Phylogenetic analyses reveal high levels of polyphyly among pleurocarpous lineages as well as novel clades

SANNA OLSSON

Institute of Botany, Plant Phylogenetics and Phylogenomics Group, Dresden University of Technology, 01062 Dresden, Germany and Botanical Museum and Department of Biological and Environmental Sciences, University of Helsinki, P.O. Box 7, FIN-00014 Helsinki, Finland e-mail: sanna.olsson@helsinki.fi

VOLKER BUCHBENDER

Institute of Botany, Plant Phylogenetics and Phylogenomics Group, Dresden University of Technology, 01062 Dresden, Germany e-mail: volker.buchbender@tu-dresden.de

JOHANNES ENROTH

Botanical Museum and Department of Biological and Environmental Sciences, University of Helsinki, P.O. Box 7, FIN-00014 Helsinki, Finland e-mail: johannes.enroth@helsinki.fi

Lars Hedenäs

Department of Cryptogamic Botany, Swedish Museum of Natural History, Box 50007, SE-104 05 Stockholm, Sweden e-mail: lars.hedenas@nrm.se

SANNA HUTTUNEN

Department of Cryptogamic Botany, Swedish Museum of Natural History, Box 50007, SE-104 05 Stockholm, Sweden. Current address: Laboratory of Genetics, Department of Biology, FI-20014 University of Turku, Finland e-mail: shuttu@utu.fi

DIETMAR QUANDT

Institute of Botany, Plant Phylogenetics and Phylogenomics Group, Dresden University of Technology, 01062 Dresden, Germany. Current address: Nees Institute for Biodiversity of Plants, Rheinische Friedrich-Wilhelms-Universität Bonn, Meckenheimer Allee 170, 53115 Bonn, Germany e-mail: quandt@uni-bonn.de

ABSTRACT. Phylogenetic analyses of the Hypnales usually show the same picture of poorly resolved trees with a large number of polyphyletic taxa and low support for the few reconstructed clades. One odd clade, however, consisting of three genera that are currently

treated either within the Leskeaceae (Miyabea) or Neckeraceae (Homaliadelphus and Bissetia), was retrieved in a previously published phylogeny based on chloroplast rbcL. In order to elucidate the reliability of the observed Homaliadelphus - Miyabea - Bissetia -clade (HMB-clade) and to reveal its phylogenetic relationships a molecular study based on a representative set of hypnalean taxa was performed. Sequence data from all three genomes, namely the ITS1 and 2 (nuclear), the trnS-rps4-trnT-trnL-trnF cluster (plastid), the nad5 intron (mitochondrial), were analyzed. Although the phylogenetic reconstruction of the combined data set was not fully resolved regarding the backbone it clearly indicated the polyphyletic nature of various hypnalean families, such as the Leskeaceae, Hypnaceae, Hylocomiaceae, Neckeraceae, Leptodontaceae and Anomodontaceae with respect to the included taxa. In addition the results favor the inclusion of the Leptodontaceae and Thamnobryaceae in the Neckeraceae. The maximally supported HMB-clade consisting of the three genera Homaliadelphus (2-3 species), Miyabea (3 species) and Bissetia (1 species) is resolved sister to a so far unnamed clade comprising Taxiphyllum aomoriense, Glossadelphus ogatae and Leptopterigynandrum. The well-resolved and supported HMBclade, here formally described as the Miyabeaceae, fam. nov. is additionally supported by morphological characters such as strongly incrassate, porose leaf cells, a relatively weak and diffuse costa and the presence of dwarf males. The latter are absent in the Neckeraceae and the Leskeaceae. It is essentially an East Asian family, with one species occurring in North America.

Keywords. *Glossadelphus, Taxiphyllum*, taxonomy, evolution, dwarf males, Miyabeaceae, Hypnales, phylogeny, *Homaliadelphus, Miyabea, Bissetia*.

• • •

Although the monophyly of pleurocarpous mosses (homocostate pleurocarps sensu Bell et al. 2007) is beyond doubt and consistently resolved with moderate to high support in multigene analyses (e.g., Beckert et al. 2001; Bell et al. 2007; Cox et al. 2000; Cox & Hedderson 1999; Quandt et al. 2007) we observe a considerable lack of resolution and support among the various pleurocarpous lineages (e.g., Buck et al. 2000; Goffinet et al. 2001; Ignatov et al. 2007; Tsubota et al. 2002). This is especially evident in species-rich and/or single marker analyses where phylogenies of homocostate pleurocarps notoriously turn out as bushes instead of trees. However, the problem of identifying natural groups is not unique to molecular systematics as bryologists throughout the last century consistently faced this challenge while recognizing lineages solely based on the interpretation of morphological traits (e.g., Buck & Vitt 1986; Hedenäs 1995). The classification of

pleurocarpous mosses even at the family level is in fact difficult, due to convergent evolution and homoplasy of morphological characters (Hedenäs 2007; Huttunen et al. 2004; Quandt et al. 2009).

Even if some families are reliably resolved through recent phylogenetic analyses (Huttunen et al. 2004; Quandt et al. 2003b, 2009; Vanderpoorten et al. 2002a), many inter- and intrafamilial relationships remain unknown, especially considering the bryological "dust bins," such as the Hypnaceae. Hence, although the new molecular tools boosted phylogenetic reconstructions, and therefore systematics, the prominent challenge in pleurocarpous moss systematics remains to identify and characterize natural higher order groups among the ca. 5000 pleurocarpous species and to relate these to each other (compare Shaw & Renzaglia 2004). This is complicated by the fact that sequence variation of the currently known markers among hypnalean taxa is extremely low, even if non-coding regions are applied. Therefore, in order to obtain a reliable backbone of pleurocarpous mosses it seems that a high sequencing effort is required and/or new markers containing better phylogenetic signals need to be applied (see www.pleurocarps.eu).

Among the few reported pleurocarpous clades receiving considerable support, a curious one was evident in the analysis based on the plastid *rbc*L gene by Tsubota et al. (2002). In this analysis, *Miyabea fruticella, Homaliadelphus targionianus* and *Bissetia lingulata* (HMB-clade), taxa that have never been considered related and are currently placed in different families, unexpectedly formed a clade with high bootstrap-support. Preliminary examination of these taxa, however, revealed that they share a number of morphological features, some of which hint at affinities to the Anomodontaceae. This suggested that the clade could be natural and inspired us to perform the present molecular study.

Bissetia was placed in the Neckeraceae since its inception by Brotherus (1906), but Enroth (1992) suggested a close relation to *Anomodon* and thus transferred it into the Anomodontaceae, a treatment that was neither reflected in the classification of mosses by Goffinet and Buck (2004) nor in the most recent classification by Goffinet et al. (2009). The genus has only one species, *B. lingulata*, distributed in Japan and South Korea (Noguchi 1989).

The genus Homaliopsis was established by Dixon and Potier de la Varde (Dixon 1928), but the generic name was recognized to be a later homonym, and the taxon was renamed Homaliadelphus by Dixon and Potier de la Varde (Dixon 1931). It has been consistently placed in the Neckeraceae, mainly due to the wide and roundish, strongly complanate leaves and a very short or absent costa. Iwatsuki (1958) revised the genus and recognized three species, but Noguchi's (1989) treatment implies he thought there were only two, the generitype, H. targionianus, with three varieties, and H. sharpii. The former has a relatively wide distribution in SE Asia, ranging from Japan and Korea to India, while the latter is restricted to North America, or, if Iwatsuki's concept of H. sharpii var. rotundatus (= H. targionianus var. rotundatus) is accepted, then it also occurs in Japan.

Miyabea has three species that are narrowly distributed in Japan, Korea and the eastern provinces of China (Noguchi 1991; Watanabe 1972; Wu et al. 2002). Brotherus (1907) originally placed the genus in the Leskeaceae "Gruppe" Anomodonteae, which was later transferred to the Thuidiaceae as the subfamily Anomodontoideae (Brotherus 1925). That placement was accepted by Watanabe (1972), although in his treatment the generic contents of the subfamily differed somewhat from Brotherus' (1925). Some authors, such as Wu et al. (2002) have recognized that taxon as an independent family, the Anomodontaceae, and included Miyabea in it. However, Buck and Goffinet (2000) as well as Goffinet et al. (2009) followed Brotherus's original concept and thought Miyabea was best placed in the Leskeaceae, even if the family's definition and circumscription differed considerably from Brotherus' concept.

In order to elucidate the reliability of the *Homaliadelphus - Miyabea - Bissetia -* clade (in the following referred to as HMB-clade) and its phylogenetic position we used a molecular approach based on sequence data from all three genomes. We combined sequence data of the ITS1-5.8S-ITS2 region (nuclear ribosomal DNA), the *nad*5-intron (mitochondrial DNA) and the *trnS-rps4-trnT-trnL-trnF* cluster (plastidal DNA). Finally, after showing the monophyly of the group, we will discuss the morphological synapomorphies distinguishing this clade.

MATERIALS AND METHODS

Taxon sampling and molecular markers. Fiftyeight taxa from 50 different genera representing 20 families of homocostate pleurocarps (Amblystegiaceae, Anomodontaceae, Brachytheciaceae, Entodontaceae, Calliergonaceae, Cryphaeaceae, Hookeriaceae, Calliergonaceae, Hypnaceae, Lembophyllaceae, Leptodontaceae, Leskeaceae, Meteoriaceae, Neckeraceae, Plagiotheciaceae, Pterobryaceae, Ptychomniaceae, Rigodiaceae, Thuidiaceae, Trachylomataceae) were included in the analyses, plus two additional outgroup taxa from the Aulacomniaceae and Hypnodendraceae. Sampling was guided by previously suggested phylogenetic affinities of *Homaliadelphus, Bissetia* and *Miyabea*, including the *rbc*L analysis of Tsubota et al. (2002). Family level treatment of the sampled taxa follows the most recent comprehensive classification of mosses by Goffinet et al. (2009).

