroids may be as large as several undred micrometres.

External water conduction by capillarity is of utmost mportance in mosses (Proctor 1979). Nevertheless, umerous experiments using a range of different echniques have demonstrated that the inner strand of ydroids, when present, is an effective preferential way for 'ater transport (Hébant (1977) and literature therein). Vater supply to the growing capsule must usually rely ntirely on internal conduction along the seta, as this lacks n external capillary system and is covered by a waterepellent cuticle. However, the capillary system formed by the ensheathing leaves in *Grimmia pulvinata* may have a ole in water supply to the growing sporophyte, thus ccounting for the lack of hydroids in the seta of this hoss.

In a distinct position relative to the other mosses stands *Takakia*, an enigmatic taxon recently transferred from verworts, where it had been placed since the discovery f its gametophyte by Mitten (1861) and the circumcription of the genus (Hattori et al. 1968). With the recent iscovery of the sporophytic generation and antheridial lants, it has become undeniable that Takakia is a moss /ith affinities with the Andreaeidae (Smith & Davison 993), a conclusion also supported by ultrastructural tudies (Garbary et al. 1993; Ligrone et al. 1993; Renzaglia Ut al. 1997). As first reported by Hébant (1973), and in Oharp contrast to other mosses, however, Takakia possesses VCCs with plasmodesmata-derived pores, a feature prmerly considered to support its classification in the verwort group of Calobryales (Schuster 1984; Schofield 985). These cells form a small, central strand both in the ametophyte shoot and in the sporophyte seta (Renzaglia t al. 1997). They are of about the same size and shape as οhe adjoining cortical parenchyma cells, i.e. ca. 12 μm in vidth and 80 µm in length, but have much thinner walls a. $0.3 \,\mu\text{m}$) (figure 3*d*,*e*). The terminal walls are nearly erpendicular or slightly oblique to the long cellular axis. like the hydroids, the central strand cells in Takakia ndergo cytoplasmic autolysis, with a strong peak of acid-phosphatase activity (Hébant 1975), and have no cytoplasmic contents at maturity. The plasmodesmata show no apparent modification during differentiation and when disrupted leave perforations in the walls, which do not normally exceed 120 nm in diameter (figure 3f). Most perforations are concentrated in the terminal walls but are also frequent in longitudinal walls. The plasmodesmata connecting WCCs with adjoining cortical cells break down only on the side of the dead cell, while on the other side remains a clearly recognizable desmotubule (figure 3g).

(b) Liverworts

Among the liverworts, an internal strand of specialized WCCs occurs in the Calobryales and in few genera of the Metzgeriales (simple thalloid liverworts), in both cases being restricted to the gametophyte (Burr *et al.* 1974; Smith 1966; Hébant 1977, 1978, 1980). The Marchantiales (complex thalloid liverworts) and Jungermanniales (leafy liverworts) have no internal strand of water-conducting cells (Hébant 1977; Kobiyama & Crandall-Stotler 1999). Like hydroids and tracheids, WCCs in liverworts are dead and lack cytoplasmic contents at maturity. However, they are distinct from both as their walls contain numerous pores arising from plasmodesmata.

As in *Takakia*, WCCs in calobryalean liverworts (figure 4) are similar in shape to ordinary parenchyma cells (*ca.* 20 μ m wide and 50–60 μ m long) with thin walls (0.25–0.50 μ m) and transverse or slightly oblique terminal septa. Different from *Takakia*, at a late stage of differentation swelling of the middle region of plasmodesmata is observed (figure 4*d*,*e*). Moreover, the wall material immediately adjoining the plasmodesmata is removed during terminal cytoplasmic dissolution, thus producing pores much wider (300–600 nm) than those in *Takakia*.

