APPENDIX TO BE MADE AVAILABLE ONLINE

APPENDIX 3. (All references are in Literature Cited of printed manuscript, except for those that appear only in Appendix 3—see below). Characters and character states considered here, but excluded from this study either for lack of accessible information, highly ambiguous homology concepts, or because they were not informative in a maximum parsimony analysis. Characters derived from published studies are indicated in parentheses by an author abbreviation followed by the character number. New characters are indicated by "New" following the character number. Unless otherwise stated, for characters and states defined in the same manner as in Pryer et al. (1995), the same character state arguments and definitions apply. References useful in coding taxa and defining characters are given if appropriate. Abbreviated publications: D - Doyle (1996); GRD - Garbary et al. (1993); GR - Garbary and Renzaglia (1998); KC - Kenrick and Crane (1997); M et al. - Mishler et al. (1994; table 3); NCSF - Nixon et al. (1994); PSS - Pryer et al. (1995); R - Rothwell (1999); S - Schneider (1996); SL - Stevenson and Loconte (1996). Additional characters excluded from this study, but pertinent only to seed plants or "bryophytes", are not listed below. For examples, see Garbary et al. (1993), Mishler et al. (1994), Nixon et al. (1994), Doyle (1996), Kenrick and Crane (1997), and Garbary and Renzaglia (1998).

I. SPOROPHYTE

I. A. LEAF
137. (SL 34) **Ligule**: absent (0); present (1). Not informative with this dataset. The presence of a scale-like structure on the adaxial surface of the leaf is known only from Selaginellaceae and Isoëtaceae in living plants. The structure is also known from fossil Lycopsida (Kenrick and Crane 1997). Structures on leaves of some monocots (e.g., palms, grasses) that are also called ligules are not homologous to ligules of *Isoëtes* and *Selaginella*. Goebel (1930), Bierhorst (1971), Kubitzki (1990), Kenrick and Crane (1997).

138. (PSS 10) **Blade hairs**: absent (0); present (1). In nearly all euphyllophytes, young euphylls possess hairs, but the adult leaves vary from naked to densely covered with hairs. The density of hairs may be relatively constant for species and sometimes for genera too, but higher taxa, such as families, are mostly polymorphic. The persistence of hairs probably reflects adaptations to dry environments and other environmental stresses. Hairs are of interest for studies focusing on the phylogeny of genera and families, but the character does not offer suitable information for deep level phylogeny. Kubitzki (1990).

139. (PSS 18; S 112) **Trophopods**: absent (0); present (1). This character is informatively variable at the genus level. Wagner and Johnson (1983), Kubitzki (1990).

140. (PSS 86) **Non-appendicular leaves**: absent (0); present (1). Not informative because non-appendicular leaves occur in very few species that are not closely related. Bierhorst (1968, 1969, 1973, 1974), Kaplan (1977).

141. (R 18) **Planation of vegetative leaves**: absent (0); present (1). Not informative with this dataset.

142. (R 19) **Biseriate vegetative leaves**: absent (0); present (1). Not informative with this dataset.
143. (R20) **Quadriseriate branching of vegetative leaves**: absent (0); present (1). Not informative with this dataset.

144. (PSS 88) **Winged petiole and/or rachis**: absent (0); present (1). Not informative with this dataset. Kubitzki (1990).

145. (SL 33) **Laminar buds**: absent (0); present (1). This character is variable within genera and may reflect ecologically-induced convergence. Kubitzki (1990).

146. (PSS 6) **Secondary vein form**: isotomous, dichotomous (0); anisotomous, dichotomous (1). Secondary and tertiary veins tend to be dichotomously branched. This character may be highly influenced by environmental and developmental constraints. Bower (1923, 1926).

147. (SL 39) **Paracostal commissure**: absent (0); present (1). Not informative with this dataset. Kubitzki (1990).

148. (SL 50) **Submarginal commissure**: absent (0); present (1). Not informative with this dataset. Bower (1923, 1926, 1928), Kubitzki (1990).

149. (SL 53) **Vascular commissure**: absent (0); present (1). Not informative with this dataset. Bower (1923, 1926, 1928), Kubitzki (1990).

