PHYLOGENETIC RELATIONSHIPS OF THE "GREEN ALGAE" AND "BRYOPHYTES"1

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ABSTRACT

Considerable progress has been made recently, based on classical morphological characters and newly described ultrastructural features, in understanding the phylogenetic relationships of the tracheophytes to the green algae and bryophytes. Recent technological advances in molecular biology, particularly the advent of the polymerase chain reaction (PCR), have allowed nucleotide sequence data relevant to such large-scale phylogenetic questions to accumulate, especially ribosomal RNA gene sequences (both the large and small subunits) from the nucleus and the chloroplast. We present synthetic cladistic analyses of the green plants that combine and compare available morphological and molecular data sets. Although the resulting phylogenies are poorly resolved in some areas at present, certain conclusions are supported: (1) The green plants are composed of two major monophyletic groups, one containing the "charophyte" green algae and the land plants (i.e., "bryophytes" plus tracheophytes), the other containing the bulk of the classically delimited "green algae" (chlorophytes, pleurastrophytes, and ulvophytes). (2) The land plants are a well-supported monophyletic group, but neither the specific outgroup for the land plants nor the precise relationships among basal lineages of land plants are all lineages of land plants are all morphological analysis). lineages of land plants are clear. In many analyses (including the combined molecular and morphological analysis) the three major lineages (i.e., liverworts, hornworts, and mosses) appear to be paraphyletic with respect to the tracheophytes, with an indication that the mosses alone may be the sister group of the tracheophytes; however, in other analysis and the sister group of the tracheophytes, and the sister group of the tracheophytes are group of the sister group of the tracheophytes are group of the sister group other analyses the "bryophytes" are supported as a monophyletic group. (3) The ulvophytes, chlorophytes, and pleurastrophytes are each supported as monophyletic (with the exception of a few taxa that may be misplaced in the current classification), with the topology: [ulvophytes [chlorophytes + pleurastrophytes]]. Combined analyses of molecular and morphological data offer the greatest potential for resolving these relationships.

Reconstruction of the broad-scale phylogenetic relationships of green plants is important to our understanding of major evolutionary events such as the origin of multicellularity, diversification of life-history strategies, and the conquest of land (Graham, 1985; Mishler & Churchill, 1985). In addition, availability of a well-supported framework of "deep" relationships is necessary for purposes of outgroup comparison in studies of tracheophyte phylogeny (Crane, 1990; Gensel, 1992).

Considerable morphological and ultrastructural data have accumulated over the last two decades that bear on the question of phylogenetic relationships of the green algae and bryophytes to the tracheophytes (e.g., Stewart & Mattox, 1975; Hébant, 1977; Pickett-Heaps, 1979; Crandall-Stotler, 1980, 1981; Brown & Lemmon, 1988; Carothers & Rushing, 1988; Duckett & Renzaglia, 1988; Ligrone & Gambardella, 1988). More recently, comparative molecular data have become

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available as well (e.g., Kantz et al., 1990; Zechman et al., 1990; Lewis et al., 1992; Mishler et al., 1992; Waters et al., 1992; Wilcox et al., 1992). A few attempts have been made to synthesize cladistically this growing database (Mishler & Churchill, 1984, 1985; Sluiman, 1985; Theriot, 1988; Graham et al., 1991; Garbary et al., 1993); none of these, however, have incorporated the newly available molecular data.

Cladistic studies to date suggest that neither the "green algae" nor the "bryophytes" are monophyletic. The green plants appear to be composed of two major lineages and a residuum of unicellular micromonadophytes. One of these major lineages contains the bulk of the classical green algae (Chlorophyceae, Pleurastrophyceae, and Ulvophyceae sensu Mattox & Stewart, 1984). There is an indication that the ulvophytes are basal to the chlorophytes plus pleurastrophytes based on morphological and ultrastructural data (Stewart & Mattox, 1975; Mattox & Stewart, 1984; O'Kelly & Floyd, 1984; Sluiman, 1985). Mishler & Churchill (1985) questioned the monophyly of the ulvophytes because of a lack of morphological synapomorphies to unite that group. This result was supported by a recent molecular study (Zechman et al., 1990) that also found non-monophyly of the chlorophytes. The pleurastrophytes have been treated in three ways: (1) as a separate sister class to the Chlorophyceae (Mattox & Stewart, 1984); (2) as part of the Chlorophyceae (Melkonian, 1990); or (3) as part of the Ulvophyceae (Sluiman, 1989).

The other major lineage of green plants includes the charophycean green algae plus the land plants (i.e., bryophytes plus tracheophytes). The genus Coleochaete (or even some part of it alone) appears to be the closest extant sister group of land plants (Graham et al., 1991). The bryophytes are composed of three distinctive lineages (i.e., liverworts, hornworts, mosses) that may be paraphyletic with respect to the tracheophytes. For example, early analyses of morphological data (Mishler & Churchill, 1984), as well as two recent molecular analyses (Mishler et al., 1992; Waters et al., 1992), concluded that the liverworts alone appear to be the basal lineage within extant land plants, i.e., the sister group to the hornworts, mosses, and tracheophytes. General morphological data (Mishler & Churchill, 1984), ultrastructural data (Theriot, 1988), and one molecular data set (Mishler et al., 1992) support a topology with the mosses alone as the sister group of tracheophytes. The phylogenetic placement of the hornworts is not clearly resolved by any of the published data sets. On the other hand, monophyly of the bryophytes (including Selaginella) was suggested by a recent cladistic analysis based on sperm ultrastructural data (Garbary et al., 1993).

Molecular sequence data have shown considerable promise for phylogenetic analysis, but they provide no panacea (despite overly optimistic claims in recent literature, e.g., Graur, 1993). In fact, theoretical considerations predict that DNA sequence characters (given their quasi-clocklike evolution and limited number of character states) could be especially problematical in "deep" phylogenetic reconstructions, where considerable asymmetry in branch lengths exists (Felsenstein, 1978; Mishler et al., 1988; Albert et al., 1992; Donoghue & Sanderson, 1992; Albert et al., 1993; Mishler, 1994). Careful evaluation of all potential characters is required; it is necessary to apply to molecular data basic principles of character analysis (for deriving strong, independent hypotheses of character homology) and cladistic analysis (for evaluating the phylogenetic "signal," if any, present in the resulting data set). Theoretical issues that must be faced in large-scale, synthetic analyses include further development of methods for: (1) combining/ comparing data sets of fundamentally different natures (including issues of character and characterstate weighting; Miyamoto, 1985; Kluge, 1989; Albert & Mishler, 1992; Albert et al., 1992; Donoghue & Sanderson, 1992; Albert et al., 1993); (2) assessing support for clades (e.g., bootstrap vs. the decay index; Mishler et al., 1991; Källersjö et al., 1992); and (3) representing diverse, yet clearly monophyletic, clades (e.g., the exemplar method vs. "compartmentalization" - a new approach involving substituting an inferred "archetype" or hypothetical ancestor for a clade accepted as monophyletic a priori in an inclusive analysis: Mishler, 1994).

When examined carefully, it is evident that different genes are phylogenetically useful at different hierarchical levels. Nuclear-encoded ribosomal RNA gene sequences (both the 26S and 18S subunits) provide data of apparent utility at the highest level of green plant phylogeny (Zimmer et al., 1989; Kantz et al., 1990; Zechman et al., 1990; Lewis et al., 1992; Waters et al., 1992; Wilcox et al., 1992). On the other hand, the 5S rRNA nuclearencoded gene appears to be too small, and has had too many substitutions (in the positions that vary), to be of use at this level (Bremer et al., 1987; Mishler et al., 1988; Steele et al., 1991). Among chloroplast genes, it appears that the 16S rRNA gene is the most conservative, followed by the 235 rRNA gene (Palmer et al., 1988). Combined data from these two genes have provided corroboration of some previous morphologically based hypotheses about the relative branching order of the major lineages of land plants (Mishler et al., 1992). In contrast, the protein-coding gene rbcL has proven too divergent to be useful among these lineages (Manhart, pers. comm.; Mishler, unpublished data), but it has provided consistent systematic characters within major lineages. For example, within the mosses the relatively basal position of Sphagnum and Andreaea is supported, as is the monophyly of the peristomate mosses (with the nematodontous taxa in a more basal position within) and monophyly of the haplolepidious mosses (Mishler, unpublished data).

It is likely that a robust and highly resolved phylogeny can be produced for the green plants in the next few years by integrating classical morphological characters with newly described ultrastructural features and various sets of molecular sequence data from the nuclear and chloroplast genomes. Careful choice of characters and application of proper methods of analysis will be essential, however. The co-authors do not agree unanimously on the homology of all characters as used here, but have found cladistic analysis to be an excellent way to frame arguments objectively. This paper attempts to synthesize data published to date, as a guide (and a target) for ongoing projects in our own and other laboratories. More data are certainly needed, but a comparison of currently available data is of interest in its own right and can assist in the identification of characters and taxa that are crucial for future research. The data sets presented here will be made generally available (annotated MacCLADE files will be sent on request) and, thus, will provide a basis for future synthetic studies

MATERIALS AND METHODS

DATA SETS

Six data sets were assembled in pairs (focused on different phylogenetic levels): two morphological data sets, one large and one small molecular data set, and two combined data sets. The sources and characteristics of these data sets are listed below:

(1) A large-scale green plant morphological data set (GP-MORPH) is focused primarily on the "deep" relationships of the green plants as a whole (thus using many characters from "green algal" systematics). Choice of OTUs was dictated by availability of molecular data for comparative purposes (see GP-MOLEC below). Several representative land plants were included (14 in all). Choice of characters was based on standard criteria of homology

and independence (Mishler, 1994); a list of characters and character states can be found in Table 1. Sources include Stewart & Mattox (1975), Mattox & Stewart (1984), Mishler & Churchill (1984, 1985), and Graham et al. (1991). All 110 characters were treated as unordered. A hypothetical ancestor was coded for rooting purposes, based on generalized states in presumed protistan outgroups. The data matrix is shown in Table 2.

(2) A more focused land plant morphological data set (LP-MORPH) is based on the set of OTUs scored by Garbary et al. (1993) for ultrastructural characters of spermatozoids. A list of characters and character states can be found in Table 3. To a modified and somewhat reduced set of their "sperm" characters (numbers 1–65), we added a set of "general morphological" characters (numbers 66–113), beginning with the Mishler & Churchill (1984, 1985) characters (modified). Of these 113 characters, only two (7 and 50) were treated as ordered. Two charophytes were included as outgroups. The data matrix is shown in Table 4.

(3) A large-scale molecular data set (GP-MO-LEC) represents a realignment of published nuclear rDNA sequences from selected taxa across the green plants. Full small subunit (18S) sequences were taken from Rausch et al. (1989), Huss & Sogin (1990), Lewis et al. (1992), Wilcox et al. (1992), and others. Partial 18S and large subunit (26S) sequences were taken from Buchheim et al. (1990), Kantz et al. (1990), Zechman et al. (1990), and Chapman & Buchheim (1991). OTUs and sources of data are shown in Table 5. All intron sequences were excluded. Those taxa that seemed repetitive at this phylogenetic level (i.e., multiple, very similar species from the same genus) were deleted, and the alignment was adjusted by eye to take into account this full set of taxa. The data matrix had 61 OTUs, including Emiliana and Anemonia as outgroups, and 2179 characters (of these, 1833 are from 18S and 346 are from 26S). Only 37 of the OTUs had 26S data available; the remainder were coded with all question marks for this region. A number of OTUs have considerable missing data between primer regions; such positions were coded with a question mark. To be conservative, gaps were also coded as missing data. For the 18S data, 17 regions (totaling 206 positions) were excluded from analysis because they could not be aligned unequivocally. The data set is too large to publish, but an aligned and annotated PAUP file is available from the authors on request.