By contrast, WCCs in the metzgerialean genera Symphyogyna (figure 5), Hymenophyton and Pallavicinia are thin and elongate (*ca.* $8 \,\mu m$ wide and up to $300 \,\mu m$ long) with tapering ends and thickened walls $(1-1.7 \,\mu m)$ perforated throughout by numerous pits (250-600 µm in diameter). The walls are strongly electron-opaque (figure 5g,h) and probably contain polyphenolic compounds because they are autofluorescent when examined in blue light, though tests to reveal lignin are negative (Smith 1966). The pits arise from plasmodesmata (figure 5c,d) through a mechanism closely recalling the genesis of pores in sieve elements (Esau & Thorsh 1985; Lucas et al. 1993). This process has recently been described in detail in Symphyogyna (Ligrone & Duckett 1996). Following an elongation phase that terminates when the cells have reached their definitive sizes, the cell walls are thickened by deposition of electron-opaque material, on extraplasmodesmatal areas, and of electron-transparent material forming collars around plasmodesmata (figure 5d). The newly deposited wall material differs cytochemically and structurally from the original wall (Smith 1966; Ligrone & Duckett 1996) and is therefore to be considered a secondary wall. Eventually, the cells undergo cytoplasmic dissolution and the electrontransparent wall material around the plasmodesmata is removed. Only a minority of the pits are completely open, while most appear to be occluded by a plug in the middle (Frey et al. 1996). This arises from the thin,



igure 6. Cytological details of moss food-conducting cells. (a) Cytoplasmic polarity in a leafy stem of *Plagiomnium undulatum*; ost of the organelles are at the distal (top) end of the lower cell. (b) *Mnium hornum*, seta showing the longitudinal alignment of longated plastids (p) and the highly elongated nucleus (n). (c,d) Longitudinal arrays of microtubules associated with tubules nd vesicles in a leafy stem of *Plagiomnium undulatum* (c) and *Polytrichum juniperinum* (d). (e) Transverse section of leptoids and djacent hydroids (h) in a stem of *Polytrichum commune*. Note the numerous vesicles and membrane profiles throughout the lectron-transparent cytoplasm. Bar lines: $a = 4.0 \,\mu\text{m}$; $b, e = 2.0 \,\mu\text{m}$; $c, d = 0.5 \,\mu\text{m}$.

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igure 7. Cytological details of leptoids in *Polytrichum juniperinum* sporophytes. (*a*) Cytoplasmic polarity in a foot. The istal cellular end of the lower cell is packed with endoplasmic reticulum. (*b*) Longitudinally aligned plastids (p) and a pindle-shaped nucleus in a seta. (*c*) Sheets of endoplasmic reticulum adjacent to the nucleus. (*d*) Endoplasmic microtubules sociated with elongate mitochondria. (*e*) Conducting parenchyma in a seta. Note the thick walls and intercellular spaces. ar lines: $a-c = 2.0 \mu m$; $d = 10 \mu m$; $e = 4.0 \mu m$.



igure 8. Cytological details of moss food-conducting cells. (a) Abundant endoplasmic reticulum in a leafy shoot of *Polytrichum iniperinum*. (b) Degenerating nucleus in a shoot of *Polytrichum juniperinum*. (c,d) Abundant plasmodesmata in the end walls etween food-conducting cells in *Neckera crispa* and *Sphagnum cuspidatum*. Note the trumpet-shaped cell ends in the latter. ar lines: $a = 5 \mu m$; $b-d = 10 \mu m$.

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primary wall area being not completely removed along with the electron-transparent collar lying above it. However, the dissolution of the desmotubule and plasmaemma leaves an open, albeit small, pore in the primary vall. The pits show a spiral arrangement that mirrors hat of cortical microtubules present during cell differeniation, and often merge together to form larger ompound pits. Because of the spiral arrangement of pits nd their tendency to fuse during development, the 'ymphyogyna-type fully differentiated WCCs exhibit a triking similarity to tracheids (figure 5e, f) (Frey *et al.* 996). Experimental evidence using cosin as a tracer in ving plants has demostrated that water moves in the entral strand at a much faster rate than in neighbouring 'arenchyma cells (Smith 1966).

WCCs in *Moerckia*, a genus closely allied to *Pallavicinia* Renzaglia 1982), appear to stand apart from those in ther metzgerialean liverworts, for these cells show wollen, apparently hydrolysed walls with no pits Hébant 1973). However, different from moss hydroids, races of plasmodesmata have occasionally been encounered in these walls (Hébant 1977). It is not possible to stablish from the data available whether WCCs in *Moerckia* are a variant of perforate WCCs or represent a istinct type.