150. (PSS 8; SL 38) **Vein areoles**: with free included veinlets (0); without free included veinlets (1). Not informative with this dataset. Bower (1923, 1926), Kubitzki (1990).

151. (PSS 40) **Hypodermis**: absent (0); present (1). Insufficient information. This character is difficult to discern. Davis (1991), Ogura (1972), Payne and Peterson (1973).
152. (PSS 87; SL 41) **Hippocampus-shaped bundles in petiole**: absent (0); present (1). Interesting character for derived fern groups only. Ogura (1972).

153. (New) **Aerophores**: absent (0); present (1). Aerophores are modified pneumathodes that extend as knob-like to peg-like structures from the petiole and rachis. They are often not well-preserved in herbarium material. Aerophores are reported for various groups of leptosporangiate ferns (e.g., Blechnaceae, Cyatheaceae, Plagiogyriaceae, Thelypteridaceae). Knoblike structures in *Culcita* and *Thyrsopteris* may be related. However, the homology of all these structures is unclear and needs further investigation. Bower (1923, 1926), Kubitzki (1990), Davis (1991).

154. (PSS 20) **Sclerenchyma coloration on petiole and rachis**: not dark-pigmented (0); dark-pigmented (1). Not informative with this dataset; probably correlated with environmental constraints. Ogura (1972), Kubitzki (1990).

155. (New) **Palisade parenchyma in mesophyll**: absent (0); present (1). In contrast to angiosperms, the mesophyll is not always differentiated into palisade and spongy parenchyma in ferns. The suppression of palisade parenchyma may be induced by environmental constraints. Ogura (1972).

156. (PSS 78; SL 29) **Blade thickness**: three or more cells thick (0); one to two (rarely three) cells thick (excluding veins) (1). The second character state is correlated with the absence of a differentiated epidermis and the absence or strong degradation of the cuticle. Reduction of blade thickness (cell layers) occurs in several groups of ferns (e.g., Hymenophyllaceae, *Hymenophyllopsis* in Cyatheaceae, Marattiacae, and Osmundaceae). This is combined with the loss of differentiation of blade tissues (e.g., palisade parenchyma, spongy parenchyma), but the lamina is usually not so extremely
reduced to three or fewer layers. Ferns with reduced but three or more cell layers are
found in Aspleniaceae (few species), Marattiaceae (*Danaea* pro parte), Osmundaceae
(*Leptopteris*), Pteridaceae (*Adiantum diaphanum*). Absence of stomata in
Hymenophyllaceae, *Hymenophyllopsis* (Cyatheaceae), and *Leptopteris* (Osmundaceae) is
correlated with the reduction of blade thickness and cuticle loss. In contrast to seed
plants, blade tissue structure is very variable in ferns. Information is incomplete, but the
presence or absence of palisade parenchyma may reflect ecologically induced

157. (NCSF 22; PSS 89; R 26; SL 28) **Stipules:** absent (0); present (1). Stipules
are found in many angiosperms, and similar structures are found also in Ophioglossaceae
and Marattiaceae. Their homology is doubtful and the structures may simply be
analogous. Bower (1923, 1926), Goebel (1930), Eames (1936), Hill and Camus (1986),

158. (PSS 9; S 98; SL 55) **Hydathodes:** absent (0); present (1). Not informative
with this dataset. Presence and absence of hydathodes may correlate with ecological

159. (PSS 90; SL 36) **Idioblasts:** absent (0); present (1). The term idioblast is
used for various kinds of specialized cells. In ferns, spicular cells are the most
remarkable type of idioblast. These are sclereids or sclereid-like cells in the epidermis of
the leaf. Such cells are found in different, not closely related, groups of ferns
(Hymenophyllaceae, Marsileaceae, Pteridaceae—including vittarioids). Esau (1977),
I. B. SHOOT

160. (R 4) **Model of growth:** psilotoid (0); selaginelloid (1); cotyledonoid (2).
Composite character based on five criteria (Rothwell 1995); each criterion is reflected by
a separate character in this study.