(4) A smaller "land plant only" molecular data set (LP-MOLEC) was excerpted from the larger set above (GP-MOLEC); the 16 OTUs (including TABLE 1. List of characters and character states used for data matrix GP-MORPH. See text for source of these characters and Table 2 for the data matrix.

- 1. Habitat of free-living vegetative stage: 0, freshwater; 1, brackish or marine; 2, terrestrial.
- 2. Life history: 0, haplontic; 1, diplontic; 2, isomorphic alternation; 3, heteromorphic alternation.
- 3. Vegetative cell or thallus attached to substrate: 0, no; 1, yes.
- 4. Radial symmetry, if multicellular: 0, no; 1, yes.
- 5. Growth form: 0, unicellular or coccoid; 1, multicellular; 2, coenobic.
- 6. Vegetative cells contiguous in multicellular organism: 0, no; 1, yes.
- 7. Multinucleate vegetative cells: 0, no; 1, yes.
- 8. Coenocytic: 0, no; 1, yes.
- 9. Distromatic foliar thalli: 0, absent; 1, present.
- 10. Plasmodesmata: 0, absent; 1, present.
- 11. Parenchyma: 0, absent; 1, present.
- 12. Vegetative cells form filaments: 0, no; 1, yes, unbranched; 2, yes, branched; 3, yes, multi-axial.
- 13. Filaments with acuminate tips: 0, no; 1, yes.
- 14. Vegetative cells or zoospores spindle-shaped: 0, no; 1, ves.
- 15. Zoospores: 0, absent; 1, present; 2, present, flattened.
- 16. Autospores/colonies: 0, no; 1, yes.
- 17. Vegetative cell with flagella: 0, no; 1, yes.
- 18. Gamete production: 0, holocarpic; 1, heterocarpic.
- 19. Multiple sporulation/fission: 0, no; 1, yes.
- 20. Type of sex: 0, isogamy; 1, anisogamy; 2, oogamy.
- 21. Chloroplast shape: 0, cup; 1, reticulated; 2, lateral cup; 3, H-shaped; 4, bi-polar; 5, sets of complete rings; 6, incomplete rings; 7, multiple disks; 8, spiral; 9, stellate; 10, plate; 11, axile.
- 22. Pyrenoids: 0, absent; 1, present.
- 23. Thylakoid membranes traverse pyrenoid: 0, no; 1, yes.
- 24. Number of flagella on vegetative cells or zoospores: 0, 2; 1, 4; 2, 1; 3, 4+; 4, 0.
- 25. Retraction of flagella during division: 0, no; 1, yes.
- 26. Angle of basal bodies relative to direction of motion: 0, angled; 1, perpendicular; 2, parallel.
- 27. Flagellar beat: 0, trailing-undulating; 1, breast stroke.
- 28. Basal bodies distant via migration in development: 0, no; 1, yes.
- 29. Flagella extend to right on motile cells: 0, no; 1, yes.
- 30. Flagellar apparatus displaying 180 degree rotational symmetry: 0, no; 1, yes.
- 31. Absolute orientation: 0, counterclockwise; 1, clockwise; 2, direct opposite.
- 32. Basal body overlap in motile cells: 0, absent; 1, present.
- 33. Basal body core connection: 0, absent; 1, present.
- 34. Mitotic spindle type: 0, metacentric; 1, centric.
- 35. Mitotic spindle closed: 0, absent; 1, present.
- 36. Spindle collapsing at telophase: 0, absent; 1, present.
- 37. Cupping microtubules surround centrioles during mitosis: 0, no; 1, yes.
- 38. Microtubules forming in plane of cell division: 0, absent; 1, present.
- 39. Phragmoplast: 0, no; 1, yes.
- 40. Cell plate in cytokinesis: 0, no; 1, yes.
- 41. Centrioles between nucleus and plane of cleavage: 0, no; 1, yes.
- 42. Lactate fermentation: 0, absent; 1, present.
- 43. Chaetophoralean autolysin lyses sporangium: 0, no; 1, yes.
- 44. Hydrogenase produced by vegetative cells: 0, no; 1, yes.
- 45. Secondary carotenoids: 0, no; 1, yes.
- 46. Siphonoxanthin: 0, absent; 1, present.
- 47. Gelatin liquifaction: 0, no; 1, yes.
- 48. Photosystem II light harvesting complex: 0, low molecular weight; 1, high molecular weight.
- 49. Dormant zygote produced: 0, no; 1, yes.
- 50. Sporulation: 0, absent; 1, present.
- 51. Zellteilung [vs. sporulation]: 0, no; 1, yes.
- 52. Common matrix surrounds cells: 0, no; 1, yes.
- 53. Papillae on vegetative cells: 0, no; 1, yes.
- 54. Crystalline cell wall: 0, no; 1, yes.
- 55. Stigma: 0, no; 1, yes.
- 56. Number of contractile vacuoles: 0, 2; 1, 2+; 2, 1; 3, absent.
- 57. Apical insertion of flagella: 0, no; 1, yes.
- 58. Zoosporangia abscise: 0, no; 1, yes.
- 59. Zoosporangia operculate: 0, no; 1, yes.
- 60. Zoosporangial exit plug: 0, no; 1, yes.
- 61. Keeled flagella: 0, no; 1, yes.
- 62. Urea amidolyase produced: 0, no; 1, yes.

TABLE 1. Continued.

TA	BLE 1. Continued.
63.	Terminal cap: 0, absent; 1, bilobed; 2, platelike.
64.	Prominent proximal sheath: 0, no; 1, yes.
	Organic scales/covering: 0, no; 1, yes.
	Transverse septum: 0, absent; 1, present.
67.	Proximal septum: 0, absent; 1, present.
	SMAC or system 1 fiber: 0, absent; 1, present.
69.	Diaphasis: 0, absent; 1, present.
70.	Distal fiber in motile cell: 0, absent; 1, present.
71.	Specialized zoosporangia: 0, absent; 1, present.
72.	MLS present: 0, no; 1, yes.
74	Glycollate oxidase: 0, no; 1, yes.
75	Oogonium associated with sterile cells: 0, no; 1, yes.
76	Eggs retained in oogonium: 0, no; 1, yes.
77	Apical cell growth: 0, no; 1, yes.
78	Flavonoids: 0, no; 1, yes. Zygote retained: 0, no; 1, yes.
79.	Placental transfer cells: 0, no; 1, yes.
80.	True antheridia: 0, no; 1, yes.
81.	Archegonia: 0, no; 1, yes.
82.	Embryo: 0, no; 1, yes.
83.	Cuticle: 0, no; 1 yes.
84.	Monoterpenes: 0, no; 1, yes.
85.	Lunularic acid: 0, no; 1, yes.
80.	Elaters: 0, no; 1, yes.
87.	Oil bodies: 0, no; 1, yes.
88.	D-Methionine distinguished: 0, no; 1, yes.
09.	Stomata: 0, no; 1, yes.
90.	Vertical division of zygote: 0, no; 1, yes.
92	Pseudoelaters: 0, no; 1, yes.
93.	Xylem: 0, no; 1, yes. Phloem: 0, no; 1, yes.
94.	Perine on spores: 0, no; 1, yes.
95.	Aerial sporophyte axis: 0, no; 1, yes.
20.	Columella in sporangium: () no: 1 ves
21.	Multicellular rhizoids: 0, no: 1, ves.
70.	Leaves on gametophyte (of moss type): U. no: 1. ves.
77.	Articulated peristome: () no: ves.
100.	independent sporophyte: () no: ves
LUI.	Dranched sporophyte: (), no: 1, ves.
	Ornamented tracheid walls: () no: 1 ves.
100.	True lignin: () no: 1 ves
104.	Megaphylls: (), no: 1 ves
106	Trichomes: 0, no; 1, yes.
107	Vascular cambium: 0, no; 1, yes. Eustele: 0, no; 1, yes.
108.	Seeds: 0, no; 1, yes.

two charophyte outgroups) used are marked in Table 5. The alignment was adjusted based on the land plant sequences alone. This was done because the phylogenetic resolution was poor with the larger GP-MOLEC data set in this part of the green plants. An aligned and annotated PAUP file is available from the authors on request.

108. Seeds: 0, no; 1, yes.

110. Flowers: 0, no; 1, yes.

109. Axillary branching: 0, no; 1, yes.

(5) A combined data set (GP-COMB) was produced by combining GP-MOLEC and GP-MORPH. This data set omitted the outgroups *Emiliana* and *Anemonia* from GP-MOLEC and the hypothetical

ancestor from GP-MORPH, but otherwise used the full combined data as described separately above for 59 OTUs. No weighting was done.

(6) A smaller combined data set (LP-COMB) that focused on the land plants was produced by combining morphological data from LP-MORPH with sequence data from GP-MOLEC. These data sets are unfortunately severely nonoverlapping; only nine taxa have data in both sets when "composite" OTUs are constructed pairing presumably related, but not identical, taxa. The composition of the nine

	1 0	2	3	4 0	District Control of the		3 9	1
Glycine max	2310110001						TO BE TO SERVE	
Oryza sativa	2310110001							
Zamia floridana	2310110001							
Psilotum sp.	2310110001							
Equisetum hymale	2310110001							
Atrichum angustatum	2310110001							
Fissidens taxifolius	2310110001							
Plagiomnium cuspidatum	2310110001							
Notothylas breutellii	2310110001							
Phaeoceros laevis	2310110001 1							
Porella pinnata	2310110001 1							
Conocephalum conicum	2310110001 1							
Asterella tenella	2310110001 1							
Riccia austinii	2310110001 1							
Coleochaete nitellarum	0010110001 1							
Klebsormidium flaccidum	00001?0001 0							
Micromonas pusila	1?00000000 0							
Mantoniella squamata	1?00000000 0							
Nephroselmis pyriformis	1700000000 0							
Pedinomonas minutissima	1?00000000 0							
Tetraselmis carteriiformis	1?00000000							
Enteromorpha intestinalis	1210110010							
Ulva fasciata	1210110010							
Ulothrix zonata	0310110000							

TABLE 2. Data matrix GP-MORPH. See Table 1 for list of characters and states, and Table 5 for list of taxa.

Cymopolia barbata Batophora oerstedtii Codium decorticatum Cladophoropsis membranosa Blastophysa rhizopus Trentepohlia sp. Cephaleuros parasiticus Characium vacuolatum Dunaliella parva Chlamydomonas reinhardtii Volvox carteri Chlorococcopsis min Draparnaldia plumosa Uronema belkae Chlamydomonas moewusii Stephanosphaera pluvialis Carteria radiosa Gonium pectorale Chlorella kessleri Chlorella vulgaris Prototheca wickerhamii Chlorella protothecoides Chlorella minutissima Neochloris aquaticus

TABLE 2. Continued.