Sequencing was performed for three genomic regions: i) the internal transcribed spacers of nuclear ribosomal DNA (ITS1 & 2), including the 5.8S gene, ii) the group I intron residing in the mitochondrial *nad5* gene (and parts of the adjacent 5' and 3' exons of the gene), and, iii) the plastidal *trnS-rps4-trnT-trnL-trnF* cluster, including four tRNAs (*trnS* (partial), *trnT*, *trnL*, *trnF* (partial)), a fast evolving gene (*rps4*), four spacers separating the coding regions, as well as one group I intron. Voucher details together with EMBL and GenBank accession numbers are listed in **Table 1**.

In addition to the material used for molecular work, several specimens were thoroughly screened for the presence of dwarf males, because they had previously been reported for two species of *Homaliadelphus* (Iwatsuki 1958; Sharp et al. 1994) but were unknown for *Bissetia* and *Miyabea*.

DNA isolation, PCR amplification and sequencing. Prior to DNA extraction, the dried specimens were cleaned with distilled water under a dissection microscope. Remaining contaminations were removed mechanically. Cleaned plant material was dried in an incubator at 70-80°C over night in a 2 ml cap with round bottom. Afterwards two stainless steel beads (5 mm) were added to each sample and crushed at 30 Hz for two times 1 min using a Mixer Mill (Retsch TissueLyser, Qiagen). From the resulting plant powder DNA was extracted using the DNeasy® Plant Mini Kit from Qiagen (Qiagen) following the manufacturer's protocol. Alternatively the CTAB-method described in Doyle and Doyle (1990) was employed. PCR amplifications (T3 Thermocycler and TGradient96, Biometra) were performed in 50 µl-reactions containing 1 U Taq DNA polymerase (peqGOLD Taq-Polymerase, peqlab Biotechnologie or Eppendorf), 1 mM dNTP mix of each 0.25 mM, $1 \times$ buffer, 1.25–2.5 mM MgCl₂ and 20 pmol of each amplification primer. Amplification of the plastid region was generally performed in three sets following the approach described in Hernández-Maqueda et al. (2008). However, primer P6/7 was generally substituted with

a new primer trnL110Rbryo, a modification of trnL110 (Borsch et al. 2003), and a new C-primer (modified from Taberlet et al. 1991) was designed. In addition two internal sequencing primers were newly designed (see Table 2) for sequencing of the rps4trnL region. PCR settings were as follows: trnS-rps4: 3 min 94°C, 35 cycles (15 s 94°C, 30 s 50°C, 1 min 72°C), 7 min 72°C; rps4-trnL: 2 min 94°C, 30 cycles (1 min 94°C, 1 min 52°C, 1 min 30 s 68°C), 5 min 68°C; trnL-F: 2 min 94°C, 35 cycles (1 min 94°C, 1 min 55°C, 1 min 68°C), 5 min 68°C. A modification of the rps4-trnL PCR-program with an increased number of cycles (to 40) was frequently used for obtaining stronger products. Amplification of the *nad5* intron was performed using a (nested) approach described in Buchbender (2009) with the following PCR profile: 1 min 30 s 96°C, 35 cycles (45 s 96°C, 1 min 55°C, 1 min 68°C), 7 min 68°C. The internal transcribed spacer of nuclear ribosomal DNA were amplified using the primers ITS5OW (Spagnuolo et al. 1999) and ITS4bryo (Stech et al. 2003) with an amplification profile of: 5 min 94°C, 40 cycles (1 min 94°C, 1 min 48°C, 45 s 68°C) with a time-increment of +4°C/cycle in the extension step, 7 min 68°C. In rare cases nested approaches were chosen using the internal primers SeqITS1 and SeqITS2. All primer sequences and references are given in Table 2. Generally multiple PCR products were pooled, concentrated and subsequently cleaned by running on 1.2% agarose gels. The excised PCR products were afterwards recovered by using the NucleoSpin Extract II kit (Macherey-Nagel) following the manufacturer's instructions. Sequencing reactions were performed using the DTCS QuickStart Reaction Kit (Beckman Coulter), applying the standard protocol supplied by the manufacturer for all reactions, using the PCR or internal primers. Extension products were run on a Beckman Coulter CEQ 8000. Alternatively, cleaned PCR products were sequenced by Macrogen Inc., South Korea (www.macrogen.com). Most sequences were generated by the authors, with some complementary sequences obtained from GenBank. Sequences were edited manually with PhyDE® v0.995 (Müller et al. 2005) and primer sequences eliminated. All generated sequences are deposited in EMBL, accession numbers are listed in Table 1.

Table 1. Taxa used in the study with EMBI submitted to GenBank from previous studie	L and GenBank accession nur es. Therefore accession numb	nbers for the sequenced or downloaded reg ers for <i>trnS-rps4-trnT-trnL-trn</i> F may be co	gions and voucher details if available. omposed of as many as three accessio	In some cases seque on numbers.	nce data have been
Species	Herbarium	Voucher ID	trnS-rps4-trnT-trnL-trnF	nad5	ITS
Anomodon giraldii	Н	H3194078	AM990342	FM161240	FM161075
Anomodon viticulosus	BUCHBENDER	Buchbender 449	AM990343	FM161241	FM161076
Aulacomnium androgynum	BM	Bell 1299	<i>rps</i> 4: AF023811	AJ291564	FM161077
			rps4-trnL: AM990344		
			<i>trn</i> L-F: AY857795		
Bissetia lingulata	Н	H3194160	AM990346	FM161243	FM161079
			rps4: AY908352		
Boulaya mittenii	HIRO	Tanaka 7308	AM990347	FM161244	FM161080
Brachythecium rivulare	Н	Parnela s.n.	AM990348	FM161245	FM161081
			<i>trn</i> L-F: AF397866		
Callicostella cf. africana	ENROTH	Rikkinen et al. 21	AM990350	FM161247	FM161085
Cratoneuropsis relaxa	MA	Musci 15238	rps4: AY908244	FM161250	FM161089
			<i>rps</i> 4- <i>trn</i> L: AM990354		
			<i>trn</i> L-F: AY429494		
Cryphaea amurensis	ENROTH	Ignatov 97–269	AM990355	FM161251	FM161090
Dichelodontium nitidum	CHR	MacMillan, BH 99/14	1754: AY449664	AY452347	ı
			rps4-trnL: AM990359		
			<i>trn</i> L-F: AY449670		
Distichophyllum crispulum	Н	H3207110	AM990360	FM161255	FM161096
Dolichomitriopsis diversiformis	Н, МНА	Nedoluzhko s.n.	rps4: AY908329	FM161257	FM161098
			<i>rps</i> 4-trnL: AM990362;		
			<i>trn</i> L-F: AF397777		
Entodon dregeanus	QUANDT	Vanderpoorten FSA AM990363	FM161258	FM161100	
Forsstroemia trichomitria	BUCHBENDER	Streimann 65120A	AM990365	FM161260	FM161103
Giraldiella levieri = Pylasia levieri	Н	Enroth 70085	AM990366	FM161261	FM161104
Glossadelphus glossoides	S	B57848	AM990368	FM161263	FM161106
Glossadelphus ogatae	Н	H3065706	AM990369	FM161264	FM161107
Gollania ruginosa	Н	Buck 23760	AM990370	FM161265	FM161108
Hampeella pallens	Н	H3205692	AM990371	FM161266	FM161109
Haplohymenium longinerve	Н	H3069640	AM990372	FM161267	FM161111
Haplohymenium pseudotriste	Н	H3069653	AM990373	FM161268	FM161112

Olsson et al.: Pleurocarpous polyphyly

Table 1. Continued.					
Species	Herbarium	Voucher ID	trnS-rps4-trnT-trnL-trnF	nad5	STT
Haplohymenium triste	Н	Enroth 63154	AM990374	FM161269	FM161113
Herpetineuron toccoae	Н	Enroth 70687	AM990375	FM161270	FM161114
Hildebrandtiella guyanensis	DREHWALD	Drehwald 4425	rps4: AY306927	FM161275	FM161119
			rps4-trnL: AM990380		
			<i>trn</i> L-F: AF509559		
Homaliadelphus targionianus	Н	Koponen et al. 55009	AM990388	FM161283	FM161129
			rps4: AY908552		
Homaliodendron exiguum	В	B263509	AM990389	FM161284	FM161130
Hookeria acutifolia	Н	Virtanen 61857	AM990393	FM161288	FM161137
Hylocomiastrum pyrenaicum	Н, МНА	Ignatov & Bezgodov 773	AM990395	FM161290	FM161140
Hylocomiastrum umbratum	Н, МНА	Ignatov & Bezgodov 81	AM990396	FM161291	FM161141
Hypnodendron vitiense	BM	Bell 480	rps4: AY524471	AY524526	FM161142
			rps4-trnL: AM990397		
			<i>trn</i> L-F: AY524499		
Hypnum cupressiforme	QUANDT	Quandt s.n.	AM990398	FM161292	FM161143
Lembophyllum divulsum	FRAHM	Frahm 8–25	AM990402	FM161296	FM161146
Leptodon smithii	В	B268385	AM990403	FM161297	FM161147
			rps4: AY908261		
Leptopterigynandrum sp.	Н	Koponen 46079	AM990404	FM161298	FM161148
Limbella tricostata	Н	H3089826	AM990406	FM161299	FM161150
			rps4: AY908572		
Lindbergia brachyptera	Н	H3194519	AM990407	FM161300	FM161151
Macrothamnium hylocomioides	Н	Sloover 42870	AM990408	FM161301	FM161152
Meteorium polytrichum	Н	Streimann 57477	AM990410	ı	FM161153
			<i>trn</i> L-F: AY044073		
Meteorium polytrichum	BUCHBENDER	Streimann 64800	AM990409	FM161302	
Miyabea fruticella	Н	Koponen 45838	AM990411	FM161303	FM161154
Miyabea rotundifolia	Н	Tan 93–771	AM990412	FM161304	FM161155
Neckera complanata	BUCHBENDER	Buchbender 204	AM990413	FM161305	FM161158
Papillaria crocea	BUCHBENDER	Streimann 47187	AM990420	FM161313	FM161186
Divilladan linarlatus		H3065601	trnL-F: AF509555 AM990367	C7C17L17C2	FM161105
F 11/11/04/011 411/8/444143	ц	TENENNETT	INCOLLING	707TOTAL.T	LIVITULIU