161. (PSS 91; R 9; SL 10) **Shoot branching:** dichotomous (0); pseudodichotomous (1); anisotomous, sympodial (2); anisotomous, monopodial (3).
Branching patterns are very variable in ferns and are often correlated with growth form
and ecological constraints. The particular pattern does not appear to be informative,
although careful comparative observations are lacking. Bell (1994), Bierhorst (1971),
Bower (1923), Hagemann (1980), Goebel (1930), Kaplan (1977), Kubitzki (1990), Roth
(1963).

162. (New) **Origin of lateral branches:** exogenous (0); endogenous (1). An
unusual origin of the lateral branches, which can be named *endogenous*, was described
for *Equisetum* by Hofmeister (1851) and recently confirmed by Stützel and Jädicke
(2000). Lateral branches are developed exogenously in all other vascular plants, but the
sporophytes of some vascular plants, such as in Marattiales and Ophioglossales, are
generally unbranched. So far, endogenous branching appears to be an apomorphy for
*Equisetum*; therefore the character was excluded as uninformative for judging
relationships among vascular plants. Hofmeister (1851), Stützel and Jädicke (2000).

163. (R 7; SL 7) **Trunk:** absent (0); present (1). Trunk-like shoots have evolved
in various unrelated groups of vascular plants. The character reflects ecological
constraints. Presence of trunks is usually correlated with secondary growth, although
trunks of palms and ferns show that this is not always true. Rothwell (1999) used the
term arborescent for the same character. Kubitzki (1990), Kubitzki et al. (1993).

164. (SL 9) **Stem filiform:** absent (0); present (1). Filiform stems are present in
various unrelated groups of vascular plants. The character reflects ecological constraints.

165. (R 6) **Growth as a liana:** absent (0); present (1). The liana growth form has
evolved independently in various groups of seed plants. Gifford and Foster (1988),
Goebel (1930).

166. (R 14) **Unvascularized enations:** absent (0); present (1). Not informative
with this dataset. Various structures such as scales and thorns are classified as

167. (PSS 32; S 83; SL 44) **Shoot scales:** uniformly colored (0); sharply
bicolored (1); clathrate (2). Not informative with this dataset. Kubitzki (1990).

168. (R 36) **Peripheral loops in the shoot vascular tissue:** absent (0); present
(1). Not informative with this dataset. Stein (1993).

169. (PSS 82; R 52) **Vascular cambium directionality:** unifacial (0); bifacial
(1). Not informative with this dataset. Unifacial cambia are described for Botrychium,
Isoëtes, and some fossil tracheophytes (non-seed plants) with secondary growth. Ogura
Bhambie (1994).

170. (R 51) **Xylem rays:** absent (0); present (1). Not informative with this
dataset. Martens (1971), Esau (1977), Fahn (1990), Kubitzki (1990), Stevenson (1990),
Norstog and Nicholls (1997).
171. (NCSF 11) **Xylem rays structure:** uniseriate/biseriate (0); some multiseriate (1). Not informative with this dataset. Esau (1977), Fahn (1990), Kubitzki (1990), Stevenson (1990), Norstog and Nicholls (1997).

172. (R 47) **Cortical sclerenchyma:** continuous cylinder (0); absent (1); scattered nests or bundles (2); discontinuous cylinder (3). Insufficient information. Ogura (1972).

173. (R 48) **Pith sclerenchyma:** absent (0); present (1). Insufficient information. Ogura (1972).

174. (R 49) **Sclerenchyma accompanying vascular tissue of stele:** absent (0); present (1). Insufficient information. Ogura (1972).

175. (R 12) **Branches alternating with leaves at nodes:** absent (0); present (1). Not informative with this dataset. Bierhorst (1971).

**I. C. ROOT**

176. (PSS 95) **Root hair distribution pattern:** scattered (0); densely matted (1). Not informative with this dataset.

177. (PSS 96) **Sporophyte fungal associations:** absent (0); present, facultative (1); present, obligate (2). Few careful observations are published. Mycorrhizae appear to be correlated with habitat and are absent mainly in epiphytes, aquatics, and xerophytes. New information is also required to improve the definition of character states. Boullard (1957, 1979), Gemma et al. (1992), Moteete et al. (1996), Schmid et al. (1996).