A recent study in the marchantialean liverwort genus *onocephalum* has produced a detailed description of itted cells scattered in the parenchyma tissue in the hallus nerve and in the stalk of archegoniophores Kobiyama & Crandall-Stotler 1999). These cells ossess reticulate primary wall thickenings of cellulosic ature and prominent pit fields, the latter occurring in oth lateral and terminal walls in C. conicum but only in ateral walls in C. japonicum. The cells have large central acuole(s) and very thin peripheral cytoplasm. Nuclear oss and partial cytoplasmic lysis are reported to occur n some of the cells, though no ultrastructural evidence produced. Kobiyama & Crandall-Stotler (1999) also laim that tests with methylene blue suggest that these ells facilitate absorption of water from the rhizoids but gain supporting micrographic data are lacking. The uthors' claim that these cells are a unique kind of vater-conducting element must therefore be viewed vith considerable caution; Kobiyama & Crandalltotler's micrographic evidence indicates that these cells re much more likely to be the food-conducting lements previously described in the centre of the thalli f complex thalloid liverworts (Ligrone & Duckett 994b).

(c) Anthocerotes

The anthocerotes contain no specialized WCCs in the ametophyte nor in the sporophyte. In some genera, e.g. *haeoceros* and *Anthoceros*, the sporophyte contains a entral strand of elongate cells, referred to as the colunella, which bears a superficial resemblance to the stele n lower tracheophytes (Campbell 1925). Associated with pores are elaters with spirally arranged wall thickenings imilar to those in tracheids but lacking lignin (Proskauer 960). The columella cells lack cytoplasmic contents at naturity but apparently have no special role in water ransport (Isaac 1941). No ultrastructural study of these ells has been reported to date.

3. FOOD-CONDUCTING TISSUES

(a) Mosses

The Polytrichales have long been known to possess, besides hydroids, specialized cells with marked morphological similarity to the protophloem sieve cells in tracheophytes. These cells were called 'leptoids' by Tansley & Chick (1901), who apparently derived this term from the term 'leptom', introduced by Haberlandt in 1879 for the whole system of phloem-like tissue in polytrichaceous mosses (table 2).

Leptoids in polytrichaceous mosses (figures 6–8) occur both in the gametophyte and the sporophyte seta. They attain the highest degree of structural specialization in the leafy stem (table 3), where they are intermingled with elongate parenchyma cells to form a ring around the central strand of hydroids. The leptoids are several hundred micrometres long, up to $500 \,\mu\text{m}$ in *Dendroligotrichum* (Scheirer 1990), have enlarged extremities and oblique end walls. Major features of mature leptoids that recall protophloem sieve cells are listed in table 4.

The pores in the terminal walls of leptoids are modified plasmodesmata (figure 9) with a desmotubule forming a median enlargement *ca*. 100–200 nm wide but usually still constricted at both ends (Hébant 1976; Scheirer 1978, 1990; cf. Lucas *et al.* 1993). The desmotubule is continuous with tubular endoplasmic reticulum elements forming a prominent network in the adjacent cytoplasm. Tubules of endoplasmic reticulum traversing the sieve area pores are of common occurrence in ferns and *Psilotum* (Evert 1990*b*) as well as in conifers (Schulz 1990), but these do not form desmotubule-type constrictions; the sieve pores are generally completely open in other vascular cryptogams (Evert 1990*b*) and in angiosperms (Eleftheriou 1990, 1996; Evert 1990*a*).

With the use of radioactive tracers, leptoids have been demonstrated to be a preferential route for the translocation of organic nutrients, with translocation rates of up to several tens of centimetres per hour (Eschrich & Steiner 1967; Collins & Oechel 1974; Thomas *et al.* 1988; Schmid 1998).

In all other moss groups apart from the Polytrichales, the cortical tissue of the leafy stem and seta consists of elongate parenchyma cells $(150-250 \,\mu\text{m})$. These are likely to function in symplasmic transport of assimilates because of the presence of numerous plasmodesmata (figure 8c) in their end walls $(10-30 \,\mu\text{m}^{-2})$. As in leptoids, the plasmodesmata in these cells present a median enlargement (figure 9) and often show continuity with endoplasmic reticulum. However, the internal parenhyma cells of nonpolytrichaceous mosses lack most of the distinctive features of leptoids and appear to be structurally much less specialized. According to Hébant (1977) they should be referred to as 'conducting parenchyma cells', while the use of the term leptoid should be restricted to the specialized conducting cells of Polytrichales.