										1	1
	1 0	2 0	3 0	4 0	5	6		8	9	0	1
Neochloris vigenis	0000001000	0000100010	?1000??001	2011110100	?????0??01	0000??1000	0?00010?01	00?000000	000000000	000000000	0000000000
Pediastrum duplex	0000210000	0000100010	?1?0011001	2011110100	1????0??11	00001?1000	0?01010101	00?0000000	000000000	0000000000	0000000000
Scenedesmus obliquus	0000210000	0000010010	?1?40???0?	???1110100	???1101??1	0000???000	01000???0?	0000000000	000000000	0000000000	0000000000
Characium hindakii	0010001000	0001100010	?1000?1001	20?1110100	?????0??01	0000??1000	0?00010?01	002000000	000000000	0000000000	0000000000
Chlorella fusca	0000000000	0000010?1?	01??0?????	??????0100	?1?1101??1	000??3?000	????0???0?	0??0000000	0000000000	0000000000	0000000000
Ankistrodesmus falcatus	0000000000	0001010?1?	?1??0?????	??????0100	?????0???1	000??3?000	????0???0?	0??0000000	000000000	0000000000	0000000000
Pseudotrebouxia gigantea	0000000000	0000200707	?1000?1001	0?00111100	0????0???1	0000?31000	0?00010001	002000000	0000000000	000000000	0000000000
Pleurastrum terrestre	0000110000	0100200707	?1000?1001	0?00111100	0????0???1	0000?31000	0?00010001	00?000000	0000000000	000000000	0000000000
Characium perforatum	0010000000	0001100010	?1100?1001	010?11?000	?????0??01	0000??1000	0?00010?01	00?000000	0000000000	0000000000	0000000000
Parietochloris pseudo	0000000000	0000100010	?1100?1001	010?11?000	?????0??01	0000??1000	0?20010001	00?0000000	0000000000	000000000	0000000000
Friedmannia israelensis	0/200000000	000020000?	?100011001	0100111100	0????0???1	0000?31000	0?00010001	000000000	000000000	000000000	000000000
Hypothetical ancestor	1000000000	0000101010	2000000000	2100000000	0????0?001	0000030000	0000000000	0100000000	000000000	000000000	0000000000

OTUs (including *Coleochaete* as outgroup) in this data set is shown in Table 6.

ANALYSES

A number of parsimony analyses were carried out to examine the phylogenetic implications of these data sets alone and in combination. Data sets were compiled and phylogenetic trees examined using MacCLADE, version 3.01 (Maddison & Maddison, 1992). All analyses were carried out using PAUP, version 3.1 (Swofford, 1991) on an Apple Macintosh Quadra 700 with 20 megabytes of RAM. With the larger data sets, heuristic search algorithms were necessary, thus finding of all the maximum-parsimony (MP) trees cannot be guaranteed.

The analyses are presented in eight groups below, using standard numbers that are used for reference in the Results and the Discussion. The PAUP commands (Swofford, 1991) employed in each search are shown. "Decay analysis" performed in some cases refers to the saving of trees longer than the MP tree(s) by the specified number of steps (Bremer, 1988; Graham et al., 1991; Mishler et al., 1991; Donoghue et al., 1992; Källersjö et al., 1992)—the "decay index" is the number of steps parsimony must be relaxed to cause a particular clade to lose its support. For example, a decay index of 2 for a clade means that it is present in the semi-strict consensus (Bremer, 1990) of the MP trees plus those that are one step longer (called decay class "1"), but absent in the semi-strict consensus of the MP trees plus those that are one or two steps longer (called decay class "2").

- Analysis 1. Land plant morphological data alone (LP-MORPH):
- la. All characters; 100 repetitions of RAN-DOM taxon addition, TBR branch swapping with MULPARS and STEEPEST DESCENT option; decay analysis to 3 steps.
- lb. Sperm data alone (i.e., characters 1-65); 100 repetitions of RANDOM taxon addition, TBR branch swapping with MULPARS and STEEPEST DESCENT option; decay analysis to 3 steps. CONSTRAINT analyses were also done (using the same PAUP options), to find the shortest topologies consistent with [mosses + tracheophytes] and [hornworts + mosses + tracheophytes] as monophyletic groups.
- lc. General morphological data alone (i.e., characters 66-113); 100 repetitions of RANDOM

- taxon addition, TBR branch swapping with MUL-PARS and STEEPEST DESCENT option; decay analysis to 3 steps. CONSTRAINT analyses were also done (using the same PAUP options), to find the shortest topologies consistent with monophyly of the bryophytes.
- ld. Sperm data alone from the nine OTUs selected as part of LP-COMB; branch-and-bound search.
- 1e. General morphological data alone from the nine OTUs selected as part of LP-COMB; branch-and-bound search.
- 1f. All characters from the nine OTUs selected as part of LP-COMB; branch-and-bound search.
- Analysis 2. Green plant morphological data alone (GP-MORPH). CLOSEST taxon addition, TBR branch swapping with MULPARS and STEEPEST DESCENT option.
- Analysis 3. Green plant molecular data alone (GP-MOLEC). 18S and 26S data combined:
- 3a. All OTUs except two outgroups (59 OTUs); CLOSEST taxon addition, NNI branch swapping with MULPARS, followed by TBR branch swapping with MULPARS on the shortest trees found by NNI swapping.
- 3b. Land plants alone plus two charophyte outgroups (16 OTUs); 10 repetitions of RANDOM taxon addition, TBR branch swapping with MUL-PARS.
- 3c. The nine land plant OTUs selected as part of LP-COMB; branch-and-bound search.
- 3d. Green algae minus charophytes, land plants, *Emiliana*, and *Anemonia* (43 OTUs); CLOSEST taxon addition, NNI branch swapping with MULPARS, followed by TBR branch swapping with MULPARS and STEEPEST DESCENT on the shortest trees found by NNI swapping.
- Analysis 4. Green plant molecular data (GP-MO-LEC). 18S data alone:
- 4a. All OTUs including the two outgroups, Emiliana and Anemonia (61 OTUs); 10 repetitions of RANDOM taxon addition, NNI branch swapping with MULPARS, followed by TBR branch swapping with MULPARS on the shortest trees found by NNI swapping; decay analysis to 2 steps.
 - 4b. Land plants only plus Coleochaete as out-

TABLE 3. List of characters and character states used for data matrix LP-MORPH. See text for source of these characters and Table 4 for the data matrix. All characters are considered unordered, except as noted.

1. Apical cell in antheridia: 0, absent; 1, present.

2. Division pattern in young antheridia: 0, four-celled; 1, two-celled.

3. Endogenous antheridia: 0, absent; 1, present.

4. Antheridial stalk: 0, absent; 1, present. 5. Operculum cells: 0, absent; 1, present.

6. Sperm in pollen tube: 0, absent; 1, present.

7. Number of sperm per male structure (ordered character): 0, 1000+; 1, 100-1000; 2, 16-24; 3, 2.

8. Nascent spermatids: 0, paired; 1, not paired.

9. Diagonal spindle in final mitotic division: 0, absent; 1, present.

10. Replication of the centrioles: 0, present; 1, absent.

- 11. Time of origin of centrioles: 0, always present; 1, sperm mother cells; 2, sperm mother cell progenitor; 3, earlier.
- 12. Basal bodies (BB) and flagella: 0, two; 1, more than two.

13. Bicentrioles: 0, present; 1, absent.

- 14. Basal body position: 0, right angles; 1, side-by-side; 2, staggered anterior-posterior; 3, staggered continuous.
- 15. Proximal extension A: 0, absent; 1, long; 2, short. 16. Proximal extension B: 0, ventral-dorsal; 1, ventral.

17. Stellate transition: 0, present; 1, absent.

18. Connecting fibers between BBs: 0, present; 1, absent; 2, fine filaments with centrin.

19. Basal body structure: 0, monomorphic; 1, dimorphic.

20. BB staggering associated with microtubule growth: 0, absent; 1, present.

21. Regression of lamellar strip: 0, absent; 1, complete; 2, partial.

22. Lamellar strip/anterior mitochondrion elongation: 0, parallel; 1, perpendicular.

23. Spline aperture: 0, absent; 1, present.

- 24. Spline aperture location: 0, left of center; 1, right of center. 25. Position of developing MLS: 0, adjacent to BBs; 1, beneath BBs.
- 26. Plaque stratified between blepharoplast: 0, absent; 1, present.

27. Spline/lamellar strip orientation: 0, 90°; 1, 45°.

28. Posterior notch to lamellar strip: 0, absent; 1, present.

29. Lamellar strip position: 0, under all BBs; 1, under anterior BB only; 2, under some BBs.

30. Stray spline microtubule: 0, absent; 1, present; 2, develops late.

31. Accessory band of microtubules: 0, absent; 1, present.

32. Maturational elongation of anterior mitochondrion: 0, absent; 1, posterior.

33. Spline shank: 0, wide; 1, less than 4 tubules.

34. Osmiophilic crest: 0, absent; 1, present.

35. Anterior osmiophilic ridge: 0, absent; 1, present.

36. Changes in BBs at maturity: 0, absent; 1, dense material at tip; 2, BB cartwheel with plug; 3, BB triplets impregnated with matrix.

37. Matrix around BBs: 0, homogenous; 1, mottled.

38. Posterior of the stellate pattern: 0, extracellular or partly; 1, entirely intracellular.

39. Flagellar scales: 0, present; 1, absent.

40. Late blepharoplast with transient core: 0, yes; 1, no.

41. Direction of flagellar emergence: 0, toward side; 1, toward rear; 2, toward anterior.

42. Nuclear shape at maturity: 0, ovoid; 1, elongate.

43. Nuclear posterior shape: 0, not expanded; 1, expanded.

44. Median constriction: 0, absent; 1, present.

45. Spline attached to nucleus: 0, yes; 1, detached at maturity; 2, never attached.

46. Spline growth associated with nuclear shaping: 0, absent; 1, present.

- 47. Direction of nuclear compaction: 0, outer shell; 1, anterior to posterior; 2, at equal rates along nucleus; 3, general increase in density.
- 48. Condensed chromatin strands: 0, spaghettilike; 1, perpendicular to spline; 2, spiral-central strand; 3, general compaction; 4, spikes; 5, irregular plates; 6, solid mass from anterior tip.

49. Diverticulum during shaping: 0, absent; 1, present.

50. Number of gyres of nucleus (ordered character): 0, not coiled; 1, 0.5-3; 2, greater than 3.

51. Dense body in anterior mitochondrion: 0, absent; 1, present.

52. Mitochondrion associated with plastids in spermatogenous tissue: 0, absent; 1, present. 53. Mitochondrion associated with plastids in young spermatids: 0, absent; 1, present.

54. Specialized anterior mitochondrion: 0, present; 1, absent.

55. Specialized posterior mitochondrion: 0, present; 1, absent. 56. Additional mitochondrion in anterior of cell: 0, absent; 1, row of mitochondria behind anterior mitochondrion; 2, numerous unspecialized.

57. Origin of anterior mitochondrion: 0, fusion; 1, elongation.

58. Osmophilic material underneath anterior mitochondrion: 0, absent; 1, present.

TABLE 3. Continued.

59. Change from cristae sacs to baffles: 0, absent; 1, present.

60. Monoplastidic sperm: 0, present; 1, absent.

- Plastid determines division polarity: 0, present—at poles; 1, present—asymmetrical; 2, absent.
- 62. Starch grains in single plastid: 0, more than one; 1, one. 63. Sperm plastid contacting nucleus: 0, absent; 1, present.