178. (S 17) **Root cortex cells with spiral cell wall thickenings:** absent (0); present (1). Not informative with this dataset. Schneider (1996).
I. D. ANATOMICAL AND MORPHOLOGICAL CHARACTERS THAT ARE APPLICABLE TO MORE THAN ONE SPOROPHYTE ORGAN

179. (PSS 36, R 1, 2, 3) Habitat: terrestrial (0); epiphytic (1); rooted aquatic (2); floating aquatic (3). Each character state reflects a suite of anatomical and morphological characters that may be the result of adaptation. The character is not independent from other characters that are included in this study (e.g., 36, 46). Scoring of actual morphological features rather than ecological features is preferred here for phylogenetic studies because they reflect inherited genetic information and are less likely to be excluded due to assumed ecological functions. Kubitzki (1990).

180. (D 14; NCSF 3) Apical meristems divided into corpus and tunica: absent (0); present (1). Loconte and Stevenson (1990) and Doyle and Donoghue (1992) discuss this character, which is not applicable to meristems with a single apical cell. It is not known if layers in the apical meristems of Lycopsida correspond to the corpus/tunica divisions in seed plants. Guttenberg (1960, 1961), Gifford and Corson (1977), Halperin (1978), Gifford and Foster (1988).


183. (New) Structure of starch grains: pteridophyte-type (0); other types (1). Starch grains of the pteridophyte-type are known for all groups of "pteridophytes" but also for Gingkoatae, Coniferidra, and Gnetidra (Czaja 1978). They are rare in
Magnolidra, which possess starch grains with a more complex structure. Complex starch grains are also found in Cycadaceae. Little is known about the ultrastructure of starch grains in mosses. Further studies are needed because this character may be phylogenetically informative.

184. (New) **Secondary xylem differentiated into tracheary elements and fibers**: absent (0); present (1). This character is informative only for plants with secondary growth. The primary difference between angiosperm wood and all other woods is the differentiation of two cell types (vessels and fibers). Martens (1971), Esau (1977), Fahn (1990).

185. (D 19; NCSF 12; PSS 38, 93) **Vessel perforation (= true vessels or tracheary elements)**: imperforate (0); scalariform or foraminate (1). It has long been accepted that tracheary elements are almost restricted to angiosperms. However, recent studies (e.g., Carlquist and Schneider 1999) have shown their widespread distribution in ferns. Tracheary elements may be absent in only a few groups of tracheophytes, but many taxa are insufficiently studied. Absence of tracheary elements is certain only for *Ginkgo* and *Pinus*. The degree of specialization in tracheary elements may not correlate with phylogenetic position as much as it depends on ecological and physiological factors (Carlquist and Schneider 1999, 2001). Bierhorst (1958, 1971), White (1961, 1963a, b), Martens (1971), Esau (1977), Fahn (1990), Kim et al. (1993), Li et al. (1999), Schneider and Carlquist (1999, 2000).

186. (R 54) **Periderm**: absent (0); present (1). The formation of a periderm is correlated with secondary growth. Peterson (1971), Esau (1977), Khandelwal and Goswami (1979), Fahn (1990), Bhambie (1994).
187. (New) **Sieve tubes:** absent (0); present (1). Sieve tubes are reported only from angiosperms, but detailed studies may find similar structures in other groups of land plants. As with vessels/tracheids the distinction between sieve cells and sieve tubes is not clear. Esau (1969), Behnke and Sjolund (1990).

188. (PSS 37; S 20; SL 14) **Mucilage canals and/or cavities:** absent (0); present (1). Stevenson and Loeconte (1996) combined mucilage-producing hairs and schizogenous tissues into one character, but they are certainly not homologues. Cavities and canals occur in Marattiaceae and some seed plants. The differences between mucilage ducts and laticifers are rather unclear. Ogura (1972), Esau (1977), Hill and Camus (1986), Fahn (1990), Kubitzki (1990), Mahlberg (1993), Datta and Iqbal (1994).