Recent ultrastructural research on members of the Sphagnales, Polytrichales and bryalean mosses (Ligrone & Duckett 1994*a*, 1998*a*), as well as preliminary observations on *Takakia*, have revealed that—irrespective of the level of morphological specialization—putative food-conducting cells share a series of morphological characteristics that enable them to be distinguished from normal parenchyma cells at first sight (table 2). This extremely



igure 9. Plasmodesmata between food-conducting cells in mosses. (a-c) Longitudinal sections showing median enlargements in leafy shoot of *Mnium hornum* (a) and *Aulocomnium palustre* (b,c). Note the frequent continuity with endoplasmic reticulum. t-f Transverse sections of Polytrichum formosum (d) and Sphagnum recurvum (e, f) showing median chambers usually containing no is cernible desmotubule, but occasionally irregular tubular structures (arrowed in d and f), and subterminal constrictions with learly defined plasmalemmal and desmotubule profiles. Bar lines: $a-c = 0.2 \mu m$; $e-f = 0.1 \mu m$.

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able 2. Cytological characteristics of putative food-conducting cells in bryophytes

, Constant feature; -, not observed; \pm , observed occasionally.)

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CCES	Takakia	Sphagnidae	Bryidae	liverworts
ongitudinal arrays of endoplasmic				
microtubules associated with:				
spindle-shaped nucleus	+	+	+	±
elongate plastids	-	—	+	_
elongate mitochondria	+	+	+	_
elongate microbodies	+	+	+	_
pleomorphic vacuoles	+	+	+	+
membrane-bound tubules and vesicles	+	+	+	+
> endoplasmic reticulum	_	_	+	_
ytoplasmic polarization: the bulk of				
organelles at the sink end of each cell				
(i.e. towards the apex of the leafy shoot				
() and capsule in the sporophyte); granular	+	+	+	±
cytoplasm at the opposite end				
isappearance of large vacuoles	+	+	+	+
(present during differentiation)				
• Ligh frequency of plasmodesmatain	+	+	+	+
end walls: desmotubules forming a	I	1	·	, i
modian anlangement				
median emargement	1	1	1	
ggregation of free ribosomes	- -	+	т ,	_
o bundant dictyosomes and	+	+	+	+
trans Golgi network				

istinctive organization is found not only in the cortical ells of the leafy shoot and seta but also in other types of issues and organs, including mature caulonemal and hizoidal cells (Duckett et al. 1998), where a need for ong-distance symplasmic transport of solutes can be ostulated (table 5).

An experimental study (Ligrone & Duckett 1996) has hown that polarized organization is suppressed following emoval of the source/sink gradient. The longitudinal lignment of the nucleus and organelles is disrupted by ryzalin, a microtubule inhibitor, while it is insensitive to ytochalasin, a drug affecting the actin microfilaments.

Of special interest is the situation recently discovered n the Sphagnales, where the internal parenchyma cells of he leafy stem are morphologically similar to the onducting parenchyma cells of other mosses but, unlike hese, arise from a subapical secondary meristem Ligrone & Duckett 1998a, b). The internal parenchyma ells of Sphagnum are most likely the main route for the ymplasmic longitudinal transport of nutrients shown xperimentally and thought to reflect nutrient recycling rom old to new parts of the plant (Rydin & Clymo 0989). From a developmental standpoint, the conducting arenchyma cells of Sphagnales are not homologous to he equivalent cells in other mosses. The presence of a 5

econdary meristem appears to be a characteristic unique b the Sphagnales, being probably related to the equally nique ecology of these mosses and their success in waterbgged oligotrophic habitats.

Major morphological similarities and differences etween sieve elements of tracheophytes and foodonducting cells of mosses are summarized in table 5.