Species	Herbarium	Voucher ID	trnS-rps4-trnT-trnL-trnF	nad5	STI
Pinnatella minuta	Н	Rikkinen et al. 32	AM990424	FM161316	FM161194
Porotrichodendron robustum	Н	B264620	AM990426	FM161318	FM161197
Pseudoleskeopsis zippelii	Н	Enroth 71165	AM990433	FM161324	FM161206
Pseudotaxiphyllum fauriei	Н	Enroth 70134	AM990434	FM161325	FM161207
Pterobryopsis hoehnelii	QUANDT	FSA 246	AM990435	FM161326	FM161208
Rigodium implexum	QUANDT	Quandt A 10008	AM990436	FM161327	FM161209
			trnL-F: AY429499		
Scleropodium purum	QUANDT	Quandt s.n.	AM990439	FM161329	FM161211
Straminergon stramineum	DR	DR028753	AM990351	FM161330	FM161213
Taiwanobryum robustum	Н	Taiwan 1544	AM990441	FM161331	FM161215
Taiwanobryum speciosum	Н	Enroth 64877	AM990442	FM161332	FM161216
			rps4: AY908272		
Taxiphyllum aomoriense	Н	Koponen 37279	AM990443	FM161333	FM161217
Thamnobryum alopecurum	BUCHBENDER	Buchbender s.n.	AM990444	FM161334	FM161218
			rps4: AF023834		
Thamnobryum subserratum	Н	Enroth 64595	AM990446	FM161336	FM161230
Trachyloma planifolium	BONN	Frahm No. 3-12	AM990449	FM161338	FM161234
Weymouthia cochlearifolia	CHR, QUANDT	99-Mo1	AM990451	FM161340	FM161236
Weymouthia mollis	CHR, QUANDT	99-Mo2	AM990452		FM161237
			rps4: AY307014		
Zelometeorium patulum	QUANDT	Quandt A 10005	AM990453	FM161342	FM161238
			trnL-F: AF397787		

Table	2.	Primers	used	in	the	study	r. 1	Modified	nucl	eotides	are	printed	in	bold	l.
-------	----	---------	------	----	-----	-------	------	----------	------	---------	-----	---------	----	------	----

Name	Sequenz	Direction	Author	Region
trnS-F	TAC CGA GGG TTC GAA TC	F	Souza-Chies et al. (1997)	trnS-rps4
rps5rev	ATG TCC CGT TAT CGA GG	R	Nadot et al. (1994)	trnS-rps4
rps4-166F	CCA TAA TGA AAA CGT AAT TTT TG	F	Hernández-Maqueda et al. (2008)	rps4-trnL
trnL_P6/7Rbryo	CAT TGA GTC TCT GCA CCT	R	Quandt et al. (2004)	rps4-trnL
trnL110Rbryo	ATT TGG CTC AGG ATT R CT Y AT	R	modified from Borsch et al. (2003)	. rps4-trnL
trnL-A-Rbryo	AGA GCA CCG CAC TTG TAA TG	R	Hernández-Maqueda et al. (2008)	rps4-trnT spacer
trnL-A-Fbryo	CAT TAC AAG TGC GGT GCT CT	F	Hernández-Maqueda et al. (2008)	<i>trn</i> T-t <i>rn</i> L spacer
trnT_154R	AGT TTT AAG GCA ACA CTT TAT G	R	this study	rps4-trnT spacer & trnT-
				t <i>rn</i> L spacer (partial)
trnT_154F	CAT AAA GTG TTG CCT TAA AAC T	F	this study	trnT-trnL spacer (partial)
				& trnL intron
trnL-C_diplo	CG R AAT T GG TAG ACG CTA CG	F	This study modified from Taberlet et al. (1991)	<i>trn</i> L-F
trnL-F	ATT TGA ACT GGT GAC ACG AG	R	Taberlet et al. (1991)	<i>trn</i> L-F
ITS5OW	GGA GAA GTC GTA ACA AGG TTT CCG	F	Spagnuolo et al. (1999)	ITS1&2
ITS4_bryo	TCC TCC GCT TAG TGA TAT GC	R	Stech et al. (2003)	ITS1&2
SeqITS1	TTG CGT TCA AAG ACT CGA TGA	R	this study	ITS1
SeqITS2	AAC AAC TCT CAG CAA CGG	F	this study	ITS2
nad5_4F	GAA GGA GTA GGT CTC GCT TCA	F	Shaw et al. (2003a)	nad5 intron
nad5_2220R	ATA TTC CAG TGG TTG CCG CG	R	Buchbender et al. (2009)	nad5 intron
nad5_3R	AAA ACG CCT GCT GTT ACC AT	R	Shaw et al. (2003a)	nad5 intron
nad5_IF2	CTT TTG TCG TGA AGA TTC G	F	Buchbender et al. (2009)	nad5 intron

Sequence analyses and phylogenetic analyses. Alignment of the sequence data was done manually with PhyDE® v0.995, based on the criteria laid out in Kelchner (2000), Borsch et al. (2003) and Quandt and Stech (2005). Simple sequence repeats were isolated based on strict motif recognition (compare Kelchner 2000). Overlapping motifs that superficially contained identical motifs but deviated in length were considered non-homologous if the motifs could be derived independently from the adjacent region (compare tab. 4 in Quandt & Stech 2005). Following the approach in Quandt et al. (2003a) and Quandt and Stech (2004, 2005), the data matrix was screened for inversions using secondary structure models calculated with RNAstructure 4.2 (Mathews et al. 2004). Detected inversions were positionally separated in the alignment. As discussed in Quandt et al. (2003a) and Quandt and Stech (2004), presence or absence of detected inversions was not coded for the phylogenetic analyses. However, in order to gain

information from substitutions within detected inversions, a second alignment file for the phylogenetic analyses was generated with the inversions included as reversed and complemented sequences. Regions of ambiguous alignment (hotspots) were exclued from phylogenetic analyses (Table 3). Hotspots in the data matrix were defined as positions with a high degree of length mutations where homology of sequence motifs could not be assessed. This is also true for poly-mononucleotide stretches as well as other microsatellite-like areas (e.g., $(AAT)_n$) that are prone to a high variation even at the population level (Provan et al. 2001 and references therein). As indel coding approaches on these areas are likely to result in a scoring of nonhomologous events, poly-mononucleotide stretches longer than four nucleotides (nts) showing a length variation of > 1 nt were excluded from the analyses. Locations of hotspots are listed in Table 3. Alignments are available from the authors on request.

15 menude	a as reverse compleme	III.).			
No.	Position	Region (plastid)	No.	Position	Region (nuclear)
H1	701–703	rps4-trnT IGS	H16*	3925–3931	ITS 1
H2	720-722	rps4-trnT IGS	H17	3980-3982	ITS 1
H3	739–768	rps4-trnT IGS	H18	4044-4805	ITS 1
H4	843-848	rps4-trnT IGS	H19	4833-4873	ITS 1
H5	878-882	rps4-trnT IGS	H20	5013-5049	ITS 1
H6	947-953	rps4-trnT IGS	H21	5054-5127	ITS 1
H7	994–998	rps4-trnT IGS	H22	5231-5246	ITS 1
H8	1059-1064	rps4-trnT IGS	H23	5416-5421	ITS 1
H9	1221-1225	rps4-trnT IGS	H24	5659-5663	ITS 1
H10	1549-1556	trnT-trnL IGS	H25	5829-5832	ITS 2
H11	1698-1701	trnT-trnL IGS	H26	6126-6349	ITS 2
H12	1832-1837	trnT-trnL IGS	H27	6410-6509	ITS 2
H13	1864-1868	trnT-trnL IGS	H28	6664-7055	ITS 2
H14	1902-1906	trnT-trnL IGS			
H15	2547-2550	trnL-trnF IGS			
I1 [§]	2496-2501	trnL-trnF IGS			

Table 3. Location (i.e., absolute position in the combined data set) and corresponding region of mutational hotspots (H), including the observed inversion (I). * autapomorphic insertion of 709 nts in *Hypnodendron vitiense* as well as 28 nts in *Aulacomnium androgynum.* [§] Location of the inversion is given with respect to the corrected and analyzed matrix (i.e., the inversion is included as reverse complement).

Both parsimony and Bayesian analyses were performed using the information provided from indels and without indel coding. When indel coding was used, indels were incorporated in the analyses as binary data using a simple indel coding (SIC) strategy (Simmons & Ochoterena 2000) as implemented in SeqState (Müller 2005). SeqState generates a readyto-use Nexus formatted data file containing the sequence alignment with an automatically generated indel matrix appended. Command files for using the parsimony ratchet (Nixon 1999) were generated using the program PRAP2 (Müller 2007) and executed in PAUP 4.0b10 (Swofford 2002). Ratchet settings were as follows: 10 random addition cycles of 200 iterations each, with 25% up-weighting of the characters in the iterations. Heuristic bootstrap searches under parsimony were performed with 500 replicates and 10 random addition cycles per bootstrap replicate.