189. (KC 3.15) **Stomata:** absent (0); present (1). Loss of stomata is a sporadic, but widespread, phenomenon in ferns and other land plants. In filmy ferns (Hymenophyllaceae), the character appears to be related to reduction of blade thickness and loss of cuticle. Kubitzki (1990).

190. (PSS 104; S 79, 80, 81, SL 114) **Chromosome numbers:** These are highly variable in some groups while constant in others. No obvious groupings could be made. Duncan and Smith (1977), Löve et al. (1977).

191. (PSS 115) **Apogamous life cycle:** absent (0); present (1). Apogamous life cycles have evolved independently in various groups of vascular plants. Asexual life cycles may be an advantage in different ecological conditions. Sheffield and Bell (1987), Bell (1979).

I. E. **SORI/SPORANGIA/SPORES**
192. (M et al. 106; R 56) **Sporangia borne on leaves:** absent (0), present (1). Character correlated with euphylls. Crane and Kenrick (1997), Kenrick and Crane (1997).

193. (SL 46) **Sporophyll shape:** bifacial (0); peltate or pseudopeltate (1). Not informative with this dataset. Character correlated with presence of lycophylls or euphylls. Kenrick and Crane (1997).


195. (R 63, 64) **Sporangia attached to a modified part of adaxial leaf surface:** absent (0); present (1). All sporangia of Ophioglossaceae are attached to a stalk-like structure (sporangiospore) that has its origin in an early division of the apical meristem of the leaf (Imaichi and Nishida 1986). A very similar condition is found in Psilotaceae (Sieger 1964, 1965, 1967, 1969, 1970, 1973; Rouffa 1978). This structure may be a synapomorphy for the Ophioglossaceae/Psilotaceae clade, but this arrangement of sporangia could also be the basal condition for the Euphyllophytina. It is not possible to interpret the arrangement of the sporangia of extant Equisetopsida without a detailed study that includes the sporangia of fossil sphenopsid groups. Similar studies are needed to determine the condition of early-diverging extinct seed plants. Bierhorst (1971), Page (1972), Taylor (1981), Kato (1988), Stewart and Rothwell (1993), Crane and Kenrick (1997), Gifford and Foster (1988), Grauvogel-Stamm and Ash (1999).

196. (New) **Schizaeoid sorophores:** absent (0); present (1). Sporangia assembled on strongly modified leaf segments or leaf lobes. The character appears to be a putative synapomorphy of schizaeoid ferns (Anemiaceae, Lygodiaceae, Schizaeaceae).
Sori on modified segments occur, however, sporadically in many families, e.g., Tectariaceae, Dryopteridaceae, Woodsiaceae (Deparia), and Polypodiaceae. Bower (1926), Kubitzki (1990), Gandolfo et al. (2000).

197. (R 58) **Sporocarps:** absent (0); present (1). Critical investigations have led to a reassessment of the homology of structures associated with sporangia and sori in heterosporous ferns (Marsileaceae, Salviniaceae), leading to a redefinition of sporocarps. The character is an apomorphy of the heterosporous fern clade. Nagalingum et al. (2006).

198. (R 59) **Monosporangiate sporocarps:** absent (0); present (1). Character based on non-homologous structures (Nagalingum et al. 2006).

199. (SL 62) **Sporocarp germination:** fragmentation (0); gelatinous (1). Informative only for heterosporous ferns (Marsileaceae, Salviniaceae).

200. (SL 58) **Acrostichoid sporangia:** absent (0); present (1). Not informative with this dataset. Kubitzki (1990).

201. (SL 59) **Involucre:** absent (0); present (1). An involucre sensu stricto is restricted to Hymenophyllaceae, but assessment of its homology relative to true indusia, or false indusia, is unclear. Bower (1926). Bierhorst (1971).