(b) Liverworts

Until very recently no specialized food-conducting issue had been described in liverworts. The internal cells elongate and were typically thought to contain a large central vacuole. In numerous thallose liverworts, the ventral cells may be colonized by endophytic fungi (Read et al., this issue). However, a recent study of Asterella (Ligrone & Duckett 1994) revealed that the parenchyma cells of the midrib, between the dorsal photosynthetic tissue and the ventral fungus-containing tissue, are polarized and contain highly pleomorphic vacuoles associated with endoplasmic microtubules. Similar microtubulevacuole associations have now been detected in other liverworts (table 5), including Pellia and Haplomitrium (figure 10).

of the thallus or of the stem in leafy liverworts are slightly

(c) Anthocerotes

No morphological specialization for symplasmic transport has been reported in the anthocerotes. However, very little ultrastructural research has been done on this group to date. Considering that the highly distinctive cellular organization of polarized food-conducting cells in mosses has escaped attention until very recently, in spite of the considerable interest focused on this group, careful investigation of the anthocerotes must now be given a high priority.

4. DISCUSSION: EVOLUTIONARY AND FUNCTIONAL CONSIDERATIONS

Making holes in the walls by disruption of plasmodesmata together with the total loss of cytoplasmic contents is perhaps the easiest way to form a WCC under a selective pressure for more efficient apoplastic water transport. Possible homology of perforate WCCs in the calobryalean and metzgerialean liverworts should therefore be considered with caution, especially when set against the multiple origin of vessel elements, a much more complex cell type. WWCs in the two liverwort

'able 3. Additional features of food-conducting cells in olytrichaceous mosses (leptoids)

nacreous walls

endopolypoidy (also documented in caulonemal cells;
Kingham et al. 1995; Duckett et al. 1998)
partial degeneration of the nucleus
refractive spherules
large endoplasmic reticulum stacks
abundant tubular endoplasmic reticulum associated
with plasmodesmata
callose associated with plasmodesmata
-

able 4. Sieve elements versus putative food-conducting cells in

losses

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milarities absence of vacuoles aggregation of ribosomes nacreous walls^a nuclear degeneration^a refractive spherules^a callose associated with plasmodesmata^b

ifferences (in sieve elements): no cytoplasmic polarization disappearance of desmotubules oryzalin-insensitive at maturity actin bundles prominent during differentiation phloem protein total nuclear degeneration

Restricted to the Polytrichales in mosses. Restricted to the Polytrichales in mosses; absent in some lower 'acheophytes.

roups present marked structural differences, those in the

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Ietzgeriales being far more specialized than those in the alobryales. Moreover, while an internal strand of WCCs a generalized feature of the Calobryales—13 species in he only genus Haplomitrium (Bartholomew-Began 1991)his is restricted to a minority of relatively advanced becies within the much larger order Metzgeriales Renzaglia 1982; Schuster 1984). Basal taxa in liverworts ack WCCs, and the closely allied families Pallavicinia-(Pallovicinia, Symphygyne) and Hymenophytaceae eae Hymenophyton) have only a very remote link with the alobryales (Mehra 1968; Schuster 1984; Schofield 1985). ven though Bartholomew-Began (1990) places the Caloryales (Haplomitriales) as an order within the subclass Ufetzgeriidae, our conclusion is that perforate WCCs have nost probably evolved independently in the Calobryales nd metzgerialean liverworts.

The general appearance of WCCs in metzgerialean verworts, notably their thickened walls with elicoidally rranged pits, is strongly reminiscent of tracheids. Iowever, pits in the former develop by removal of econdary wall material closely associated with modified lasmodesmata, while in the latter they arise from the /sis of primary unlignified walls with no direct relation p plasmodesmata, albeit in both cases cortical microibules appear to have a prominent role in morphogenesis of. Ligrone & Duckett 1996; McCann 1997; Chaffey *et al.* 997, 1999; Seagull & Falconer 1991). The two cell types, Table 5. The occurrence of 'food-conducting cytology' in bryophytes

mosses

- inner parenchyma cells in the stem of the leafy shoot
- inner parenchyma cells in the sporophyte foot and seta
- parenchyma cells in the leaf nerve
- rhizoids
- caulonemata
- basal cells of mucilage hairs
- stalks of cauline gemmae

hepatics

- inner parenchyma cells in the central thallus of *Conocephalum* and *Asterella* (Marchantiales) and *Pellia* (Metzgeriales)
- inner parenchyma cells surrounding water-conducting cells in the leafy shoot and 'roots' of *Haplomitrium* (Calobryales)

anthocerotes

unknown

therefore, have sharply different developmental designs and homology between them is highly unlikely. The same applies to perforate WCCs in other bryophyte groups.