64. Fibrillenscheide: 0, absent; 1, present.

- 65. Cytoplasmic loss: 0, absent; 1, partial; 2, complete (or tiny remnant).
- 66. Embryo: 0, absent; 1, present. 67. Cuticle: 0, absent; 1, present.
- 68. Lunularic acid: 0, absent; 1, present.
- 69. Elaters: 0, absent; 1, present.
- 70. Oil bodies: 0, absent; 1, present.
- 71. D-Methionine: 0, not recognize; 1, recognize.
- 72. Stomates: 0, absent; 1, present.
- 73. Vertical division of zygote: 0, absent; 1, present.
- 74. Xylem: 0, absent; 1, present. 75. Phloem: 0, absent; 1, present.
- 76. Perine layer on spores: 0, absent; 1, present.
- 77. Aerial sporophyte axis of moss/tracheophyte type: 0, absent; 1, present.
- 78. Columella: 0, absent; 1, present.
- 79. Multicellular rhizoids: 0, absent; 1, present.
- 80. Leaves on gametophyte (of moss type): 0, absent; 1, present.
- 81. Independent sporophyte: 0, absent; 1, present.
- 82. Branched sporophyte with multiple sporangia: 0, absent; 1, present.
- 83. Tracheids: 0, absent; 1, present.
- 84. Lignin: 0, absent; 1, present.
- 85. Long exserted seta of the liverwort type: 0, absent; 1, present.
- 86. Oil body cells: 0, absent; 1, present.
- 87. Spore mother cells lobed: 0, absent; 1, present.
- 88. Capsule 2-4 valved on a regular basis: 0, absent; 1, present.
- 89. Capsule wall cells with transverse thickenings: 0, absent; 1, present.
- 90. Thick capsule wall: 0, absent; 1, present.
- 91. Elongate antheridia: 0, absent; 1, present. 92. Aerial calyptra: 0, absent; 1, present.
- 93. Paraphyses: 0, absent; 1, present. 94. Costate leaves: 0, absent; 1, present.
- 95. Peristome: 0, absent; 1, present.
- 96. Operculum: 0, absent; 1, present.
- 97. Cylindrical sporogenous layer: 0, absent; 1, present.
- 98. Transfer cells on gametophyte side: 0, absent; 1, present.
- 99. Transfer cells on sporophyte side: 0, absent; 1, present.
- 100. Seed: 0, absent; 1, present.
- 101. Megaphyll: 0, absent; 1, present.
- 102. Microphyll: 0, absent; 1, present.
- 103. Lateral, broad sporangia: 0, absent; 1, present.
- 104. Exarch maturation of xylem: 0, absent; 1, present.
- 105. Chloroplast DNA inversion: 0, absent; 1, present.
- 106. Sporangia borne on leaves: 0, absent; 1, present.
- 107. Trichomes: 0, absent; 1, present.
- 108. Flavonoids: 0, absent; 1, present.
- 109. Retention of zygote: 0, absent; 1, present.
- 110. Sheathed hairs: 0, absent; 1, present.
- 111. Polyphenolics induced by sex: 0, absent; 1, present.
- 112. Parenchyma: 0, absent; 1, present.
- 113. Beaked mucilage papillae: 0, absent; 1, present.

group (15 OTUs); 10 repetitions of RANDOM taxon addition, TBR branch swapping with MUL-PARS and STEEPEST DESCENT option.

4c. Land plants only plus Coleochaete and Klebsormidium as outgroups (16 OTUs); CLOS- EST taxon addition, TBR branch swapping with MULPARS and STEEPEST DESCENT option.

Land plants only plus Coleochaete and Klebsormidium; Pedinomonas as outgroup (17 OTUs); CLOSEST taxon addition, TBR branch

TABLE 4. Data matrix LP-MORPH. See Table 3 for list of characters and states. Characters 1-65 are from sperm ultrastructure; 66-113 are from general morphology. The asterisk (*) for character 65, Lycopodium, represents a polymorphism between states 1 and 2 (Renzaglia, unpublished); the asterisk for character 86, Sphaerocarpos, represents a polymorphism between states 0 and 1 within the order Sphaerocarpales (Riella has differentiated cells).

			sper	m character					gene	eral morpho	logy: 1		
	0	2 0	3 0	4 0	5 0	6		7 0	0	0	0	0	
CHARA	?????0??00	30?00?020?	??0??0???0	0?0000000?	1100000001	0??1?1?011	20102	00000	0000000000	0000000000	2000000000	0000700100	000
NITELLA	7777707700	30?00?020?	??0?000??0	0?0000000?	1100000001	0001???0?1	20102	00000	0000000000	0000000000	200000000	0000700100	000
COLEOC PUL	2222202220	00?00?0?0?	0007000770	0000000002	0070077700	0000?1??00	00701	00000	0000000000	0000000000	2000000000	0000700011	100
COLECC ORB	2222207210	0020020202	0002000220	0000000002	0020022200	0000717700	00701	00000	0000000000	0000000000	7000000100	0000700011	110
PHAEOCEROS	0011000001	1011111100?	2007101000	0100030012	1101001101	1110000110	11102	11000	1110000100	0000000000	0000000100	0000700010	110
NOTOTHYLAS			2007101000					11000	1110000100	0000000000	0000000100	0000700010	110
MARCHANTIA			2011101100									0000000110	
			2011101100									0000700110	
PELLIA			2011101000									0000700110	1144
BLASIA			2011101100									0000700110	
			2011101000									0000700110	
HAPLOMITRIUM			2010101000									0000700110	
			2010101000										
TREUBIA			1011101012									0000700110	
SPHAGNUM													
ANDREAEA			2011101010									0000700010	
POLYTRICHUM			2011101011									0000700010	
HYPNUM			2011101011									0000000110	
TAKAKIA			7011101017									0000700110	
LYCOPODIELLA			0107101000									0111000110	
LYCOPODIUM			010?101000									0111000110	100
SELAGINELLA			200?101000									0111000110	
EQUISETUM			010?111020									0700110110	
PTERIDIUM			0107111020									1000111110	
MARSILEA			0 0107111020		A BOOK OF							1000?11110	E LE
OSMUNDA		The State of the S	0 0107111020									1000111110	
ZAMIA			00 010717102									1000111110	

swapping with MULPARS and STEEPEST DE-SCENT option.

4e. The nine land plant OTUs selected as part of LP-COMB; branch-and-bound search.

Analysis 5. Green plant molecular data (GP-MO-LEC). 26S data alone.

5a. All OTUs that have 26S data; *Pedinomonas* as outgroup (37 OTUs); 10 repetitions of RANDOM taxon addition, TBR branch swapping with MULPARS and STEEPEST DESCENT option.

5b. Land plants that have 26S data; no outgroups (11 OTUs); CLOSEST taxon addition, TBR branch swapping with MULPARS.

5c. The seven land plant OTUs selected as part of LP-COMB that have 26S data; branch-and-bound search.

Analysis 6. Land plant molecular data alone (LP-MOLEC). Each with 10 repetitions of RANDOM taxon addition, TBR branch swapping with MUL-PARS and STEEPEST DESCENT option:

6a. All data; Coleochaete and Klebsormidium as outgroups (16 OTUs).

6b. All data; Coleochaete only as outgroup (15 OTUs).

6c. All data; *Klebsormidium* only as outgroup (15 OTUs).

6d. All data; no outgroups (14 OTUs).

6e. 18S data only; no outgroups (14 OTUs).

6f. 26S data only; no outgroups (11 OTUs).

Analysis 7. Land plant combined data (LP-COMB). Each a branch-and-bound search:

7a. All data.

7b. General morphological data plus all molecular data.

7c. All morphological data plus 18S molecular data.

7d. All morphological data plus 26S molecular data.

7e. General morphological data plus 18S molecular data; decay analysis to 2 steps.

7f. General morphological data plus 26S molecular data; decay analysis to 5 steps.

Analysis 8. Green plant combined data (GP-COMB). CLOSEST taxon addition, NNI branch swapping with MULPARS, followed by TBR branch swapping with MULPARS on the shortest trees found by NNI swapping; decay analysis to 3 steps.

RESULTS

Analysis 1a. Only one tree island was found, of 9 MP trees at 208 steps (CI = 0.678; RI = 0.877). The strict consensus tree is shown in Figure 1, along with the decay index for each informative branch (obviously, here and in later figures, the terminal branches leading to OTUs cannot decay). Several traditional groups (seed plants, tracheophytes, mosses, hornworts, liverworts, land plants) were well supported by this data set, but note that the resolution among these major lineages was unresolved. Note also that the lycophytes were not supported as monophyletic.

Analysis 1b. Only one tree island was found, of 56 MP trees at 130 steps (CI = 0.708; RI = 0.882). The strict consensus tree is shown in Figure 2, along with the decay index for each informative branch. The decay analysis was completed at 1 step. However, not all trees at 2 and 3 steps less parsimonious could be saved with available RAM, thus decay classes "2," "3," and "4+" in Figure 2 are estimates. Note that relatively few clades were well supported by this data set, but a monophyletic bryophyte clade was moderately well supported. Selaginella was placed weakly with the bryophytes. CONSTRAINT analyses showed that MP topologies with a forced Mishler & Churchill (1984, 1985) resolution, i.e., [liverworts [hornworts [mosses + tracheophytes]]], were 133 steps long with this data set.

Analysis 1c. Only one tree island was found, of 8 MP trees at 67 steps (CI = 0.731; RI = 0.919). The strict consensus tree is shown in Figure 3, along with the decay index for each informative branch. The decay analysis was completed at 2 steps. However, not all trees at 3 steps less parsimonious could be saved with available RAM; thus decay classes "3" and "4+" in Figure 3 are estimates. The Mishler and Churchill topology was supported by this data set, but not strongly since trees one step longer have other paraphyletic arrangements of the three major bryophyte groups; note, however, that CONSTRAINT analyses showed that MP topologies with forced bryophyte monophyly were 70 steps long with this data set. Note that the tracheophytes were strongly supported as monophyletic, as were the lycophytes. Thus it is

TABLE 5. Taxa included in the data sets GP-MOLEC, GP-MORPH, and GP-COMB, with source for rDNA sequence data (GENBANK accession number given, if known). The taxa selected for the data set LP-MOLEC are marked with an asterisk.