Bayesian analyses were performed with MrBayes v3.1.2 (Huelsenbeck & Ronquist 2001; Ronquist & Huelsenbeck 2003), applying the GTR+ Γ +I model for the sequence data (as proposed by AIC in Modeltest 3.7) and the restriction site model for the binary indel partition (partition 4), with the ascertainment (coding) bias set to variable (lset

coding = variable). To allow for possible deviating substitution models for the different regions, the sequence alignment was divided into three partitions (partition 1: chloroplast DNA; partition 2: mitochondrial DNA; partition 3: nuclear DNA). The specified prior probabilities supplied were those supplied by the default settings of the program. Posterior probability (PP) distributions of trees were created using the Metropolis-coupled Markov chain Monte Carlo (MCMCMC) method and the following search strategies suggested by Huelsenbeck et al. (2001, 2002). Ten runs with four chains (1.5×10^6) generations each) were run simultaneously (mcmc nruns = 10 nchains = 4 ngens = 1,500,000). Chains were sampled every ten generations and the respective trees written to a tree file. The ten runs mixed properly and the acceptances were within appropriate bounds. The program Tracer v1.4 (Rambaut & Drummond 2007) was used to calculate the burnin point and to examine the log likelihoods, ensuring that the runs were in the stationary phase. Since the first run was reaching its stationary phase later than the rest of the runs (at generation 650,000), this was set as the burnin point. The log likelihood values (lnL) were between 103 (run 8) and 333 (run 7), and the standard deviation varied from 0.521 to

0.962. Calculations of the consensus tree and of the posterior probability of clades were performed based upon the trees sampled after the chains converged. Consensus topologies and support values from the different methodological approaches were compiled and drawn using TreeGraph (Müller & Müller 2004).

RESULTS

Alignment and sequence analyses. The original combined and aligned sequence matrix contained 7054 positions of which 2550 positions belong to the plastid partition, 1290 positions to the mitochondrial partition and 3214 positions to the nuclear ribosomal partition. In total 28 hotspots were assigned that were almost equally distributed between the plastid region (H1-15) and the nrDNA (H16-28), with no hotspots in nad5. As most of the hotspots in the plastid data were composed of poly-mononucleotide stretches that occasionally reached the critical amount of >10 nts, in some taxa sequencing problems were encountered. However, additional sequencing with internal primers generally solved this problem. Whereas hotspots in the plastid region exclusively consisted of poly-mononucleotide stretches or microsatellite-like repetitive elements, hotspots in the ITS region often consisted of complex motifs of varying length and uncertain homology assessment. This is reflected by more than double the amount of indels compared to the chloroplast (cp) data, although the nrDNA amplicon is only half the size. In addition large autapomorphic sequence stretches were observed in the ITS region such as a putative 709 nts insertion in the ITS1 of Hypnodendron vitiense. Length mutations in the nad5 intron were rather limited and therefore alignment of nad5 was straightforward. After exclusion of the hotspots and reverse complementing of the hairpinassociated inversion in front of the trnF gene as described by Quandt et al. (2003a, 2004) and Quandt and Stech (2004), 5575 nucleotide positions could be used in the phylogenetic analyses. Of these positions 21% were variable and 11% parsimony-informative. The plastid region provided slightly more variation (27%; 14.5% parsimony-informative (p.i.) sites) compared to the nuclear region (24%; 13.5% p.i. sites), whereas the mitochondrial data showed considerably lower variation (19%, 8% p.i. sites).

Since the ITS region (1874 positions) provided only three quarters and nad5 only half (1290 positions) the number of positions compared to the cp data (2437 positions), the plastid region contributed the most to the analyses. Among the plastid partitions rps4 contained as high levels of variation and p.i. sites as the non-coding regions that were even higher than in the ITS data. This is almost reversed once indels are taken into account. 851 indels of which 268 were parsimony-informative were coded and used in the analyses. Here the nuclear indels (589 with 207 p.i. sites (35%)) vastly outnumbered the other regions (cpDNA: 227 indels containing 46 p.i. sites (20%); nad5: 34 indels containing 25 p.i. sites (74%)), although the nad5 indels provided a higher degree of p.i. sites. Detailed statistics considering the alignment, with the contribution of each region included, are listed in Table 4.

Phylogenetic analyses. The parsimony analysis including indel coding retained one most parsimonious tree (MPT, length 4848, CI= 0.557, RI= 0.515), while the analysis not including indels retained two MPTs (length 1385, CI=0,608, RI=0.724). Both of the parsimony consensus trees remained with a considerable lack of supported resolution. The MPTs showed no conflict with the results from the Bayesian inference. The results from analyses where indels were coded did not show any incongruence with the results from analyses without indel-coding, but resulted in slightly better resolved and supported trees. Therefore, only the analyses including indel-coding are discussed and only the MrBayes tree including indel-coding is illustrated in Fig. 1, complemented with bootstrap values (BS) of the parsimony analysis including indel-coding when applicable. Among homocostate pleurocarp species, the Ptychomniaceae (Ptychomniales) were resolved as the first branching clade and the Hookeriaceae (Hookeriales) sister to the Hypnales. Among the Hypnales (core ingroup) branching order is as follows: Trachylomataceae, Plagiotheciaceae, Cryphaeaceae, Pterobryaceae and Calliergonaceae. The relationships among these have moderate to high support. Although the backbone of the core ingroup is not fully resolved and lacks support in various parts, two main results are evident: i) the tree clearly indicates the polyphyletic nature of several hypnalean

Table 4. Sequence length, divergence and proportional contribution of the different regions to the data matrix as well as ti/tv ratios, number and distribution of indels. Number of characters, p-distance (p-dist.), transition/transversion ratio (ti/tv), variable sites, parsimony informative sites (p.i.) and number of indels are presented based on the data set with the hotspots excluded, whereas the length range together with the mean and the standard deviation (S.D.) are provided from the original alignment.

		length range					variable	p.i. sites	
character set	No. chars.	[nt]	Mean [nt]	S.D.	p-dist. [%]	ti/tv	sites [%]	[%]	No. indels
<i>trn</i> S-F	2437	1671–1787	1710.90	22.868	4.345	2.667	27.235	14.602	227
nad5	1290	1098-1233	1201.08	31.438	1.439	6.748	18.837	7.984	34
ITS	1847	0-1379	705.15	129.683	9.815	1.424	23.714	13.481	589
trnS-rps4 IGS	60	16-46	32.383	3.755	9.187	1.821	41.667	23.333	9
rps4	609	609	609	-	3.092	6.41	30.328	16.066	0
rps4-trnT IGS	480	265-335	303.517	11.342	5.536	2.339	29.792	15	62
<i>trn</i> T	72	72	72	-	0.366	-	8.333	1.389	0
trnT-trnL IGS	582	252-336	276.717	12.897	7.5	1.848	26.976	14.433	98
<i>trn</i> L	85	85-85	85	-	0.23	-	3.529	2.353	0
trnL intron	463	254-345	270.667	16.452	4.071	2.294	24.19	13.391	47
trnL-trnF IGS	86	47-68	60.6	3.094	8.157	2.15	38.372	26.744	11
nad5 exon1	285	276-285	284.55	1.962	1.274	2.841	12.281	7.018	0
nad5 intron	899	821-842	830.933	3.27	1.528	5.243	22.024	8.899	34
nad5 exon2	106	0-106	n.a.	n.a.	1.109	0.564	9.434	2.83	0
ITS1	863	0-979	268.95	100.278	13.82	1.487	23.523	14.137	303
5.8S	162	0-161	157.383	20.492	1.102	0.647	11.111	4.321	3
ITS2	814	0-376	271.433	41.786	12.724	1.526	26.658	14.742	283

families, such as the Leskeaceae, Hypnaceae, Hylocomiaceae, Neckeraceae, Leptodontaceae and Anomodontaceae and ii) the maximally supported HMB-clade is resolved sister to a clade consisting of *Leptopterigynandrum, Glossadelphus ogatae* and *Taxiphyllum aomoriense* with affinities to the Anomodontaceae. Besides several expected clades, unexpected but well-supported ones were found. These will be described below.

Three main clades were resolved, although support at their basal nodes is often lacking. The first clade comprises a heterogeneous group of almost as many species as traditional families with an unsupported sister group relation to the rest of the core ingroup and can be divided into two sister groups. The first group within this clade contains *Cratoneuropsis relaxa* (Amblystegiaceae), *Lindbergia brachyptera*, *Pseudoleskeopsis zippelii* (both Leskeaceae), *Boulaya mittenii* (Thuidiaceae), *Entodon dregeanus* (Entodontaceae), *Giraldiella levieri* (*Pylaisia levieri*, see Arikawa 2004, Hypnaceae), *Macrothamnium hylocomioides* and *Gollania ruginosa* (both Hylocomiaceae) sister to a clade with Phyllodon lingulatus (syn. Glossadelphus baldwinii), Glossadelphus glossoides (both Hypnaceae) and Herpetineuron toccoae (Anomodontaceae). The third Glossadelphus s.l. (incl. Phyllodon) species, G. ogatae is resolved as sister to Taxiphyllum aomoriense and Leptopterigynandrum turning Glossadelphus polyphyletic. However, within this clade a close relationship of Pseudoleskeopsis zippelii with Boulaya mittenii as well as Giraldiella levieri with Macrothamnium hylocomioides and Gollania ruginosa is suggested, whereas Hylocomiastrum (Hylocomiaceae) is resolved elsewhere rendering the Hylocomiaceae polyphyletic. Entodon dregeanus together with the aforementioned species pairs forms a significantly supported grouping.

The second main clade received a posterior probability (PP) of 92% and contains on the one hand *Hypnum cupressiforme* sister to the highly supported Anomodontaceae s. str. (*Anomodon* and *Haplohymenium*). However, *Anomodon* itself is resolved as polyphyletic, with *A. giraldii* being deeply nested among the neckeraceous taxa. On the other hand, the maximally supported *Homaliadelphus* -



Figure 1. Majority consensus of trees sampled after stationarity in the Bayesian analysis of the matrix including indels, with posterior probabilities for individual clades above the branches. Values below the branches refer to bootstrap support values.