Reports of cells with helical thickenings from Lower Devonian Rhyniophytina (sensu Edwards 1993), referred to as *Sennicaulis*-type or S-type WCCs, emphasize alleged similarities with metzgerialean WCCs, notably the presence of 'micropores' scattered thoroughout a continuous inner wall layer (Kenrick *et al.* 1991; Kenrick & Crane 1991). As noted by Frey *et al.* (1996) and much earlier by Smith (1966), a more careful comparison of the structure of the two cellular types reveals that the two cellular types have basically different designs.

The WCCs in Haplomitrium are reminiscent of those in the moss Takakia. Consideration of the possible relationships of these cells becomes highly pertinent to the still unresolved question of phyletic interrelationships between mosses and liverworts. Thus it is intriguing, in the present context, to observe that similar WCCs is but one of several characters that provided the basis for the grouping Haplomitrium and Takakia in the liverwort order Calobryales (Schuster 1984; Gradstein 1990). The assumption of homology of WCCs in Takakia and Haplomitrium is consistent with topologies in which mosses and liverworts form a monophyletic group (e.g. Garbary & Renzaglia 1998), with the additional implication of mosses sharing a common ancestor with Haplomitrium. Once again, however, it must be emphasized that because of the extreme structural simplicity of WCCs in the two taxa, independent origin of these cells is quite possible. The lack of sporophytic WCCs in the liverworts is a major character separating liverworts from mosses, including Takakia. Most likely this is functionally related to the fact that the sporophyte seta in liverworts elongates by cell expansion (Schnepf & Deichgräber 1979) only after spore maturation, while in mosses the seta completes development before capsule maturation. WCCs are dead and no longer capable of expanding at maturity; hence they can develop in the seta of mosses but not of liverworts.



igure 10. Details of food-conducting cells in liverworts. (a-c) Asterella wilmsii. (d,e) Pellia epiphylla. (f,g) Haplomitrium hookeri. [] a,b] Light and transmission electron micrographs, longitudinal sections of inner thallus cells showing their thickened, pitted valls and cytoplasmic polarity (b). (c) Details of the pits showing numerous plasmodesmata with enlarged median chambers ontaining desmotubules. (d) trans Golgi network. (e-h) Endoplasmic microtubules associated with a mitochondrion (e) and rembranous vesicles and tubules (f-h). Bar lines: $a = 20 \,\mu\text{m}$; $b = 2 \,\mu\text{m}$; $c, f-h = 0.2 \,\mu\text{m}$; $d, e = 0.4 \,\mu\text{m}$.

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The hydroids in bryoid mosses are a highly distinctive ype of WCC. Unlike WCCs in other bryophytes, hydroids ave imperforate walls, the plasmodesmata being obliterted during development. Preliminary experimental bservations have shown that when dehydrated, hydroids end to collapse, unlike tracheary elements, and are ighly resistant to cavitation (Schmid 1998; A. M. chmid, unpublished data). Most likely this remarkable roperty has made possible the reconciliation of dead VCCs with desiccation tolerance, which is a major hysiological characteristic of mosses (Proctor 1982; Oliver & Bewley 1984). Desiccation tolerance not only Cours in the Bryidae, but also in the Sphagnidae and -Indreaeidae. There are at present no clues as to whether he absence of hydroids in these two groups is derived or rimitive, though primitiveness is consistent with the asal position assigned to both groups. The perforate **O**VCCs in *Takakia* might be interpreted either as a derived \checkmark ature (autapomorphy) of this taxon or as a primitive ature (plesiomorphy) lost in most living mosses.

Nothing is known of the effects of dehydration on erforate WCCs. It is interesting, however, that neither *Iaplomitrium* nor *Takakia*, is tolerant to desiccation (Grubb 970). The same is probably true for the Pallaviciniaceae. mong the liverworts, desiccation tolerance is a widepread feature of the Jungermanniales and also occurs in nembers of the Marchantiales (Proctor 1982), both of thich lack internal WCCs.