Emiliana huxleyi (Lohm.) Hay & Mohler	Bhattacharya et al., 1992
Anemonia sulcata L.	Hendriks et al., 1990 (X53498)
*Glycine max (L.) Merr.	Eckenrode et al., 1985
*Oryza sativa L.	Takaiwa et al., 1984
*Zamia floridana L.	Arnold, unpublished
*Psilotum sp.	Zimmer et al., 1989
*Equisetum hymale L.	Hamby, unpublished
*Atrichum angustatum (Brid.) Bruch & Schimp.	Waters et al., 1992
*Notothylas breutelii Gott.	Waters et al., 1992
*Phaeoceros laevis (L.) Prosk.	Waters et al., 1992
*Porella pinnata L.	Buchheim & Chapman, 1992
*Conocephalum conicum (L.) Lindb.	Waters et al., 1992
*Asterella tenella (L.) P. Beauv.	Waters et al., 1992
*Riccia austinii Steph. *Vlokoomidium flooridum (A. B.) Silve Metter & Plealerell	Waters et al., 1992
*Klebsormidium flaccidum (A. Br.) Silva, Mattox & Blackwell	Waters et al., 1992
*Coleochaete nitellarum Jost.	Waters et al., 1992
*Fissidens taxifolius Hedw.	Waters et al., 1992
*Plagiomnium cuspidatum (Hedw). T. Kop.	Waters et al., 1992
Micromonas pusilla (Butcher) Manton & Parke	Kantz et al., 1990
Mantoniella squamata (Manton & Parke) Desikachary	Kantz et al., 1990
Nephroselmis pyriformis (Butcher) Rayns	Kantz et al., 1990
"Pedinomonas minutissima Skuja"	Kantz et al., 1990
Tetraselmis carteriiformis Butcher	Kantz et al., 1990
Enteromorpha intestinalis (L.) Link	Kantz et al., 1990
Ulva fasciata Delile	Zechman et al., 1990
Ulothrix zonata (Weber & Mohr) Kutz	Zechman et al., 1990
Cymopolia barbata (L.) Lamour.	Zechman et al., 1990
Bathophora oerstedii J. Ag.	Zechman et al., 1990
Codium decorticatum (Woodward) Howe	Zechman et al., 1990
Cladophoropsis membranacea (C. Ag.) Borg.	Zechman et al., 1990
Blastophysa rhizopus Reinke	Zechman et al., 1990
Trentepohlia sp.	Zechman et al., 1990
Cephaleuros parasiticus Karsten	Chapman, unpublished
Characium vacuolatum Lee & Bold	Lewis et al., 1992 (M63001)
Dunaliella parva Lerche	Lewis et al., 1992 (M62998)
Chlamydomonas reinhardtii Dangeard	Gunderson et al., 1987
Volvox carteri f. nagariensis	Rausch et al., 1989
Chlorococcopsis minuta (= Ettlia minuta (Arce & Bold) Komarek)	Lewis et al., 1992 (M62996)
Draparnaldia plumosa (Vauch.) Ag.	Buchheim & Chapman, 1992
Uronema belkae Mattox & Bold	Zechman et al., 1990
Chlamydomonas moewusii Gerloff	Buchheim et al., 1990
Stephanosphaera pluvialis Cohn	Buchheim & Chapman, 1991
Carteria radiosa Korschikoff	Buchheim & Chapman, 1992
Gonium pectorale Muller	Buchheim & Chapman, 1991
Chlorella kessleri Fott & Novakova	Huss & Sogin, 1990
Chlorella vulgaris Beij.	Huss & Sogin, 1990
Prototheca wickerhamii Soneda & Tubaki	Huss & Sogin, 1990
Chlorella protothecoides Krug.	Huss & Sogin, 1990
Chlorella minutissima Fott & Novakova	Huss & Sogin, 1990
Neochloris aquatica Starr	Lewis et al., 1992 (M62801)
Neochloris vigenis Archibald	Wilcox et al., 1992 (M74496)
Pediastrum duplex Meyen	Lewis et al., 1992 (M62997)
Scenedesmus obliquus (Turp.) Kutz	Huss & Sogin, 1990
Characium hindakii Lee & Bold	Lewis et al., 1992 (M63000)
Chlorella fusca var. vacuolata Shihira & Krauss	Huss & Sogin, 1990
Jack dar. Cacaotata Simila & Krauss	Huss & Sogni,

TABLE 5. Continued.

Ankistrodesmus stipitatus (= A. falcatus var. stipitatus (Chodat) Lemm.)	Huss & Sogin, 1990
Pseudotrebouxia gigantea Hildreth & Ahmadj.	Kantz et al., 1990
Pleurastrum terrestre Fritsch & John	Kantz et al., 1990
Characium perforatum Lee & Bold	Lewis et al., 1992 (M62999)
Parietochloris pseudoalveolaris (Deason & Bold) Watanabe & Floyd	Lewis et al., 1992 (M63002)
Friedmannia israelensis Chantanachat & Bold	Lewis et al., 1992 (M62995)

clear that a major conflict exists between the sperm data (see results of analysis 1b) and the general morphological data.

Analysis 1d. Three MP trees were found, at 65 steps (CI = 0.862; RI = 0.870), the strict consensus of which is shown in Figure 10. Note that as in analysis 1b, bryophyte monophyly was supported.

Analysis 1e. Two MP trees were found, at 35 steps (CI = 0.857; RI = 0.886), the strict consensus of which is shown in Figure 10. Note that, whereas the mosses alone were supported as sister group to the tracheophytes as in analysis 1c, the relative position of liverworts and hornworts was not resolved.

Analysis 1f. Two MP trees were found, at 105 steps (CI = 0.819; RI = 0.832), the strict consensus of which is shown in Figure 10. Note that bryophyte monophyly was supported, unlike the result of analysis 1a.

Analysis 2. 26,300 MP trees were found at 231 steps (CI = 0.550; RI = 0.895), but the search could not be completed because the RAM of the computer was exceeded. Thus, the effectiveness of the heuristic search was diminished and more trees at 231 steps undoubtedly exist than could be saved. A strict consensus of the trees that were saved is shown in Figure 4. Note the poor resolution among green algal and land plant groups.

Analysis 3a. 32 MP trees were found at 2245 steps (CI = 0.458; RI = 0.589). A strict consensus of the trees that were saved is shown in Figure 5. The main groups of land plants were unresolved (in fact, neither the tracheophytes nor the liverworts were supported as monophyletic). Tetraselmis appeared as basal to the green algal clade, while the Trentepohliales + Ulvophyceae (minus Cymopolia and Batophora) formed a monophyletic group basal to the chlorophytes plus pleurastrophytes. Pleurastrum was widely separated from the other pleurastrophytes.

Analysis 3b. Only one island of 5 MP trees at 304 steps was found (CI = 0.612; RI = 0.600). In the consensus (not shown), neither the land plants, tracheophytes, nor liverworts were monophyletic.

Analysis 3c. Two MP trees at 127 steps (CI = 0.630; RI = 0.466) were found, the strict consensus of which is shown in Figure 10. The mosses and hornworts formed a monophyletic group, and the liverworts appeared paraphyletic.

Analysis 3d. Only one MP tree, of 1597 steps, was found (CI = 0.501; RI = 0.518; shown in Fig. 6). None of the three classes of green algae sensu Mattox & Stewart (1984: chlorophytes, pleurastrophytes, and ulvophytes) included in this analysis appeared to be strictly monophyletic. The ulvophytes were paraphyletic near the base, and

Table 6. Composition of the OTUs used in data set LP-COMB. Morphological data were taken from selected taxa in LP-MORPH (Table 4) and molecular data were taken from selected taxa in GP-MOLEC (Table 5).

OTU	LP-MORPH	GP-MOLEC
COLEOCHAETE PHAEOCEROS NOTOTHYLAS MARCHANTIALES JUNGERMANNIALES POLYTRICHALES	C. orbicularis P. laevis Notothylas Marchantia Jungermannia Polytrichum	C. nitellarum P. laevis N. breutelei Asterella Porella Atrichum
RYALES QUISETUM AMIA	Hypnum Equisetum Zamia	Plagiomnium Equisetum Zamia

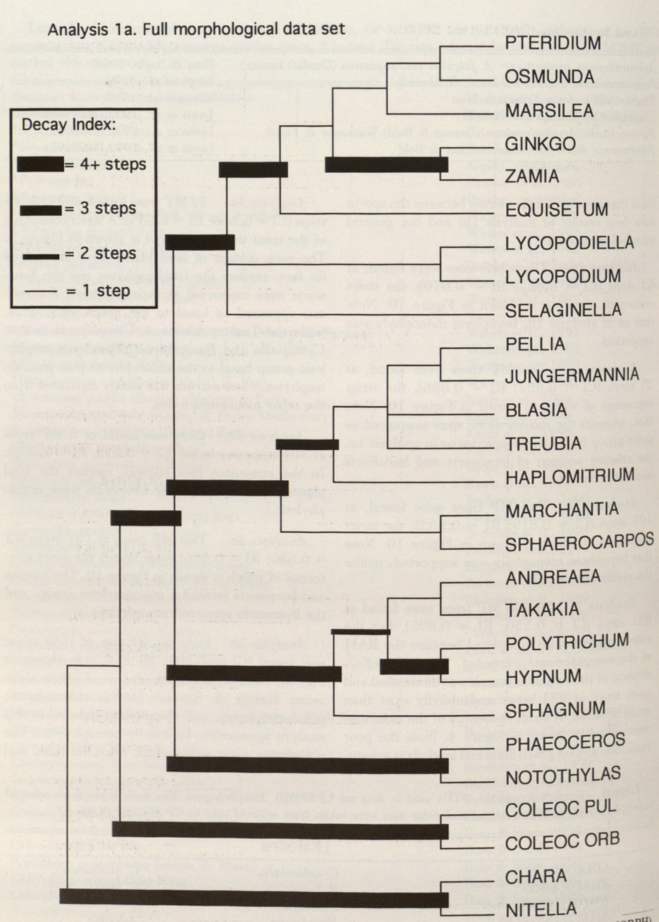


FIGURE 1. Semi-strict consensus of nine most parsimonious trees for the full morphological data set (LP-MORPH). The decay index is shown for each informative branch; thicker branches are better supported.

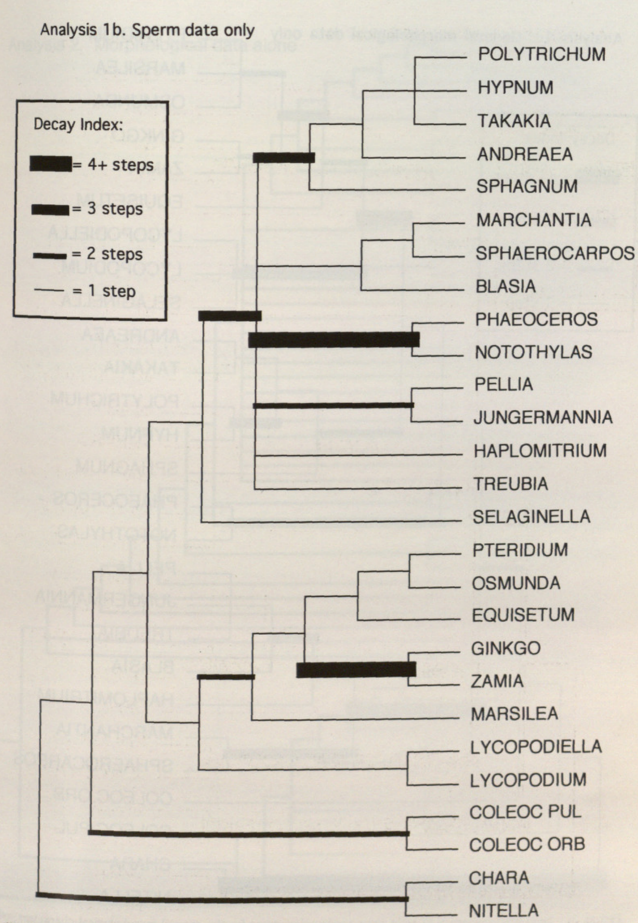


FIGURE 2. Semi-strict consensus of 56 most parsimonious trees for the sperm characters only (characters 1-65, LP-MORPH). The decay index is shown for each informative branch; thicker branches are better supported.