Figure 2. Dwarf males. a. Homaliadelphus targionianus (Redfearn Jr. 35536, s). Scale bar = 0.3 mm. b. Bissetia lingulata (Mayebara s.n., s: B121919). Scale bar = 0.2 mm. c. Miyabea fruticella (Ando s.n., s: B121920). Scale bar = 0.3 mm.

Miyabea - Bissetia - clade is sister to a small and morphologically ill-defined group consisting of *Taxiphyllum aomoriense, Leptopterigynandrum* sp. and *Glossadelphus ogatae.*

The third main clade consists of: i) a wellsupported (PP 100, BS 97) Meteoriaceae-Brachytheciaceae sister group that clusters with *Limbella tricostata*, albeit with no support, and ii) a strongly supported (PP 100, BS 98), Lembophyllaceae/Rigodiaceae/Neckeraceae/ Thamnobryaceae/Leptodontaceae-clade also including *Anomodon giraldii*. Among the latter the Rigodiaceae are resolved nested within the maximally supported (PP 100, BS 100) Lembophyllaceae sister to the highly supported (PP 100, BS 97) Neckeraceae/ Thamnobryaceae/Leptodontaceae. The former Thamnobryaceae are nested among the representatives of the polyphyletic Neckeraceae and Leptodontaceae.

Dwarf males (Fig. 2a-c). Within the HMBclade, specimens with dwarf males were found in Homaliadelphus sharpii [U.S.A. Tennessee, 15 Mar 1931, Sharp, North American Musci Perfecti 232 (s)], Homaliadelphus targionianus var. targionianus [China. Sichuan, Redfearn Jr. 35536 (s)], Bissetia lingulata [Japan. Kyushū: Kumamoto, K. Mayebara (s; reg. no. B121918); Kyushū: Kumamoto, K. Mayebara (s; reg. no. B121919)] and Miyabea fruticella [Japan. Hiroshima Pref.: Sandan-kyo, H. Ando (s; reg. no. B121920)].

DISCUSSION

Sequence variation of molecular markers.

Although rps4 as well as trnL-F are classic markers in molecular phylogenetics of bryophytes, the two spacers separating rps4 from trnT and trnT from trnL have been largely ignored. Only the trnT-L IGS has been occasionally used with varying success exclusively on generic or population levels (e.g., Frey et al. 1999; Pfeiffer et al. 2004; Stech 2004). On deeper levels, however, Hernández-Maqueda et al. (2008) were the first to successfully use both spacers combined with rps4 and trnL-F in a phylogenetic study on the Grimmiaceae, an approach that was followed here. Reported sequence variation by Hernández-Maqueda et al. (2008) of the trnS-F region was similar to the values observed in our analyses (25% variable sites, 16.4% p.i. sites versus 27.2% variable sites; 14.6 p.i. sites), although their study only dealt with intrafamily level relationships. In contrast to Hernández-Maqueda et al. (2008) who reported various inversions often combined with a complex structural evolution of the *trn*L intron, only the common inversion in front of *trn*F was observed in the data set. Sequence characteristics (length, number of characters, p.i. sites, etc.) of both noncoding plastid spacers as well as the *trn*L intron were quite similar, with the variability of the intron being relatively slightly smaller (see Table 4). The second included group I intron (nad5-intron), however, was more than double the size of the *trn*L intron, but

contained roughly 30% less indels and a lower relative amount of variable and parsimonyinformative sites. As in Quandt et al. (2007) the highest relative amount of parsimony-informative sites was observed in rps4, illustrating the fastevolving nature of this gene. In terms of sequence divergence, ITS clearly diverged more than the organellar regions (see Table 4) which is surprisingly not reflected in the relative amount of p.i. sites that are comparable to the non-coding plastid regions. Although the ITS region represents a relatively short amplicon the alignment resulted in a fairly high number of positions attributed to the high number of indels that additionally displayed a high length variation. The largest indel (autapomorphic) with 709 nts was found in Hypnodendron vitiense. The high amount of indels together with the fact that one third of the indels were parsimony-informative, in contrast to one fifth in the cp data, almost doubled the p.i. sites of the nrDNA partition. In terms of parsimony information obtained from indels the nad5 intron is the most efficient, as 74% of the indels were p.i. sites, although only a few indels were recorded (34). However, as considerable parts of the length mutations in the plastid as well as in the nuclear data were excluded from the analyses (excluded hotspots), the number of length mutations, i.e., indels, represents only a proportion of those actually present.

In comparison with a recent phylogenetic study addressing the evolution of diplolepideous-alternate mosses and applying almost the same marker combinations (Quandt et al. 2007), we observe only half the sequence variability and p.i. sites in our data set. Whereas the nad5 intron displayed a p-distance of 4.4% with 32.5% of the characters being variable and 18.8% parsimony-informative, among a representative set of diplolepideous-alternate mosses, the same marker in our data set displays a p-distance of 1.4% with only 18.8% variable and 8% informative sites. In addition, the number of indels is only half as large (34) compared to a representative set of diplolepideous-alternate mosses (63). Similarly, the sequence variation (p-distance) and content of p.i. sites drops in the plastid markers from 6.7% (29.1%) to 3.1% (16.1%) in rps4 and from 8.6% (19.7%) to 4.1% (13.4%) in the trnL-intron. One reason for this phenomenon could be that the Hypnales represent the derived and rapidly radiated branch of diplolepideous-alternate mosses (cf. Shaw et al. 2003b) that has not allowed the accumulation/ fixation of synapomorphic mutations. As mentioned above, the low sequence variation among the hypnalean taxa is pronounced in the mitochondrial nad5 where sequence variation merely reaches 1.5% and the percentage of parsimony-informative sites is only half of the values found in the plastid or nuclear markers. Whereas nad5 contained several large indels characteristic for the different groupings among hypnodendroid pleurocarps (Bell et al. 2007), indels in the present data set usually comprise small simple sequence repeats of only 2-8 nts. Despite its great use among early diverging diplolepideous-alternate mosses or hypnodendroid pleurocarps (Bell et al. 2007; Quandt et al. 2007) nad5 seems to perform worse than plastid or nuclear regions in the Hypnales. This is nicely illustrated by the fact that nad5 contains only 4.1% p.i. sites (overall variability = 9.8%) in the Hypnales, whereas the plastid as well as the nuclear data set contained 11.8-13.7% p.i. sites (overall variability = 21.6–22.8%). Again, rps4 performed better compared to all other regions, even within the Hypnales (21.6% variable sites; 11.8% p.i. sites). To conclude, the observed minimal inter- and intrafamilial sequence divergence as well as the low content of p.i. sites among hypnalean nad5 sequences rejects nad5 as a cost-efficient marker for inferring relationships among the Hypnales. Moreover, because overall sequence divergence as well as phylogenetic signal of the traditional markers is faint in the Hypnales the sequencing effort needs to be extended compared to previous studies among diplolepideous taxa and/or new markers are urgently needed in order to gain a well-resolved and supported tree of the Hypnales.

Phylogenetic analyses. It is not surprising that several families included in the analyses are resolved polyphyletic, since the discrepancy between molecular phylogenetic results and previous morphological concepts of pleurocarpous mosses, which is due to morphological convergence or plasticity, is evident from several recent phylogenetic analyses (e.g., Ignatov et al. 2007; Quandt et al. 2009; Quandt & Huttunen 2004; Vanderpoorten et al.

2002a, b). However, among the Hypnales only a few families, such as the Amblystegiaceae, Brachytheciaceae, Lembophyllaceae, Meteoriaceae and Leskeaceae have been revised recently with the aid of molecular data (e.g., Huttunen et al. 2004; Huttunen & Quandt 2007; Ignatov et al. 2007; Quandt et al. 2003b, 2009; Vanderpoorten et al. 2002a, b). In contrast to previous molecular studies on other pleurocarpous families the Leskeaceae have been reported scattered all over the trees suggesting that "the Leskeaceae in the traditional circumscription is rather a concept than a taxon" (Ignatov et al. 2007), which is also indicated in the present analysis. Few molecular-based attempts have been made to elucidate the relationships among hypnalean families, and with limited success due to the low phylogenetic signal of the traditional markers (Buck et al. 2000; Ignatov et al. 2007; Tsubota et al. 2002).

Lembophyllaceae/Rigodiaceae/Neckeraceae/ Thamnobryaceae/Leptodontaceae-clade (clade A). Following the classification of Goffinet and Buck (2004) we have maintained the Rigodiaceae so far, although recent studies have already transferred Rigodium and the Rigodiaceae to the Lembophyllaceae (Quandt et al. 2009; Stech et al. 2008). The polyphyletic nature of the Neckeraceae and Leptodontaceae previously indicated by the analyses of Ignatov et al. (2007) and Tsubota et al. (2002) is supported in our analyses based on a somewhat broader sampling of both families. Our results indicate that the Leptodontaceae should be merged with the Neckeraceae. The highly supported monophyletic Thamnobryaceae (cf. Buck & Vitt 1986) are nested among the traditional Neckeraceae and Leptodontaceae and should therefore also be included in the Neckeraceae as already suggested by Enroth and Tan (1994) and Buck (1998). The placement of Anomodon giraldii within the Neckeraceae was already suggested by Tsubota et al. (2002), but we refrain from transferring the species to a new or existing Neckeraceae genus as the sampling of the Neckeraceae is presently too small and the phylogenetic position therefore too uncertain. The generic concepts of the Neckeraceae and the phylogenetic position of A. giraldii will be discussed in detail in later papers. However, it is

already clear that a more broadly defined Neckeraceae have a highly supported sister group relationship with the Lembophyllaceae.