202. (PSS 99) **Stomium cell differentiation:** Insufficient information. In derived leptosporangiate ferns, the annulus does not form a continuous bow. It is interrupted by parenchymatous cells that are arranged in three groups: hypostomium, stomium, epistomium. The number of stomium cells differs among the major groups of derived leptosporangiate ferns and this character may be very important in a phylogenetic analysis directed at their relationships. Wilson (1959), Kubitzki (1990), Edwards (1996).
203. (PSS 102) **Spore dimensions:** Dimension of spores is usually correlated with ploidy and no trend was observed that might be pertinent at deeper phylogenetic levels. Tryon and Lugardon (1991).

204. (PSS 100; R 86; SL 70) **Spore shape:** globose (0); tetrahedral (1); monolete (1). Correlated with the shape of laesura. Monolete spores have a linear laesura, while tetrahedral spores have a triradiate laesura. Globose spores have triradiate, circular, or sulcate laesura, but this is not informative for this dataset. Tryon and Lugardon (1991).

205. (PSS 101) **Laesura length:** A highly variable character. Tryon and Lugardon (1991).

206. (SL 80) **Exine (= exospore) type:** Lycopodium-type (0); Selaginella-type (1); Psilotum-type (2); Equisetum-type (3); Ophioglossum-type (4); Osmunda-type (5); Gleichenia-type (6); Filicean-type (7). Insufficient information. Tryon and Lugardon (1991).

207. (R 91) **Fine structure of meio/microspore perine (= perispore):** solid (0); filamentous (1); lamellar (2). Insufficient information. Tryon and Lugardon (1991).

208. (PSS 62; S 43; SL 82) **Spore equatorial flange (cingulum):** absent (0); present (1). Not informative with this dataset. Tryon and Lugardon (1991).

209. (R 90) **Perine forming elaters:** absent (0); present (1). This character is an autapomorphy of the extant genus *Equisetum*. The fossil record of the *Equisetum* lineage indicates that not all members of the lineage formed elaters. In general, elaters were present in various genera of the horsetail lineage but these structures were lacking in others (Taylor and Taylor 1990). The evolution of this character in the horsetail lineage requires further study. Tryon and Lugardon (1991).
210. (SL 83) **Spore proximal flange**: absent (0); present (1). Not informative with this dataset. Tryon and Lugardon (1991).

211. (R 92) **Meio/microspore massulae**: absent (0); present (1). Informative only for heterosporous ferns (Marsileaceae, Salvinia).

212. (PSS 103) **Intine**: absent (0); present (1). Intine (= endospore) appears to be present in all vascular plants, but its development may differ in ferns. Tryon and Lugardon (1991).

213. (New) **Mesospore**: absent (0); present (1). Among extant plants, a true mesospore is known only from megaspores of *Selaginella*. The character is of great interest for studies focused on the relationships of Lycopsida. Taylor (1994).

214. (D 29) **Ovule-bearing structure**: pinnate (0); simple (1). Informative only for seed plants. Gifford and Foster (1988), Kubitzki (1990).

215. (D 33) **Closed carpel with stigmatic pollen germination**: absent (0); present (1). Informative only for seed plants. Gifford and Foster (1988).

216. (D 55) **Integument**: free from nucellus (0); fused more than halfway from the base of the ovule (1). Informative only for seed plants. Gifford and Foster (1988).

II. **GAMETOPHYTE**


218. (SL 87) **Gametophyte development**: *Marattia*-type (0); *Osmunda*-type (1); *Drynaria*-type (2); *Adiantum*-type (3); *Ceratopteris*-type (4); *Aspidium*-type (5); polar

219. (PSS 106) Gametophyte duration: Insufficient information. This character may be correlated with growth form and various other characters (e.g., 97, 102, 103, 131).

220. (PSS 107) Gametophyte growth conditions: This character may be correlated with growth form and various other characters (e.g., 97, 102, 103, 131).

221. (PSS 69; S 137; SL 92) Gametophyte hairs: absent (0); present (1). This character includes various types of hairs, perhaps not homologous, found on gametophytes. Two specialized types of gametophyte hairs are scored here as independent characters (characters 100-101), but a general classification of gametophyte hairs is not possible without detailed ontogenetic evidence. The presence of unicellular or multicellular hairs may reflect certain environmental conditions, although multicellular hairs are restricted to few fern families. Gametophytes of many early-diverging fern lineages are naked. Nayar and Kaur (1971), Schuster (1967, 1992).