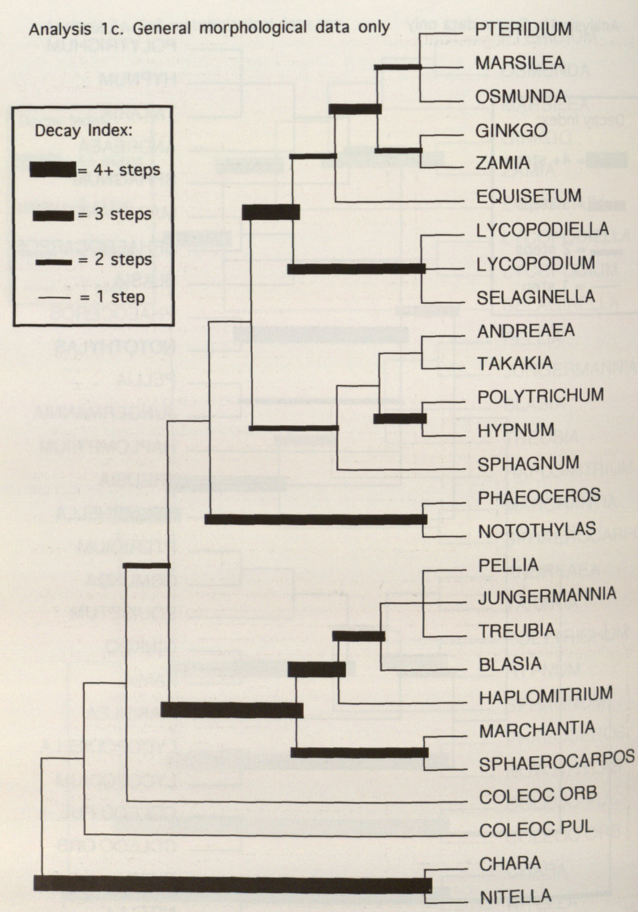


FIGURE 3. Semi-strict consensus of eight most parsimonious trees for the general morphological characters only (characters 66-113, LP-MORPH). The decay index is shown for each informative branch; thicker branches are better supported.

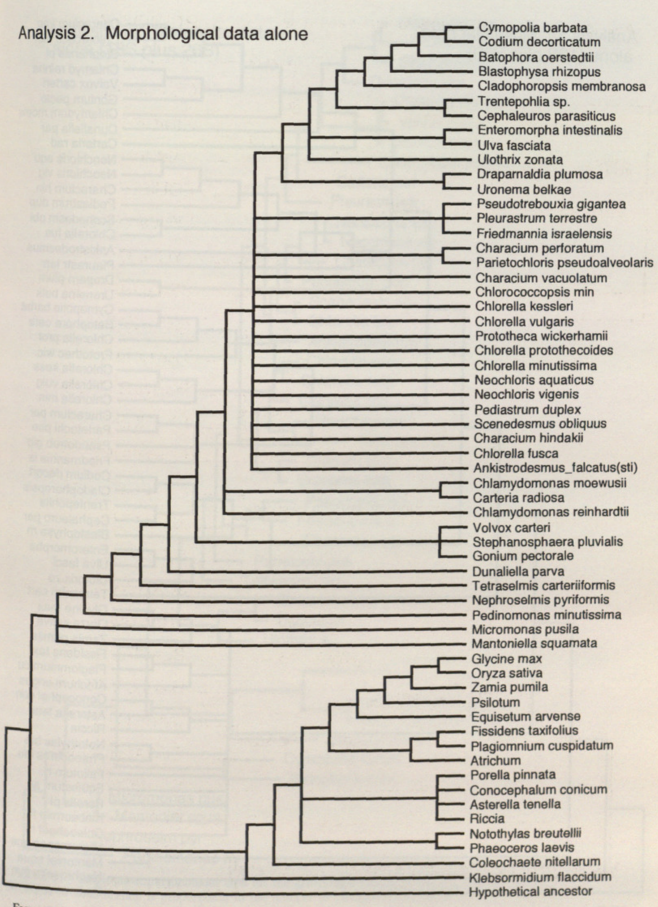


FIGURE 4. Strict consensus of 26,300 most parsimonious trees for the large-scale morphological data set (GP-MORPH). More trees at this length existed that could not be saved.

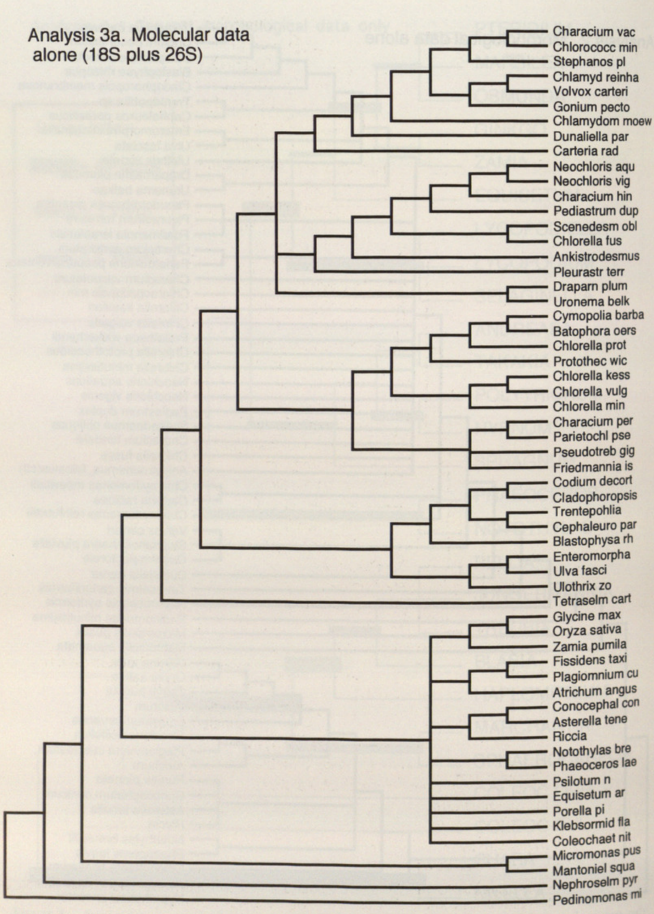


FIGURE 5. Strict consensus of 32 most parsimonious trees for the full molecular data set (GP-MOLEC).

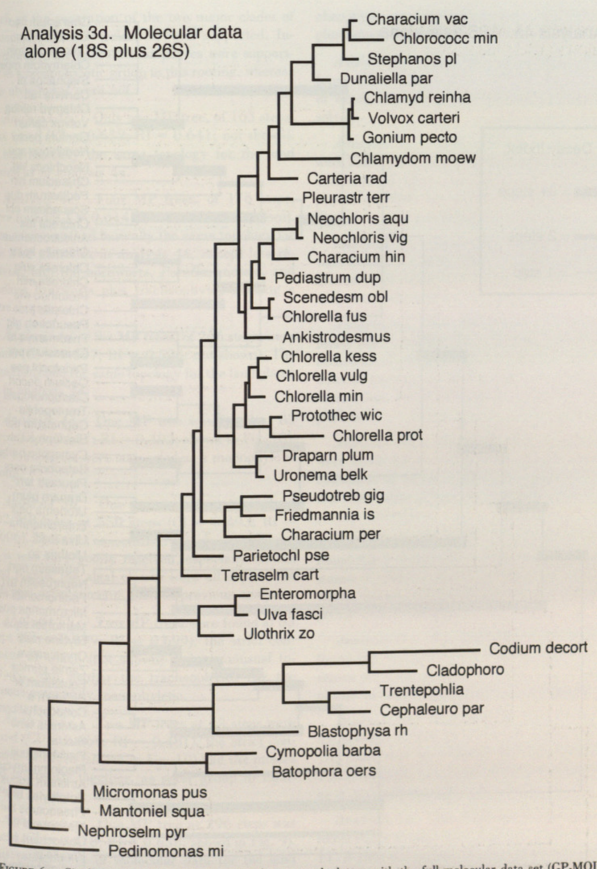


FIGURE 6. Single most parsimonious tree for the green algal taxa with the full molecular data set (GP-MOLEC). The length of each branch is proportional to the number of changes under ACCTRAN optimization.

the pleurastrophytes were paraphyletic above to the chlorophytes + Pleurastrum.

Analysis 4a. 84 MP trees at 1888 steps were found (CI = 0.450; RI = 0.589), the strict con-

sensus of which is shown in Figure 7. The land plants were resolved in an unusual topology, with the thallose liverworts alone as the sister group to the tracheophytes and the mosses as the sister group to that clade. The deep branches in the tree,

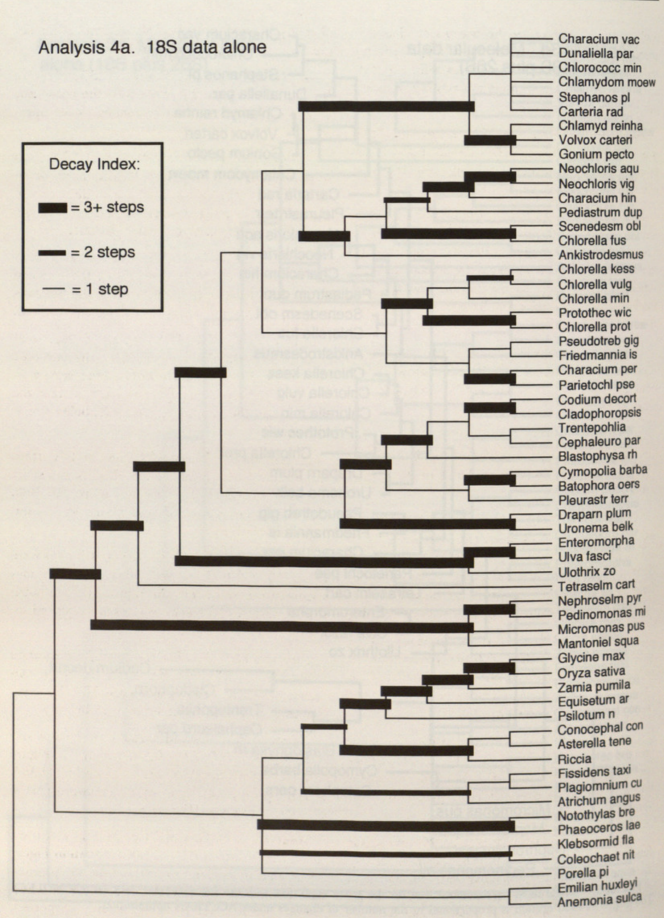


FIGURE 7. Strict consensus of 84 most parsimonious trees for the 18S characters only (GP-MOLEC). The decay index is shown for each informative branch; thicker branches are better supported.

such as the separation of the two major clades of green plants, were relatively well supported. Interestingly, the micromonadophytes were supported as a monophyletic group in this rooting, whereas the ulvophytes were not.

Analysis 4b. Only one MP tree, of 163 steps, was found (CI = 0.632; RI = 0.641; not shown). It had basically the same topology for the land plants as in analysis 4a.

Analysis 4c. Four MP trees, of 176 steps, were found (CI = 0.614; RI = 0.630; not shown). The consensus had basically the same topology for the land plants as in analysis 4a, except less resolved, with the hornworts, *Porella*, mosses, and thallose liverworts plus tracheophytes forming a tetrachotomy.