In addition to the confusion within this clade, several members of the Neckeraceae are resolved outside of clade A, including Homaliadelphus, Bissetia and Limbella tricostata. Whereas, Homaliadelphus and Bissetia largely constitute the HMB-clade (see below), Limbella tricostata clusters with the Brachytheciaceae and Meteoriaceae. A detailed taxonomical and nomenclatural treatment of Limbella (consisting of the Hawaiian endemic L. tricostata and the closely similar L. fryei from Oregon) was provided by Ochyra (1987), who placed the genus in the Thamnobryaceae (= Neckeraceae in our concept). There is, however, a third species, currently called Limbella bartlettii, which differs clearly from the two above mentioned ones and was treated as Vittia bartlettii, within the Amblystegiaceae (Hedenäs 2003), the family where it was also placed by, e.g., Buck (1998: 211) and Goffinet and Buck (2004). The correct use of the generic name Limbella needs further clarification but we will not address the associated nomenclatural problems in the present paper, since it has no bearing on our study. In our analysis L. tricostata and, by implication, very probably also L. fryei, are related to the Brachytheciaceae-Meteoriaceae clade. It should be noted, however, that Arikawa and Higuchi (1999) found that L. tricostata (as Sciaromium tricostatum) formed a clade with Pleuroziopsis ruthenica, the single species in the family Pleuroziops(id)aceae (Goffinet & Buck 2004), although the support for the clade was quite low.

Taxiphyllum-Glossadelphus-Leptopterigynandrum-Miyabea-Bissetia-Homaliadelphus *clade* (clade B). Tsubota et al. (2002) reported an odd "*Taxiphyllum-Glossadelphus-Miyabea-Bissetia-Homaliadelphus*-clade," but with no further discussion, which basically set the stage for the present analyses. In the analyses by Tsubota et al. (2002), a clade formed by *Taxiphyllum aomoriense* and *Glossadelphus ogatae* (both illustrated in Noguchi 1994) was sister to the HMB-clade that is here formally recognized as the Miyabeaceae. As mentioned above, *Glossadelphus* is resolved as polyphyletic in the present analysis, something that was not observed in previous studies due to limited sample size. A detailed screening of the literature revealed numerous systematic and taxonomic problems associated with this genus. When the type of Glossadelphus was transferred to Phyllodon by Buck (1987) the generic name Glossadelphus became redundant. However, only a limited set of Glossadelphus species were moved to other genera. The names *Glossadelphus ogatae* and *G. glossoides* are therefore still used here, whereas G. baldwinii was synonymized with Phyllodon lingulatus by Kis (2002), a concept which is adopted here. Phyllodon was placed in the Hypnaceae by Buck and Goffinet (2000). Regardless of whether the genus is named Phyllodon or Glossadelphus, it is polyphyletic according to our analysis. While G. ogatae groups with Taxiphyllum aomoriense, Phyllodon lingulatus and G. glossoides form a clade with Herpetineuron toccoae. This is highly interesting since based on our sampling the proposed affinity of *Phyllodon* with Taxiphyllum (Buck 1987) seems to be true only for Glossadelphus ogatae. Much additional work seems to be warranted to solve the systematic and taxonomic problems within this group.

From a morphological point of view, a sister group relationship between the Miyabeaceae and the Taxiphyllum-Glossadelphus clade is difficult to sustain. Both the latter genera have homotropous to orthogonal or antitropous (terms adopted from Hedenäs 2007), more or less asymmetric capsules with an essentially unreduced peristome. The leaf cells are clearly elongate and not nearly as strongly incrassate as in the Miyabeaceae. In our analysis, an unidentified Chinese species of Leptopterigynandrum is nested in the Taxiphyllum-Glossadelphus clade, which makes this assemblage more difficult to circumscribe morphologically. However, already Ignatov et al. (2007) noticed that, e.g., Leptopterigynandrum austro-alpinum clusters with Taxiphyllum and Glossadelphus ogatae. Leptopterigynandrum is currently placed in the Leskeaceae (Buck & Goffinet 2000; Goffinet & Buck 2004) and it resembles members of the Miyabeaceae in the orthotropous capsules and reduced peristome. However, its leaf characters, including the only somewhat decurrent bases, lanceolate and acute to acuminate apices, distinctly bifurcate costa and only slightly incrassate, minutely multipapillose leaf cells (e.g., Crum & Buck 1994), bear no resemblance to the Miyabeaceae. As far as we know, dwarf males have not been reported for any species placed in *Taxiphyllum, Glossadelphus/Phyllodon* or *Leptopterigynandrum.* The sister group of the Miyabeaceae is thus morphologically heterogeneous and in need of further analyses.

Anomodontaceae (clade C). The polyphyly of Anomodon is consistent with the results of Tsubota et al. (2002). Both analyses show A. giraldii nested within the Neckeraceae. As the type species of Herpetineuron is forming a maximally supported branch with Phyllodon s. l. (see above) outside the Anomodontaceae, Herpetineuron should be excluded from the family, even if its family level relationship remains uncertain. This is in sharp contrast to the analyses by Tsubota et al. (2002) where Herpetineuron toccoae is clearly resolved within the Anomodontaceae based on rbcL.

Morphologically the Anomodontaceae sensu Goffinet and Buck (2004) represent the closest match for the HMB-clade which is to some extent supported by the molecular analyses (Fig. 1). Several species of Anomodon and Haplohymenium (the latter was included in Anomodon by Granzow-de la Cerda 1997) have orthotropous capsules with basically similarly reduced peristomes as in the Miyabeaceae, although the exostomes of Miyabea and Bissetia differ in their strongly lamellate dorsal plates, strongly trabeculate ventral plates and cristate tooth margins. Haplohymenium and species such as Anomodon viticulosus and A. rugelii have leaf shapes reminiscent of the Miyabeaceae, having decurrent bases and obtuse to rounded apices. A further similarity is the strongly incrassate leaf cells, at least partly porose, found in both the Anomodontaceae and Miyabeaceae. The main differences between the Anomodontaceae and Miyabeaceae are as follows. In the Anomodontaceae the leaf cells are strongly papillose to prorulose, but in the Miyabeaceae they are smooth. Those taxa of the Anomodontaceae that have character states resembling the Miyabeaceae mentioned above, have a strong and well-defined costa almost reaching the leaf apex or at least above mid-leaf; in the Miyabeaceae, the costa is absent (Homaliadelphus) or, when present, weak and diffuse (not sharply defined from the adjacent laminal cells) and mostly reaching to about midleaf at most, but usually ending well below it. Also, to our knowledge, dwarf males have not been reported for any species in the Anomodontaceae. Considering the fact that the *Anomodon-Haplohymenium* clade shares more morphological characters with the Miyabeaceae than the sister group of the latter does, but molecular data suggest that it is more distantly related, the Miyabeaceae obviously represent a morphologically very well-defined clade sharply delimited from its nearest relatives.

Dwarf males (Fig. 2). One of the most striking characters defining the Miyabeaceae within the context suggested by our results is the presence of dwarf males, or phyllodioicy, in all genera (although not confirmed for every species). Dwarf males were reported for Homaliadelphus laevidentatus by Iwatsuki (1958) and for *H. sharpii* (var. sharpii) by Sharp et al. (1994), but they have so far gone unnoticed for Bissetia and Miyabea. Noguchi (1989) considered B. lingulata as dioicous and stated that all examined herbarium material of this species comprised female plants. In addition, he found no male plants despite thorough investigation. Watanabe (1972) stated that species of Miyabea are dioicous, but failed to describe male plants or perigonia, as did also Noguchi (1991) and Wu et al. (2002). Watanabe (1972), however, described the spores of Miyabea fruticella and M. rotundifolia as dimorphic, that is, falling in two distinct size-classes and thus exhibiting anisospory, which is often "correlated with presence of dwarf males" in mosses (Mogensen 1983; see also Ramsay 1979). In M. fruticella the smaller spores are 8-16 µm and the larger 25-40 µm, while in M. rotundifolia the respective ranges are 12-22 and 29-38 µm. Sporophytes of the third species, M. thuidioides, are unknown. Based on measurements of 50 spores from both of the specimens 3011293 (H) and 0317006 (H-BR), we observed a basically similar but slightly less pronounced anisospory in Bissetia lingulata. The spores largely fall in two size-classes, 15-22 and 25-31 μ m, most of the spores being 20–22 or 25–27 μ m. In Homaliadelphus targionianus (specimen H3071598) the spores are very similar, 11–13 μm in diameter. Based on our own observations and on the literature cited above, the genus *Homaliadelphus* is facultatively phylloautoicous, while *Bissetia* and *Miyabea* are obligately so. The fact that *Homaliadelphus* shares a sister group relation to *Bissetia* and *Miyabea* might indicate that the latter condition evolved from a falcultative one.

DESCRIPTION OF THE MIYABEACEAE

Miyabeaceae Enroth, S. Olsson, Buchbender,

Hedenäs, Huttunen & D. Quandt, fam. nov. Plantae huius familiae foliis basi decurrentibus vel lobatis, apice late acutis, obtusis vel rotundatis, cellulis foliorum laevibus, parietibus cellularum praecipue ad basim mediumque folii valde incrassatis et porosis, costa nulla vel invalida, brevi et diffusa, plantis masculinis pumilibus praesentibus in generibus omnibus, seta longa, capsula erecta, peristomio reducto cum endostomio rudimentali vel nullo proprio.

Type GENUS: *Miyabea* Broth., Nat. Pflanzenfam. 1(3): 984. 1907.

OTHER GENERA INCLUDED: Bissetia, Homaliadelphus.