222. (PSS 70) Gametophytes green: no (0); yes (1). This character is correlated with gametophyte dependence. Independent gametophytes are green, whereas gametophytes dependent on mycorrhizae, a megaspore, or a sporophyte, are not green. Absence of chlorophyll in the gametophyte signals a change from an autotrophic to a heterotrophic life style. Bierhorst (1971), Nayar and Kaur (1971), Tryon and Tryon (1982), Gifford and Foster (1988), Kubitzki (1990), Tryon and Lugardon (1991).

223. (GR 10; KC 3.24; M et al. 79; PSS 108; SL 99) Multicellular rhizoids: absent (0); present (1). Insufficient information. Multicellular rhizoids are reported to
occur occasionally on the gametophytes of *Botrychium*, *Psilotum*, and *Stromatopteris*. Their occurrence may be related to the obligate mycorrhizae of these gametophytes. Similarly, the lack of early rhizoid development may be correlated with the presence of mycorrhizae (e.g., young gametophytes of *Botrychium*, *Huperzia*, *Ophioglossum*, *Psilotum*, *Stromatopteris*, *Tmesipteris*), or to endospory (e.g., Marsileaceae). Insufficient information precludes the use of other characters of rhizoids, such as forked rhizoids of Hymenophyllaceae. Atkinson and Stokey (1964), Bierhorst (1971), Nayar and Kaur (1971), Whittier and Pintaud (1999).

224. (PSS 109; SL 92, 93, 94) **Gametophyte hair types:** Insufficient information. Atkinson and Stokey (1964), Nayar and Kaur (1971).

225. (PSS 77; SL 89) **Gametophytes gemma-producing:** absent (0); borne from gametophyte thallus (1); borne from gametophyte rhizoids (2). In land plants, gametophytic gemmae differ in their structure and location. Gametophytic gemmae need to be studied using a comparative approach so as to clarify their homologies. Bierhorst (1971), Farrar (1967, 1974), Nayar and Kaur (1971), Farrar and Johnson-Groh (1990), Kubitzki (1990), Dassler and Farrar (1997).


227. (SL 95) **Antheridium size:** massive (0); minute (1). Needs re-evaluation. This character is likely correlated with the number of cells making up the antheridium and the number of sperm cells per antheridium. Atkinson and Stokey (1964), Nayar and Kaur (1971).
228. (SL 97) **Antheridial basal cell origin:** antheridial initial (0); thallus (1).

229. (GRD 72; M et al. 53) **Mitochondria associated with plastids in young spermatids:** absent (0); present (1). Insufficient information. Duckett and Bell (1977), Garbary et al. (1993), Maden et al. (1997), Garbary and Renzaglia (1998), Renzaglia et al. (1999).

230. (GRD 74; M et al. 54) **Specialized anterior mitochondrion in spermatids:**
present (0); absent (1). Insufficient information for this dataset. Duckett and Bell (1977), Garbary et al. (1993), Maden et al. (1997), Garbary and Renzaglia (1998), Renzaglia et al. (1999).

231. (M et al. 55) **Specialized posterior mitochondrion in spermatids:** present (0); absent (1). Insufficient information for this dataset.

232. (GRD 75; M et al. 56) **Additional mitochondria in anterior of cell in spermatids:**
absent (0); row of mitochondria behind anterior mitochondrion (1); numerous, unspecialized mitochondria (2). Insufficient information. Duckett and Bell (1977), Garbary et al. (1993), Maden et al. (1997), Garbary and Renzaglia (1998), Renzaglia et al. (1999).

233. (M et al. 57) **Origin of anterior mitochondrion:** fusion (0); elongation (1).

234. (GRD 77; M et al. 58) **Osmophilic material underneath anterior mitochondrion:**
absent (0); present (1). Insufficient information. Duckett and Bell
(1977), Garbary et al. (1993), Maden et al. (1997), Garbary and Renzaglia (1998),
Renzaglia et al. (1999).