Analysis 4d. Six MP trees, of 206 steps, were found (CI = 0.607; RI = 0.595; not shown). The consensus had the same topology for the land plants as in analysis 4c.

Analysis 4e. One MP tree was found at 83 steps (CI = 0.627; RI = 0.492; shown in Fig. 10). The bryophytes were supported as a monophyletic group.

Analysis 5a. One tree island was found, of 462 MP trees at 550 steps (CI = 0.493; RI = 0.606). The strict consensus tree is shown in Figure 8; it was very poorly resolved, especially for the land plants; the algal classes were all depicted in different arrangements than in previous analyses.

Analysis 5b. Two MP trees were found at 83 steps (CI = 0.699; RI = 0.699), the strict consensus of which (not shown) had an unusual topology, with neither the tracheophytes nor the mosses potentially monophyletic.

Analysis 5c. Two MP trees at 42 steps were found (CI = 0.667; RI = 0.481), the strict consensus of which (shown in Fig. 10) had the mosses and hornworts together, as sister group to Equisetum.

Analysis 6a. One MP tree at 296 steps was found (CI = 0.618; RI = 0.638; shown in Fig. 9). This realignment of molecular data for the land plants alone gave an unusual topology, with the two putative algal outgroups widely separated, and the tracheophytes polyphyletic.

Analysis 6b. Three MP trees at 271 steps were found (CI = 0.642; RI = 0.664), the consensus of which (not shown) had the topology predictable from a re-rooting of Figure 9. The tra-

cheophytes were still polyphyletic, and the mosses plus hornworts were monophyletic.

Analysis 6c. Four MP trees at 250 steps were found (CI = 0.636; RI = 0.669), the consensus of which (not shown) had all major groups unresolved.

Analysis 6d. Three MP trees at 225 steps were found (CI = 0.662; RI = 0.702), the unrooted consensus of which (not shown) had the tracheophytes together, but neither the mosses nor the liverworts were potentially monophyletic.

Analysis 6e. Seven MP trees at 146 steps were found (CI = 0.662; RI = 0.702), the unrooted consensus of which (not shown) had the topology of that in analysis 6d except even less well resolved.

Analysis 6f. Two MP trees at 78 steps were found (CI = 0.692; RI = 0.700), the unrooted consensus of which (not shown) had the same very unusual topology of analysis 5b, with neither the mosses nor the tracheophytes potentially monophyletic.

Analysis 7a. Three MP trees at 240 steps were found (CI = 0.692; RI = 0.632), the strict consensus of which is shown in Figure 10. Note that the major groups of land plants were completely unresolved.

Analysis 7b. Four MP trees at 168 steps were found (CI = 0.655; RI = 0.561), the strict consensus of which is shown in Figure 10. The hornworts, mosses, and tracheophytes were placed together in a trichotomy.

Analysis 7c. Two MP trees at 191 steps were found (CI = 0.723; RI = 0.695), the strict consensus of which is shown in Figure 10. The bryophytes were placed together in a trichotomy.

Analysis 7d. One MP tree at 150 steps was found (CI = 0.760; RI = 0.743), shown in Figure 10. Note that the Mishler & Churchill (1984, 1985) topology, i.e., [liverworts [hornworts [mosses + tracheophytes]]], was present.

Analysis 7e. One MP tree at 124 steps was found (CI = 0.661; RI = 0.600), shown in Figure 11. A similar topology to analysis 4a was present, with [hornworts [liverworts [mosses + tracheophytes]]], but it was relatively weakly supported.

Analysis 7f. One MP tree at 78 steps was found (CI = 0.744; RI = 0.718), shown in Figure 11. Note that the Mishler & Churchill (1984, 1985) topology was again present and was relatively well supported.

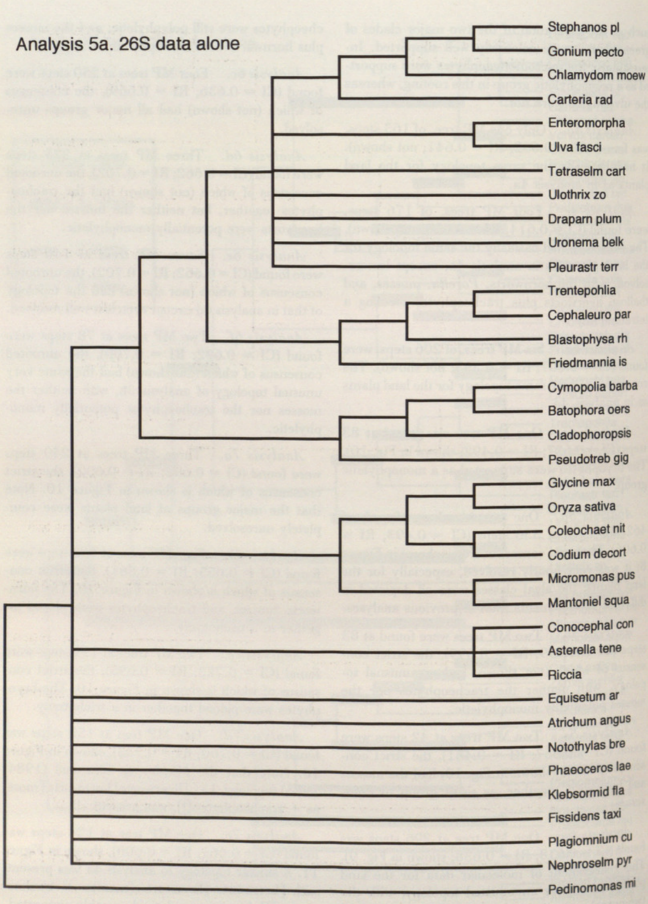


FIGURE 8. Strict consensus of 462 most parsimonious trees for the 26S characters only (GP-MOLEC).

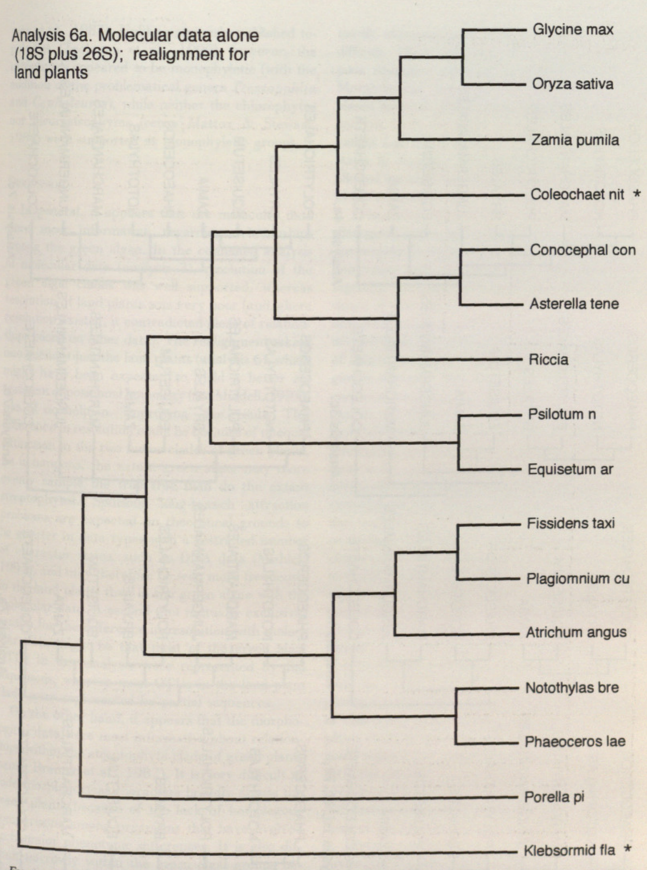
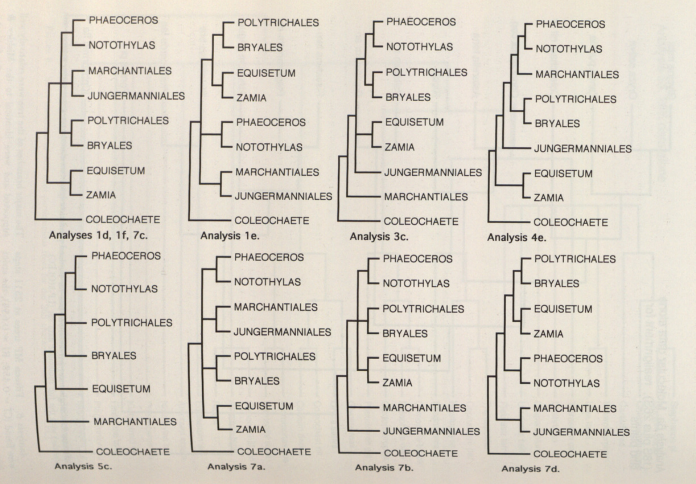


FIGURE 9. Single most parsimonious tree for the land plant taxa (plus two charophyte green algae, marked with asterisks) with the full molecular data set (LP-MOLEC).

Analysis 8. Fifteen MP trees at 2511 steps were found (CI = 0.458; RI = 0.654), the strict consensus of which is shown in Figure 12, along with the decay index for each informative branch.

The major branches of the tree were relatively well supported and were identical to the Mishler & Churchill (1984, 1985) topology in the "streptophyte" clade (i.e., charophytes + land plants; Bre-



mer et al., 1987). Unlike previously published topologies (Zechman et al., 1990), however, the ulvophytes appeared to be monophyletic (with the addition of the problematical genera *Trentopohlia* and *Cephaleuros*), while neither the chlorophytes nor pleurastrophytes (sensu Mattox & Stewart, 1984) were supported as monophyletic groups.

DISCUSSION

In general, it appears that the molecular data were most informative regarding relationships among the green algae. In the combined analysis of molecular data (analysis 3), resolution of the green algal clades was well supported, whereas resolution of land plants was very poor (and where resolution existed, it contradicted ideas of relationships based on other data). The realignment taking into account just the land plants (analysis 6), which might have been expected to yield a better assessment of positional homology (see Mindell, 1991), was of no help in "improving" the results. This difference in resolution could be because of unequal extinction in the two major clades of green plants. As it happens, the extant green algae may more evenly sample the true tree than do the extant streptophytes. Spurious long-branch attraction problems are expected on theoretical grounds to be greater in data types with a restricted number of character states, such as DNA data (Mishler, 1994), and may therefore be seen more frequently in the land plants than in the green algae with the molecular data. A second (not mutually exclusive) reason for the difference in resolution with molecular data might be that most of the green algal OTUs in this analysis were represented by full sequences, whereas most OTUs in the land plant clade were represented by partial sequences.

On the other hand, it appears that the morphological data were most informative about relationships within the streptophyte clade of green plants (sensu Bremer et al., 1987). It is very difficult to code morphological characters broadly across the green plants because of the lack of homologous comparisons among organisms that have evolved such major phenotypic differences. It is also difficult narrowly within the green algal groups because the characters useful at that level are pri-

marily ultrastructural, making sufficient sampling difficult. These difficulties are evidenced by the poor resolution obtained in analysis 2 (Fig. 4). Morphological data coded narrowly for the land plants, however, give a better supported result (e.g., analysis 1). In the latter case, the greater conservatism and larger number of potential character states in complex morphological data may have allowed the recovery of a historical signal obscured in the molecular data.