Description. Plants small to medium-sized. Main stems creeping, without a central strand, producing irregularly to subpinnately branched aerial stems with larger leaves. Paraphyllia absent. Leaves appressed-imbricate to complanate and ± homomallous when dry, ovate to ligulate or nearly rounded, base distinctly decurrent or lobed; leaf apices broadly acute to obtuse or rounded; leaf margins entire below and crenulate to toothed near apex, or entire throughout; costa absent or diffuse and ill-defined, reaching to mid-leaf or rarely to 3/4 of leaf length; laminal cells smooth, incrassate, especially so in central parts from midleaf to leaf base, where also distinctly porose; marginal cells not differentiated, but in Bissetia towards base rather transverse in several rows; alar cells indistinct. Dioicous and phyllodioicous. Setae elongate, 3-12 mm long, smooth, twisted or not; capsules orthotropous, symmetric, cylindric to obovoid; apophysal stomata few, phaneropore, round-pored; annulus absent or very poorly defined; operculum conical, obliquely long-rostrate; peristome reduced; exostome teeth smooth to papillose, not striate, in Bissetia and Miyabea lamellate at front, strongly trabeculate at back and with cristate margins;

endostome fragmentary (Noguchi 1991) or absent (*Miyabea*) to strongly reduced with fragile segments often adhering to exostome (*Homaliadelphus, Bissetia*). Calyptra cucullate, naked or with few hairs. Spores 11–13 μ m (*Homaliadelphus*) or anisosporous and ca. 15–22 and 25–31 μ m (*Bissetia*) or 8–22 and 25–40 μ m in diam. (*Miyabea*).

Discussion. The family is characterized by decurrent to lobed leaf bases, smooth, thick-walled, often porose laminal cells, especially in the median parts of the leaves, broadly acute to obtuse or rounded leaf apices, absence of costa or presence of a weak, short and rather diffuse one, presence of dwarf males in all genera, elongate seta, orthotropous, symmetrical capsules and a reduced peristome with endostome absent or rudimentary.

ACKNOWLEDGMENTS

We thank Mr. Heino Vänskä, Lic. Phil., for correcting the Latin diagnosis. SO acknowledges financial support by the Helsingin Sanomat Centennial Foundation, the University of Helsinki, Societas pro Fauna et Flora Fennica and the Finnish Biological Society Vanamo. Furthermore, the authors received support from two researcher exchange grants from the Finnish Academy/DAAD (JE, DQ) and DAAD/STINT (VB, LH, SH, SO, DQ), which is highly acknowledged. We are grateful to Cymon Cox and Bill Buck for comments on the manuscript. Research was funded by the Deutsche Forschungsgemeinschaft (DFG QU 153/3-1) and SYNTHESYS (VB, JE, SO), which is financed by the European Community Research Infrastructure Action under the FP6 "Structuring the European Research Area" Program (http://www.sysnthesys.info).

LITERATURE CITED

Arikawa, T. 2004. A taxonomic study of the genus *Pylaisia* (Hypnaceae, Musci). Journal of the Hattori Botanical Laboratory 95: 71–154.

& M. Higuchi. 1999. Phylogenetic analysis of the Plagiotheciaceae (Musci) and its relatives based on *rbcL* sequences. Cryptogamie: Bryologie et Lichénologie 20: 231–245.

- Beckert, S., H. Muhle, D. Pruchner & V. Knoop. 2001. The mitochondrial *nad*2 gene as a novel marker locus for phylogenetic analysis of early land plants: a comparative analysis in mosses. Molecular Phylogenetics and Evolution 18: 117–126.
- Bell, N., D. Quandt, T. O'Brien & A. Newton. 2007. Taxonomy and phylogeny in the earliest diverging pleurocarps: square holes and bifurcating pegs. The Bryologist 110: 533–560.
- Borsch, T., K. W. Hilu, D. Quandt, V. Wilde, C. Neinhuis & W. Barthlott. 2003. Non-coding plastid trnT-trnF sequences

reveal a well resolved phylogeny of basal angiosperms. Journal of Evolutionary Biology 16: 558–576.

- Brotherus, V. F. 1906. Neckeraceae. IX. Neckereae. In A. Engler & K. Prantl (eds.), Die natürlichen Pflanzenfamilien 1(3): 835–851. W. Englemann, Leipzig.
 - —. 1907. Leskeaceae. III. Anomodonteae. *In* A. Engler & K. Prantl (eds.), Die natürlichen Pflanzenfamilien 1(3): 984–990. W. Englemann, Leipzig.

—. 1925. Musci (Laubmoose). 2. Hälfte. In A. Engler (ed.), Die natürlichen Pflanzenfamilien, ed. 2, 11: 1–542. W. Engelmann, Leipzig.

- Buchbender, V., H. Hespanhol, C. Sérgio, A. Séneca, L. Hedenäs & D. Quandt. 2009. Phylogenetic reconstructions of the Hedwigiaceae (Bryophyta) reveal cryptic speciation and hybridisation in *Hedwigia*. Submitted to Australian Systematic Botany.
- Buck, W. R. 1987. Notes on Asian Hypnaceae and associated taxa. Memoirs of The New York Botanical Garden 45: 519–527.
 - —. 1998. Pleurocarpous mosses of the West Indies. Memoirs of The New York Botanical Garden 82: 1–400.
 - & B. Goffinet. 2000. Morphology and classification of mosses. Pages 71–123. *In* J. Shaw & B. Goffinet (eds.), Bryophyte Biology. Cambridge University Press, Cambridge, U.K.
- —, —— & A. J. Shaw. 2000. Testing morphological concepts of orders of pleurocarpous mosses (Bryophyta) using phylogenetic reconstructions based on *trnL-trnF* and *rps4* sequences. Molecular Phylogenetics and Evolution 16: 180–198.
- & D. H. Vitt. 1986. Suggestions for a new familial classification of pleurocarpous mosses. Taxon 35: 21–60.
- Cox, C. J., B. Goffinet, A. E. Newton, A. J. Shaw & T. A. Hedderson. 2000. Phylogenetic relationships among the diplolepideous-alternate mosses (Bryidae) inferred from nuclear and chloroplast DNA sequences. The Bryologist 103: 224–241.
- & T. A. Hedderson. 1999. Phylogenetic relationships among the ciliate arthrodontous mosses: evidence from chloroplast and nuclear DNA sequences. Plant Systematics and Evolution 215: 119–139.
- Crum, H. & W. R. Buck. 1994. Leskeaceae. In A. J. Sharp, H. Crum & P. M. Eckel (eds.), The moss flora of Mexico. Part II. Memoirs of The New York Botanical Garden 69: 847–860.
- Dixon, H. N. 1928. *Homaliopsis* Dix. & Potier de la Varde, gen. nov. muscorum. Annales Bryologici 1: 47–48.
 - ——. 1931. Homaliadelphus Dix. & P. de la Varde, nom. nov. Revue Bryologique et Lichénologique 4: 142.
- Doyle, J. J. & J. L. Doyle. 1990. Isolation of plant DNA from fresh tissue. Focus 12: 13–15.
- Enroth, J. 1992. Notes on the Neckeraceae (Musci). 11–12. The taxonomic position of *Pinnatella callicostelloides* and

Bissetia lingulata, with the description of *Chileobryon* (Anomodontaceae). Nova Hedwigia 54: 137–146.

- & B. Tan. 1994. Contributions to the bryoflora of China 10. The identity of *Homaliodendron neckeroides* (Neckeraceae, Musci). Annales Botanica Fennici 31: 53–57.
- Frey, W., M. Stech & K. Meißner. 1999. Chloroplast DNArelationship in palaeoaustral *Lopidium concinnum* (Hypopterygiaceae, Musci). An example of steno-evolution in mosses. Studies in austral temperate rain forest bryophytes 2. Plant Systematics and Evolution 218: 67–75.
- Goffinet, B. & W. R. Buck. 2004. Systematics of the Bryophyta (mosses): from molecules to a revised classification. Monographs in Systematic Botany from the Missouri Botanical Garden 98: 270–289.
 - —, —— & A. J. Shaw. 2009 [2008]. Morphology, anatomy and classification of the Bryophyta. *In* B. Goffinet & A. J. Shaw (eds.), Bryophyte Biology, ed. 2. Cambridge University Press, Cambridge, U.K.
- —, C. J. Cox, A. J. Shaw & T. A. Hedderson. 2001. The Bryophyta (mosses): systematic and evolutionary inferences from an *rps*4 gene (cpDNA) phylogeny. Annals of Botany 87: 191–208.
- Granzow-de la Cerda, Í. 1997. Revision and phylogeny of Anomodon and Herpetineuron (Anomodontaceae, Musci).
 Contributions from the University of Michigan Herbarium 21: 205–275.
- Hedenäs, L. 1995. Higher taxonomic level relationships among diplolepidous pleurocarpous mosses —a cladistic overview. Journal of Bryology 18: 723–781.
 - 2003. Amblystegiaceae (Musci). Flora Neotropica Monograph 89: 1–107.
- ———. 2007. Morphological characters and their use in pleurocarpous moss systematics. *In* A. E. Newton & R. Tangney (eds.), Pleurocarpous mossessystematics and evolution. The Systematics Association Special Volume Series 71: 227–245.
- Hernández-Maqueda, R., D. Quandt, O. Werner & J. Muñoz. 2008. Phylogeny and classification of the Grimmiaceae/ Ptychomitriaceae complex (Bryophyta) inferred from cpDNA. Molecular Phylogenetics and Evolution 46: 863–877.
- Huelsenbeck, J. P., B. Larget, R. E. Miller & F. Ronquist. 2002. Potential applications and pitfalls of Bayesian inference of phylogeny. Systematic Biolology 51: 673–688.
- & F. Ronquist. 2001. MRBAYES: Bayesian inference of phylogeny. Bioinformatics 17: 754–755.
 - , _____, R. Nielsen & J. P. Bollback. 2001. Bayesian inference of phylogeny and its impact on evolutionary biology. Science 294: 2310–2314.
- Huttunen, S., M. S. Ignatov, K. Müller & D. Quandt. 2004. Phylogeny and evolution of epiphytism in the three moss families Meteoriaceae, Brachytheciaceae and Lembophyllaceae. Monographs in Systematic Botany from the Missouri Botanical Garden 98: 328–361.