**236. (R 81) Coenocytic development of megagametophyte:** absent (0); present (1). Not informative with this dataset.

**237. (R 82) Number of megaspores:** more than two (0); one to two (1). Not informative with this dataset.

**238. (R 84) Nonhydrasperman pollen chamber:** present (0); absent (1). Not informative with this dataset.

**239. (R 86) Integument with micropyle:** absent (0); present (1). Not informative with this dataset.

**240. (D 83) Fusion of sperm cells:** fusion of only one sperm cell with a female gametophyte nucleus (0); regular fusion of both sperm cells (1). These processes appear to be quite distinct in angiosperms and Gnetales and various authors have expressed their doubts about a common origin of the character in both seed plant lineages (Doyle 1998b, Friedman and Floyd 2001). Friedman and Carmichael (1996), Gifford and Foster (1988).

### III. EMBRYO

**241. (New) Polyembryony:** present (0); absent (1). Insufficient information. Bierhorst (1971), Nayar and Kaur (1971).

**242. (M et al. 98, 99) Gametophyte-sporophyte junction:** Insufficient information. The presence and structure of transfer cells on the gametophyte and/or
sporophyte may be important conditions in the evolution of land plants. Wardlaw (1965), Bierhorst (1971), Gifford and Foster (1988), Ligrone et al. (1993).

IV. LIFE CYCLE

243. (New) **Life cycle length (spore to spore):** more than two years (0); one to two years (1); less than one year (2). Some early-diverging fern groups have a very long life cycle length with more than three to five years (e.g., *Botrychium*) required from spore to spore. This is also the case for Lycopodiaceae. In contrast, most leptosporangiate ferns complete their life cycle in one or two years. Not enough data are available at this time, and taxa with long life cycles may have a heterotrophic gametophyte with obligate mycorrhizae. The length of the life cycle is influenced by the plant habit, especially in large arborescent sporophytes, which need years before they produce spores. The time-span of the sporophytic generation is very variable in some lineages of seed plants. Kubitzki (1990).

V. BIOCHEMISTRY

244. (New) **Auxin metabolism:** low IAA conjugate level (0); high IAA conjugate level (1). Insufficient information. Modifications of the auxin metabolism pathway may be important in the evolution of plant development. Stein et al. (1999).

245. (New) **Product of chlorophyll degradation:** RCCR-3 (0); RCCR-0, RCCR-1 or RCCR-2 (1). Insufficient information. Present data indicate RCCR-3 to be the product in Lycophyta, several fern lineages (including Equisetopsida, Marattiales, Psilotales), and some lineages of seed plants. RCCR-0 and RCCR-2 are found in
leptosporangiate ferns (Polypodiopsida), several genera of conifers (Coniferidra), and angiosperms (Magnolidra). Hoertensteiner et al. (2000).

VI. MOLECULAR DATA

246. (New) *ndhB duplication in cpDNA*: absent (0); present (1). Insufficient information. Raubeson and Stein (1995).

247. (New) *psbA plus two inversions in cpDNA*: absent (0); present (1). Insufficient information. Stein et al. (1992), Raubeson and Stein (1995).

248. (New) *chlL partial duplication in cpDNA*: absent (0); present (1). Insufficient information. Raubeson and Stein (1995).

249. (New) *cox2.i3 intron in mtDNA*: absent (0); present (1). Insufficient information. Qiu et al. (1998) and Qiu pers. comm.

250. (New) *cox2.i4 intron in mtDNA*: absent (0); present (1). Insufficient information. Qiu et al. (1998) and Qiu pers. comm.

251. (New) *Group II intron in nad5 gene in mtDNA*: absent (0); present (1). Insufficient information. Vangerow et al. (1999) and Vangerow pers. comm.

252. (New) *Second intron in nad1 gene in mtDNA*: absent (0); present (1). Insufficient information. Gugerli et al. (2001).


256. (New) **Nuclear actin genes**: Insufficient information to define homology and occurrence of gene copies. Bhattacharya et al. (2000).


**Literature Cited**

[For complete Literature Cited see printed manuscript. References below appear only in Appendix 3].


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