It has been suggested that even complex morphological character systems may be subject to convergent evolution when under strong selective constraints and, thus, could give misleading phylogenetic reconstructions because of non-independence of characters. It is possible that the conflict observed in analysis 1 between the set of characters derived from the spermatozoid and the general set of characters derived from many parts of the organism (compare Figs. 2 and 3) is due to selective constraints imposed on the streamlined, swimming gamete (cf. Garbary et al., 1993) or on the other morphological features such as conducting tissues. Ultimately, the only means of resolving such conflicts is by investigation of many different characters and character systems. Any one character system (or maybe all) are influenced by constraints that tend to bias phylogeny reconstruction one way or another. The hope is that if one looks at enough character systems the various noise-producing factors will "cancel-out" and a common historical signal can be detected. There is only one known process that can impose a common pattern across all these widely different character systems: phylogeny.

It is difficult to draw any definitive conclusions from the combination of molecular data with morphological data in analysis 7, because the sampling of characters is so uneven that only eight land plants could be included. The combination of both genes together, as well as 18S alone, with the LP-MORPH data set tended to favor a monophyletic bryophyte lineage, whereas the combination with 26S alone tended to favor a paraphyletic arrangement of bryophytes in the same pattern as Figure 3: [liverworts [hornworts [mosses + tracheophytes]]]. The last result is especially intriguing, as neither the morphological data alone (analysis

FIGURE 10. Strict consensus trees resulting from maximum parsimony analyses of various character combinations from the reduced data set (LP-COMB) that contains land plants which had both molecular and morphological data available. See text for explanation of which characters were used in each analysis. Coleochaete was used for outgroup rooting.

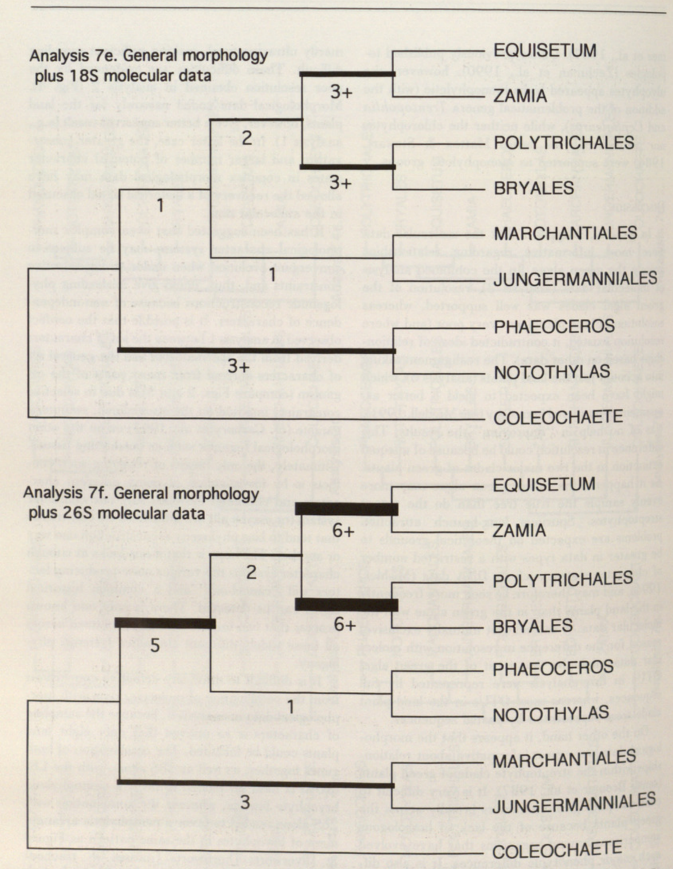


FIGURE 11. Single most parsimonious trees from two different combined analyses of the reduced data set (LP-COMB) that contains land plants which had both molecular and morphological data available. The decay index is shown for each informative branch; thicker branches are better supported.

1f) nor the 26S data alone (analysis 5c) gave such a pattern.

The overall combined analysis of GP-MOLEC and GP-MORPH (analysis 8—Fig. 12) produced

a topology that was better resolved than the consensus of the individual analyses of each data set alone. This property is often the case with such "total evidence" analyses (Kluge, 1989). The

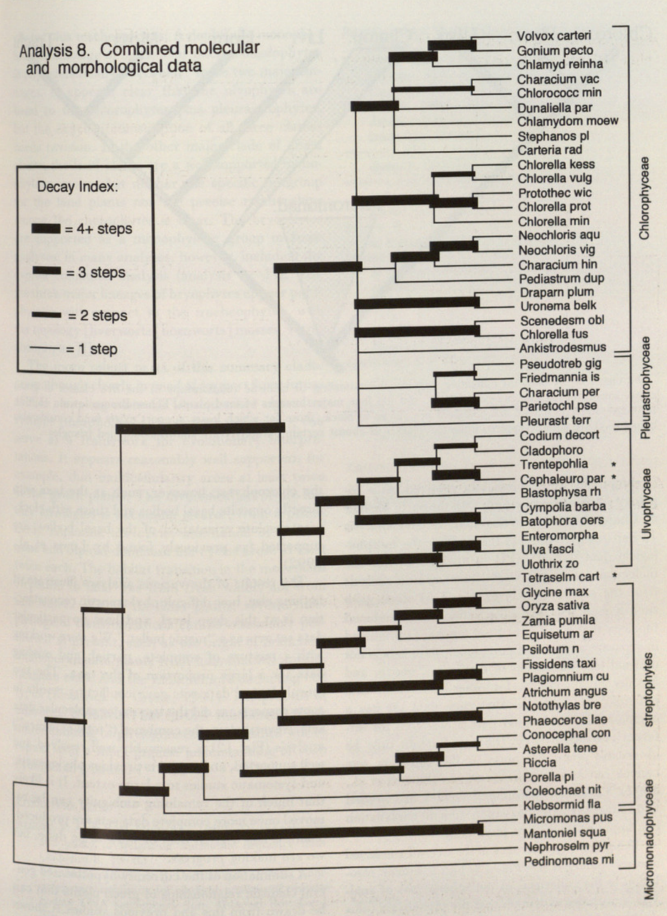


FIGURE 12. Strict consensus of 15 most parsimonious trees for the large-scale combined data set (GP-COMB). The decay index is shown for each informative branch; thicker branches are better supported. The assignment of green algae to classes according to the concept of Mattox & Stewart (1984) is shown; "streptophyte" indicates charophyte green algae plus embryophytes (sensu Bremer et al., 1987). The unusual placement of three enigmatic taxa is indicated with an asterisk.

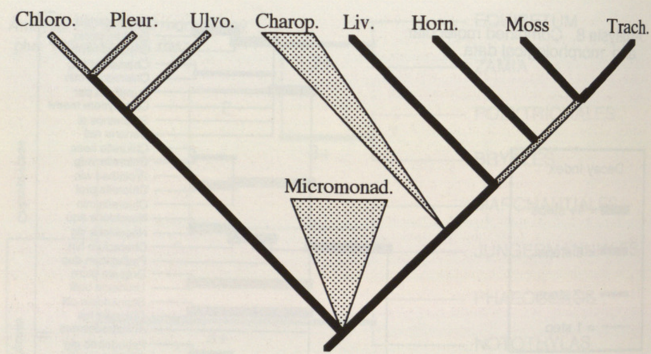


FIGURE 13. A diagram indicating current understanding of cladistic relationships of the five classes of green algae, sensu Mattox and Stewart (1984), and the four major lineages of land plants, sensu Bremer et al., (1987). Branches about which we are confident are shown in black, those for which lesser support exists (and considerable homoplasy) are shown stippled, while two classes of almost certain paraphyly are shown as stippled triangles.

streptophytes were completely resolved, in a topology completely consistent with the early Mishler & Churchill (1984, 1985) analyses and analysis la here, despite the fact that neither the morphological data (GP-MORPH in analysis 2) nor the molecular data (GP-MOLEC as a whole in analysis 3 or the separate analyses 4 and 5) alone gave such a topology. The observation of a combined topology that is different from topologies produced in separate analyses of component data sets is not unusual (see discussion of analysis 7 above, and Mishler et al., 1992, for other examples). One potential explanation is that each data set has a common historical signal that is obscured for different reasons in each; the signal might only be observable when the noise in the two data sets cancels out in a combined analysis (Barrett et al., 1991). Carefully coordinated studies are needed to discover the extent to which such an explanation can apply to land plant phylogeny.

The green algal lineages in the overall combined analysis (analysis 8—Fig. 12) had a similar resolution to the consensus of the subanalyses of analysis 3, with the exception that the ulvophytes (plus Trentopohlia and Cephaleuros) were supported as monophyletic. Several OTUs with long terminal branches (such as Pleurastrum and Ankistrodesmus) "jump around" in equally parsimonious positions, thus lowering support for the monophyly of both the Pleurastrophyceae and Chlorophyceae as currently circumscribed. Major clades within

the chlorophytes, however, such as the taxa with directly opposite basal bodies and those with clockwise absolute orientation of the basal bodies, are supported (as previously found by Lewis et al., 1992).

The results of the various analyses illustrate, if nothing else, how difficult phylogenetic reconstruction is at this deep level, and how no particular data set acts as a "magic bullet." We were working with a mixture of complete, partial, and missing data for a large proportion of the taxa. The two morphological data sets gave conflicting results in some respects, as did the two major molecular data sets. Nevertheless, the combined ("total evidence") analysis (Fig. 12) is reasonably well resolved and well supported, and supports previous phylogenetic and systematic studies to a large extent. It is likely that much of the remaining ambiguity can be removed once more complete data sets are produced. There is still a great deal of work to be done, but we are making progress.

A summation of the currently hypothesized general cladistic relationships of green plants that can be drawn from this and previous studies is shown in Figure 13, along with an indication of places in the phylogeny where uncertainty is greatest. The green plants are composed of two major monophyletic groups, one containing the bulk of the classical "green algae" (chlorophytes, pleurastrophytes, and ulvophytes), the other containing "charophyte" green algae along with the bryo-

phytes plus tracheophytes. A doubtfully monophyletic assemblage of unicellular micromonadophytes is phylogenetically "between" these two major lineages. It appears clear that the ulvophytes are basal to the chlorophytes plus pleurastrophytes, but the exact circumscription of all three classes needs revision. In the other major clade of green plants, the land plants are a well-supported monophyletic group, but neither the specific outgroup for the land plants nor the precise relationships among the charophytes is clear. The bryophytes are supported as a monophyletic group in some analyses; in many analyses, however, including the overall combined analysis (analysis 8-Fig. 12), the three major lineages of bryophytes appear paraphyletic with respect to the tracheophytes, with the topology [liverworts [hornworts [mosses + tracheophytes]]].

The more robust parts of this summary cladogram, though clearly in need of support from future studies sampling more OTUs and more character systems (morphological as well as molecular), can serve as a framework for evolutionary interpretations. It appears reasonably well supported, for example, that multicellularity arose at least twice in the green plants. The diversification of life-history strategies is becoming clearer; from a primitively haplontic life cycle, alternation of generations and diploid-dominant life cycles arose at least twice each. The habitat transition in the movement of plants to land was from fresh water, not from salt water. Within the land plants, several morphological transformations can be reasonably postulated at present, such as the origin of branched, multisporangiate plants from unbranched, unisporangiate ones, and the radiation of conducting cell types (Kendrick & Crane, 1991). The process of inference is difficult, but further refinement of our understanding of phylogenetic relationships will be repaid with a more precise understanding of such evolutionary issues.

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