- & D. Quandt. 2007. Phylogenetic relationships within the moss family Meteoriaceae in the light of different datasets, alignment and analysis methods. *In* A. E. Newton & R. Tangney (eds.), Pleurocarpous mossessystematics and evolution. The Systematics Association Special Volume Series 71: 145–162.
- Ignatov, M. S., A. A. Gardiner, V. K. Bobrova, I. A. Milyutina, S. Huttunen & A. V. Troitsky. 2007. On the relationships of mosses of the order Hypnales, with special reference to taxa traditionally classified in the Leskeaceae. *In* A. E. Newton & R. Tangney (eds.), Pleurocarpous mossessystematics and evolution. The Systematics Association Special Volume Series 71: 177–214.
- Iwatsuki, Z. 1958. Review of the genus *Homaliadelphus*. The Bryologist 61: 68–78.
- Kelchner, S. A. 2000. The evolution of non-coding chloroplast DNA and its application in plant systematics. Annals of the Missouri Botanical Garden 87: 482–498.
- Kis, G. 2002. Comments on some African species of the moss genus *Glossadelphus* M. Fleisch. Cryptogamie: Bryologie 23: 157–169.
- Mathews, D. H., M. D. Disney, J. L. Childs, S. J. Schroeder, M. Zuker & D. H. Turner. 2004. Incorporating chemical modification constraints into a dynamic programming algorithm for prediction of RNA secondary structure. Proceedings of the National Academy of Sciences U.S.A. 101: 7287–7292.
- Mogensen, G. S. 1983. The spore. *In* R. M. Schuster (ed.), New Manual of Bryology 1: 325–342. Hattori Botanical Laboratory, Nichinan, Japan.
- Müller, J. & K. Müller. 2004. TreeGraph: automated drawing of complex tree figures using an extensible tree description format. Molecular Ecology Notes 4: 786–788.
- Müller, K. F. 2005. SeqState—primer design and sequence statistics for phylogenetic DNA data sets. Applied Bioinformatics 4: 65–69.
- 2007. PRAP2—likelihood and parsimony ratchet analysis, v. 0.9. Available at www.phyde.de.
- —, D. Quandt, J. Müller & C. Neinhuis. 2005. PhyDE [®] 0.995: Phylogenetic Data Editor, Dresden. Available at www.phyde.de.
- Nadot, S., R. Bajon & B. Lejeune. 1994. The chloroplast gene rps4 as a tool for the study of Poaceae phylogeny. Plant Systematics and Evolution 191: 27–38.
- Nixon, K. C. 1999. The parsimony ratchet, a new method for rapid parsimony analysis. Cladistics 15: 407–411.
- Noguchi, A. (supplemented by Z. Iwatsuki). 1989. Illustrated Moss Flora of Japan. Part 3. Hattori Botanical Laboratory, Nichinan, Japan.
- ——— (supplemented by Z. Iwatsuki & T. Yamaguchi). 1991. Illustrated Moss Flora of Japan. Part 4. Hattori Botanical Laboratory, Nichinan, Japan.
- ——. 1994. Illustrated Moss Flora of Japan. Part 5. Hattori Botanical Laboratory, Nichinan, Japan.

Ochyra, R. 1987. On the taxonomy and family placement of the moss genus *Limbella* (C. Müll.) Broth. Journal of Bryology 14: 465–485.

Pfeiffer, T., F. Schaumann, G. G. Hässel de Menéndez & W. Frey. 2004. Inter- and infraspecific relationships in the Gondwanan liverwort genus *Hymenophyton* (Hymenophytaceae, Hepaticophytina). Studies in austral temperate rain forest bryophytes 23. Australian Systematic Botany 17: 407–421.

Provan, J., W. Powell & P. M. Hollingsworth. 2001. Chloroplast microsatellites: new tools for studies in plant ecology and systematics. Trends in Ecology and Evolution 16: 142–147.

- Quandt, D., N. Bell & M. Stech. 2007. Unravelling the knot: the Pulchrinodus fam. nov. (Bryales). Nova Hedwigia Beiheft 131: 21–39.
- & S. Huttunen. 2004. Evolution of pendent life-forms in bryophytes. Journal of the Hattori Botanical Laboratory 95: 206–217.

—, —, R. Tangney & M. Stech. 2009. A generic revision of the Lembophyllaceae (Bryopsida) based on molecular data. Systematic Botany 34: in press.

- —, K. Müller & S. Huttunen. 2003a. Characterisation of the chloroplast DNA *psb*T-H region and the influence of dyad symmetrical elements on phylogenetic reconstructions. Plant Biology 5: 400–410.
- —, —, M. Stech, K. W. Hilu, W. Frey, J.-P. Frahm & T. Borsch. 2004. Molecular evolution of the chloroplast *trn*L-F region in land plants. Monographs in Systematic Botany from the Missouri Botanical Garden 98: 13–37.
- _____, _____, _____, H. Streimann, J.-P. Frahm & W. Frey. 2003b. Molecular phylogenetics of the Meteoriaceae s. str.: focusing on the genera *Meteorium* and *Papillaria*. Molecular Phylogenetics and Evolution 32: 435–461.
- & M. Stech. 2004. Molecular evolution of the trnT_{UGU}trnF_{GAA} region in bryophytes. Plant Biology 6: 545–554.
- & ______. 2005. Molecular evolution and secondary structure of the chloroplast *trn*L intron in bryophytes. Molecular Phylogenetics and Evolution 36: 429–443.

Rambaut, A. & A. J. Drummond. 2007. Tracer v1.4. Available from http://beast.bio.ed.ac.uk/Tracer.

Ramsay, H. P. 1979. Anisospory and sexual dimorphism in the Musci. Pages 281–316. *In* G. C. S. Clarke & J. G. Duckett (eds.), Bryophyte Systematics. Academic Press, London.

Ronquist, F. & J. P. Huelsenbeck. 2003. MRBAYES 3: Bayesian phylogenetics inference under mixed models. Bioinformatics 19: 1572–1574.

- Sharp, A. J., H. Crum & P. M. Eckel (eds.). 1994. The moss flora of Mexico. Vol. 2. Memoirs of The New York Botanical Garden 69: 581–1113.
- Shaw, A. J., C. J. Cox & S. B. Boles. 2003a. Polarity of peatmoss (*Sphagnum*) evolution: who says bryophytes have no roots? American Journal of Botany 90: 1777–1787.
 - —, —, B. Goffinet, W. R. Buck & S. B. Boles. 2003b. Phylogenetic evidence for a rapid radiation of pleurocarpous mosses (Bryopsida). Evolution 57: 2226–2241.

— & K. Renzaglia. 2004. Phylogeny and diversification of Bryophytes. American Journal of Botany 91: 1557–1581.

Simmons, M. P. & H. Ochoterena. 2000. Gaps as characters in sequence-based phylogenetic analyses. Systematic Biology 49: 369–381.

Souza-Chies, T. T., G. Bittar, S. Nadot, L. Carter, E. Besin & B. Lejeune. 1997. Phylogenetic analysis of Iridaceae with parsimony and distance methods using the plastid gene *rps*4. Plant Systematics and Evolution 204: 109–123.

Spagnuolo, V., S. Cozzolino, R. Castaldo & P. De Luca. 1999. Patterns of relationships in Trichostomoideae (Pottiaceae, Musci). Plant Systematics and Evolution 216: 69–79.

Stech, M. 2004. Supraspecific circumscription and classification of *Campylopus* (Dicranaceae, Bryopsida) based on inferences from sequence data. Systematic Botany 29: 817–824.

—, D. Quandt, A. Lindlar & J.-P. Frahm. 2003. The systematic position of *Pulchrinodus inflatus* (*Eucamptodon inflatus*) based on molecular data. Studies in austral temperate rain forest bryophytes 14. Australian Systematic Botany 16: 561–568.

- , M. Sim-Sim, G. Esquível, S. Fontinha, R. Tangney, C. Lobo, R. Gabriel & D. Quandt. 2008. Explaining the "anomalous" distribution of *Echinodium* Jur. (Bryopsida): independent evolution in Macaronesia and Australasia. Organisms, Diversity & Evolution 8: 282–292.
- Swofford, D. L. 2002. PAUP*. Phylogenetic Analysis Using Parsimony (*and Other Methods). Version 4. Sinauer Associates, Sunderland, MA.

Taberlet, P., L. Gielly, G. Pautou & J. Bouvet. 1991. Universal primers for amplification of 3 noncoding regions of chloroplast DNA. Plant Molecular Biology 17: 1105–1109.

Tsubota, H., T. Arikawa, H. Akiyama, E. De Luna, D. González, M. Higuchi & H. Deguchi. 2002. Molecular phylogeny of hypnobryalean mosses as inferred from a large-scale dataset of chloroplast *rbcL*, with special reference to the Hypnaceae and possibly related families. Hikobia 13: 645–665.

Vanderpoorten, A., L. Hedenäs, C. J. Cox & A. J. Shaw. 2002a. Phylogeny and morphological evolution of the Amblystegiaceae (Bryopsida). Molecular Phylogenetics and Evolution 23: 1–21.

, _____, ____ & _____. 2002b. Circumscription, classification, and taxonomy of Amblystegiaceae (Bryopsida) inferred from nuclear and chloroplast sequence data and morphology. Taxon 51: 115–122.

- Watanabe, R. 1972. A revision of the family Thuidiaceae in Japan and adjacent areas. Journal of the Hattori Botanical Laboratory 36: 171–320.
- Wu, P. C., J. Yu & M. Z. Wang. 2002. Anomodontaceae. *In* P. C. Wu, M. R. Crosby & H. Si (eds.), Moss Flora of China (English Version) 6(Hookeriaceae–Thuidiaceae): 131–149.
 Science Press (Beijing, New York) & Missouri Botanical Garden Press (St. Louis).

ms. received July 14, 2008; accepted February 19, 2009.