Similarities between hydroids and tracheids have been mphasized to support the contention that these two cell ypes are homologous (Scheirer 1980). In particular, attenon has been focused on the mechanism of wall modificaon, allegedly involving in both cases partial hydrolysis of ertain areas of the primary wall, with polyphenols laying a protective role in hydroids similar to that postuted for lignin in tracheids (O'Brien 1974). The assumpon of homology of hydroids and tracheids is fundamental cladograms in which the mosses are the sister group to cacheophytes (Mishler & Churchill 1984, 1985; Mishler et l. 1994). A closer look at the developmental design of the wo cell types, however, shows several major differences. 'he deposition of secondary lignified walls is a fundanental event in the morphogenesis of tracheids and is receded by the establishment of a cortical microtubule rray that accurately predicts the sites of formation of wall nickenings and pits (Hogetsu 1991; McCann 1997; haffey et al. 1997, 1999). No immunocytochemical study f differentiating hydroids has hitherto been carried out, ut transmission electron microscopy shows no pre-Oatterned arrangement of microtubules in the cortical Oytoplasm underlying expanding walls. Cell wall thickning, most likely of primary nature, only occurs in the ydroids of certain polytrichaceous mosses and here it only ffects longitudinal walls. In the large majority of mosses, he hydroids undergo no wall thickening. Moreover, cell all alteration indiscriminately affects all unthickened zalls. This suggests that, at least in part, cell wall loosning results from passive stretching of differentiating and ead hydroids by adjoining living cells. We conclude from

nosses between hydroids and tracheids. The discovery of unornamented WCCs, referred to as nydroids', in the branched sporophyte of the Lower

his that hydroids are a specialized type of WCC unique to

Devonian plant Aglaophyton major (Edwards 1986) is unlikely to support the hypothesis of homology. These cells are much wider (*ca.* 40 μ m) than moss hydroids and, as Edwards (1993) properly remarks, have no exact counterpart in extant bryophytes.

Indirect support for the notion of independent origin for hydroids and tracheids comes from studies of putative food-conducting cells in mosses. The discovery of a common structural pattern in leptoids and conducting parenchyma cells of mosses, independent of the diverse morphology and varying degree of cytological specialization that may be found in different species and organs, introduces a fundamental distinction from sieve cells of tracheophytes. In the latter cytoplasmic polarity has never been observed, while microtubules, present in the form of conspicuous cortical arrays during development, do not persist beyond maturation of the cell (Eleftheriou 1990, 1996; Evert 1990a,b; Schulz 1990). The two cell types, therefore, are unlikely to be homologous. The striking similarity of leptoids in polytrichaceous mosses to sieve cells is probably an instance of homoplasy related to the relatively large sizes attained by these mosses and consequent evolutionary pressure for a more efficient transport of assimilates. A similar example of homoplasy is found in the brown algae, notably the Laminariales, which contain highly specialized food-conducting cells bearing striking similarity to sieve elements of tracheophytes (Schmitz 1990). The occurrence in liverworts of cells structurally similar to food-conducting cells of mosses raises the possibility that this type of cell is a plesiomorphy of the liverwort-moss clade. This notion cannot be reconciled with topologies where mosses alone are resolved as the sister group to tracheophytes.

While pointing to conducting tissues as a major issue in embryophyte phylogeny, the present analysis introduces more new questions than answers. A more precise assessment of homology/homoplasy requires further investigation. In particular, useful information is expected from a better characterization of conducting tissues in the fossil record of early embryophytes as well as from comparative cytochemical and immunocytochemical studies of conducting tissues in extant bryophytes. Also required are physiological experiments that compare rates of water and solute transport in the diverse conducting elements found in bryophytes with those in the xylem and phloem of lower tracheophytes.

5. CONCLUSIONS

Perforate WCCs in the Calobryales and metzgerialean liverworts have probably evolved independently.

Perforate WCCs in *Takakia* have no counterpart in other mosses nor in tracheophytes. Similarities with WCCs in Calobryales are most likely a homoplasy.

The imperforate hydroids in bryoid mosses have no counterpart in other living embryophytes and must be considered as an autapomorphy of this group; absence of hydroids in one or both generations in numerous bryoid mosses is due to reduction.

Polarized food-conducting cells of mosses (including leptoids in polytrichaceous mosses) are not homologous with sieve cells of tracheophytes.

Further studies are needed to clarify possible relationships between (i) the WCCs in *Moerckia* and in other netzgerialean liverworts; (ii) perforate WCCs of *Takakia* nd imperforate hydroids in the Bryidae; (iii) cells with food-conducting cytology' in liverworts and in mosses.

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